

Technical Report

Volume 2b Appendixes (question 3)

Introduction

This volume deals with question 3 of the systematic review for perioperative patient blood management.

In patients undergoing surgery, what is the effect of perioperative strategies that minimise blood loss on morbidity, mortality and blood transfusion?

This volume is accompanied by Volume 1b, which presents the systematic review of the evidence and the evidence-based recommendations for this question. Two other volumes – 1a and 2a – cover questions 1, 2 and 4–9. Dates covered by the literature searches are given in Chapter 2 of Volume 1a (see Table 2.1.1).

Question 3 includes the following 10 interventions:

- Intervention 1 acute normovolemic haemodilution (ANH)
- Intervention 2 intraoperative cell salvage
- Intervention 3 perioperative acute normovolemic haemodilution combined with intraoperative cell salvage
- Intervention 4 postoperative cell salvage
- Intervention 5 deliberate induced hypotension
- Intervention 6 prevention of hypothermia
- Intervention 7 point-of-care testing using thromboelastography
- Intervention 8 administration of antifibrinolytics (aprotinin, tranexamic acid, εaminocaproic acid) and desmopressin (1-deamino-8-D-arginine vasopressin, DDAVP)
- Intervention 9 appropriate patient positioning
- Intervention 10 preoperative autologous donation (PAD).

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Appendix A: Literature searches

EMBASE.com

Intervention 1 - Acute normovolemic haemodilution: Level I evidence

Search conducted 21 December 2009 (1966 to 30 July 2009)

#	Search	Results
#1	'surgery' OR 'surgery'/exp OR surgery OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR peroperat* OR postoperat* OR 'peroperative period'/exp OR 'peroperative period' OR 'postoperative period'/exp OR 'preoperative period' NOT [30-7-2009]/sd	13076417
#2	'blood'/exp OR 'blood' OR 'serum'/exp OR 'serum' OR 'plasma'/exp OR 'plasma' OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR 'bleeding' OR bleed* OR hemarthros* OR haemarthros* OR haemat* OR haemoperi* OR haemoperi* OR 'anemia'/exp OR 'anemia' OR 'anaemia'/exp OR 'anaemia' NOT [30-7-2009]/sd	2398903
#3	'meta analysis'/exp OR 'meta analysis' OR 'systematic review'/exp OR 'systematic review' OR 'pooled analysis' OR ('review'/exp OR 'review' AND ('meta analysis'/exp OR 'meta analysis' OR systemat* OR pool*)) NOT [30-7-2009]/sd	111681
#4	#1 AND #2 AND #3	20746
#5	'hemodilution' OR 'hemodilution'/exp OR hemodilution OR 'haemodilution' OR 'haemodilution'/exp OR haemodilution OR 'blood dilution'/exp OR 'blood dilution' NOT [30-7-2009]/sd	7601
#6	#21 AND #30	69

Intervention 1 – Acute normovolemic haemodilution: Level II evidence

Search conducted 3 January 2010 (January 2002 to 30 July 2009)

#	Search	Results
#1	'surgery' OR 'surgery'/exp OR surgery OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR peroperat* OR postoperat* OR 'peroperative period'/exp OR 'peroperative period' OR 'postoperative period'/exp OR 'preoperative period' NOT [30-7-2009]/sd AND [2002-2010]/py	4357522
#2	'blood'/exp OR 'blood' OR 'serum'/exp OR 'serum' OR 'plasma'/exp OR 'plasma' OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR 'bleeding' OR bleed* OR hemarthros* OR haemarthros* OR haemarthros* OR haemarthros* OR haemarthros* OR 'anaemia' OR 'anaemia'/exp OR 'anaemia'	2471241
#3	'clinical trial'/exp OR 'clinical trial' OR 'randomized controlled trial'/exp OR 'randomized controlled trial' OR 'randomization'/exp OR randomization OR 'single blind procedure'/exp OR 'single blind procedure' OR 'double blind procedure'/exp OR 'triple blind procedure'/exp OR 'triple blind procedure'/exp OR 'crossover procedure'/exp OR 'crossover procedure' OR 'placebo'/exp OR placebo OR randomi?ed:ab,ti OR rct:ab,ti OR 'random allocation':ab,ti OR 'randomly allocated':ab,ti OR 'allocated randomly':ab,ti OR (allocated NEAR/2 random*):ab,ti OR 'single blind':ab,ti OR 'single blinded':ab,ti OR 'double blinded':ab,ti OR 'treble blind':ab,ti OR 'treble blinded':ab,ti OR 'triple blind':ab,ti OR 'triple blinded':ab,ti OR placebo*:ab,ti OR 'prospective study'/exp OR 'prospective study' NOT ('case study'/exp OR 'case study' OR 'case report':ab,ti OR 'abstract report'/exp OR 'abstract report' OR 'letter'/exp OR letter)	1116315
#4	#1 AND #2 AND #3	123429
#5	'hemodilution'/exp OR hemodilution OR 'haemodilution'/exp OR haemodilution OR 'blood dilution'/exp OR 'blood dilution'	7739
#6	#8 AND #9	393

Interventions 2-4 - Intraoperative and postoperative cell salvage: Level I evidence

Search conducted 22 December 2009 (1966 to 30 July 2009)

Note: This search was used to locate Level I evidence for perioperative question 3, interventions 2–4.

#	Search	Results
#1	'surgery' OR 'surgery'/exp OR surgery OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR peroperat* OR postoperat* OR 'peroperative period'/exp OR 'peroperative period' OR 'postoperative period'/exp OR 'preoperative period' NOT [30-7-2009]/sd	13076417
#2	'blood'/exp OR 'blood' OR 'serum'/exp OR 'serum' OR 'plasma'/exp OR 'plasma' OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR 'bleeding' OR bleed* OR hemarthros* OR haemarthros* OR hemat* OR haemat* OR haemoperi* OR haemoperi* OR 'anemia'/exp OR 'anemia' OR 'anaemia'/exp OR 'anaemia' NOT [30-7-2009]/sd	2398903
#3	'meta analysis'/exp OR 'meta analysis' OR 'systematic review'/exp OR 'systematic review' OR 'pooled analysis' OR ('review'/exp OR 'review' AND ('meta analysis'/exp OR 'meta analysis' OR systemat* OR pool*)) NOT [30-7-2009]/sd	111681
#4	'blood salvage'/exp OR 'blood salvage' OR 'salvage therapy'/exp OR 'salvage therapy' OR 'cell salvage' OR 'erythrocyte salvage' OR 'cell saver' OR 'cell savers' NOT [30-7-2009]/sd	12817
#5	#1 AND #2 AND #3 AND #4	129

Intervention 2 - Intraoperative cell salvage: Level II evidence

Search conducted 3 January 2010 (January 2004 to 30 July 2009)

#	Search	Results
#1	'blood'/exp OR 'blood' OR 'serum'/exp OR 'serum' OR 'plasma'/exp OR 'plasma' OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR 'bleeding' OR bleed* OR hemarthros* OR haemarthros* OR haemat* OR haemat* OR haemoperi* OR haemoperi* OR 'anaemia'/exp OR 'anaemia' OR 'anaemia'/exp OR 'anaemia'	2471241
#2	'clinical trial'/exp OR 'clinical trial' OR 'randomized controlled trial'/exp OR 'randomized controlled trial' OR 'randomization'/exp OR randomization OR 'single blind procedure'/exp OR 'single blind procedure' OR 'double blind procedure'/exp OR 'double blind procedure'/exp OR 'triple blind procedure'/exp OR 'triple blind procedure'/exp OR 'crossover procedure'/exp OR 'crossover procedure'/exp OR placebo OR randomi?ed:ab,ti OR rct:ab,ti OR 'random allocation':ab,ti OR 'randomly allocated':ab,ti OR 'allocated randomly':ab,ti OR (allocated NEAR/2 random*):ab,ti OR 'single blind':ab,ti OR 'single blinded':ab,ti OR 'double blinded':ab,ti OR 'treble blind':ab,ti OR 'treble blinded':ab,ti OR 'triple blind':ab,ti OR 'triple blinded':ab,ti OR placebo*:ab,ti OR 'prospective study'/exp OR 'prospective study' NOT ('case study'/exp OR 'case study' OR 'case report':ab,ti OR 'abstract report'/exp OR 'abstract report' OR 'letter'/exp OR letter)	1116315
#3	'surgery' OR 'surgery'/exp OR surgery OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR peroperat* OR postoperat OR 'peroperative period'/exp OR 'peroperative period' OR 'postoperative period'/exp OR 'preoperative period' NOT [30-7-2009]/sd	13076415
#4	'blood salvage'/exp OR 'blood salvage' OR 'salvage therapy'/exp OR 'salvage therapy' OR 'cell salvage' OR 'erythrocyte salvage' OR 'cell saver' OR 'cell savers' AND [2004-2010]/py	6710
#5	#1 AND #2 AND #3 AND #4	971

Intervention 3 – Perioperative acute normovolemic haemodilution combined with intraoperative cell salvage: Level II evidence

Search conducted 3 January 2010 (1966 to 30 July 2009)

#	Search	Results
#1	'blood'/exp OR 'blood' OR 'serum'/exp OR 'serum' OR 'plasma'/exp OR 'plasma' OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR 'bleeding' OR bleed* OR hemarthros* OR haemarthros* OR haemat* OR haemat* OR haemoperi* OR haemoperi* OR 'anaemia'/exp OR 'anaemia' OR 'anaemia'/exp OR 'anaemia'	2471241
#2	'clinical trial'/exp OR 'clinical trial' OR 'randomized controlled trial'/exp OR 'randomized controlled trial' OR 'randomization'/exp OR randomization OR 'single blind procedure'/exp OR 'single blind procedure' OR 'double blind procedure'/exp OR 'double blind procedure' OR 'triple blind procedure'/exp OR 'triple blind procedure' OR 'crossover procedure'/exp OR 'crossover procedure' OR 'placebo'/exp OR placebo OR randomi?ed:ab,ti OR rct:ab,ti OR 'random allocation':ab,ti OR 'randomly allocated':ab,ti OR 'allocated randomly':ab,ti OR (allocated NEAR/2 random*):ab,ti OR 'single blind':ab,ti OR 'single blinded':ab,ti OR 'double blinded':ab,ti OR 'treble blind':ab,ti OR 'treble blinded':ab,ti OR 'triple blind':ab,ti OR 'triple blinded':ab,ti OR placebo*:ab,ti OR 'prospective study'/exp OR 'prospective study' NOT ('case study'/exp OR 'case study' OR 'case report':ab,ti OR 'abstract report'/exp OR 'abstract report' OR 'letter'/exp OR letter)	1116315
#3	'hemodilution'/exp OR hemodilution OR 'haemodilution'/exp OR haemodilution OR 'blood dilution'/exp OR 'blood dilution'	7739
#4	'surgery' OR 'surgery'/exp OR surgery OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR perioperat* OR postoperat* OR 'peroperative period'/exp OR 'peroperative period' OR 'postoperative period'/exp OR 'preoperative period' NOT [30-7-2009]/sd	13076415
#5	'blood salvage'/exp OR 'blood salvage' OR 'salvage therapy'/exp OR 'salvage therapy' OR 'cell salvage' OR 'erythrocyte salvage' OR 'cell saver' OR 'cell savers'	13585
#6	#1 AND #2 AND #3 AND #4 AND #5	60

Intervention 3 – Perioperative acute normovolemic haemodilution combined with intraoperative cell salvage: Level III and Level IV evidence

Search conducted 11 February 2010 (1966 to 30 July 2009)

#	Search	Results
#1	'blood'/exp OR 'blood' OR 'serum'/exp OR 'serum' OR 'plasma'/exp OR 'plasma' OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR 'bleeding' OR bleed* OR hemarthros* OR haemarthros* OR peroperat* OR peroperat* OR peroperat* OR peroperative period'/exp OR 'prospective period' OR 'postoperative period'/exp OR 'prospective period'/exp OR 'salvage therapy'/exp OR 'salvage therapy'/exp OR 'salvage 'OR 'cell	60
#2	'blood'/exp OR 'serum'/exp OR 'plasma'/exp OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR bleed* OR hemarthros* OR haemarthros* OR hemat* OR haemat* OR hemoperi* OR haemoperi* OR 'anemia'/exp OR 'anaemia'/exp	2459023
#3	'hemodilution'/exp OR 'haemodilution'/exp OR 'blood dilution'/exp	5588
#4	'surgery'/exp OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR perioperat* OR postoperat* OR 'perioperative period'/exp OR 'postoperative period'/exp OR 'preoperative period'/exp NOT [30-7-2009]/sd	13076029
#5	'blood salvage'/exp OR 'salvage therapy'/exp OR 'cell salvage' OR 'erythrocyte salvage' OR 'cell saver' OR 'cell savers'	11918
#6	#2 AND #3 AND #4 AND #5	130
#7	#6 NOT #1	85

Intervention 4 - Postoperative cell salvage: Level II evidence

Search conducted 3 January 2010 (1966 to 30 July 2009)

#	Search	Results
#1	'blood'/exp OR 'blood' OR 'serum'/exp OR 'serum' OR 'plasma'/exp OR 'plasma' OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR 'bleeding' OR bleed* OR hemarthros* OR haemarthros* OR haemarthros* OR haemart* OR haemoperi* OR haemoperi* OR 'anaemia'/exp OR 'anaemia' OR 'anaemia'/exp OR 'anaemia'	2471241
#2	'clinical trial'/exp OR 'clinical trial' OR 'randomized controlled trial'/exp OR 'randomized controlled trial' OR 'randomization'/exp OR randomization OR 'single blind procedure'/exp OR 'single blind procedure' OR 'double blind procedure'/exp OR 'triple blind procedure'/exp OR 'triple blind procedure'/exp OR 'crossover procedure'/exp OR 'crossover procedure' OR 'placebo'/exp OR placebo OR randomi?ed:ab,ti OR rct:ab,ti OR 'random allocation':ab,ti OR 'randomly allocated':ab,ti OR 'allocated randomly':ab,ti OR (allocated NEAR/2 random*):ab,ti OR 'single blind':ab,ti OR 'single blinded':ab,ti OR 'double blinded':ab,ti OR 'treble blind':ab,ti OR 'treble blinded':ab,ti OR 'triple blind':ab,ti OR 'triple blinded':ab,ti OR placebo*:ab,ti OR 'prospective study'/exp OR 'prospective study' NOT ('case study'/exp OR 'case study' OR 'case report':ab,ti OR 'abstract report'/exp OR 'abstract report' OR 'letter'/exp OR letter)	1116315
#3	'surgery' OR 'surgery'/exp OR surgery OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR peroperat* OR postoperat OR 'peroperative period'/exp OR 'peroperative period' OR 'postoperative period'/exp OR 'preoperative period' NOT [30-7-2009]/sd	13076415
#4	'blood salvage'/exp OR 'blood salvage' OR 'salvage therapy'/exp OR 'salvage therapy' OR 'cell salvage' OR 'erythrocyte salvage' OR 'cell saver' OR 'cell savers'	13585
#5	postoperative OR 'post operative'	691807
#6	#1 AND #2 AND #3 AND #4 AND #5	292

Intervention 5 – Deliberate induced hypotension: Level I evidence

Search conducted 21 December 2009 (1966 to 30 July 2009)

#	Search	Results
#1	'surgery' OR 'surgery'/exp OR surgery OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR peroperat* OR postoperat* OR 'peroperative period'/exp OR 'peroperative period' OR 'postoperative period'/exp OR 'postoperative period' OR 'preoperative period'/exp OR 'preoperative period' NOT [30-7-2009]/sd	13076417
#2	'blood'/exp OR 'blood' OR 'serum'/exp OR 'serum' OR 'plasma'/exp OR 'plasma' OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR 'bleeding' OR bleed* OR hemarthros* OR haemarthros* OR hemat* OR haemat* OR hemoperi* OR haemoperi* OR 'anemia'/exp OR 'anemia' OR 'anaemia'/exp OR 'anaemia' NOT [30-7-2009]/sd	2398903
#3	'meta analysis'/exp OR 'meta analysis' OR 'systematic review'/exp OR 'systematic review' OR 'pooled analysis' OR ('review'/exp OR 'review' AND ('meta analysis'/exp OR 'meta analysis' OR systemat* OR pool*)) NOT [30-7-2009]/sd	111681
#4	#1 AND #2 AND #3	20746
#5	'induced hypotension'/exp OR 'induced hypotension' OR 'controlled hypotension'/exp OR 'controlled hypotension' OR 'hypotensive anesthesia' OR 'hypotensive epidural anesthesia' OR 'hypotensive epidural anaesthesia' OR 'iatrogenic hypotension'/exp OR 'iatrogenic hypotension' NOT [30-6-2009]/sd	71749
#6	#4 AND #5	909

Intervention 5 - Deliberate induced hypotension: Level II evidence

Search conducted 5 January 2010 (1966 to 30 July 2009)

#	Search	Results
#1	'surgery' OR 'surgery'/exp OR surgery OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR peroperat* OR postoperat* OR 'peroperative period'/exp OR 'peroperative period' OR 'postoperative period' OR 'preoperative period'/exp OR 'preoperative period'	13433297
#2	'blood'/exp OR 'blood' OR 'serum'/exp OR 'serum' OR 'plasma'/exp OR 'plasma' OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR 'bleeding' OR bleed* OR hemarthros* OR haemarthros* OR haemat* OR haemoperi* OR haemoperi* OR 'anemia'/exp OR 'anemia' OR 'anaemia'/exp OR 'anaemia'	2471869
#3	'clinical trial'/exp OR 'clinical trial' OR 'randomized controlled trial'/exp OR 'randomized controlled trial' OR 'randomization'/exp OR randomization OR 'single blind procedure'/exp OR 'single blind procedure' OR 'double blind procedure'/exp OR 'double blind procedure' OR 'triple blind procedure'/exp OR 'triple blind procedure' OR 'crossover procedure'/exp OR 'crossover procedure' OR 'placebo'/exp OR placebo OR randomi?ed:ab,ti OR rct:ab,ti OR 'random allocation':ab,ti OR 'randomly allocated':ab,ti OR 'allocated randomly:ab,ti OR (allocated NEAR/2 random*):ab,ti OR 'single blind':ab,ti OR 'single blinded':ab,ti OR 'double blinded':ab,ti OR 'treble blind':ab,ti OR 'treble blinded':ab,ti OR 'triple blinded':ab,ti OR 'treble blinded':ab,ti OR 'prospective study'/exp OR 'prospective study' NOT ('case study'/exp OR 'case study' OR 'case report':ab,ti OR 'abstract report'/exp OR 'abstract report' OR 'letter'/exp OR letter)	1116696
#4	'induced hypotension'/exp OR 'induced hypotension' OR 'induced hypotension':ab,ti OR 'controlled hypotension':ab,ti OR 'iatrogenic hypotension':ab,ti OR 'hypotensive anaesthesia':ab,ti OR 'hypotensive epidural anaesthesia':ab,ti OR 'hypotensive epidural anaesthesia':ab,ti NOT [30-7-2009]/sd	4580
#5	#1 AND #2 AND #3 AND #4	257

Intervention 6 - Prevention of hypothermia: Level I evidence

Search conducted 21 December 2009 (1966 to 30 July 2009)

#	Search	Results
#1	'surgery' OR 'surgery'/exp OR surgery OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR peroperat* OR postoperat* OR 'peroperative period'/exp OR 'peroperative period' OR 'postoperative period'/exp OR 'postoperative period' OR 'preoperative period'/exp OR 'preoperative period' NOT [30-7-2009]/sd	13076417
#2	'blood'/exp OR 'blood' OR 'serum'/exp OR 'serum' OR 'plasma'/exp OR 'plasma' OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR 'bleeding' OR bleed* OR hemarthros* OR haemarthros* OR hemat* OR haemat* OR hemoperi* OR haemoperi* OR 'anemia'/exp OR 'anemia' OR 'anaemia'/exp OR 'anaemia' NOT [30-7-2009]/sd	2398903
#3	'meta analysis'/exp OR 'meta analysis' OR 'systematic review'/exp OR 'systematic review' OR 'pooled analysis' OR ('review'/exp OR 'review' AND ('meta analysis'/exp OR 'meta analysis' OR systemat* OR pool*)) NOT [30-7-2009]/sd	111681
#4	#1 AND #2 AND #3	20746
#5	'normothermia'/exp OR 'normothermia' OR 'thermoregulation'/exp OR 'thermoregulation' OR 'warming'/exp OR 'warming' OR 'hypothermia'/exp OR 'hypothermia' AND ('blood' OR 'blood'/exp OR blood OR hemorrhag* OR haemorrhag* OR 'anaemia' OR 'anaemia'/exp OR anaemia OR 'anemia' OR 'anemia' OR 'anaemia'/exp OR anaemia) NOT [30-7-2009]/sd	35595
#6	#4 and #5	158

Intervention 6 – Prevention of hypothermia: Level II evidence

Search conducted 5 January 2010 (1966 to 30 July 2009)

#	Search	Results
#1	'surgery' OR 'surgery'/exp OR surgery OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR peroperat* OR postoperat* OR 'peroperative period'/exp OR 'peroperative period' OR 'postoperative period' OR 'preoperative period'/exp OR 'preoperative period'	13433297
#2	'blood'/exp OR 'blood' OR 'serum'/exp OR 'serum' OR 'plasma'/exp OR 'plasma' OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR 'bleeding' OR bleed* OR hemarthros* OR haemarthros* OR haemat* OR haemoperi* OR haemoperi* OR 'anemia'/exp OR 'anaemia' OR 'anaemia'/exp OR 'anaemia'	2471869
#3	'clinical trial'/exp OR 'clinical trial' OR 'randomized controlled trial'/exp OR 'randomized controlled trial' OR 'randomization'/exp OR randomization OR 'single blind procedure'/exp OR 'single blind procedure' OR 'double blind procedure'/exp OR 'double blind procedure' OR 'triple blind procedure'/exp OR 'triple blind procedure' OR 'crossover procedure'/exp OR 'crossover procedure' OR 'placebo'/exp OR placebo OR randomi?ed:ab,ti OR rct:ab,ti OR 'random allocation':ab,ti OR 'randomly allocated':ab,ti OR 'allocated randomly:ab,ti OR (allocated NEAR/2 random*):ab,ti OR 'single blind':ab,ti OR 'double blind':ab,ti OR 'double blinded':ab,ti OR 'treble blind':ab,ti OR 'treble blind':ab,ti OR 'triple blinded':ab,ti OR 'triple blinded':ab,ti OR placebo*:ab,ti OR 'prospective study'/exp OR 'prospective study' NOT ('case study'/exp OR 'case study' OR 'case report':ab,ti OR 'abstract report'/exp OR 'abstract report' OR 'letter'/exp OR letter)	1116696
#4	'hypothermia'/exp/dm_pc OR 'hypothermia' OR 'normothermia':ab,ti OR 'thermoregulation':ab,ti OR 'warming':ab,ti OR 'hypothermia':ab,ti OR ('hypothermia'/exp OR 'hypothermia' AND ('perioperative complication'/exp/dm_pc OR 'perioperative complication' OR 'peroperative complication'/exp/dm_pc OR 'peroperative complication' OR 'postoperative complication' OR 'prevention'/exp OR 'prevention' OR 'prevention' OR 'primary prevention'/exp OR 'primary prevention')) NOT [30-7-2009]/sd	44694
#5	#1 AND #2 AND #3 AND #4	1002

Intervention 7 - Point-of-care testing: Level I and Level II evidence

Search conducted 22 December 2009 for any type of point-of-care test (1966 to 30 July 2009)

#	Search	Results
#1	'comparative study'/exp OR 'randomised controlled trial'/exp OR 'crossover procedure'/exp OR 'double blind procedure'/exp OR 'parallel design'/exp OR 'single blind procedure'/exp OR 'placebo'/exp OR comparative OR 'open label' OR placebo OR randomi* OR 'double blind' OR 'single blind' OR controlled OR single OR (double AND dummy)	6119239
#2	'meta analysis'/exp OR 'meta analysis' OR 'systematic review'/exp OR 'systematic review' OR 'pooled analysis' OR ('review'/exp OR 'review' AND ('meta analysis'/exp OR 'meta analysis' OR systemat* OR pool*))	118272
#3	#1 OR #2	6172408
#4	'blood'/exp OR 'blood' OR 'serum'/exp OR 'serum' OR 'plasma'/exp OR 'plasma' OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR 'bleeding' OR bleed* OR hemarthros* OR haemarthros* OR haemat* OR haemat* OR haemoperi* OR haemoperi* OR 'anemia'/exp OR 'anemia' OR 'anaemia'/exp OR 'anaemia' NOT [30-7-2009]/sd	2398903
#5	'point of care testing'/exp OR 'point of care testing' OR 'point of care' OR 'bedside' NEAR/3 'testing' OR 'bed side' NEAR/3 'testing' OR 'bedside' NEAR/3 'test' OR 'bedside' NEAR/3 'tests' OR 'bedside' NEAR/3 'tests' OR 'bed side' NEAR/3 'monitoring' OR 'bedside' NEAR/3 'computing' OR 'bedside' NEAR/3 'technology' OR 'bed side' NEAR/3 'technology' NOT [30-7-2009]/sd	5169
#6	#3 AND #4 AND #5	786

The preliminary literature search above found a limited body of comparative evidence for the effect of point-of-care testing other than thromboelastography (TEG) on mortality, morbidity and the need for allogeneic blood transfusion. A more focused literature search for specific tests was then conducted. The CRG subsequently made a decision to limit the scope of this intervention to comparative studies of TEG and TEG-based point-of-care tests.

Intervention 7 - Point-of-care testing: Level I-III evidence

Search conducted 2 February 2010 for specific point-of-care tests (1966 to 30 July 2009)

#	Search	Results
#1	'comparative study'/exp OR 'comparative study' OR 'randomised controlled trial'/exp OR 'randomised controlled trial' OR 'crossover procedure'/exp OR 'crossover procedure' OR 'double blind procedure'/exp OR 'double blind procedure' OR 'parallel design'/exp OR 'parallel design' OR 'single blind procedure'/exp OR 'single blind procedure' OR comparative OR 'open label' OR 'placebo'/exp OR 'placebo' OR randomi* OR 'double blind' OR 'single blind' OR controlled OR single OR (double AND dummy) OR 'meta analysis'/exp OR 'meta analysis' OR 'systematic review'/exp OR 'systematic review' OR 'pooled analysis' OR ('review'/exp OR 'review' AND ('meta analysis'/exp OR 'meta analysis' OR systemat* OR pool*)) AND ('blood'/exp OR 'blood' OR 'serum'/exp OR 'serum' OR 'plasma'/exp OR 'plasma' OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR 'bleeding' AND bleed* OR hemarthros* OR haemarthros* OR haemat* OR haemoperi* OR haemoperi* OR 'anemia'/exp OR 'anaemia' OR 'anaemia'/exp OR 'anaemia' OR 'bedside' NEAR/3 'testing' OR 'bedside' NEAR/3 'testing' OR 'bedside' NEAR/3 'test' OR 'bedside' NEAR/3 'tests' OR 'bedside' NEAR/3 'test' OR 'bedside' NEAR/3 'monitoring' OR 'bedside' NEAR/3 'tests' OR 'bedside' NEAR/3 'computing' OR 'bedside' NEAR/3 'technology' OR 'b	786
#2	'blood'/exp OR 'serum'/exp OR 'plasma'/exp OR hemorrh* OR haemorrh* OR 'bleeding'/exp AND bleed* OR hemarthros* OR haemarthros* OR haemat* OR haemat* OR haemoperi* OR haemoperi* OR 'anemia'/exp OR 'anaemia'/exp AND ('blood clotting parameters'/exp OR 'blood analysis'/exp OR 'blood examination'/exp)	100275
#3	'surgery'/exp OR 'surgery' OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR peroperat* OR postoperat* OR 'peroperative period'/exp OR 'peroperative period' OR 'postoperative period' OR 'preoperative period'/exp OR 'preoperative period' NOT [30-7-2009]/sd	13076023
#4	'teg':ab,ti OR 'sonoclot':ab,ti OR 'rotem':ab,ti OR 'roteg':ab,ti OR hemocue OR 'international normalised ratio':ab,ti OR 'hemoglobin test':ab,ti OR 'hb test':ab,ti OR 'thromboelastograph':ab,ti OR 'thromboelastography':ab,ti OR 'thromboelastography':ab,ti OR 'hemoglobin blood level'/exp OR 'hemoglobin blood level' OR 'hemoglobin blood level':ab,ti OR 'thrombelastography':ab,ti OR 'haemoglobin blood level'/exp OR 'haemoglobin blood level'	11233
#5	'meta analysis'/exp OR 'meta analysis' OR 'systematic review'/exp OR 'systematic review' OR 'pooled analysis' OR ('review'/exp OR 'review' AND ('meta analysis'/exp OR 'meta analysis' OR systemat* OR pool*)) OR 'comparative study'/exp OR 'comparative study' OR 'randomised controlled trial' OR 'crossover procedure'/exp OR 'crossover procedure' OR 'double blind procedure'/exp OR 'double blind procedure' OR 'parallel design'/exp OR 'parallel design' OR 'single blind procedure'/exp OR 'single blind procedure' OR comparative OR 'open label' OR 'placebo'/exp OR 'placebo' OR randomi* OR 'double blind' OR 'single blind' OR controlled OR single OR (double AND dummy)	6309867
#6	#1 OR (#2 AND #3 AND #4 AND #5)	1921

Intervention 8 - Administration of antifibrinolytics and DDAVP: Level I evidence

Search conducted 22 December 2009 (1966 to 30 July 2009)

#	Search	Results
#1	'surgery' OR 'surgery'/exp OR surgery OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR peroperat* OR postoperat* OR 'peroperative period'/exp OR 'peroperative period' OR 'postoperative period'/exp OR 'preoperative period' NOT [30-7-2009]/sd	13076417
#2	'blood'/exp OR 'blood' OR 'serum'/exp OR 'serum' OR 'plasma'/exp OR 'plasma' OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR 'bleeding' OR bleed* OR hemarthros* OR haemarthros* OR haemat* OR haemat* OR haemoperi* OR haemoperi* OR 'anemia'/exp OR 'anaemia' NOT [30-7-2009]/sd	2398903
#3	'meta analysis'/exp OR 'meta analysis' OR 'systematic review'/exp OR 'systematic review' OR 'pooled analysis' OR ('review'/exp OR 'review' AND ('meta analysis'/exp OR 'meta analysis' OR systemat* OR pool*)) NOT [30-7-2009]/sd	111681
#4	'antifibrinolytic agent'/exp OR 'antifibrinolytic agent' OR antifibrinolytic* OR 'anti fibrinolytic' OR 'anti fibrinolytics' OR antiplasmin* OR 'anti plasmin' OR 'anti plasmins' OR antifibrinolysin* OR 'anti fibrinolysis inhibitor'/exp OR 'fibrinolysis inhibitor' OR 'fibrinolysis inhibitors' OR 'plasmin inhibitor'/exp OR 'plasmin inhibitor'/exp OR 'plasmin inhibitor'/exp OR 'plasmin inhibitor'/exp OR 'glasmin inhibitor'/exp OR 'aminocaproic acid'/exp OR 'aminocaproic acid' OR 'eaca'/exp OR 'glasmin inhibitor'/exp OR 'aminocaproic acid' OR 'glasmin inhibitor'/exp OR 'aminocaproic acid'/exp OR 'aminocaproic acid' OR 'glasmin inhibitor'/exp OR 'aminocaproic acid' OR 'glasmin inhibitor'/exp OR 'aminocaproic acid' OR 'glasmin inhibitor'/exp OR 'aminocaproic acid'/exp OR 'aminocaproic acid'/exp OR 'aminocaproic acid' OR 'glasmin inhibitor'/exp OR 'aminocaproic acid'/exp OR 'aminocaproic acid' OR 'glasmin inhibitor'/exp OR 'aminocaproic acid'/exp OR 'aminocaproic acid' OR 'glasmin inhibitor'/exp OR 'aminocaproic acid' OR 'glasmin inhibitor'/exp OR 'aminocaproic acid'/exp OR 'a	33573
#5	#1 AND #2 AND #3 AND #4	388

Intervention 8 - Administration of antifibrinolytics and DDAVP: Level II evidence for aprotinin

Search conducted 21 April 2010 (1 July 2006 to 30 July 2009)

#	Search	Results
#1	'clinical trial'/exp OR 'clinical trial' OR 'randomized controlled trial'/exp OR 'randomized controlled trial' OR 'randomization'/exp OR randomization OR 'single blind procedure'/exp OR 'single blind procedure' OR 'double blind procedure'/exp OR 'triple blind procedure'/exp OR 'triple blind procedure' OR 'crossover procedure'/exp OR 'crossover procedure' OR 'placebo'/exp OR placebo OR randomi?ed:ab,ti OR rct:ab,ti OR 'random allocation':ab,ti OR 'randomly allocated':ab,ti OR 'allocated randomly':ab,ti OR (allocated NEAR/2 random*):ab,ti OR 'single blind:ab,ti OR 'single blinded':ab,ti OR 'double blinded':ab,ti OR 'treble blind':ab,ti OR 'treble blinded':ab,ti OR 'triple blinded':ab,ti OR 'treble blinded':ab,ti OR 'triple blinded':ab,ti OR 'prospective study'/exp OR 'prospective study' NOT ('case study'/exp OR 'case study' OR 'case report':ab,ti OR 'abstract report'/exp OR 'abstract report' OR 'letter'/exp OR letter)	1093468
#2	'surgery'/exp OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR perioperat* OR postoperat* OR 'peroperative period'/exp OR 'postoperative period'/exp OR 'preoperative period'/exp NOT [30-7-2009]/sd	13076116
#3	'blood'/exp OR 'serum'/exp OR 'plasma'/exp OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR bleed* OR hemarthros* OR haemarthros* OR hemat* OR haemat* OR hemoperi* OR haemoperi* OR 'anaemia'/exp OR 'anaemia'/exp	3541273
#4	'meta analysis'/exp OR 'meta analysis' OR 'systematic review'/exp OR 'systematic review' OR 'pooled analysis' OR ('review'/exp OR 'review' AND ('meta analysis'/exp OR 'meta analysis' OR systemat* OR pool*))	104108
#5	'aprotinin'/exp OR 'aprotinin' OR 'trasylol'/exp OR 'trasylol' OR 'antilysin'/exp OR 'antilysin' OR '11004 21 0':rn OR '12407 79 3':rn OR '50936 63 5':rn OR '52229 70 6':rn OR '58591 29 0':rn OR '9050 74 2':rn OR '9075 10 9':rn OR '9087 70 1':rn AND [1-1-2006]/sd	1728
#6	#1 AND #2 AND #3 AND #5	371
#7	#2 AND #3 AND #4 AND #5	81
#8	#6 NOT #7	301

Intervention 8 – Administration of antifibrinolytics and DDAVP: Level II evidence for tranexamic acid and ϵ -aminocaproic acid

Search conducted 24 February 2010 (1 July 2006 to 30 July 2009)

#	Search	Results
#1	'clinical trial'/exp OR 'clinical trial' OR 'randomized controlled trial'/exp OR 'randomized controlled trial' OR 'randomization'/exp OR randomization OR 'single blind procedure'/exp OR 'single blind procedure' OR 'double blind procedure'/exp OR 'triple blind procedure'/exp OR 'triple blind procedure' OR 'crossover procedure'/exp OR 'crossover procedure' OR 'placebo'/exp OR placebo OR randomi?ed:ab,ti OR rct:ab,ti OR 'random allocation':ab,ti OR 'randomly allocated':ab,ti OR 'allocated randomly:ab,ti OR (allocated NEAR/2 random*):ab,ti OR 'single blind:ab,ti OR 'single blinded':ab,ti OR 'double blind':ab,ti OR 'double blinded':ab,ti OR 'treble blind':ab,ti OR 'treble blinded':ab,ti OR 'triple blind':ab,ti OR 'triple blinded':ab,ti OR 'prospective study'/exp OR 'prospective study' NOT ('case study'/exp OR 'case study' OR 'case report':ab,ti OR 'abstract report'/exp OR 'abstract report' OR 'letter'/exp OR letter)	1136181
#2	'surgery'/exp OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR perioperat* OR perioperat* OR perioperat* OR 'postoperative period'/exp OR 'postoperative period'/exp NOT [30-7-2009]/sd	13076029
#3	'blood'/exp OR 'serum'/exp OR 'plasma'/exp OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR bleed* OR hemarthros* OR haemarthros* OR hemat* OR haemat* OR hemoperi* OR haemoperi* OR 'anaemia'/exp OR 'anaemia'/exp	3491096
#4	'meta analysis'/exp OR 'meta analysis' OR 'systematic review'/exp OR 'systematic review' OR 'pooled analysis' OR ('review'/exp OR 'review' AND ('meta analysis'/exp OR 'meta analysis' OR systemat* OR pool*))	121839
#5	'1319 82 0':rn OR '60 32 2':rn OR 'aminocaproic acid'/exp OR 'aminocaproic acid' OR 'eaca'/exp OR 'eaca' OR 'amicar'/exp OR 'amicar' OR '1197 18 8':rn OR '701 54 2':rn OR 'tranexamic acid'/exp OR 'tranexamic acid' OR 'cyklokapron'/exp OR 'cyklokapron' AND [1-7-2006]/sd	1600
#6	#1 AND #2 AND #3 AND #5	394
#7	#2 AND #3 AND #4 AND #5	88
#8	#6 NOT #7	321

Intervention 8 - Administration of antifibrinolytics and DDAVP: Level II evidence for DDAVP

Search conducted 16 February 2010 (January 2008 to 30 July 2009)

#	Search	Results
#1	'desmopressin'/exp OR 'desmopressin' OR 'ddavp'/exp OR 'ddavp' OR '16679 58 6':rn AND [2008-2010]/py	845
#2	'surgery'/exp OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR perioperat* OR postoperat* OR 'peroperative period'/exp OR 'postoperative period'/exp OR 'preoperative period'/exp NOT [30-7-2009]/sd	13076029
#3	'blood'/exp OR 'serum'/exp OR 'plasma'/exp OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR bleed* OR hemarthros* OR haemarthros*	3478770
#4	'clinical trial'/exp OR 'clinical trial' OR 'randomized controlled trial'/exp OR 'randomized controlled trial' OR 'randomization'/exp OR randomization OR 'single blind procedure'/exp OR 'single blind procedure' OR 'double blind procedure'/exp OR 'double blind procedure' OR 'triple blind procedure'/exp OR 'triple blind procedure' OR 'crossover procedure'/exp OR 'crossover procedure' OR 'placebo'/exp OR placebo OR randomi?ed:ab,ti OR rct:ab,ti OR 'random allocation':ab,ti OR 'randomly allocated':ab,ti OR 'allocated randomly':ab,ti OR (allocated NEAR/2 random*):ab,ti OR 'single blind':ab,ti OR 'single blinded':ab,ti OR 'double blinded':ab,ti OR 'treble blind':ab,ti OR 'triple blinded':ab,ti OR 'triple blinded':ab,ti OR 'triple blinded':ab,ti OR placebo*:ab,ti OR 'prospective study'/exp OR 'prospective study' NOT ('case study'/exp OR 'case study' OR 'case report':ab,ti OR 'abstract report'/exp OR 'abstract report' OR 'letter'/exp OR letter)	1132142
#5	'meta analysis'/exp OR 'systematic review'/exp OR 'pooled analysis' OR ('review'/exp AND ('meta analysis'/exp OR systemat* OR pool*))	100954
#6	#1 AND #2 AND #3 AND #4	84
#7	#1 AND #2 AND #3 AND #5	8
#8	#6 NOT #7	78

Intervention 9 – Appropriate patient positioning: Level I evidence

Search conducted 22 December 2009 (1966 to 30 July 2009)

#	Search	Results
#1	'surgery' OR 'surgery'/exp OR surgery OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR peroperat* OR postoperat* OR 'peroperative period'/exp OR 'peroperative period' OR 'postoperative period'/exp OR 'preoperative period' NOT [30-7-2009]/sd	13076417
#2	'blood'/exp OR 'blood' OR 'serum'/exp OR 'serum' OR 'plasma'/exp OR 'plasma' OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR 'bleeding' OR bleed* OR hemarthros* OR haemarthros* OR haemarthros* OR haemarthros* OR haemarthros* OR 'anaemia' OR 'anaemia'/exp OR 'anaemia' NOT [30-7-2009]/sd	2398903
#3	'meta analysis'/exp OR 'meta analysis' OR 'systematic review'/exp OR 'systematic review' OR 'pooled analysis' OR ('review'/exp OR 'review' AND ('meta analysis'/exp OR 'meta analysis' OR systemat* OR pool*)) NOT [30-7-2009]/sd	111681
#4	'patient positioning'/exp OR 'patient positioning' OR 'position'/exp OR 'position' OR 'semi sitting' OR 'operative positioning' OR 'lateral position' OR (patient* AND position*) NOT [30-7-2009]/sd	323990
#5	#1 AND #2 AND #3 AND #4	468

Intervention 9 – Appropriate patient positioning: Level II evidence

Search conducted 3 January 2010 (1966 to 30 July 2009)

#	Search	Results
#1	'blood'/exp OR 'blood' OR 'serum'/exp OR 'serum' OR 'plasma'/exp OR 'plasma' OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR 'bleeding' OR bleed* OR hemarthros* OR haemarthros* OR haemarthros* OR haemarthros* OR haemarthros* OR 'anaemia' OR 'anaemia' OR 'anaemia'	2471241
#2	'clinical trial'/exp OR 'clinical trial' OR 'randomized controlled trial'/exp OR 'randomized controlled trial' OR 'randomization'/exp OR randomization OR 'single blind procedure'/exp OR 'single blind procedure' OR 'double blind procedure'/exp OR 'triple blind procedure'/exp OR 'triple blind procedure'/exp OR 'triple blind procedure'/exp OR 'crossover procedure'/exp OR 'crossover procedure'/exp OR placebo OR randomi?ed:ab,ti OR rct:ab,ti OR 'random allocation':ab,ti OR 'randomly allocated':ab,ti OR 'allocated randomly':ab,ti OR (allocated NEAR/2 random*):ab,ti OR 'single blind':ab,ti OR 'single blinded':ab,ti OR 'double blinded':ab,ti OR 'treble blind':ab,ti OR 'treble blinded':ab,ti OR 'triple blind':ab,ti OR 'triple blinded':ab,ti OR placebo*:ab,ti OR 'prospective study'/exp OR 'prospective study' NOT ('case study'/exp OR 'case study' OR 'case report':ab,ti OR 'abstract report'/exp OR 'abstract report' OR 'letter'/exp OR letter)	1116315
#3	'surgery' OR 'surgery'/exp OR surgery OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR peroperat* OR postoperat OR 'peroperative period'/exp OR 'peroperative period' OR 'postoperative period'/exp OR 'preoperative period' NOT [30-7-2009]/sd	13076415
#4	'patient positioning'/exp OR 'patient positioning' OR 'position'/exp OR 'position' OR 'semi sitting' OR 'operative positioning' OR 'lateral position' OR (patient* AND position*) AND ('blood' OR 'blood'/exp OR blood OR hemorrhag* OR haemorrhag* OR 'anaemia' OR 'anaemia'/exp OR anaemia OR 'anemia' OR 'anaemia'/exp OR anaemia)	56266
#5	#1 AND #2 AND #3 AND #4	1640

Intervention 10 - Preoperative autologous donation: Level I evidence

Search conducted 22 December 2009 (1966 to 30 July 2009)

#	Search	Results
#1	'surgery' OR 'surgery'/exp OR surgery OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR peroperat* OR postoperat* OR 'peroperative period'/exp OR 'peroperative period' OR 'postoperative period'/exp OR 'preoperative period' NOT [30-7-2009]/sd	13076417
#2	'blood'/exp OR 'blood' OR 'serum'/exp OR 'serum' OR 'plasma'/exp OR 'plasma' OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR 'bleeding' OR bleed* OR hemarthros* OR haemarthros* OR hemat* OR haemat* OR haemoperi* OR haemoperi* OR 'anaemia'/exp OR 'anaemia' OR 'anaemia'/exp OR 'anaemia' NOT [30-7-2009]/sd	2398903
#3	'meta analysis'/exp OR 'meta analysis' OR 'systematic review'/exp OR 'systematic review' OR 'pooled analysis' OR ('review'/exp OR 'review' AND ('meta analysis'/exp OR 'meta analysis' OR systemat* OR pool*)) NOT [30-7-2009]/sd	111681
#4	#1 AND #2 AND #3	20746
#5	'autohemotransfusion' OR 'autohemotransfusion'/exp OR autohemotransfusion OR autohaemotransfusion OR 'autotransfusion' OR 'autotransfusion'/exp OR autotransfusion OR autotransfus* OR ('blood' OR 'blood'/exp OR blood OR 'plasma' OR 'plasma'/exp OR plasma AND (autologous* OR predonat* OR donat* OR predeposit*)) NOT [30-7-2009]/sd	101841
#6	#4 AND #5	479

Intervention 10 - Preoperative autologous donation: Level II evidence

Search conducted 3 January 2010 (January 2004 to 30 July 2009)

#	Search	Results
#1	'blood'/exp OR 'blood' OR 'serum'/exp OR 'serum' OR 'plasma'/exp OR 'plasma' OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR 'bleeding' OR bleed* OR hemarthros* OR haemarthros* OR haemat* OR haemoperi* OR haemoperi* OR 'an	2471241
#2	'clinical trial'/exp OR 'clinical trial' OR 'randomized controlled trial'/exp OR 'randomized controlled trial' OR 'randomization'/exp OR randomization OR 'single blind procedure'/exp OR 'single blind procedure' OR 'double blind procedure'/exp OR 'double blind procedure' OR 'triple blind procedure'/exp OR 'triple blind procedure' OR 'crossover procedure'/exp OR 'crossover procedure' OR 'placebo'/exp OR placebo OR randomi?ed:ab,ti OR rct:ab,ti OR 'random allocation':ab,ti OR 'randomly allocated':ab,ti OR 'allocated randomly:ab,ti OR (allocated NEAR/2 random*):ab,ti OR 'single blind':ab,ti OR 'single blinded':ab,ti OR 'double blinded':ab,ti OR 'treble blind':ab,ti OR 'treble blinded':ab,ti OR 'triple blinded':ab,ti OR 'triple blinded':ab,ti OR placebo*:ab,ti OR 'prospective study'/exp OR 'prospective study' NOT ('case study'/exp OR 'case study' OR 'case report':ab,ti OR 'abstract report'/exp OR 'abstract report' OR 'letter'/exp OR letter)	1116315
#3	'surgery' OR 'surgery'/exp OR surgery OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR perioperat* OR postoperat* OR 'peroperative period'/exp OR 'peroperative period' OR 'postoperative period'/exp OR 'preoperative period' NOT [30-7-2009]/sd	13076415
#4	'autohemotransfusion' OR 'autohemotransfusion'/exp OR autohemotransfusion OR 'autotransfusion' OR 'autotransfusion'/exp OR autotransfusion OR autotransfus* OR (autologous* AND transfus*) AND [2004-2010]/py	4413
#5	#1 AND #2 AND #3 AND #4	927

Perioperative Question 3 – Quality of life: not limited by study type

Search conducted 14 February 2010 (1966 to 30 July 2009)

Note: Higher levels of evidence did not capture quality of life as an outcome for any of the interventions in perioperative question 3. This literature search was intended to identify clinical studies of any type that are relevant to perioperative question 3 and report quality of life as a key outcome.

#	Search	Results
#1	'quality of life'/exp	144031
#2	'surgery'/exp OR surger* OR surgical* OR transplant* OR reconstruct* OR procedur* OR operat* OR preoperat* OR intraoperat* OR perioperat* OR perioperat* OR postoperat* OR 'peroperative period'/exp OR 'postoperative p	13076029
#3	'blood'/exp OR 'serum'/exp OR 'plasma'/exp OR hemorrh* OR haemorrh* OR 'bleeding'/exp OR bleed* OR hemarthros* OR haemarthros* OR haemat* OR haemat* OR haemoperi* OR 'anemia'/exp OR 'anaemia'/exp	3475166
#4	#1 AND #2 AND #3	17977
#5	'hemodilution'/exp OR 'haemodilution'/exp OR 'blood dilution'/exp	5588
#6	'blood salvage'/exp OR 'salvage therapy'/exp OR 'cell salvage' OR 'erythrocyte salvage' OR 'cell saver' OR 'cell savers'	11923
#7	'induced hypotension'/exp OR 'controlled hypotension'/exp OR 'hypotensive anesthesia' OR 'hypotensive anaesthesia' OR 'hypotensive epidural anesthesia' OR 'hypotensive epidural anaesthesia' OR 'iatrogenic hypotension'/exp	74657
#8	'normothermia'/exp OR 'thermoregulation'/exp OR 'warming'/exp OR 'hypothermia'/exp	84849
#9	'antifibrinolytic agent'/exp OR antifibrinolytic* OR 'anti fibrinolytic' OR 'anti fibrinolytics' OR antiplasmin* OR 'anti plasmin' OR 'anti plasmins' OR antifibrinolysin* OR 'anti fibrinolysin' OR 'anti fibrinolysin' OR 'anti fibrinolysin' OR 'anti plasmin' OR 'fibrinolysis inhibitors' OR 'plasmin inhibitor'/exp OR 'plasmin inhibitor'/exp OR 'plasmin inhibitor'/exp OR 'amicar'/exp OR 'tranexamic acid'/exp OR 'cyklokapron'/exp OR 'aminocaproic acid'/exp OR 'eaca'/exp OR 'amicar'/exp OR 'aprotinin'/exp OR 'trasylol'/exp OR 'antilysin'/exp OR 'desmopressin'/exp OR 'ddavp'/exp OR '1197 18 8':rn OR '701 54 2':rn OR '1319 82 0':rn OR '60 32 2':rn OR '11004 21 0':rn OR '12407 79 3':rn OR '50936 63 5':rn OR '52229 70 6':rn OR '58591 29 0':rn OR '9050 74 2':rn OR '9075 10 9':rn OR '9087 70 1':rn OR '16679 58 6':rn OR '62288 83 9':rn OR '62357 86 2'	30091
#10	'patient positioning'/exp OR 'position'/exp OR 'semi sitting' OR 'operative positioning' OR 'lateral position' OR (patient* AND position*)	156770
#11	'autohemotransfusion'/exp OR autohaemotransfusion OR 'autotransfusion'/exp OR autotransfus* OR ('blood'/exp OR 'plasma'/exp AND (autologous* OR predonat* OR donat* OR predeposit* OR deposit*))	39281
#12	#5 OR #6 OR #7 OR #8 OR #9 OR #10 OR #11	393199
#13	#4 AND #12	1173

Cochrane Library

Intervention 1 – Acute normovolemic haemodilution: Level I evidence

Search conducted 21 December 2009

ID	Search	Results
#1	MeSH descriptor Hemodilution , this term only	341
#2	(acute AND (normovolemic OR normovolaemic))	157
#3	(acute AND ("normo volemic" OR "normo volaemic"))	0
#4	(acute NEAR/2 ("normovolemic hemodilution" OR "normovolemic haemodilution"))	102
#5	(acute NEAR/2 ("normovolaemic hemodilution" OR "normovolaemic haemodilution"))	50
#6	(acute NEAR/2 ("normo volemic hemodilution" OR "normo volemic haemodilution"))	0
#7	(acute NEAR/2 ("normo volaemic hemodilution" OR "normo volaemic haemodilution"))	0
#8	#1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7	408

Citations identified in 'Cochrane reviews', 'Database of systematic reviews' and 'Technological assessments' were exported into reference manager. Total number of citations exported: 25

Intervention 1 - Acute normovolemic haemodilution: Level II evidence

Search conducted 3 January 2010

ID	Search	Results
#1	MeSH descriptor Hemodilution , this term only	346
#2	acute AND (normovolemic OR normovolaemic OR 'normo volemic' OR 'normo volaemic')	158
#3	(acute NEAR/2 ("normovolemic hemodilution" OR "normovolemic haemodilution"))	103
#4	(acute NEAR/2 ("normovolaemic hemodilution" OR "normovolaemic haemodilution"))	51
#5	(acute NEAR/2 ("normo volemic hemodilution" OR "normo volemic haemodilution"))	0
#6	(acute NEAR/2 ("normo volaemic hemodilution" OR "normo volaemic haemodilution"))	0
#7	(#1 OR #2 OR #3 OR #4 OR #5 OR #6), from 2002 to 2010	141

Citations identified in 'Central register of clinical trials' exported into reference manager. Number of citations exported: 116

Interventions 2-4 - Intraoperative and postoperative cell salvage: Level I evidence

Search conducted 22 December 2009

Note: This search was used to locate Level I evidence for Perioperative question 3 interventions 2 to 4.

ID	Search	Results
#1	MeSH descriptor Salvage Therapy explode all trees	365
#2	"blood salvage" OR "salvage therapy" OR "cell salvage" OR "erythrocyte salvage" OR "cell saver" OR "Cell savers"	742
#3	#1 OR #2	742

Citations identified in 'Cochrane Reviews', 'Database of systematic reviews' and 'Technological assessments' were exported into reference manager. Total number of citations exported: 72

Interventions 2-4 - Intraoperative and postoperative cell salvage: Level II evidence

Search conducted 3 January 2010

ID	Search	Results
#1	MeSH descriptor Salvage Therapy explode all trees	365
#2	"blood salvage" OR "salvage therapy" OR "cell salvage" OR "erythrocyte salvage" OR "cell saver" OR "Cell savers"	742
#3	(#1 OR #2), from 2004 to 2010	314

Citations identified in 'Central register of clinical trials' were exported into reference manager. Total number of citations exported: 228

Intervention 3 – Perioperative acute normovolemic haemodilution combined with intraoperative cell salvage: Level II evidence

Search conducted 3 January 2010

ID	Search	Results
#1	MeSH descriptor Salvage Therapy explode all trees	365
#2	"blood salvage" OR "salvage therapy" OR "cell salvage" OR "erythrocyte salvage" OR "cell saver" OR "Cell savers"	742
#3	#1 OR #2	742
#4	MeSH descriptor Hemodilution , this term only	341
#5	(acute AND (normovolemic OR normovolaemic))	157
#6	(acute AND ("normo volemic" OR "normo volaemic"))	0
#7	(acute NEAR/2 ("normovolemic hemodilution" OR "normovolemic haemodilution"))	102
#8	(acute NEAR/2 ("normovolaemic hemodilution" OR "normovolaemic haemodilution"))	50
#9	(acute NEAR/2 ("normo volemic hemodilution" OR "normo volemic haemodilution"))	0
#10	(acute NEAR/2 ("normo volaemic hemodilution" OR "normo volaemic haemodilution"))	0
#11	#4 OR #5 OR #6 OR #7 OR #8 OR #9 OR #10	408
#12	#3 AND #11	31

Citations identified in 'Central register of clinical trials' were exported into reference manager. Total number of citations exported: 17

Intervention 5 - Deliberate induced hypotension: Level I and Level II evidence

Search conducted 21 December 2009

ID	Search	Results
#1	MeSH descriptor Hypotension explode all trees with qualifier: PC	258
#2	"induced hypotension"	305
#3	"controlled hypotension" OR "iatrogenic hypotension"	119
#4	(#1 OR #2 OR #3)	614

Citations identified in 'Cochrane reviews', 'Database of systematic reviews', and 'Central register of clinical trials' were exported into reference manager. Total number of citations exported: 613

Intervention 6 - Prevention of hypothermia: Level I and Level II evidence

Search conducted 22 December 2009

#	Search	Results
#1	MeSH descriptor Hypothermia explode all trees with qualifier: PC	141
#2	(hypothermia AND prevent*):ti	57
#3	(prevent* NEAR/20 hypothermia)	347
#4	(#1 OR #2 OR #3)	347

Citations identified in 'Cochrane reviews', 'Database of systematic reviews', and 'Central register of clinical trials' were exported into reference manager. Total number of citations exported: 342

Intervention 7 - Point-of-care testing: Level I and Level II evidence

Search conducted 22 December 2009 for any point-of-care tests

ID	Search	Results
#1	MeSH descriptor Point-of-Care Systems, this term only	234
#2	"point of care"	343
#3	(bedside OR "bed side") NEAR/3 (test OR tests OR testing)	51
#4	(bedside OR "bed side") NEAR/3 monitoring	36
#5	(bedside OR "bed side") NEAR/3 computing	0
#6	(bedside OR "bed side") NEAR/3 technology	9
#7	#1 OR #2 OR #3 OR #4 OR #5 OR #6	405

Citations identified in 'Cochrane Reviews', 'Database of systematic reviews', 'Central register of clinical trials', and 'Technological assessments' were exported into reference manager. Total number of citations exported: 64

Intervention 7 – Point-of-care testing using thromboelastography: Level I and Level II evidence

Search conducted 2 February 2010 for specific point-of-care tests

ID	Search	Results
#1	MeSH descriptor Thrombelastography explode all trees	114
#2	Sonoclot	8
#3	rotem	20
#4	roteg	5
#5	"international normalized ratio"	536
#6	"international normalised ratio"	137
#7	"haemoglobin test"	2
#8	"hemoglobin test"	4
#9	surgery	84598
#10	transplant*	17598
#11	reconstruct*	3403
#12	operat*	33199
#13	preoperat*	13310
#14	intraoperat*	8780
#15	perioperat*	5575
#16	peroperat*	506
#17	postoperat*	43291
#18	(#9 OR #10 OR #11 OR #12 OR #13 OR #14 OR #15 OR #16 OR #17)	119171
#19	(#1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7 OR #8)	780
#20	(#18 AND #19)	265

Citations identified in 'Cochrane Reviews', 'Database of systematic reviews', 'Central register of clinical trials', and 'Technological Assessments' were exported into reference manager. Total number of citations exported: 243

Intervention 8 – Administration of antifibrinolytics & DDAVP: Level I evidence

Search conducted 22 December 2009

ID	Search	Results
#1	MeSH descriptor Antifibrinolytic Agents, this term only	299
#2	MeSH descriptor Tranexamic Acid, this term only	253
#3	MeSH descriptor Aprotinin , this term only	497
#4	MeSH descriptor Deamino Arginine Vasopressin, this term only	274
#5	(antifibrinolytic* OR "anti fibrinolytic" OR "anti fibrinolytics")	495
#6	(antiplasmin* OR "anti plasmin" OR "anti plasmins")	260
#7	(antifibrinolysin* OR "anti fibrinolysin" OR "anti fibrinolysins")	3
#8	"fibrinolysis inhibitor" OR "fibrinolysis inhibitors"	33
#9	"plasmin inhibitor" OR "plamin inhibitors"	54
#10	"tranexamic acid" OR Cyklokapron	450
#11	"aminocaproic acid" OR eaca OR Amicar	172
#12	(aprotinin OR Trasylol OR antilysin)	783
#13	(desmopressin OR ddavp)	454
#14	#1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #8 OR #9 OR #10 OR #11 OR #12 OR #13	2072

Citations identified in 'Cochrane Reviews', 'Database of systematic reviews' and 'Technological assessments' were exported into reference manager. Total number of citations exported: 106

Intervention 8 - Administration of antifibrinolytics & DDAVP: Level II evidence for aprotinin

Search conducted 22 April 2010

ID	Search	Results
#1	aprotinin OR trasylol OR antilysin, from 2006 to 2009	78

Citations identified in 'Central register of clinical trials' were exported into reference manager. Total number of citations exported: 49

Intervention 8 – Administration of antifibrinolytics & DDAVP: Level II evidence for tranexamic acid and ε-aminocaproic acid

Search conducted 17 February 2010

ID	Search	Results
#1	"tranexamic acid" OR Cyklokapron OR "aminocaproic acid" OR eaca OR Amicar, from 2006 to 2009	125

Citations identified in 'Central register of clinical trials' were exported into reference manager. Total number of citations exported: 82

Intervention 8 - Administration of antifibrinolytics & DDAVP: Level II evidence for DDAVP

Search conducted 17 February 2010

ID	Search	Results
#1	MeSH descriptor Deamino Arginine Vasopressin , this term only	275
#2	desmopressin OR ddavp	457
#3	(#1 OR #2), from 2008 to 2010	39

Citations identified in 'Central register of clinical trials' were exported into reference manager. Total number of citations exported: 17

Intervention 9 - Appropriate patient positioning: Level I evidence

Search conducted 22 December 2009

ID	Search	Results
#1	MeSH descriptor Posture explode all trees	2746
#2	(patient OR patients) AND position*:ti	732
#3	(patient OR patients) NEAR/20 position*	2585
#4	(#35 OR #36 OR #37)	4856

Citations identified in 'Cochrane Reviews', 'Database of systematic reviews' and 'Technological assessments' were exported into reference manager. Total number of citations exported: 257

Intervention 9 – Appropriate patient positioning: Level II evidence

Search conducted 3 January 2010

ID	Search	Results
#1	MeSH descriptor Posture explode all trees	2746
#2	(patient OR patients) AND position*:ti	732
#3	(patient OR patients) NEAR/20 position*	2585
#4	(#1 OR #2 OR #3)	4856
#5	blood OR serum OR plasma OR hemorrh* OR haemorrh* OR bleed* OR hemarthros* OR haemorthros* OR hemat* OR haemat* or hemoperi* OR haemoperi* OR anemia	190593
#6	(#4 AND #5)	1807
#7	posture	3447
#8	(minimis* NEAR/5 ("blood loss" OR transfusion*))	28
#9	(minimiz* NEAR/5 ("blood loss" OR transfusion*))	53
#10	(reduc* NEAR/5 ("blood loss" OR transfusion*))	1529
#11	(minimis* AND ("blood loss" OR transfusion*)):ti	12
#12	(minimiz* AND ("blood loss" OR transfusion*)):ti	14
#13	(reduc* AND ("blood loss" OR transfusion*)):ti	391
#14	(#8 OR #9 OR #10 OR #11 OR #12 OR #13)	1595
#15	(#4 AND #14)	10

Citations identified in 'Central register of clinical trials' were exported into reference manager. Total number of citations exported: 6

Intervention 10 - Preoperative autologous donation: Level I and Level II evidence

Search conducted 22 December 2009

ID	Search	Results
#1	MeSH descriptor Blood Transfusion, Autologous explode all trees	586
#2	autologous	3986
#3	MeSH descriptor Blood Donors, this term only	267
#4	donor* OR donation* OR deposit* OR collection	17709
#5	#3 OR #4	17709
#6	#5 AND #2	787
#7	autologous NEAR/1 donor*	16
#8	(autologous NEAR/1 (predeposit* OR "pre deposit" OR "pre deposits"))	17
#9	(Autologous NEAR/1 Predonation*)	10
#10	("predeposit autologous" OR "pre deposit autologous") NEAR/1 donation	0
#11	"predonated autologous blood" OR "pre donated autologous blood"	11
#12	(predonation OR "pre donation") NEAR/1 "autologous blood"	1
#13	("preoperative autologous" OR "pre operative autologous") NEAR/1 deposit*	0
#14	("preoperative autologous" OR "pre operative autologous") NEAR/1 donation	1
#15	("preoperative donation" OR "pre operative donation") NEAR/1 "autologous blood"	0
#16	"preoperatively donated autologous blood" OR "pre operatively donated autologous blood"	4
#17	(#6 OR #7 OR #8 OR #9 OR #10 OR #11 OR #12 OR #13 OR #14 OR #15 OR #16), from 2004 to 2010	260

Citations identified in 'Cochrane Reviews', 'Database of systematic reviews', 'Technological Assessments', and 'Central register of clinical trials' were exported into reference manager. Total number of citations exported: 293

CINAHL (Nursing and Allied Health)

Search conducted 19 June 2009 (1982 to present)

ID	Query	Results
S114	S32 or S43 or S51 or S56 or S60 or S68 or S82 or S87 or S113	293
S113	S17 and S112	51
S112	S96 or S97 or S98 or S99 or S100 or S101 or S102 or S103 or S104 or S105 or S106 or S107 or S108 or S109 or S110 or S111	95
S111	TI ("preoperatively donated autologous blood" OR "pre operatively donated autologous blood") or AB ("preoperatively donated autologous blood")	1
S110	TI "pre operative donation" N1 "autologous blood" or AB "pre operative donation" N1 "autologous blood"	0
S109	TI "preoperative donation" N1 "autologous blood" or AB "preoperative donation" N1 "autologous blood"	2
S108	TI "pre operative autologous" N1 donation or AB "pre operative autologous" N1 donation	1
S107	TI "preoperative autologous" N1 donation or AB "preoperative autologous" N1 donation	32
S106	TI "pre operative autologous" N1 deposit* or AB "pre operative autologous" N1 deposit*	0
S105	TI "preoperative autologous" N1 deposit* or AB "preoperative autologous" N1 deposit*	1
S104	TI "pre donation" N1 "autologous blood" or AB "pre donation" N1 "autologous blood"	1
S103	TI predonation N1 "autologous blood" or AB predonation N1 "autologous blood"	0
S102	TI ("predonated autologous blood" OR "pre donated autologous blood") or AB ("predonated autologous blood" OR "pre donated autologous blood")	0
S101	TI "pre deposit autologous" N1 donation or AB "pre deposit autologous" N1 donation	1
S100	TI "predeposit autologous" N1 donation or AB "predeposit autologous" N1 donation	0
S99	TI Autologous N1 Predonation* or AB Autologous N1 Predonation*	5
S98	TI (autologous N1 (predeposit* OR "pre deposit" OR "pre deposits")) or AB (autologous N1 (predeposit* OR "pre deposit" OR "pre deposit" OR "pre deposit")	4
S97	TI autologous N1 donor* or AB autologous N1 donor*	18
S96	S90 and S93 and S95	73
S95	S7 or S9 or S94	12748
S94	(MH "Preoperative Care+")	6957
S93	S91 or S92	24193
S92	TI (donor* OR donation* OR deposit* OR collection) or AB (donor* OR donation* OR deposit* OR collection)	23720
S91	(MH "Blood Donors")	1344
S90	S88 or S89	1992
S89	TI autologous or AB autologous	1723
S88	(MH "Blood Transfusion, Autologous")	470
S87	S18 and S86	11
S86	S83 or S84 or S85	11301
S85	TI (patients AND position*) or AB (patients AND position*)	5798
S84	TI (patient AND position*) or AB (patient AND position*)	3424
S83	(MH "Patient Positioning+")	4774
S82	S18 and S81	100
S81	S69 or S70 or S71 or S72 or S73 or S74 or S75 or S76 or S77 or S78 or S79 or S80	722
S80	TI (desmopressin OR ddavp) or AB (desmopressin OR ddavp)	128

S79	TI (aprotinin OR Trasylol OR antilysin) or AB (aprotinin OR Trasylol OR antilysin)	202
S78	TI ("aminocaproic acid" OR eaca OR Amicar) or AB ("aminocaproic acid" OR eaca OR Amicar)	54
S77	TI ("tranexamic acid" OR Cyklokapron) or AB ("tranexamic acid" OR Cyklokapron)	82
S76	TI ("plasmin inhibitor" OR "plamin inhibitors") or AB ("plasmin inhibitor" OR "plamin inhibitors")	9
S75	TI ("fibrinolysis inhibitor" OR "fibrinolysis inhibitors") or AB ("fibrinolysis inhibitor" OR "fibrinolysis inhibitors")	21
S74	TI (antifibrinolysin* OR "anti fibrinolysin" OR "anti fibrinolysins") or AB (antifibrinolysin* OR "anti fibrinolysin" OR "anti fibrinolysins")	0
S73	TI (antiplasmin* OR "anti plasmin" OR "anti plasmins") or AB (antiplasmin* OR "anti plasmin" OR "anti plasmins")	41
S72	TI (antifibrinolytic* OR "anti fibrinolytic" OR "anti fibrinolytics") or AB (antifibrinolytic* OR "anti fibrinolytic" OR "anti fibrinolytics")	76
S71	(MH "Desmopressin")	177
S70	(MH "Aprotinin")	207
S69	(MH "Antifibrinolytic Agents")	167
S68	S18 and S67	55
S67	S61 or S62 or S63 or S64 or S65 or S66	76034
S66	TI ("bed side" N3 (monitoring OR computing OR technology)) or AB ("bed side" N3 (monitoring OR computing OR technology))	22000
S65	TI (bedside N3 (monitoring OR computing OR technology)) or AB (bedside N3 (monitoring OR computing OR technology))	22108
S64	TI ("bed side" N3 (test OR tests OR testing)) or AB ("bed side" N3 (test OR tests OR testing))	53967
S63	TI (bedside N3 (test OR tests OR testing)) or AB (bedside N3 (test OR tests OR testing))	54015
S62	TI "point of care" or AB "point of care"	1030
S61	(MH "Point-of-Care Testing")	909
S60	S18 and S59	10
S59	S57 or S58	516
S58	TI (hypothermia AND prevent*) or AB (hypothermia AND prevent*)	215
S57	(MH "Hypothermia/PC")	382
S56	S18 and S55	10
S55	S52 or S53 or S54	256
S54	TI ("controlled hypotension" OR "iatrogenic hypotension") or AB ("controlled hypotension" OR "iatrogenic hypotension")	13
S53	TI "induced hypotension" or AB "induced hypotension"	55
S52	(MH "Hypotension+/CI")	203
S51	S17 and S50	73
S50	S46 or S47 or S48 or S49	200
S49	TI ("postoperative blood salvage" OR "post operative blood salvage") or AB ("postoperative blood salvage" OR "post operative blood salvage")	14
S48	TI "post operative" N2 "cell salvage" or AB "post operative" N2 "cell salvage"	1
S47	TI postoperative N2 "cell salvage" or AB postoperative N2 "cell salvage"	3
S46	S35 and S45	199
S45	S3 or S4 or S12 or S44	35418
S44	(MH "Postoperative Care+")	7051
S43	S17 and S42	87

S42	S38 or S39 or S40 or S41	167
S41	TI ("intraoperative blood salvage" OR "intra operative blood salvage") or AB ("intraoperative blood salvage" OR "intra operative blood salvage")	10
S40	TI "intra operative" N2 "cell salvage" or AB "intra operative" N2 "cell salvage"	2
S39	TI intraoperative N2 "cell salvage" or AB intraoperative N2 "cell salvage"	6
S38	S35 and S37	164
S37	S5 or S6 or S10 or S36	6697
S36	(MH "Intraoperative Care+")	2766
S35	S33 or S34	1006
S34	(MH "Salvage Therapy")	461
S33	(MH "Blood Salvage+")	546
S32	S18 and S31	13
S31	S23 or S24	31
S30	TI acute N2 "normo volaemic haemodilution" or AB acute N2 "normo volaemic haemodilution"	0
S29	TI acute N2 "normo volaemic hemodilution" or AB acute N2 "normo volaemic hemodilution"	0
S28	TI (acute N2 ("normo volaemic hemodilution" OR "normo volaemic haemodilution") or AB (acute N2 ("normo volaemic hemodilution" OR "normo volaemic haemodilution")	0
S27	TI acute N2 "normo volemic haemodilution" or AB acute N2 "normo volemic haemodilution"	0
S26	TI acute N2 "normovolaemic hemodilution" or AB acute N2 "normovolaemic hemodilution"	0
S25	TI (acute N2 ("normovolaemic hemodilution" OR "normovolaemic haemodilution") or AB (acute N2 ("normovolaemic hemodilution" OR "normovolaemic haemodilution")	0
S24	TI (acute N2 ("normovolemic hemodilution" OR "normovolemic haemodilution")) or AB (acute N2 ("normovolemic hemodilution" OR "normovolemic haemodilution"))	27
S23	S19 and S22	27
S22	S20 or S21	36
S21	TI (acute AND ("normo volemic" OR "normo volaemic")) or AB (acute AND ("normo volemic" OR "normo volaemic"))	0
S20	TI (acute AND (normovolemic OR normovolaemic)) or AB (acute AND (normovolemic OR normovolaemic))	36
S19	(MH "Hemodilution")	158
S18	S13 and S17	839
S17	S14 or S15 or S16	4531
S16	TI (reduc* n5 ("blood loss" OR transfusion*)) or AB (reduc* N5 ("blood loss" OR transfusion*))	4510
S15	TI (minimiz* N5 ("blood loss" OR transfusion*)) or AB (minimiz* N5 ("blood loss" OR transfusion*))	4409
S14	TI (minimis* N5 ("blood loss" OR transfusion*)) or AB (minimis* N5 ("blood loss" OR transfusion*))	4384
S13	S1 or S2 or S3 or S4 or S5 or S6 or S7 or S8 or S9 or S10 or S11 or S12	54847
S12	TI (postoperative OR "post operative") or AB (postoperative OR "post operative")	14541
S11	TI (peroperative OR "per operative") or AB (peroperative OR "per operative")	51
S10	TI (intraoperative OR "intra operative") or AB (intraoperative OR "intra operative")	2995
S9	TI (preoperative OR "pre operative") or AB (preoperative OR "pre operative")	7264
S8	TI (perioperative OR "peri operative") or AB (perioperative OR "peri operative")	5356
S7	MH "Preoperative Period+"	725
S6	MH "Intraoperative Period"	367
S5	MH "Intraoperative Complications+"	1821

S4	MH "Postoperative Period"	1923
S3	MH "Postoperative Complications+"	21486
S2	MH "Perioperative Nursing"	8857
S1	MH "Perioperative Care+"	16222

AMI

Search conducted 19 June 2009 (1968 to present)

ID	Search Terms	Results
#7	((TI=(reduc* AND ("blood loss" OR transfusion*))) OR (TI=(minimiz* AND ("blood loss" OR transfusion*))) OR (TI=(minimis* AND ("blood loss" OR transfusion*))) OR (TI=(minimis* AND ("blood loss" OR transfusion*))) OR (TI=(minimiz* %5 ("blood loss" OR transfusion*))) OR (TI=(minimiz* %5 ("blood loss" OR transfusion*))) OR (TI=(minimis* %5 ("blood loss" OR transfusion*))) OR (TI=(minimis* %5 ("blood loss" OR transfusion*))) OR AB=(minimis* %5 ("blood loss" OR transfusion*))))	48
#6	TI=(reduc* AND ("blood loss" OR transfusion*))	7
#5	TI=(minimiz* AND ("blood loss" OR transfusion*))	0
#4	TI=(minimis* AND ("blood loss" OR transfusion*))	1
#3	TI=(reduc* %5 ("blood loss" OR transfusion*)) OR AB=(reduc* %5 ("blood loss" OR transfusion*))	43
#2	TI=(minimiz* %5 ("blood loss" OR transfusion*)) OR AB=(minimiz* %5 ("blood loss" OR transfusion*))	0
#1	TI=(minimis* %5 ("blood loss" OR transfusion*)) OR AB=(minimis* %5 ("blood loss" OR transfusion*))	4

Appendix B: Excluded studies

This appendix documents studies that met inclusion criteria determined by PICO criteria, but were later excluded. These studies, and their reasons for exclusion, are listed below.

Intervention 8 – Administration of antifibrinolytics and DDAVP

Only three studies that met inclusion criteria were subsequently excluded from the evaluation; these are noted in the technical report and shown below. The reason for exclusion was the lack of pooling of data in the systematic review. Due to the large number of identified systematic reviews which included pooled data (30 systematic reviews), it was decided that these additional studies would not add substantial additional data to the evaluation.

Erstad BL. Systemic hemostatic medications for reducing surgical blood loss. *Ann Pharmacother* 35:925–934, 2001.

Fergusson D, VanWalraven C, Coyle D, Laupacis A. Economic evaluations of technologies to minimize perioperative transfusion: A systematic review of published studies. *Transfus Med Rev* 13:106–117, 1999.

Thiagarajamurthy S, Levine A, Dunning J. Does prophylactic tranexamic acid safely reduce bleeding without increasing thrombotic complications in patients undergoing cardiac surgery? *Interact Cardiovasc Thorac Surg* 3:489-494, 2004.

Appendix C: Literature search results

Intervention 1 – Acute normovolemic haemodilution

LEVEL I EVIDENCE: SYSTEMATIC REVIEWS	
REASON FOR EXCLUSION	NUMBER OF CITATIONS EXCLUDED
Initial number of citations (EMBASE.com)	69
Initial number of citations (Cochrane Library)	25
Number of duplicates	5
Number of citations searched by title/abstract	89
Non-duplicate citations identified in CINAHL & AMI	0
Title/abstract	
Not a clinical study	34
Wrong intervention	45
Wrong comparator	0
Wrong indication	0
Wrong outcome	1
Number of citations retrieved	9
Citations retrieved from manual search	1
Full paper	
Not a clinical study	3
Wrong intervention	2
Wrong comparator	0
Wrong indication	0
Wrong outcome	0
Number of citations included	5

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS	
REASON FOR EXCLUSION	Number of citations excluded
Initial number of citations (EMBASE.com)	393
Initial number of citations (Cochrane Library)	116
Number of duplicates	25
Number of citations searched by title/abstract	484
Non-duplicate citations identified in CINAHL & AMI	0
Title/abstract	
Not a clinical study	205
Wrong intervention	234
Wrong comparator	3
Wrong indication	3
Wrong outcome	13
Not an RCT	4
Not in English	8
Number of citations retrieved	14
Citations retrieved from manual search	2

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS	
REASON FOR EXCLUSION	NUMBER OF CITATIONS EXCLUDED
Full paper	
Not a clinical study	0
Wrong intervention	2
Wrong comparator	0
Wrong indication	0
Wrong outcome	0
Number of citations included	14

Intervention 2 – Intraoperative cell salvage

Level I evidence: Systematic reviews	
REASON FOR EXCLUSION	NUMBER OF CITATIONS EXCLUDED
Initial number of citations (EMBASE.com)	129
Initial number of citations (Cochrane Library)	72
Number of duplicates	41
Number of citations searched by title/abstract	160
Non-duplicate citations identified in CINAHL & AMI	0
Title/abstract	
Not a clinical study	27
Wrong intervention	115
Wrong comparator	0
Wrong indication	8
Wrong outcome	0
Number of citations retrieved	10
Citations retrieved from manual search	1
Full paper	
Not a clinical study	3
Wrong intervention	2
Wrong comparator	0
Wrong indication	0
Wrong outcome	0
Not a SR	1
Number of citations included	5

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS	
REASON FOR EXCLUSION	Number of citations excluded
Initial number of citations (EMBASE.com)	971
Initial number of citations (Cochrane Library)	228
Number of duplicates	84
Number of citations searched by title/abstract	1115
Non-duplicate citations identified in CINAHL & AMI	0
Title/abstract	
Not a clinical study	329
Wrong intervention	741
Wrong comparator	0
Wrong indication	0
Wrong outcome	2
Not an RCT	14
Number of citations retrieved	29
Citations retrieved from manual search	4
Full paper	
Not a clinical study	1
Wrong intervention	18
Wrong comparator	1
Wrong indication	0

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS	
REASON FOR EXCLUSION	NUMBER OF CITATIONS EXCLUDED
Wrong outcome	1
Not in English	1
Not an RCT	2
Number of citations included	9

Intervention 3 – Perioperative acute normovolemic haemodilution combined with intraoperative cell salvage

LEVEL I EVIDENCE: SYSTEMATIC REVIEWS		
REASON FOR EXCLUSION	Number of citations excluded	
Initial number of citations (EMBASE.com)	129	
Initial number of citations (Cochrane Library)	72	
Number of duplicates	41	
Number of citations searched by title/abstract	160	
Non-duplicate citations identified in CINAHL & AMI	0	
Title/abstract		
Not a clinical study	27	
Wrong intervention	115	
Wrong comparator	0	
Wrong indication	8	
Wrong outcome	0	
Number of citations retrieved	10	
Citations retrieved from manual search	1	
Full paper		
Not a clinical study	3	
Wrong intervention	7	
Wrong comparator	0	
Wrong indication	0	
Wrong outcome	0	
Not a SR	1	
Number of citations included	0	

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS	
REASON FOR EXCLUSION	Number of citations excluded
Initial number of citations (EMBASE.com)	60
Initial number of citations (Cochrane Library)	228
Number of duplicates	14
Number of citations searched by title/abstract	274
Non-duplicate citations identified in CINAHL & AMI	0
Title/abstract	
Not a clinical study	45
Wrong intervention	216
Wrong comparator	2
Wrong indication	4
Wrong outcome	0
Not an RCT	3
Not in English	1
Number of citations retrieved	3
Citations retrieved from manual search	0
Full paper	
Not a clinical study	0
Wrong intervention	0

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS		
REASON FOR EXCLUSION	NUMBER OF CITATIONS EXCLUDED	
Wrong comparator	0	
Wrong indication	0	
Wrong outcome	0	
Not in English	0	
Not an RCT	0	
Number of citations included	3	

Level III and IV evidence	
REASON FOR EXCLUSION	Number of citations excluded
Initial number of citations (EMBASE.com)	85
Initial number of citations (Cochrane Library)	0
Number of duplicates	2
Number of citations searched by title/abstract	83
Non-duplicate citations identified in CINAHL & AMI	0
Title/abstract	
Not a clinical study	49
Wrong intervention	30
Wrong comparator	0
Wrong indication	0
Wrong outcome	0
Not an RCT	0
Not in English	2
Individual case study	2
Number of citations retrieved	0
Citations retrieved from manual search	0

Intervention 4 – Postoperative cell salvage

LEVEL EVIDENCE: SYSTEMATIC REVIEWS	
REASON FOR EXCLUSION	Number of citations excluded
Initial number of citations (EMBASE.com)	129
Initial number of citations (Cochrane Library)	72
Number of duplicates	41
Number of citations searched by title/abstract	160
Non-duplicate citations identified in CINAHL & AMI	0
Title/abstract	
Not a clinical study	27
Wrong intervention	115
Wrong comparator	0
Wrong indication	8
Wrong outcome	0
Number of citations retrieved	10
Citations retrieved from manual search	1
Full paper	
Not a clinical study	3
Wrong intervention	2
Wrong comparator	0
Wrong indication	0
Wrong outcome	0
Not a SR	1
Number of citations included	5

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS		
REASON FOR EXCLUSION	NUMBER OF CITATIONS EXCLUDED	
Initial number of citations (EMBASE.com)	292	
Initial number of citations (Cochrane Library)	17	
Number of duplicates	9	
Number of citations searched by title/abstract	300	
Non-duplicate citations identified in CINAHL & AMI	0	
Title/abstract		
Not a clinical study	70	
Wrong intervention	213	
Wrong comparator	1	
Wrong indication	0	
Wrong outcome	1	
Not in English	1	
Not an RCT	12	
Number of citations retrieved	2	
Citations retrieved from manual search	1	
Full paper		
Not a clinical study	0	
Wrong intervention	0	
Wrong comparator	0	

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS	
REASON FOR EXCLUSION	NUMBER OF CITATIONS EXCLUDED
Wrong indication	0
Wrong outcome	0
Not in English	0
Not an RCT	0
Number of citations included	3

Intervention 5 – Deliberate induced hypotension

LEVEL EVIDENCE: SYSTEMATIC REVIEWS	
REASON FOR EXCLUSION	NUMBER OF CITATIONS EXCLUDED
Initial number of citations (EMBASE.com)	909
Initial number of citations (Cochrane Library)	16
Number of duplicates	6
Number of citations searched by title/abstract	919
Non-duplicate citations identified in CINAHL & AMI	0
Title/abstract	
Not a clinical study	12
Wrong intervention	895
Wrong comparator	0
Wrong indication	0
Wrong outcome	0
Number of citations retrieved	12
Citations retrieved from manual search	0
Full paper	
Not a clinical study	7
Wrong intervention	1
Wrong comparator	0
Wrong outcome	1
Not in English	1
Insufficient data	1
Number of citations included	1

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS		
REASON FOR EXCLUSION	NUMBER OF CITATIONS EXCLUDED	
Initial number of citations (EMBASE.com)	257	
Initial number of citations (Cochrane Library)	597	
Number of duplicates	107	
Number of citations searched by title/abstract	747	
Non-duplicate citations identified in CINAHL & AMI	0	
Title/abstract		
Not a clinical study	58	
Wrong intervention	524	
Wrong comparator	33	
Wrong indication	0	
Wrong outcome	17	
Not in English	77	
Citations included in Level I evidence	11	
Number of citations retrieved	27	
Citations retrieved from manual search	0	
Full paper		
Not a clinical study	2	
Wrong intervention	2	
Wrong comparator	7	

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS	
REASON FOR EXCLUSION	NUMBER OF CITATIONS EXCLUDED
Wrong outcome	1
Not in English	4
Publication not available	1
Number of citations included	10

Intervention 6 – Prevention of hypothermia

LEVEL EVIDENCE: SYSTEMATIC REVIEWS		
REASON FOR EXCLUSION	Number of citations excluded	
Initial number of citations (EMBASE.com)	158	
Initial number of citations (Cochrane Library)	23	
Number of duplicates	1	
Number of citations searched by title/abstract	180	
Non-duplicate citations identified in CINAHL & AMI	0	
Title/abstract		
Not a clinical study	1	
Wrong intervention	0	
Wrong comparator	0	
Wrong indication	171	
Wrong outcome	0	
Number of citations retrieved	8	
Citations retrieved from manual search	0	
Full paper		
Not a clinical study	2	
Wrong intervention	0	
Wrong comparator	0	
Wrong indication	2	
Wrong outcome	0	
Not in English	1	
Number of citations included	3	

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS	
REASON FOR EXCLUSION	NUMBER OF CITATIONS EXCLUDED
Initial number of citations (EMBASE.com)	1002
Initial number of citations (Cochrane Library)	319
Number of duplicates	41
Citations retrieved from manual search	6
Number of citations searched by title/abstract	1286
Non-duplicate citations identified in CINAHL & AMI	0
Title/abstract	
Not a clinical study	518
Wrong intervention	607
Wrong comparator	42
Wrong indication	1
Wrong outcome	45
Not in English	26
Number of citations retrieved	47
Full paper	
Not a clinical study	1
Wrong intervention	1
Wrong comparator	9
Wrong indication	0

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS	
REASON FOR EXCLUSION	NUMBER OF CITATIONS EXCLUDED
Wrong outcome	13
Not in English	0
Citations included in Level I evidence	16
Number of citations included	5

Intervention 7 - Point-of-care testing using thromboelastography

LEVEL I-IV EVIDENCE: ALL STUDY TYPES	
REASON FOR EXCLUSION	Number of citations excluded
Initial number of citations (EMBASE.com)	1921
Initial number of citations (Cochrane Library)	307
Number of duplicates	40
Number of citations searched by title/abstract	2188
Non-duplicate citations identified in CINAHL & AMI	0
Title/abstract	
Not a clinical study	157
Wrong intervention	2001
Wrong comparator	0
Wrong indication	0
Wrong outcome	13
Number of citations retrieved	17
Citations retrieved from manual search	0
Full paper	
Not a clinical study	0
Wrong intervention	10
Wrong comparator	0
Wrong indication	0
Wrong outcome	0
Not in English	0
Number of citations included	7

Intervention 8 – Administration of antifibrinolytics & DDAVP

LEVEL EVIDENCE: SYSTEMATIC REVIEWS		
REASON FOR EXCLUSION	NUMBER OF CITATIONS EXCLUDED	
Initial number of citations (EMBASE.com)	388	
Initial number of citations (Cochrane Library)	106	
Number of duplicates	56	
Number of citations searched by title/abstract		438
Non-duplicate citations identified in CINAHL & AMI		0
Title/abstract	·	
Not a clinical study	243	
Wrong intervention/comparator	84	
Wrong indication	59	
Wrong outcome	2	
Not a SR	3	
Other ^a	1	
Not in English	2	
Number of citations retrieved		44
Citations retrieved from manual search		3
Full paper	·	
Not a clinical study	8	
Wrong intervention/comparator	2	
Wrong indication	0	
Wrong outcome	1	
Not a SR	3	
Other ^b	3	
Number of citations included		30

^a Other = review does not exist. ^b Other = data not pooled x 3.

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS (APROTININ)		
REASON FOR EXCLUSION	NUMBER OF CITATIONS EXCLUDED	
Initial number of citations (EMBASE.com)	301	
Initial number of citations (Cochrane Library)	49	
Number of duplicates	50	
Number of citations searched by title/abstract		300
Title/abstract		
Not a clinical study	201	
Wrong intervention/comparator	53	
Wrong indication	1	
Wrong outcome	6	
Not an RCT	22	
Othera	3	
Not in English	3	
Number of citations retrieved		11
Citations retrieved from manual search		0
Full paper		
Not a clinical study	0	
Wrong intervention/comparator	0	
Wrong indication	2	
Wrong outcome	0	
Not an RCT	0	
Other ^b	2	
Number of citations included		7

^a Other = abstract only x 2; < 10 patients per treatment arm.
^b Other = abstract only; < 10 patients per treatment arm.

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS (TR	ANEXAMIC ACID AND E-AMINOCAPROIC ACID)
REASON FOR EXCLUSION	NUMBER OF CITATIONS EXCLUDED
Initial number of citations (EMBASE.com)	321
Initial number of citations (Cochrane Library)	78
Number of duplicates	59
Number of citations searched by title/abstract	340
Title/abstract	
Not a clinical study	221
Wrong intervention/comparator	53
Wrong indication	18
Wrong outcome	0
Not an RCT	9
Not in English	7
Number of citations retrieved	32
Citations retrieved from manual search	
Full paper	
Not a clinical study	0
Wrong intervention/comparator	0
Wrong indication	1
Wrong outcome	0
Not an RCT	2
Othera	11
Number of citations included	18

^a Other = abstract only x 8; duplicate data x 2; < 10 patients per treatment arm.

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS (D	ESMOPRESSIN)	
REASON FOR EXCLUSION	Number of citations excluded	
Initial number of citations (EMBASE.com)	78	
Initial number of citations (Cochrane Library)	17	
Number of duplicates	3	
Number of citations searched by title/abstract		92
Title/abstract	·	
Not a clinical study	66	
Wrong intervention/comparator	8	
Wrong indication	18	
Wrong outcome	0	
Not an RCT	0	
Not in English	0	
Number of citations retrieved		0
Citations retrieved from manual search		0
Number of citations included		0

Intervention 9 – Appropriate patient positioning

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS		
REASON FOR EXCLUSION	Number of citations excluded	
Initial number of citations (EMBASE.com)	1641	
Initial number of citations (Cochrane Library)	6	
Number of duplicates	52	
Citations retrieved from manual search	0	
Number of citations searched by title/abstract		1589
Non-duplicate citations identified in CINAHL & AMI		0
Title/abstract		
Not a clinical study	95	
Wrong intervention	1413	
Wrong comparator	0	
Wrong indication	34	
Wrong outcome	43	
Number of citations retrieved		14
Full paper		
Not a clinical study	2	
Wrong intervention	0	
Wrong comparator	0	
Wrong indication	1	
Wrong outcome	3	
Not in English	2	
Number of citations included		6

Intervention 10 – Preoperative autologous donation

LEVEL EVIDENCE: SYSTEMATIC REVIEWS	
REASON FOR EXCLUSION	NUMBER OF CITATIONS EXCLUDED
Initial number of citations (EMBASE.com)	479
Initial number of citations (Cochrane Library)	146
Number of duplicates	2
Number of citations searched by title/abstract	623
Non-duplicate citations identified in CINAHL & AMI	0
Title/abstract	
Not a clinical study	167
Wrong intervention	445
Wrong comparator	0
Wrong indication	1
Wrong outcome	0
Number of citations retrieved	10
Citations retrieved from manual search	0
Full paper	
Not a clinical study	0
Wrong intervention	2
Wrong comparator	0
Wrong indication	0
Wrong outcome	0
Not a SR	0
Number of citations included	8

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS	
REASON FOR EXCLUSION	Number of citations excluded
Initial number of citations (EMBASE.com)	927
Initial number of citations (Cochrane Library)	147
Number of duplicates	56
Number of citations searched by title/abstract	1018
Non-duplicate citations identified in CINAHL & AMI	0
Title/abstract	
Not a clinical study	463
Wrong intervention	524
Wrong comparator	5
Wrong indication	0
Wrong outcome	0
Not an RCT	22
Not in English	3
Number of citations retrieved	1
Citations retrieved from manual search	1
Full paper	
Not a clinical study	0
Wrong intervention	0
Wrong comparator	0

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS					
REASON FOR EXCLUSION	NUMBER OF CITATIONS EXCLUDED				
Wrong indication	0				
Wrong outcome	0				
Not in English	0				
Not an RCT	0				
Number of citations included	2				

Perioperative Question 3 – Quality of life

LEVEL II EVIDENCE: RANDOMISED CONTROLLED TRIALS	
REASON FOR EXCLUSION	Number of citations excluded
Initial number of citations (EMBASE.com)	1173
Initial number of citations (Cochrane Library)	NA a
Number of duplicates	0
Number of citations searched by title/abstract	1173
Non-duplicate citations identified in CINAHL & AMI	0
Title/abstract	
Not a clinical study	375
Wrong intervention	798
Wrong comparator	0
Wrong indication	0
Wrong outcome	0
Number of citations retrieved	0
Citations retrieved from manual search	0

^a The clinical trials database in the Cochrane Library does not contain Level III and Level IV evidence

Appendix D: Evidence matrixes

Evidence matrixes are presented below for each intervention within Perioperative Question 3. A separate evidence statement form is shown for each primary outcome and also any of the secondary outcomes that were co-reported. Each evidence statement form is accompanied by an evidence summary table which summarises the evidence base for that particular outcome.

For each intervention, the complete set of evidence statement forms is followed by a separate form that contains any recommendations which were formulated from the evidence base for that intervention.

Intervention 1 – Acute normovolemic haemodilution

Key question(s): In patients undergoing surgery, what is the effect of ANH on transfusion incidents.	dence	e?	Evidence table ref*: POQ3.I1.P1			
1. Evidence base			1			
Level I evidence: Carless 2004 (fair quality ¹ ; 25 trials, N=1081; adults undergoing any type of surgery) and	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias			
Gurusamy 2009 (good quality, 3 trials, N=233; adults undergoing liver resection). Level II evidence published after the Carless 2004 literature search: 12 RCTs: Bennett 2006 (fair quality;	В	One or two Level II studies with a low risk of bias or SR/several Le	evel III studies with a low risk of bias			
N=155); Casati 2002 (poor quality; N=204); Casati 2004 (fair quality; N=100); Friesen 2006 (fair quality; N=32);	С	One or two Level III studies with a low risk of bias or Level I or II s				
Hohn 2002 (poor quality; N=80); Jarnagin 2008 (fair quality; N=130); Juelsgaard 2002 (fair quality; N=28); Lim 2003 (fair quality; N=30); Matot 2002 (fair quality; N=78); Sanders 2004 (fair quality; N=160); Saricaoglu 2005 (good quality?: N=30); Wolowczyk 2003 (fair quality; N=36).	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	S			
2. Consistency						
The meta-analysis conducted herein showed a significant degree of heterogeneity (P< 0.0001; I²=83%). The	Α	All studies consistent				
eterogeneity remains significant when assessed by surgery type.		Most studies consistent and inconsistency can be explained				
		Some inconsistency, reflecting genuine uncertainty around quest	ion			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact						
Meta-analysis of systematic review and 11 of the 12 RCTs (except Wolowczyk 2003); Patients transfused	А	Very large				
with allogeneic blood; see Technical Report. All surgery types – RR 0.71 (0.61, 0.84); 37 trials; N=2098	В	Substantial				
Cardiac surgery – RR 0.84 (0.70, 1.02); 14 trials; N=940	С	Moderate				
Orthopaedic surgery – RR 0.76 (0.58, 1.00); 9 trials; N=467 Miscellaneous surgery ⁴ – RR 0.57 (0.43, 0.76); 14 trials; N=691	D	Slight/Restricted				
Results from Gurusamy 2009 (liver resection): RR 0.41 (0.25, 0.66); 3 trials; N=233						
4. Generalisability						
The evidence is generalisable to an adult population who are undergoing elective surgery. The studies were	Α	Evidence directly generalisable to target population				
conducted in adults undergoing cardiac, orthopaedic, liver and other surgeries.	В	Evidence directly generalisable to target population with some car	veats			
	С	Evidence not directly generalisable to the target population but co	ould be sensibly applied			
	D	Evidence not directly generalisable to target population and hard	to judge whether it is sensible to			
5. Applicability						
Most of the studies were conducted in developed countries (UK, USA, Germany, France, Sweden, Turkey,	Α	Evidence directly applicable to Australian healthcare context				
Taiwan, Belgium, Egypt, South Africa, Israel). Boussofara 2002 (from the Carless 2004 systematic review) was conducted in Tunisia, but the exclusion of this study does not impact on the result. None of the studies were	В	Evidence applicable to Australian healthcare context with few cav	eats			
conducted in Funisia, but the exclusion of this study does not impact on the result. Notice of the studies were conducted in Australia.	С	Evidence probably applicable to Australian healthcare context wit	h some caveats			
	D	Evidence not applicable to Australian healthcare context				
	-					

Other factors

Carless 2004 did not report the quality of the included studies; however, Bryson 1998 conducted a quality assessment 16 of the 25 RCTs included in Carless 2004. All 16 RCTs would be considered fair or poor quality.

Friesen 2006 was not taken into account as the study was in infants.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
Evidence base	С	Two Level I studies and several Level II studies with moderate risk of bias
2. Consistency	С	Some inconsistency, reflecting genuine uncertainty around question
3. Clinical impact	В	ANH moderately reduces the incidence of allogeneic blood transfusion
4. Generalisability	В	Evidence directly generalisable to target population with some caveats. Surgery types assessed include cardiac, orthopaedic, liver resection, and others.
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, ANH reduces the incidence of allogeneic blood transfusion.

Abbreviations: ANH, acute normovolemic haemodilution; het, heterogeneity; RCT, randomised controlled trial; RR, relative risk.

Primary outcomes: P1 = transfusion incidence. P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

¹ No quality assessment of included studies were undertaken, the characteristics of the individual studies were appropriately summarised, the sources of heterogeneity were not sufficiently explored.

² Allocation to treatment groups was concealed from those responsible for recruiting subjects, patient characteristics and demographics were similar between treatment arms at baseline, all randomised patients were included in the analysis, the statistical methods were appropriate, and a transfusion protocol was reported. The study was not double-blinded.

³ Study assesses transfusion incidence of banked autologous blood rather than allogeneic blood.

⁴ Including cystectomy (Atallah 1993), prostatectomy (Boidt 1999), prostate resection (Malinovsky 1989), maxillofacial surgery (Bonnet 1986), Cervicofacial and ENT surgery (Boussofara 2002), liver resection (Jarnagin 2008; Matot 2002; von Bormann 1986), spinal surgery (Lim 2003), thoracic surgery (Moyes 1985), surgery for cancer (Rose 1981), gastrointestinal surgery (Sanders 2004), and any surgery type (Khanna 1998).

POQ3.I1.P1 Characteristics and results of studies examining the effect of acute normovolemic haemodilution on transfusion incidence.

	Level of evidence	No. of trials /	Patient population / Surgical				Results			Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Carless (2004)	Level I Fair	25 trials (quality NR¹) N=1081	Adults undergoing any type of surgery	All the studies conducted in countries with well developed healthcare systems (not specifically Aus/NZ).	ANH	Incidence of allogeneic blood transfusion	RR (95% CI): 0.69 (0.56, 0.84)		P<0.05	P<0.00001
Bennett (2006)	Level II Fair	N=155	Adults undergoing <u>elective hip</u> <u>surgery</u> . ² Anticipated blood loss between 1 to 1.5 L	Hospital in UK	Autologous blood was collected immediately before surgery, aiming to reduce haemoglobin concentration to a target of 110 g per L. All autologous blood was returned within 6 hours of collection, starting on wound closure or sooner if a transfusion trigger was reached.	Incidence of allogeneic blood transfusion	15/78 (19%)	22 /77(29%)	P=0.18	
Casati (2002)	Level II Poor	N=204	Adults undergoing <u>cardiac</u> <u>surgery</u> ³	Hospital in Italy	Low volume ANH: 5-8 mL/kg of blood withdrawn before systemic heparinisation and replaced with colloid solutions.	Incidence of allogeneic blood (including PRBC, FFP, and PLTC) transfusion	35/103 (34%)	36/101 (36%)	P=0.88	
						Incidence of PRBC transfusion	32/103 (31%)	34/101 (34%)	P=0.47	
Casati (2004)	Level II Fair	N=100	Adults undergoing off-CPB CABG	Hospital in Italy	ANH with tranexamic acid (with tranexamic acid as control)	Incidence of PRBC transfusion	2/50 (4%)	10/50 (20%)	P=0.028	
						Incidence of allogeneic blood transfusion (including PRBC, FFP, and PLTC)	2/50 (4%)	10/50 (20%)	P=0.028	

	Level of evidence	No. of trials /	Patient population / Surgical				Results			Notes							
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value								
Friesen (2006)	Level II Fair	N=32	Infants undergoing <u>non-</u> <u>complex open cardiac surgery</u>	Hospital in USA	ANH: 15 mL/kg whole blood withdrawn from the patient through the central venous catheter. Isovolemia was maintained by infusion of 1 mL of 5% albumin solution for each mL of blood withdrawn. Autologous blood retransfused postoperatively.	Incidence of FFP or platelet transfusion	1/16 (6.2%)	5/16 (31%)	P=0.06								
Hohn (2002)	Level II Poor	N=80	Adults undergoing on-CPB cardiac surgery	Hospital in Switzerland	ANH from a mean haematocrit of 43% to 28%.	Incidence of allogeneic blood transfusion	12/39 (31%)	12/41 (29%)	P=0.88								
3 ()	Level II Fair	N=130 Adults undergoing <u>major</u> <u>hepatic resection</u> (three or more liver segments) for any diagnosis, with or without any	Hospital in USA	ANH: blood was withdrawn to a target haemoglobin concentration of 8.0 g/dL, with a maximum of 3 L of	Patients undergoing any allogeneic transfusion	14/63 (22.2%)	23/67 (34%)	P=0.13									
			other planned procedures		blood removed. Normovolemia was maintained by replacing half of the removed blood volume with 5% albumin and	Incidence of allogeneic RBC transfusion (total)	8/63 (12.7%)	17/67 (25.4%)	P=0.08								
											the other half with crystalloid.		Incidence of allogeneic RBC transfusion (intraoperative)	1/63 (1.6%)	7/67 (10.4%)	P=0.07	
						Incidence of FFP transfusion	11/63 (17.5%)	19/67 (28.4%)	P=0.15								
Juelsgaard (2002)	Level II Fair	N=28	Adults undergoing <u>TKA</u>	Hospital in Denmark	20% of the total blood volume was drawn before anaesthesia. This volume was simultaneously replaced with an equal volume of HES 6%. Blood re-transfusion was completed within 6 h.	Incidence of PRBC transfusion	7/14 (50%)	6/14 (43%)	P=0.71								

0	Level of evidence	No. of trials /	Patient population / Surgical				Results			Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Lim (2003)	Level II Fair	N=30	Adults undergoing spinal surgery	Hospital in South Korea	ANH and esmolol-induced controlled hypotension (E-ANH group): for ANH autologous blood was withdrawn from the radial artery aiming for 28% haematocrit. To maintain normovolemia, the first 500 mL of blood drawn was simultaneously replaced with an equal amount of 6% HES, and the blood thereafter was replaced with three times that volume of Lactated Ringer's solution.	Incidence of PRBC transfusion	10/15 (67%)	15/15 (100%)	P=0.04	
Matot (2002)	Level II Fair	N=78	Adults undergoing <u>liver</u> resection	Hospital in Israel	ANH: Preoperatively, blood was transfused from the patients into standard citrate-phosphate-dextrose blood storage bags, and was simultaneously replaced by colloid solutions. The volume of blood collected was 2,020 ± 412 mL.	Incidence of PRBC transfusion	4/39 (10%)	14/39 (36%)	P=0.014	
Sanders (2004)	Level II Fair	N=160	Adults undergoing major gastrointestinal surgery (colorectal, gastric, or pancreatic) ⁴	Hospital in UK	Maximum 3 units of blood withdrawn and transfused into blood bags containing citrate-phosphate-dextrose (anticoagulant). Warmed cell-free fluid was administered during blood withdrawal to maintain normovolemia. At the end of the operation, all the autologous blood was retransfused.	Incidence of allogeneic blood transfusion	22/78 (28%)	25/82 (30%)	P=0.75	

Charles	Level of evidence	No. of trials /	Patient population / Surgical procedure	Setting	Intervention	Outcome		Notes		
Study	Quality	sample size					Intervention	Comparator	p-value	
Saricaoglu (2005)	Level II Good	N=30	Adults undergoing <u>hip</u> arthroplasty	Hospital in Turkey	ANH: autologous blood 15 mL/kg was withdrawn and replaced by ~15mL/kg 6% HES HHD: 15 mL/kg HES administered without removal of any autologous blood Control: no haemodilution	Incidence of allogeneic blood transfusion	2/10 (20%)	HDD: 4/10 (40%) Control: 10/10 (100%)	ANH vs. HHD P=0.35 ANH vs. control P=0.01	
Wolowczyk (2003)	aortic aneurysm repair g/kg of blood was withdra		Intraoperative transfusion of banked autologous blood	7/16 (44%)	7/18 (39%)	P=0.77				
					(including cell salvage)	Postoperative transfusion of banked autologous blood	5/16 (31%)	10/18 (56%)	P=0.18	
						Total transfusion of banked autologous blood	10/16 (63%)	13/18 (72%)	P=0.55	
Cardiac surgery	1	-1	,	•		1	1	1	•	•
Carless (2004)	Level I Fair	10 trials (quality NR) N=NR	Adult patients undergoing any type of surgery	All studies conducted in developed countries	ANH	Incidence of allogeneic blood transfusion	RR (95% CI): 0.77	(0.57, 1.04)	P>0.05	Phet=NR
Orthopaedic sur	gery				•					
Carless (2004)	Level I Fair	6 trials (quality NR) N=NR	Adult patients undergoing any type of surgery	All studies conducted in developed countries	ANH	Incidence of allogeneic blood transfusion	RR (95% CI): 0.79	(0.60, 1.06)	P>0.05	Phet=NR

Study	Level of evidence	No. of trials /	Patient population / Surgical procedure	Setting			Results			Notes	
	Quality	sample size			Intervention	Outcome	Intervention	Comparator	p-value		
Liver resection	Liver resection										
Gurusamy (2009)	Level I Fair	3 trials (fair quality) N=233	Patients undergoing <u>liver</u> resection ⁵	The RCTs were conducted in USA, Israel, and China.	ANH	Incidence of allogeneic blood transfusion	RR (95% CI): 0.41 (0.25, 0.66)	P<0.05	Phet=NR	
Miscellaneous											
Carless (2004)	Level I Fair	9 trials (quality NR) N=NR	Adult patients undergoing any type of surgery	All studies conducted in developed countries	ANH	Incidence of allogeneic blood transfusion	RR (95% CI): 0.42 (0.24, 0.74)	P<0.05		
Transfusion prot	ocol used	•					•			•	
Carless (2004)	Level I Fair	16 trials (quality NR) N=NR	Adult patients undergoing any type of surgery	All studies conducted in developed countries	ANH	Incidence of allogeneic blood transfusion	RR (95% CI): 0.81 (0.62, 1.00)	P=0.05		
No transfusion p	No transfusion protocol used										
Carless (2004)	Level I Fair	9 trials (quality NR) N=NR	Adult patients undergoing any type of surgery	All studies conducted in developed countries	ANH	Incidence of allogeneic blood transfusion	RR (95% CI): 0.53 (0.36, 0.76)	P<0.05		

Abbreviations: ANH, acute normovolemic haemodilution; CABG, coronary artery bypass surgery; CI, confidence interval; CPB, cardiopulmonary bypass; FFP, fresh frozen plasma; HES, hydroxyethyl starch; HHD, hypervolemic haemodilution; NR, not reported; PLTC, platelet concentration; PRBC, packed red blood cells; RCT, randomised controlled trial; TKA, total knee arthroplasty.

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¹ Bryson 1998 reported the quality of 16 of the 25 studies included in Carless 2004 that reported this outcome. Three studies had a Jadad score of 2 and the rest had a Jadad score of 1. Seven studies reported a transfusion protocol. Two studies (Triulzi 1995 and Von Bormann 1986) with a Jadad score of 2 reported the use of a transfusion protocol. Full texts of these two papers were retrieved. Triulzi 1995 was considered to be fair quality (not double-blinded, no allocation concealment reported, demographics similar between groups, all randomised patients included in analysis, statistical methods appropriate). Von Bormann 1986 was in German and therefore its quality was not assessed further. The other 14 studies in Bryson 1998 would have been rated as either fair or poor based on the Jadad scores and whether or not a transfusion protocol was reported.

² Most patients underwent primary total hip replacement, with 15 revision hip arthroplasties (seven in ANH and eight in standard transfusion) and one hip resurfacing procedure.

³ Procedures included single and multiple valve surgery, aortic root surgery, coronary surgery combined with valve surgery, or partial left ventriculectomy.

⁴ These operations were considered high risk (>40%) for allogeneic transfusion.

⁵ Trials were included irrespective of whether they included major or minor liver resections, normal or cirrhotic livers, vascular occlusion was used or not, and irrespective of the reason for liver resection.

Key question(s): In patients undergoing surgery, what is the effect of <u>ANH</u> on <u>transfusion vo</u>	lume?		Evidence table ref*: POQ3.I1.P2				
1. Evidence base		•					
Level I evidence: Carless 2004 (fair quality); 17 trials, N=NR; adults undergoing any type of surgery) and	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias					
Gurusamy 2009 (good quality: 2 trials, N=150; adults undergoing liver resection) Level II evidence: 11 trials: Aklagh 2007 (poor quality; N=60); Bennett 2006 (fair quality; N=155); Casati 2002	В	One or two Level II studies with a low risk of bias or SR/several Level II	I studies with a low risk of bias				
(poor quality; N=204); Casati 2004 (fair quality, N=100); Hohn 2002 (poor quality; N=80); Jarnagin 2008 (fair quality; N=130); Juelsgaard 2002 (fair quality; N=28); Lim 2003 (fair quality; N=30); Sanders 2004 (fair quality; N=20); Lim 2003 (fair quality; N=30); Sanders 2004 (fair quality; N=30);	С	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias					
N=160); Saricaoglu 2005 (good quality?; N=30); Wolowczyk 2003 (fair quality; N=36)	D	Level IV studies or Level I to III studies/SRs with a high risk of bias					
2. Consistency							
Carless 2004 did not report the level of heterogeneity. The results from Carless 2004 are not consistent with	Α	All studies consistent					
many of the subsequently published RCTs. Gurusamy 2009 found a significant degree of heterogeneity between trials of adults undergoing liver resection. There was a significant degree of heterogeneity in the	В	Most studies consistent and inconsistency can be explained					
meta-analysis conducted herein (P<0.0001; I ² = 79% for all surgery types).	С	Some inconsistency, reflecting genuine uncertainty around question					
	D	Evidence is inconsistent					
	NA	Not applicable (one study only)					
w Clinical impact							
A meta-analysis was conducted herein using results from the RCTs reported in Carless 20043 and the	Α	Very large					
subsequently published RCTs that reported sufficient information to be included in the analysis (Jarnagin 2008; Lim 2003; Saricaoglu 2005); see Technical Report.	В	Substantial					
General – mean difference (unit) -0.90 (-1.22, -0.57); 16 trials; N=817	С	Moderate					
Cardiac surgery – mean difference -1.00 (-1.48, -0.52); 10 trials; N=537	D	Slight/Restricted					
Orthopaedic surgery – mean difference -0.61 (-1.39, 0.18); 3 trials; N=70 Miscellaneous surgery – mean difference -1.14 (-2.57, 0.30); 3 trials; N=210							
4. Generalisability		<u> </u>					
The evidence is generalisable to an adult population who are undergoing elective surgery. The studies were	Α	Evidence directly generalisable to target population					
conducted in adults undergoing cardiac, orthopaedic, liver and other surgeries.	В	Evidence directly generalisable to target population with some caveats	i				
	С	Evidence not directly generalisable to the target population but could be	pe sensibly applied				
	D	Evidence not directly generalisable to target population and hard to juc	dge whether it is sensible to apply				
5. Applicability							
The studies were conducted in a wide range of countries (Germany, USA, Belgium, India, Turkey, South	А	Evidence directly applicable to Australian healthcare context					
Africa, Israel, Taipei, UK, China, Italy, Switzerland, Denmark, South Korea, Iran). All the studies were in a hospital setting. None of the studies were conducted in Australia.	В	Evidence applicable to Australian healthcare context with few caveats					
The same of the sa	С	Evidence probably applicable to Australian healthcare context with sor	me caveats				
	D	Evidence not applicable to Australian healthcare context					

Other factors

Carless 2004 did not report the quality of the included studies; however, Bryson 1998 conducted a quality assessment 13 of the 17 RCTs included in Carless 2004. All 13 RCTs would be considered fair or poor quality.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	Two Level I studies and several Level II studies with moderate risk of bias
2. Consistency	С	Some inconsistency, reflecting genuine uncertainty around question
3. Clinical impact	С	ANH moderately reduces the volume of allogeneic blood transfusion
4. Generalisability	В	Evidence directly generalisable to target population with some caveats. Surgery types assessed include cardiac, orthopaedic, liver resection, and others.
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, ANH may reduce the volume of allogeneic blood transfusion.

Abbreviations: ANH, acute normovolemic haemodilution; NR, not reported; RCT, randomised controlled trial

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

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^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission 1 No quality assessment of included studies were undertaken, the characteristics of the individual studies were appropriately summarised, the sources of heterogeneity were not sufficiently explored.

² Allocation to treatment groups was concealed from those responsible for recruiting subjects, patient characteristics and demographics were similar between treatment arms at baseline, all randomised patients were included in the analysis, the statistical methods were appropriate, and a transfusion protocol was reported. The study was not double-blinded.

³ Carless 2004 did not provide sufficient detail for the meta-analysis; therefore the original RCTs were sourced. Lilleaasen (1977) was not included because the study comparator was low volume ANH; Von Borman 1986 was excluded because the study was not in English; and Vedrinne 1992 was excluded due to insufficient detail.

POQ3.I1.P2 Characteristics and results of studies examining the effect of ANH on transfusion volume.

	Level of evidence	No. of trials /	Patient population / Surgical procedure				Results			
Study	Quality	sample size		Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Carless (2004)	Level I Fair	17 trials (quality NR¹) N=NR	Adults undergoing any type of surgery	All studies conducted in countries with well developed healthcare systems (not specifically Aus/NZ).	ANH	Mean difference (95% CI) in units of allogeneic blood transfused	-1.9 (-2.7, -1.1)		P<0.05	
Gurusamy (2009)	Level I Good	2 trials (fair quality) N=150	Patients undergoing liver resection ²	Studies conducted in USA, Israel, and China.	ANH	Mean difference (95% CI) in units of RBCs transfused	-0.09 (-0.48, 0.29)		P>0.05	
Akhlagh (2007)	Level II Poor	N=60	Adults undergoing on-CPB CABG	Hospital in Iran	ANH and re-transfusion of autologous blood after separating the patient from the cardiopulmonary machine.	Mean (SD) volume of allogeneic blood transfused, mL	870 (NR)	2010 (NR)	P=0.024	
Bennett (2006)	Level II Fair	N=155	Adults undergoing <u>elective hip</u> <u>surgery</u> . Anticipated blood loss between 1 to 1.5 L.	Hospital in UK	Autologous blood was collected immediately before surgery, aiming to reduce Hb concentration to a target of 110 g per L. All autologous blood was returned within 6 hours of collection, starting on wound closure or sooner if a transfusion trigger was reached.	Mean (SD) units of allogeneic blood transfused			NR	
Casati (2002)	Level II Poor	N=204	Adults undergoing <u>cardiac surgery</u> ³	Hospital in Italy	Low volume ANH: 5-8 mL/kg of blood withdrawn before systemic heparinisation and replaced with colloid solutions.	Mean (SD) units of PRBCs transfused	3.8 (NR)	3.7 (NR)	P=0.47	
Casati (2004)	Level II Fair	N=100	Adults undergoing off-CPB CABG	Hospital in Italy	ANH with tranexamic acid (with tranexamic acid as control)	Mean (SD) units of PRBCs transfused	2.5 (NR)	2.4 (NR)	P<0.001	
Hohn (2002)	Level II Poor	N=80	Adults undergoing on-CPB cardiac surgery	Hospital in Switzerland	ANH from a mean haematocrit of 43% to 28%.	Median (range) units of allogeneic blood transfused	2 (1 to 5)	2 (1 to 3)	P=0.219	
Jarnagin (2008)	Level II Fair	N=130	Adults undergoing <u>major hepatic</u> <u>resection</u> (three or more liver segments) for any diagnosis, with	Hospital in USA	ANH: blood was withdrawn to a target Hb concentration of 8.0 g/dL, with a maximum of 3 L of	Mean (SE) units of allogeneic PRBC transfused	3.5 (1.3)	2.1 (0.5)	P=0.6	

Appendix D: Evidence matrixes – Intervention 1 (Acute normovolemic haemodilution)

	Level of evidence	No. of trials /	Patient population / Surgical	Setting			Results			
Study	Quality	sample size	procedure		Intervention	Outcome	Intervention	Comparator	p-value	
			or without any other planned procedures		blood removed. Euvolemia was maintained by replacing half of the removed blood volume with 5% albumin and the other half with crystalloid.	Mean (SE) units of any allogeneic transfusion (PRBC or FFP)	5.6 (1.7)	6.9 (2.7)	P=0.72	
Juelsgaard (2002)	Level II Fair	N=28	Adults undergoing TKA	Hospital in Denmark	20% of the total blood volume was drawn before anaesthesia. This volume was simultaneously replaced with an equal volume of HES 6%. Blood re-transfusion was completed within 6 h.	Mean (SD) volume of allogeneic blood transfused, mL	386	343	P=0.85	
Lim (2003)	Level II Fair	N=30	Adults undergoing spinal surgery	Hospital in South Korea	ANH and esmolol-induced controlled hypotension (E-ANH group): for ANH autologous blood was withdrawn from the radial artery aiming for 28% haematocrit. To maintain normovolemia, the first 500 mL of blood drawn was simultaneously replaced with an equal amount of 6% HES, and the blood thereafter was replaced with three times that volume of Lactated Ringer's solution.	Mean (SE) units of PRBCs transfused	2.2 (2.3)	4.3 (1.5)	P<0.01	
Sanders (2004)	Level II Fair	N=160	Adults undergoing <u>major</u> <u>gastrointestinal surgery</u> (colorectal, gastric, or pancreatic) ⁴	Hospital in UK	ANH: Maximum 3 units of blood withdrawn preoperatively and replaced with warmed cell-free fluid to maintain normovolemia. At the end of the operation, all the autologous blood was retransfused.	Mean (SD) units of allogeneic blood transfused	4.1 (NR)	3.7 (NR)	P>0.05	
Saricaoglu (2005)	Level II Good	N=30	Adults undergoing hip arthroplasty	Hospital in Turkey	ANH: autologous blood 15 mL/kg was withdrawn and replaced by ~15mL/kg 6% HES HHD: 15 mL/kg HES administered without removal of any autologous blood Control: no haemodilution	Mean (SD) units of allogeneic PRBCs transfused	1.5 (0.7)	HHD: 1.25 (0.5) Control: 1.3 (0.5)	ANH vs. HHD P=0.33 ANH vs. Control P=0.33	

Appendix D: Evidence matrixes – Intervention 1 (Acute normovolemic haemodilution)

Level of evidence		No. of trials /	Patient population / Surgical				Results			
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Wolowczyk (2003)	Level II Fair	N=36	Adults undergoing <u>abdominal aortic</u> aneurysm repair	Hospital in UK	of blood was withdrawn and replaced with a similar volume	Median (IQR) units of allogeneic blood transfused intraoperatively	0 (0 to 4)	0 (0 to 2)	P=0.51	
				(including cell salvage) ui	(including cell salvage) unit		Median (IQR) units of allogeneic blood transfused postoperatively	0 (0 to 2)	1 (0 to 2)	P=0.33
				Median (IQR) units of allogeneic blood transfused intra- and postoperatively	2 (0 to 5)	2.5 (0 to 5)	P=0.68			
Carless (2004)	Level I Fair	NR	Adults undergoing <u>any type of</u> <u>surgery</u>	All studies conducted in countries with well developed healthcare systems (not specifically Aus/NZ).	ANH	Mean difference (95% CI) in units of allogeneic blood transfused.	-1.0 (-1.7, -0.4)		P<0.05	
Carless (2004)	Level I Fair	NR	Adults undergoing <u>any type of</u> <u>surgery</u>	All studies conducted in countries with well developed healthcare systems (not specifically Aus/NZ).	ANH	Mean difference (95% CI) in units of allogeneic blood transfused.	-3.0 (-4.9, -1.1)		P<0.05	

Abbreviations: ANH, acute normovolemic haemodilution; CABG, coronary artery bypass graft; CI, confidence interval; CPB, cardiopulmonary bypass; FFP, fresh frozen plasma; Hb, haemoglobin; HES, hydroxyethyl starch; het, heterogeneity; HHD, hypervolemic haemodilution; IQR, interquartile range; NR, not reported; PRBC, packed red blood cells; RCT, randomised controlled trial; SD, standard deviation; TKA, total knee arthroplasty.

¹Bryson 1998 reported the quality of 13 of the 17 studies included in Carless 2004 that reported this outcome. Five studies had a Jadad score of 2 and the rest had a Jadad score of 1. Six studies reported the use of a transfusion protocol. Three studies (Kochamba 1996, Triulzi 1995, and Von Borman 1986) with a Jadad score of 2 reported the use of a transfusion protocol. Full texts of these three papers were retrieved. Kochamba 1996 and Triulzi 1995 were considered to be fair quality (not double-blinded, no allocation concealment reported, demographics similar between groups, all randomised patients included in analysis, statistical methods appropriate). Von Bormann 1986 was in German and therefore its quality was not assessed further. The other 10 studies in Bryson 1998 would have been rated as either fair or poor based on the Jadad scores and whether or not a transfusion protocol was reported.

²Trials were included irrespective of whether they included major or minor liver resections, normal or cirrhotic livers, vascular occlusion was used or not, and irrespective of the reason for liver resection.

³ Most patients underwent primary total hip replacement, with 15 revision hip arthroplasties (seven in ANH and eight in standard transfusion) and one hip resurfacing procedure.

⁴ These operations were considered high risk (>40%) for allogeneic transfusion.

Key question(s): In patients undergoing surgery, what is the effect of <u>ANH</u> on <u>blood loss?</u>			Evidence table ref*: POQ3.I1.P3		
1. Evidence base			1		
Level I evidence: Bryson 1998 (good quality; 13 trials, all fair to poor quality, N=500; any surgery type) and Gurusamy 2009 (good quality; 2 trials; N=98; adults undergoing liver resection)	Α	One or more level I studies with a low risk of bias or several level II st	udies with a low risk of bias		
Level II evidence: 11 RCTs: Bennett 2006 (fair quality; N=155); Casati 2002 (poor quality; N=204); Casati	В	One or two Level II studies with a low risk of bias or SR/several Level	III studies with a low risk of bias		
2004 (fair quality; N=100); Friesen 2006 (fair quality; N=32); Jarnagin 2008 (fair quality; N=130); Juelsgaard	С	One or two Level III studies with a low risk of bias or Level I or II studi	es with a moderate risk of bias		
2002 (fair quality; N=28); Lim 2003 (fair quality; N=30); Matot 2002 (fair quality; N=78); Sanders 2004 (fair quality; N=160); Saricaoglu 2005 (good quality ¹ ; N=30); Wolowczyk 2003 (fair quality; N=36)	D	Level IV studies or Level I to III studies/SRs with a high risk of bias			
2. Consistency					
Within Bryson 1998, five studies found that ANH was associated with a significant decrease in blood loss and	Α	All studies consistent			
eight studies found no significant difference. The degree of heterogeneity in the Bryson 1998 studies is statistically significant (P<0.001). The meta-analysis in Bryson 1998 found a small, but statistically significant	В	Most studies consistent and inconsistency can be explained			
impact in the trials with patients undergoing cardiac surgery but not trials of patients undergoing orthopaedic	С	Some inconsistency, reflecting genuine uncertainty around question			
surgery or other surgery types. Of the RCTs published after Bryson 1998, two found a significant association	D	Evidence is inconsistent			
between ANH and decreased blood loss and the other nine studies found no statistically significant	NA	Not applicable (one study only)			
3. Clinical impact					
Results from Bryson 1998	Α	Very large			
All surgery types – mean difference -117 mL (-292, 58); 13 trials; N=500 Cardiac surgery – mean difference -233 mL (-459, -5); 7 trials; N=350 (of the two RCTs published	В	Substantial			
subsequently, only one reported lower blood loss in the intervention arm, and neither reported a significant difference)	С	Moderate			
Orthopaedic surgery – mean difference 33 mL (-512, 578); 1 trial; N=31	D	Slight/Restricted			
Miscellaneous surgery – mean difference -97 mL (-339, 145); 5 trials; N=119 Results from Gurusamy 2009 (liver resection) – mean difference 1.53 (-102, 105); 2 trials; N=98					
Results from Level II studies – see evidence summary table POQ3.I1.P3					
4. Generalisability The evidence is generalisable to an adult population who are undergoing elective surgery. The studies were		Terr non north control			
conducted in adults undergoing cardiac, orthopaedic, liver and other surgeries.	A	Evidence directly generalisable to target population			
ornation in data and going statuto, ornopastary, mor and ornor starger to	B C	Evidence directly generalisable to target population with some caveat Evidence not directly generalisable to the target population but could			
	C	70 011			
	D	Evidence not directly generalisable to target population and hard to ju	udge whether it is sensible to apply		
5. Applicability	,	T			
The studies in Bryson 1998 were conducted in Europe with the exception of two studies conducted in USA and one in South Africa. The studies in Gurusamy 2009 were conducted in Israel and China. The RCTs not	Α	Evidence directly applicable to Australian healthcare context			
included in either Bryson 1998 or Gurusamy 2009 were conducted in UK, Italy, USA, Denmark, South Korea,	В	Evidence applicable to Australian healthcare context with few caveats			
and Turkey.	С	Evidence probably applicable to Australian healthcare context with some caveats			
	D	Evidence not applicable to Australian healthcare context			

Although Bryson 1998 was a good quality systematic review, all the included RCTs were fair to poor quality.

Friesen 2006 was not taken into account as the study was in infants.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	Two good quality Level I studies and several Level II studies with moderate risk of bias
2. Consistency	С	Some inconsistency, reflecting genuine uncertainty around question
3. Clinical impact	D	No statistically significant impact
4. Generalisability	В	Evidence directly generalisable to target population with some caveats
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of ANH on blood loss is uncertain.

Abbreviations: ANH, acute normovolemic haemodilution; RCT, randomised controlled trial

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission ¹ Allocation to treatment groups was concealed from those responsible for recruiting subjects, patient characteristics and demographics were similar between treatment arms at baseline, all randomised patients were included in the analysis, the statistical methods were appropriate, and a transfusion protocol was reported. The study was not double-blinded.

^{* &}lt;u>Interventions</u>: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I1.P3 Characteristics and results of studies examining the effect of acute normovolemic haemodilution on blood loss.

	Level of evidence	No. of trials /	Patient population / Surgical					Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Bryson (1998)	Level I Good	13 trial (fair and poor quality ¹) N=500	Adults undergoing <u>any surgery</u> type	All studies conducted in developed countries	ANH	Mean difference (95% CI) in perioperative blood loss, mL	-117 (-292, 58)		P>0.05	Phet<0.001
Bennett (2006)	Level II Fair	N=155	Adults undergoing <u>elective hip</u> <u>surgery</u> .1 Anticipated blood loss between 1 to 1.5 L.	Hospital in UK	Autologous blood collected immediately before surgery, aiming to reduce Hb	Median (IQR) intraoperative blood loss, mL	692 (452, 1019)	641 (477, 1007)	P=0.82	
	concentration to a target of 110 g per L. All autologous blood returned within 6 hours of collection, starting on wound closure or sooner if a transfusion trigger was reached. N=204 Adults undergoing cardiac Hospital in Italy Low yolume ANH: 5-8 mL/kg	Median (IQR) total blood loss, mL	1182 (840, 1646)	1210 (816, 1545)	P=0.82					
Casati (2002)	Level II Poor	N=204	Adults undergoing <u>cardiac</u> <u>surgery</u> ²	Hospital in Italy	Low volume ANH: 5-8 mL/kg of blood withdrawn before systemic heparinisation and replaced with colloid solutions.	Median (IQR) bleeding 0-4 hours after surgery, mL	158 (106, 305)	172 (117.5, 265)	P=0.93	
						Mean (IQR) total postoperative bleeding (mL)	374 (255, 704)	412 (313, 552)	P=0.94	
Casati (2004)	Level II Fair	N=100	Adults undergoing off-CPB CABG	Hospital in Italy	ANH with tranexamic acid (with tranexamic acid as control)	Median (IQR) bleeding 0-4 hours after surgery, mL	160 (110, 235)	150 (100, 220)	NS	
						Mean (IQR) total postoperative bleeding (mL)	375 (248, 475)	350 (300, 443)	NS	
Friesen (2006)	Level II Fair		Infants undergoing non- complex open cardiac surgery	Hospital in USA	ANH: 15 mL/kg whole blood withdrawn through the central venous catheter. Normovolaemia was maintained by infusion of 1 mL of 5% albumin solution for each mL of blood withdrawn. Autologous blood re-transfused postoperatively.	Postoperative 24 hour blood loss	NR	NR	when measure (P=0.036), but	e treatment group ed as mL per 24 h not significantly asured as mL/kg.24h

Study	Level of evidence	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome		Results		Notes
Study	Quality	sample size	procedure	Setting	intervention	Outcome	Intervention	Comparator	p-value	
Jarnagin (2008)	Level II Fair	N=130	Adults undergoing major hepatic resection (three or more liver segments) for any diagnosis, with or without any other planned procedures.	Hospital in USA	ANH: blood withdrawn to a target Hb concentration of 8.0 g/dL, with a maximum of 3 L of blood removed. Normovolaemia was maintained by replacing half of the removed blood volume with 5% albumin and the other half with crystalloid.	Mean (range) blood loss, mL	800 (100 to 3200)	700 (100 to 4000)	P=0.42	
Juelsgaard (2002)	Level II Fair	N=28	Adults undergoing <u>TKA</u>	Hospital in Denmark	20% of the total blood volume was drawn before anaesthesia and	Mean (SD) intraoperative blood loss, mL	131 (78)	111 (56)	P=0.45	
Lim (2003)					simultaneously replaced with an equal volume of HES 6%. Blood re- transfusion was completed within 6 h.	Mean (SD) total blood loss, mL	1306 (300)	1026 (294)	P=0.02	
(/	Level II Fair	N=30	Adults undergoing <u>spinal</u> <u>surgery</u>	Hospital in South Korea	ANH and esmolol-induced controlled hypotension (E-ANH group): for ANH autologous blood was withdrawn from the radial	Mean (SD) volume of intraoperative bleeding, mL	1600 (620)	1500 (697)	P>0.05	
					artery aiming for 28% haematocrit. To maintain normovolaemia, the first 500 mL of blood drawn was simultaneously replaced with an equal amount of 6% HES, and the blood thereafter was replaced with three times that volume of Lactated Ringer's solution.	Mean (SD) volume of postoperative bleeding, mL	600 (372)	883 (473)	P>0.05	
Matot (2002)	Level II Fair	N=78	Adults undergoing <u>liver</u> resection	Hospital in Israel	ANH: Preoperatively, blood was transfused from the patients into standard citrate-phosphate-dextrose blood storage bags, and was simultaneously replaced by colloid solutions. The volume of blood collected was 2,020 ± 412 mL.	Mean (SD) surgical blood loss (mL)	1442 (1827)	1528 (1822)	P=0.84	

Ctudu	Level of evidence	No. of trials /	Patient population / Surgical	Catting	Intervention	Outcome		Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Sanders (2004)	Level II Fair	N=160	Adults undergoing <u>major</u> gastrointestinal surgery (colorectal, gastric, or pancreatic) ³	Hospital in UK	ANH: Maximum 3 units of blood withdrawn preoperatively and replaced with warmed cell-free fluid to maintain normovolaemia. At the end of the operation, all autologous blood was re- transfused.	Median (range) blood loss, mL	750-1000 (100- 4500)	750-1000 (100- 4368)	NR	
Saricaoglu (2005)	Level II Good	N=30	Adults undergoing hip arthroplasty	Hospital in Turkey	ANH: autologous blood 15 mL/kg was withdrawn and replaced by ~15mL/kg 6% HES HHD: 15 mL/kg HES administered without removal of any autologous blood Control: no haemodilution	Median (95% CI) intraoperative blood loss	740 (600, 830)	HHD: 650 (500, 855) Control: 695 (510, 855)	P=0.275	
Wolowczyk (2003)	Level II Fair	N=36	Adults undergoing <u>abdominal</u> <u>aortic aneurysm repair</u>	Hospital in UK	ANH and cell salvage: 15 g/kg of blood was withdrawn and replaced with a similar	Median (IQR) intraoperative blood loss, mL	1780 (930, 5000)	1700 (750, 2600)	P=0.55	
					volume of 6% HES <u>Control</u> : standard care (including cell salvage)	Patients with blood loss below 1000 mL	4/16 (25%)	5/18 (28%)	P=1.0	
Cardiac surgery										
Bryson (1998)	Level I Good	7 trials (fair and poor quality ⁴) N=350	Adults undergoing <u>any surgery</u> type	All studies conducted in developed countries	ANH	Mean difference (95% CI) in perioperative blood loss, mL	-233 (-459, -5)		P<0.05	Phet<0.001
Orthopaedic surg	gery									
Bryson (1998)	Level I Good	1 trial (fair/poor quality ⁵) N=31	Adults undergoing <u>any surgery</u> type	All studies conducted in developed countries	ANH	Mean difference (95% CI) in perioperative blood loss, mL	33 (-512, 578)		P>0.05	Phet=NA

	Level of evidence	No. of trials /	Patient population / Surgical					Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Liver surgery	Liver surgery									
Gurusamy (2009)	Level I Good	2 trials (fair quality) N=98	Patients undergoing <u>liver</u> resection ⁶	Studies conducted in USA, Israel, and China.	ANH	Mean difference (95% CI) perioperative blood loss, mL	1.53 (-102, 105)		P>0.05	Phet=0.83
Miscellaneous s	urgery ⁷									
Bryson (1998)	Level I Good	5 trials (fair and poor quality ⁸) N=119	Adults undergoing <u>any surgery</u> type	All studies conducted in developed countries	ANH	Mean difference (95% CI) in perioperative blood loss, mL	-97 (-339, 145)		P>0.05	Phet=0.013

Abbreviations: ANH, acute normovolemic haemodilution; CABG; coronary artery bypass graft; CI, confidence interval; CPB, cardiopulmonary bypass; Hb, haemoglobin; HES, hydroxyethyl starch; het, heterogeneity; HHD, hypervolemic haemodilution; IQR, interquartile range; NR, not reported; NS, not significant; SD, standard deviation; TKA, total knee arthroplasty.

¹ Three studies had a Jadad score of 2 and the other studies had a Jadad score of 1. Five studies reported the use of a transfusion protocol. Two studies (Kochamba 1996 and Triulzi 1995) with a Jadad score of 2 reported the use of a transfusion protocol. Full texts of these two papers were retrived. Kochamba 1996 and Triulzi 1995 were considered to be fair quality (not double-blinded, no allocation concealment reported, demographics similar between groups, all randomised patients included in analysis, statistical methods appropriate). The other 11 studies in Bryson 1998 would have been rated as either fair or poor based on the Jadad scores and whether or not a transfusion protocol was reported.

² Procedures included single and multiple valve surgery, aortic root surgery, coronary surgery combined with valve surgery, or partial left ventriculectomy.

³ These operations were considered high risk (>40%) for allogeneic transfusion.

⁴ Two studies had a Jadad score of 2 and the other studies had a Jadad score of 1. Three studies reported the use of a transfusion protocol. Two studies (Kochamba 1996 and Triulzi 1995) with a Jadad score of 2 reported the use of a transfusion protocol. Full texts of these two papers were retrived. Kochamba 1996 and Triulzi 1995 were considered to be fair quality (not double-blinded, no allocation concealment reported, demographics similar between groups, all randomised patients included in analysis, statistical methods appropriate). The other 5 studies in Bryson 1998 in this subgroup would have been rated as either fair or poor based on the Jadad scores and whether or not a transfusion protocol was reported.

⁵ The study had a Jadad score of 1 and reported the use of a transfusion protocol.

⁶ Trials were included irrespective of whether they included major or minor liver resections, normal or cirrhotic livers, vascular occlusion was used or not, and irrespective of the reason for liver resection.

⁷ Including GI surgery (1 trial), thoracic surgery (1 trial), hepatic surgery (1 trial), ENT surgery (2 trials), urological surgery (2 trials), and yascular surgery (2 trials).

One study had a Jadad score of 2 and the other studies had a Jadad score of 1. One study reported the use of a transfusion protocol. Neither of the studies with a Jadad score of 2 reported the use of a transfusion protocol. Therefore, all the studies in this subgroup would have been rated as either fair or poor based on the Jadad scores and whether or not a transfusion protocol was reported.

Key question(s):			Evidence table ref*:				
In patients undergoing surgery, what is the effect of ANH on mortality?			POQ3.I1.P4				
1. Evidence base							
Level I evidence: Carless 2004 (fair quality ¹ ; 8 trials, N=NR; adults undergoing any type of surgery) and Gurusamy 2009 (good quality; 2 trials, N=150; adults undergoing liver resection)	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias				
Level II evidence: 6 RCTs: Bennett 2006 (fair quality; N=155); Casati 2002 (poor quality; N=204); Casati 2004	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias					
(fair quality; N=100); Hohn 2002 (poor quality; N=80); Matot 2002 (fair quality; N=78); Sanders 2004 (fair quality; N=160)	С	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias					
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	3				
2. Consistency							
All the studies are consistent in finding no significant impact. However, studies are likely underpowered. Carless 2004 did not report heterogeneity.	Α	All studies consistent					
Cariess 2004 did not report neterogeneity.	В	Most studies consistent and inconsistency can be explained					
	С	Some inconsistency, reflecting genuine uncertainty around questi	on				
	D	Evidence is inconsistent					
	NA	Not applicable					
3. Clinical impact							
Results from Carless 2004 – RR 1.16 (0.19, 7.15); 8 trials; N=NR Results from Gurusamy 2009 – RR 0.35 (0.04, 3.32); 2 trials; N=150	Α	Very large					
Results from Level II studies – see evidence summary table POQ3.11.P4	В	Substantial					
	С	Moderate					
	D	No difference/underpowered					
4. Generalisability							
The evidence is generalisable to an adult population who are undergoing elective surgery. The studies were conducted in adults undergoing cardiac, orthopaedic, liver and other surgeries.	Α	Evidence directly generalisable to target population					
conducted in addits undergoing cardiac, ormopaedic, liver and other surgenes.	В	Evidence directly generalisable to target population with some case	/eats				
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied				
	D	Evidence not directly generalisable to target population and hard	to judge whether it is sensible to				
5. Applicability							
All studies were conducted in developed countries.	Α	Evidence directly applicable to Australian healthcare context					
	В	Evidence applicable to Australian healthcare context with few cave	eats				
	С	Evidence probably applicable to Australian healthcare context with	h some caveats				
	D	Evidence not applicable to Australian healthcare context					

Included studies were underpowered to detect a mortality difference.

Quality of RCTs not reported in Carless 2004.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	Two Level I studies and several Level II studies with moderate risk of bias
2. Consistency	Α	All studies consistent in finding no difference due to being underpowered
3. Clinical impact	D	No difference/underpowered
4. Generalisability	В	Evidence directly generalisable to target population with some caveats
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of ANH on mortality is uncertain.

Abbreviations; acute normovolemic haemodilution; NR, not reported; RCT, randomised controlled trial; RR, relative risk.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission 1 No quality assessment of included studies were undertaken, the characteristics of the individual studies were appropriately summarised, the sources of heterogeneity were not sufficiently explored.

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I1.P4 Characteristics and results of studies examining the effect of acute normovolemic haemodilution on mortality.

	Level of evidence	No. of trials /	Patient population / Surgical					Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Carless (2004)	Level I Fair	8 trials (quality NR) N=NR	Adults undergoing any type of surgery	All studies conducted in countries with well developed healthcare systems (not specifically Aus/NZ).	ANH		RR (95% CI): 1.16	(0.19, 7.15)	P>0.05	Phet=NR
Gurusamy (2009)	Level I Good	2 trials (fair quality) N=150	Patients undergoing <u>liver</u> resection. ¹	Studies conducted in USA, Israel, and China.	ANH		RR (95% CI): 0.35	(0.04, 3.32)	P>0.05	Phet=1.00
Bennett (2006)	Level II Fair	N=155	Adults undergoing <u>elective hip</u> <u>surgery</u> . ² Anticipated blood loss between 1 to 1.5 L.	Hospital in UK	Autologous blood was collected immediately before surgery, aiming to reduce Hb concentration to a target of 110 g per L. All autologous blood was returned within 6 hours of collection, starting on wound closure or sooner if a transfusion trigger was reached.		1/78 (1.3%)	0/77 (0%)	P=0.50	
Casati (2002)	Level II Poor	N=204	Adults undergoing <u>cardiac</u> <u>surgery</u> ¹	Hospital in Italy	Low volume ANH: 5-8 mL/kg of blood withdrawn before systemic heparinisation and replaced with colloid solutions.		4/103 (3.9%)	4/101 (4%)	P=0.98	
Casati (2004)	Level II Fair	N=100	Adults undergoing off-CPB CABG	Hospital in Italy	ANH with tranexamic acid (with tranexamic acid as control)		0/50 (0%)	1/50 (2%)	P=0.50	
Hohn (2002)	Level II Poor	N=80	Adults undergoing on-CPB cardiac surgery	Hospital in Switzerland	ANH from a mean haematocrit of 43% to 28%.		0/39 (0%)	2/41 (5%)	P=0.31	

Appendix D: Evidence matrixes – Intervention 1 (Acute normovolemic haemodilution)

	Level of evidence	No. of trials /	Patient population / Surgical procedure					Results		Notes
Study	Quality	sample size		Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Matot (2002)	Level II Fair	N=78	Adults undergoing liver resection	Hospital in Israel	ANH: Preoperatively, blood was transfused into standard citrate-phosphate-dextrose blood storage bags, and was simultaneously replaced by colloid solutions. The volume of blood collected was 2,020 ± 412 mL.		0/39 (0%)	0/39 (0%)	Not estimable	
Sanders (2004)	Level II Fair	N=160	Adults undergoing <u>major</u> gastrointestinal surgery (colorectal, gastric, or pancreatic) ²	Hospital in UK	ANH: Maximum 3 units of blood withdrawn preoperatively and replaced with warmed cell-free fluid to maintain normovolemia. At the end of the operation, all autologous blood was retransfused.		2/78 (3%)	1/82 (1%)	P=0.54	

Abbreviations: ANH, acute normovolemic haemodilution; CABG, coronary artery bypass graft; CI, confidence interval; CPB, cardiopulmonary bypass; Hb, haemoglobin; het, heterogeneity; NR, not reported; RCT, randomised controlled trial; RR, relative risk.

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¹ Trials were included irrespective of whether they included major or minor liver resections, normal or cirrhotic livers, vascular occlusion was used or not, and irrespective of the reason for liver resection.

² Procedures included single and multiple valve surgery, aortic root surgery, coronary surgery combined with valve surgery, or partial left ventriculectomy.

³ These operations were considered high risk (>40%) for allogeneic transfusion.

Key question(s): In patients undergoing surgery, what is the effect of ANH on morbidity?			Evidence table ref*: POQ3.I1.P5				
Evidence base							
Level I evidence: Carless 2004 (fair quality)	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias				
Infection (2 trials; N=NR); thrombosis (3 trials; N=NR); non-fatal MI (3 trials; N=NR) Gurusamy 2009 (good quality)	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias				
Bile leak (1 trial; N=78); intra-abdominal bleeding (2 trials; N=208); intra-abdominal infection (1 trial; N=78); intra-							
abdominal collection req. drainage (1 trial; N=130); wound infection (2 trials; N=208); chest infection (1 trial; N=78)	С	One or two Level III studies with a low risk of bias or Level I or II st					
Level II evidence: 7 RCTs: Bennett 2006 (N=155; fair quality); Casati 2002 (N=204; poor quality); Casati 2004 (N=100; fair quality); Jarnagin 2008 (N=130; fair quality); Lim 2003 (N=30; fair quality); Matot 2002 (N=78; fair quality); Sanders 2004 (N=160; fair quality).	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	5				
2. Consistency							
Most of the studies the studies are consistent in finding no significant impact of ANH on morbidity outcomes. In	Α	All studies consistent					
Bennet 2006, significantly fewer ANH patients had at least one significant postoperative complication (Bennet 2006 was the only study to report overall incidence as an outcome). Carless 2004 found that the incidence of	В	Most studies consistent and inconsistency can be explained					
thrombosis was significantly less in ANH patients. Carless 2004 did not report the degree of heterogeneity for the	С	Some inconsistency, reflecting genuine uncertainty around questi	on				
trials that reported thrombosis as an outcome. None of the RCTs published after Careless 2004 reported	D	Evidence is inconsistent					
thrombosis as an outcome.	NA	Not applicable (one study only)					
3. Clinical impact							
Results from Carless 2004	Α	Very large					
Infection – RR 4.94 (0.61, 40.19) Thrombosis – RR 0.44 (0.21, 0.93)	В	Substantial					
Non-fatal MI – RR 3.43 (0.15, 79.74)	С	Moderate					
Results from Gurusamy 2009 and Level II studies – see evidence summary table POQ3.11.P5	D	Slight/Restricted					
4. Generalisability							
The evidence is generalisable to an adult population who are undergoing elective surgery. The studies were	Α	Evidence directly generalisable to target population					
conducted in adults undergoing cardiac, orthopaedic, liver and other surgeries.	В	Evidence directly generalisable to target population with some cav	veats				
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied				
	D	Evidence not directly generalisable to target population and hard t	to judge whether it is sensible to				
5. Applicability							
All studies were conducted in developed countries.	Α	Evidence directly applicable to Australian healthcare context					
	В	Evidence applicable to Australian healthcare context with few cave	eats				
	С	Evidence probably applicable to Australian healthcare context with	h some caveats				
	D	Evidence not applicable to Australian healthcare context					

Thrombosis in Carless 2004 review was the only significant outcome but was not defined. As thrombosis was not adequately defined, the CRG did believe it was appropriate to make an evidence statement for this outcome.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

	<u> </u>	
Component	Rating	Description
1. Evidence base	С	Two Level I studies and several Level II studies with low risk of bias
2. Consistency	С	Most studies consistent in finding no significant impact on morbidity
3. Clinical impact		A statistically significant impact on thrombosis with a confidence interval that includes clinically insignificant values. No statistically significant impact on other morbidity outcomes.
4. Generalisability	В	Evidence directly generalisable to target population with some caveats
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of ANH on morbidity is uncertain.

Abbreviations: ANH, acute normovolemic haemodilution: NR, not reported; RCT, randomised controlled trial; RR, relative risk.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission 1 No quality assessment of included studies were undertaken, the characteristics of the individual studies were appropriately summarised, the sources of heterogeneity were not sufficiently explored.

POQ3.I1.P5 Characteristics and results of studies examining the effect of acute normovolemic haemodilution on morbidity.

	Level of	evidence No. of trials / Patient population / Surgical Sotting Intervention				Results		Notes		
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Carless (2004)	Level I Fair	2 trials (quality NR) N=NR	Adults undergoing any type of surgery	All studies conducted in countries with well	ANH	Infection	RR (95% CI): 4.94	(0.61, 40.19)	P>0.05	Phet=NR
		3 trials (quality NR) N=NR		developed healthcare systems (not specifically Aus/NZ).		Thrombosis	RR (95% CI): 0.44 ((0.21, 0.93)	P<0.05	Phet=NR
		3 trials (quality NR) N=NR				Non-fatal MI	RR (95% CI): 3.43 (0.15, 79.74)		P>0.05	Phet=NR
Gurusamy (2009)	Level I Good	1 trial (fair quality) N=78	resection.1	Studies conducted in USA, Israel, and China.	in ANH	Bile leak	RR (95% CI): 1.5 (0	0.27, 8.49)	P>0.05	Phet=NA
		2 trials (fair quality) N=208				Intra-abdominal bleeding	RR (95% CI): 1.87 ((0.4, 8.67)	P>0.05	Phet=0.39
		1 trial (fair quality) N=78				Intra-abdominal infection	RR (95% CI): 0.33 ((0.04, 3.07)	P>0.05	P=NA
		1 trial (fair quality) N=130				Intra-abdominal collection requiring drainage	RR (95% CI): 1.26	(0.061, 2.60)	P>0.05	Phet=NA
		2 trials (fair quality) N=208				Wound infection	RR (95% CI): 0.84	(0.34, 2.03)	P>0.05	Phet=0.18
		1 trial (fair quality) N=78				Chest infection	RR (95% CI): 1.50 ((0.27, 8.49)	P>0.05	Phet=NA

	Level of evidence	No. of trials /	Patient population / Surgical procedure	Setting		Outcome	Results			Notes
Study	Quality	sample size			Intervention		Intervention	Comparator	p-value	
, ,	Level II Fair	N=155	Adults undergoing <u>elective hip</u> <u>surgery</u> . ² Anticipated blood loss between 1 to 1.5 L.	Hospital in UK	Autologous blood was collected immediately before surgery, aiming to reduce Hb concentration to a target of 110 g per L. All	Patients with at least one significant postoperative complication.	14/78 (18%)	30/77 (38%)	P=0.006	
					autologous blood was returned within 6 hours of collection, starting on wound closure or sooner if a	Cardiovascular event	1/78 (1%)	4/77 (5%)	P=0.21	
					transfusion trigger was reached.	Postoperative infection	7/78 (9%)	17/77 (22%)	P=0.03	
						Wound (non- infective)	2/78 (3%)	0/77 (0%)	P=0.30	
						Bleeding	0/78 (0%)	1/77 (1%)	P=0.49	
						Venous thromboembolis m	2/78 (3%)	1/77 (1%)	P=0.58	
						Urinary retention	3/78 (4%)	3/77 (4%)	P=0.99	
						Transfusion reaction	0/78 (0%)	1/77 (1%)	P=0.49	
Casati (2002)	Level II Poor	N=204	Adults undergoing <u>cardiac</u> <u>surgery</u> ³	Hospital in Italy	Low volume ANH: 5-8 mL/kg of blood withdrawn before systemic heparinisation and	MI	2/103 (2%)	1/101 (1%)	P=0.58	
					replaced with colloid solutions.	Renal failure	3/103 (2.9%)	4/101 (4%)	P=0.68	
						Minor neurological complications	7/103 (6.9%)	8/101 (8%)	P=0.86	
						Stroke	2/103 (2%)	1/101 (1%)	P=0.58	
						Pulmonary embolism	0/103 (0%)	1/101 (1%)	P=0.49	

Charder	Level of evidence	No. of trials /	Patient population / Surgical	Setting	lado montino	Outcome		Results	_	Notes	
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value		
Casati (2004)	Level II Fair	N=100	Adults undergoing <u>off-CPB</u> <u>CABG</u>	Hospital in Italy	ANH with tranexamic acid (with tranexamic acid as control)	Respiratory failure	1/50 (2%)	1/50 (2%)	P=1.00		
						Atrial fibrillation	5/50 (10%)	6/50 (12%)	P=0.75		
						Major ventricular arrhythmia	1/50 (2%)	1/50 (2%)	P=1.00		
				Myocardial infarction	1/50 (2%)	1/50 (2%)	P=1.00				
			Creatinine double the baseline	1/50 (2%)	2/50 (4%)	P=0.57					
					Minor neurological complications	2/50 (4%)	1/50 (2%)	P=0.57			
Jarnagin (2008)	Level II Fair	N=130	Adults undergoing <u>major</u> <u>hepatic resection</u> (three or more liver segments) for any diagnosis, with or without any other planned procedures	Hospital in USA	ANH: blood was withdrawn to a target Hb concentration of 8.0 g/dL, with a maximum	Overall morbidity	28/63 (44%)	22/67 (33%)	P=0.17		
					of 3 L of blood removed. Normovolemia was maintained by replacing half of the removed blood volume with 5% albumin and the other half with crystalloid.	Grade ≥ 3 morbidity	19/63 (30%)	19/67 (28%)	P=0.82		
Lim (2003)	Level II Fair	N=30	Adults undergoing <u>spinal</u> <u>surgery</u>	Hospital in South Korea	ANH and esmolol-induced controlled hypotension (E-ANH group): for ANH autologous blood was withdrawn from the radial artery aiming for 28% haematocrit. To maintain normovolemia, the first 500 mL of blood drawn was simultaneously replaced with an equal amount of 6% HES, and the blood thereafter was replaced with three times that volume of Lactated Ringer's solution.	General morbidity			after the operation of postoperation (thromboembo	ts were evaluated 1 week operation, and there were perative complications embolism, neurologic or wound infection) in https://www.sembolism.	

Appendix D: Evidence matrixes – Intervention 1 (Acute normovolemic haemodilution)

	Level of evidence	No. of trials /	Patient population / Surgical procedure	Setting	lata a satis a			Results		Notes
Study	Quality	sample size			Intervention	Outcome	Intervention	Comparator	p-value	
Matot (2002)	Level II Fair	N=78	Adults undergoing <u>liver</u> resection	Hospital in Israel	ANH: Preoperatively, blood was transfused into standard citrate-phosphate-dextrose blood storage bags, and was simultaneously replaced by colloid solutions. The volume of blood collected was 2,020 ± 412 mL.	Adverse cardiac, renal, or neurological outcomes	0/39 (0%)	0/39 (0%)	Not estimable	
Sanders (2004)	Level II Fair	N=160	Adults undergoing <u>major</u> gastrointestinal surgery (colorectal, gastric, or	Hospital in UK	ANH: Maximum 3 units of blood withdrawn preoperatively and replaced	Pyrexia	0/78 (0%)	3/82 (4%)	P=0.21	
	pancreatic) ⁴				with warmed cell-free fluid to maintain normovolemia. At the end of the operation, all	UTI	8/78 (10%)	7/82 (9%)	P=0.71	
			autologous blood was retransfused.	RTI	2/78 (3%)	1/82 (1%)	P=0.54			
						Wound infection	3/78 (4%)	6/82 (7%)	P=0.35	
						Deep infection	1/78 (1%)	0/78 (0%)	P=0.48	
						Septicaemia	1/78 (1%)	1/82 (1%)	P=0.97	
						DVT	2/78 (3%)	2/82 (2%)	P=0.96	
						PE	0/78 (0%)	2/82 (2%)	P=0.31	
						Anastomotic leak	0/78 (0%)	3/82 (4%)	P=0.21	
	1	1						1		

Abbreviations: ANH, acute normovolemic haemodilution; CABG, coronary artery bypass graft; CI, confidence interval; CPB, cardiopulmonary bypass; DVT, deep vein thrombosis; Hb, haemoglobin; het, heterogeneity; NA, not applicable; NR, not reported; MI; myocardial infarction; PE, pulmonary embolism; RCT, randomised controlled trial; RR, relative risk; RTI, respiratory tract infection.

¹ Trials were included irrespective of whether they included major or minor liver resections, normal or cirrhotic livers, vascular occlusion was used or not, and irrespective of the reason for liver resection.

² Most patients underwent primary total hip replacement, with 15 revision hip arthroplasties (seven in ANH and eight in standard transfusion) and one hip resurfacing procedure.

³ Procedures included single and multiple valve surgery, aortic root surgery, coronary surgery combined with valve surgery, or partial left ventriculectomy.

⁴ These operations were considered high risk (>40%) for allogeneic transfusion

Key question(s): In patients undergoing surgery, what is the effect of <u>ANH</u> o	n <u>quality of life?</u>	Evidence table ref*: POQ3.I1.P6
1. Evidence base		,
No evidence found	A	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias
	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias
	С	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias
2. Consistency		
NA	А	All studies consistent
	В	Most studies consistent and inconsistency can be explained
	С	Some inconsistency, reflecting genuine uncertainty around question
	D	Evidence is inconsistent
	NA	Not applicable (one study only)
3. Clinical impact	·	
NA	A	Very large
	В	Substantial
	С	Moderate
	D	Slight/Restricted
4. Generalisability	1	
NA	А	Evidence directly generalisable to target population
	В	Evidence directly generalisable to target population with some caveats
	С	Evidence not directly generalisable to the target population but could be sensibly applied
	D	Evidence not directly generalisable to target population and hard to judge whether it is sensible to
5. Applicability	T	T
NA	A	Evidence directly applicable to Australian healthcare context
	В	Evidence applicable to Australian healthcare context with few caveats
	С	Evidence probably applicable to Australian healthcare context with some caveats
	D	Evidence not applicable to Australian healthcare context

Other factors			
EVIDENCE STA	TEMENT	MATRIX	
Please summarise t	the develop	ment group's synthesis of the evidence relating to t	he key question, taking all the above factors into account.
Component	Rating	Description	
Evidence base	NA		
2. Consistency	NA		
3. Clinical impact	NA		
4. Generalisability	NA		
5. Applicability	NA		
DRAFT EVIDEN Based on the body of			
		dergoing surgery in which substantial blood I	oss is anticipated, the effect of ANH on quality of life is unknown.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coaquilation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Key question(s): In patients undergoing surgery, what is the effect of <u>ANH</u> on <u>haemoglobin co</u>	ncen	tration?	Evidence table ref*: POQ3.I1.S1				
1. Evidence base			I				
Level II evidence: 8 RCTs: Akhlagh 2007 (poor quality; N=60); Friesen 2006 (fair quality; N=32); Hohn 2002 (poor quality; N=80); Lim 2003 (fair quality; N=30); Matot 2002 (fair quality; N=78); Obasi 2006 (poor quality;	Α	One or more level I studies with a low risk of bias or several level	n a low risk of bias or several level II studies with a low risk of bias				
N=62); Saricaoglu 2005 (good quality¹; N=30); Wolowczyk 2003 (fair quality; N=36).	В	One or two Level II studies with a low risk of bias or SR/several Level	vel III studies with a low risk of bias				
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	3				
2. Consistency	<u> </u>						
With the exception of Wolowczyk 2003 and Obasi 2006, all of the studies are consistent in finding no significant		All studies consistent					
association between ANH and haemoglobin concentration. Wolowczyk 2003 found that ANH was significantly associated with a lower median haemoglobin concentration at aortic clamping and clamp release but a	В	Most studies consistent and inconsistency can be explained					
significantly higher median haemoglobin concentration at 7 days postoperative. Obasi 2006 found significantly higher Hb concentration in the ANH group at 6-h post-surgery but not immediately post-surgery.	С	Some inconsistency, reflecting genuine uncertainty around questi	enuine uncertainty around question				
nigher no concentration in the ANN group at 6-11 post-surgery but not infinediately post-surgery.	D	Evidence is inconsistent					
	NA	Not applicable (one study only)					
3. Clinical impact							
See evidence summary table POQ3.I1.S1	Α	Very large					
Evidence inconsistent	В	Substantial					
	С	Moderate					
	D	Underpowered/inconsistent					
4. Generalisability							
The evidence is generalisable to an adult population who are undergoing elective surgery. The studies were conducted in adults undergoing cardiac, orthopaedic, liver and spinal surgery.	Α	Evidence directly generalisable to target population					
conducted in addits differ going cardiac, orthopaedic, liver and spirial surgery.	В	Evidence directly generalisable to target population with some cave	veats				
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied				
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to				
5. Applicability							
Countries included Iran, USA, Switzerland, South Korea, Israel, Poland, Turkey, and UK.	А	Evidence directly applicable to Australian healthcare context					
	В	Evidence applicable to Australian healthcare context with few cave	applicable to Australian healthcare context with few caveats				
		, , , , , , , , , , , , , , , , , , , ,	probably applicable to Australian healthcare context with some caveats				
	D	Evidence not applicable to Australian healthcare context					

There is a large variability in the reported timepoints and methods of analysis. The CRG considered outcomes at 7 days post operative most relevant. All the studies were underpowered. Intervention has a direct effect on Hb concentration so only trials that controlled would provide evidence.

The CRG considered that Obasi 2006 was not comparable because they did not report transfusion incidence or volume (ie, patients may not have been transfused).

Friesch 2006 was not taken into account as the study was in infants

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description						
1. Evidence base	С	Several Level II studies with a moderate risk of bias						
2. Consistency	С	Most studies consistent and inconsistency can be explained						
3. Clinical impact	D	Underpowered/inconsistent						
4. Generalisability	В	Evidence directly generalisable to target population with some caveats						
5. Applicability	В	Evidence probably applicable to Australian healthcare context with some caveats						

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of ANH on postoperative haemoglobin concentration is uncertain.

Abbreviations: ANH, acute normovolemic haemodilution; RCT, randomised control trial.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission ¹ Allocation to treatment groups was concealed from those responsible for recruiting subjects, patient characteristics and demographics were similar between treatment arms at baseline, all randomised patients were included in the analysis, the statistical methods were appropriate, and a transfusion protocol was reported. The study was not double-blinded.

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I1.S1 Characteristics and results of studies examining the effect of acute normovolemic haemodilution on haemoglobin concentration.

	Level of evidence	No. of trials /	Patient population / Surgical procedure	Setting		Outcome		Results		Notes
Study	Quality	sample size			Intervention		Intervention	Comparator	p-value	
Akhlagh (2007)	Level II Poor	N=60	Adults undergoing on-CPB CABG	Hospital in Iran	ANH and re-transfusion of autologous blood after separating the patient from the cardiopulmonary machine.	Mean (SD) haematocrit at 24 h postoperative, %	36.5 (1.5)	37 (2)	P=0.21	
Friesen (2006)	Level II Fair	N=32	Infants undergoing non- complex open cardiac surgery	Hospital in USA	ANH: 15 mL/kg whole blood withdrawn through the central venous catheter. Normovolemia was	Mean (SD) haematocrit at T1 (baseline), %	32 (3)	32 (4)	P=1.00	
					maintained by infusion of 1 mL of 5% albumin solution for each mL of blood withdrawn. Autologous blood re-transfused postoperatively.	Mean (SD) haematocrit at T2 (following conclusion of CPB and modified ultrafiltration), %	32 (8)	34 (6)	P=0.42	
						Mean (SD) haematocrit at T3 (20 minutes after T2), %	33 (7)	34 (6)	P=0.66	
						Mean (SD) haematocrit at T4 (after 2 hours in the ICU), %	35 (8)	34 (5)	P=0.67	
						ΔT2 – T3, %	1 (2)	1 (1)	P=1.00	
						ΔT2 – T4, %	3 (4)	0 (3)	P=0.009	
Hohn (2002)	Level II Poor	N=80	Adults undergoing on-CPB cardiac surgery	Hospital in Switzerland	ANH from a mean haematocrit of 43% to 28%.	Mean (SD) haematocrit baseline, %	43.3 (3.9)	43.2 (2.4)	P=0.89	
	ſ					Mean (SD) haematocrit immediate postoperative, %	25 (3.5)	25.7 (3.3)	P=0.36	

	Level of evidence	No. of trials /	Patient population / Surgical					Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Lim (2003)	Level II Fair	N=30	Adults undergoing <u>spinal</u> <u>surgery</u>	Hospital in South Korea	ANH and esmolol-induced controlled hypotension (E-ANH group): for ANH autologous blood was withdrawn from the radial artery aiming for 28% haematocrit. To maintain normovolemia, the first 500 mL of blood drawn was simultaneously replaced with an equal amount of 6% HES, and the blood thereafter was replaced with three times that volume of Lactated Ringer's solution.	Mean (SD) Hb one week postoperative, g/dL	11.3 (1.16)	11.3 (0.77)	P>0.05	
Matot (2002)	Level II Fair	N=78	Adults undergoing <u>liver</u> resection	Hospital in Israel	ANH: Preoperatively, blood was transfused into standard citrate-phosphate-dextrose blood storage bags, and was simultaneously replaced by colloid solutions. The volume of blood collected was 2,020 ± 412 mL.	Mean (SD) haematocrit (%) (before vs after)	40.8 ± 2.7 vs 23.5 ± 1.2 (P<0.05)	41.6 ± 3.2 vs 40.9 ± 2.8 (P>0.05)		
Obasi (2006)	Level II Poor	N=62	Adults undergoing <u>surgical</u> <u>procedures</u>	Hospital in Poland	Before the administration of anaesthesia, 500 to 800 mL of blood was effused from	Mean (SD) Hb preoperative, mmol/L	8.37 (0.43)	8.37 (0.63)	P=1.00	
					the patients (depending on body weight, values of Hb and haematocrit) with the simultaneous infusion of 6% HES in the ratio of 1:1 in an	Mean (SD) Hb immediately postoperative, mmol/L	6.45 (0.52)	6.46 (0.56)	P=0.94	
					aseptic and closed circuit.	Mean (SD) Hb 6 hours postoperative, mmol/L	7.20 (0.53)	6.48 (0.56)	P<0.00001	
Saricaoglu (2005)			Adults undergoing <u>hip</u> arthroplasty		ANH: autologous blood 15 mL/kg was withdrawn and replaced by ~15 mL/kg 6% HES	Median (95% CI) haematocrit preoperative, %	39.2 (34.6, 46.0)	HHD: 41.1 (37, 45.3) Control: 43.2 (35.8, 45.8)	P=0.5	
					HHD: 15 mL/kg HES administered without removal of any autologous blood	Median (95% CI) haematocrit postoperative , %	32.7 (26.5, 38.6)	HHD: 29.1 (26.5, 38.6) Control: 32.3 (26.5, 38.6)	P=0.398	

Appendix D: Evidence matrixes – Intervention 1 (Acute normovolemic haemodilution)

Charles	Level of evidence	No. of trials /	Patient population / Surgical procedure	Setting	lukaman Man	Outcome		Results		Notes
Study	Quality	sample size			Intervention	Outcome	Intervention	Comparator	p-value	
					Control: no haemodilution	Median (95% CI) haematocrit 24 h postoperative, %	32.7 (30.1, 40.1)	HHD: 34.9 (30.2, 36.7) Control: 32.9 (30, 36.5)	P=0.89	
Wolowczyk (2003)	Level II Fair	N=36	Adults undergoing <u>abdominal</u> <u>aortic aneurysm repair</u>	Hospital in UK	ANH and cell salvage: 15 g/kg of blood was withdrawn and replaced with a similar volume of 6% HES	Median (range) Hb preoperative, g/dL	14.2 (12.1 to 16.5)	13.8 (12.1 to 15.6)	P=0.57	
		(including cell salvage)	Median (range) Hb post-ANH, g/dL	9.4 (7.0 to 12.1)	NA	NA				
				Median (range) Hb at aortic clamping, g/dL	9.2 (6.8 to 10.6)	11.3 (7.2 to 14.5)	P=0.001			
						Median (range) Hb at clamp release, g/dL	7.7 (6.6 to 9.3)	9.1 (5.1 to 11.9)	P=0.004	
						Median (range) Hb at 1-2 hours postoperative, g/dL	10.8 (8.8 to 13.3)	10.3 (8.1 to 12.7)	P=0.68	
				Median (range) Hb at 1 day postoperative, g/dL	10.4 (8.3 to 12.4)	10.4 (8.2 to 12.8)	P=0.68			
						Median (range) Hb at 2 days postoperative, g/dL	10.6 (8.2 to 13.3)	9.7 (8.5 to 13.7)	P=0.60	
Abbas dations CAD						Median (range) Hb at 7 days postoperative, g/dL	11.5 (10.2 to 12.4)	10.7 (9.1 to 11.9)	P=0.021	

Abbreviations: CABG, coronary artery bypass graft; CI, confidence interval; CPB, cardiopulmonary bypass; Hb, haemoglobin; HES, hydroxyethyl starch; HHD, hypervolemic haemodilution; NA, not applicable; SD, standard deviation.

Key question(s):			Evidence table ref*:			
In patients undergoing surgery, what is the effect of ANH on reoperation for the	oleed	ing?	POQ3.I1.S2			
1. Evidence base						
Level I evidence: Carless 2004 (fair quality ¹ ; 7 trials, N=NR; adults undergoing any type of surgery)	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias			
Level II evidence: Hohn 2002 (poor quality; N=80)	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	3			
2. Consistency						
All studies are consistent in finding no significant impact. The studies were underpowered to find a significant difference.	Α	All studies consistent				
uniciones.	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact						
Results from Carless 2004: RR 1.59 (0.20, 12.53); 7 trials; N=NR	Α	Very large				
Results from Hohn 2002: RR 7.35 (0.39, 137.84)	В	Substantial				
	С	Moderate				
	D	No difference/underpowered				
4. Generalisability						
The evidence is generalisable to an adult population who are undergoing elective surgery.	Α	Evidence directly generalisable to target population				
	В	Evidence directly generalisable to target population with some cave	veats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard	to judge whether it is sensible to			
5. Applicability						
All studies were conducted in developed countries.	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few cave	eats			
		Evidence probably applicable to Australian healthcare context with some caveats				
	D	Evidence not applicable to Australian healthcare context				

Quality of RCTs in Carless 2004 not reported.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

		0 1 9
Component	Rating	Description
1. Evidence base	С	One Level I study and one Level II study with a low risk of bias
2. Consistency	Α	All studies consistent
3. Clinical impact	D	No difference/underpowered
4. Generalisability	В	Evidence directly generalisable to target population with some caveats
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of ANH on risk of reoperation for bleeding is uncertain.

Abbreviations: ANH, acute normovolemic haemodilution; NR, not reported; RR, relative risk.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission 1 No quality assessment of included studies were undertaken, the characteristics of the individual studies were appropriately summarised, the sources of heterogeneity were not sufficiently explored.

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I1.S2 Characteristics and results of studies examining the effect of acute normovolemic haemodilution on reoperation for bleeding.

	Level of evidence Ouality No. of trials / sample size Patient population / Surgical procedure Setting Intervention Outcome	No. of trials /	Patient population / Surgical	Calling	Intervention (0.1	Results			Notes
Study		Outcome	Intervention	Comparator	p-value					
Carless (2004)	Level I Fair	7 trials (quality NR) N=NR	Adults undergoing any type of surgery	All studies conducted in countries with well developed healthcare systems (not specifically Aus/NZ).	ANH	Proportion of patients who underwent reoperation for bleeding	RR (95% CI):1.59 (0.20, 12.53)	P>0.05	Phet=NR
Hohn (2002)	Level II Poor	N=80	Adults undergoing <u>on-CPB</u> cardiac surgery	Hospital in Switzerland	ANH from a mean haematocrit of 43% to 28%.	Proportion of patients who underwent reoperation for bleeding	3/39 (8%)	0/41 (0%)	P=0.18	

Abbreviations: ANH, acute normovolemic haemodilution; CI, confidence interval; CPB, cardiopulmonary bypass; het, heterogeneity; NR, not reported; RR, relative risk.

Key question(s):			Evidence table ref*:			
In patients undergoing surgery, what is the effect of ANH on correction/preve	ntion	of DIC and coagulopathy?	POQ3.I1.S3			
1. Evidence base						
Level II evidence: Saricaoglu 2005 (good quality ¹ ; N=30; adults undergoing hip arthroplasty).	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias			
Friesen 2006 was not taken into account as the study was in infants. (fair quality; N=32; infants undergoing non-	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias			
complex open cardiac surgery)	С	One or two Level III studies with a low risk of bias or Level I or II st	tudies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	S			
2. Consistency						
The two trials do not report the same coagulopathy outcomes; therefore the results are non-comparable.	Α	All studies consistent				
	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around questi	ion			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact						
See evidence summary table POQ3.I1.S3	Α	Very large				
	В	Substantial				
	С	Moderate				
	D	Slight/Restricted				
4. Generalisability						
Saricaoglu 2005 was in adults undergoing hip arthroplasty.	Α	Evidence directly generalisable to target population				
	В	Evidence directly generalisable to target population with some case	veats			
	С	Evidence not directly generalisable to the target population but co	ould be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to	to judge whether it is sensible to			
5. Applicability						
Saricaoglu 2005 was conducted in Turkey.	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few cave				
	С	Evidence probably applicable to Australian healthcare context with	h some caveats			
	D	Evidence not applicable to Australian healthcare context				

Friesen 2006 was not taken into account as the study was in infants.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	Two Level II studies with moderate risk of bias
2. Consistency	NA	The two trials do not report the same coagulopathy outcomes; therefore the results are non-comparable
3. Clinical impact	D	Slight/Restricted clinical impact
4. Generalisability	С	Evidence is not directly generalisable to target population. One trial included 30 adult patients
5. Applicability	С	Evidence probably applicable to Australian healthcare context with some caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of ANH on coagulation parameters is uncertain.

Abbreviations: ANH, acute normovolemic haemodilution; DIC, disseminated intravascular coagulation.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

¹ Allocation to treatment groups was concealed from those responsible for recruiting subjects, patient characteristics and demographics were similar between treatment arms at baseline, all randomised patients were included in the analysis, the statistical methods were appropriate, and a transfusion protocol was reported. The study was not double-blinded.

POQ3.I1.S3 Characteristics and results of studies examining the effect of acute normovolemic haemodilution on correction/prevention of DIC and coagulopathy.

	Level of evidence	No. of trials /	Patient population /					Results					
Study	Quality	sample size Surgical procedure Setting Intervention Outcome		Outcome	Intervention	Comparator	p-value						
Friesen (2006)	Level II	N=32	Infants undergoing	Hospital in	ANH: 15 mL/kg whole blood	Mean (SD) platelet count at T1 (baseline), 109/L	353 (92)	335 (92)	P=0.58				
	Fair		non-complex open cardiac surgery	USA	was maintained by infusion of 1 mL	venous catheter. Normovolemia	venous catheter. Normovolemia was maintained by infusion of 1 mL	venous catheter. Normovolemia was maintained by infusion of 1 mL	Mean (SD) platelet count at T2 (following conclusion of CPB and modified ultrafiltration), 10°/L	126 (49)	140 (47)	P=0.42	
					of blood withdrawn. Autologous blood re-transfused	Mean (SD) platelet count at T3 (20 minutes after T2), 109/L	161 (55)	158 (57)	P=0.88				
				postoperatively.	postoperatively.	Mean (SD) platelet count at T4 (after 2 hours in the ICU), 10 ⁹ /L	207 (53)	217 (59)	P=0.62				
						Mean (SD) platelet count, ΔT2 – T3 (platelet count), 10°/L	36 (22)	18 (17)	P=0.018				
						Mean (SD) platelet count, ΔT2 – T4, 109/L	82 (43)	70 (42)	P=0.43				
								Mean (SD) platelet aggregation at T1 (baseline), seconds	205 (62)	189 (54)	P=0.44		
							Mean (SD) platelet aggregation at T2 (following conclusion of CPB and modified ultrafiltration), seconds	222 (71)	210 (70)	P=0.63			
										Mean (SD) platelet aggregation at T3 (20 minutes after T2), seconds	144 (58)	159 (72)	P=0.52
										Mean (SD) platelet aggregation at T4 (after 2 hours in the ICU), seconds	112 (23)	113 (32)	P=0.92
							Mean (SD) platelet aggregation, ΔT2 – T3 (platelet count), seconds	-78 (53)	-49 (77)	P=0.22			
						Mean (SD) platelet aggregation, ΔT2 – T4, seconds	-109 (67)	-97 (64)	P=0.61				
						Mean (SD) prothrombin time at T1 (baseline), seconds	13.4 (0.9)	14.1 (1.1)	P=0.058				
								Mean (SD) prothrombin time at T2 (following conclusion of CPB and modified ultrafiltration), seconds	20.4 (4.3)	19.9 (3.8)	P=0.73		
				Mean (SD) prothrombin time at T3 (20 minutes after T2), seconds	18.1 (3.1)	18.9 (3.6)	P=0.51						
						Mean (SD) prothrombin time at T4 (after 2 hours in the ICU), seconds	15.9 (2.1)	16.8 (2.0)	P=0.22				

Appendix D: Evidence matrixes – Intervention 1 (Acute normovolemic haemodilution)

	Level of evidence	No. of trials /	Patient population /					Results	
Study	Quality	sample size	Surgical procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value
						Mean (SD) prothrombin time, ΔT2 – T3 (platelet count), seconds	-2.3 (1.9)	-0.9 (1.2)	P=0.015
						Mean (SD) prothrombin time, ΔT2 – T4, seconds	-4.5 (3.2)	-3.0 (2.7)	P=0.16
						Mean (SD) aPTT at T1 (baseline), seconds	35.9 (9.3)	36.9 (8.7)	P=0.76
						Mean (SD) aPTT at T2 (following conclusion of CPB and modified ultrafiltration), seconds	46.7 (14.2)	44.1 (12.6)	P=0.59
						Mean (SD) aPTT at T3 (20 minutes after T2), seconds	42.2 (14.1)	43.7 (13.1)	P=0.76
						Mean (SD) aPTT at T4 (after 2 hours in the ICU), seconds	37.8 (13.2)	41.9 (17.2)	P=0.46
						Mean (SD) aPTT, ΔT2 – T3 (platelet count), seconds	-4.4 (7.7)	-0.4 (9.6)	P=0.20
						Mean (SD) aPTT, ΔT2 – T4, seconds	-8.9 (11.0)	-2.3 (16.7)	P=0.20
						Mean (SD) fibrinogen concentration at T1 (baseline), mg/dL	235 (63)	215 (55)	P=0.35
						Mean (SD) fibrinogen concentration at T2 (following conclusion of CPB and modified ultrafiltration), mg/dL	109 (37)	129 (38)	P=0.14
						Mean (SD) fibrinogen concentration at T3 (20 minutes after T2), mg/dL	132 (44)	128 (32)	P=0.77
						Mean (SD) fibrinogen concentration at T4 (after 2 hours in the ICU), mg/dL	152 (51)	146 (36)	P=0.70
						Mean (SD) fibrinogen concentration, ΔT2 – T3 (platelet count), seconds	14 (9)	-1 (16)	P=0.0027
						Mean (SD) fibrinogen concentration ΔT2 – T4, seconds	35 (18)	17 (20)	P=0.019
Saricaoglu (2005)	Level II Good	N=30	Adults undergoing <u>hip</u> arthroplasty	Hospital in Turkey	ANH: autologous blood 15 mL/kg was withdrawn and replaced by ~15mL/kg 6% HES HHD: 15 mL/kg HES administered	Median (95% CI) preoperative platelet count, 1000/mm ³	280 (132, 367)	HHD: 286 (240, 387) Control: 285 (240, 387)	P=0.98
					without removal of any autologous blood <u>Control</u> : no haemodilution	Median (95% CI) postoperative platelet count, 1000/mm ³	258 (123, 354)	HHD: 204 (167, 300) Control: 241 (175, 310)	P=0.96

Appendix D: Evidence matrixes – Intervention 1 (Acute normovolemic haemodilution)

	Level of evidence	No. of trials /	Patient population /				Results			
Study	Quality	sample size	Surgical procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
						Median (95% CI) 24 h postoperative platelet count, 1000/mm ³	283 (138, 356)	HHD: 195 (163, 300) <u>Control</u> : 283 (190, 356)	P=0.010 (HHD)	
						Median (95% CI) preoperative INR	1.1 (0.92, 1.3)	HHD: 1.15 (0.95, 1.4) Control: 1.15 (0.92, 1.14)	P=0.6	
						Median (95% CI) postoperative INR	1.2 (1.1, 2.3)	HHD: 1.4 (1.2, 1.5) Control: 1.35 (1.2, 1.5)	P=0.052	
						Median (95% CI) 24 h postoperative INR	1.2 (1.1, 1.87)	HHD: 1.2 (1.1, 1.3) Control: 1.2 (1.1, 1.3) 1.3)	P=0.68	
						Median (95% CI) preoperative aPTT, seconds	27.6 (26.4, 35.9)	HHD: 28.5 (26.8, 32.1) Control: 27.6 (26.4, 32.1)	P=0.4	
						Median (95% CI) postoperative aPTT, seconds	26.75 (23.8, 32.3)	HHD: 33.8 (30.1, 35.6) Control: 27.5 (24.7, 34.2)	P=0.01 P(ANH v HHD)<0.008	
						Median (95% CI) 24 h postoperative aPTT, seconds	26.5 (24.7, 30.1)	HHD: 30.1 (24.7, 34.2) <u>Control</u> : 24.2 (24.2, 34.7)	P=0.182	

Abbreviations: ANH, acute normovolemic haemodilution; aPTT, activated partial thromboplastin time; CI, confidence interval; CPB, cardiopulmonary bypass; HES, hydroxyethyl starch; HHD, hypervolemic haemodilution; ICU, intensive care unit; INR, international normalised ratio; SD, standard deviation.

Key question(s): In patients undergoing surgery, what is the effect of ANH on hospital length of	of sta	γ?	Evidence table ref*: POQ3.I1.S5			
1. Evidence base		-	-			
Level I evidence: Carless 2004 (fair quality; 3 trials ¹ , N=96; adults undergoing any type of surgery) and Gurusamy 2009 (good quality; 1 trial, N=130; adults undergoing liver resection)	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bi				
Level II evidence: 6 RCTs: Bennett 2006 (fair quality; N=155); Casati 2002 (poor quality; N=204); Casati 2004 (fair quality; N=100); Hohn 2002 (poor quality; N=80); Jarnagin 2008 (fair quality; N=130); Sanders 2004 (fair	В	One or two Level II studies with a low risk of bias or SR/several Le	evel III studies with a low risk of bias			
quality; N=160)	С	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bia	S			
2. Consistency						
All the studies except Bennett (2006) are consistent in finding no significant impact.	Α	All studies consistent				
	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around question				
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact	•					
Results from Carless 2004: Mean difference (95% CI), days: 0.21 (-1.26, 1.68)	Α	Very large				
Results from Gurusamy 2009: Mean difference (95% CI), days: 0.0 (-2.66, 2.66)	В	Substantial				
Results from Bennett (2006): ANH vs. control; median (IQR), days: 7 (6, 9) vs. 8 (6, 11); P=0.03	С	Moderate				
Results from Level II studies – see evidence summary table POQ3.I1.P5	D	Slight/Restricted				
4. Generalisability						
The evidence is generalisable to an adult population who are undergoing elective surgery. The studies were conducted in adults undergoing cardiac, orthopaedic, liver and other surgeries.	Α	Evidence directly generalisable to target population				
conducted in addits undergoing cardiac, orthopaedic, liver and other surgenes.	В	Evidence directly generalisable to target population with some ca	veats			
	С	Evidence not directly generalisable to the target population but co	ould be sensibly applied			
	D	Evidence not directly generalisable to target population and hard	to judge whether it is sensible to			
5. Applicability						
All the studies were conducted in developed countries.	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few cav				
	С		vidence probably applicable to Australian healthcare context with some caveats			
	D	Evidence not applicable to Australian healthcare context				

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EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	Two Level I studies and several Level II studies with a low risk of bias
2. Consistency	С	All studies consistent
3. Clinical impact	D	No statistically significant impact
4. Generalisability	В	Evidence directly generalisable to target population with some caveats
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of ANH on length of hospital stay is uncertain.

Abbreviations: ANH, acute normovolemic haemodilution; CI, confidence interval; RCT, randomised controlled trial.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission ¹ Fair to poor quality.

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I1.S5 Characteristics and results of studies examining the effect of acute normovolemic haemodilution on hospital length of stay.

	Level of evidence	No. of trials /	Patient population / Surgical					Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Carless (2004)	Level I Fair	3 trials (quality NR) N=96	Adults undergoing any type of surgery	All studies conducted in countries with well developed healthcare systems (not specifically Aus/NZ).	ANH	Mean difference (95% CI) hospital length of stay, days	0.21 (-1.26, 1.68)		P>0.05	Phet=NR
Gurusamy (2009)	Level I Good	1 trial (fair quality) N=130	Patients undergoing liver resection ¹	Studies conducted in USA, Israel, and China.	ANH	Mean difference (95% CI) length of hospital stay, days	0.0 (-2.66, 2.66)		P>0.05	Phet=NA
Bennett (2006)	Level II Fair	N=155	Adults undergoing <u>elective hip</u> <u>surgery.</u> ² Anticipated blood loss between 1 to 1.5 L	Hospital in UK	Autologous blood was collected immediately before surgery, aiming to reduce haemoglobin concentration to a target of 110 g per L. All autologous blood was returned within 6 hours of collection, starting on wound closure or sooner if a transfusion trigger was reached.	Median (IQR) length of hospital stay, days	7 (6, 9)	8 (6,11)	P=0.03	
Casati (2002)	Level II Poor	N=204	Adults undergoing <u>cardiac</u> <u>surgery</u> ³	Hospital in Italy	Low volume ANH: 5-8 mL/kg of blood withdrawn before systemic heparinisation and replaced with colloid solutions.	Median (IQR) postoperative hospital stay, days	7 (6, 9)	7 (6, 8.25)	P=0.54	
Casati (2004)	Level II Fair	N=100	Adults undergoing off-CPB CABG	Hospital in Italy	ANH with tranexamic acid (with tranexamic acid as control)	Mean (IQR) postoperative hospital stay, days	6 (6, 7)	6 (6, 7)	NR	
Hohn (2002)	Level II Poor	N=80	Adults undergoing <u>on-CPB</u> cardiac surgery	Hospital in Switzerland	ANH from a mean haematocrit of 43% to 28%.	Mean (SD) postoperative length of hospital stay, days	13.1 (3.7)	13.4 (8.3)	P=0.83	

Appendix D: Evidence matrixes – Intervention 1 (Acute normovolemic haemodilution)

	Level of evidence	No. of trials /	Patient population / Surgical	Catting		Outcomo		Notes		
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Jarnagin (2008)	Level II Fair	N=130	Adults undergoing major hepatic resection (three or more liver segments) for any diagnosis, with or without any other planned procedures	Hospital in USA	ANH: blood was withdrawn to a target Hb concentration of 8.0 g/dL, with a maximum of 3 L of blood removed. Normovolemia was maintained by replacing half of the removed blood volume with 5% albumin and the other half with crystalloid.	Median (range) length of hospital stay, days	7 (5 to 50)	7 (4 to 26)	P=0.33	
Sanders (2004)	Level II Fair	N=160	Adults undergoing <u>major</u> gastrointestinal surgery (colorectal, gastric, or pancreatic) ⁴	Hospital in UK	Maximum 3 units of blood withdrawn and transfused into blood bags containing citrate-phosphate-dextrose (anticoagulant). Warmed cell-free fluid was administered during blood withdrawal to maintain normovolemia. At the end of the operation, all autologous blood was re-transfused.	Median (range) length of hospital stay, days	8 (5 to 110)	10 (5 to 92)	NS	

Abbreviations: ANH, acute normovolemic haemodilution; CABG, coronary artery bypass graft; CI, confidence interval; CPB, cardiopulmonary bypass; IHb, haemoglobin; het, heterogeneity; NA, not applicable; NR, not reported; NS, not significant; QR, interquartile range; RCT, randomised controlled trial; SD, standard deviation.

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¹ Trials were included irrespective of whether they included major or minor liver resections, normal or cirrhotic livers, vascular occlusion was used or not, and irrespective of the reason for liver resection.

² Most patients underwent primary total hip replacement, with 15 revision hip arthroplasties (seven in ANH and eight in standard transfusion) and one hip resurfacing procedure.

³ Procedures included single and multiple valve surgery, aortic root surgery, coronary surgery combined with valve surgery, or partial left ventriculectomy.

⁴ These operations were considered high risk (>40%) for allogeneic transfusion.

Key question(s):			Evidence table ref*:		
In patients undergoing surgery, what is the effect of ANH on ICU admission a	ind le	ength of stay?	POQ3.I1.S6		
1. Evidence base					
Level II evidence: 3 RCTs: Casati 2002(poor quality; N=204); Casati 2004 (fair quality; N=100); Hohn 2002 (poor	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias		
quality; N=80)	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias		
	С	One or two Level III studies with a low risk of bias or Level I or II st	tudies with a moderate risk of bias		
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	S		
2. Consistency					
All studies are consistent in finding no significant impact	Α	All studies consistent			
	В	Most studies consistent and inconsistency can be explained			
	С	Some inconsistency, reflecting genuine uncertainty around questi	ion		
	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact					
ANH vs. control Casati 2002: Median (IQR) ICU stay, days: 1 (1, 1) vs. 1 (1, 2); P=0.49; N=204	A Very large				
Casati 2004: Mean (IQR) ICU stay, days: 1 (1, 1) vs. 1 (1, 1); P=1; N=100	В	Substantial			
Hohn 2002: Mean (SD) length of ICU stay, days: 3.1 (1.3) vs. 3.0 (1.3); P=0.73; N=80	С	Moderate			
	D	No difference/underpowered			
4. Generalisability					
All studies were in adults undergoing cardiac surgery.	Α	Evidence directly generalisable to target population			
	В	Evidence directly generalisable to target population with some case	veats		
	С	Evidence not directly generalisable to the target population but co	ould be sensibly applied		
	D	Evidence not directly generalisable to target population and hard	to judge whether it is sensible to		
5. Applicability					
The studies were conducted in Italy (Casati 2002 and 2004) and Switzerland (Hohn 2002)	Α	Evidence directly applicable to Australian healthcare context			
	В	Evidence applicable to Australian healthcare context with few cave			
	С	Evidence probably applicable to Australian healthcare context with some caveats			
	D	Evidence not applicable to Australian healthcare context			

Other factors		

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence	С	Three Level II studies with moderate risk of bias
2. Consistency	А	All studies consistent
3. Clinical impact	D	No difference/underpowered
4. Generalisability	В	Evidence directly generalisable to target population with some caveats
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of ANH on length of ICU stay is uncertain.

Abbreviations: ANH, acute normovolemic haemodilution; ICU, intensive care unit; IQR, interquartile range; RCT, randomised controlled trial.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Characteristics and results of studies examining the effect of acute normovolemic haemodilution on ICU admission and length of POQ3.I1.S6 stay.

	Level of evidence	No. of trials /	Patient population / Surgical	0 111		Outcome	Results			
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Casati (2002)	Level II Poor	N=204	Adults undergoing <u>cardiac</u> <u>surgery</u> ¹	Hospital in Italy	Low volume ANH: 5-8 mL/kg of blood withdrawn before systemic heparinisation and replaced with colloid solutions.	Median (IQR) ICU stay, days	1 (1, 1)	1 (1, 2)	P=0.49	
Casati (2004)	Level II Fair	N=100	Adults undergoing off-CPB CABG	Hospital in Italy	ANH with tranexamic acid (with tranexamic acid as control)	Mean (IQR) ICU stay, days	1 (1, 1)	1 (1, 1)	P=1	
Hohn (2002)	Level II Poor	N=80	Adults undergoing on-CPB cardiac surgery	Hospital in Switzerland	ANH from a mean haematocrit of 43% to 28%.	Mean (SD) length of ICU stay, days	3.1 (1.3)	3.0 (1.3)	P=0.73	

Abbreviations: ANH, acute normovolemic haemodilution; CABG, coronary artery bypass graft; CPB, cardiopulmonary bypass; ICU, intensive care unit; IQR, interquartile range; SD, standard deviation.

1 Procedures included single and multiple valve surgery, aortic root surgery, coronary surgery combined with valve surgery, or partial left ventriculectomy.

Recommendation(s) for acute normovolemic haemodilution

RECOMMENDATION What recommendation(s) does the guideline development group draw from this evidence? Use action statements where possible.	GRADE	GRADE RELEVENCE FOR TABLET					
In adult patients undergoing surgery in which substantial blood loss is anticipated, the use of acute	С	PO3.I1.P1,					
normovolemic haemodilution should be considered.							
IMPLEMENTATION OF RECOMMENDATION Please indicate yes or no to the following questions. Where the answer is yes please provide explanatory information about this. This information will be used to develop the implementation plan for the quidelines.							
Will this recommendation result in changes in usual care?		YES	NO				
Use of ANH will increase (not widely used at present).		-					
Are there any resource implications associated with implementing this recommendation?		YES	NO				
Training and equipment costs.							
Will the implementation of this recommendation require changes in the way care is currently organised?		YES	NO				
Increased preoperative time; placement of bigger neck (jugular) lines; increased vascular complications (due to neck lines); th	eatre sche	duling;				
extension of theatre utilisation time; requirement for a protocol.							
Are the guideline development group aware of any barriers to the implementation of this recommendation		YES	NO				
Appropriate venesection equipment; blood bag requirements from Blood Bank.							
What could help to facilitate implementation of the recommendation?							
Development of a local ANH protocol.							

Intervention 2 – Intraoperative cell salvage

Key question(s):			Evidence table ref*:				
In patients undergoing surgery, what is the effect of intraoperative cell salvag	<u>e</u> on	<u>transfusion incidence?</u>	POQ3.I2.P1				
1. Evidence base							
Pivotal evidence: 1 level I SR (Carless 2006): good quality; adults undergoing any type of surgery; all of the studies used a transfusion protocol; includes 5 RCTs (N=382): 2 vascular (1 fair and 1 poor quality), 2 cardiac	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias				
(fair quality), 1 orthopaedic (fair quality)	В	One or two Level II studies with a low risk of bias or SR/several Leve	el III studies with a low risk of bias				
Supportive data published after Carless 2006 from 6 level II studies: Damgaard 2006 (cardiac; good quality;	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias				
N=60); Goel 2007 (cardiac; fair quality; N=50); Mercer 2004 (vascular; good quality; N=81); Murphy 2005 (cardiac; fair quality; N=61); Wiefferink 2007 (cardiac; fair quality; N=30); Zhang 2004 (orthopaedic; poor quality; N=48)	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	3				
2. Consistency							
The results are not consistent by surgery type (See meta-analysis in technical report).	Α	All studies consistent					
Within surgery types The two trials assessing cell salvage in orthopaedic surgery are consistent in reporting a significantly lower	В	Most studies consistent and inconsistency can be explained					
transfusion incidence in the cell salvage groups. There is inconsistency in the trials assessing cell salvage in cardiac and vascular surgery.	С	Some inconsistency, reflecting genuine uncertainty around question					
cardiac and vascular surgery.	D	Evidence is inconsistent	inconsistent				
	NA	Not applicable (one study only)					
3. Clinical impact							
Meta-analysis of systematic review and 4 of the RCTs (Goel 2007; Mercer 2004; Murphy 2005; Zhang	Α	Very large					
2004) ¹ . Patients transfused with allogeneic blood. All surgery types – RR 0.61 (0.46, 0.81); 9 RCTs (N=621)	В	Substantial					
Cardiac surgery – RR 0.63 (0.41, 0.98); 4 RCTs (N=316)	С	Moderate					
Orthopaedic surgery – RR 0.33 (0.22, 0.49); 2 RCTs (N=88)	D	Slight/Restricted					
Vascular surgery – RR 0.83 (0.67, 1.03); 3 RCTs (N=217) 4. Generalisability							
The evidence is generalisable to an adult population who are undergoing elective surgery.	Α	Evidence directly generalisable to target population					
	В	Evidence directly generalisable to target population with some cav	veats.				
	C	Evidence not directly generalisable to the target population but co					
			,				
E. A P L. P.	D	Evidence not directly generalisable to target population and hard t	o juage whether it is sensible to				
5. Applicability		I					
Most of the studies were conducted in developed countries. Goel 2007 was conducted in India and Zhang was conducted in China.	Α	Evidence directly applicable to Australian healthcare context					
	В	Evidence applicable to Australian healthcare context with few cav					
	С	Evidence probably applicable to Australian healthcare context with	n some caveats				
	D	Evidence not applicable to Australian healthcare context					

Cardiac surgery included one good quality study; however, it couldn't be included in the meta-analysis because it reported transfusion of allogeneic blood components rather than allogeneic blood. Therefore this study was not taken into account when rating the evidence base for cardiac surgery.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	В	One level I study and several level II studies with low risk of bias
2. Consistency	С	Some inconsistency, reflecting genuine uncertainty around question
3. Clinical impact	В	There is a substantial clinical impact
4. Generalisability	В	Evidence directly generalisable to target population with some caveats
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, intraoperative cell salvage reduces the incidence of allogeneic blood transfusion.

Abbreviations: RCT, randomised controlled trial; RR, relative risk; SR, systematic review.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission ¹ Damgaard 2006 and Wiefferink 2007 were not included in the meta-analysis, because instead of measuring the incidence of allogeneic blood transfusion they measured transfusion of blood components and transfusion of packed RBCs respectively. See Summary Table I2.P1.

POQ3.I2.P1 Characteristics and results of studies examining the effect of intraoperative cell salvage on transfusion incidence.

0	Level of evidence	No. of trials /	Patient population / Surgical procedure	Setting				Results		Notes
Study	Quality	sample size			Intervention	Outcome	Intervention	Comparator	p-value	
Carless (2006)	Level I Good	5 trials (1 poor quality, 4 fair quality) N=382	Adults undergoing any type of surgery.	All studies were conducted in developed countries.	Intraoperative cell salvage	Allogeneic blood	74/191 (41%)	113/191 (59%)	P=0.03	Phet=0.01
Damgaard (2006)	Level II Good	N=60	Adults undergoing sub-acute coronary bypass surgery without heart-lung machine.	Hospital in Denmark The continuously heparinised suction and reservoir belonging to the cell saver were used for all patients in both groups. The suctioned blood from patients in the cell saver group was processed and autotransfused before the patient was transferred to the ICU.		Allogeneic blood components	17/30 (57%)	21/29 (72%)	P=0.21	
Goel (2007)	Level II Fair	N=50	Adults undergoing off-pump coronary artery bypass grafting.	Hospital in India	Intraoperative cell salvage and autotransfusion of washed shed blood and transfusion of allogeneic blood if required.	Allogeneic blood	20/24 (83%)	25/25 (100%)	P=0.07	
Mercer (2004)	Level II Good	N=81	Adults undergoing surgery for abdominal aortic aneurysm.	Hospital in UK	Intraoperative cell salvage. Processed blood was returned to the patient as soon as haemostasis had been achieved.	Allogeneic blood	21/40 (53%)	31/41 (76%)	P=0.04	
Murphy (2005)	Level II Fair	N=61	Patients scheduled for non- emergency first-time CABG (off-pump)	Hospital in UK	Intraoperative cell salvage, with Autotransfusion of washed, salvaged red blood	Allogeneic blood components	5/30 (17%)	11/31 (36%)	P=0.11	
					cells at the completion of the operative procedure. Salvaged washed RBCs	Allogeneic blood	4/30 (13%)	7/31 (23%)	P=0.36	
					were autotransfused at the time of skin closure.	Platelets	2/30 (7%)	6/31 (19%)	P=0.17	
Wiefferink (2007)	Level II Fair	N=30	Adults undergoing CABG with CPB	Hospital in the Netherlands	Intraoperative cell salvage: The mediastinal and residual CPB blood was processed by a continuous autotransfusion system before reinfusion.	Allogeneic packed RBCs	8/15 (54%)	10/15 (67%)	P=0.46	

	Level of evidence	No. of trials /	Patient population / Surgical			Outcomo		Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Zhang (2004)	Level II Poor	N=48	Adults undergoing operation for scoliosis.	Hospital in China	Intraoperative cell salvage and retransfusion of washed autologous blood.	Allogeneic blood	11/36 (31%)	12/12 (100%)	P<0.00001	
Cardiac surgery										
Carless (2006)	Level I Good	2 trials (fair quality) N=206	Adult patients undergoing any type of surgery.	All studies were conducted in developed countries.	Intraoperative cell salvage	Allogeneic blood	31/103 (30%)	56/103 (54%)	P=0.0009	Phet=0.32
Orthopaedic sur	gery					•		•		•
Carless (2006)	Level I Good	1 trial (fair quality) N=40	Adult patients undergoing any type of surgery.	All studies were conducted in developed countries.	Intraoperative cell salvage	Allogeneic blood	6/20 (30%)	18/20 (90%)	P=0.002	Phet=NA
Vascular surger	у					•		•		•
Carless (2006)	Level I Good	2 trials (1 poor, 1 fair quality) N=136	Adult patients undergoing any type of surgery.	All studies were conducted in developed countries.	Intraoperative cell salvage	Allogeneic blood	37/68 (54%)	39/68 (57%)	P=0.58	Phet=0.58

Abbreviations: CABG, coronary artery bypass graft; CPB, cardiopulmonary bypass; ICU, intensive care unit; NA, not applicable; RBC, red blood cells.

Key question(s):			Evidence table ref*:				
In patients undergoing surgery, what is the effect of intraoperative cell salvage	<u>ge</u> on	transfusion volume?	POQ3.I2.P2				
1. Evidence base							
Pivotal evidence: 1 level 1 SR (Carless 2006); good quality; adults undergoing any type of surgery; includes 6	Α	One or more level I studies with a low risk of bias or several level	as or several level II studies with a low risk of bias				
RCTs (N=432): 2 cardiac (fair quality), 1 orthopaedic (fair quality), 3 vascular (2 fair quality, 1 poor quality) Supportive published after Carless 2006 from 7 level II studies: Bowley 2006 (trauma; fair quality; N=44); Damgaard 2006 (cardiac; good quality; N=60); Goel 2007 (cardiac; fair quality; N=50); Mercer 2004 (vascular;	В	One or two Level II studies with a low risk of bias or SR/several Level	ias or SR/several Level III studies with a low risk of bias				
good quality; N=81); Niranjan 2006 (cardiac; good quality; N=80); Selo-Ojeme 2007 (ruptured ectopic pregnancy fair quality; N=112); Wiefferink 2007 (cardiac; fair quality; N=30)	; C	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias					
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	:				
2. Consistency							
The results are not consistent by surgery type. Within surgery types (of the RCTs included in the updated meta-analysis¹ conducted herein)	Α	All studies consistent					
The results are consistently significant in cardiac surgery (3 RCTs), and consistently insignificant in vascular	В	Most studies consistent and inconsistency can be explained					
surgery (3 RCTs) (see meta-analysis). Consistency with RCTs not included in the updated meta-analysis	С	Some inconsistency, reflecting genuine uncertainty around question	tainty around question				
The results from Mercer 2004 are inconsistent with the results from the updated meta-analysis. Selo-Ojeme 2005 found that cell salvage in ruptured ectopic pregnancy significantly reduced the proportion of women requiring	D D	Evidence is inconsistent					
transfusion of > 1000 mL of blood. The results of the other RCTs agreed with those of the updated meta-analysis	s. NA	Not applicable (one study only)					
3. Clinical impact	•						
Meta-analysis of Carless 2006 and 2 RCTs (Goel 2007 and Bowley 2006) was conducted. All surgery types – mean difference -0.86 (-1.54, -0.18); 8 RCTs	Α	Very large					
Cardiac surgery – mean difference -0.58 (-0.93, -0.23); 3 RCTs (N=256)	В	Substantial					
Orthopaedic surgery – mean difference -2.04 (-2.58, -1.50); 1 RCT (N=40) Vascular surgery – mean difference 0.02 (-0.34, 0.38); 3 RCTs (N=186)	С	Moderate					
Penetrating trauma- mean difference -4.70 (-8.01, -1.39); 1 RCT (N=44)	D	No difference					
4. Generalisability							
Both elective (cardiac, orthopaedic, and vascular) surgery and surgery for traumatic injury are included in the review. The efficacy of intraoperative cell salvage is dependent on surgery type.	Α	Evidence directly generalisable to target population					
review. The efficacy of infraoperative cell salvage is dependent on surgery type.	В	Evidence directly generalisable to target population with some cave	veats				
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied				
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to				
5. Applicability							
Most of the studies were conducted in developed countries. Goel 2007 was conducted in India; however, the exclusion of this study does not impact the results. Bowley 2006 (trauma) was conducted in Johannesburg.	Α	Evidence directly applicable to Australian healthcare context					
enclusion of this study does not impact the results. Downey 2000 (trauma) was conducted in Johannesburg.	В	Evidence applicable to Australian healthcare context with few cave	eats (cardiac, ortho, vascular)				
	С	Evidence probably applicable to Australian healthcare context with	n some caveats (trauma)				
	D	Evidence not applicable to Australian healthcare context					

Further studies required to strengthen evidence base. Selo-Ojeme 2007 was a Nigerian study in women with ruptured ectopic pregnancy and the CRG decided to not make a separate Evidence Statement based on this study.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	В	In each surgery subgroup (except trauma) there are several Level II studies with a moderate risk of bias. In trauma surgery there is one Level II study with a moderate risk of bias.
2. Consistency	С	In vascular and orthopaedic surgery all studies are consistent. In cardiac surgery most studies are consistent and inconsistency can be explained
3. Clinical impact		Intraoperative cell salvage substantially reduces mean transfusion volume in orthopaedic and traumatic surgeries, moderately reduces mean transfusion in cardiac surgery, and does not significantly reduce mean transfusion in vascular surgery.
4. Generalisability		Evidence directly generalisable to target population with some caveats. Surgery types assessed include cardiac, orthopaedic, vascular, and surgery for penetrating trauma.
5. Applicability	В	In cardiac, vascular, and orthopaedic surgery evidence is applicable to Australian healthcare context with few caveats. The one study assessing trauma was conducted in Johannesburg.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, intraoperative cell salvage may reduce the volume of allogeneic blood transfused.

Abbreviations: RCT, randomised controlled trial; SR, systematic review.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{* &}lt;u>Interventions</u>: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hosp

²Mercer 2004 found that intraoperative cell salvage significantly reduces the median volume of allogeneic blood transfused in patients undergoing surgery for abdominal aortic aneurysm; a finding that is inconsistent with the meta-analytical results that reveal no significant impact of intraoperative cell salvage on volume transfused in patients undergoing vascular surgery.

POQ3.I2.P2 Characteristics and results of studies examining the effect of intraoperative cell salvage on transfusion volume.

	Level of evidence	No. of trials /	Patient population / Surgical		latem ention	Outcome		Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention		Intervention	Comparator	p-value	
Carless (2006)	Level I Good	6 trials (1 poor quality, 5 fair quality) N=432	Adults undergoing any type of surgery.	All studies were conducted in developed countries.	Intraoperative cell salvage	Units of allogeneic blood transfused Mean difference (95% CI)	-0.69 (-1.47, 0.08)		P=0.08	Phet<0.0001
Bowley (2006)	Level II Fair	N=44	Adults with penetrating torso injury requiring a laparotomy and who had exhibited hypotension either pre-hospital or an arrival and in whom there was considered to be a significant loss.	Hospital in Johannesburg	Intraoperative blood salvage with transfusion of both allogeneic and washed autologous blood.	Units of allogeneic blood transfused Mean (SD)	6.47 (5.14)	11.17 (6.06)	P=0.005	
Damgaard Level I (2006) Good	Level II Good		N=60 Adults undergoing sub-acute coronary bypass surgery without heart-lung machine.	Hospital in Denmark	heparinised suction and reservoir belonging to the cell saver were used for all patients in both groups. The suctioned blood from patients in the cell saver group was processed and autotransfused before the patient was transferred to the ICU.	Units of allogeneic blood components transfused Median (IQR)	1 (0 to 2)	2 (0 to 7)	P=0.06	
						Units of allogeneic packed RBCs transfused Median (IQR)	1 (0 to 2)	2 (0 to 5)	P=0.07	
						Units of FFP transfused (ICU) Median (IQR)	0 (0 to 0) Range: 0 to 4	0 (0,0) Range: 0 to 22	P=0.40	
						Units of FFP transfused (ward) Median (IQR)	0 (0 to 0) Range: 0 to 0	0 (0, 0) Range: 0 to 1	P=0.31	
				Units of platelets transfused Median (IQR)	0 (0 to 0) Range: 0 to 1	0 (0 to 0) Range: 0 to 1	P=NR			

	Level of evidence	No. of trials /	Patient population / Surgical		Interception			Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Goel (2007)	Level II Fair	N=50	Adults undergoing off-pump coronary artery bypass grafting.	Hospital in India	Intraoperative cell salvage and autotransfusion of washed shed blood and transfusion of allogeneic blood if required.	Units of allogeneic blood transfused Mean (SD)	1.5 (1.1)	2.4 (1.3)	P=0.008	
Mercer (2004)	Level II Good	N=81	Adults undergoing surgery for abdominal aortic aneurysm.	Hospital in UK	Intraoperative cell salvage. Processed blood was returned to the patient as soon as haemostasis had been achieved.	Units of allogeneic blood transfused Median (IQR)	1 (0 to 3)	3 (1 to 5)	P=0.012	
Niranjan (2006)	Level II Good	N=80	Adults undergoing first-time, isolated CABG	Hospital in UK	Intraoperative cell salvage, with Autotransfusion of washed, salvaged red blood cells at the conclusion of the procedure.	Volume of allogeneic blood transfused, mL Mean (SD)	On-CPB: 179 (214) Off-CPB: 141 (183) Combined: 159 (196)	On-CPB: 230 (240) Off-CPB: 595 (438) Combined: 413 (394)	On-CPB: P=0.048 Off-CPB: P<0.0001 Combined: P=0.0003	
Selo-Ojeme (2007)	Level II Fair	N=112	Women undergoing surgery for ruptured ectopic pregnancy	Hospital in Nigeria	Intraoperative cell salvage with transfusion of filtered autologous blood.	Patients transfused with ≥ 1000 mL with blood	34/56 (60%)	11/56 (20%)	P=0.0001	
Wiefferink (2007)	Level II Fair	N=30	Adults undergoing CABG with CPB	Hospital in the Netherlands	Intraoperative cell salvage: The mediastinal and residual CPB blood was processed by a continuous autotransfusion system before reinfusion.	Patients transfused with ≥ 2 units of allogeneic packed RBCs	2/15 (13%)	7/15 (47%)	P=0.08	
Cardiac surgery	1	1	-	<u> </u>		1	<u> </u>	1	1	Т
Carless (2006)	Level I Good	2 trials (fair quality) N=206	Adult patients undergoing any type of surgery.	All studies were conducted in developed countries.	Intraoperative cell salvage	Allogeneic blood Mean difference (95% CI)	-0.46 (-0.86, -0.05)		P=0.03	Phet=0.58
Orthopaedic sur	gery									•
Carless (2006)	Level I Good	1 trial (fair quality) N=40	Adult patients undergoing any type of surgery.	All studies were conducted in developed countries.	Intraoperative cell salvage	Allogeneic blood Mean difference (95% CI)	-2.04 (-2.58, -1.50)		P<0.00001	Phet=NA

Appendix D: Evidence matrixes – Intervention 2 (Intraoperative cell salvage)

Study	Level of evidence No. of trials / sample size	No. of trials /	Patient population / Surgical procedure	Setting	Intervention	Outcome	Results			Notes		
							Intervention	Comparator	p-value			
Vascular surgery	Vascular surgery											
Carless (2006)	Level I Good	3 trials (1 poor quality, 2 fair quality) N=186	Adult patients undergoing any type of surgery.	All studies were conducted in developed countries.	Intraoperative cell salvage	Allogeneic blood Mean difference (95% CI)	0.02 (-0.32, 0.52)		P=0.91	Phet=0.42		

Abbreviations: CABG, coronary artery bypass graft; CI, confidence interval; CPB, cardiopulmonary bypass; FFP, fresh frozen plasma; ICU, intensive care unit; IQR, interquartile range; NR, not reported; RBCs, red blood cells; SD, standard deviation.

Key question(s): In patients undergoing surgery, what is the effect of intraoperative cell salvage.	operative blood loss?	Evidence table ref*: POQ3.I2.P3				
1. Evidence base			,			
Pivotal evidence: 1 level 1 SR (Carless 2006); good quality; adults undergoing any type of surgery; includes 6 RCTs (N=431): 2 cardiac (fair quality), 1 orthopaedic (fair quality), 3 vascular (2 fair quality, 1 poor quality).	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias			
Supportive evidence published after Carless 2006 from 4 level II studies: Damgaard 2006 (cardiac; good quality; N=60); Mercer 2004 (vascular; good quality; N=81); Niranjan 2006 (cardiac; good quality; N=80); Zhang	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias				
2004 (orthopaedic; poor quality; N=48)	С	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	3			
2. Consistency						
Orthopaedic Intervention was associated with a significant reduction in blood loss in the 1 study included in Carless (2006) but	Α	All studies consistent				
there was no significant difference in Zhang 2004.	В	Most studies consistent and inconsistency can be explained				
Other surgery The results from Niranjan 2006 for on-CPB CABG patients are inconsistent with the results from Carless 2006.	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
The results from Mercer 2004 and the off-CPB CABG patients in the Niranjan 2006 trial are consistent with the finding from Carless 2006 in finding no significant impact of cell salvage on blood loss.	D	Evidence is inconsistent				
maing from Cariess 2000 in finding no significant impact of cell salvage on blood loss.	NA	Not applicable (one study only)				
3. Clinical impact						
Carless (2006)	Α	Very large				
All surgery types – mean difference -108.47 mL (-407.53, 190.58); 6 RCTs (N=431) Cardiac surgery – mean difference 27.17 mL (-102.74, 157.08); 2 RCTs (N=206)	В	Substantial				
Orthopaedic surgery – mean difference -736.00 mL (-1054.00, -418.00); 1 RCT (N=39) Vascular surgery – mean difference 34.62 mL (-268.98, 338.21); 3 trials (N=186)	С	Moderate				
	D	Slight				
4. Generalisability						
The results are generalisable for patients undergoing elective surgery.	Α	Evidence directly generalisable to target population				
	В	Evidence directly generalisable to target population with some care	veats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard	to judge whether it is sensible to			
5. Applicability						
Orthopaedic One study was in China and the other in Europe	Α	Evidence directly applicable to Australian healthcare context				
Other surgery	В	Evidence applicable to Australian healthcare context with few caveats				
All the studies were conducted in developed countries.	С	Evidence probably applicable to Australian healthcare context wit	h some caveats			
	D	Evidence not applicable to Australian healthcare context				

There is heterogeneity in the blood loss measures. Intuitively, cell salvage will not be expected to be a determinant of blood loss.

Damgaard 2006 only reported net blood loss.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	Two level II studies with moderate risk of bias in orthopaedic surgery. Several level II studies with low risk of bias for other surgery types.
2. Consistency	С	Some inconsistency, reflecting genuine uncertainty around question
3. Clinical impact	D	Moderate impact in orthopaedic surgery. No difference in other surgery
4. Generalisability	В	Evidence directly generalisable to target population with some caveats
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats (one orthopaedic study conducted in China)

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intraoperative cell salvage on operative blood loss is uncertain.

Abbreviations: CABG, coronary artery bypass graft; CPB, cardiopulmonary bypass; RCT, randomised controlled trial; SR, systematic review.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hosp

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I2.P3 Characteristics and results of studies examining the effect of intraoperative cell salvage on blood loss.

	Level of evidence	No. of trials /	Patient population / Surgical			_		Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Carless (2006)	Level I Good	6 trials (1 poor quality, 5 fair quality) N=431	Adults undergoing any type of surgery.	All studies were conducted in developed countries.	Intraoperative cell salvage	Mean difference (95% CI), mL	-108.47 (-407.53, 1	190.58)	P=0.48	Phet=0.001
Damgaard (2006)	Level II Good	N=60	Adults undergoing sub-acute coronary bypass surgery without heart-lung machine.	Hospital in Denmark	The continuously heparinised suction and reservoir belonging to the cell saver were used for all patients in both groups. The suctioned blood from patients in the cell saver group was processed and autotransfused before the patient was transferred to the ICU.	Median net blood loss (IQR), mL	300 (193 to 403)	610 (450 to 928)	P<0.001	
Mercer (2004)	Level II Good	N=81	Adults undergoing surgery for abdominal aortic aneurysm.	Hospital in UK	Intraoperative cell salvage. Processed blood was returned to the patient as soon as haemostasis had been achieved.	Median (IQR) blood loss, mL	1950 (775 to 285)	1270 (775 to 2850)	P=0.140	
Niranjan (2006)	Level II Good	N=80	Adults undergoing first-time, isolated CABG	Hospital in UK	Intraoperative cell salvage, with Autotransfusion of washed, salvaged red blood cells at the conclusion of the procedure.	Mean (SD) blood loss, mL	On-CPB: 842 (276) Off-CPB: 869 (286)	On-CPB: 1023 (291) Off-CPB: 903 (315)	On-CPB: P=0.04 Off-CPB: P=0.72	
Zhang (2004)	Level II Poor	N=48	Adults undergoing operation for scoliosis.	Hospital in China	Intraoperative cell salvage and retransfusion of washed autologous blood.	Mean (SD) blood loss	NR	NR	P=NS	
Cardiac surgery										
Carless (2006)	Level I Good	2 trials (fair quality) N=206	Adults undergoing any type of surgery.	All studies were conducted in developed countries.	Intraoperative cell salvage	Mean difference (95% CI), mL	27.17 (-102.74, 157	7.08)	P=0.68	Phet=0.96
Orthopaedic sur	gery									
Carless (2006)	Level I Good	1 trial (fair quality) N=39	Adults undergoing any type of surgery.	All studies were conducted in developed countries.	Intraoperative cell salvage	Mean difference (95% CI), mL	-736.00 (-1054.00,	-418.00)	P<0.00001	Phet=NA

	Level of evidence	No. of trials /	Patient population / Surgical			Outcome		Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention		Intervention	Comparator	p-value	
Vascular surgery	Vascular surgery									
Carless (2006)	Level I Good	3 trials (1 poor quality, 2 fair quality) N=186	Adults undergoing any type of surgery.	All studies were conducted in developed countries.	Intraoperative cell salvage	Mean difference (95% CI), mL	34.62 (-268.98, 338.21)		P=0.82	Phet=0.83

Abbreviations: CABG, coronary artery bypass graft; CI, confidence interval; ICU, intensive care unit; IQR, interquartile range; NR, not reported; NS, not significant; SD, standard deviation.

Key question(s): In patients undergoing surgery, what is the effect of intraoperative cell salvag	<u>e</u> on	mortality?	Evidence table ref*: POQ3.I2.P4
1. Evidence base			
Pivotal evidence: 1 level 1 SR (Carless 2006); good quality; adults undergoing any type of surgery; includes 3	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias
RCTs (N=186): all vascular (1 poor quality, 2 fair quality) Supportive published after Carless 2006 from 7 level II studies: Bowley 2006 (trauma; fair quality; N=44);	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias
Damgaard 2006 (cardiac; good quality; N=60); Goel 2007 (cardiac; fair quality; N=50); Mercer 2004 (vascular; good quality; N=81); Murphy 2005 (cardiac; fair quality; N=61); Selo-Ojeme 2007 (ruptured ectopic pregnancy;	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias
fair quality; N=112); Zhang 2004 (orthopaedic; poor quality; N=48)	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	3
2. Consistency			
All the studies are consistent in finding no significant association between intraoperative cell salvage mortality.	Α	All studies consistent	
	В	Most studies consistent and inconsistency can be explained	
	С	Some inconsistency, reflecting genuine uncertainty around questi	on
	D	Evidence is inconsistent	
	NA	Not applicable (one study only)	
3. Clinical impact			
Meta-analysis of Carless 2006 and the 7 RCTs was conducted	Α	Very large	
All surgery types – RR 1.01 (0.68, 1.50); 10 RCTs (N=641) Cardiac surgery – RR 0.20 (0.01, 4.00); 3 RCTs (N=170)	В	Substantial	
Trauma surgery – RR 1.02 (0.67, 1.56); 1 RCT (N=29)	С	Moderate	
Vascular surgery – RR 1.16 (0.33, 4.09); 4 RCTs (N=267)	D	No difference/underpowered	
4. Generalisability			
Both elective (cardiac, orthopaedic, and vascular) surgery and surgery for traumatic injury are included in the review. The lack of effect of intraoperative cell salvage on mortality is not dependent on surgery type.	Α	Evidence directly generalisable to target population	
Teview. The lack of effect of intraoperative cell salvage on mortality is not dependent of surgery type.	В	Evidence directly generalisable to target population with some call	/eats
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to
5. Applicability			
Most of the studies were conducted in developed countries. Goel 2007 was conducted in India, Selo-Ojeme 2007 in Nigeria, and Zhang 2004 in China. The exclusion of these studies does not impact the results.	Α	Evidence directly applicable to Australian healthcare context	
in ringona, and zhang 2004 in Ohina. The exclusion of these studies does not impact the results.	В	Evidence applicable to Australian healthcare context with few cave	
	С	Evidence probably applicable to Australian healthcare context with	n some caveats
	D	Evidence not applicable to Australian healthcare context	

Included studies were underpowered to detect a mortality difference. The high rates of mortality in Bowley 2006 was due to the study population (patients undergoing surgery for penetrating torso injury)

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	В	Two good quality level II studies with a low risk of bias and several other level II studies with moderate risk of bias
2. Consistency	Α	All studies consistent because studies are underpowered
3. Clinical impact	D	There is no statistically significant association between intraoperative cell salvage and mortality
4. Generalisability	В	Evidence directly generalisable to target population with some caveats
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intraoperative cell salvage on mortality is uncertain.

Abbreviations: RCT, randomised controlled trial; RR, relative risk; SR, systematic review.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I2.P4 Characteristics and results of studies examining the effect of intraoperative cell salvage on mortality.

0	Level of evidence	No. of trials /	Patient population / Surgical	0.111				Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Carless (2006)	Level I Good	3 trials (1 poor quality, 2 fair quality) N=186	Adults undergoing any type of surgery. ¹	All studies were conducted in developed countries.	Intraoperative cell salvage		4/93 (4%)	4/93 (4%)	P=0.93	Phet=0.18
Bowley (2006)	Level II Fair	N=44	Adults with penetrating torso injury requiring a laparotomy and who had exhibited hypotension either pre-hospital or an arrival and in whom there was considered to be a significant loss.	Hospital in Johannesburg	Intraoperative blood salvage with transfusion of both allogeneic and washed autologous blood.		14/21 (67%)	15/23 (65%)	P=0.92	
Damgaard (2006)	Level II Good	N=60	Adults undergoing sub-acute coronary bypass surgery without heart-lung machine.	Hospital in Denmark	The continuously heparinised suction and reservoir belonging to the cell saver were used for all patients in both groups. The suctioned blood from patients in the cell saver group was processed and autotransfused before the patient was transferred to the ICU.		0/30 (0%)	2/30 (7%)	P=0.29	
Goel (2007)	Level II Fair	N=50	Adults undergoing off-pump coronary artery bypass grafting.	Hospital in India	Intraoperative cell salvage and autotransfusion of washed shed blood and transfusion of allogeneic blood if required.		0/24 (0%)	0/25 (0%)	Not estimable	
Mercer (2004)	Level II Good	N=81	Adults undergoing surgery for abdominal aortic aneurysm.	Hospital in UK	Intraoperative cell salvage. Processed blood was returned to the patient as soon as haemostasis had been achieved.		1/40 (3%)	1/41 (2%)	P=1.000	
Murphy (2005)	Level II Fair	N=61	Patients scheduled for non- emergency first-time CABG (off-pump)	Hospital in UK	Intraoperative cell salvage, with Autotransfusion of washed, salvaged red blood cells at the completion of the operative procedure. Salvaged washed RBCs were autotransfused at the time of skin closure.		0/30 (0%)	0/31 (0%)	Not estimable	

Study	Level of evidence	No. of trials / sample size	Patient population / Surgical procedure	Setting	Intervention	Outcome		Notes		
	Quality						Intervention	Comparator	p-value	
Selo-Ojeme (2007)	Level II Fair	N=112	Women undergoing surgery for ruptured ectopic pregnancy	Hospital in Nigeria	Intraoperative cell salvage with transfusion of filtered autologous blood.		0/56 (0%)	0/56 (0%)	Not estimable	
Zhang (2004)	Level II Poor	N=48	Adults undergoing operation for scoliosis.	Hospital in China	Intraoperative cell salvage and retransfusion of washed autologous blood.		0/36 (0%)	0/12 (0%)	Not estimable	

Abbreviations: CABG, coronary artery bypass graft; ICU, intensive care unit; RBCs, red blood cells.

¹ All in vascular surgery

Key question(s): n patients undergoing surgery, what is the effect of <u>intraoperative cell salvag</u>	ıo on	morhidity?	Evidence table ref*: POQ3.12.P5			
1. Evidence base	<u>e</u> on	inorpiaity:	1 0 20.12.1 0			
Pivotal evidence: 1 level 1 SR (Carless 2006); good quality. Infection, 2 trials (1 cardiac, 1 vascular, both fair quality; N=268); wound complication, 1 trial (vascular, fair	А	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias			
quality; N=100); any thrombosis, 2 trials (1 orthopaedic, 1 vascular, both fair quality; N=139); stroke, 2 trials (1 cardiac, 1 vascular, both fair quality; N=268); non-fatal MI, 3 trials (2 in vascular, 1 poor and 1 fair quality; 1 in	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias			
cardiac; N=304); DVT, 1 trial (orthopaedic, fair quality N=39) Supportive evidence published after Carless 2006 from 7 level II studies: Damgaard 2006 (cardiac; good quality; N=60); Goel 2007 (cardiac, fair quality; N=50); Mercer 2004 (vascular, good quality; N=81); Murphy 2004	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
(vascular, fair quality; N=61); Niranjan 2006 (vascular, good quality; N=80); Selo-Ojeme 2007 (ruptured ectopic pregnancy, fair quality; N=112); Zhang 2004 (orthopaedic, poor quality; N=48)	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency						
There was no association between intraoperative cell salvage and any reported adverse events in either Carless	Α	All studies consistent				
 2006 or the subsequent 7 RCTs with the following exceptions: In Mercer 2004 there was a significantly lower rate of infection and SIRS in the cell salvage group. 	В	Most studies consistent and inconsistency can be explained				
 In Zhang 2004 there was a significantly lower rate of allergic reactions in the cell salvage group. 	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
Infection in vascular surgery Carless 2006 reported no significant difference (10% vs 20%; P>0.05; N=100). In Mercer 2004 cell salvage was	D	Evidence is inconsistent				
associated with a significant reduction in infection (13% vs 34%; P=0.03; N=81)	NA	Not applicable (one study only)				
3. Clinical impact						
Infection (a meta–analysis was conducted combining the results from Carless 2006 with those from Selo-Ojeme	Α	Very large				
2007, Damgaard 2006, Goel 2007, Murphy 2005, and Mercer 2004) Vascular - RR 0.42 (0.21, 0.83); 2 trials (N=181); Cardiac - RR 1.40 (0.62, 3.13); 4 trials (N=338); Trauma - RR	В	Substantial				
0.75 (0.18, 3.20); 1 trial (N=112)	С	Moderate				
<u>SIRS</u> (Mercer 2004; N=81) – RR 0.46 (0.24, 0.89) <u>Allergic reaction</u> (Zhang 2004; N=48) – RR 0.05 (0.00, 0.91)	D	Slight/Restricted				
4. Generalisability						
The results are generalisable for elective, non-emergency surgery.	Α	Evidence directly generalisable to target population				
	В	Evidence directly generalisable to target population with some cav	reats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to			
5. Applicability						
Most of the studies were conducted in developed countries. Goel 2007 was conducted in India, Selo-Ojeme 2007	Α	Evidence directly applicable to Australian healthcare context				
in Nigeria, and Zhang 2004 in China. The exclusion of these studies does not impact the results.	В	Evidence applicable to Australian healthcare context with few cave	eats			
	С	Evidence probably applicable to Australian healthcare context with	n some caveats			

Zhang 2004 studies T-cell sets etc, and so was not considered relevant. The CRG noted that the definitions of infection in Mercer 2004 were not clear (and therefore it was not appropriate to make an evidence statement for this outcome).

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

	<u> </u>	
Component	Rating	Description
1. Evidence base	В	One level I study and several level II studies with low risk of bias
2. Consistency	В	Most studies consistent and inconsistency can be explained
3. Clinical impact		Cell salvage is associated with a moderate reduction in the risk of infection. No statistically significant association was found between intraoperative cell salvage and other morbidity outcomes
4. Generalisability	В	Evidence directly generalisable to target population with some caveats
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intraoperative cell salvage on morbidity is uncertain.

Abbreviations: DVT, deep vein thrombosis; MI, myocardial infarction; RCT, randomised controlled trial; RR, relative risk; SIRS, systemic inflammatory response syndrome; SR, systematic review.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I2.P5 Characteristics and results of studies examining the effect of intraoperative cell salvage on morbidity.

	Level of evidence	No. of trials /	Patient population / Surgical			_		Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Carless (2006)	Level I Good	2 trials (fair quality) N=268	Adults undergoing any type of surgery.	All studies were conducted in	Intraoperative cell salvage	Infection	16/134 (12%)	17/134 (13%)	P=0.86	Phet=0.09
		1 trial (fair quality) N=100		developed countries.		Wound complication	3/50 (6%)	3/50 (6%)	P=1.00	Phet=NA
		2 trials (fair quality) N=139				Any thrombosis	3/69 (4%)	2/70 (3%)	P=0.59	Phet=NA
		2 trials (fair quality) N=268				Stroke	1/134 (1%)	3/134 (2%)	P=0.39	Phet=0.84
	3 trials (1 poor, 2 fair quality) N=304 1 trial (fair quality)		Non fatal MI	5/152 (3%)	13/152 (9%)	P=0.09	Phet=0.84			
	1 trial (fair quality) N=39				DVT	3/19 (16%)	2/20 (10%)	P=0.59	Phet=NA	
Damgaard Level II	Level II	N=60	Adults undergoing sub-acute		The continuously heparinised suction and reservoir belonging to the cell saver were used for all patients in both groups. The suctioned blood from patients in the cell saver group was processed and autotransfused before the patient was transferred to the ICU.	Stroke	0/30 (0%)	1/30 (3%)	P=0.50	
(2006) Good	Good		coronary bypass surgery without heart-lung machine.			MI	0/30 (0%)	1/30 (3%)	P=0.50	
			Without Hourt larg machine.			Pneumonia	2/30 (7%)	3/30 (10%)	P=0.64	
						GI bleeding	0/30 (0%)	3/30 (10%)	P=0.19	
						Deep sterna wound infection	0/30 (0%)	1/30 (3%)	P=0.50	
						Leg wound infection	0/30 (0%)	1/30 (3%)	P=0.50	
						Dialysis	1/30 (3%)	2/30 (7%)	P=0.56	
						Low cardiac output syndrome	0/30 (0%)	6/30 (20%)	P=0.08	
						Atrial arrhythmia	14/30 (47%)	20/30 (67%)	P=0.13	
						Ventricular arrhythmia	0/30 (0%)	3/30 (10%)	P=0.19	
Goel (2007)	Level II Fair	N=50	Adults undergoing off-pump coronary artery bypass grafting.	Hospital in India	Intraoperative cell salvage and autotransfusion of washed shed blood and transfusion of allogeneic blood if required.	Deep sterna wound infection	0/24 (0%)	0/25 (0%)	Not estimable	

0	Level of evidence	No. of trials /	Patient population / Surgical	0.111				Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Mercer (2004)	Level II	N=81	Adults undergoing surgery for	Hospital in UK	Intraoperative cell salvage.	Infection	5/40 (13%)	14/41 (34%)	P=0.03	
	Good		abdominal aortic aneurysm.		Processed blood was returned to the patient as	Sepsis	4/40 (10%)	8/41 (20%)	P=0.49	
					soon as haemostasis had been achieved.	SIRS	9/40 (23%)	20/41 (49%)	P=0.02	
Murphy (2005)	Level II Fair	N=61	Patients scheduled for non- emergency first-time CABG	Hospital in UK	Intraoperative cell salvage, with Autotransfusion of	Stroke	0/30 (0%)	0/31 (0%)	Not estimable	
			(off-pump)		washed, salvaged red blood cells at the completion of the	MI	2/30 (7%)	0/31 (0%)	P=0.28	
					operative procedure. Salvaged washed RBCs	Pulmonary complications	0/30 (0%)	4/31 (13%)	P=0.11	
					were autotransfused at the time of skin closure.	Infection	2/30 (7%)	1/31 (3%)	P=0.54	
				time of skin closure.	Renal complications	0/30 (0%)	2/31 (7%)	P=0.49		
					Arrhythmia	6/30 (20%)	7/31 (23%)	P=0.81		
Niranjan (2006) Level II Good	Level II Good	N=80	Adults undergoing first-time, isolated CABG	Hospital in UK	Intraoperative cell salvage, with Autotransfusion of washed, salvaged red blood cells at the conclusion of the procedure.	Cardiovascular accident	On-CPB: 0/20 (0%) Off-CPB: 1/20 (5%)	On-CPB: 1/20 (5%) Off-CPB: 0/20 (0%)	On-CPB: P=0.49 Off-CPB: P=0.49	
					procedure.	pulmonary complications	On-CPB: 4/20 (20%) Off-CPB: 2/20 (10%)	On-CPB: 3/20 (15%) Off-CPB: 1/20 (5%)	On-CPB: P=0.68 Off-CPB: P=0.56	
						Renal complications	On-CPB: 2/20 (10%) Off-CPB: 1/20 (5%)	On-CPB: 1/20 (5%) Off-CPB: 0/20 (0%)	On-CPB: P=0.56 Off-CPB: P=0.49	
						Arrhythmia	On-CPB: 7/20 (35%) Off-CPB: 3/20 (25%)	On-CPB: 5/20 (25%) Off-CPB: 4/20 (20%)	On-CPB: P=0.49 Off-CPB: P=0.68	
Selo-Ojeme	Level II		Intraoperative cell salvage	Infection	3/56 (5%)	4/56 (7%)	P=0.70			
2007)	Fair		ruptured ectopic pregnancy		with transfusion of filtered	Postoperative fever	20/56 (36%)	21/56 (38%)	P=0.84	
Zhang (2004)	Level II Poor	N=48	Adults undergoing operation for scoliosis.	Hospital in China	Intraoperative cell salvage and retransfusion of washed autologous blood.	Allergic reaction	0/36 (0%)	3/12 (25%)	P=0.04	

	Level of evidence	No. of trials /	Patient population / Surgical					Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Cardiac surgery										
Carless (2006)	Level I Good	1 trial (fair quality) N=168	Adults undergoing any type of surgery.	All studies were conducted in	Intraoperative cell salvage	Infection	11/84 (13%)	7/84 (8%)	P=0.32	Phet=NA
		1 trial (fair quality) N=168		developed countries.		Non fatal MI	5/84 (6%)	10/84 (12%)	P=0.19	Phet=NA
Orthopaedic sur	gery									
Carless (2006)	Level I Good	1 trial (fair quality) N=39	Adults undergoing any type of surgery.	All studies were conducted in	Intraoperative cell salvage	Any thrombosis	3/19 (16%)	2/20 (10%)	P=0.59	Phet=NA
		1 trial (fair quality) N=168	developed countries.		Stroke	1/84 (1%)	2/84 (2%)	P=0.57	Phet=NA	
Vascular surger	у									
Carless (2006)	Level I Good	` ' ' ' '	Adults undergoing any type of surgery.	All studies were conducted in	Intraoperative cell salvage	Infection	5/50 (10%)	10/50 (20%)	P=0.17	Phet=NA
		1 trial (fair quality) N=100		developed countries.		Wound complication	3/50 (6%)	3/50 (6%)	P=1.00	Phet=NA
		1 trial (fair quality) N=100				Any thrombosis	0/50 (0%)	0/50 (0%)	Not estimable	Phet=NA
		1 trial (fair quality) N=100				Stroke	0/50 (0%)	1/50 (2%)	P=0.50	Phet=NA
Abbreviotione, C		2 trials (1 poor, 1 fair quality) N=136)			Non fatal MI	0/68 (0%)	3/68 (4%)	P=0.22	Phet=0.84

Abbreviations: CABG, coronary artery bypass graft; CPB, cardiopulmonary bypass; DVT; deep vein thrombosis; GI, gastrointestinal; ICU, intensive care unit; NA, not applicable; MI, myocardial infarction; RBCs, red blood cells; SIRS; systemic inflammatory response syndrome.

Key question(s): In patients undergoing surgery, what is the effect of intraoperative cell salvage	<u>e</u> on	quality of life?	Evidence table ref*: POQ3.I2.P6	
1. Evidence base				
No evidence found	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias	
	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias	
	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias	
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias		
2. Consistency		L		
	Α	All studies consistent		
	В	Most studies consistent and inconsistency can be explained		
	С	Some inconsistency, reflecting genuine uncertainty around question	on	
	D	Evidence is inconsistent		
	NA	Not applicable (one study only)		
3. Clinical impact				
	Α	Very large		
	В	Substantial		
	С	Moderate		
	D	Slight/Restricted		
4. Generalisability				
	Α	Evidence directly generalisable to target population		
	В	Evidence directly generalisable to target population with some cav	eats	
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied	
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to	
5. Applicability		[
	A	Evidence directly applicable to Australian healthcare context		
	В	Evidence applicable to Australian healthcare context with few cave		
	С	Evidence probably applicable to Australian healthcare context with	some caveats	
	D	Evidence not applicable to Australian healthcare context		

Other factors		
EVIDENCE STA	TEMENT	MATRIX
Please summarise	the develor	oment group's synthesis of the evidence relating to the key question, taking all the above factors into account.
Component	Rating	Description
1. Evidence base	NA	
2. Consistency	NA	
3. Clinical impact	NA	
4. Generalisability	NA	
5. Applicability	NA	
DRAFT EVIDEN	ICE STA	ΓΕΜΕΝΤ
Daniel on the back	-£ : -	
Based on the body	or evidence	above.
In adult p	atients un	dergoing surgery in which substantial blood loss is anticipated, the effect of intraoperative cell salvage on quality of life is unknown.
		ative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for

coagulation status and haemoglobin, 18 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, 19 = appropriate patient positioning, 110 = PAD

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission

Key question(s):	has madahin as nontration?	Evidence table ref*: POQ3.I2.S1			
In patients undergoing surgery, what is the effect of intraoperative cell salvag	<u>e</u> 011	<u>Itaernogiobili concentration?</u>	1 003.12.31		
1. Evidence base	Ι.	To			
Five Level II studies: Damgaard 2006 (cardiac; good quality; N=60); Goel 2007 (cardiac; fair quality; N=50); Murphy 2005 (cardiac; fair quality; N=61); Niranjan 2006 (cardiac; good quality; N=80); Selo-Ojeme 2007	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias		
(ruptured ectopic pregnancy; fair quality; N=112) Murphy 2005 reports the postoperative Hb concentration but not the preoperative concentration. Goel 2007 and	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias		
Niranjan 2006 report the change in Hb concentration from preoperative to postoperative. Damgaard 2006 reports	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias		
preoperative and postoperative Hb concentration values. Murphy 2005 and Selo-Ojeme 2007 report the postoperative, but not the preoperative, concentration of haematocrit in the cell salvage and control groups. Damgaard 2006 reports both pre and postoperative haematocrit concentration values.	D	Level IV studies or Level I to III studies/SRs with a high risk of bias			
2. Consistency					
Haemoglobin concentration: Goel 2007 reported a lesser decrease in Hb from pre- to postoperative levels, for cell salvage compared with control but Niranjan 2006 found no significant difference.	Α	All studies consistent			
Murphy 2005 found that the cell salvage group had significantly higher Hb concentration at 24 hours	В	Most studies consistent and inconsistency can be explained			
postoperative compared with control. Damgaard 2006 found no significant difference in Hb concentration either pre- or postoperatively.	С	Some inconsistency, reflecting genuine uncertainty around question	on		
Haematocrit concentration: Damgaard 2006 found no significant difference in haematocrit concentration between the cell salvage and control groups either pre- or postoperatively. Murphy 2005 and Selo-Ojeme 2007	D	Evidence is inconsistent			
found that the haematocrit concentration was significantly greater in the cell salvage group 24 hours postoperative and immediately postoperative respectively.	NA	Not applicable (one study only)			
3. Clinical impact					
Goel 2007 (cell salvage vs. control; mean difference in the decrease in Hb concentration from preoperative to immediately postoperative; g/dL) – mean difference -0.90 (-1.68, -0.12).	Α	Very large			
Niranjan 2006 (cell salvage vs. control; mean difference in the Hb concentration from preoperative to 24 hours	В	Substantial			
postoperative; g/dL): On-CPB – mean difference 0.55 (-0.07, 1.17), Off-CPB – mean difference -0.05 (-1.01, 0.91).	С	Moderate			
<u>Damgaard 2006</u> found no significant difference in Hb concentration between the cell salvage and the control groups either pre- or postoperatively. ^a <u>Murphy 2005</u> found that the cell salvage group had significantly higher Hb concentration at 24 hours postoperative compared with control. ^b See Summary Table POQ3.I2.S1 for haematocrit concentration results. ^c	D	Slight/Restricted			
4. Generalisability					
Damgaard 2006, Goel 2007, and Murphy 2005 were conducted in patients undergoing off-CPB CABG. Niranjan	Α	Evidence directly generalisable to target population			
2006 was conducted in patients undergoing on- and off-CPB CABG. Selo-Ojeme 2007 was assessed cell salvage in women with ruptured ectopic pregnancy.	В	Evidence directly generalisable to target population with some cav	eats		
in women with raptarea ectopic pregnancy.	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied		
	D	Evidence not directly generalisable to target population & hard to j	udge whether it is sensible to apply		
5. Applicability					
Murphy 2005 and Niranjan 2006 were conducted in UK, Damgaard 2006 in Denmark, Goel 2007 in India, and Selo-Ojeme 2007 in Nigeria.	Α	Evidence directly applicable to Australian healthcare context			
Selo-Ojenie 2007 in Nigeria.	В	Evidence applicable to Australian healthcare context with few cave	ats		
	С	Evidence probably applicable to Australian healthcare context with	some caveats		
	D	Evidence not applicable to Australian healthcare context			

The results were not able to be meta-analysed because of insufficient details in the reported data and differences in the measurement timing of haemoglobin concentration.

The CRG based their decision making on the results from Goel 2007 and Murphy 2005, which reported the most relevant timepoints.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base		Five fair-to-good quality level II studies reported haemoglobin/haematocrit concentration as a clinical outcome, but only three level II studies (one fair, two good quality) reported both pre- and postoperative haemoglobin concentration values.
2. Consistency	С	Some inconsistency, reflecting genuine uncertainty around question.
3. Clinical impact	D	Some of the trials found that intraoperative cell salvage had a moderate impact and some found no impact.
4. Generalisability	В	Evidence directly generalisable to target population with some caveats
5. Applicability	В	Evidence probably applicable to Australian healthcare context with some caveats.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing off-pump coronary artery surgery, intraoperative cell salvage may increase postoperative haemoglobin concentration and haematocrit.

Abbreviations: CABG, coronary artery bypass graft; CPB, cardiopulmonary bypass; Hb, haemoglobin; RR, relative risk.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission

Results from Murphy 2005 (mean difference [g/dL]): immediately postoperative, 0.00 (-0.02, 0.02); 1 hour postoperative, 0.01 (-0.01, 0.02); 24 hours postoperative, 0.03 (0.01, 0.05).

Results from Selo-Ojeme: mean difference not reported, see Summary Table POQ3.12.S1 for median [IQR] values and P-values.

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

^a Baseline mean difference not reported. See Summary Table P0Q3.12.S1 for median [IQR] values and P-values.

^b Results from Murphy 2005 (mean difference [q/dL]): immediately postoperative, -0.11 (-0.69, 0.47); 1 hour postoperative, 0.15 (-0.42, 0.72); 24 hours postoperative, 1.02 (0.45, 1.59).

^c Results from Damgaard 2006: mean difference not reported, see Summary Table P0Q3.I2.S1 for median [IQR] values and P-values.

POQ3.I2.S1 Characteristics and results of studies examining the effect of intraoperative cell salvage on haemoglobin concentration.

0	Level of evidence	ce No. of trials / Sample size Patient population / Surgical Setting Intervention				Results			Notes			
Study	Quality		Intervention	Outcome	Intervention	Comparator	p-value					
Damgaard (2006)	Level II Good	N=60	Adults undergoing sub-acute coronary bypass surgery without heart-lung machine.	Hospital in Denmark	The continuously heparinised suction and reservoir belonging to the cell saver were used for all	Baseline Hb concentration (median [IQR]), mmol/L	7.9 (7.4 to 8.7)	8.2 (7.4 to 8.9)	P=0.43			
			patients in the cell saver	Lowest Hb concentration in ICU (median [IQR]), mmol/L	5.9 (5.3 to 6.6)	5.8 (5.2 to 6.7)	P=0.97					
			patie		patient was transferred to the ICU.	patient was transferred to	patient was transferred to the ICU.	Lowest Hb concentration in ward (median [IQR]), mmol/L	6.4 (5.9 to 6.8)	6.6 (5.8 to 7.1)	P=0.58	
								Lowest Hb concentration at hospital discharge (median [IQR]), mmol/L	7.1 (6.5 to 7.4)	7.2 (6.5 to 8.1)	P=0.25	
						Baseline haematocrit concentration (median [IQR]), %	39 (36 to 42)	41 (38 to 44)	P=0.21			
			Lowest haematocrit concentration in ICU (median [IQR]), mmol/L	29 (27 to 33)	29 (25 to 33)	P=0.69						
Goel (2007)	Level II Fair	N=50	Adults undergoing off-pump coronary artery bypass grafting.	Hospital in India	Intraoperative cell salvage and autotransfusion of washed shed blood and transfusion of allogeneic blood if required.	Decrease in Hb (from preoperative to immediately postoperative) (mean [SD]), g/dL	1.8 (1.2)	2.7 (1.6)	P=0.02			

Level of evidence		No. of trials /	Patient population / Surgical				Results			Notes	
Study evidence ample size ample s	sample size	sample size procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value			
Murphy (2005)	Fair emergency first-time CABG with autotransfusion of washed, salvaged red by cells at the completion of operative procedure.	washed, salvaged red blood cells at the completion of the operative procedure. Salvaged washed RBCs	Hb concentration immediately postoperative (mean [SD]), g/dL	11.14 (1.15)	11.25 (1.17)	P=0.71					
		were autotransfused at the time of skin closure.	Hb concentration 1 hour postoperative (mean [SD]), g/dL	10.55 (1.15)	10.40 (1.11)	P=0.60					
						Hb concentration 24 hours postoperative (mean [SD]), g/dL	11.71 (1.15)	10.69 (1.11)	P=0.0007		
								Haematocrit concentration immediately postoperative (mean [SD]), %	0.345 (0.033)	0.344 (0.033)	P=0.91
						Haematocrit concentration 1 hour postoperative (mean [SD]), %	0.312 (0.033)	0.305 (0.033)	P=0.46		
				Haematocrit concentration 24 hour postoperative (mean [SD]), %	0.350 (0.033)	0.319 (0.033)	P=0.0008				
Niranjan (2006)	Level II Good	N=80	Adults undergoing first-time, isolated CABG	Hospital in UK	Intraoperative cell salvage, with Autotransfusion of washed, salvaged red blood cells at the conclusion of the procedure.	Decrease in Hb (from preoperative to 24 hours postoperative) (mean [SD]), g/dL	On-CPB: 4.95 (1.1) Off-CPB: 4.95 (1.5)	On-CPB: 4.4 (0.9) Off-CPB: 5.0 (1.6)	On-CPB: P=0.08 Off-CPB: P=0.92		

0	Level of evidence	No. of trials /	Patient population / Surgical	Setting Intervention Outcome	Results	Results		Notes		
Study evidence Quality		sample size	procedure		Intervention	Outcome	Intervention	Comparator	p-value	
Selo-Ojeme (2007)	Level II Fair	N=112	Women undergoing surgery for ruptured ectopic pregnancy	Hospital in Nigeria	Intraoperative cell salvage with transfusion of filtered autologous blood.	Haematocrit concentration immediately postoperative, %	29	26	P<0.01	

Abbreviations: CABG, coronary artery bypass graft; CPB, cardiopulmonary bypass; ICU, intensive care unit; IQR, interquartile range; HB, haemoglobin; SD, standard deviation.

Cey question(s): n patients undergoing surgery, what is the effect of <u>intraoperative cell salva</u>	ge on	the rate of reoperation for bleeding?	Evidence table ref*: POQ3.12.S2			
1. Evidence base						
1 SR (Carless 2006); good quality; includes 2 RCTs (N=218): 1 cardiac (fair quality), 1 vascular (fair quality)	А	One or more level I studies with a low risk of bias or seve	everal level II studies with a low risk of bias			
2 level II studies: Damgaard 2006 (cardiac; good quality; N=60); Goel 2007 (cardiac; fair quality; N=50)	В	One or two Level II studies with a low risk of bias or SR/s	/several Level III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Leve	el I or II studies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high ri	sk of bias			
2. Consistency						
All the studies are consistent in finding no significant association between intraoperative cell salvage and reoperation for bleeding.	Α	All studies consistent				
eoperation for bleeding.	В	Most studies consistent and inconsistency can be explain	ned			
	С	Some inconsistency, reflecting genuine uncertainty around question				
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact		,				
Carless 2006 – RR 0.57 (0.12, 2.63) Damgaard 2006 – RR 0.33 (0.04, 3.03)	А	Very large				
Goel 2007 – No patients in either treatment arms required reoperation for bleeding.	В	Substantial				
Meta-analysed results – RR 0.48 (0.13, 1.68)	С	Moderate				
	D	No difference/underpowered				
4. Generalisability						
One of the trials in Carless 2006 was in cardiac surgery, and the other in vascular surgery. Damgaard 2006	А	Evidence directly generalisable to target population				
assessed cell salvage in off-CPB CABG.	В	Evidence directly generalisable to target population with	some caveats			
	С	Evidence not directly generalisable to the target population	on but could be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to judge whether it is sensible to				
5. Applicability						
All the studies in Carless 2006 were conducted in developed countries, Damgaard 2006 was conducted in Denmark, and Goel 2007 was conducted in India. The exclusion of Goel 2007 does not impact the results.	А	Evidence directly applicable to Australian healthcare con	text			
Definitions, and Goet 2007 was conducted in India. The exclusion of Goet 2007 does not impact the results.	В	Evidence applicable to Australian healthcare context with	n few caveats			
	С	Evidence probably applicable to Australian healthcare co	ntext with some caveats			
	D	Evidence not applicable to Australian healthcare context				

Included studies were underpowered to detect a difference in reoperation for bleeding.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	One level I study including two RCTs and an additional two level II studies with moderate risk of bias published subsequently
2. Consistency	Α	All studies consistent
3. Clinical impact	D	No difference/underpowered
4. Generalisability	В	Evidence directly generalisable to target population with some caveats. Surgery types assessed are all cardiovascular.
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intraoperative cell salvage on risk of reoperation for bleeding is uncertain.

Abbreviations: CABG, coronary artery bypass graft; CPB, cardiopulmonary bypass; RCT, randomised controlled trial; RR, relative risk; SR, systematic review.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I2.S2 Characteristics and results of studies examining the effect of intraoperative cell salvage on reoperation for bleeding.

	Level of evidence	ence No. of trials / Patient population / Surgical Setting Intervention	Patient population / Surgical					Results		Notes
Study	Quality		Outcome	Intervention	Comparator	p-value				
Carless (2006)	Level I Good	2 trials (fair quality) N=218	Adults undergoing any type of surgery.	All studies were conducted in developed countries.	Intraoperative cell salvage		2/109 (2%)	4/109 (4%)	P=0.47	Phet=0.71
Damgaard (2006)	Level II Good	N=60	Adults undergoing sub-acute coronary bypass surgery without heart-lung machine.	Hospital in Denmark	The continuously heparinised suction and reservoir belonging to the cell saver were used for all patients in both groups. The suctioned blood from patients in the cell saver group was processed and autotransfused before the patient was transferred to the ICU.		1/30 (3%)	3/30 (10%)	P=0.35	
Goel (2007)	Level II Fair	N=50	Adults undergoing off-pump coronary artery bypass grafting.	Hospital in India	Intraoperative cell salvage and autotransfusion of washed shed blood and transfusion of allogeneic blood if required.		0/24 (0%)	0/25 (0%)	Not estimable	
Cardiac surgery										
Carless (2006)	Level I Good	1 trial (fair quality) N=168	Adults undergoing any type of surgery.	All studies were conducted in developed countries.	Intraoperative cell salvage		2/84 (2%)	3/84 (4%)	P=0.65	Phet=NA
Vascular surgery	/									
Carless (2006)	Level I Good	1 trial (fair quality) N=50	Adults undergoing any type of surgery.	All studies were conducted in developed countries.	Intraoperative cell salvage		0/25 (0%)	1/25 (4%)	P=0.49	Phet=NA

Abbreviations: ICU, intensive care unit; NA, not applicable.

Key question(s):		Evidence table ref*:				
In patients undergoing surgery, what is the effect of intraoperative cell salvag	<u>e</u> on	coagulation status?	POQ3.I2.S3			
1. Evidence base						
Two Level II studies: Murphy 2005 (cardiac; fair quality; N=61); Niranjan 2006 (cardiac; good quality; N=80)	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias			
Murphy 2005 reports platelet count, prothrombin ratio, APTT, and fibrinogen concentration. Niranjan 2006 only reports prothrombin time	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	3			
2. Consistency						
The studies were consistent in finding no significant impact.	Α	All studies consistent				
	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact						
Neither of the studies found any significant difference between cell salvage and control for any of the coagulation parameters.	Α	Very large				
See Summary Table POQ3.I2.S3	В	Substantial				
	С	Moderate				
	D	No difference				
4. Generalisability						
Both studies were conducted in patients undergoing first-time CABG.	Α	Evidence directly generalisable to target population				
	В	Evidence directly generalisable to target population with some cave	veats .			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to	to judge whether it is sensible to			
5. Applicability						
Both studies were conducted in the UK.	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few cave				
	С	Evidence probably applicable to Australian healthcare context with	h some caveats			
	D	Evidence not applicable to Australian healthcare context				

The reported platelet count in Murphy 2005 is not clinically significant. The reported standard deviations in Murphy 2005 may actually be standard errors.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description		
1. Evidence base	С	Level II studies with a low risk of bias		
2. Consistency	Α	All studies consistent		
3. Clinical impact	D	lifference		
4. Generalisability	Generalisability B Both studies were in adults undergoing first-time CABG			
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats (both studies conducted in UK)		

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing off-pump coronary artery surgery, the effect of intraoperative cell salvage on coagulation status is uncertain.

Abbreviations: APTT, activated partial thromboplastin time; CABG, coronary artery bypass graft.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I2.S3 Characteristics and results of studies examining the effect of intraoperative cell salvage on coagulation status.

	Level of evidence	No. of trials /	Patient population / Surgical			Outcomo		Results		Notes	
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value		
Murphy (2005)	Fair emergency first-time CABG (off-pump) with Autotrar washed, salv cells at the coperative pro Salvaged wa were autotra	Intraoperative cell salvage, with Autotransfusion of washed, salvaged red blood cells at the completion of the operative procedure.	Platelet count 1 hour postoperative (mean [SD]), X109/L	192.8 (0.15)	189.7 (0.14)	NR					
		were autotransfused at the time of skin closure.	Platelet count 24 hour postoperative (mean [SD]), X109/L	225.4 (0.15)	218.2 (0.14)	NR					
			im	Prothrombin ratio immediately postoperative (mean [SD])	1.27 (0.07)	1.27 (0.07)	NR				
						Prothrombin ratio 1 hour postoperative (mean [SD])	1.19 (0.06)	1.19 (0.06)	NR		
					Prothrombin ratio 24 hour postoperative (mean [SD])	1.15 (0.07)	1.15 (0.07)	NR			
			APTT ratio immediately postoperative (mean [SD])	1.17 (0.13)	1.14 (0.12)	NR					
						APTT ratio 1 hour postoperative (mean [SD])	1.08 (0.12)	1.13 (0.12)	NR		
								APTT ratio 24 hours postoperative (mean [SD])	1.08 (0.12)	1.11 (0.12)	NR

	Level of evidence	No. of trials /	Patient population / Surgical		lada mana Mana		Results			Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
						Fibrinogen concentration immediately postoperative (mean [SD]), g/L	2.59 (0.20)	2.68 (0.18)	NR	
						Fibrinogen concentration 1 hour postoperative (mean [SD]), g/L	2.21 (0.19)	2.34 (0.18)	NR	
						Fibrinogen concentration 24 hours postoperative (mean [SD]), g/L	4.92 (0.19)	5.04 (0.19)	NR	
Niranjan (2006)	Level II Good	N=80	Adults undergoing first-time, isolated CABG (on-and off-CBP)	Hospital in UK	Intraoperative cell salvage, with Autotransfusion of washed, salvaged red blood	Prothrombin time	NR	NR	NS	
		9.05			cells at the conclusion of the procedure.	Partial thromboplastin time (ratio)	NR	NR	NS	

Abbreviations: CABG, coronary artery bypass graft; CPB, cardiopulmonary bypass; NR, not reported; NS, not significant; RBC, red blood cells; SD, standard deviation.

Key question(s):						
In patients undergoing surgery, what is the effect of intraoperative cell salvag	<u>e</u> on	hospital length of stay?	POQ3.I2.S5			
1. Evidence base						
1 SR Careless (2006): good quality; includes 1 RCT (N=100): vascular (fair quality)	А	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias			
6 level II studies: Bowley 2006 (trauma; fair quality; N=44); Damgaard 2006 (cardiac; good quality; N=60); Mercer 2004 (vascular; good quality; N=81); Murphy 2005 (cardiac; fair quality; N=61); Niranjan 2006 (cardiac;	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias			
good quality; N=80); Selo-Ojeme 2007 (ruptured ectopic pregnancy; fair quality; N=112)	С	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	3			
2. Consistency						
All the studies are consistent in finding no significant association between intraoperative cell salvage and length of hospital stay.	Α	All studies consistent				
or nospital stay.	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact						
Carless 2006 – mean difference -0.50 (-2.46, 1.46); 1 trial (N=100) Bowley 2006 – mean difference 1.10 (-3.71, 5.91), N=44	Α	Very large				
Niranjan 2006 – mean difference (on-CPB) -0.20 (-1.82, 1.42); mean difference (off-CPB) -0.20 (-1.56, 1.16)	В	Substantial				
Selo-Ojeme 2007: Length of hospital stay > 7 days; RR 1.33 (0.49, 3.59)	С	Moderate				
	D	No difference				
4. Generalisability						
The studies were conducted in patients undergoing cardiovascular surgery, surgery for penetrating trauma, and surgery for ruptured ectopic pregnancy.	Α	Evidence directly generalisable to target population				
Surgery for replaced ectopic programoy.	В	Evidence directly generalisable to target population with some cave	veats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to			
5. Applicability	1					
Most of the studies were conducted in developed countries. Selo-Ojeme 2007 was conducted in Nigeria; however, the exclusion of this study does not impact the results.	Α	Evidence directly applicable to Australian healthcare context				
nowever, the exclusion of this study does not impact the results.	В	Evidence applicable to Australian healthcare context with few cave	eats			
	С	Evidence probably applicable to Australian healthcare context with some caveats				
	D	Evidence not applicable to Australian healthcare context				

Other factors

Length of stay is uncertain because of the range of surgeries.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	Several level II studies with low risk of bias
2. Consistency	Α	All studies consistent
3. Clinical impact	D	No difference
4. Generalisability		Evidence directly generalisable to target population with some caveats. Included studies assessed cardiovascular surgery, and surgery for trauma, and surgery for ruptured ectopic pregnancy.
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intraoperative cell salvage on length of hospital stay is uncertain.

Abbreviations: CPB, cardiopulmonary bypass; RCT, randomised controlled trial; RR, relative risk; SR, systematic review.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: 11 = ANH, 12 = intraoperative cell salvage, 13 = perioperative ANH and intraoperative cell salvage, 14 = postoperative cell salvage, 15 = deliberate induced hypotension, 16 = prevention of hypothermia, 17 = point-of-care testing for coagulation status and haemoglobin, 18 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, 19 = appropriate patient positioning, 110 = PAD

POQ3.I2.S5 Characteristics and results of studies examining the effect of intraoperative cell salvage on hospital length of stay.

	Level of evidence	No. of trials /	Patient population / Surgical	Catting		Outcomo		Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Carless (2006)	Level I Good	1 trial (fair quality) N=100	Adults undergoing any type of surgery.1	All studies were conducted in developed countries.	Intraoperative cell salvage	Mean (SD), days	12.2 (4.7)	12.7 (5.3)	P=0.62	
Bowley (2006)	Level II Fair	N=44	Adults with penetrating torso injury requiring a laparotomy and who had exhibited hypotension either pre-hospital or an arrival and in whom there was considered to be a significant loss.	Hospital in Johannesburg	Intraoperative blood salvage with transfusion of both allogeneic and washed autologous blood.	Mean (SD), days	15.7 (9.17)	14.6 (6.8)	P=0.65	
Damgaard (2006)	Level II Good	N=60	Adults undergoing sub-acute coronary bypass surgery without heart-lung machine.	Hospital in Denmark	The continuously heparinised suction and reservoir belonging to the cell saver were used for all patients in both groups. The suctioned blood from patients in the cell saver group was processed and autotransfused before the patient was transferred to the ICU.	Median (IQR)	7 (6 to 8)	7 (6 to 9)	P=NS	
Mercer (2004)	Level II Good	N=81	Adults undergoing surgery for abdominal aortic aneurysm.	Hospital in UK	Intraoperative cell salvage. Processed blood was returned to the patient as soon as haemostasis had been achieved.	Median (IQR)	12 (8 to 19)	13 (10 to 19)	P=0.385	
Murphy (2005)	Level II Fair	N=61	Patients scheduled for non- emergency first-time CABG (off-pump)	Hospital in UK	Intraoperative cell salvage, with Autotransfusion of washed, salvaged red blood cells at the completion of the operative procedure. Salvaged washed RBCs were autotransfused at the time of skin closure.	Median (IQR)	6.0 (5.0, 8.3)	6.0 (5.0, 8.0)	P=0.73	
Niranjan (2006)	Level II Good	N=80	Adults undergoing first-time, isolated CABG	Hospital in UK	Intraoperative cell salvage, with Autotransfusion of washed, salvaged red blood cells at the conclusion of the procedure.	Mean (SD), days	On-CPB: 8.1 (2) Off-CPB: 7.2 (2.3)	On-CPB: 8.3 (3.1) Off-CPB: 7.4 (2.1)	On-CPB: P=0.81 Off-CPB: P=0.77	

	Level of evidence	No. of trials /	Patient population / Surgical			Outcome		Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Selo-Ojeme (2007)	Level II Fair		Women undergoing surgery for ruptured ectopic pregnancy	Hospital in Nigeria	Intraoperative cell salvage with transfusion of filtered autologous blood.	Length of hospital stay > 7 days	8/56 (14%)	6/56 (11%)	P=0.57	

Abbreviations: CABG, coronary artery bypass graft; CPB, cardiopulmonary bypass; ICU, intensive care unit; IQR, interquartile range; RBC, red blood cells; SD, standard deviation.

¹ The study was conducted in patients undergoing vascular surgery.

Key question(s): In patients undergoing surgery, what is the effect of intraoperative cell salvag	ICU admission and length of stay?	Evidence table ref*: POQ3.I2.S6			
1. Evidence base					
Evidence for ICU readmission – 1 level II RCT: Murphy 2005 (cardiac; fair quality; N=61) Evidence for length of ICU stay – 3 level II studies: Damgaard 2006 (cardiac; good quality; N=60); Murphy 2009	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias			
(cardiac; fair quality; N=61); Niranjan 2006 (cardiac; good quality; N=80)	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias		
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	3		
2. Consistency					
Evidence for ICU readmission – Not applicable (one study only)	Α	All studies consistent			
Evidence for length of ICU stay – All the studies are consistent in finding no significant association between	В	Most studies consistent and inconsistency can be explained			
intraoperative cell salvage and length of ICU stay.		Some inconsistency, reflecting genuine uncertainty around questi	on		
	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact					
Evidence for ICU readmission Murphy 2005 – RR 0.33 (0.04, 3.03)	А	Very large			
Evidence for length of ICU stay	В	Substantial			
Damgaard 2006 (length of ICU stay > 24 hours) – RR 0.17 (0.02, 1.30) Murphy 2005 (cell salvage vs. control; median [IQR] length of ICU stay, days): 1.0 (1.0, 1.0) vs. 1.0 (1.0, 1.0)	С	Moderate			
Niranjan 2006 (length of ICU stay in days): mean difference (on-CPB) -0.10 (-0.35, 0.15); mean difference (off-CPB) 0.10 (-0.10, 0.30)	D	No difference/underpowered			
4. Generalisability					
All three studies were conducted in patients undergoing CABG.	Α	Evidence directly generalisable to target population			
	В	Evidence directly generalisable to target population with some cave	veats		
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied		
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to		
5. Applicability		T			
Two of the studies were conducted in the UK and the other in Denmark.	A	Evidence directly applicable to Australian healthcare context			
	В	Evidence applicable to Australian healthcare context with few cave			
	С	vidence probably applicable to Australian healthcare context with some caveats			
	D	Evidence not applicable to Australian healthcare context			

Other factors	Other factors							
EVIDENCE STA	TEMEN	IT MATRIX						
Please summarise	the devel	opment group's synthesis of the evidence relating to the key question, taking all the above factors into account.						
Component	Rating	Description						
Evidence base	В	Three level II studies with fair to good quality, with low risk of bias.						
2. Consistency	Α	All studies consistent						
3. Clinical impact	D	No difference/underpowered						
4. Generalisability	В	All studies are in patients undergoing CABG						
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats. Studies conducted in UK and Denmark.						
DRAFT EVIDEN	ICE STA	ATEMENT						

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, the effect of intraoperative cell salvage on ICU admission and length of stay is uncertain.

Abbreviations: CABG, coronary artery bypass graft; CPB, cardiopulmonary bypass; ICU, intensive care unit; IQR, interguartile range; RR, relative risk; SR, systematic review.

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

POQ3.I2.S6 Characteristics and results of studies examining the effect of intraoperative cell salvage on ICU admission and length of stay.

	Level of evidence	No. of trials /	Patient population / Surgical	:				Notes		
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Damgaard (2006)	Level II Good	N=60	Adults undergoing sub-acute coronary bypass surgery without heart-lung machine.	Hospital in Denmark	The continuously heparinised suction and reservoir belonging to the cell saver were used for all patients in both groups. The suctioned blood from patients in the cell saver group was processed and autotransfused before the patient was transferred to the ICU.	Length of ICU stay > 24 hours	1/30 (3%)	6/30 (21%)	P=0.09	
Murphy (2005)	Level II Fair	N=61	Patients scheduled for non- emergency first-time CABG (off-pump)	Hospital in UK	Intraoperative cell salvage, with Autotransfusion of washed, salvaged red blood	Median (IQR) length of ICU stay, days	1.0 (1.0, 1.0)	1.0 (1.0, 1.0)	P=0.50	
					cells at the completion of the operative procedure. Salvaged washed RBCs were autotransfused at the time of skin closure.	Readmission to ICU	1/30 (3%)	1/31 (3%)	P=0.98	
Niranjan (2006)	Level II Good	N=80	Adults undergoing first-time, isolated CABG	Hospital in UK	Intraoperative cell salvage, with Autotransfusion of washed, salvaged red blood cells at the conclusion of the procedure.	Mean (SD) length of ICU stay, days	On-CPB: 0.9 (0.4) Off-CPB: 1 (0.4)	On-CPB: 1 (0.4) Off-CPB: 0.9 (0.2)	On-CPB: P=0.43 Off-CPB: P=0.32	

Abbreviations: CABG, coronary artery bypass graft; CPB, cardiopulmonary bypass; ICU, intensive care unit; IQR, interquartile range; RBCs, red blood cells; SD, standard deviation.

Recommendation(s) for intraoperative cell salvage

RECOMMENDATION What recommendation(s) does the guideline development group draw from this evidence? Use action statements where possible.	GRADE	RADE RELEVA EVIDEN TABL		
In adult patients undergoing surgery in which substantial blood loss is anticipated, intraoperative cell salvage is recommended.				
IMPLEMENTATION OF RECOMMENDATION Please indicate yes or no to the following questions. Where the answer is yes please provide explanatory information about this. This information will be used to develop the implementation plan for the guidelines.		I		
Will this recommendation result in changes in usual care?		YES	NO	
Use of intraoperative cell salvage will increase.				
Are there any resource implications associated with implementing this recommendation?		YES	NO	
Significant costs relating to equipment, training, and staffing (technicians).				
Will the implementation of this recommendation require changes in the way care is currently organised?		YES	NO	
Workforce issues; organisational structure implications (scheduling use of the device).				
Are the guideline development group aware of any barriers to the implementation of this recommendation		YES	NO	
Initial capital investment; recurrent expenditure (but preservation of blood supply at ARCBS level, which r	esults in cost-shiftin	g from		
Commonwealth to State).				
What could help to facilitate implementation of the recommendation?		YES	NO	
Development of local policies for intraoperative cell salvage; lobbying for funding of cell salvage device.				

Intervention 3 – Perioperative acute normovolemic haemodilution combined with intraoperative cell salvage

Key question(s): In patients undergoing surgery, what is the effect of perioperative ANH and in	perative cell salvage on transfusion incidence?	Evidence table ref*: POQ3.I3.P1				
1. Evidence base						
Two level II studies: McGill 2002 (cardiac; fair quality; N=254); Wong 2002 (vascular; fair quality; N=145)	Α	One or more level I studies with a low risk of bias or several level	ral level II studies with a low risk of bias			
	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	3			
2. Consistency						
Both studies reported allogeneic blood transfusion incidence. The studies agreed in direction. In McGill 2002, ANH and cell salvage was associated with a significant reduction in transfusion incidence. In Wong 2002 the	Α	All studies consistent				
association was not significant, however this may be due to study size. A meta-analysis of the two studies found	В	Most studies consistent and inconsistency can be explained				
that the heterogeneity was not significant (P=0.54; I²=0%).		Some inconsistency, reflecting genuine uncertainty around questi	on			
	D	Evidence is inconsistent				
	NA	Not applicable (one cardiac study and one vascular surgery study)				
3. Clinical impact						
McGill 2002 (ANH + cell salvage vs. control)¹ Patents transfused with any allogeneic blood product – RR 0.69 (0.49, 0.95)	Α	Very large				
Patients transfused with allogeneic blood – RR 0.66 (0.46, 0.95)	В	Substantial				
Patients transfused with FFP – RR 0.98 (0.48, 1.98) Patients transfused with platelets – RR 0.98 (0.51, 1.87)	С	Moderate				
Patients transfused with allogeneic blood (Wong 2002) – RR 0.77 (0.55, 10.07) Meta-analysed value for transfusion of allogeneic blood – RR 0.72 (0.56, 0.91)	D	Slight/Restricted/No difference				
4. Generalisability						
McGill 2002 was conducted in patients undergoing cardiac surgery and Wong 2002 was conducted in patients	Α	Evidence directly generalisable to target population				
undergoing aortic surgery.	В	Evidence directly generalisable to target population with some caveats				
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard	to judge whether it is sensible to			
5. Applicability						
Both studies were conducted in a UK hospital setting.	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few cave	eats			
	С	Evidence probably applicable to Australian healthcare context with some caveats				
	D	Evidence not applicable to Australian healthcare context				

Other factors

The inconsistencies between McGill 2002 and Wong 2002 are likely to be due to the slightly different outcomes reported. McGill 2002 reported perioperative transfusion, and Wong 2002 reported intraoperative transfusion.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	Two fair quality level II studies – one in patients undergoing cardiac surgery and one in patients undergoing aortic surgery.
2. Consistency	В	Most studies consistent and inconsistency can be explained.
3. Clinical impact	В	Substantial clinical impact
4. Generalisability	С	Evidence not directly generalisable to the target population but could be sensibly applied.
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats. Both studies were conducted in the UK.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, a combination of ANH and intraoperative cell salvage may reduce the incidence of allogeneic blood transfusion.

Abbreviations: ANH, acute normovolemic haemodilution: FFP, fresh frozen plasma: RR, relative risk.

Primary outcomes: P1 = transfusion incidence. P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission ¹ See summary table POQ3.13.P1 for values comparing ANH + cell salvage vs. cell salvage alone.

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I3.P1 Characteristics and results of studies examining the effect of perioperative ANH and intraoperative cell salvage on transfusion incidence.

	Level of evidence	No. of trials /	Patient population / Surgical				Results			Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
McGill (2002)	Level II Fair	N=254	Adults undergoing cardiac surgery	UK hospital	ANH and intraoperative cell salvage compared with cell salvage alone or control	Patients transfused with any allogeneic blood product (n/N [%])	33/86 (38%)	<u>Cell salvage</u> 32/84 (38%) <u>Control</u> 47/84 (56%)	Cell salvage P=0.97 Control P=0.02	
	tr. al	Patients transfused with allogeneic blood (n/N [%])	29/86 (34%)	<u>Cell salvage</u> : 26/84 (31%) <u>Control</u> : 43/84 (51%)	Cell salvage P=0.70 Control P=0.02					
			Patients transfused with FFP (n/N [%])	13/86 (15%)	<u>Cell salvage</u> : 14/84 (17%) <u>Control</u> : 13/84 (15%)	Cell salvage P=0.78 Control P=0.95				
						Patients transfused with platelets (n/N [%])	15/86 (17%)	<u>Cell salvage</u> : 11/84 (13%) <u>Control</u> : 15/84 (18%)	Cell salvage P=0.43 Control P=0.94	
Wong (2002)	Level II Fair	N=145	Adults undergoing aortic surgery	UK hospital	Before skin incision, sufficient blood was taken to reduce the haemoglobin concentration to 11 g/dL. Blood volume was replaced simultaneously with crystalloids, maintaining a steady central venous pressure during blood collection. Blood lost during the procedure was salvaged. All autologous blood (ANH and salvaged) was reinfused within 6 hours of withdrawal.	Patients transfused with allogeneic blood during surgery (n/N [%])	32/74 (43%)	40/71 (56%)	P=0.12	

Abbreviations: ANH, acute normovolemic haemodilution; FFP, fresh frozen plasma.

Key question(s): In patients undergoing surgery, what is the effect of perioperative ANH and in	ntraoj	oerative cell salvage on transfusion volume?	Evidence table ref*: POQ3.I3.P2		
1. Evidence base					
Two level II studies: McGill 2002 (cardiac surgery; fair quality; N=254); Wong 2002 (aortic surgery; fair quality; N=145)	Α	One or more level I studies with a low risk of bias or several lev	rel II studies with a low risk of bias		
N=145)	В	One or two Level II studies with a low risk of bias or SR/several	Level III studies with a low risk of bias		
	С	One or two Level III studies with a low risk of bias or Level I or II	I studies with a moderate risk of bias		
	D	Level IV studies or Level I to III studies/SRs with a high risk of b	ias		
2. Consistency	ı	1			
Both studies found that ANH + cell salvage significantly reduced allogeneic blood transfusion volume.	Α	All studies consistent			
	В	Most studies consistent and inconsistency can be explained			
	С	Some inconsistency, reflecting genuine uncertainty around que	estion		
	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact					
McGill 2002 (ANH + cell salvage vs. control)¹ Units of allogeneic blood transfused – mean difference -0.44 (-0.86, -0.02); P=0.04	Α	Very large			
Units of FFP transfused – mean difference -0.06 (-0.42, 0.30); P=0.74	В	Substantial			
Units of platelets transfused – mean difference 0.02 (-0.20, 0.24); P=0.86 Wong 2002	С	Moderate			
Median (IQR) units of allogeneic blood transfused, ANH + cell salvage vs. control: 0 (0 to 2) vs. 2 (0 to 4); P=0.008	D	Slight/Restricted			
4. Generalisability					
McGill 2002 was conducted in patients undergoing cardiac surgery and Wong 2002 was conducted in patients undergoing aortic surgery.	Α	Evidence directly generalisable to target population			
undergoing aonte surgery.	В	Evidence directly generalisable to target population with some of	caveats		
	С	Evidence not directly generalisable to the target population but	could be sensibly applied		
	D	Evidence not directly generalisable to target population and har	rd to judge whether it is sensible to		
5. Applicability	1	T			
Both studies were conducted in a UK hospital setting.	Α	Evidence directly applicable to Australian healthcare context			
	В	Evidence applicable to Australian healthcare context with few context with			
	С	Evidence probably applicable to Australian healthcare context with some caveats			
	D	Evidence not applicable to Australian healthcare context			

Other	factors	
Other	ractors	

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	Two fair quality level II studies.
2. Consistency	Α	The results from both studies are consistent.
3. Clinical impact	С	Moderate clinical impact.
4. Generalisability	С	Evidence not directly generalisable to the target population but could be sensibly applied. Both studies were conducted in patients undergoing cardiovascular surgery.
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats. Both studies were conducted in the UK.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, a combination of ANH and intraoperative cell salvage may reduce the volume of allogeneic blood transfusion.

Abbreviations: ANH, acute normovolemic haemodilution; FFP, fresh frozen plasma; IQR, interquartile range.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission 1 See summary table POQ3.I3.P1 for values comparing ANH + cell salvage vs. cell salvage vs.

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I3.P2 Characteristics and results of studies examining the effect of perioperative ANH and intraoperative cell salvage on transfusion volume.

	Level of evidence	No. of trials /	Patient population / Surgical				Results			Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
McGill (2002)	Level II Fair	N=254	Adults undergoing cardiac surgery	UK hospital	ANH and intraoperative cell salvage compared with cell salvage alone or control	Units of allogeneic blood transfused during surgery ¹ (mean [SD])	0.63 (1.22)	cell salvage – 0.68 (1.55) control – 1.07 (1.56)	Cell salvage P=0.82 Control P=0.04	
		Units of FFP transfused (mean [SD])	0.43 (1.12)	Cell salvage: 0.57 (1.47) Control: 0.49 (1.25)	Cell salvage P=0.49 Control P=0.74					
						Units of platelets transfused (mean [SD])	0.31 (0.81)	Cell salvage: 0.20 (0.62) Control: 0.29 (0.67)	Cell salvage P=0.32 Control P=0.86	
Wong (2002)	Level II Fair	N=145	Adults undergoing aortic surgery	UK hospital	ANH and intraoperative cell salvage ²	Median units of allogeneic blood transfused during surgery (median [IQR])	0 (0 to 2)	2 (0 to 4)	P=0.008	
					Total units of allogeneic blood transfused during surgery (aneurysm patients) ³	102	201	NR		
						Median units of allogeneic blood transfused during surgery (for all occlusive disease patients) ⁴	0 (0 to 2)	0 (0 to 2)	Mean difference: NR; P=0.87	

		Level of evidence	No. of trials /	Patient population / Surgical				Results			Notes
St	tudy	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
							Total units of allogeneic blood transfused (occlusive disease patients) ⁴	15	50	NR	

Abbreviations: ANH, acute normovolemic haemodilution; FFP, fresh frozen plasma; IQR, intensive care unit; NR, not reported; SD, standard deviation.

¹ Nine patients needed a markedly higher amount of transfused blood (≥3 units). These patients were returned to the operating theatre for re-exploration of the mediastinum. A surgical cause of bleeding was found in seven of these patients (three in the control group and two each in the cell salvage and combined treatment groups).

² Before skin incision, sufficient blood was taken to reduce the haemoglobin concentration to 11 g/dL. Blood volume was replaced simultaneously with crystalloids, maintaining a steady central venous pressure during blood collection. Blood lost during the procedure was salvaged. All autologous blood (ANH and salvaged) was reinfused within 6 hours of withdrawal.

³Two of the patients required a laparotomy (one for massive bleeding from the proximal aortic anastomosis, one for upper gastrointestinal haemorrhage).

⁴Three patients had intraoperative bleeding and a further five required reoperation for intra-abdominal bleeding.

Key question(s): In patients undergoing surgery, what is the effect of <u>perioperative ANH and in</u>	ntraoj	perative cell salvage on blood loss?	Evidence table ref*: POQ3.I3.P3		
1. Evidence base					
One level II study: Wong 2002 (aortic surgery; fair quality; N=145)	А	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias		
	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias		
	С	One or two Level III studies with a low risk of bias or Level I or II st	tudies with a moderate risk of bias		
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	S		
2. Consistency					
	Α	All studies consistent			
	В	Most studies consistent and inconsistency can be explained			
	С	Some inconsistency, reflecting genuine uncertainty around questi	ion		
	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact	•				
Median [IQR] intraoperative blood loss (ANH + cell salvage vs. control), mL: 921 (661 to 1374) vs 1000 (688 to 1734); P=0.37	Α	Very large			
1734), P=0.37	В	Substantial			
	С	Moderate			
	D	No difference			
4. Generalisability					
Wong 2002 was conducted in patients undergoing aortic surgery.	Α	Evidence directly generalisable to target population			
	В	Evidence directly generalisable to target population with some case	veats		
	С	Evidence not directly generalisable to the target population but co	ould be sensibly applied		
	D	Evidence not directly generalisable to target population and hard	to judge whether it is sensible to		
5. Applicability					
Wong 2002 was conducted in a UK hospital setting.	Α	Evidence directly applicable to Australian healthcare context			
	В	Evidence applicable to Australian healthcare context with few cav-			
	С	Evidence probably applicable to Australian healthcare context wit	h some caveats		
	D	Evidence not applicable to Australian healthcare context			

Other	factors
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EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	One fair quality level II study with moderate risk of bias.
2. Consistency	NA	Not applicable (one study only).
3. Clinical impact	D	No difference.
4. Generalisability	С	Evidence not directly generalisable to the target population but could be sensibly applied. The study was conducted in patients undergoing aortic surgery.
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats. The study was conducted in the UK.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of a combination of ANH and intraoperative cell salvage on blood loss is uncertain.

Abbreviations: ANH, acute normovolemic haemodilution: IQR, interquartile range.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I3.P3 Characteristics and results of studies examining the effect of perioperative ANH and intraoperative cell salvage on blood loss.

		Level of evidence	No. of trials /	Patient population / Surgical	0 111			Outcome		Results		Notes
Study	Quality	sample size		Setting	Intervention	Outcome	Intervention	Comparator	p-value			
W	ong (2002)	Level II Fair	N=145	Adults undergoing aortic surgery	UK hospital	ANH and intraoperative cell salvage ¹	Intraoperative blood loss (median [IQR]), mL	921 (661 to 1374)	1000 (688 to 1734)	P=0.37		

Abbreviations: ANH, acute normovolemic haemodilution; IQR, interquartile range.

¹ Before skin incision, sufficient blood was taken to reduce the haemoglobin concentration to 11 g/dL. Blood volume was replaced simultaneously with crystalloids, maintaining a steady central venous pressure during blood collection. Blood lost during the procedure was salvaged. All autologous blood (ANH and salvaged) was reinfused within 6 hours of withdrawal.

Key question(s): In patients undergoing surgery, what is the effect of <u>periopera</u>	itive ANH and intrao	perative cell salvage on mortality?	Evidence table ref*: POQ3.I3.P4
1. Evidence base			
One level II study: Wong 2002 (fair quality; N=145)	А	One or more level I studies with a low risk of bias or se	everal level II studies with a low risk of bias
	В	One or two Level II studies with a low risk of bias or SR	R/several Level III studies with a low risk of bias
	С	One or two Level III studies with a low risk of bias or Le	evel I or II studies with a moderate risk of bias
	D	Level IV studies or Level I to III studies/SRs with a high	risk of bias
2. Consistency			
-	A	All studies consistent	
	В	Most studies consistent and inconsistency can be expl	lained
	С	Some inconsistency, reflecting genuine uncertainty are	ound question
	D	Evidence is inconsistent	
	NA	Not applicable (one study only)	
3. Clinical impact	·		
RR 1.13 (0.54, 2.36); P=0.91; N=145	А	Very large	
	В	Substantial	
	С	Moderate	
	D	No difference/underpowered	
4. Generalisability			
Wong 2002 was conducted in patients undergoing aortic surgery.	A	Evidence directly generalisable to target population	
	В	Evidence directly generalisable to target population wi	th some caveats
	С	Evidence not directly generalisable to the target popular	ation but could be sensibly applied
	D	Evidence not directly generalisable to target population	n and hard to judge whether it is sensible to
5. Applicability			
Wong 2002 was conducted in a UK hospital setting.	А	Evidence directly applicable to Australian healthcare of	
	В	Evidence applicable to Australian healthcare context w	
	С	Evidence probably applicable to Australian healthcare	
	D	Evidence not applicable to Australian healthcare conte	ext

Other factors

The study was underpowered to show a significant difference in this outcome.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
Evidence base	С	One fair quality level II study with moderate risk of bias.
2. Consistency	NA	Not applicable (one study only).
3. Clinical impact	D	No difference/underpowered.
4. Generalisability	С	Evidence not directly generalisable to the target population but could be sensibly applied. The study was conducted in patients undergoing aortic surgery.
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats. The study was conducted in the UK.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of a combination of perioperative ANH and intraoperative cell salvage on mortality is uncertain.

Abbreviations: ANH, acute normovolemic haemodilution; RR, relative risk.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I3.P4 Characteristics and results of studies examining the effect of perioperative ANH and intraoperative cell salvage on mortality.

	Level of evidence No. of trials / Patient population / Surgical Sotting Intervention Outcome		Results		Notes					
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Wong (2002)	Level II Fair	N=145	Adults undergoing aortic surgery	UK hospital	ANH and intraoperative cell salvage ¹	Mortality (n/N [%])	13/74 (18%)	11/71 (15%)	P=0.91	

Abbreviations: ANH, acute normovolemic haemodilution.

Before skin incision, sufficient blood was taken to reduce the haemoglobin concentration to 11 g/dL. Blood volume was replaced simultaneously with crystalloids, maintaining a steady central venous pressure during blood collection. Blood lost during the procedure was salvaged. All autologous blood (ANH and salvaged) was reinfused within 6 hours of withdrawal.

Key question(s): In patients undergoing surgery, what is the effect of <u>perioperative ANH and ir</u>	<u>ntraoj</u>	perative cell salvage on morbidity?	Evidence table ref*: POQ3.I3.P5			
1. Evidence base			<u> </u>			
Two level II studies: McGill 2002 (cardiac surgery; fair quality; N=254); Wong 2002 (aortic surgery; fair quality;	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias			
N=145)	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	3			
2. Consistency	ı					
The studies are consistent in finding no significant impact of ANH + cell salvage on morbidity.	Α	All studies consistent in finding no difference				
	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around question				
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact						
McGill 2002 found no significant difference between ANH + cell salvage and control for the following morbidity outcomes: all perioperative complications; haemorrhagic complications; cerebrovascular accident; arrhythmias;	Α	Very large				
renal failure; infection; or MI. McGill 2002 found no significant difference in adverse events between ANH + cell	В	Substantial				
salvage and cell salvage alone. Wong 2002 found no significant difference between ANH + cell salvage and control for infection, minor	С	Moderate				
transfusion reaction, cardiac events, and haemorrhagic complications. See Summary Table POQ.13.P5 for more details.	D	No difference/underpowered				
4. Generalisability						
McGill 2002 was conducted in patients undergoing cardiac surgery and Wong 2002 was conducted in patients undergoing aortic surgery.	Α	Evidence directly generalisable to target population				
undergoing dorne surgery.	В	Evidence directly generalisable to target population with some cave	veats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to judge whether it is sensible to				
5. Applicability	1	T				
Both studies were conducted in a UK hospital setting.	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few caveats				
	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
	D	Evidence not applicable to Australian healthcare context				

Other factors

The studies were underpowered to show a significant difference in morbidity outcomes.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account

Component	Rating	Description
1. Evidence base	С	Two fair quality level II studies.
2. Consistency	Α	Both studies are consistent.
3. Clinical impact	D	No difference/underpowered
4. Generalisability	С	Both studies were conducted in patients undergoing cardiovascular surgery.
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats. Both studies were conducted in the UK.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of a combination of perioperative ANH and intraoperative cell salvage on morbidity is uncertain.

Abbreviations: ANH, acute normovolemic haemodilution; MI, myocardial infarction.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I3.P5 Characteristics and results of studies examining the effect of perioperative ANH and intraoperative cell salvage on morbidity.

Charles	Level of evidence	No. of trials /	Patient population / Surgical	Callian	1	0.4		Results		Notes				
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value					
McGill (2002) Level II Fair		N=254	Adults undergoing cardiac surgery	UK hospital	ANH and intraoperative cell salvage compared with cell salvage alone or control	All perioperative complications (n/N [%])	46/86 (53%)	Cell salvage: 46/84 (55%) Control: 42/84 (50%)	Cell salvage P=0.87 Control P=0.65					
						Haemorrhagic complications (n/N [%])	2/86 (2%)	Cell salvage: 2/84 (2%) Control: 3/84 (4%)	Cell salvage P=0.98 Control P=0.63					
						Cerebrovascula r accident (n/N [%])	1/86 (1%)	Cell salvage: 1/84 (1%) Control: 2/84 (2%)	Cell salvage P=0.99 Control P=0.56					
										Arrhythmias (n/N [%])	20/86 (23%)	Cell salvage: 17/84 (20%) Control: 27/84 (32%)	Cell salvage P=0.63 Control P=0.20	
						Renal failure (n/N [%])	2/86 (2%)	Cell salvage: 1/84 (1%) Control: 0/84 (0%)	Cell salvage P=0.58 Control P=0.30					
						Infection (n/N [%])	7/86 (8%)	Cell salvage: 11/84 (13%) Control: 7/84 (8%)	Cell salvage P=0.30 Control P=0.96					
						MI (n/N [%])	4/84 (5%)	Cell salvage: 5/84 (6%) Control: 10/84 (12%)	Cell salvage P=0.97 Control P=0.17					
Wong (2002)	Level II Fair	N=145	Adults undergoing aortic surgery	UK hospital	ANH and intraoperative cell salvage [*]	Infection (n/N [%])	16/74 (22%)	19/71 (27%)	P=0.6					
						Minor transfusion reaction (n/N [%])	0/74 (0%)	1/71 (1%)	P=0.48					

Appendix D: Evidence matrixes – Intervention 3 (Perioperative acute normovolemic haemodilution and intraoperative cell salvage)

	Level of evidence	No. of trials /	Patient population / Surgical			Results				Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
						Cardiac events (n/N [%])	13/74 (18%)	8/71 (11%)	P=0.4	
						Haemorrhagic complications (n/N [%])	5/74 (7%)	8/71 (11%)	P=0.35	

Abbreviations: ANH, acute normovolemic haemodilution; MI, myocardial infarction.

Before skin incision, sufficient blood was taken to reduce the haemoglobin concentration to 11 g/dL. Blood volume was replaced simultaneously with crystalloids, maintaining a steady central venous pressure during blood collection. Blood lost during the procedure was salvaged. All autologous blood (ANH and salvaged) was reinfused within 6 hours of withdrawal.

Key question(s): In patients undergoing surgery, what is the effect of perioperative ANH and in	traoi	perative cell salvage on quality of life?	Evidence table ref*: POQ3.I3.P6		
1. Evidence base					
No evidence found	۸	0	- - - - - - - - - -		
No evidence round	А	One or more level I studies with a low risk of bias or several level I			
	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias		
	С	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias			
2. Consistency		1			
-	Α	All studies consistent			
	В	Most studies consistent and inconsistency can be explained			
	С	Some inconsistency, reflecting genuine uncertainty around question			
	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact					
	Α	Very large			
	В	Substantial			
	С	Moderate			
	D	Slight/Restricted			
4. Generalisability		1			
·	Α	Evidence directly generalisable to target population			
	В	Evidence directly generalisable to target population with some cav	eats		
	С	Evidence not directly generalisable to the target population but cou	uld be sensibly applied		
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to		
5. Applicability					
	Α	Evidence directly applicable to Australian healthcare context			
	В	Evidence applicable to Australian healthcare context with few cave			
	С	Evidence probably applicable to Australian healthcare context with	some caveats		
	D	Evidence not applicable to Australian healthcare context			

Other factors		
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EVIDENCE STA	TEMENT	MATRIX
Please summarise t	he develop	oment group's synthesis of the evidence relating to the key question, taking all the above factors into account.
Component	Rating	Description
Evidence base	NA	
2. Consistency	NA	
3. Clinical impact	NA	
4. Generalisability	NA	
5. Applicability	NA	
DRAFT EVIDEN	CE STA	ΓΕΜΕΝΤ
Based on the body		
		dergoing surgery in which substantial blood loss is anticipated, the effect of a combination of ANH and intraoperative cell salvage on quality of life
is unknow	n.	
Abbreviations: ANH acute	normovolomi	c haamadilution

^{*} Interventions: 11 = ANH, 12 = intraoperative cell salvage, 13 = perioperative ANH and intraoperative cell salvage, 14 = postoperative cell salvage, 15 = deliberate induced hypotension, 16 = prevention of hypothermia, 17 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Key question(s): In patients undergoing surgery, what is the effect of perioperative ANH and i	ntranı	perative cell salvage on change in haemoglobin	Evidence table ref*: POQ3.I3.S1			
in patients undergoing surgery, what is the effect of perioperative Aivit and i	nuau	ocrative cell salvage on change in nacmoglobin	1 0 20110101			
1. Evidence base						
One level II study: McGill 2002 (fair quality; N=254)	А	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias			
	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency		,				
	А	All studies consistent				
	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around question	on			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact						
Median (IQR) haemoglobin concentration; ANH + cell salvage vs. control ¹	Α	A Very large				
Preoperative: 145 (138 to 150) vs. 142 (135 to 150); P=NR At admission to ICU: 108 (99 to 116) vs. 100 (91 to 107); P=NR	В	Substantial				
24 hours after surgery: 105 (96 to 113) vs. 100 (94 to 109); P=NR Three days after surgery: 108 (100 to 119) vs. 106 (98 to 112); P=NR	С	Moderate				
Tillee days after Surgery. 100 (100 to 119) vs. 100 (46 to 112), P=NR	D	Slight/Restricted				
4. Generalisability						
McGill 2002 was conducted in patients undergoing cardiac surgery.	А	Evidence directly generalisable to target population				
	В	Evidence directly generalisable to target population with some cave	reats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to			
5. Applicability	_					
McGill 2002 was conducted in a UK hospital setting.	A	Evidence directly applicable to Australian healthcare context				
	В					
	С	Evidence probably applicable to Australian healthcare context with some caveats				
	D	Evidence not applicable to Australian healthcare context				

Other factors

24 hours post-op was considered by the CRG to be most relevant.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	One fair quality level II study with moderate risk of bias.
2. Consistency	NA	Not applicable (one study only).
3. Clinical impact	D	No significant impact.
4. Generalisability	С	The study was conducted in patients undergoing cardiac surgery.
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats. The study was conducted in the UK

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of a combination of ANH and intraoperative cell salvage on postoperative haemoglobin concentration is uncertain.

Abbreviations: ANH, acute normovolemic haemodilution; ICU, intensive care unit; IQR, interquartile range; NR, not reported.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission

1 See summary table POQ3.13.P1 for values comparing ANH + cell salvage vs. cell salvage alone.

^{* &}lt;u>Interventions</u>: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I3.S1 Characteristics and results of studies examining the effect of perioperative ANH and intraoperative cell salvage on change in haemoglobin concentration.

	Level of evidence	No. of trials /	Patient population / Surgical		Intervention Outcome		Notes			
Study	Quality	sample size	procedure	Setting		Outcome	Intervention	Comparator	p-value	
McGill (2002)	Level II Fair	N=254	Adults undergoing cardiac surgery	UK hospital	ANH and intraoperative cell salvage compared with cell salvage alone or control	Preoperative Hb concentration (median [IQR]), g/dL	145 (138 to 150)	Cell salvage: 145 (136 to 150) Control: 142 (135 to 150)	NR	
		C O IC	Hb concentration on admission to ICU (median [IQR]), g/dL	108 (99 to 116)	Cell salvage: 105 (98 to 116) Control: 100 (91 to 107)	NR				
						Hb concentration 24 hours after surgery (median [IQR]), g/dL	105 (96 to 113)	Cell salvage: 104 (95 to 115) Control: 100 (94 to 109)	NR	
			Hb concentration 3 days after surgery (median [IQR]), g/dL	108 (100 to 119)	Cell salvage: 105 (98 to 115) Control: 106 (98 to 112)	NR				

Abbreviations: ANH, acute normovolemic haemodilution; Hb, haemodilution; ICU, intensive care unit; IQR, interquartile range; NR, not reported.

Key question(s): In patients undergoing surgery, what is the effect of periopera	ntive ANH and intrao	perative cell salvage on reoperation for bleeding? Evidence table ref*: POQ3.I3.S2
1. Evidence base		·
One level II study: Wong 2002 (fair quality; N=145)	А	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias
	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias
	С	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias
2. Consistency		
	A	All studies consistent
	В	Most studies consistent and inconsistency can be explained
	С	Some inconsistency, reflecting genuine uncertainty around question
	D	Evidence is inconsistent
	NA	Not applicable (one study only)
3. Clinical impact		
RR 1.37 (0.55, 3.40); P=0.50	А	Very large
	В	Substantial
	С	Moderate
	D	No difference/underpowered
4. Generalisability	·	
Wong 2002 was conducted in patients undergoing aortic surgery.	А	Evidence directly generalisable to target population
	В	Evidence directly generalisable to target population with some caveats
	С	Evidence not directly generalisable to the target population but could be sensibly applied
	D	Evidence not directly generalisable to target population and hard to judge whether it is sensible to
5. Applicability		
Wong 2002 was conducted in a UK hospital setting.	Α	Evidence directly applicable to Australian healthcare context
	В	Evidence applicable to Australian healthcare context with few caveats
	С	Evidence probably applicable to Australian healthcare context with some caveats
	D	Evidence not applicable to Australian healthcare context

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EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description					
1. Evidence base	С	One fair quality level II study with moderate risk of bias.					
2. Consistency	NA	Not applicable (one study only).					
3. Clinical impact	D	No difference/underpowered					
4. Generalisability	С	The study was conducted in patients undergoing aortic surgery.					
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats. The study was conducted in the UK.					

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of a combination of ANH and intraoperative cell salvage on risk of reoperation for bleeding is uncertain.

Abbreviations: ANH, acute normovolemic haemodilution; RR, relative risk.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I3.S2 Characteristics and results of studies examining the effect of perioperative ANH and intraoperative cell salvage on reoperation for bleeding.

Study	Level of evidence Quality	No. of trials / sample size	Patient population / Surgical procedure	Setting	Intervention	Outcome	Results			Notes
							Intervention	Comparator	p-value	
Wong (2002)	Level II Fair	N=145	Adults undergoing aortic surgery	UK hospital	ANH and intraoperative cell salvage ¹	Reoperation (n/N [%])	10/74 (14%)	7/71 (10%)	P=0.50	

Abbreviations: ANH, acute normovolemic haemodilution.

¹ Before skin incision, sufficient blood was taken to reduce the haemoglobin concentration to 11 g/dL. Blood volume was replaced simultaneously with crystalloids, maintaining a steady central venous pressure during blood collection. Blood lost during the procedure was salvaged. All autologous blood (ANH and salvaged) was reinfused within 6 hours of withdrawal.

Key question(s):	Evidence table ref*:					
In patients undergoing surgery, what is the effect of perioperative ANH and in	POQ3.I3.S5					
1. Evidence base						
Two level II studies: McGill 2002 (fair quality; N=254); Wong 2002 (fair quality; N=145)	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias				
	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias				
	С	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency	1	1				
Both studies are consistent in finding no significant difference between treatment arms.	Α	All studies consistent				
	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around question				
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact						
McGill 2002 (median [IQR]) Length of hospital stay (ANH + cell salvage vs. control): 170 (147.1 to 221.6) vs. 168.9 (140.3 to 219.3); P=NR	Α	Very large				
Length of hospital stay (ANH + cell salvage vs. collitor): 170 (147.1 to 221.6) vs. 168.9 (140.3 to 219.3); P=NR Length of hospital stay (ANH + cell salvage vs. cell salvage alone): 170 (147.1 to 221.6) vs. 160.7 (145.5 to	В	Substantial				
198.8); P=NR Kruskal-Wallis P-value=0.724	С	Moderate				
Wong (2002) (median [IQR], ANH +cell salvage vs. control)	D	No difference				
Length of hospital stay: 10 (8 to 13) vs. 9 (7 to 12); P=0.17		<u> </u>				
4. Generalisability McGill 2002 was conducted in patients undergoing cardiac surgery and Wong 2002 was conducted in patients	Ι ,	Cridence discetty removalisable to toward new detica				
undergoing aortic surgery.	Α	Evidence directly generalisable to target population				
	В	Evidence directly generalisable to target population with some caveats				
	С	Evidence not directly generalisable to the target population but could be sensibly applied				
	D	Evidence not directly generalisable to target population and hard	hard to judge whether it is sensible to			
5. Applicability						
Both studies were conducted in a UK hospital setting.	Α	Evidence directly applicable to Australian healthcare context				
		Evidence applicable to Australian healthcare context with few caveats				
	С	Evidence probably applicable to Australian healthcare context wit	h some caveats			
	D	Evidence not applicable to Australian healthcare context				

rr-	
Other factors	
EVIDENCE STATEMENT MATRIX	

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	Two fair quality level II studies, with a moderate risk of bias.
2. Consistency	Α	Both studies consistent.
3. Clinical impact	D	No statistically significant impact.
4. Generalisability	С	Evidence not directly generalisable to the target population but could be sensibly applied. Both studies were conducted in patients undergoing cardiac surgery.
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats. Both studies were conducted in the UK.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of a combination of ANH and intraoperative cell salvage on hospital length of stay is uncertain.

Abbreviations: ANH, acute normovolemic haemodilution; IQR, interguartile range.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I3.S5 Characteristics and results of studies examining the effect of perioperative ANH and intraoperative cell salvage on hospital length of stay.

	Level of evidence	No. of trials / sample size	Patient population / Surgical procedure	Setting			Results			Notes
Study	Quality				Intervention	Outcome	Intervention	Comparator	p-value	
McGill (2002)	Level II Fair	N=254	Adults undergoing cardiac surgery	UK hospital	ANH and intraoperative cell salvage compared with cell salvage alone or control	Length of hospital stay (median [IQR]), days	170 (147.1 to 221.6)	Cell salvage: 160.7 (145.5 to 198.8) Control: 168.9 (140.3 to 219.3)	Kruskal- Wallis P- value=0.724	
Wong (2002)	Level II Fair	N=145	Adults undergoing aortic surgery	UK hospital	ANH and intraoperative cell salvage ¹	Length of hospital stay (median [IQR]), days	10 (8 to 13)	9 (7 to 12)	P=0.17	

Abbreviations: ANH, acute normovolemic haemodilution; IQR, interquartile range.

¹ Before skin incision, sufficient blood was taken to reduce the haemoglobin concentration to 11 g/dL. Blood volume was replaced simultaneously with crystalloids, maintaining a steady central venous pressure during blood collection. Blood lost during the procedure was salvaged. All autologous blood (ANH and salvaged) was reinfused within 6 hours of withdrawal.

Key question(s): In patients undergoing surgery, what is the effect of perioperative ANH and in	perative cell salvage on ICU length of stay?	Evidence table ref*: POQ3.I3.S6				
1. Evidence base			1			
Two level II studies: McGill 2002 (fair quality; N=254); Wong 2002 (fair quality; N=145)	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias				
	В	One or two Level II studies with a low risk of bias or SR/several Le	evel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II s	tudies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	S			
2. Consistency	I					
Both studies are consistent in finding no significant difference between treatment arms.	Α	All studies consistent				
	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around quest	ion			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact	•					
McGill 2002 (median [IQR])	Α	Very large				
Length of ICU stay (ANH + cell salvage vs. control): 23.3 (22.5 to 25.0) vs. 22.9 (21.8 to 24.5); P=NR Length of ICU stay (ANH + cell salvage vs. cell salvage alone): 23.3 (22.5 to 25.0) vs. 22.7 (22.0 to 24.6); P=NR	В	Substantial				
Kruskal-Wallis P-value=0.249 Wong (2002) (median [IQR], ANH +cell salvage vs. control)	С	Moderate				
Length of ICU stay: 1 (0 to 25) vs. 1 (0 to 25); P=0.89	D	No difference				
4. Generalisability						
McGill 2002 was conducted in patients undergoing cardiac surgery and Wong 2002 was conducted in patients undergoing aortic surgery.	Α	Evidence directly generalisable to target population				
undergoing aonite surgery.	В	Evidence directly generalisable to target population with some ca	veats			
	С	Evidence not directly generalisable to the target population but co	ould be sensibly applied			
	D	Evidence not directly generalisable to target population and hard	to judge whether it is sensible to			
5. Applicability						
Both studies were conducted in a UK hospital setting.	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few cav				
	С	Evidence probably applicable to Australian healthcare context wit	h some caveats			
	D	Evidence not applicable to Australian healthcare context				

Other factors		
EVIDENCE STA	ATEMEN	IT MATRIX
Please summarise	the devel	opment group's synthesis of the evidence relating to the key question, taking all the above factors into account.
Component	Rating	Description
Evidence base	С	Two fair quality level II studies, with a moderate risk of bias.
2. Consistency	Α	Both studies consistent.
3. Clinical impact	D	No statistically significant impact.
4. Generalisability	С	Evidence not directly generalisable to the target population but could be sensibly applied. One study was in cardiac study and the other in aortic surgery.
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats. Both studies were conducted in the UK.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of a combination of ANH and intraoperative cell salvage on ICU length of stay is uncertain.

Abbreviations: ANH, acute normovolemic haemodilution; ICU, intensive care unit; IQR, interquartile range.

^{* &}lt;u>Interventions</u>: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD <u>Primary outcomes</u>: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

POQ3.I3.S6 Characteristics and results of studies examining the effect of perioperative ANH and intraoperative cell salvage on length of ICU stay.

	Level of	evidence No. of trials / sample size	Patient population / Surgical procedure	Setting	Intervention		Results			Notes
Study	Quality					Outcome	Intervention	Comparator	p-value	
McGill (2002)	Level II Fair	N=254	Adults undergoing cardiac surgery	UK hospital	ANH and intraoperative cell salvage compared with cell salvage alone or control	Length of ICU stay (median [IQR]), days	23.3 (22.5 to 25.0)	Cell salvage: 22.7 (22.0 to 24.6) Control: 22.9 (21.8 to 24.5)	Kruskal- Wallis P- value=0.249	
Wong (2002)	Level II Fair	N=145	Adults undergoing aortic surgery	UK hospital	ANH and intraoperative cell salvage ¹	Length of ICU stay (median [IQR]), days	1 (0 to 25)	1 (0 to 25)	P=0.89	

Abbreviations: ANH, acute normovolemic haemodilution; ICU, intensive care unit; IQR, interquartile range.

Before skin incision, sufficient blood was taken to reduce the haemoglobin concentration to 11 g/dL. Blood volume was replaced simultaneously with crystalloids, maintaining a steady central venous pressure during blood collection. Blood lost during the procedure was salvaged. All autologous blood (ANH and salvaged) was reinfused within 6 hours of withdrawal.

Recommendation(s) for acute normovolemic haemodilution combined with intraoperative cell salvage

RECOMMENDATION What recommendation(s) does the guideline development group draw from this evidence? Use action statements where possible.					
No recommendation made for this combined intervention. See individual recommendations for					
intervention 1 (acute normovolemic haemodilution) and intervention 2 (intraoperative cell salvage).					
IMPLEMENTATION OF RECOMMENDATION					
Please indicate yes or no to the following questions. Where the answer is yes please provide explanatory information about this. This information will be used to develop the implementation plan for the guidelines.					
Will this recommendation result in changes in usual care?		YES	NO		
Are there any resource implications associated with implementing this recommendation?		YES	NO		
Will the implementation of this recommendation require changes in the way care is currently organised?		YES	NO		
Are the guideline development group aware of any barriers to the implementation of this recommendation		YES	NO		
What could help to facilitate implementation of the recommendation?		YES	NO		

Intervention 4 – Postoperative cell salvage

In patients undergoing surgery, what is the effect of postoperative cell salvage on transfusion incidence? 1. Evidence base 1. Evidence base 1. Evidence base 1. Event study controls 2006 good quality includes 18 RCTs* (N=146/): 10 cardiac and 8 orthopaedic (all bit hough 1), and 1) and 1). The evidence base is a significant for Cardess 2006 search date. Amin 2008 (orthopaedic surgery; bir quality, N=46/): Acharopoutos 2007 (orthopaedic surgery; paragraphy, N=46/): A surgery paragraphy (N=46/): A surgery par	Key question(s):			Evidence table ref*:		
New I study. Cariess 2006, good quality, includes 18 RCTs' (N-1462): 10 cardiac and 8 orthopsedic salleys reposition of suddes published after the Carless 2006 search date. Amin 2008 (orthopsedic salgery) progradily N-19), Chong 2005 (orthopsedic salgery); the operation of suddes published after the Carless 2006 search date. Amin 2008 (orthopsedic salgery); progradily N-19), Chong 2005 (orthopsedic salgery); the operation of suddes published after the Carless 2006 search date. Amin 2008 (orthopsedic salgery); progradily N-19), Chong 2005 (orthopsedic salgery); the operation of suddes salgery); the operation of suddes salgery in the Irist in Carless 2006 (Phel-0.00001). The impact of prosperative call is studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/sever	In patients undergoing surgery, what is the effect of postoperative cell salvage	<u>e</u> on	transfusion incidence?	POQ3.I4.P1		
Sevel Its Judies published after the Carless 2006 search date. Amin 2008 (orthopaedic surgery; fair quality, N-60); Zacharopoulos 2007 (orthopaedic surgery; fair quality, N-60); Zachar	1. Evidence base					
Second Studies published after the Carless 2006 search date. Armin 2008 (orthopaedic surgery, fair quality, N=40) 2cd actanopoulos 2007 (orthopaedic surgery per		Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias			
Consistency	3 level II studies published after the Carless 2006 search date: Amin 2008 (orthopaedic surgery; fair quality;	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias		
Consistency There is a significant degree of heterogeneity within the trials in Carless 2006 (Rhet-Concount). The impact of postoperative cell salvage is significant for all of the subgroups analysed in Carless 2006 except studies without a transfusion protocol. In all three of the RCTs uncovered in the SR conducted herein, cell salvage had no many analysed in Carless 2006 except studies without a transfusion incidence. A Ill studies consistent and inconsistency can be explained (orthopaedic)		С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias		
There is a significant degree of heterogeneity within the Italia in Carless 2006 (Phete-Cu0001). The impact of postoperalive cell salvage is significant facil of the Subgroups analysed in Carless 2006 except studies without transfusion protocol. In all three of the RCTs uncovered in the SR conducted herein, cell salvage had no impact on transfusion incidence. A Mature of the RCTs uncovered in the SR conducted herein, cell salvage had no impact on transfusion incidence. A Meta-analysis was conducted using the data from Carless 2006 and the three RCTs uncovered in the systematic review conducted herein. All surgery types — RR 0.0 (0.7, 0.77): 21 Irials (N=1760) Cardiac surgery — RR 0.8 (0.7, 1.00): 01 trials (N=1780) Studies with a transfusion protocol — RR 0.21 (0.03, 1.70): 4 trials (N=1375) Studies with a transfusion protocol — RR 0.21 (0.03, 1.70): 4 trials (N=385) 4. Ceneralisability A Evidence directly generalisable to target population with some caveats Evidence with approximately 8000 inhabitants. Exclusion of this study from the meta-analysis does not impact in Greece with approximately 8000 inhabitants. Exclusion of this study from the meta-analysis does not impact in forece with approximately 8000 inhabitants. Exclusion of this study from the meta-analysis does not impact the results. A All studies consistent and inconsistency can be explained (orthopaedic) Some inconsistent (Some inconsistent) Most of the studies were conducted using the data from Carless 2006 and the three RCTs uncovered in the studies were conducted in developed countries. Zacharopoulos 2007 was conducted in a small town in Greece with approximately 8000 inhabitants. Exclusion of this study from the meta-analysis does not impact the results. E Vidence directly generalisable to Australian healthcare context with few caveats E Vidence directly applicable to Australian healthcare context with few caveats E Vidence directly applicable to Australian healthcare context with some caveats		D	Level IV studies or Level I to III studies/SRs with a high risk of bias			
postperative cell sahage is significant for all of the subgroups analysed in Carless 2006 except studies without a transfusion incidence. A meta-analysis was conducted using the data from Carless 2006 and the three RCTs uncovered in the systematic review conducted herein, C77.72 trials (N-1737) A meta-analysis was conducted using the data from Carless 2006 and the three RCTs uncovered in the systematic review conducted herein. A meta-analysis was conducted herein. A meta-analysis was conducted herein. A laurger types - RR 0.05 (0.74, 1.07): 21 trials (N-1760) Cardiac surgery - RR 0.05 (0.74, 1.00): 10 trials (N-173) Studies without a transfusion protocol - RR 0.66 (0.52, 0.85): 17 trials (N-1375) Studies without a transfusion protocol - RR 0.21 (0.03, 1.70): 4 trials (N-385) A Substantial A Very large (orthopaedic) B Substantial C Moderate D Slight/Restricted (cardiac) A Seneralisability A Evidence directly generalisable to target population with some caveats C Evidence not directly generalisable to target population and hard to judge whether it is sensible to be reviewed to face the target population and hard to judge whether it is sensible to receive the face office the studies were conducted in developed countries. Zacharopoulos 2007 was conducted in a small town in Greece with approximately 8000 inhabitants. Exclusion of this study from the meta-analysis does not impact the results. A Evidence directly applicable to Australian healthcare context with few caveats C Evidence probably applicable to Australian healthcare context with some caveats C Evidence applicable to Australian healthcare context with some caveats C Evidence probably applicable to Australian healthcare context with some caveats C Evidence probably applicable to Australian healthcare context with some caveats	2. Consistency					
In ansilusion protocol. In all three of the RCTs uncovered in the SR conducted herein, cell salvage had no impact on transfusion incidence. C	There is a significant degree of heterogeneity within the trials in Carless 2006 (Phet<0.00001). The impact of	Α	All studies consistent			
C Some inconsistency, reflecting genuine uncertainty around question (cardiac)	transfusion protocol. In all three of the RCTs uncovered in the SR conducted herein, cell salvage had no impact	В	Most studies consistent and inconsistency can be explained (ortho	ppaedic)		
3. Clinical impact A meta-analysis was conducted using the data from Carless 2006 and the three RCTs uncovered in the systematic review conducted herein. All surgery Npcs - RR 0.60 (0.47, 0.77); 21 trials (N=1760) Cardiac surgery - RR 0.86 (0.74, 1.00); 10 trials (N=1760) Cardiac surgery - RR 0.37 (0.24, 0.55); 11 trials (N=1017) Studies with a transfusion protocol - RR 0.65 (0.52, 0.85); 17 trials (N=1375) Studies without a transfusion protocol - RR 0.21 (0.03, 1.70); 4 trials (N=385) 4. Generalisability All the studies were conducted in patients undergoing cardiac surgery or orthopaedic surgery. A Evidence directly generalisable to target population B Evidence directly generalisable to target population with some caveats C Evidence not directly generalisable to target population and hard to judge whether it is sensible to be sensibly applied D Evidence unit directly generalisable to Australian healthcare context B Evidence directly applicable to Australian healthcare context B Evidence applicable to Australian healthcare context with some caveats C Evidence probably applicable to Australian healthcare context with some caveats C Evidence applicable to Australian healthcare context with some caveats C Evidence probably applicable to Australian healthcare context with some caveats C Evidence applicable to Australian healthcare context with some caveats C Evidence probably applicable to Australian healthcare context with some caveats	on transfusion incidence.	С	Some inconsistency, reflecting genuine uncertainty around question	on (cardiac)		
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A meta-analysis was conducted using the data from Carless 2006 and the three RCTs uncovered in the systematic review conducted herein. All surgery types – RR 0.60 (0.47, 0.77); 21 trials (N=1760) Cardiac surgery – RR 0.80 (0.47, 0.55); 11 trials (N=1017) Studies with a transfusion protocol – RR 0.66 (0.52, 0.85); 17 trials (N=1375) Studies without a transfusion protocol – RR 0.21 (0.03, 1.70); 4 trials (N=385) 4. Generalisability All the studies were conducted in patients undergoing cardiac surgery or orthopaedic surgery. All the studies were conducted in patients undergoing cardiac surgery or orthopaedic surgery. All the studies were conducted in patients undergoing cardiac surgery or orthopaedic surgery. A Evidence directly generalisable to target population B Evidence directly generalisable to the target population but could be sensibly applied C Evidence not directly generalisable to target population and hard to judge whether it is sensible to be sensible to the target population and hard to judge whether it is sensible to be sensible to a sensible to target population and hard to judge whether it is sensible to the results. 5. Applicability Most of the studies were conducted in developed countries. Zacharopoulos 2007 was conducted in a small town in Greece with approximately 8000 inhabitants. Exclusion of this study from the meta-analysis does not impact in Greece with approximately 8000 inhabitants. Exclusion of this study from the meta-analysis does not impact the results. A Evidence directly applicable to Australian healthcare context with few caveats C Evidence applicable to Australian healthcare context with some caveats		NA	Not applicable (one study only)			
systematic review conducted herein. All surgery types – RR 0.60 (0.47, 0.77); 21 trials (N=1760) Cardiac surgery – RR 0.80 (0.74, 1.00); 10 trials (N=743) Orthopaedic surgery – RR 0.37 (0.24, 0.55); 11 trials (N=1017) Studies with a transfusion protocol – RR 0.66 (0.52, 0.85); 17 trials (N=1375) Studies without a transfusion protocol – RR 0.21 (0.03, 1.70); 4 trials (N=385) 4. Generalisability All the studies were conducted in patients undergoing cardiac surgery or orthopaedic surgery. All the studies were conducted in patients undergoing cardiac surgery or orthopaedic surgery. All the studies were conducted in patients undergoing cardiac surgery or orthopaedic surgery. Be Evidence directly generalisable to target population with some caveats C Evidence not directly generalisable to target population but could be sensibly applied D Evidence orthic trictly generalisable to target population and hard to judge whether it is sensible to surgery or the studies were conducted in developed countries. Zacharopoulos 2007 was conducted in a small town in Greece with approximately 8000 inhabitants. Exclusion of this study from the meta-analysis does not impact the results. All Evidence directly applicable to Australian healthcare context with few caveats C Evidence applicable to Australian healthcare context with some caveats	3. Clinical impact					
All surgery types – RR 0.60 (0.47, 0.77); 21 trials (N=1760) Cardiac surgery – RR 0.86 (0.74, 1.00); 10 trials (N=1743) Orthoppaedic surgery – RR 0.86 (0.74, 1.00); 10 trials (N=1743) Studies with a transfusion protocol – RR 0.66 (0.52, 0.85); 17 trials (N=1375) Studies without a transfusion protocol – RR 0.21 (0.03, 1.70); 4 trials (N=385) 4. Generalisability All the studies were conducted in patients undergoing cardiac surgery or orthopaedic surgery. All the studies were conducted in patients undergoing cardiac surgery or orthopaedic surgery. A Evidence directly generalisable to target population B Evidence not directly generalisable to the target population but could be sensibly applied C Evidence not directly generalisable to target population and hard to judge whether it is sensible to the surger population and hard to judge whether it is sensible to the surger population and hard to judge whether it is sensible to the surger population and hard to judge whether it is sensible to the surger population and hard to judge whether it is sensible to the surger population and hard to judge whether it is sensible to access with approximately 8000 inhabitants. Exclusion of this study from the meta-analysis does not impact the results. A Evidence directly applicable to Australian healthcare context with few caveats C Evidence applicable to Australian healthcare context with some caveats	A meta-analysis was conducted using the data from Carless 2006 and the three RCTs uncovered in the	Α	Very large (orthopaedic)			
Orthopaedic surgery – RR 0.37 (0.24, 0.55); 11 trials (N=1017) Studies with a transfusion protocol – RR 0.66 (0.52, 0.85); 17 trials (N=375) Studies without a transfusion protocol – RR 0.21 (0.03, 1.70); 4 trials (N=385) 4. Generalisability All the studies were conducted in patients undergoing cardiac surgery or orthopaedic surgery. All the studies were conducted in patients undergoing cardiac surgery or orthopaedic surgery. All the studies were conducted in patients undergoing cardiac surgery or orthopaedic surgery. A Evidence directly generalisable to target population B Evidence not directly generalisable to target population but could be sensibly applied C Evidence not directly generalisable to target population and hard to judge whether it is sensible to surgery population and hard to judge whether it is sensible to surgery population and hard to judge whether it is sensible to surgery population and hard to judge whether it is sensible to surgery population and hard to judge whether it is sensible to surgery population and hard to judge whether it is sensible to surgery population and hard to judge whether it is sensible to surgery population and hard to judge whether it is sensible to surgery population and hard to judge whether it is sensible to surgery population and hard to judge whether it is sensible to surgery population and hard to judge whether it is sensible to surgery population and hard to judge whether it is sensible to surgery population and hard to judge whether it is sensible to surgery population and hard to judge whether it is sensible to surgery population and hard to judge whether it is sensible to surgery population and hard to judge whether it is sensible to a surgery population and hard to judge whether it is sensible to a surgery population and hard to judge whether it is sensible to a surgery population and hard to judge whether it is sensible to a surgery population and hard to judge whether it is sensible to a surgery population and hard to judge whether it is sensible to	All surgery types – RR 0.60 (0.47, 0.77); 21 trials (N=1760)	В	Substantial			
Studies with a transfusion protocol – RR 0.66 (0.52, 0.85): 17 trials (N=1375) Studies without a transfusion protocol – RR 0.21 (0.03, 1.70): 4 trials (N=385) 4. Generalisability All the studies were conducted in patients undergoing cardiac surgery or orthopaedic surgery. All the studies were conducted in patients undergoing cardiac surgery or orthopaedic surgery. A Evidence directly generalisable to target population B Evidence directly generalisable to target population with some caveats C Evidence not directly generalisable to target population but could be sensibly applied D Evidence not directly generalisable to target population and hard to judge whether it is sensible to the studies were conducted in developed countries. Zacharopoulos 2007 was conducted in a small town in Greece with approximately 8000 inhabitants. Exclusion of this study from the meta-analysis does not impact the results. A Evidence directly applicable to Australian healthcare context with few caveats C Evidence applicable to Australian healthcare context with some caveats C Evidence probably applicable to Australian healthcare context with some caveats		С	Moderate			
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B Evidence directly generalisable to target population with some caveats C Evidence not directly generalisable to the target population but could be sensibly applied D Evidence not directly generalisable to target population and hard to judge whether it is sensible to 5. Applicability Most of the studies were conducted in developed countries. Zacharopoulos 2007 was conducted in a small town in Greece with approximately 8000 inhabitants. Exclusion of this study from the meta-analysis does not impact the results. A Evidence directly generalisable to target population and hard to judge whether it is sensible to Evidence directly applicable to Australian healthcare context B Evidence applicable to Australian healthcare context with few caveats C Evidence probably applicable to Australian healthcare context with some caveats	4. Generalisability					
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Most of the studies were conducted in developed countries. Zacharopoulos 2007 was conducted in a small town in Greece with approximately 8000 inhabitants. Exclusion of this study from the meta-analysis does not impact the results. A Evidence directly applicable to Australian healthcare context with few caveats B Evidence applicable to Australian healthcare context with some caveats C Evidence probably applicable to Australian healthcare context with some caveats		D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to		
in Greece with approximately 8000 inhabitants. Exclusion of this study from the meta-analysis does not impact the results. B Evidence applicable to Australian healthcare context with few caveats C Evidence probably applicable to Australian healthcare context with some caveats	5. Applicability					
the results. B Evidence applicable to Australian healthcare context with few caveats C Evidence probably applicable to Australian healthcare context with some caveats	Most of the studies were conducted in developed countries. Zacharopoulos 2007 was conducted in a small town in Greece with approximately 2000 inhabitants. Exclusion of this study from the meta analysis does not impact	Α	Evidence directly applicable to Australian healthcare context			
		В	Evidence applicable to Australian healthcare context with few cave	eats		
D Evidence not applicable to Australian healthcare context		С	Evidence probably applicable to Australian healthcare context with	some caveats		
		D	Evidence not applicable to Australian healthcare context			

Other factors

Carless 2006 included several older studies. The CRG noted that cell salvage technology has changed over time. The transfusion triggers within these studies have also changed over this time.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

			· · · · · · · · · · · · · · · · · · ·
Component	Cardiac	TKA	Description
1. Evidence base	С	С	One good quality level I study of fair quality level II studies and three subsequently published fair-to-poor quality level II studies.
2. Consistency	С	В	In cardiac evidence there is some inconsistency reflecting genuine uncertainty. In TKA most studies consistent and inconsistency can be explained
3. Clinical impact	D	Α	Slight impact in cardiac surgery and a very large impact in TKA.
4. Generalisability	В	В	Evidence directly generalisable to target population (with some caveats for cardiac).
5. Applicability	В	В	Evidence applicable to Australian healthcare context with few caveats.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing total knee arthroplasty, postoperative cell salvage reduces the incidence of allogeneic blood transfusion.

In adult patients undergoing cardiac surgery, postoperative cell salvage may reduce the incidence of allogeneic blood transfusion.

Abbreviations: RCT, randomised controlled trial; RR, relative risk; SR, systematic review; TKA, total knee arthroplasty.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I4.P1 Characteristics and results of studies examining the effect of postoperative cell salvage on transfusion incidence.

	Level of evidence	No. of trials /	Patient population / Surgical					Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Carless (2006)	Level I Good	18 trials (fair quality) N=1462	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Patients transfused with allogeneic blood	287/738 (39%)	473/724 (65%)	P=0.0002	Phet<0.00001
Amin (2008)	Level II Fair	N=178	Adults undergoing unilateral TKA	Hospital in UK	Postoperative cell salvage: patients in the reinfusion group had their blood reinfused from drains within 6 hours of surgery.	Patients transfused with allogeneic blood	12/92 (13%)	13/86 (15%)	P=0.69	
Cheng (2005)	Level II Fair	N=60	Adults undergoing TKA	Hospital in Hong Kong	Postoperative cell salvage: patients in the reinfusion group had their blood reinfused from drains within 6 hours of surgery.	Patients transfused with allogeneic blood	4/26 (15%)	13/34 (38%)	P=0.07	
Zacharopoulos (2007)	Level II Poor	N=60	Adults undergoing TKA	The study was conducted in a small town of ~8000 inhabitants in Greece	Postoperative cell salvage and reinfusion of washed blood within 6 hours of operation.	Patients transfused with allogeneic blood	5/30 (17%)	10/30 (33%)	P=0.15	
Cardiac surgery	,							•		
Carless (2006)	Level I Good	10 trials (fair quality) N=743	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Patients transfused with allogeneic blood	232/375 (62%)	275/368 (75%)	P=0.05	Phet=0.0001
Orthopaedic sur	gery	1				•			•	
Carless (2006)	Level I Good	8 trials (fair quality) N=719	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Patients transfused with allogeneic blood	55/363 (15%)	198/356 (56%)	P<0.00001	Phet=0.002
Studies with a tra	ansfusion proto	col								
Carless (2006)	Level I Good	15 trials (fair quality) N=1137	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Patients transfused with allogeneic blood	233/576 (40%)	348/561 (62%)	P=0.002	Phet<0.00001

	Level of evidence	No. of trials /	Patient population / Surgical				Results			Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Studies without a	transfusion pro	otocol								
Carless (2006)	Level I Good	3 trials (fair quality) N=179	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Patients transfused with allogeneic blood	54/162 (33%)	125/163 (77%)	P=0.27	Phet<0.00001

Abbreviations: TKA, total knee arthroplasty

Key question(s):	Evidence table ref*:								
In patients undergoing surgery, what is the effect of <u>postoperative cell salvage</u> on <u>transfusion volume?</u> POQ3.I4.P2									
1. Evidence base									
1 level I study: Carless 2006: good quality; includes 9 RCTs1 (N=689): 7 cardiac and 2 orthopaedic (all fair quality)	А	One or more level I studies with a low risk of bias or several level	Il studies with a low risk of bias						
3 level II studies published after the Carless 2006 search date: Amin 2008 (orthopaedic; fair quality; N=178); Cheng 2005 (orthopaedic; fair quality; N=60); Zacharopoulos 2007 (orthopaedic; poor quality; N=60).	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias						
NB: none of the three RCTs above provided enough information to conduct a meta-analysis. Similarly none of the	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias						
studies provided enough detail to determine whether the differences in the point estimates between the treatment arms is statistically significant.	D	Level IV studies or Level I to III studies/SRs with a high risk of bias							
2. Consistency									
There is a significant degree of heterogeneity between the studies (Phet=0.03). When the studies are limited to those using a transfusion protocol the heterogeneity is no longer significant (Phet=0.09). The impact of	А	All studies consistent							
postoperative cell salvage is significant for all of the subgroups analysed in Carless 2006. The results of the	В	Most studies consistent and inconsistency can be explained							
RCTs published after Carless 2006 are consistent in direction.	С	Some inconsistency, reflecting genuine uncertainty around question	on						
	D	Evidence is inconsistent							
	NA	Not applicable (one study only)							
3. Clinical impact									
Carless 2006 All surgery types – mean difference -0.82 units (-1.12, -0.51); 9 trials (N=689)	А	Very large							
Cardiac surgery – mean difference -0.83 units (-1.25, -0.40); 7 trials (N=580)	В	Substantial							
Orthopaedic surgery – mean difference -0.80 units (-1.17, -0.43); 2 trials (N=109)	С	Moderate							
	D	Slight/Restricted							
4. Generalisability									
The studies were conducted in patients undergoing cardiac and orthopaedic surgery.	Α	Evidence directly generalisable to target population							
	В	Evidence directly generalisable to target population with some cav	reats						
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied						
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to						
5. Applicability									
All the studies were conducted in developed countries.	Α	Evidence directly applicable to Australian healthcare context							
	В	Evidence applicable to Australian healthcare context with few cave							
	С	Evidence probably applicable to Australian healthcare context with	n some caveats						
	D	Evidence not applicable to Australian healthcare context							

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EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description				
1. Evidence base	ence base C One good quality level I study (included studies were of fair quality)					
2. Consistency	В	Most studies consistent and inconsistency can be explained				
3. Clinical impact	С	Moderate impact in cardiac and orthopaedic surgery				
4. Generalisability	В	Evidence directly generalisable to target population with some caveats				
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats				

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery or total knee arthroplasty, postoperative cell salvage reduces the volume of allogeneic blood transfusion.

Abbreviations: RCT, randomised controlled trial.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hospital readmissi

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I4.P2 Characteristics and results of studies examining the effect of postoperative cell salvage on transfusion volume.

	Level of evidence	No. of trials /	Patient population / Surgical					Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Carless (2006)	Level I Good	9 trials (fair quality) N=689	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Mean difference (95% CI), units	-0.82 (-1.12, -0.51)		P<0.00001	Phet=0.03
Amin (2008)	Level II Fair	N=178	Adults undergoing unilateral TKA	Hospital in UK	Postoperative cell salvage: patients in the reinfusion group had their blood reinfused from drains within 6 hours of surgery.	Total units of allogeneic blood transfused	22	26	NR	
Cheng (2005)	Level II Fair	N=60	Adults undergoing TKA	Hospital in Hong Kong	Postoperative cell salvage: patients in the reinfusion group had their blood reinfused from drains within 6 hours of surgery.	Mean (SD) units of allogeneic blood transfused	0.15 (0 to 1)	0.46 (0 to 4)	P=0.033	
Zacharopoulos (2007)	Level II Poor	N=60	Adults undergoing TKA	The study was conducted in a small town of -8000 inhabitants in Greece	Postoperative cell salvage and reinfusion of washed blood within 6 hours of operation.	Median (IQR) units of allogeneic blood transfused	0.3 (NR)	1.5 (NR)	NR	
Cardiac surgery							•			
Carless (2006)	Level I Good	7 trials (fair quality) N=580	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Mean difference (95% CI), units	-0.83 (-1.25, -0.40)		P=0.0001	Phet=0.01
Orthopaedic sur	gery	•	1	•		•			<u>'</u>	•
Carless (2006)	Level I Good	2 trials (fair quality) N=109	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Mean difference (95% CI), units	-0.80 (-1.17, -0.43)		P<0.0001	Phet=1.00
Studies with a tr	ansfusion proto	ocol							•	
Carless (2006)	Level I Good	6 trials (fair quality) N=398	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Mean difference (95% CI), units	-0.75 (-1.02, -0.47)		P<0.00001	Phet=0.09
Studies without	a transfusion p	rotocol								
Carless (2006)	Level I Good	3 trials (fair quality) N=291	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Mean difference (95% CI), units	-1.64 (-2.96, -0.33)		P=0.01	Phet=0.05

Abbreviations: CI, confidence interval; IQR, interquartile range; NR, not reported; SD, standard deviation; TKA, total knee arthroplasty.

Key question(s): In patients undergoing surgery, what is the effect of <u>postoperative cell salvagents</u>	Evidence table ref*: POQ3.I4.P3					
1. Evidence base						
1 level I study: Carless 2006 (good quality; includes 8 RCTs ¹ ; N=555) 1 level II studies published after the Carless 2006 search date: Cheng 2005 (fair quality; N=60; patients	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias			
undergoing TKA)	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency	•					
The degree of heterogeneity within the trials in Carless 2006 is not significant (Phet=0.12). The lack of impact of postoperative cell salvage on blood loss is consistent across cardiac and orthopaedic surgery. Carless 2006	Α	All studies consistent in finding no difference				
reported total blood loss whilst Cheng 2005 reported operative blood loss.	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around question	on			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact						
Carless 2006 All surgery types – mean difference -56.97 mL (-152.05, 38.12); 8 trials (N=555)	Α	Very large				
Cardiac surgery – mean difference -85.04 mL (-212.50, 42.41); 5 trials (N=366)	В	Substantial				
Orthopaedic surgery – mean difference -21.74 mL (-164.51, 121.04); 2 trials (N=189) Cheng 2005	С	Moderate				
Median (IQR), cell salvage vs. control: 273 (100 to 600) vs. 280 (100 to 800); P=0.84	D	No difference				
4. Generalisability	•					
All the studies were conducted in patients undergoing cardiac surgery or orthopaedic surgery.	Α	Evidence directly generalisable to target population				
	В	Evidence directly generalisable to target population with some cav	eats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
		Evidence not directly generalisable to target population and hard to judge whether it is sensible to				
5. Applicability						
All the studies were conducted in developed countries.	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few cave				
	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
	D	Evidence not applicable to Australian healthcare context				

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EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	One good quality level I study (included studies were of fair quality) and one subsequently published fair quality level II study.
2. Consistency	Α	All studies consistent in finding no difference.
3. Clinical impact	D	No statistically significant impact.
4. Generalisability	В	Evidence directly generalisable to target population with some caveats.
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery or total knee arthroplasty, postoperative cell salvage does not appear to have an effect on total blood loss.

Abbreviations: RCT, randomised controlled trial; TKA, Total knee arthroplasty.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hosp

POQ3.I4.P1 Characteristics and results of studies examining the effect of postoperative cell salvage on blood loss.

	Level of	No. of trials / Patient population / Surgical		Results			Notes			
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Carless (2006)	Level I Good	8 trials (fair quality) N=555	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Total blood loss, mL Mean difference (95% CI)	-56.97 (-152.05, 38.	12)	P=0.24	Phet=0.12
Cheng (2005)	Level II Fair	N=60	Adults undergoing TKA	Hospital in Hong Kong	Postoperative cell salvage: patients in the reinfusion group had their blood reinfused from drains within 6 hours of surgery.	Operative blood loss, mL Median (IQR)	273 (100 to 600)	280 (100 to 800)	P=0.84	
Cardiac surgery										
Carless (2006)	Level I Good	5 trials (fair quality) N=366	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Total blood loss, mL Mean difference (95% CI)	-85.04 (-212.50, 42.	41)	P=0.19	Phet=0.03
Orthopaedic surg	gery	•							•	
Carless (2006)	Level I Good	3 trials (fair quality) N=189	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Total blood loss, mL Mean difference (95% CI)	-21.74 (-164.51, 12	1.04)	P=0.77	Phet=0.81

Abbreviations: CI, confidence interval; IQR, interquartile range; TKA, total knee arthroplasty.

Key question(s): In patients undergoing surgery, what is the effect of postoperative cell salvage.	Evidence table ref*: POQ3.I4.P4					
1. Evidence base						
1 level I study: Carless 2006 (good quality; includes 5 RCTs all of fair quality; N=471). All the studies were in patients undergoing cardiac surgery.	Α	One or more level I studies with a low risk of bias or several level	I studies with a low risk of bias			
patients undergoing cardiac surgery.	В	One or two Level II studies with a low risk of bias or SR/several Lev	/el III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency	<u>I</u>					
There is no significant heterogeneity between the studies (Phet=0.92). All of the studies are consistent in finding that cell salvage had no significant impact.	Α	All studies consistent in finding no difference				
triat cell sarvage nau no significant impact.	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around question	on			
	D	Evidence is inconsistent				
	NA	NA Not applicable (one study only)				
3. Clinical impact						
RR 1.64 (0.52, 5.17)	Α	Very large				
	В	Substantial				
	С	Moderate				
	D	No difference/underpowered				
4. Generalisability All of the five studies that constant metality as an extreme were conducted in national undergoing condicated.	Ι ,	Fridan distribution of the state of the stat				
All of the five studies that reported mortality as an outcome were conducted in patients undergoing cardiac surgery.	A	Evidence directly generalisable to target population				
	В	Evidence directly generalisable to target population with some cave				
	С	Evidence not directly generalisable to the target population but co	,			
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to			
5. Applicability						
All of the studies were conducted in developed countries.	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few cave				
	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
	D	Evidence not applicable to Australian healthcare context				

Other factors

Included studies were underpowered to detect a mortality difference. Transfusion is a confounder for this outcome.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	One good quality level I study (included studies were of fair quality).
2. Consistency	Α	All studies consistent in finding no difference.
3. Clinical impact	D	No statistically significant impact.
4. Generalisability	В	Evidence not directly generalisable to the target population but could be sensibly applied. Only includes patients undergoing cardiac surgery.
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, the effect of postoperative cell salvage on mortality is uncertain.

Abbreviations: RCT, randomised controlled trial; RR, relative risk.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{* &}lt;u>Interventions</u>: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I4.P4 Characteristics and results of studies examining the effect of postoperative cell salvage on mortality.

	Level of evidence	No. of trials /	Patient population / Surgical			Outcome	Results				Notes
Study evidence Quality		sample size	procedure	Setting	Intervention		Intervention	Comparator	p-value		
Carless (2006)	Level I Good	5 trials N=471	Adults undergoing any elective surgery. All of the 5 studies that reported mortality as an outcome were conducted in patients undergoing cardiac surgery.	All studies conducted in developed countries	Postoperative cell salvage	Mortality	8/246 (3%)	4/225 (2%)	P=0.40	Phet=0.92	

Key question(s):			Evidence table ref*:			
In patients undergoing surgery, what is the effect of postoperative cell salvage	<u>e</u> on_	morbidity?	POQ3.I4.P5			
1. Evidence base						
1 level I study: Carless (good quality) – infection (5 RCTs; 3 cardiac and 2 orthopaedic; N=429); wound complication (6 RCTs; 4 cardiac and 2 orthopaedic; N=404); any thrombosis (4 RCTs; all orthopaedic; N=240);	Α	One or more level I studies with a low risk of bias or several level I	studies with a low risk of bias			
stroke (1 RCT; cardiac; N=30); non-fatal MI (2 RCTs; both cardiac; N=144); DVT (3 RCTs; all orthopaedic;	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias			
N=210). All the RCTs were fair quality 2 level II studies published after the Carless 2006 search date:	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias			
Amin 2008 (orthopaedic; fair quality; N=178) – wound infection; infections other than wound infection; DVT; persistent wound drainage (no infection) Cheng 2005 (orthopaedic; fair quality; N=60) – febrile complications	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency	I	l				
The heterogeneity within trials in Carless 2006 is not significant. The lack of impact of postoperative cell salvage	Α	All studies consistent in finding no difference.				
is consistent across patients undergoing cardiac surgery and orthopaedic surgery.	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around question	n			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact						
Carless 2006 Infection: RR 0.60 (0.17, 2.15); wound complication: RR 0.84 (0.37, 1.92); any thrombosis: RR 1.41 (0.43, 4.57);	Α	Very large				
stroke: RR 3.00 (0.13, 68.26); non-fatal MI: RR 0.85 (0.25, 2.93); DVT: RR 0.64 (0.15, 2.66)	В	Substantial				
Amin 2008 Wound infection: RR 1.40 (0.24, 8.19); infections other than wound infection: RR 0.93 (0.13, 6.49); DVT: RR 0.47 (0.04, 5.06); persistent wound drainage (no infection): RR 1.87 (0.17, 20.25)	С	Moderate				
(0.04, 5.06); persistent wound drainage (no infection): RR 1.87 (0.17, 20.25) Cheng 2005	D	No difference/underpowered				
Febrile complications: RR 2.62 (0.25, 27.30)						
4. Generalisability						
All the studies were conducted in patients undergoing cardiac surgery or orthopaedic surgery.	Α	Evidence directly generalisable to target population				
	В	Evidence directly generalisable to target population with some cav	eats			
	С	Evidence not directly generalisable to the target population but cou	ıld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to	judge whether it is sensible to			
5. Applicability						
All of the studies were conducted in developed countries.	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few cave	ats			
	С	Evidence probably applicable to Australian healthcare context with	some caveats			
	D	Evidence not applicable to Australian healthcare context				
	•					

Other factors

CRG considered washed vs. unwashed to be important for this outcome. Carless 2006 did not report washed vs. unwashed for morbidity.

Carless 2006 included several older studies. The CRG noted that cell salvage technology has changed over time.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	One good quality level I study (included studies were of fair quality) and two subsequently published fair quality level II studies.
2. Consistency	Α	All studies consistent in finding no difference
3. Clinical impact	D	No difference/underpowered
4. Generalisability	В	Evidence directly generalisable to target population with some caveats.
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery or total knee arthroplasty, postoperative cell salvage does not appear to have an effect on morbidity, including infection.

Abbreviations: DVT, deep vein thrombosis; MI, myocardial infarction; RCT, randomised controlled trial; RR, relative risk.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I4.P5 Characteristics and results of studies examining the effect of postoperative cell salvage on morbidity.

0	Level of evidence	No. of trials /	Patient population / Surgical	0 111				Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Carless (2006)	Level I Good	5 trials (fair quality) N=429	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Infection	7/210 (3%)	17/219 (8%)	P=0.43	Phet=0.26
		6 trials (fair quality) N=404	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Wound complication	11/213 (5%)	11/191 (6%)	P=0.69	Phet=0.63
		4 trials (fair quality) N=240	Adults undergoing any elective surgery.	All studies conducted in developed countries	Postoperative cell salvage	Any thrombosis	6/120 (5%)	4/120 (3%)	P=0.57	Phet=0.83
		1 trial (fair quality) N=30	Adults undergoing any elective surgery.	All studies conducted in developed countries	Postoperative cell salvage	Stroke	1/15 (7%)	0/15 (0%)	P=0.49	Phet=NA
		2 trials (fair quality) N=144	Adults undergoing any elective surgery.	All studies conducted in developed countries	Postoperative cell salvage	Non-fatal MI	5/71 (7%)	6/73 (8%)	P=0.80	Phet=0.94
		3 trials (fair quality) N=210	Adults undergoing any elective surgery.	All studies conducted in developed countries	Postoperative cell salvage	DVT	3/105 (3%)	5/105 (5%)	P=0.54	Phet=0.46
Amin (2008)	Level II Fair	T//A		Hospital in UK	Postoperative cell salvage: patients in the reinfusion group had their blood	Wound infection	3/92 (3%)	2/86 (2%)	P=0.71	
				reinfused from drains within 6 hours of surgery.	Infections other than wound infection	2/92 (2%)	2/86 (2%)	P=0.95		
						DVT	1/92 (1%)	2/86 (2%)	P=0.53	
				Persistent wound drainage (no infection)	2/92 (2%)	1/86 (1%)	P=0.61			
Cheng (2005)	Level II Fair	N=60	Adults undergoing TKA	Hospital in Hong Kong	Postoperative cell salvage: patients in the reinfusion group had their blood reinfused from drains within 6 hours of surgery.	Febrile complications	2/26 (8%)	1/34 (3%)	P=0.403	

Study	Level of evidence	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcomo		Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Cardiac surgery	,									
Carless (2006)	Level I Good	3 trials (fair quality) N=259	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Infection	4/125 (3%)	13/134 (10%)	P=0.53	Phet=0.14
		4 trials (fair quality) N=264	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Wound complication	6/143 (4%)	5/121 (4%)	P=0.88	Phet=0.33
		1 trial (fair quality) N=30	Adults undergoing any elective surgery. The only trial that reported incidence of stroke as an outcome was in patients undergoing cardiac surgery.	All studies conducted in developed countries	Postoperative cell salvage	Stroke	1/15 (7%)	0/15 (0%)	P=0.49	Phet=NA
		2 trials (fair quality) N=144	Adults undergoing any elective surgery. All of the trials that reported incidence of non-fatal MI as an outcome were in patients undergoing cardiac surgery.	All studies conducted in developed countries	Postoperative cell salvage	Non-fatal MI	5/71 (7%)	6/73 (8%)	P=0.80	Phet=0.94
Orthopaedic sur	rgery									
Carless (2006)	Level I Good	2 trials (fair quality) N=170	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Infection	3/85 (4%)	4/85 (5%)	P=0.78	Phet=0.28
		2 trials (fair quality) N=140	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Wound complication	5/70 (7%)	6/70 (9%)	P=0.73	Phet=0.88
		4 trials (fair quality) N=240	Adults undergoing any elective surgery. All of the trials that reported incidence of thrombosis as an outcome were in patients undergoing orthopaedic surgery.	All studies conducted in developed countries	Postoperative cell salvage	Any thrombosis	6/120 (5%)	4/120 (3%)	P=0.57	Phet=0.83
		3 trials (fair quality) N=210	Adults undergoing any elective surgery. All of the trials that reported incidence of DVT as an outcome were in patients undergoing orthopaedic surgery.	All studies conducted in developed countries	Postoperative cell salvage	DVT	3/105 (3%)	5/105 (5%)	P=0.54	Phet=0.46

Abbreviations: DVT, deep vein thrombosis; MI, myocardial infarction; NA, not applicable; TKA, total knee arthroplasty.

Key question(s): In patients undergoing surgery, what is the effect of postoperative cell salvage	<u>e</u> on	quality of life?	Evidence table ref*: POQ3.I4.P5				
1. Evidence base							
	А	One or more level I studies with a low risk of bias or several level I	risk of bias or several level II studies with a low risk of bias				
	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias				
	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias					
2. Consistency							
	Α	All studies consistent					
	В	Most studies consistent and inconsistency can be explained					
	С	Some inconsistency, reflecting genuine uncertainty around question	on				
	D	Evidence is inconsistent					
	NA	Not applicable (one study only)					
3. Clinical impact							
	Α	Very large					
	В	Substantial	ntial				
	С	Moderate					
	D	Slight/Restricted					
4. Generalisability	•						
-	Α	Evidence directly generalisable to target population					
	В	Evidence directly generalisable to target population with some cav	reats				
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied				
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to				
5. Applicability							
	Α	Evidence directly applicable to Australian healthcare context					
	В	Evidence applicable to Australian healthcare context with few cave	eats				
	С	Evidence probably applicable to Australian healthcare context with	n some caveats				
	D	Evidence not applicable to Australian healthcare context					

Other factors		
EVIDENCE STA	TEMENT	MATRIX
Please summarise t	the develop	ment group's synthesis of the evidence relating to the key question, taking all the above factors into account.
Component	Rating	Description
Evidence base	NA	
2. Consistency	NA	
3. Clinical impact	NA	
4. Generalisability	NA	
5. Applicability	NA	
DRAFT EVIDEN	CE STA	TEMENT
Based on the body	of evidence	e above.
In adult pa	atients un	dergoing surgery in which substantial blood loss is anticipated, the effect of postoperative cell salvage on quality of life is unknown.

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Key question(s):			Evidence table ref*:			
In patients undergoing surgery, what is the effect of postoperative cell salvage	<u>e</u> on_	<u>change in haemoglobin concentration?</u>	POQ3.I4.S1			
1. Evidence base						
3 level II studies: Amin 2008 (orthopaedic; fair quality; N=178); Cheng 2005 (orthopaedic; fair quality; N=60); Zacharopoulos 2007 (orthopaedic; poor quality; N=60)	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias				
Zacitatopodios 2007 (ortitopacato, poor quality, N=00)	В	One or two Level II studies with a low risk of bias or SR/several Level	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	•			
2. Consistency	ı					
All three RCTs are consistent in finding that postoperative cell salvage had no significant impact on change in Hb concentration.	Α	All studies consistent				
Concentration.	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact						
Amin 2008 – mean (SD), treatment vs. control: 2.2 (0.7) vs. 2.6 (0.8); P=0.354 Cheng 2005 – Median (IQR): 101 (84 to 128) vs. 104 (87 vs. 137); P=0.332	Α	Very large				
Zacharopoulos 2007 – No significant difference (no more detail provided)	В	Substantial				
	С	Moderate				
	D	No difference				
4. Generalisability						
All three studies were conducted in patients undergoing TKA.	Α	Evidence directly generalisable to target population				
	В	Evidence directly generalisable to target population with some cave	veats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to			
5. Applicability	•					
Amin 2008 was conducted in the UK, Cheng 2005 was conducted in Hong Kong, and Zacharopoulos 2007 was conducted in a small town in Greece with approximately 8000 inhabitants. The exclusion of Zacharopoulos 2007	Α	Evidence directly applicable to Australian healthcare context				
does not influence the outcome.	В	Evidence applicable to Australian healthcare context with few caveats				
	С	Evidence probably applicable to Australian healthcare context with some caveats				
	D	Evidence not applicable to Australian healthcare context				

ner factors

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	Three fair-to-poor quality level II studies.
2. Consistency	Α	All studies consistent.
3. Clinical impact	D	No statistically significant impact.
4. Generalisability	Α	Evidence directly generalisable to target population
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing total knee arthroplasty, the effect of postoperative cell salvage on haemoglobin concentration is uncertain.

Abbreviations: Hb, haemoglobin; IQR, interquartile range; RCT, randomised controlled trial; SD, standard deviation; TKA, total knee arthroplasty.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I4.S1 Characteristics and results of studies examining the effect of postoperative cell salvage on change in haemoglobin.

	Level of evidence	No. of trials /	Patient population / Surgical			_		Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Amin (2008)	Level II Fair	N=178	Adults undergoing unilateral TKA	Hospital in UK	Postoperative cell salvage: patients in the reinfusion group had their blood reinfused from drains within 6 hours of surgery.	Mean (SD) change in Hb concentration (pre- vs. postoperative 24, 48, 72 hr), g/dL	2.2 (0.7)	2.6 (0.8)	P=0.354	
Cheng (2005)	Level II Fair	N=60	Adults undergoing TKA	Hospital in Hong Kong	Postoperative cell salvage: patients in the reinfusion group had their blood reinfused from drains within 6 hours of surgery.	Mean (range) haemoglobin level immediately postoperative, g/dL	101 (84 to 128)	104 (87 to 137)	P=0.332	
						Mean (range) haemoglobin level 3 days postoperative, g/dL	98 (77 to 130)	101 (77 to 130)	P=0.401	
Zacharopoulos (2007)	Level II Poor	N=60	Adults undergoing TKA	The study was conducted in a small town of ~8000 inhabitants in Greece	Postoperative cell salvage and reinfusion of washed blood within 6 hours of operation.	Mean (SD) change in Hb concentration (pre- vs. postoperative Day 1, 5, 15)	NR	NR	NS	

Abbreviations: Hb, haemoglobin concentration; NR, not reported; NS, not significant; SD, standard deviation: TKA, total knee arthroplasty.

Key question(s): In patients undergoing surgery, what is the effect of postoperative cell salve	reoperation for bleeding?	Evidence table ref*: POQ3.I4.S2			
1. Evidence base					
1 level I study: Carless 2006 (good quality; includes 6 RCTs all fair quality, all cardiac surgery; N=374)	А	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias		
No Level II evidence published after Carless 2006.	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias		
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias		
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	3		
2. Consistency	ı				
The degree of heterogeneity within the trials in Carless 2006 is not significant (Phet=0.54).	Α	All studies consistent in finding no difference			
	В	Most studies consistent and inconsistency can be explained			
	С	Some inconsistency, reflecting genuine uncertainty around questi	on		
	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact	'				
Carless 2006 – RR 1.41 (0.53, 3.78)	А	Very large			
	В	Substantial			
	С	Moderate			
	D	No difference/underpowered			
4. Generalisability					
All the trials in Carless 2006 were in patients undergoing cardiac surgery.	А	Evidence directly generalisable to target population			
	В	Evidence directly generalisable to target population with some call	/eats		
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied		
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to		
5. Applicability					
All of the studies were conducted in developed countries.	Α	Evidence directly applicable to Australian healthcare context			
	В	Evidence applicable to Australian healthcare context with few cav			
	С	Evidence probably applicable to Australian healthcare context wi	th some caveats		
	D	Evidence not applicable to Australian healthcare context			

Other factors		
Other factors		
Other factors		

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
Evidence base	С	One good quality level I study (included studies were of fair quality) and one subsequently published fair quality level II study.
2. Consistency	Α	All studies consistent in finding no difference
3. Clinical impact	D	No statistically significant impact.
4. Generalisability	В	Evidence directly generalisable to target population with some caveats.
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, the effect of postoperative cell salvage on risk of reoperation for bleeding is uncertain.

Abbreviations: RCT, randomised controlled trial: RR, relative risk: TKA, total knee arthroplasty.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I4.S2 Characteristics and results of studies examining the effect of postoperative cell salvage on reoperation for bleeding.

Study	Level of evidence Quality	No. of trials / sample size	Patient population / Surgical procedure	Setting In	Intervention Outcome		Notes			
						Outcome	Intervention	Comparator	p-value	
Carless (2006)	Level I Good	6 trials (fair quality) N=374	Adults undergoing any elective surgery. All of the trials that reported reoperation for bleeding as an outcome were conducted in patients undergoing cardiac surgery.	All studies conducted in developed countries	Postoperative cell salvage	Patients who underwent reoperation for bleeding	11/193 (6%)	6/181 (3%)	P=0.50	Phet=0.54

Abbreviations: TKA, total knee arthroplasty.

Key question(s): In patients undergoing surgery, what is the effect of <u>postoperative cell salvagents</u>	length of hospital stay?	Evidence table ref*: POQ3.I4.S5				
1. Evidence base						
1 level I study: Carless 2006 (good quality; includes 4 RCTs; 3 cardiac and 1 orthopaedic (all fair quality); N=297 Level II study published after the Carless 2006 search date: Amin 2008 (fair quality; N=178)	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias				
Level II study published after the Carless 2000 Search date. Allin 2000 (fall quality, N=170)	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias				
	С	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency						
The degree of heterogeneity within the trials in Carless 2006 is not significant (Phet=0.11). The results in Carless 2006 are consistent across both cardiac and orthopaedic surgery. The results from Amin 2008 are not consistent	Α	All studies consistent				
with the results from Carless 2006.		Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around question				
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact						
Carless 2006 mean difference -1.72 days (-2.82, -0.62)	Α	Very large				
Amin 2008		Substantial				
Median (IQR), cell salvage: 6.6 days (3 to 14) vs. 7.0 days (3 to 16); P=0.54	С	Moderate				
		No difference				
4. Generalisability						
The trials in Carless 2006 were conducted in patients undergoing cardiac and patients undergoing orthopaedic surgery. Amin 2008 was conducted in patients undergoing TKA.		Evidence directly generalisable to target population				
		Evidence directly generalisable to target population with some caveats				
		Evidence not directly generalisable to the target population but could be sensibly applied				
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to			
5. Applicability	ı					
All the studies were conducted in developed countries.	A	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few cave				
	С	Evidence probably applicable to Australian healthcare context with some caveats				
	D	Evidence not applicable to Australian healthcare context				

Other factors

Applicability depends on regional practice (rehabilitation varies between institutions).

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	Three level II studies with a moderate risk of bias
2. Consistency	С	Some inconsistency, reflecting genuine uncertainty around question
3. Clinical impact	С	Moderate clinical impact
4. Generalisability	В	Evidence directly generalisable to target population with some caveats
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery and total knee arthroplasty, postoperative cell salvage may reduce length of hospital stay.

Abbreviations: IQR, interguartile range; RCT, randomised controlled trial; TKA, total knee arthroplasty.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I4.S5 Characteristics and results of studies examining the effect of postoperative cell salvage on length of hospital stay.

Study Level of evidence Ouality No. of trials / sample size Patient population / Surgical procedure Setting Intervention		No of trials /	Patient population / Surgical				Results			Notes
	Intervention	Outcome	Intervention	Comparator	p-value					
Carless (2006)	Level I Good	4 trials (fair quality) N=297	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Mean difference (95% CI), days	-1.72 (-2.82, -0.62)		P=0.002	Phet=0.11
Amin (2008)	Level II Fair	N=178	Adults undergoing unilateral TKA	Hospital in UK	Postoperative cell salvage: patients in the reinfusion group had their blood reinfused from drains within 6 hours of surgery.	Median (IQR) difference, days	6.6 (3 to 14)		7.0 (3 to 16)	P=0.54
Cardiac surgery										
Carless (2006)	Level I Good	3 trials (fair quality) N=227	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Mean difference (95% CI), days	-1.41 (-2.69, -0.13)		P=0.03	Phet=0.08
Orthopaedic surgery										
Carless (2006)	Level I Good	1 trial (fair quality) N=70	Adults undergoing any elective surgery	All studies conducted in developed countries	Postoperative cell salvage	Mean difference (95% CI), days	-2.60 (-4.76, -0.44)		P=0.02	Phet=NA

Abbreviations: CI, confidence interval; IQR, interquartile range; NA, not applicable; TKA, total knee arthroplasty.

Recommendation(s) for postoperative cell salvage

In adult patients undergoing cardiac surgery or total knee arthroplasty, in whom significant postoperative blood loss is anticipated, postoperative cell salvage should be considered. IMPLEMENTATION OF RECOMMENDATION Please indicate yes or no to the following questions. Where the answer is yes please provide explanatory information about this. This information will be used to develop the implementation plan for the guidelines. Will this recommendation result in changes in usual care? Postoperative cell salvage use will increase (not widely used at present).	PO3 PO3	E TABLE 3.14.P1, 3.14.P2, 3.14.P5, 3.14.S5
IMPLEMENTATION OF RECOMMENDATION Please indicate yes or no to the following questions. Where the answer is yes please provide explanatory information about this. This information will be used to develop the implementation plan for the guidelines. Will this recommendation result in changes in usual care? Postoperative cell salvage use will increase (not widely used at present).	PO3	3.I4.P2, 3.I4.P5,
IMPLEMENTATION OF RECOMMENDATION Please indicate yes or no to the following questions. Where the answer is yes please provide explanatory information about this. This information will be used to develop the implementation plan for the guidelines. Will this recommendation result in changes in usual care? Postoperative cell salvage use will increase (not widely used at present).	POS	3.I4.P5,
Please indicate yes or no to the following questions. Where the answer is yes please provide explanatory information about this. This information will be used to develop the implementation plan for the guidelines. Will this recommendation result in changes in usual care? Postoperative cell salvage use will increase (not widely used at present).		•
Please indicate yes or no to the following questions. Where the answer is yes please provide explanatory information about this. This information will be used to develop the implementation plan for the guidelines. Will this recommendation result in changes in usual care? Postoperative cell salvage use will increase (not widely used at present).		•
Please indicate yes or no to the following questions. Where the answer is yes please provide explanatory information about this. This information will be used to develop the implementation plan for the guidelines. Will this recommendation result in changes in usual care? Postoperative cell salvage use will increase (not widely used at present).		
This information will be used to develop the implementation plan for the guidelines. Will this recommendation result in changes in usual care? Postoperative cell salvage use will increase (not widely used at present).		
Will this recommendation result in changes in usual care? Postoperative cell salvage use will increase (not widely used at present).		
Postoperative cell salvage use will increase (not widely used at present).		
	YES	NO
Are there any resource implications associated with implementing this recommendation?		
Are there any resource implications associated with implementing this recommendations	YES	NO
Training and equipment costs.		
Will the implementation of this recommendation require changes in the way care is currently organised?	YES	NO
Changes in organisation of postoperative care which will have nursing resource implications in postoperative wards.		
Are the guideline development group aware of any barriers to the implementation of this recommendation	YES	NO
Initial capital investment; recurrent expenditure (but preservation of blood supply at ARCBS level, which results in cost-shifting from Commo	nonwea	alth to
State); only applicable when postoperative drainage utilised; may not be widely supported by orthopaedic surgeons.		
What could help to facilitate implementation of the recommendation?	YES	NO
Development of local policies for postoperative cell salvage; lobbying for funding of cell salvage device.		

Intervention 5 – Deliberate induced hypotension

Key question(s): In patients undergoing surgery, what is the effect of deliberate induced hypote	n on transfusion incidence?	Evidence table ref*: POQ3.I5.P1				
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)					
Five level II studies: 3 good quality RCTs, 2 fair quality RCTs.	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk or				
		One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of				
	С	One or two Level III studies with a low risk of bias or Level I or II st	rudies with a moderate risk of bias			
		Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
4 RCTs examined prostatectomy patients and reported a significant effect. Important to note that 3 of the RCTs were conducted at the same institution, not possible to ascertain if there was an overlap in study population.	Α	All studies consistent				
1 RCT examined lienorenal shunt surgery and did not observe a significant effect, however the sample size was	В	Most studies consistent and inconsistency can be explained				
small (N=18). Test of heterogeneity across the 5 RCTs was not significant (P=0.14).	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
	D	Evidence is inconsistent				
		Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>ur</u>	nknowr	n factor (not simply study quality or sample size) and thus the clinical impa	ct of the intervention could not be			
Meta-analysis of 5 RCTs revealed a risk ratio of 0.38 (95%CI 0.19, 0.75), P=0.005.	Α	Very large				
This shows that the incidence of blood transfusion was 62% lower in patients with induced hypotension.	В	Substantial				
	С	Moderate				
		Slight/Restricted				
4. Generalisability (How well does the body of evidence match the population and clinical set	tings b	eing targeted by the Guideline?)				
As 4 of the 5 studies examined patients undergoing prostatectomy. Consequently, the evidence is likely	Α	Evidence directly generalisable to target population				
generalisable to patients undergoing this surgical procedure.	В	Evidence directly generalisable to target population with some caveats				
	С	Evidence not directly generalisable to the target population but could be sensibly applied				
	D	Evidence not directly generalisable to target population and hard	to judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of healt	th services/delivery of care and cultural factors?)				
The 4 RCTs examining prostatectomy were conducted in Germany and Canada, these findings are likely applicable to Australia.	Α	Evidence directly applicable to Australian healthcare context				
The RCT examining Lienorenal shunt surgery was conducted in India, which limits its applicability in the	В	Evidence applicable to Australian healthcare context with few caveats				
Australian context.	С	Evidence probably applicable to Australian healthcare context with some caveats				
	D	Evidence not applicable to Australian healthcare context				

Three publications (Boldt et al., Piper et al. and Suttner et al.) were conducted at the same institution. However, based on information in the publications, it is not possible to determine if there was an overlap in the study populations. Consequently, lack of independence should be considered during the interpretation of the results from these three studies.

Given the concerns regarding generalisability and applicability, the recommendation was graded 'C'.

The Clinical/Consumer Reference Group (CRG) noted the study by Sood et al. had a small sample size, was conducted in India, and examined lienorenal shunt surgery. Consequently, the findings from this study were not considered by the CRG in assessing the effect of induced hypotension on transfusion incidence.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	В	Three RCTs of good quality and two of fair quality.
2. Consistency	В	Test of heterogeneity across the five RCTs was not significant.
3. Clinical impact	Α	Meta-analysis of the five RCTs revealed a risk ratio of 0.38 (95%CI 0.19, 0.75), P=0.005.
4. Generalisability	С	The evidence is likely generalisable to patients undergoing prostatectomy.
5. Applicability	С	The four studies that examined prostatectomy patients were conducted in Germany and Canada, as such the findings are likely applicable in the Australian context. The study that examined lienorenal shunt surgery was conducted in India, which limits the applicability of the evidence in Australia.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing radical prostatectomy, deliberate induced hypotension (mean arterial pressure 50–60 mmHg) reduces the incidence of allogeneic RBC transfusion.

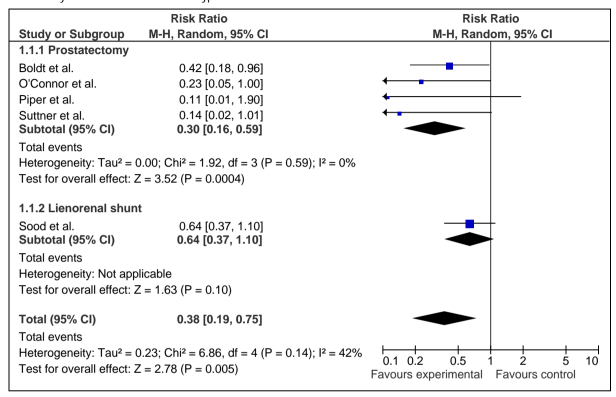
POQ3.I5.P1 Characteristics and results of studies examining the effect of <u>deliberate induced hypotension</u> on <u>transfusion incidence</u>

	Level of evidence	No. of trials	Patient population /		Internation	Outcomo	Results			
Study	Quality	(N)	Surgical procedure	Setting Intervention (Outcome	Intervention	Comparator	p-value	Notes
Prostatectomy		•								
O'Connor et al. (2006)	Level II	N=99	Patients with adenocarcinoma of the prostate to undergo radical retropubic prostatectomy.	Medical Institution in Canada.	Hypotension induced using epidural and ropivacaine (MAP 55-60mmHg)	Incidence of blood transfusion n/N (%)	2/49 (4%)	9/50 (18%)	0.028	
Piper et al. (2002)	Level II Fair	N=30	Patients undergoing elective radical prostatectomy (ASA class II and III only).	Hospital in Germany ^a	Controlled hypotension (MAP ~50mmHg) using sodium nitroprusside	Incidence of blood transfusion n/N (%)	0/15 (0%)	4/15 (27%)	<0.05	
Suttner et al. (2001)	Level II Good	N=28	Patients undergoing elective radical prostatectomy.	Hospital in Germany ^a	Controlled hypotension (MAP ~50mmHg) using sodium nitroprusside	Incidence of blood transfusion n/N (%)	1/14 (7%)	7/14 (50%)	<0.05	See meta-analysis of effect
Boldt et al. (1999)	Level II Good	N=40	Patients under the age of 75 years undergoing retropubic radical prostatectomy with bilateral pelvic lymphadenectomy.	Hospital in Germany ^a	Controlled hypotension (MAP ~50mmHg) using sodium nitroprusside	Incidence of blood transfusion n/N (%)	5/20 (25%)	12/20 (60%)	<0.05	
Lienorenal shunt	surgery								•	
Sood et al. (1987)	Level II Fair	N=18	Patients undergoing elective, proximal, lienorenal shunts for portal hypertension.	Hospital in India.	Controlled hypotension (Systolic BP 90-95mmHg) using sodium nitroprusside	Incidence of blood transfusion n/N (%)	5/8 (63%)	10/10 (100%)	NR	

Abbreviations: ASA, American Society of Anaesthesiologists; BP, blood pressure; MAP, mean arterial pressure; NR, not reported.

^a Three publications (Boldt et al., Piper et al. and Suttner et al.) were conducted at the same institution. However, based on information in the publications, it is not possible to determine if there was an overlap in the study populations. Consequently, lack of independence should be considered during the interpretation of the results from these three studies.

Meta-analysis of the effect of induced hypotension on the incidence of blood transfusion.



Meta-analysis shows that the incidence of blood transfusion was 62% lower in patients with induced hypotension, compared to patients with normotension.

Key question(s):			Evidence table ref*:			
In patients undergoing surgery, what is the effect of <u>deliberate induced hypot</u>	<u>ensio</u>	n on transfusion volume?	POQ3.I5.P2			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)					
One level I study: systematic review with good quality rating, low risk of bias. 7 level II studies: 3 good quality RCTs, 4 fair quality RCTs.	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias				
7 lever it studies. 3 good quality RC15, 4 fair quality RC15.	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias				
		One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
		Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Systematic review conducted a meta-analysis and found a significant reduction in transfusion volume in patients with induced hypotension. Test of heterogeneity was significant (P<0.05).	А	All studies consistent				
4 RCTs examined prostatectomy patients and reported a significant reduction in transfusion volume. Important to note that 3 of the RCTs were conducted at the same institution (see other factors). 2 RCTs examined patients undergoing hip arthroplasty, one observed a significant reduction. 1 RCT examined patients undergoing lienorenal shunt surgery and observe a significant reduction.	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>undetermined</u>)	nknowr	factor (not simply study quality or sample size) and thus the clinical impact	ct of the intervention could not be			
Meta-analysis: Systematic review estimated that in orthopaedic surgical patients, blood transfusion was 667mL (95%CI 370, 963) lower in patients with induced hypotension.	Α	Very large				
Use of different units and measurements made it difficult to synthesize a single effect estimate across the RCTs	В	Substantial				
identified.	С	Moderate				
	D	Slight/Restricted				
4. Generalisability (How well does the body of evidence match the population and clinical sea	ttings b	eing targeted by the Guideline?)				
The evidence is generalisable to patients undergoing major joint replacement surgery and prostatectomy.	Α	Evidence directly generalisable to target population				
	В	Evidence directly generalisable to target population with some cave	reats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to judge whether it is sensible to				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of healt					
Most of the studies included in the systematic review were conducted in Europe, US and Canada.	Α	Evidence directly applicable to Australian healthcare context				
The 2 RCTs examining hip arthroplasty were conducted in Turkey and Sweden, while the 4 RCTs examining prostatectomy were conducted in Germany and Canada.	В	Evidence applicable to Australian healthcare context with few cave	eats			
As such it is likely that these findings are applicable in the Australian context.	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
	D	Evidence not applicable to Australian healthcare context				

Three publications (Boldt et al., Piper et al. and Suttner et al.) were conducted at the same institution. However, based on information in the publications, it is not possible to determine if there was an overlap in the study populations. Consequently, lack of independence should be considered during the interpretation of the results from these three studies.

The Clinical/Consumer Reference Group (CRG) noted the study by Sood et al. had a small sample size, was conducted in India, and examined lienorenal shunt surgery. Consequently, the CRG did not consider the findings from this study in assessing the effect of induced hypotension on transfusion incidence.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	One systematic review of good quality. Three RCTs of good quality and four RCTs of fair quality.
2. Consistency	Α	Induced hypotension was found to significantly reduce transfusion volume in all studies except for one RCT, conducted in Turkey with a small sample size (N=20).
3. Clinical impact	В	Meta-analysis estimated that in major joint surgery, blood transfusion was 667mL lower in patients with induced hypotension.
4. Generalisability	В	The evidence is likely generalisable to patients undergoing major joint replacement surgery and prostatectomy.
5. Applicability	В	The studies identified were mostly conducted in developed western countries, similarly developed to Australia. As such, the findings are likely applicable in the Australian context.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing radical prostatectomy or major joint replacement, deliberate induced hypotension (MAP 50–60 mmHg) reduces the volume of allogeneic blood transfusion.

POQ3.I5.P2 Characteristics and results of studies examining the effect of <u>deliberate induced hypotension</u> on <u>transfusion volume</u>

0	Level of evidence	No. of trials	Patient population / Surgical	0				Results		Notes					
Study	Quality	(N)	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	Notes					
Orthopaedic surg	ery	•													
Paul et al. (2007)	Level I Good	6 (N=222)	Patients undergoing major joint replacement surgery	NR	Deliberate induced hypotension by any method	Blood transfused (mL) WMD (95%CI)	WMD: -667 ml	. (-963, -370)	NR	Search date: Up to Jan 2006					
Karakaya et al. (1999)	Level II Fair	N=20	ASA class I and II patients undergoing primary total hip arthroplasty	Medical Institution in Turkey	Nitroglycerine induced hypotension (MAP 60- 65mmhg)	Blood transfused per patient (Units) Mean (SD)	2.3 (0.8)	2.7 (1.1)	NS	Not included in Paul et al. as haemodilution was used concurrently					
Fredin et al.	Level II	N=57	Patients undergoing total hip	Hospital in	Controlled Hypoterision (3DI	Controlled Hypoterision (3D)	Controlled Hypoterision (3D)	CONTROLLED ITANOGENSION CODE	Controlled Hypotension (3DI	Controlled Hypotension (SDI		Intraop: 580 (380)	1210 (620)	<0.01	Not included in Paul et al. as all patients were also
(1984)	Fair	14-37	arthroplasty.	Sweden	nitroprusside	Mean (SD)	Total: 920 (580)	1540 (1050)	<0.01	given blood thinners					
Prostatectomy															
O'Connor et al. (2006)	Level II Good	N=99	Patients with adenocarcinoma of the prostate to undergo radical retropubic prostatectomy.	Medical Institution in Canada.	Hypotension induced using epidural and ropivacaine (MAP 55-60mmHg)	Total volume of RBC transfused (Units)	3	24	NR	-					
Piper et al. (2002)	Level II Fair	N=30	Patients undergoing elective radical prostatectomy (ASA class II and III only).	Hospital in Germany ^a	Controlled hypotension (MAP ~50mmHg) using sodium nitroprusside	Total volume of RBC transfused (Units)	0	10	P<0.05	-					
Suttner et al. (2001)	Level II Good	N=28	Patients undergoing elective radical prostatectomy.	Hospital in Germany ^a	Controlled hypotension (MAP ~50mmHg) using sodium nitroprusside	Total volume of RBC transfused (Units)	3	17	P<0.05	-					
Boldt et al. (1999)	Level II Good	N=40	Patients under the age of 75 years undergoing retropubic radical prostatectomy with bilateral pelvic lymphadenectomy.	Hospital in Germany ^a	Controlled hypotension (MAP ~50mmHg) using sodium nitroprusside	Total volume of RBC transfused (Units)	14	28	P<0.05	-					

	Level of evidence	No. of trials	Patient population / Surgical					Results			
Study	Quality	(N)	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	Notes	
Lienorenal shunt :	surgery										
Sood et al. (1987)	Level II Fair	N=18	Patients undergoing elective, proximal, lienorenal shunts for portal hypertension.	Hospital in India	Controlled hypotension (SBP 90-95mmHg) using sodium nitroprusside	Blood transfused per patient (Units) Mean (SD)	0.88 (0.9)	3.0 (1.2)	P<0.01	-	

Abbreviations: ASA, American Society of Anaesthesiologists; BP, blood pressure; CI, confidence interval; MAP, mean arterial pressure; NR, not reported; NS, not statistically significant; RBC, red blood cells; SD, standard deviation; WMD, weighted mean difference.

^a Three publications (Boldt et al., Piper et al. and Suttner et al.) were conducted at the same institution. However, based on information in the publications, it is not possible to determine if there was an overlap in the study populations. Consequently, lack of independence should be considered during the interpretation of the results from these three studies.

Key question(s):			Evidence table ref*:			
In patients undergoing surgery, what is the effect of <u>deliberate induced hypote</u>	ensio	n on <u>blood loss</u> ?	POQ3.I5.P3			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)					
One level I study: 1 systematic review with good quality rating. The systematic review reported non-significant bias (Egger's test P=0.955).	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias				
Nine level II studies: 5 good quality RCTs, 4 fair quality RCTs.	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias				
		One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
The test of heterogeneity conducted by the systematic review was significant; this suggests that there may be differences between surgical methods and the methods of inducing hypotension.	Α	All studies consistent				
	В	Most studies consistent and inconsistency can be explained				
All RCTs, except 1 (Jacobi et al.), reported a significant reduction in blood loss among patients with induced hypotension. The study by Jacobi et al. examined patients undergoing endoscopic sinus surgery and had a small sample size (N=32), which may have contributed to the lack of a significant finding in the study.	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>ur</u>	nknown	factor (not simply study quality or sample size) and thus the clinical impac	ct of the intervention could not be			
Systematic review by Paul et al. estimated that in orthopaedic surgical patients, blood loss was lower by 286mL (95%CI 127, 447) in patients with induced hypotension.	Α	Very large				
(93%CF127, 447) in patients with induced hypotension.	В	Substantial				
8 RCTs provided sufficient data for meta-analysis, which showed that induced hypotension reduced blood loss by an average of 460mL (95%CI 210.9, 709.8), P=0.0003.	С	Moderate				
	D	Slight/Restricted				
4. Generalisability (How well does the body of evidence match the population and clinical set	tings be	eing targeted by the Guideline?)				
The evidence is generalisable to patients undergoing orthopaedic surgery and prostatectomy. While the evidence is likely generalisable to all surgical procedures, the effect (reduction in blood loss) of the intervention would likely	Α	Evidence directly generalisable to target population				
vary.	В	Evidence directly generalisable to target population with some cave	reats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
		Evidence not directly generalisable to target population and hard to judge whether it is sensible to				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of healt	h services/delivery of care and cultural factors?)				
The systematic review included studies conducted mostly in Europe, US and Canada.	Α	Evidence directly applicable to Australian healthcare context				
The RCTs were conducted in Germany, Canada, Sweden, the Netherlands, Egypt and India. As such it is likely that these findings are applicable in the Australian context.	В	Evidence applicable to Australian healthcare context with few caveats				
ů	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
	D	Evidence not applicable to Australian healthcare context				

Three publications (Boldt et al., Piper et al. and Suttner et al.) were conducted at the same institution. However, based on information in the publications, it is not possible to determine if there was an overlap in the study populations. Consequently, lack of independence should be considered during the interpretation of the results from these three studies.

The Clinical/Consumer Reference Group noted the study by Sood et al. had a small sample size, was conducted in India and examined lienorenal shunt surgery. Consequently, the findings from this study were not considered in assessing the effect of induced hypotension on transfusion incidence.

The CRG also noted that the studies by Elsharnouby et al. and Jacobi et al. examined blood loss during endoscopic sinus surgery. In these studies, the importance of blood loss is related more to the obstruction of surgical field rather than to issues relating to blood transfusion requirements. Consequently, less emphasis has been placed on the findings from these studies.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	One good quality systematic review. Five RCTs of good quality and four RCTs of fair quality.
2. Consistency	Α	All except one study showed that induced hypotension significantly reduced blood loss.
3. Clinical impact	В	The systematic review showed that induced hypotension reduces blood loss by 286mL (95%Cl 127, 447) during orthopaedic surgery. Meta-analysis of the RCTs showed that induced hypotension reduced blood loss by 460mL (95%Cl 210, 709).
4. Generalisability	В	The evidence is likely generalisable to patients undergoing major joint replacement surgery, breast reduction surgery and prostatectomy.
5. Applicability	В	The studies were mostly conducted in developed western countries, as such the evidence is likely applicable in the Australian context.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing radical prostatectomy, major joint replacement or breast reduction surgery, deliberate induced hypotension (MAP 50–60 mmHg) reduces the volume of blood loss.

POQ3.I5.P3 Characteristics and results of studies examining the effect of <u>deliberate induced hypotension</u> on <u>blood loss</u>

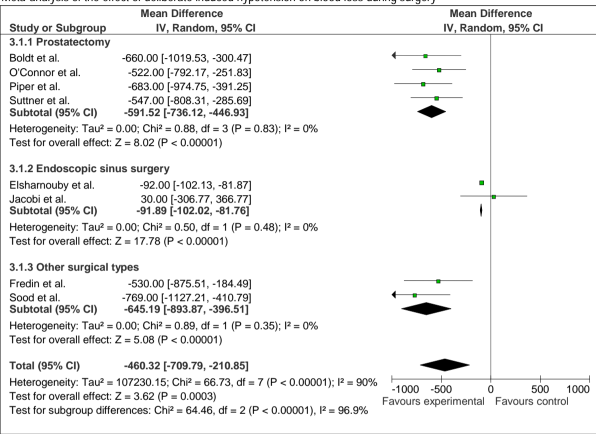
	Level of evidence	No. of trials	Patient population /					Results		
Study	Quality	(N)	Surgical procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	Notes
Orthopaedic surg	gery									
Paul et al. (2007)	Level I Good	17 (N=586)	Patients undergoing major joint replacement surgery.	NR	Deliberate induced hypotension by any method	Blood loss (mL) WMD (95%CI)	WMD: -286	(-447, -127)	NR	Search date: Up to Jan 2006
Fredin et al.	et al. Level II Patients undergoing total Hospital in Controlled hypotension (SBP Blo		Blood loss (mL)	Intraop: 620 (240)	1070 (630)	<0.001	Not included in Paul et al. as all patients were also			
(1984)	Fair	N=57	hip arthroplasty.	Sweden.	70-80mmHg) using sodium nitroprusside	Mean (SD)	Total: 1170 (395)	1700 (860)	<0.01	given blood thinners
Prostatectomy										
O'Connor et al. (2006)	Level II Good	N=99	Patients with adenocarcinoma of the prostate to undergo radical retropubic prostatectomy.	Medical Institution in Canada.	Hypotension induced using epidural and ropivacaine (MAP 55-60mmHg)	Blood loss (mL) Mean (SD)	955 (517)	1477 (823)	<0.001	
Piper et al. (2002)	Level II Fair	N=30	Patients undergoing elective radical prostatectomy (ASA class II and III only).	Hospital in Germany ^a	Controlled hypotension (MAP ~50mmHg) using sodium nitroprusside	Blood loss (mL) Mean (SD)	843 (233)	1526 (409)	<0.05	Con mate analysis
Suttner et al. (2001)	Level II Good	N=28	Patients undergoing elective radical prostatectomy.	Hospital in Germany ^a	Controlled hypotension (MAP ~50mmHg) using sodium nitroprusside	Blood loss (mL) Mean (SD)	788 (193)	1335 (460)	<0.05	See meta-analysis
Boldt et al. (1999)	Level II Good	N=40	Patients under the age of 75 years undergoing retropubic radical prostatectomy with bilateral pelvic lymphadenectomy.	Hospital in Germany ^a	Controlled hypotension (MAP ~50mmHg) using sodium nitroprusside	Intraoperative and Total Blood loss (mL)	1260 (570)	1920 (590)	<0.05	

	Level of evidence	No. of trials	Patient population /		Intervention	_			Notes	
Study	Quality	(N)	Surgical procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	Notes
Endoscopic sinus	Endoscopic sinus surgery									
Elsharnouby et al. (2006)	Level II Good	N=60	Patients undergoing functional endoscopic sinus surgery.	Hospital in Egypt.	Controlled hypotension (MAP 50-60mmHg) using magnesium sulphate	Blood loss (mL) Mean (SD)	165 (19)	257 (21)	<0.05	See meta-analysis
Jacobi et al. (2000)	Level II Fair	N=32	Patients undergoing endoscopic sinus surgery.	Hospital in Germany.	Controlled hypotension (MAP 65-75mmHg) using sodium nitroprusside	Blood loss (mL) Mean (SD)	278 (528)	245 (440)	NS	
Other surgical pr	rocedures									
Kop et al. (2009)	Level II Good	N=85	Patients (<60 years, ASA I and II) undergoing bilateral breast reduction surgery.	Hospital in the Netherlands.	Controlled hypotension (MAP >50mmHg) using sodium nitroprusside	Blood loss (mL) Mean (range)	316 (133–560)	598 (250–1335)	<0.001	
Sood et al. (1987)	Level II Fair	N=18	Patients undergoing elective, proximal, lienorenal shunts for portal hypertension.	Hospital in India.	Controlled hypotension (SBP 90-95mmHg) using sodium nitroprusside	Blood loss (mL) Mean (SD)	517 (220)	1286 (523)	<0.01	See meta-analysis

Abbreviations: ASA, American Society of Anaesthesiologists; BP, blood pressure; MAP, mean arterial pressure; NR, not reported; NS, not statistically significant; SD, standard deviation.

^a Three publications (Boldt et al., Piper et al. and Suttner et al.) were conducted at the same institution. However, based on information in the publications, it is not possible to determine if there was an overlap in the study populations. Consequently, lack of independence should be considered during the interpretation of the results from these three studies.

Meta-analysis of the effect of deliberate induced hypotension on blood loss during surgery



Meta-analysis shows that patients with induced hypotension on average lose 460mL less blood. Test of heterogeneity showed that the volume of blood loss varied significantly between surgical procedures.

Key question(s): In patients undergoing surgery, what is the effect of deliberate induced hypot	Evidence table ref*: POQ3.15.P4					
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)					
One level II study of good quality.	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias				
	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias				
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
NA	Α	All studies consistent				
		Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>undetermined</u>)	nknown	factor (not simply study quality or sample size) and thus the clinical impac	ct of the intervention could not be			
No deaths occurred in either patient group.	Α	Very large				
	В	Substantial				
	С	Moderate				
	D	No difference/underpowered				
4. Generalisability (How well does the body of evidence match the population and clinical sea	ttings be	eing targeted by the Guideline?)				
This study was conducted in patients undergoing prostatectomy, as such, the generalisability of the evidence would be limited to patients undergoing this surgical procedure.	Α	Evidence directly generalisable to target population				
would be limited to patients differential this surgical procedure.	В	Evidence directly generalisable to target population with some cave	reats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of healt					
The RCT was conducted in Canada, as such, the evidence is likely applicable to Australia.	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few cave				
	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
	D	Evidence not applicable to Australian healthcare context				

Other factors (Indicate here any other factors that you took into account when assessing the evidence base (for example, issues that might cause the group to downgrade or upgrade the recommendation)							
EVIDENCE STA	TEMENT	T MATRIX					
Please summarise ti	he develop	oment group's synthesis of the evidence relating to the key question, taking all the above factors into account.					
Component	Rating	Description					
1. Evidence base	В	One good quality RCT					
2. Consistency	NA	Not applicable					
3. Clinical impact	D	No deaths occurred in either patient group.					
4. Generalisability	С	This study was conducted in patients undergoing prostatectomy, as such, the evidence would be most generalisable to patients undergoing this surgical procedure.					
5. Applicability	В	This study was conducted in Canada, as such, the evidence is likely applicable to Australia.					
DRAFT EVIDEN	CE STA	TEMENT					
Based on the body of evidence above.							
In adult nations, undergoing surgery in which substantial blood loss is anticipated, the effect of deliberate induced hypotension (MAP 50–60 mmHg) on mortality is							

uncertain.

POQ3.I5.P4 Characteristics and results of studies examining the effect of <u>deliberate induced hypotension</u> on <u>mortality</u>

Study	Level of evidence	No. of trials	S Patient population / Surgical procedure		Intervention		Results			
	Quality	(N)		Setting		Outcome	Intervention	Comparator	p-value	Notes
O'Connor et al. (2006)	Level II Good	N=99	Patients with adenocarcinoma of the prostate to undergo radical retropubic prostatectomy.	Medical Institution in Canada.	Hypotension induced using epidural and ropivacaine (MAP 55-60mmHg)	Serious adverse events	0	0	NA	Includes death, myocardial infarction, stroke, renal impairment, DVT, PE

Abbreviations: DVT, deep vein thrombosis; NA, not applicable; PE, pulmonary embolism.

Key question(s):			Evidence table ref*:				
In patients undergoing surgery, what is the effect of <u>deliberate induced hypot</u>	<u>ensio</u>	<u>n</u> on <u>morbidity</u> ?	POQ3.I5.P5				
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)						
Three level II studies: 2 good quality RCTs, 1 fair quality RCT.	Α	One or more level I studies with a low risk of bias or several level	Il studies with a low risk of bias				
	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias				
		One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias					
2. Consistency (If only one study was available, rank this component as 'not applicable')							
Several different morbidity outcomes were examined. None of the RCTs observed a significant difference between treatment groups for the morbidity outcomes	Α	All studies consistent					
examined (serious adverse events, incidence of DVT and PE).	В	Most studies consistent and inconsistency can be explained					
	С	Some inconsistency, reflecting genuine uncertainty around question					
	D	Evidence is inconsistent					
	NA	Not applicable (one study only)					
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u>	nknown	factor (not simply study quality or sample size) and thus the clinical impac	ct of the intervention could not be				
No significant difference between patient groups	Α	Very large					
	В	Substantial					
	С	Moderate					
	D	No difference/underpowered					
4. Generalisability (How well does the body of evidence match the population and clinical sea	ttings be	eing targeted by the Guideline?)					
The generalisability of the evidence is likely limited to the morbidity outcome in the specific surgical patient populations examined by the respective studies.	Α	Evidence directly generalisable to target population					
populations examined by the respective studies.	В	Evidence directly generalisable to target population with some call	reats				
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied				
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of healt						
The RCTs were conducted in Germany, Canada and Sweden. As such the findings from these studies are likely applicable to Australia.	Α	Evidence directly applicable to Australian healthcare context					
applicable to haditalia.	В	Evidence applicable to Australian healthcare context with few cave					
	С	Evidence probably applicable to Australian healthcare context with	n some caveats				
	D	Evidence not applicable to Australian healthcare context					

The Clinical/Consumer Reference Group noted that the study by Fredin et al. was conducted in 1983; as such differences in surgical practices and patient management, relating to pulmonary embolism, may reduce the applicability of the study.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	В	Two RCTs of good quality and one RCT of fair quality.
2. Consistency	В	Several different morbidity outcomes were examined. None of the studies observed a significant effect of induced hypotension on the incidence of morbid events.
3. Clinical impact	D	No significant difference between patient groups. Underpowered.
4. Generalisability	С	The generalisability of the evidence is likely limited to the morbidity outcome in the specific surgical patient populations examined by the respective studies.
5. Applicability	С	The RCTs were conducted in Germany, Canada and Sweden. As such the findings from these studies are likely applicable to Australia.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of deliberate induced hypotension (MAP 50–60 mmHg) on morbidity is uncertain.

Abbreviations: DVT, deep vein thrombosis; PE, pulmonary embolism; RCT, randomised clinical trial.

POQ3.I5.P5 Characteristics and results of studies examining the effect of <u>deliberate induced hypotension</u> on <u>morbidity</u>

Study	Level of evidence	No. of trials	Patient population / Surgical procedure							
	Quality	(N)		Setting	Intervention	Outcome	Intervention	Comparator	p-value	Notes
O'Connor et al. (2006)	Level II	N=99	Patients with adenocarcinoma of the prostate to undergo radical retropubic prostatectomy.	Medical Institution in Canada.	Hypotension induced using epidural and ropivacaine (MAP 55-60mmHg)	Serious adverse events	0	0	NA	Includes death, myocardial infarction, stroke, renal impairment, DVT, PE
Fredin et al.	Level II	N 57	Patients undergoing total		Controlled hypotension (SBP	Incidence of DVT n/N (%)	11/24 (46%)	10/26 (38%)	NS	-
(1984)	Fair	N=57		Sweden.	70-80mmHg) using sodium nitroprusside	Incidence of PE n/N (%)	6/26 (26%)	1/28 (4%)	NS	-

Abbreviations: DVT, deep vein thrombosis; NA, not applicable; NR, not reported; NS, not statistically significant; PE, pulmonary embolism.

Key question(s):		W 6 N6 O	Evidence table ref*: POQ3.I5.P6				
In patients undergoing surgery, what is the effect of <u>deliberate induced hypote</u>	<u>ensioi</u>	on quality of life?	POQ3.13.P0				
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies)						
No studies identified in literature search	Α	One or more level I studies with a low risk of bias or several level	more level I studies with a low risk of bias or several level II studies with a low risk of bias				
	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias				
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias					
2. Consistency (If only one study was available, rank this component as 'not applicable')	•						
NA	Α	All studies consistent					
	В	Most studies consistent and inconsistency can be explained					
	С	Some inconsistency, reflecting genuine uncertainty around questi	on				
	D	Evidence is inconsistent					
	NA	Not applicable (one study only)					
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>ur</u>	nknown	factor (not simply study quality or sample size) and thus the clinical impac	ct of the intervention could not be				
NA NA	Α	Very large					
	В	Substantial					
	С	Moderate					
	D	Slight/Restricted					
4. Generalisability (How well does the body of evidence match the population and clinical set	tings be	eing targeted by the Guideline?)					
NA	Α	Evidence directly generalisable to target population					
	В	Evidence directly generalisable to target population with some cave	reats				
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied				
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of healt						
NA	Α	Evidence directly applicable to Australian healthcare context					
	В	Evidence applicable to Australian healthcare context with few cave					
	С	Evidence probably applicable to Australian healthcare context with	n some caveats				
	D	Evidence not applicable to Australian healthcare context					

Other factors (Ind	dicate here a	any other factors that you took into account when assessing the evidence base (for example, issues that might cause the group to downgrade or upgrade the recommendation)
EVIDENCE STA	TEMENT	MATRIX
Please summarise t	the develop	oment group's synthesis of the evidence relating to the key question, taking all the above factors into account.
Component	Rating	Description
Evidence base	NA	
2. Consistency	NA	
3. Clinical impact	NA	
4. Generalisability	NA	
5. Applicability	NA	
DRAFT EVIDEN	ICE STA	ΓΕΜΕΝΤ
Based on the body	of evidence	e above.
In adult patie unknown.	nts under	going surgery in which substantial blood loss is anticipated, the effect of deliberate induced hypotension (MAP 50–60 mmHg) on quality of life is

Key question(s):			Evidence table ref*:			
In patients undergoing surgery, what is the effect of <u>deliberate induced hypote</u>	ensior	on <u>haemoglobin concentration</u> ?	POQ3.I5.S1			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies)					
Two level II studies: both of fair quality.	Α	One or more level I studies with a low risk of bias or several level	I studies with a low risk of bias			
	В	One or two Level II studies with a low risk of bias or SR/several Level	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
The findings of the two RCTs differed. Piper et al. examined patients undergoing prostatectomy and found that patients with induced hypotension had	Α	All studies consistent				
significantly higher haemoglobin concentration during and after surgery.	В	Most studies consistent and inconsistency can be explained				
Krakaya et al. examined patients undergoing hip arthroplasty and reported that there was no significant difference in haemoglobin concentration between patient groups.	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
ano en es maemogosm con es manon son con panon y cape.	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>ur</u>	known	factor (not simply study quality or sample size) and thus the clinical impac	ct of the intervention could not be			
	Α	Very large				
	В	Substantial				
	С	Moderate	pact of the intervention could not be			
	D	Conflicting evidence				
4. Generalisability (How well does the body of evidence match the population and clinical set	tings be	eing targeted by the Guideline?)				
The effects observed are likely generalisable to patients undergoing the surgical procedures examined in each of the respective studies.	Α	Evidence directly generalisable to target population				
the respective studies.	В	Evidence directly generalisable to target population with some cave	reats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of healti					
Piper et al. was conducted in Germany, as such the findings are likely applicable in the Australian context.	Α	Evidence directly applicable to Australian healthcare context				
Karakaya et al. was conducted in Turkey. Additional information on the healthcare system would allow an	В	Evidence applicable to Australian healthcare context with few cave				
assessment of the applicability of findings.	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
	D	Evidence not applicable to Australian healthcare context				

The Clinical/Consumer Reference Group noted that the measure of haemoglobin concentration was a surrogate for blood loss. Please see POQ3.15.P3 for the evidence on blood loss.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
Evidence base	С	Two RCTs of fair quality.
2. Consistency	D	The studies had inconsistent findings.
3. Clinical impact	D	Only one study observed a significant effect. The haemoglobin concentrations in patients with induced hypotension appeared to be ~1.5g/dL higher, compared to patients with normotension.
4. Generalisability	С	The evidence is likely generalisable to patients undergoing the surgical procedures examined in each of the respective studies.
5. Applicability	С	The study by Piper et al. was conducted in Germany, as such the evidence is probably relevant in Australia. In contrast, the study by Karakaya et al. was conducted in Turkey, which may limit the applicability of the evidence.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of deliberate induced hypotension (MAP 50–60 mmHg) on haemoglobin concentration is uncertain.

POQ3.I5.S1 Characteristics and results of studies examining the effect of <u>deliberate induced hypotension</u> on <u>haemoglobin concentration</u>

	Level of evidence	No. of trials (N)	Patient population /							
Study	Quality		Surgical procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	Notes
Piper et al. (2002)	Level II Fair	N=30	Patients undergoing elective radical prostatectomy (ASA class II and III only).	Hospital in Germany.	Controlled hypotension (MAP ~50mmHg) using sodium nitroprusside	Postoperative haemoglobin concentration	Significantly higher in intervention group (see figure below)		P<0.05	-
							After intubation: 11.6 (0.4)	11.9 (0.8)		
Karakaya et al. (1999)	Level II Fair	N=20 undergoi	ASA class I and II patients undergoing primary total hip arthroplasty Medical Insti in Turkey.	in Turkey hypotension (MAP 60-65mmha)	Haemoglobin concentrations (g/dL)	After operation: 9.2 (0.19)	9.7 (0.2)	NS		
						(3, 4-)	After 5 days: 10.2 (0.3)	10.3 (0.5)		

Abbreviations: ASA, American Society of Anaesthesiologists; MAP, mean arterial pressure; NS, not statistically significant.

Figure from Piper et al. 2002.

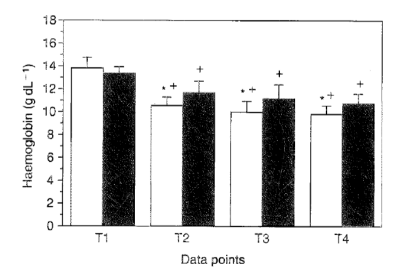


Figure 1. Changes in haemoglobin concentration with time. T1: after induction of general anaesthesia; T2: at the end of surgery; T3: 2 h postoperatively; T4: 24 h postoperatively. Data are the mean \pm SD. *P < 0.05 different versus the other group. +P < 0.05 different versus the baseline. \Box : Control group; : hypotension group.

Key question(s): In patients undergoing surgery, what is the effect of deliberate induced hypoteness.	n on coagulation status?	Evidence table ref*: POQ3.I5.S3					
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)						
One good quality RCT.	Α	One or more level I studies with a low risk of bias or several level	Il studies with a low risk of bias				
	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias				
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	;				
2. Consistency (If only one study was available, rank this component as 'not applicable')							
NA	Α	All studies consistent					
	В	Most studies consistent and inconsistency can be explained					
	С	Some inconsistency, reflecting genuine uncertainty around questi	on				
	D	Evidence is inconsistent					
	NA	Not applicable (one study only)					
3. Clinical impact (Indicate in the space below if the study results varied according to some undetermined)	nknown	factor (not simply study quality or sample size) and thus the clinical impac	ct of the intervention could not be				
No significant difference in coagulation data (aPTT, AT III, fibrinogen, platelet count) were observed between patient groups	Α	Very large					
patient groups	В	Substantial					
	С	Moderate					
	D	No difference					
4. Generalisability (How well does the body of evidence match the population and clinical set	ttings be	eing targeted by the Guideline?)					
The generalisability of the evidence is likely limited to patients undergoing prostatectomy.	Α	Evidence directly generalisable to target population					
	В	Evidence directly generalisable to target population with some cave	reats				
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied				
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of healt						
The study was conducted in Germany, as such the findings are likely applicable to Australia.	Α	Evidence directly applicable to Australian healthcare context					
	В	Evidence applicable to Australian healthcare context with few cave					
	С	Evidence probably applicable to Australian healthcare context with	n some caveats				
	D	Evidence not applicable to Australian healthcare context					

The Clinical/Consumer Reference Group noted that the study by Fredin et al. was conducted in 1983, as such differences in surgical practices and patient management, relating to pulmonary embolism, may reduce the applicability of the study.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	В	One RCT of good quality
2. Consistency	NA	Only one included study.
3. Clinical impact	D	No significant difference between patient groups.
4. Generalisability	С	The generalisability of the evidence is likely to patients undergoing prostatectomy.
5. Applicability	С	The study was conducted in Germany, as such the findings are likely applicable to Australia.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of deliberate induced hypotension (MAP 50–60 mmHg) on coagulation status is uncertain.

Abbreviations: aPTT, activated partial thromboplastin time; AT, antithrombin III; RCT, randomised clinical trial.

POQ3.I5.S3 Characteristics and results of studies examining the effect of <u>deliberate induced hypotension</u> on <u>coagulation status</u>

Study	Level of evidence	No. of trials (N)	Patient population / Surgical procedure			Outcome				
	Quality			Setting	Intervention		Intervention	Comparator	p-value	Notes
(1000)			Patients under the age of 75 years undergoing retropubic radical prostatectomy with bilateral pelvic lymphadenectomy.	Hospital in Germany.	Controlled hypotension (MAP ~50mmHg) using sodium nitroprusside	Coagulation status	Pre-op: 34.1 (2.7) Post-op: 42.3 (5.4)	Pre-op: 34.3 (2.3) Post-op: 52.2 (12.1)	NS	-
	Level II	N. 40					Pre-op: 78.7 (5.5) Post-op: 58.7 (4.3)	Pre-op: 81.5 (7.8) Post-op: 60.1 (12.1)	NS	
	Good	nod N=40					Pre-op: 308 (39) Post-op: 181 (37)	Pre-op: 318 (44) Post-op: 145 (22)	NS	
							Pre-op: 209 (30) Post-op: 166 (35)	Pre-op: 221 (36) Post-op: 119 (33)	NS	

Abbreviations: aPTT, activated partial thromboplastin time; AT, antithrombin III; NS, not statistically significant.

Key question(s): In patients undergoing surgery, what is the effect of deliberate induced hypot	Evidence table ref*: POQ3.I5.S5						
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies)							
One level II study, of good quality was identified. The study examined patients undergoing prostatectomy.	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias					
		One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias					
	С	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias					
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	6				
2. Consistency (If only one study was available, rank this component as 'not applicable')							
NA	Α	All studies consistent					
	В	Most studies consistent and inconsistency can be explained					
		Some inconsistency, reflecting genuine uncertainty around question					
		Evidence is inconsistent					
	NA	Not applicable (one study only)					
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>undetermined</u>)	nknown	factor (not simply study quality or sample size) and thus the clinical impa	ct of the intervention could not be				
There was no significant difference in the number of patients who stayed in hospital for more than 5 days between treatment groups.	Α	Very large					
between treatment groups.	В	Substantial					
	С	Moderate					
	D	Slight/Restricted					
4. Generalisability (How well does the body of evidence match the population and clinical sea	ttings be	eing targeted by the Guideline?)					
The effects are likely generalisable to only patients undergoing prostatectomy.	Α	Evidence directly generalisable to target population					
	В	Evidence directly generalisable to target population with some case	veats				
	С	Evidence not directly generalisable to the target population but could be sensibly applied					
	D	Evidence not directly generalisable to target population and hard	to judge whether it is sensible to				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of healt						
This study was conducted in Canada, as such the findings are likely applicable in the Australian context.	Α	Evidence directly applicable to Australian healthcare context					
	В	Evidence applicable to Australian healthcare context with few caveats					
	С	Evidence probably applicable to Australian healthcare context with some caveats					
	D	Evidence not applicable to Australian healthcare context					

The CRG noted that the use of epidural in the study by O'Connor may have affected the hospital length of stay (eg, due to reduced mobility). Consequently, the findings may not be relevant for the purposes of this review.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	В	One RCT of good quality.
2. Consistency	NA	Not applicable
3. Clinical impact	D	There was no significant difference between patient groups.
4. Generalisability	С	The effects are likely generalisable to only patients undergoing prostatectomy.
5. Applicability	В	This study was conducted in Canada, as such the findings are likely applicable in the Australian context.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of deliberate induced hypotension (MAP 50–60 mmHg) on length of hospital stay is uncertain.

POQ3.I5.S5 Characteristics and results of studies examining the effect of <u>deliberate induced hypotension</u> on <u>hospital length of stay</u>

	Level of evidence No. of trials Patient population / Softing Intervention Outcome			Results						
Study Evidence	Surgical procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	Notes		
O'Connor et al. (2006)	Level II Good	N=99	Patients with adenocarcinoma of the prostate to undergo radical retropubic prostatectomy.	Medical Institution in Canada.	Hypotension induced using epidural and ropivacaine (MAP 55-60mmHg)	Hospital stay > 5 days n/N (%)	24/49 (49%)	34/50 (68%)	P=0.055	-

Abbreviations: MAP, mean arterial pressure.

Recommendation(s) for deliberate induced hypotension

RECOMMENDATION	GRADE	RELE	EVANT	
What recommendation(s) does the guideline development group draw from this evidence? Use action statements where possible.		EVIDENC	CE TABLE	
In patients undergoing radical prostatectomy or major joint replacement, if substantial blood loss is anticipated,	С	PC)3.I5.P1,	
deliberate induced hypotension (mean arterial blood pressure 50–60 mmHg) should be considered, balancing	I	PO3.I5.P2,		
the risk of blood loss and the preservation of vital organ perfusion.		PC	PO3.I5.P3	
IMPLEMENTATION OF RECOMMENDATION				
Please indicate yes or no to the following questions. Where the answer is yes please provide explanatory information about this. This information will be used to develop the implementation plan for the guidelines.				
Will this recommendation result in changes in usual care?		YES	NO	
Are there any resource implications associated with implementing this recommendation?		YES	NO	
Will the implementation of this recommendation require changes in the way care is currently organised?		YES	NO	
will the implementation of this recommendation require changes in the way care is currently organised:		1 113	IVO	
Are the guideline development group aware of any barriers to the implementation of this recommendation		YES	NO	
What could help to facilitate implementation of the recommendation?		YES	NO	

Intervention 6 – Prevention of hypothermia

Key question(s):	Evidence table ref*:					
In patients undergoing surgery, what is the effect of the prevention of hypothe	POQ3.I6.P1					
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies)						
Two level I studies: 1 good quality (Rajagopalan et al. 2008), 1 fair quality (Scott et al. 2006). Publication bias was assessed in the review by Rajagopalan et al. and found to be low.		One or more level I studies with a low risk of bias or several level II studies with a low risk of bias				
One level II study: One RCT was identified and considered to be of fair quality (Yau et al. 1992).	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias				
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Test of heterogeneity across the 10 RCTs in the review by Rajagopalan et al. was not significant (P=0.25).	Α	All studies consistent				
Both Rajagopalan et al. and Scott et al. observed a significant reduction in transfusion incidence. It is important note that two of the three studies included by Scott et al. for this outcome, were also included in the review by	В	Most studies consistent and inconsistency can be explained				
Rajagopalan et al. The level II study showed a non-significant effect. The study had a small sample size (N=20) and examined the	С	Some inconsistency, reflecting genuine uncertainty around question				
incidence of red blood cell transfusion rather than blood transfusion, these factors may have contributed to the inconsistent finding.	D	Evidence is inconsistent				
inconsistent intuing.		Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>unknown</u> factor (not simply study quality or sample size) and thus the clinical impact of the intervention could not be determined)						
Meta-analysis of the level I and II studies revealed a risk ratio of 0.78 (0.63, 0.96), P=0.021). This indicates that	Α	Very large				
the overall incidence of transfusion is 22% lower in patients when hypothermia prevention strategies are used.	В	Substantial				
	С	Moderate				
	D	Slight/Restricted				
4. Generalisability (How well does the body of evidence match the population and clinical set	tings be	eing targeted by the Guideline?)				
The systematic reviews by Rajagopalan et al. and Scott et al. included studies which examined patients undergoing hip surgery, cardiac surgery, abdominal and colorectal surgery. As such, the evidence is likely	Α	Evidence directly generalisable to target population				
generalisable to a broad range of surgical procedures.	В	Evidence directly generalisable to target population with some caveats				
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of healt	h services/delivery of care and cultural factors?)				
The studies included in the systematic review by Rajagopalan et al. were mainly conducted in Europe and the US, while the RCT was conducted in Canada. As such, the evidence is likely applicable in the Australian	Α	Evidence directly applicable to Australian healthcare context				
context.	В	Evidence applicable to Australian healthcare context with few caveats				
	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
	D	Evidence not applicable to Australian healthcare context				

The Clinical/Consumer Reference Group suggested caution in the interpretation of the results from the study by Yau et al. due to the method of hypothermia prevention used (warming of systemic perfusion) and the small sample size (N=20).

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
Evidence base	Α	1 good and 1 fair systematic review that comprised 11 RCTs from a range of surgical procedures
2. Consistency	В	The results from the studies were largely consistent.
3. Clinical impact	В	Meta-analysis of all the studies identified revealed a significant reduction in transfusion incidence, with the use of hypothermia prevention strategies (RR 0.78, 95%CI 0.63, 0.93)
4. Generalisability	В	The studies included patients from a variety of surgical procedures and should be generalisable to a broad patient population.
5. Applicability	В	Studies were conducted in countries with a comparable healthcare system to Australia.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

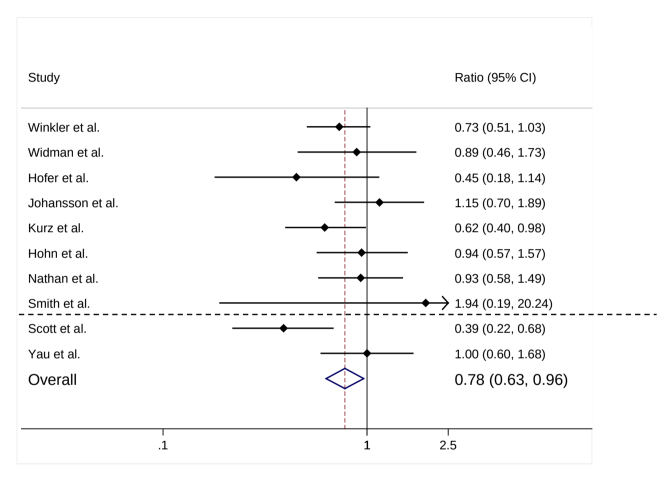
In adult patients undergoing surgery in which substantial blood loss is anticipated, the prevention of hypothermia reduces the incidence of transfusion.

Characteristics and results of studies examining the effect of the <u>prevention of hypothermia</u> during surgery on <u>transfusion incidence</u> POQ3.I6.P1

	Level of evidence No. of trials / Patient population / Surgical Setting Intervention Out										
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	Notes	
Rajagopalan et al. (2008)	Level I Good	10 (N=895)	Patients undergoing any surgical procedure	NR	Maintenance of normothermia, compared to patients with non-induced mild hypothermia	Ratio for need of blood transfusion (intervention vs control)	0.78 (0.63, 0.97)		P=0.027	Search date: 1996 to Oct 2006 Test of heterogeneity (P=0.25). Low publication bias observed.	
Scott et al. (2006)	Level I Fair	3 (N=250) ^a	Patients undergoing any surgical procedure (except cardiac procedures) under regional or general anaesthesia.	NR	Maintenance of normothermia, compared to patients with non-induced hypothermia	Ratio for need of blood transfusion (intervention vs control)	0.39 (0.22	2, 0.68)	NR	Search date: 1948 to May 2003 No test of heterogeneity or bias conducted.	
Yau et al. (1992)	Level II Fair	N=20	Patients undergoing isolated primary CABG.	Hospital in Canada	Warm systemic perfusion	Red blood cell transfusion incidence n/N (%)	6/8 (75)	9/12 (75)	NS	1	

Abbreviations: CABG, coronary artery bypass graft; NR, not reported; NS, not statistically significant; SD, standard deviation. ^a Two of the three studies identified by Scott et al. were also included in the review by Rajagopalan et al.

Meta-analysis of effect of the prevention of hypothermia on transfusion incidence



Studies included in Rajagopalan et al. (excluding those included in Scott et al.)

Meta-analysis (random effects model used) of the two Level I studies and one RCTs revealed a risk ratio of 0.78 (95% CI 0.63, 0.96), P=0.021. This indicates that the incidence of transfusion was 22% lower in patients when hypothermia prevention strategies were used.

Key question(s): In patients undergoing surgery, what is the effect of the prevention of hypothe	Evidence table ref*: POQ3.16.P2						
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies)							
One level I study: 1 poor quality review (Mahoney et al. 1999) Two level II studies: 1 fair quality (Zhao et al. 2005), 1 poor quality (Jeong et al. 2008)		One or more level I studies with a low risk of bias or several level II studies with a low risk of bias					
Two lever it studies: 1 fall quality (zhao et al. 2005), 1 pool quality (seong et al. 2006)	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias					
		One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias					
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	5				
2. Consistency (If only one study was available, rank this component as 'not applicable')							
No test of heterogeneity was conducted by the systematic review.	Α	All studies consistent					
Although a significant reduction in transfusion volume was observed in the systematic review, the two other	В	Most studies consistent and inconsistency can be explained					
RCTs (from South Korea and China) failed to detect a significant effect. This may be due to differences in the surgical procedures examined, or the small sample sizes of the RCTs identified (N=40 each).	С	Some inconsistency, reflecting genuine uncertainty around question					
	D	Evidence is inconsistent					
	NA	Not applicable (one study only)					
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>ur</u>	nknown	n factor (not simply study quality or sample size) and thus the clinical impa	ct of the intervention could not be				
The systematic review reported that the prevention of hypothermia significantly reduced transfusion volume of red blood cells (0.12 Units vs 1.17 Units, P<0.05).	Α	Very large					
	В	Substantial					
The Level II studies reported that the prevention of hypothermia actually increased transfusion volume, although the difference was not statistically significant.	С	Moderate					
, ,	D	Slight/Restricted					
4. Generalisability (How well does the body of evidence match the population and clinical set	tings be						
The systematic review identified five studies which examined RBC transfusion, this comprised patients from any surgical procedure. As such, the results is likely generalisable to a general surgical patient population.	Α	Evidence directly generalisable to target population					
The two RCTs identified examined patients undergoing CABG and abdominal surgery; consequently, the evidence	В	Evidence directly generalisable to target population with some care	veats				
is likely limited to patients undergoing such surgical procedures.	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied				
	D	Evidence not directly generalisable to target population and hard apply	to judge whether it is sensible to				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of healt						
The systematic review included studies mainly conducted in western countries, and as such, the evidence is likely applicable in Australia.	Α	Evidence directly applicable to Australian healthcare context					
In contrast, the two RCTs were conducted in Korea and China. Consequently, the applicability of the evidence	В						
may be more limited.		Evidence probably applicable to Australian healthcare context with some caveats					
	D	Evidence not applicable to Australian healthcare context					

The five studies identified in the systematic review included one non-randomised study (N=262), which could potentially affect the accuracy of the pooled effect estimate.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	1 poor quality systematic review. 1 poor and 1 fair quality RCT.
2. Consistency	С	Significant effect reported in systematic review, but not in 2 other RCTs. Differences in study design, sample size and population make it difficult to rationalise the inconsistency.
3. Clinical impact	D	A significant difference in transfusion volume was observed in the systematic review. However, the inclusion of a non-randomised study diminished confidence in the findings of the study.
4. Generalisability	С	The studies identified comprise patients undergoing a variety of surgery.
5. Applicability	С	The systematic review comprised studies conducted mostly in western countries, which are similarly developed to Australia. As such, the evidence is likely applicable in the Australian context. The two RCTs were conducted in South Korea and China, which may limit the applicability of the findings from those studies.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the prevention of hypothermia may reduce the volume of transfusion.

Abbreviations: CABG, coronary artery bypass graft; RBC, red blood cell.

POQ3.l6.P2 Characteristics and results of studies examining the effect of the <u>prevention of hypothermia</u> during surgery on <u>transfusion volume</u>

	Level of evidence	No. of trials /	Patient population /				Results				
Study	Quality	sample size	Surgical procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	Notes	
Mahoney et al. (1999)	Level I Poor	5 (N=859) a	Patients undergoing any surgical procedure	NR	Maintenance of normothermia, compared to patients with non-induced hypothermia	Units of RBC transfused Mean (SD)	0.12 (0.02)	1.17 (0.09)	P<0.05	Search Date: 1989 to 1997 No test of heterogeneity or bias conducted.	
Jeong et al. (2008)	Level II Poor	N=40	Patients undergoing isolated off-pump coronary artery bypass surgery.	Hospital in South Korea	Warming of all intravenous fluids	Blood transfused (mL) Mean (SD)	400.5 (622.8)	365.0 (437.1)	NS	-	
Zhao et al. (2005)	Level II	N=40	ASA class I and II patients undergoing abdominal	Hospital in	Warming from a forced-air blanket and warming of all intravenous	RBC transfused (Units) Mean (SD)	2.6 (2.5)	1.6 (2.4)	NS		
(2003)	Fair	N=40	surgery lasting at least 2 hours.	China	fluids.	Plasma transfused (mL) Mean (SD)	220 (460)	240 (480)	NS	_	

Abbreviations: ASA, American Society of Anaesthesiologists; CABG, coronary artery bypass graft; CI, confidence interval; NR, not reported; NS, not statistically significant; RBC, red blood cells; SD, standard deviation.

^a Includes one non-randomised controlled trial (N=262).

Key question(s):		Evidence table ref*:				
In patients undergoing surgery, what is the effect of the prevention of hypothermia on blood loss? POQ3.16.P3						
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)					
One level I study: The systematic review by Rajagopalan et al. (2008) was considered to be of good quality. Publication bias was assessed using a funnel plot and determined to be low.	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias			
Two level II studies: 2 RCTs of fair quality (Zhao et al. 2005; Yau et al. 1992).	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
		Level IV studies or Level I to III studies/SRs with a high risk of bias	3			
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Test of heterogeneity across the 14 RCTs in the review by Rajagopalan et al. was significant (P<0.001). Accordingly, a random effects model was used in the meta-analysis.	Α	All studies consistent				
The meta-analysis conducted by Rajagopalan et al. showed a significant reduction in blood loss.	В	Most studies consistent and inconsistency can be explained				
None of the level II studies showed a significant effect on blood loss. The variability of the observed effects may be due to the different hypothermia prevention methods employed,	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
and the different surgical procedures examined by the different studies.	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>ur</u>	nknown	factor (not simply study quality or sample size) and thus the clinical impac	ct of the intervention could not be			
A meta-analysis of the two Level II studies, together with the 14 studies included in the systematic review by Rajagopalan 2008 was conducted. The meta-analysis estimate indicates that the use of hypothermia prevention	Α	Very large				
strategies resulted in a 14% lower average blood loss in patients as compared to patients where hypothermia	В	Substantial				
was not prevented (Ratio: 0.86 (95%CI 0.76, 0.98), P=0.021).	С	Moderate				
	D	Slight/Restricted				
4. Generalisability (How well does the body of evidence match the population and clinical set	tings be	eing targeted by the Guideline?)				
The level I and II studies identified include studies which examined patients undergoing a variety of surgical procedures. As such, the results are likely to be generalisable to a broad range of surgical patient populations.	Α	Evidence directly generalisable to target population				
procedures. As such, the results are likely to be generalisable to a broad range of surgical patient populations.	В	Evidence directly generalisable to target population with some call	veats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to judge whether it is sensible to				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of healti	h services/delivery of care and cultural factors?)				
The studies included in the systematic review by Rajagopalan et al. were mainly conducted in Europe and the US, consequently, the findings are likely applicable in the Australian context.	Α	Evidence directly applicable to Australian healthcare context				
One of the RCTs was conducted in Canada, and is also likely applicable in the Australian context.	В	Evidence applicable to Australian healthcare context with few cave	eats			
The other RCT was conducted in China; as such, the evidence from these studies may not be as applicable in Australia.	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
Australia.		Evidence not applicable to Australian healthcare context				

The Clinical/Consumer Reference Group suggested caution in the interpretation of the results from the study by Yau et al. due to the method of hypothermia prevention used (warming of systemic perfusion) and the small sample size (N=20).

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
Evidence base	Α	1 good quality systematic review. 2 fair quality RCTs.
2. Consistency	С	Some variability between studies was observed, this may be due to the different hypothermia prevention methods used and different surgical procedures examined.
3. Clinical impact	В	A meta-analysis of the level I and II studies showed that blood loss was reduced by 14% when hypothermia was prevented (Ratio: 0.86 (95%CI 0.76, 0.98)).
4. Generalisability	В	The level I and II studies identified include studies which examined patients undergoing a variety of surgical procedures.
5. Applicability	В	The systematic review included studies mainly conducted in Europe and the US, as such the evidence is likely applicable in Australia.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the prevention of hypothermia reduces blood loss.

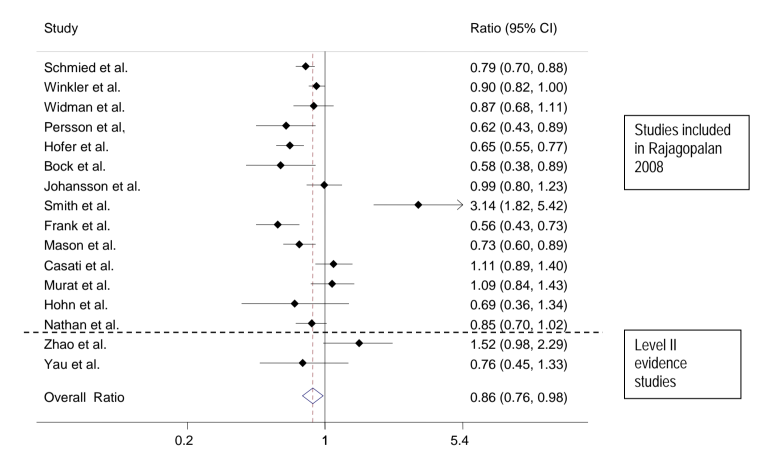
POQ3.l6.P3 Characteristics and results of studies examining the effect of the <u>prevention of hypothermia</u> during surgery on <u>blood loss</u>

Level of evidence		No. of trials /	Patient population /				Results			
Study	Quality	sample size	Surgical procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	Notes
Rajagopalan et al. (2008)	Level I Good	14 (N=1249)	Patients undergoing any surgical procedure	NR	Maintainence of normothermia, compared to patients with non-induced mild hypothermia	Ratio of blood loss (intervention vs control)	0.84 (95%C	10.74, 0.96)	0.009	Search date: 1996 to Oct 2006 Test of heterogeneity: P<0.001 Low publication observed
Zhao et al. (2005)	Level II Fair	N=40	ASA class I and II patients undergoing abdominal surgery lasting at least 2 hours.	Hospital in China	Warming from a forced-air blanket and warming of all intravenous fluids.	Blood loss (mL) Mean (SD)	639 (441)	421 (249)	NS	-
Yau et al. (1992)	Level II Fair	N=20	Patients undergoing isolated primary CABG.	Hospital in Canada	Warm systemic perfusion	Blood loss (mL) Mean (SD)	949 (427)	1253 (796)	NS	-

Abbreviations: ASA, American Society of Anaesthesiologists; CABG, coronary artery bypass graft; CI, confidence interval; NR, not reported; NS, not statistically significant; SD, standard deviation.

Note: To clarify the effect of hypothermia prevention on blood loss, a meta-analysis of the two Level II studies, together with the 14 studies included in the systematic review by Rajagopalan 2008 was conducted.

Meta-analysis of ratio of blood loss (Hypothermia prevention: No prevention)



Treatment effect is expressed as a ratio of the mean blood loss between treatment groups rather than a difference in mean as this allows a more intuitive comparison of studies examining different surgical procedures with varying volumes of blood loss.

The meta-analysis estimate (random effects model) indicates that the use of hypothermia prevention strategies resulted in a 14% lower average blood loss in patients as compared to patients where hypothermia was not prevented (Ratio: 0.86 (95%CI 0.75, 0.98), P=0.021).

Key question(s):		Evidence table ref*:				
In patients undergoing surgery, what is the effect of the prevention of hypothe	on <u>mortality</u> ?	POQ3.I6.P4				
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)					
Only one level I study, of poor quality, was identified (Mahoney et al. 1999).	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias			
	В	One or two Level II studies with a low risk of bias or SR/several Level	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	;			
2. Consistency (If only one study was available, rank this component as 'not applicable')						
No test of heterogeneity was reported in the systematic review.	Α	All studies consistent				
The systematic review identified two studies which reported on transfusion dose, one of which was a non-	В	Most studies consistent and inconsistency can be explained				
randomised controlled trial. The pooled effect estimates from the two studies showed a significant reduction in mortality rate when	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
hypothermia was prevented (2.7% vs 6.01%, P<0.05). However, data retrieved from the only the RCT did not	D	Evidence is inconsistent				
show a significant effect of hypothermia prevention on mortality (1.3% vs 1.4%, P=0.91).	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u>	nknowr	factor (not simply study quality or sample size) and thus the clinical impac	ct of the intervention could not be			
The systematic review found that the prevention of hypothermia significantly reduced mortality in patients undergoing surgery (2.7% vs 6.01%, P<0.05), however, the pooled effect was derived from two studies, of	Α	Very large				
which, only one was randomised.	В	Substantial				
Data from only the randomised study showed hypothermia prevention did not have an effect on mortality rate (1.3% vs 1.4%, P=0.91).	С	Moderate				
	D	Slight/Restricted				
4. Generalisability (How well does the body of evidence match the population and clinical se	ttings b	eing targeted by the Guideline?)				
The systematic review included two studies that reported mortality rates in patients undergoing abdominal, vascular and thoracic surgery. As such the evidence is likely generalisable to patients undergoing such surgical	Α	Evidence directly generalisable to target population				
procedures.	В	Evidence directly generalisable to target population with some cave	veats			
	С	Evidence not directly generalisable to the target population but could be sensibly applied				
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of healt					
The studies included in the systematic review were conducted in the US, as such the findings are probably applicable in the Australian context.	Α	Evidence directly applicable to Australian healthcare context				
аррисамо ит ите лазнанан сотнем.	В	Evidence applicable to Australian healthcare context with few cave				
	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
	D	Evidence not applicable to Australian healthcare context				

The systematic review by Mahoney et al. included one RCT and one non-randomised study to derive the mortality rate estimate.

Using only the data from the randomised study showed that prevention of hypothermia did not have an effect on mortality rate (1.3% vs 1.4%, P=0.91).

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	One level 1 study, of poor quality. Only two of the studies identified in the review reported mortality rates.
2. Consistency	NA	Only 1 systematic review was identified, no test of heterogeneity was reported.
3. Clinical impact	D	There was no evidence for an effect on mortality rate.
4. Generalisability	В	The studies identified included patients undergoing abdominal vascular and thoracic surgery. As such, the evidence is likely relevant to patients undergoing such procedures.
5. Applicability	С	Studies identified were conducted in the US, as such, the evidence is probably applicable in Australia.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of the prevention of hypothermia on mortality is uncertain.

POQ3.I6.P4 Characteristics and results of studies examining the effect of the prevention of hypothermia during surgery on mortality

	Level of evidence	No. of trials /	Patient population /					Notes		
Study	Quality Settir		Setting Intervention 0		Outcome	Intervention	Comparator	p-value		
					Maintananaa of		Pooled Incidence	e (RCT & non-RCT)	D .0 0E	Search Date:
Mahoney et	Level I	2 (2) 5 (2)	Patients undergoing any	ing any ND Maintenance of normothermia, compared to		Mortality rate	2.70%	6.01%	P<0.05	1989 to 1997
al. (1999) <i>Poor</i>	Poor	2 (N=562) ^a	surgical procedure	NR	patients with non-induced hypothermia	Pooled mean % (SD)	Incidence fro	m RCT (N=300)	D 0.04	No test of
							2/158 (1.3%)	2/142 (1.4%)	P=0.91	heterogeneity or bias conducted

Abbreviations: NR, not reported; SD, standard deviation.

^a Includes an RCT (N=300) and a non-randomised controlled trial (N=262).

Key question(s):	Evidence table ref*:				
In patients undergoing surgery, what is the effect of the prevention of hypothermia on morbidity? POQ3.I6.P5					
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)				
Two level I studies: 1 of fair quality (Scott et al.) and 1 of poor quality (Mahoney et al.) Two level II studies: 1 good quality RCT (Melling et al.), 1 fair quality RCT (Kim et al.).	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias			
	В	One or two Level II studies with a low risk of bias or SR/several Level	vel III studies with a low risk of bias		
Note: The systematic review by Mahoney et al. (1999) included two studies which reported on morbidity outcomes, one of the studies was also included in the more recent review by Scott et al. (2006).	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias		
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias			
2. Consistency (If only one study was available, rank this component as 'not applicable')					
Several different morbidity outcomes were examined. The prevention of hypothermia was found to significantly reduce the incidence of outcomes such as morbid cardiac events and wound infections.	Α	All studies consistent			
The studies by Scott et al. and Kim et al. failed to find a significant effect of the intervention on pain.	В	Most studies consistent and inconsistency can be explained			
	С	Some inconsistency, reflecting genuine uncertainty around questi	on		
	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>undetermined</u>)	<u>nknown</u>	factor (not simply study quality or sample size) and thus the clinical impact	ct of the intervention could not be		
Meta-analysis by Scott et al. showed that primary complication (morbid cardiac events, wound infection) were reduced by 63% in patients where hypothermia was prevented (P<0.00001). Mahoney et al. also reported	Α	Very large			
significantly lower rates of myocardial infarction when hypothermia was prevented in patients (2.3% vs 4.1%,	В	Substantial			
P<0.05). The study by Melling et al. also showed that wound infection was reduced (5% vs 14%, P=0.001).	С	Moderate			
	D	Slight/Restricted			
4. Generalisability (How well does the body of evidence match the population and clinical set	tings be	eing targeted by the Guideline?)			
The level I and II studies identified examined patients undergoing a variety of surgical procedures. As such, the evidence is likely generalisable to a broad range of surgical patient populations.	Α	Evidence directly generalisable to target population			
evidence is intoly generalisable to a block range of surgical patient populations.	В	Evidence directly generalisable to target population with some cave	reats		
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied		
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to		
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of healt				
The systematic reviews and the RCT by Melling et al. were conducted in countries in Europe, US and the UK. As such, the evidence is likely applicable in the Australian context.	Α	Evidence directly applicable to Australian healthcare context			
The study by Kim et al. was conducted in a military hospital in South Korea, as such, the findings of that study	В	Evidence applicable to Australian healthcare context with few cave			
are probably not applicable.	С	Evidence probably applicable to Australian healthcare context with	n some caveats		
	D	Evidence not applicable to Australian healthcare context			

The systematic review by Mahoney et al. included one non-randomised study (N=262), which may have affected the accuracy and reliability of the effect estimate.

The Clinical/Consumer Reference Group (CRG) suggested caution in the interpretation of the study by Melling et al. as it included operations with short durations and warming was done only preoperatively.

The CRG also noted that the morbid cardiac events, as reported by Scott et al., did not include the morbid outcomes of interest for this review, but rather haemodynamic changes (eg. tachycardia and hypotension)

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	В	2 systematic reviews comprising 8 studies in total. In addition to 1 good quality and 1 fair quality RCT.
2. Consistency	В	The systematic review found that hypothermia prevention significantly reduced the incidence of cardiac events and wound infection. However, this was derived from data of two studies only.
3. Clinical impact	С	Meta-analysis showed that primary complications (morbid cardiac events, wound infections) were reduced by 63% in patients where hypothermia was prevented
4. Generalisability	В	The studies identified included a patients undergoing any surgery, as such, the evidence is likely generalisable to patients undergoing a range of procedures.
5. Applicability	В	Except for the RCT by Kim et al., which was conducted in a military hospital in South Korea, the evidence from the other studies is probably applicable in Australia.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the prevention of hypothermia reduces the incidence of wound infection.

POQ3.I6.P5 Characteristics and results of studies examining the effect of the <u>prevention of hypothermia</u> during surgery on <u>morbidity</u>

	Level of evidence	No. of trials /	Patient population / Surgical					Results		Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	Notes
		7 (N=1061)				All primary complications	0.37 (0.2	27, 0.51)	<0.00001	
Scott et al. (2006)	Level I	2 (N=287)	Patients undergoing any surgical procedure (except cardiac procedures) under regional or	NR	Maintenance of normothermia, compared to patients with non-induced hypothermia	Morbid cardiac events ^a RR (95%CI)	0.34 (0.20, 0.57) NR		NR	Search date: 1948 to May 2003 No test of heterogeneity or bias conducted
(2000)	Fair	2 (N=284)	general anaesthesia.			Wound infection RR (95%CI)	0.26 (0.12, 0.58)		NR	
		3 (N=131)				Pain	No significant di between treat		NS	
Mahoney et al. (1999)	Level I Poor	2 (N=562) b	Patients undergoing any surgical procedure	NR	Maintenance of normothermia, compared to patients with non-induced hypothermia	Myocardial infarction Pooled mean % (SD)	2.3% (0.88)	4.1% (1.34)	P<0.05	Search Date: 1989 to 1997 No test of heterogeneity or bias conducted
Melling et al. (2001)	Level II Good	N=421	Patients having clean surgery (e.g. breast, varicose vein, or hernia), that would result in a scar longer than 3 cm.	Hospital in UK	Preoperative warming	Wound infection n/N (%)	13/277 (5)	19/139 (14)	P=0.001	-
Kim et al. (2009)	Level II Fair	N=50	ASA I or II patients undergoing arthroscopic shoulder surgery.	Military hospital in South Korea	Use of warm irrigation fluid	Pain measured using the VAS score	5.0 (1.7)	4.9 (1.6)	P=0.927	-

Abbreviations: ASA, American Society of Anaesthesiologists; CI, confidence interval; NR, not reported; NS, not statistically significant; RR, relative risk; SD, standard deviation; VAS, visual analogue score.

^a Includes hypertension, tachycardia, angina, cardiac arrest or myocardial infarction recoded on an electrocardiograph monitor.
^b Includes one non-randomised controlled trial.

In patients undergoing surgery, what is the effect of the prevention of hypothermia on quality of life? 1. Evidence base (number of studies, level of evidence and risk of bias in the included studies) No studies identified in literature search A One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or Level I or II studies with a low risk of bias or Level I or II studies with a low risk of bias or Level I or II studies with a moderate risk of bias or Level II or II studies with a high risk of bias or Level II or II studies with a high risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a high risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or III studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II or II studies with a low risk of bias or Level II studies with a low risk of bias or Level II studies with a low risk of bias or Level II studies with a low ri	Key question(s):			Evidence table ref*:
No studies identified in literature search A One or more level if studies with a low risk of bias or several level if studies with a low risk of bias or SR/several Level if studies with a low risk of bias or SR/several Level if studies with a low risk of bias or Consistency (if only one study was available, rank this component as not applicable?) NA A All studies consistent and inconsistency can be explained C Some inconsistency, reflecting genuine uncertainty around question D Evidence is inconsistent NA Not applicable (one study only) 3. Clinical impact (Indicate in the space below if the study results varied according to some unknown factor (not simply study quality or sample size) and thus the clinical impact of the intervention could not be debatrational. NA A Very targe B Substantial C Moderate D Sight/Restricted 4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?) NA A Evidence directly generalisable to target population with some caveats C Evidence and directly generalisable to target population but could be sensibly applied D Evidence and directly generalisable to target population and hard to judge whether it is sensible to Evidence and cultural factors?) NA A Evidence directly generalisable to target population and hard to judge whether it is sensible to Evidence and directly generalisable to Australian healthcare context B Evidence and directly generalisable to Australian healthcare context B Evidence and directly generalisable to Australian healthcare context B Evidence probably applicable to Australian healthcare context with two caveats C Evidence probably applicable to Australian healthcare context with two caveats C Evidence probably applicable to Australian healthcare context with two caveats	In patients undergoing surgery, what is the effect of the prevention of hypothe	<u>ermia</u>	on quality of life?	POQ3.I6.P6
B One or two Level II studies with a low risk of bias or SR/several Level III studies with a moderate risk of bias C One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias D Level IV studies or Level I to III studies/SRs with a high risk of bias 2. Consistency (if only one study was available, rank this component as not applicable) NA A All studies consistent B Most studies consistent C Some inconsistency, reflecting genuine uncertainty around question D Evidence is inconsistent NA Not applicable (one study only) 3. Clinical impact (Indicate in the space below if the study results varied according to some unknown factor (not simply study quality or sample size) and thus the clinical impact of the intervention could not be determined. NA A Very large B Substantial C Moderate D Slight/Restricted 4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?) NA A Evidence directly generalisable to target population with some caveats C Evidence of directly generalisable to target population with some caveats C Evidence of directly generalisable to target population and hard to judge whether it is sensible to D Evidence not directly generalisable to target population and hard to judge whether it is sensible to B Evidence of directly generalisable to Australian healthcare context with few caveats C Evidence directly applicable to Australian healthcare context with some caveats Evidence probably applicable to Australian healthcare context with some caveats C Evidence probably applicable to Australian healthcare context with some caveats	1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)		
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4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?) NA A Evidence directly generalisable to target population with some caveats C Evidence not directly generalisable to the target population but could be sensibly applied D Evidence not directly generalisable to target population and hard to judge whether it is sensible to 5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of health services/delivery of care and cultural factors?) NA A Evidence directly applicable to Australian healthcare context with few caveats C Evidence applicable to Australian healthcare context with some caveats		В	Substantial	
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C Evidence probably applicable to Australian healthcare context with some caveats	NA	Α	<u> </u>	
		В		
D Evidence not applicable to Australian healthcare context		С	Evidence probably applicable to Australian healthcare context with	n some caveats
		D	Evidence not applicable to Australian healthcare context	

Other factors (Ind	dicate here	any other factors that you took into account when assessing the evidence base (for example, issues that might cause the group to downgrade or upgrade the recommendation)
EVIDENCE STA	TEMEN	T MATRIX
Please summarise t	the develo	pment group's synthesis of the evidence relating to the key question, taking all the above factors into account.
Component	Rating	Description
1. Evidence base	NA	There were no studies that reported data on quality of life.
2. Consistency	NA	
3. Clinical impact	NA	
4. Generalisability	NA	
5. Applicability	NA	
DRAFT EVIDEN	ICE STA	TEMENT
Based on the body	of evidenc	e above.
In adult patie	nts unde	rgoing surgery in which substantial blood loss is anticipated, the effect of the prevention of hypothermia on quality of life is unknown.

Key question(s):			Evidence table ref*:		
In patients undergoing surgery, what is the effect of the prevention of hypothe	<u>ermia</u>	on <u>haemoglobin concentration</u> ?	POQ3.I6.S1		
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)				
Two level II studies: both of fair quality (Kim et al. and Yau et al.)	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias			
	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias		
	С	One or two Level III studies with a low risk of bias or Level I or II s	tudies with a moderate risk of bias		
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	S		
2. Consistency (If only one study was available, rank this component as 'not applicable')					
Both level II studies did not observe a significant effect of hypothermia prevention on haemoglobin concentration.	Α	All studies consistent			
concentiation.	В	Most studies consistent and inconsistency can be explained			
	С	Some inconsistency, reflecting genuine uncertainty around question			
	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u>	nknowr	n factor (not simply study quality or sample size) and thus the clinical impa	ct of the intervention could not be		
No significant effect was reported.	Α	Very large			
	В	Substantial			
	С	Moderate			
	D	No difference			
4. Generalisability (How well does the body of evidence match the population and clinical se	ttings b				
The RCT by Kim et al. was conducted in patients undergoing arthroscopic shoulder surgery, while Yau et al. examined patients undergoing CABG surgery. Generalisability of the findings is likely limited to patients	Α	Evidence directly generalisable to target population			
undergoing the aforementioned surgical procedures.	В	Evidence directly generalisable to target population with some car	veats		
	С	Evidence not directly generalisable to the target population but co	ould be sensibly applied		
	D	Evidence not directly generalisable to target population and hard	to judge whether it is sensible to		
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of healt	•			
The study by Kim et al. was conducted in a military hospital in South Korea; accordingly, the evidence may not be applicable in the Australian context.	Α	Evidence directly applicable to Australian healthcare context			
The study by Yau et al. was conducted in a hospital in Canada, as such the findings in this study are likely	В	Evidence applicable to Australian healthcare context with few cav			
applicable in Australia.	С	Evidence probably applicable to Australian healthcare context wit	h some caveats		
	D	Evidence not applicable to Australian healthcare context			

The Clinical/Consumer Reference Group suggested caution in the interpretation of the results from the study by Yau et al. due to the method of hypothermia prevention used (warming of systemic perfusion) and the small sample size (N=20).

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	2 Level II studies of fair quality.
2. Consistency	Α	Both studies did not observe a significant effect of hypothermia prevention on haemoglobin levels.
3. Clinical impact	D	No significant effect observed.
4. Generalisability	С	Generalisability of the findings is likely limited to the two surgical procedures examined by the two RCTs identified.
5. Applicability	D	One RCT was conducted in a military hospital in South Korea, as such the evidence may not be applicable in Australia. In contrast, the other RCT was conducted in Canada, as such, the findings are probably applicable in Australia.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In patients undergoing surgery in which substantial blood loss is anticipated, the effect of the prevention of hypothermia on haemoglobin concentration is uncertain.

Abbreviations: CABG, coronary artery bypass graft.

POQ3.l6.S1 Characteristics and results of studies examining the effect of the <u>prevention of hypothermia</u> during surgery on <u>haemoglobin concentration</u>

	Study Level of evidence Ouality No. of trials / sample size	Patient population / Surgical				Results				
Study		procedure Setting		Intervention	Outcome	Intervention	Comparator	p-value	Notes	
Kim et al. (2009)	Level II Fair	N=50	ASA I or II patients undergoing arthroscopic shoulder surgery.	Military hospital in South Korea	Use of warm irrigation fluid	Decrease in haemoglobin after surgery (g/dL) Mean (SD)	1.7 (0.7)	1.4 (0.6)	0.165	-
Yau et al. (1992)	Level II Fair	N=20	Patients undergoing isolated primary CABG.	Hospital in Canada	Warm systemic perfusion	Postoperative haemoglobin concentration	o o	No significant difference between treatment groups		-

Abbreviations: ASA, American Society of Anaesthesiologists; CABG, coronary artery bypass graft; CI, confidence interval; NR, not reported; standard deviation; VAS, visual analogue score.

Key question(s):			Evidence table ref*:		
In patients undergoing surgery, what is the effect of the prevention of hypothe	<u>ermia</u>	on <u>hospital length of stay</u> ?	POQ3.I6.S5		
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)				
One level I study: 1 systematic review of poor quality (Mahoney et al.) One level II study: 1 poor quality (Jeong et al.).	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias			
One lever it study: T poor quality (Jeong et al.).	В	One or two Level II studies with a low risk of bias or SR/several Lev	/el III studies with a low risk of bias		
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias		
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias			
2. Consistency (If only one study was available, rank this component as 'not applicable')					
The review by Mahoney et al. found that the prevention of hypothermia in patients was associated with a significantly shorter hospital stay.	Α	All studies consistent			
Similarly Jeong et al. observed a shorter hospital stay in patients in the intervention group, however, the	В	Most studies consistent and inconsistency can be explained			
difference was not statistically significant (10.6 days vs 11.6 days).	С	Some inconsistency, reflecting genuine uncertainty around question			
	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>ur</u>	nknown	factor (not simply study quality or sample size) and thus the clinical impac	t of the intervention could not be		
On average, the prevention of hypothermia was estimated to reduce hospital stay by 1 to 7.7 days.	Α	Very large			
	В	Substantial			
	С	Moderate			
	D	Slight/Restricted			
4. Generalisability (How well does the body of evidence match the population and clinical set	tings be	eing targeted by the Guideline?)			
The systematic review included patients undergoing any surgical procedure, as such the findings are likely relevant to the general surgical patient population.	Α	Evidence directly generalisable to target population			
The RCT by Jeong et al.was conducted in patients undergoing cardiac surgery, as such, findings from this study	В	Evidence directly generalisable to target population with some cave	eats		
is likely limited to patients in this surgical group.	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied		
	D	Evidence not directly generalisable to target population and hard tapply	o judge whether it is sensible to		
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of healt				
The systematic review by Mahoney et al included studies mostly conducted in Europe and the US. Accordingly, the findings are likely applicable in the Australian context.	Α	Evidence directly applicable to Australian healthcare context			
The RCT by Jeong et al. were conducted in South Korea. As such, the applicability of the findings may be more	В	Evidence applicable to Australian healthcare context with few cave			
limited.	С	Evidence probably applicable to Australian healthcare context with	ı some caveats		
	D	Evidence not applicable to Australian healthcare context			

The systematic review by Mahoney et al. included one non-randomised study (N=262), which may have affected the accuracy and reliability of the effect estimate.

The CRG noted that they did not consider length of hospital stay to be a clinically relevant outcome in assessing the effect of the prevention of hypothermia during surgery.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	1 poor quality systematic review. 1 good quality RCT.
2. Consistency	С	The studies showed that patients experienced shorter hospital stays when hypothermia was prevented. The extent of the effect may vary according to the surgical procedure examined.
3. Clinical impact	NA	This outcome was not considered to be a clinically relevant outcome for this intervention.
4. Generalisability	D	The studies identified in this review included patients undergoing a range of surgical procedure. As such, the evidence likely is generalisable to a broad range of surgical patients.
5. Applicability		The studies included in the systematic review were conducted in Europe and US, as such the evidence is likely applicable to Australia. However, the RCTs by Jeong et al. were conducted in South Kore, as such the applicability of those studies may be more limited.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In patients undergoing surgery in which substantial blood loss is anticipated, the effect of the prevention of hypothermia on length of hospital stay is uncertain.

POQ3.l6.S5 Characteristics and results of studies examining the effect of the <u>prevention of hypothermia</u> during surgery on <u>hospital length of stay</u>

	Level of evidence	No of trials / Patient population /								
Study	Quality	sample size	Surgical procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	Notes
Mahoney et al. (1999)	Level I Poor	3 (N=762) a	Patients undergoing any surgical procedure	NR	Maintenance of normothermia, compared to patients with non-induced hypothermia	No. of days in hospital Pooled mean (SD)	11.77 (0.10)	19.44 (0.16)	<0.05	Search Date: 1989 to 1997 No test of heterogeneity or bias
Jeong et al. (2008)	Level II Poor	N=40	Patients undergoing isolated off-pump coronary artery bypass surgery.	Hospital in South Korea	Warming of all intravenous fluids	No of days in hospital Mean (SD)	10.6 (2.2)	11.6 (2.7)	NS	-

Abbreviations: NS, not statistically significant; NR, not reported; SD, standard deviation.

^a Includes one non-randomised controlled trial.

Key question(s):		Evidence table ref*:			
In patients undergoing surgery, what is the effect of the prevention of hypothe	<u>rmia</u>	on <u>length of ICU stay</u> ?	POQ3.I6.S6		
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	.)				
One level I study: 1 systematic review of poor quality (Mahoney et al.)	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias		
One level II studies: 1 poor quality (Jeong et al.).	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias		
	С	One or two Level III studies with a low risk of bias or Level I or II st	rudies with a moderate risk of bias		
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	3		
2. Consistency (If only one study was available, rank this component as 'not applicable')					
All studies observed shorter stays in the ICU for patients when hypothermia was prevented. However, the difference was only significant in the review by Mahoney et al.	Α	All studies consistent			
	В	Most studies consistent and inconsistency can be explained			
The small sample size in the study by Jeong et al. may have reduced study power to detect a significant difference between the treatment groups.	С	Some inconsistency, reflecting genuine uncertainty around question			
amorono sourcon una usaunom groupor	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>ur</u>	nknown	factor (not simply study quality or sample size) and thus the clinical impact	ct of the intervention could not be		
The review by Mahoney et al. estimated that hypothermia prevention strategies during surgery reduced ICU stay by an average of 4.4 hours.	Α	Very large			
by an average or 4.4 nours.	В	Substantial			
	С	Moderate			
	D	Slight/Restricted			
4. Generalisability (How well does the body of evidence match the population and clinical set	tings be	eing targeted by the Guideline?)			
The systematic review by Mahoney et al. included patients undergoing any surgical procedure; as such the findings are likely relevant to the general surgical patient population.	Α	Evidence directly generalisable to target population			
The RCT by Jeong et al.was conducted in patients undergoing cardiac surgery. As such, findings from this study	В	Evidence directly generalisable to target population with some call	veats		
is likely limited to patients in this surgical group.	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied		
	D	Evidence not directly generalisable to target population and hard tapply	to judge whether it is sensible to		
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of healti	h services/delivery of care and cultural factors?)			
The systematic review by Mahoney et al. included studies mostly conducted in Europe and the US, as such, the findings are likely applicable in the Australian context.	Α	Evidence directly applicable to Australian healthcare context			
The RCT by Jeong et al. was conducted in South Korea. As such, the applicability of the findings may be more	В	Evidence applicable to Australian healthcare context with few cave	eats		
limited.	С	Evidence probably applicable to Australian healthcare context with	h some caveats		
	D	Evidence not applicable to Australian healthcare context			

The systematic review by Mahoney et al. included one non-randomised study (N=262), which may have affected the accuracy and reliability of the effect estimate.

The CRG noted that they did not consider ICU stay to be a clinically relevant outcome in assessing the effect of the prevention of hypothermia during surgery.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
Evidence base	С	1 systematic review of fair quality, 1 RCT of poor quality.
2. Consistency	С	Patients with hypothermia prevention experienced shorter ICU stays. However, this effect was not statistically significance in one of the RCTs.
3. Clinical impact	N/A	This outcome was not considered to be a clinically relevant outcome for this intervention.
4. Generalisability	D	The systematic review examined patients undergoing a range of surgical procedures, while the RCT examined cardiac surgery patients.
5. Applicability		Most of the data were from studies conducted in similarly developed countries like Australia. However, differences in the healthcare and hospital system may affect the applicability of the evidence.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of the prevention of hypothermia on length of ICU stay is uncertain.

POQ3.l6.S6 Characteristics and results of studies examining the effect of the <u>prevention of hypothermia</u> during surgery on <u>ICU stay</u>

	Level of evidence	No of trials / Patient population /			Results					
Study	Quality	sample size	Surgical procedure	Setting Intervention Outcome	Outcome	Intervention	Comparator	p-value	Notes	
Mahoney et al. (1999)	Level I Poor	2 (N=462) a	Patients undergoing any surgical procedure	NR	Maintenance of normothermia, compared to patients with non-induced hypothermia	ICU stay (hours) Pooled mean (SD)	5.51 (0.09)	9.70 (0.17)	P<0.05	Search Date: 1989 to 1997 No test of heterogeneity and bias conducted
Jeong et al. (2008)	Level II Poor	N=40	Patients undergoing isolated off-pump coronary artery bypass surgery.	Hospital in South Korea	Warming of all intravenous fluids	ICU stay (hours) Mean (SD)	59.6 (19.6)	70.5 (17.8)	NS	-

Abbreviations: ICU, intensive care unit; NS, not statistically significant; NR, not reported; SD, standard deviation.

^a Includes one non-randomised controlled trial.

Recommendation(s) for prevention of hypothermia

RECOMMENDATION	GRADE	RELE	VANT			
What recommendation(s) does the guideline development group draw from this evidence? Use action statements where possible.						
In patients undergoing surgery, measures to prevent hypothermia should be used.	A	PC	3.I6.P1,			
		PC	3.I6.P2,			
		PC	3.I6.P3,			
		PC	3.I6.P5			
IMPLEMENTATION OF RECOMMENDATION						
Please indicate yes or no to the following questions. Where the answer is yes please provide explanatory information about this. This information will be used to develop the implementation plan for the guidelines.						
Will this recommendation result in changes in usual care?		YES	NO			
Increased use of warming methods.						
Are there any resource implications associated with implementing this recommendation?		YES	NO			
Equipment costs.		·				
Will the implementation of this recommendation require changes in the way care is currently organised?		YES	NO			
Are the guideline development group aware of any barriers to the implementation of this recommendation		YES	NO			
Cost.						
What could help to facilitate implementation of the recommendation?		YES	NO			
Targeted funding for warming devices.						

Intervention 7 – Point-of-care testing using thromboelastography

Key question(s):			Evidence table ref*:		
In patients undergoing surgery, what is the effect of TEG-based point-of-care	ng on transfusion incidence?	POQ3.I7.P1			
1. Evidence base					
Level II evidence: Ak 2009 (fair quality; N=224); Avidan 2004 (fair quality; N=102); Shore-Lesserson 1999 (fair quality; N=105); Royston 2001 (poor quality; N=60)	А	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias			
Level III evidence: Avidan 2004 (fair quality; N=159)	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias		
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias		
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	i		
2. Consistency					
Meta-analyses of the Level II evidence were conducted herein using the results from Ak 2009, Avidan 2004, and	А	All studies consistent			
Shore-Lesserson 1999 ² . The degree of heterogeneity between the trials was not significant for the transfusion incidence of PRBCs (P=0.36; l ² =1%), FFP (P=0.11; l ² =55%), or platelets (P=0.51; l ² =0%). The results from	В	Most studies consistent and inconsistency can be explained			
Royston 2001 are not significant, but agree in direction with the results of the meta-analysis. The Level III	С	Some inconsistency, reflecting genuine uncertainty around questi	on		
evidence from Avidan 2004 agrees with the meta-analysis results; however the results are only significant for PRBC transfusion, not FFP or platelet transfusion.	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact					
Meta-analysed results	Α	Very large			
Incidence of PRBC transfusion – RR 0.84 (0.71, 1.00); P=0.05; N=431 Incidence of FFP transfusion – RR 0.52 (0.34, 0.81); P=0.003; N=431	В	Substantial (FFP; PLT)			
Incidence of platelet transfusion – RR 0.56 (0.36, 0.87); P=0.01; N=431	С	Moderate			
	D	Slight/Restricted (PRBC)			
4. Generalisability					
The studies were all conducted in adults undergoing cardiac surgery.	А	Evidence directly generalisable to target population			
	В	Evidence directly generalisable to target population with some caveats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied		
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to		
5. Applicability					
The studies were conducted in Turkey (Ak 2009), UK (Avidan 2004; Royston 2001), and USA (Shore-Lesserson 1999).		Evidence directly applicable to Australian healthcare context			
		Evidence applicable to Australian healthcare context with few cave	eats		
	С	Evidence probably applicable to Australian healthcare context with	n some caveats		
	D	Evidence not applicable to Australian healthcare context			

Other factors

Avidan 2004 provides Level II information for its RCT component and Level III if using the historical control arm.

Avidan 2004 included an algorithm with TEG and other tests.

For Royston 2001, the CRG assumed that red cells are inherent in blood component transfusion.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating			Description
	FFP	PRBC	PLT	
1. Evidence base	С	С	С	Several Level II and III studies with moderate risk of bias.
2. Consistency	С	С	С	Most studies consistent and inconsistency can be explained.
3. Clinical impact	В	D	В	Statistically significant and substantial clinical impact for FFP and PLT; not statistically significant for PRBC
4. Generalisability	В	В	В	All studies conducted in adults undergoing cardiac surgery
5. Applicability	В	В	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, the use of thromboelastography may reduce the incidence of FFP and platelet transfusion; the effect on the incidence of RBC transfusion is uncertain.

Abbreviations: FFP, fresh frozen plasma; het, heterogeneity; PRBC, packed red blood cells; RR, risk ratio.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission

^{* &}lt;u>Interventions</u>: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Royston 2001 was not included in the meta-analysis as the others reported the total units of blood components transfused but not the type of components transfused.

POQ3.I7.P1 Characteristics and results of studies examining the effect of point-of-care testing on transfusion incidence.

	Level of evidence	No. of trials /	Patient population / Surgical					Results	
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value
Ak (2009)	Level II Fair	N=224	Adults undergoing elective, first-time CABG with CPB.	Hospital in Turkey	TEG- based algorithm guided transfusion (comprising kaolinactivated (k) TEG and h-kTEG	Patients transfused with PRBC	52/114 (45.6%)	60/110 (54.5%)	P=0.181
					analyses) Comparator was clinician- directed transfusion ¹	Patients transfused with FFP	19/114 (16.6%)	31/110 (28.1%)	P=0.038
						Patients transfused with platelets	17/114 (14.9%)	29/110 (26.3%)	P=0.033
Avidan (2004)	Level II (POC vs laboratory test) or Level III (POC vs clinical discretion)	POC: N=51 Laboratory: N= 51 Clinical discretion: N=108	Adults undergoing elective, first-time CABG with CPB.	Hospital in UK	Algorithm based on near-patient haemostatic testing. POC devices used include ACT+/Junior, Hepcon HMS Hemostasis Management System, PFA-100 platelet function analyser; and two dual-channel TEG coagulation	Patients transfused with PRBCs	POC: 34/51 (67%)	(69%) Clinician discretion: 92/108 (85%)	Chi-square test: P=0.01 POC vs lab P=0.83 POC vs clinician P=0.02
	Fair				analysers used in parallel. Randomised comparator was algorithm using routine laboratory haemostatic tests and historical comparator was clinician discretion.	Patients transfused with FFP	POC: 2/51 (4%)	Laboratory: 0/51 (0%) Clinician discretion: 16/108 (15%)	Chi-square test: P=0.003 POC vs lab P=0.29 POC vs clinician P=0.07
						Patients transfused with platelets	POC: 2/51 (4%)	Laboratory: 1/51 (2%) Clinician discretion: 14/108 (13%)	Chi-square test: P=0.02 POC vs lab P=0.57 POC vs clinician P=0.10
Royston (2001)	Level II Poor	N=60	Adults undergoing cardiac surgery ²	Hospital in UK	Heparinase-modified TEG- guided intraoperative algorithm. Comparator was transfusion guided by clinical criteria and laboratory-based tests	Patients transfused with blood components	5/30 (17%)	10/30 (33%)	P=0.15

	Level of evidence	No. of trials /	Patient population / Surgical					Results	
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value
Shore-Lesserson (1999)	Level II Fair	N=105	Adults undergoing cardiac surgery with a moderate to high risk for requiring a transfusion	Hospital in USA	TEG-guided transfusion algorithm compared with standard laboratory-based transfusion therapy.	Patients transfused with allogeneic blood components (total)	22/53 (42%)	34/52 (65%)	P=0.01
					Patients transfused with packed RBCs (intraoperative)	17/53 (32%)	23/52 (44%)	P=0.2	
				Patients transfused with packed RBCs (postoperative)	10/53 (19%)	16/52 (31%)	P=0.16		
				Patients transfused with packed RBCs (total)	22/53 (42%)	31/52 (60%)	P=0.06		
						Patients transfused with FFP (intraoperative)	3/53 (6%)	8/52 (44%)	P=0.12
						Patients transfused with FFP (postoperative)	2/53 (4%)	11/52 (21%)	P<0.05
						Patients transfused with FFP (total)	4/53 (8%)	16/52 (31%)	P<0.05
					Patients transfused with platelet concentrates (intraoperative)	5/53 (9%)	8/52 (15%)	P=0.4	
				Patients transfused with platelet concentrates (postoperative)	3/53 (6%)	9/52 (17%)	P=0.06		

	Level of evidence	No. of trials /	Patient population / Surgical		Intervention		Results				
Study	Quality	sample size	procedure			Outcome	Intervention	Comparator	p-value		
						Patients transfused with platelet concentrates (total)	7/53 (13%)	15/52 (29%)	P<0.05		

Abbreviations: CABG, coronary artery bypass graft; CPB, cardiopulmonary bypass; FFP, fresh frozen plasma; POC, point-of-care; PRBC, packed red blood cells; TEG, thromboelastography.

¹ Using criteria obtained from abnormal laboratory tests (PT, APTT and platelet count), absence of visible clots and presence of generalised oozing-type bleeding in the surgical field to determine blood product administration.

² Ten per cent of patients in each series had a heart transplantation and were taking aspirin and/or warfarin immediately before surgery. About 50% of the patients in each group had revascularisation and were also taking aspirin, and required multiple grafts with a bypass time estimated to be greater than 100 minutes. The remaining 40% of the patients were having the Ross procedure, multiple valve or valve and revascularisation surgery. No patients were having repeat operations and none received prophylactic aprotinin, epsilon aminocaproic acid or tranexamic acid.

Key question(s): In patients undergoing surgery, what is the effect of point-of-care t	estina (on tr	ansfi	usion volume?	Evidence table ref*: POQ3.I7.P2
Evidence base	osung .	<u></u>	<u> </u>	Sign voidino.	
Level II evidence: Ak 2009 (fair quality; N=224); Avidan 2004 (fair quality; N=102); Royston 2001	PRRC	FFP	PI T		
(poor quality; N=60); Shore-Lesserson 1999 (fair quality; N=105); Westbrook 2009 (fair quality;	Α	Α	Α	One or more level I studies with a low risk of bias or several level II s	
N=69) Level III evidence: Avidan 2004 (fair quality; N=159)	В	В	В	One or two Level II studies with a low risk of bias or SR/several Level	III studies with a low risk of bias
200 M of Mondo. 7 Madin 2001 (kill quality, 11 107)	С	С	С	One or two Level III studies with a low risk of bias or Level I or II studi	es with a moderate risk of bias
	D	D	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	
2. Consistency			•		
The studies are consistent in finding no significant impact on volume of <u>PRBC</u> transfusion. Ak	PRBC	FFP	PLT		
2009 and Shore-Lesserson 1999 both found that the TEG-based clinical algorithm resulted in a significantly lower volume of <u>FFP</u> transfusion. Westbrook 2009 found no significant impact on	Α	Α	Α	All studies consistent	
volume of FFP transfusion. Ak 2009 found that the TEG algorithm resulted in a lower volume of	В	В	В	Most studies consistent and inconsistency can be explained	
<u>platelet</u> transfusion; whereas Shore-Lesserson 1999 and Westbrook 2009 found no significant difference between arms for this outcome.	С	С	С	Some inconsistency, reflecting genuine uncertainty around question	
Royston 2001 found that TEG-based transfusion significantly reduced volume of blood	D	D	D	Evidence is inconsistent	
components (FFP and platelets) transfused.	NA	NA	NA	Not applicable (one study only)	
3. Clinical impact					
See Summary Table POQ3.17.P2	PRBC	FFP	PLT		
	Α	Α	Α	Very large	
	В	В	В	Substantial	
	С	С	С	Moderate	
	D	D	D	No difference	
4. Generalisability					
The studies were all conducted in adults undergoing cardiac surgery.	PRBC	FFP	PLT		
	Α	Α	Α	Evidence directly generalisable to target population	
	В	В	В	Evidence directly generalisable to target population with some cavea	ts
	С	С	С	Evidence not directly generalisable to the target population but could	be sensibly applied
	D	D	D	Evidence not directly generalisable to target population and hard to ju	udge whether it is sensible to
5. Applicability			•		
The studies were conducted in Australia (Westbrook 2009), Turkey (Ak 2009), UK (Avidan 2004;	PRBC	FFP	PLT		
Royston 2001), and USA (Shore-Lesserson 1999).	Α	Α	Α	Evidence directly applicable to Australian healthcare context	
	В	В	В	Evidence applicable to Australian healthcare context with few caveats	3
	С	С	С	Evidence probably applicable to Australian healthcare context with so	ome caveats
	D	D	D	Evidence not applicable to Australian healthcare context	
		1	· ·		0.75

Other factors

Generalisability rated as 'B' due to inclusion of Avidan 2004, which included TEG and other tests in their transfusion algorithm.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component		Rating	l	Description
	PRBC	FFP	PLT	
1. Evidence base	С	С	С	Five Level II studies with moderate risk of bias.
2. Consistency	Α	В	С	Evidence is inconsistent.
3. Clinical impact	D	С	С	Moderate decrease in volume of FFPand PLT transfusion. No statistically significant impact on the volume of transfusion of PRBC.
4. Generalisability	В	В	В	Evidence not directly generalisable to the target population but could be sensibly applied.
5. Applicability	В	В	В	Evidence applicable to Australian healthcare context with few caveats.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, the use of thromboelastography may reduce the volume of FFP transfusion; the effect on volume of RBC and platelet transfusion is uncertain.

Abbreviations: FFP, fresh frozen plasma; PRBC, packed red blood cells; TEG, thromboelastography.* Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission

POQ3.I7.P2 Characteristics and results of studies examining the effect of point-of-care testing on transfusion volume.

	Level of evidence	No. of trials /	Patient population /					Results	
Study	Quality	sample size	Surgical procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value
Ak (2009)	Level II	N=224	Adults undergoing elective, first-time	Hospital in Turkey	TEG-based algorithm guided transfusion (comprising kaolin-	Median (IQR) units of PRBCs transfused intraoperatively	1 (0, 1)	0 (0, 1)	P=0.581
	Fall		CABG with CPB.		activated (k) TEG and h-kTEG analyses)	Median (IQR) units of PRBCs transfused postoperatively	1 (0, 1)	1 (0, 1)	P=0.741
					Comparator was clinician-directed transfusion ¹	Median (IQR) units of PRBCs transfused both intra-and postoperatively	1 (0, 1)	1 (1, 2)	P=0.599
						Median (IQR) units of FFP transfused intraoperatively	0 (0, 1)	1 (0, 1)	P=0.008
						Median (IQR) units of FFP transfused postoperatively	1 (0, 1)	1 (0, 1)	P=0.034
						Median (IQR) units FFP transfused both intra- and postoperatively	1 (0, 1)	1 (1, 2)	P=0.001
						Median (IQR) units of platelets transfused intraoperatively	0(0, 1)	1 (0, 1)	P=0.004
						Median (IQR) units of platelets transfused postoperatively	1 (0, 1)	1 (0, 1)	P=0.028
						Median (IQR) units of platelets transfused both intra- and postoperatively	1 (1, 1)	1 (1, 2)	P=0.001
Avidan (2004)	Level II (POC vs laboratory test) or Level	POC. N=51 Laboratory test. N=51	Adults undergoing elective, first-time CABG with CPB.	Hospital in UK	Algorithm based on near-patient haemostatic testing. POC devices used include ACT+/Junior, Hepcon HMS Hemostasts/Annagement System,	Mean (SD) units of PRBCs transfused for those transfused	POC: 2.9 (NR)	Laboratory: 2.7 (NR) Clinician discretion: 3.1 (NR)	NR
	III (POC vs clinical discretion) Fair	Clinical discretion. N=108			PFA-100 platelet function analyser; and two dual-channel TEG coagulation analysers used in parallel. Randomised comparator was algorithm using routine laboratory harmostatic.	Median (IQR) volume of PRBCs transfused, mL	POC: 500 (0, 678)	Laboratory: 495 (0, 612) Clinician discretion: 512 (286, 962)	Kruskal-Wallis ANOVA: P=0.03
		tests a	using routine laboratory haemostatic lests and historical comparator was clinician discretion.	Mean (SD) units of platelets transfused for those transfused	POC: 1.5 (NR)	Laboratory: 2.0 (NR) Clinician discretion: 1.0 (NR)	NR		

	Level of evidence	No. of trials /	Patient population /					Results	
Study	Quality	sample size	Surgical procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value
						Mean (SD) units of FFP transfused for those transfused	POC: 3.0 (NR)	Laboratory: 0 (0) Clinician discretion: 4.1 (NR)	NR
Royston (2001)	Level II Poor	N=60	Adults undergoing cardiac surgery ²	Hospital in UK	Heparinase-modified TEG-guided intraoperative algorithm.	Volume of blood components transfused	5 units of FFP and 1 pool of platelets	16 units of FFP and 9 platelet pools	P<0.05
	Pool				Comparator was transfusion guided by clinical criteria and laboratory-based tests				
Shore- Lesserson	Level II	N=105	Adults undergoing cardiac surgery with a	Hospital in USA	TEG-guided transfusion algorithm compared with standard laboratory-	Mean (SD) volume of PRBCs transfused (intraoperative), mL	267 (423)	346 (449)	P=0.4
(1999)	Fair		moderate to high risk for requiring a transfusion		based transfusion therapy.	Mean (SD) volume of PRBCs transfused (postoperative), mL	103 (252)	177 (318)	P=0.27
			II dii Siu Siu II			Mean (SD) volume of PRBCs transfused (total), mL	354 (487)	475 (593)	P=0.12
					-	Mean (SD) volume of FFP transfused (intraoperative), mL	22 (101)	113 (407)	P=0.4
						Mean (SD) volume of FFP transfused (postoperative), mL	33 (169)	146 (378)	P=0.13
						Mean (SD) volume of FFP transfused (total), mL	36 (142)	217 (463)	P<0.05
						Mean (SD) volume of platelet concentrates transfused (intraoperative), mL	22 (75)	41 (122)	P=0.6
						Mean (SD) volume of platelet concentrates transfused (postoperative), mL	11 (46)	42 (107)	P=0.3
						Mean (SD) volume of platelet concentrates transfused (total), mL	34 (94)	83 (160)	P=0.16
Westbrook (2009)	Level II	N=69	Adults undergoing cardiac surgery with	Hospital in Australia	TEG-guided transfusion algorithm vs. clinician directed administration with	Units of blood products transfused intraoperatively	19	44	ns (p-value not reported)
	(2009) Fair		the exception of one patient who underwent lung transplantation		reference to laboratory coagulation tests	Units of blood products transfused in ICU	18	46	ns (p-value not reported)
						Total units of blood products transfused	37	90	ns (p-value not reported)
						Units of PRBCs transfused intraoperatively	11	15	ns (p-value not reported)

	Level of evidence	No. of trials /	Patient population /					Results	
Study	Quality	sample size	Surgical procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value
						Units of PRBCs transfused in ICU	3	18	ns (p-value not reported)
						Total units of PRBCs transfused	14	33	ns (p-value not reported)
						Units of FFP transfused intraoperatively	8	14	ns (p-value not reported)
						Units of FFP transfused postoperatively	10	8	ns (p-value not reported)
						Total units of FFP transfused	18	22	ns (p-value not reported)
						Units of platelets transfused intraoperatively	0	10	ns (p-value not reported)
						Units of platelets transfused postoperatively	5	5	ns (p-value not reported)
						Total units of platelets transfused	5	15	ns (p-value not reported)
						Units of cryoprecipitate transfused intraoperatively	0	5	ns(p-value not reported)
						Units of cryoprecipitate transfused postoperatively	0	15	ns(p-value not reported)
						Total units of cryoprecipitate transfused	0	20	ns(p-value not reported)

Abbreviations: ANOVA, analysis of variance; CABG, coronary artery bypass graft; CPB, cardiopulmonary bypass; FFP, fresh frozen plasma; ICU, intensive care unit; IQR, interquartile range; POC, point-of-care; PRBC, packed red blood cells; SD, standard deviation; TEG, thromboelastography. Using criteria obtained from abnormal laboratory tests (PT, APTT and platelet count), absence of visible clots and presence of generalized oozing-type bleeding in the surgical field to determine blood product administration.

² Ten per cent of the patients in each series had a heart transplantation and were taking aspirin and/or warfarin immediately before surgery. About 50% of the patients in each group had revascularisation and were also taking aspirin, and required multiple grafts with a bypass time estimated to be greater than 100 minutes. The remaining 40% of the patients were having the Ross procedure, multiple valve or valve and revascularisation surgery. No patients were having repeat operations and none received prophylactic aprotinin, epsilon aminocaproic or tranexamic acid.

³ Before and after cohort design, single institution.

⁴ Bedside instrument measuring clot formation and dissolution indicating changes in coagulation, platelet function, platelet-fibrinogen interaction and fibrinolysis

Key question(s):			Evidence table ref*:			
In patients undergoing surgery, what is the effect of point-of-care testing on b	lood	loss?	POQ3.I7.P3			
1. Evidence base						
Level II evidence: Ak 2009 (fair quality; N=224); Avidan 2004 (fair quality; N=102); Royston 2001 (poor quality; N=60); Shore-Lesserson 1999 (fair quality; N=105); Westbrook 2009 (fair quality; N=69)	А	One or more level I studies with a low risk of bias or several level	I studies with a low risk of bias			
Level III evidence: Avidan 2004 (fair quality; N=159)	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency						
The studies are consistent in finding no significant difference between treatment arms.	Α	All studies consistent				
	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around question	on			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact						
All studies found no significant impact.	Α	Very large				
See Summary Table POQ3.17.P3.	В	Substantial				
	С	Moderate				
	D	Slight/Restricted				
4. Generalisability						
The studies were all conducted in adults undergoing cardiac surgery.	Α	Evidence directly generalisable to target population				
	В	Evidence directly generalisable to target population with some cav	reats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to			
5. Applicability						
The studies were conducted in Australia (Westbrook 2009), Turkey (Ak 2009), UK (Avidan 2004; Royston 2001), and USA (Chara Legerran 1000)	Α	Evidence directly applicable to Australian healthcare context				
and USA (Shore-Lesserson 1999).	В	Evidence applicable to Australian healthcare context with few cave	eats			
	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
	D	Evidence not applicable to Australian healthcare context				

Other factors

Avidan 2004 included TEG and other tests in their transfusion algorithm.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	Several Level II and III studies with moderate risk of bias.
2. Consistency	В	All studies consistent
3. Clinical impact	D	No statistically significant impact.
4. Generalisability	В	Evidence not directly generalisable to the target population but could be sensibly applied.
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, the use of thromboelastography does not appear to have an effect on blood loss.

Abbreviations: NA, not applicable.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission

^{* &}lt;u>Interventions</u>: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.17.P3 Characteristics and results of studies examining the effect of point-of-care testing on blood loss.

	Level of evidence	No. of trials /	Patient population /			_	Results			
Study	Quality	sample size	Surgical procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Ak (2009)	Level II Fair	N=224	Adults undergoing elective, first-time CABG with CPB.	Hospital in Turkey	TEG-based algorithm guided transfusion (comprising kaolin-activated (k) TEG and h-kTEG analyses) Comparator was clinician-directed transfusion ¹	Mean (SD) 12- hour mediastinal chest tube drainage, mL	480.5 (351)	591.4 (339.2)	P=0.087	
Avidan (2004)	Level II (POC vs laboratory test) or Level III (POC vs clinical discretion)	POC. N=51 Laboratory. N=51 Clinical discretion. N=108	Adults undergoing elective, first-time CABG with CPB.	Hospital in UK	Algorithm based on near-patient haemostatic testing. POC devices used include ACT+/Junior, Hepcon HMS Hemostasis Management System, PFA-100 platelet function analyser; and two dual-channel TEG coagulation analysers used in parallel. Randomised comparator was algorithm using routine laboratory haemostatic tests and historical comparator was clinician discretion.	Median (IQR) 24- hour postoperative blood loss, mL	POC: 755 (606, 975)	Laboratory: 850 (688, 1095) Clinician discretion: 810 (550, 1295)	NR	
Royston (2001)	Level II Poor	N=60	Adults undergoing cardiac surgery ¹	Hospital in UK	Heparinase-modified TEG-guided intraoperative algorithm. Comparator was transfusion guided by clinical criteria and laboratory-based tests	Median (IQR) 12-hour chest tube drainage, mL	470 (295, 820)	390 (240, 820)	NR	
Shore- Lesserson (1999)	Level II Fair	N=105	Adults undergoing cardiac surgery with a moderate to high risk	Hospital in USA	TEG-guided transfusion algorithm compared with standard laboratory-based transfusion therapy.	Mean (SD) six- hour mediastinal drainage, mL	362 (274)	469 (637)	P=0.63	
			for requiring a transfusion			Mean (SD) 24- hour mediastinal drainage, mL	702 (500)	901 (847)	P=0.27	
Westbrook (2009)	Level II Fair	N=69	Adults undergoing cardiac surgery with the exception of one patient who underwent lung transplantation	Hospital in Australia	TEG-guided transfusion algorithm vs. clinician directed administration with reference to laboratory coagulation tests	Median (IQR) blood loss, mL	875 (755-1130)	960 (820-1200)	P=0.437	

Abbreviations: CABG, coronary artery bypass graft; CPB, cardiopulmonary bypass; IQR, interquartile range; POC, point-of-care testing; NR, not reported; SD, standard deviation; TEG, thromboelastography.

¹ Using criteria obtained from abnormal laboratory tests (PT, APTT and platelet count), absence of visible clots and presence of generalized oozing-type bleeding in the surgical field to determine blood product administration.

Key question(s):	lik O	Evidence table ref*: POQ3.17.P4				
In patients undergoing surgery, what is the effect of point-of-care testing on nation 1. Evidence base	IIUI (a	<u>inty ?</u>	1 0 2 3 .17 .1 4			
Level II evidence: Ak 2009 (fair quality; N=224); Shore-Lesserson 1999 (fair quality; N=105)	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias			
Level III evidence: Spalding 2007 (fair quality; N=1422)	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of				
		One or two Level III studies with a low risk of bias or Level I or II st				
	С					
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency						
All the studies are consistent in finding no significant impact	Α	All studies consistent				
	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact						
Meta-analysis (conducted herein) of Level II evidence: RR 0.75 (0.19, 3.02); P=0.69; N=329	А	Very large				
Meta-analysis (conducted herein) of Level II and III evidence: RR 1.00 (0.67, 1.49); P=1.00; N=1751	В	Substantial				
	С	Moderate				
	D	No difference				
4. Generalisability						
The studies were all conducted in adults undergoing cardiac surgery.	Α	Evidence directly generalisable to target population				
	В	Evidence directly generalisable to target population with some cav	reats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to			
5. Applicability						
The studies were conducted in Turkey (Ak 2009), USA (Shore-Lesserson 1999), and Germany (Spalding 2007).	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few cave	eats			
	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
	D	Evidence not applicable to Australian healthcare context				

Other factors

Included studies were underpowered to detect a mortality difference.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description					
1. Evidence base	С	Several Level II and III studies with a moderate risk of bias.					
2. Consistency	Α	All studies consistent					
3. Clinical impact	D	No statistically significant impact					
4. Generalisability	В	All studies conducted in adults undergoing cardiac surgery					
5. Applicability	С	Evidence probably applicable to Australian healthcare context with some caveats					

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, the effect of the use of thromboelastography on mortality is uncertain.

Abbreviations: NA, not applicable; RR, relative risk.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{* &}lt;u>Interventions</u>: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I7.P4 Characteristics and results of studies examining the effect of point-of-care testing on mortality.

Level of evidence		N = - 6 + -! = I = /	Patient population / Surgical				Results		
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value
Ak (2009)	Level II Fair	N=224	Adults undergoing elective, first-time CABG with CPB.	Hospital in Turkey	TEG-based algorithm guided transfusion (comprising kaolinactivated (k) TEG and h-kTEG analyses)	Early mortality (defined as death within 30 days of operation)	3/114 (2.6%) (low cardiac output=2, multiple organ failure=1)	2/110 (1.8%)	P=0.68
					Comparator was clinician- directed transfusion ¹				
Shore-Lesserson (1999)	Level II Fair	N=105	Adults undergoing cardiac surgery with a moderate to high risk for requiring a transfusion	Hospital in USA	TEG-guided transfusion algorithm compared with standard laboratory-based transfusion therapy.	Mortality (ITT)	0/53 (0%)	2/52 (4%)	P=0.29
Spalding (2007)	Level III Fair	N=1422	Adults undergoing cardiac surgery	Hospital in Germany	TEG vs. no TEG	Early mortality (%)	41/693 (6%)	43/729 (6%)	P=0.99

Abbreviations: CABG, coronary artery bypass graft; CPB, cardiopulmonary bypass; ITT, intention-to-treat; TEG, thromboelastography.

Key question(s):	lli o	Evidence table ref*:		
In patients undergoing surgery, what is the effect of point-of-care testing on n	norbi	<u>dity?</u>	POQ3.I7.P5	
1. Evidence base				
Level II evidence: Shore-Lesserson 1999 (fair quality: N=105)		One or more level I studies with a low risk of bias or several level	I studies with a low risk of bias	
	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias	
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias	
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias		
2. Consistency				
	Α	All studies consistent		
	В	Most studies consistent and inconsistency can be explained		
	С	Some inconsistency, reflecting genuine uncertainty around question	on	
	D	Evidence is inconsistent		
	NA	Not applicable (one study only)		
3. Clinical impact				
Cerebrovascular ischemic event – 1 event in the TEG group (N=53); no events in the control group (N=52); RR	Α	Very large		
2.94 (0.12, 70.67); P=0.51; N=105	В	Substantial		
	С	Moderate		
	D	No difference		
4. Generalisability				
The study was conducted in adults undergoing cardiac surgery.	Α	Evidence directly generalisable to target population		
	В	Evidence directly generalisable to target population with some cave	eats	
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied	
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to	
5. Applicability				
The study was conducted in the USA.	Α	Evidence directly applicable to Australian healthcare context		
	В	Evidence applicable to Australian healthcare context with few cave	eats	
	С	Evidence probably applicable to Australian healthcare context with	n some caveats	
	D	Evidence not applicable to Australian healthcare context		

Other factors		
EVIDENCE STA	TEMENT	T MATRIX
Please summarise th	he develop	oment group's synthesis of the evidence relating to the key question, taking all the above factors into account.
Component	Rating	Description
Evidence base	С	One Level II study with moderate risk of bias.
2. Consistency	NA	Not applicable (one study only)
3. Clinical impact	D	No statistically significant impact
4. Generalisability	Α	Evidence not directly generalisable to the target population but could be sensibly applied
5. Applicability	В	Evidence probably applicable to Australian healthcare context with some caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, the effect of the use of thromboelastography on morbidity is uncertain.

Abbreviations: CI; confidence interval; RR, relative risk; TEG, thromboelastography.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I7.P5 Characteristics and results of studies examining the effect of point-of-care testing on morbidity.

0	Level of evidence Ouality No. of trials / sample size Patient population / Surgical procedure Setting Intervention Outcon	Intervention			Results				
Study				Setting	Intervention	Outcome	Intervention	Comparator	p-value
Shore-Lesserson (1999)	Level II Fair	N=105	Adults undergoing cardiac surgery with a moderate to high risk for requiring a transfusion	Hospital in USA	TEG-guided transfusion algorithm compared with standard laboratory-based transfusion therapy.	Cerebrovascular ischemic event (ITT)	1/53 (2%)	0/52 (0%)	P=0.51

Abbreviations: ITT, intension-to-treat; TEG, thromboelastography.

Key question(s):		Evidence table ref*:	
In patients undergoing surgery, what is the effect of point-of-care testing on g	<u>uality</u>	<u>y of life?</u>	POQ3.I7.P6
1. Evidence base			
No evidence found	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias
	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	
2. Consistency	1		
	Α	All studies consistent	
	В	Most studies consistent and inconsistency can be explained	
	С	Some inconsistency, reflecting genuine uncertainty around question	on
	D	Evidence is inconsistent	
	NA	Not applicable (one study only)	
3. Clinical impact			
	Α	Very large	
	В	Substantial	
	С	Moderate	
	D	Slight/Restricted	
4. Generalisability			
•	Α	Evidence directly generalisable to target population	
	В	Evidence directly generalisable to target population with some cav	reats
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to
5. Applicability			
	Α	Evidence directly applicable to Australian healthcare context	
	В	Evidence applicable to Australian healthcare context with few cave	
	С	Evidence probably applicable to Australian healthcare context with	n some caveats
	D	Evidence not applicable to Australian healthcare context	

Other factors	Other factors						
EVIDENCE STA	TEMENT	MATRIX					
Please summarise ti	he develop	oment group's synthesis of the evidence relating to the key question, taking all the above factors into account.					
Component	Rating	Description					
1. Evidence base	NA						
2. Consistency	NA						
3. Clinical impact	NA						
4. Generalisability	NA						
5. Applicability	NA						
DRAFT EVIDEN	CE STAT	ΓΕΜΕΝΤ					
Based on the body of	of evidence	above.					
In adult pa	itients un	dergoing surgery in which substantial blood loss is anticipated, the effect of the use of thromboelastography on quality of life is unknown.					
Abbreviations: NA, not app	licable.						

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Key question(s):	aglahin concentration?	Evidence table ref*: POQ3.I7.S1		
In patients undergoing surgery, what is the effect of <u>point-of-care testing</u> on <u>I</u> 1. Evidence base	iaeiii	oglobili concentration?	1 003.17.51	
Level II evidence: Avidan 2004 (fair quality; N=102); Westbrook 2009 (fair quality; N=69)	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias		
Level III evidence: Avidan 2004 (fair quality; N=159)				
	В	One or two Level II studies with a low risk of bias or SR/several Lev		
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias	
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias		
2. Consistency				
The studies are consistent in finding no significant difference.	Α	All studies consistent		
	В	Most studies consistent and inconsistency can be explained		
	С	Some inconsistency, reflecting genuine uncertainty around question	on	
	D	Evidence is inconsistent		
	NA	Not applicable (one study only)		
3. Clinical impact	1			
Median (IQR); TEG vs comparator	Α	Very large		
Level II evidence: Postoperative Hb concentration (Avidan 2004), g/dL – 9.3 (8.4, 10.3) vs. 9.3 (8.5, 9.7); P=NR; N=102	В	Substantial		
24 h postoperative Hb concentration (Avidan 2004), g/dL – 10.1 (9, 10.9) vs. 9.9 (9, 10.8); P=NR; N=102	С	Moderate		
Median (IQR) minimum Hb concentration (Westbrook 2009), g/L– 87 (83, 94) vs. 86 (82, 104); P=NR; N=69 Level III evidence (historical control)	D	Slight/Restricted		
24 h postoperative Hb concentration (Avidan 2004), g/dL – 10.1 (9, 10.9) vs. 10.1 (9.6, 10.8); P=NR; N=159		3		
4. Generalisability				
Both studies were conducted in adults undergoing cardiac surgery.	Α	Evidence directly generalisable to target population		
	В	Evidence directly generalisable to target population with some cav	eats	
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied	
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to	
5. Applicability	•			
The studies were conducted in Australia and the UK.	А	Evidence directly applicable to Australian healthcare context		
	В	Evidence applicable to Australian healthcare context with few cave	eats	
	С	Evidence probably applicable to Australian healthcare context with	n some caveats	
	D	Evidence not applicable to Australian healthcare context		

Other factors

Avidan 2004 included TEG and other tests in their transfusion algorithm.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	Level II and III studies with moderate risk of bias.
2. Consistency	Α	All studies consistent
3. Clinical impact	D	No statistically significant impact
4. Generalisability	В	Evidence not directly generalisable to the target population but could be sensibly applied
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above

In adult patients undergoing cardiac surgery, the effect of the use of thromboelastography on haemoglobin concentration is uncertain.

Abbreviations: NA, not applicable; NR not reported; TEG, thromboelastography.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{* &}lt;u>Interventions</u>: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I7.S1 Characteristics and results of studies examining the effect of point-of-care testing on haemoglobin concentration.

Study	Level of	No. of trials /	Patient population / Surgical procedure	Setting		Outcome	Results		
	evidence Quality	sample size			Intervention		Intervention	Comparator	p-value
Avidan (2004)	Level II (POC vs laboratory test) or Level III (POC vs	POC. N=51 Laboratory. N= 51 Clinical discretion.	Adults undergoing elective, first-time CABG with CPB.	Hospital in UK	Algorithm based on near-patient haemostatic testing. POC devices used include ACT+/Junior, Hepcon HMS Hemostasis Management System, PFA-100 platelet function analyser; and two dual-channel TEG coagulation analysers used in parallel.	Median (IQR) postoperative Hb concentration, g/dL	POC: 9.3 (8.4, 10.3)	Laboratory: 9.3 (8.5, 9.7) Clinician discretion: Not available	NR
	clinical discretion) Fair	N=108				Median (IQR) postoperative 24- hour Hb, g/dL	POC: 10.1 (9, 10.9)	Laboratory: 9.9 (9, 10.8) Clinician discretion: 10.1 (9.6, 10.8)	NR
					Randomised comparator was algorithm using routine laboratory haemostatic tests and historical comparator was clinician discretion.				
Westbrook (2009)	Level II Fair	N=69	Adults undergoing cardiac surgery with the exception of one patient who underwent lung transplantation	Hospital in Australia	TEG-guided transfusion algorithm vs. clinician directed administration with reference to laboratory coagulation tests	Median (IQR) minimum Hb concentration, g/I	87 (83-94)	86 (82-104)	NS (p value not reported)

Abbreviations: CABG, coronary artery bypass graft; CPB, cardiopulmonary bypass; Hb, haemoglobin; IQR, interquartile range; NR, not reported; NS, not significant; POC, point-of-care testing; TEG, thromboelastography.

Key question(s):		Evidence table ref*:				
In patients undergoing surgery, what is the effect of point-of-care testing on re	<u>eope</u>	ration for bleeding?	POQ3.I7.S2			
1. Evidence base						
Level II evidence: Ak 2009 (fair quality; N=224); Avidan 2004 (fair quality; N=102); Shore-Lesserson 1999 (fair quality; N=105)	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of b				
Level III evidence: Avidan 2004 (fair quality; N=159)	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency	Į.					
All the studies are consistent in finding no significant impact.	Α	All studies consistent				
	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around question	on			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact						
Meta-analysis (conducted herein) of Level II evidence: RR 0.86 (0.33, 2.25); P=0.76; N=431	Α	Very large				
	В	Substantial				
	С	Moderate				
	D	Slight/Restricted				
4. Generalisability						
The studies were conducted in adults undergoing cardiac surgery.	Α	Evidence directly generalisable to target population				
	В	Evidence directly generalisable to target population with some cav	reats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to			
5. Applicability						
The studies were conducted in Turkey (Ak 2009), the UK (Avidan 2004), and the USA (Shore-Lesserson 1999).	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few cave	eats			
	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
	D	Evidence not applicable to Australian healthcare context				

Other factors

Avidan 2004 included TEG and other tests in their transfusion algorithm.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
Evidence base		Several Level II and III studies with moderate risk of bias
2. Consistency	Α	All studies consistent
3. Clinical impact	D	No statistically significant impact
4. Generalisability	В	Evidence not directly generalisable to the target population but could be sensibly applied
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, the effect of the use of thromboelastography on the risk of reoperation for bleeding is uncertain.

Abbreviations: CI, confidence interval; NA, not applicable; RR, relative risk.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.17.S2 Characteristics and results of studies examining the effect of point-of-care testing on reoperation for bleeding.

	Level of	No. of trials /	Patient population / Surgical				Results		
Study	evidence Quality	sample size procedure Setting Intervention	Intervention	Outcome	Intervention	Comparator	p-value		
Ak (2009)	Level II Fair	N=224	Adults undergoing elective, first-time CABG with CPB.	Hospital in Turkey	TEG-based algorithm guided transfusion (comprising kaolinactivated (k) TEG and h-kTEG analyses) Comparator was cliniciandirected transfusion ¹	Re-exploration for bleeding	6/114 (5%) (causes all surgical)	5/110 (5%) (Causes, 2= surgical, 3 inappropriate surgical intervention for bleeding)	NR
Avidan (2004)	Level II (POC vs laboratory test) or Level III (POC vs clinical discretion)	POC. N=51 Laboratory. N=51 Clinical discretion. N=108	Adults undergoing elective, first-time CABG with CPB.	Hospital in UK	Algorithm based on near-patient haemostatic testing. POC devices used include ACT+/Junior, Hepcon HMS Hemostasis Management System, PFA-100 platelet function analyser; and two dual-channel TEG coagulation analysers used in parallel.	Reoperation for bleeding	POC: 1/51 (2%)	Laboratory: 1/51 (2%) Clinician discretion: 3/108 (3%)	POC vs. laboratory RR (95% CI): 1.00 (0.06, 15.56); P=1.00 POC vs. clinician discretion
					Randomised comparator was algorithm using routine laboratory haemostatic tests and historical comparator was clinician discretion.				RR (95% CI): 0.71 (0.08, 6.62); P=0.76
Shore-Lesserson (1999)	Level II Fair	N=105	Adults undergoing cardiac surgery with a moderate to high risk for requiring a transfusion	Hospital in USA	TEG-guided transfusion algorithm compared with standard laboratory-based transfusion therapy.	Reoperation for bleeding	0/53 (0%)	2/52 (4%)²	P=0.29

Abbreviations: CI, confidence interval; POC, point-of-care testing; RR, relative risk; TEG, thromboelastography.

¹ Using criteria obtained from abnormal laboratory tests (PT, APTT and platelet count), absence of visible clots and presence of generalised oozing-type bleeding in the surgical field to determine blood product administration.

² In one patient, a specific surgical source of bleeding was discovered.

Key question(s):			Evidence table ref*: POQ3.17.S3		
In patients undergoing surgery, what is the effect of <u>point-of-care testing</u> on	coagu	JIATION STATUS?	FUU3.11.33		
1. Evidence base Level II evidence: Shore-Lesserson 1999 (fair quality; N=105)	Ι,	One or more level I studies with a low risk of bias or several level	Il studios with a low risk of higs		
Level if evidence. Shore-Lesserson 1777 (Idii quality, N=103)	A				
	В	e or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bia			
	C One or two Level III studies with a low risk of bias or Level I or II studi				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	•		
2. Consistency					
Not applicable	Α	All studies consistent			
	В	Most studies consistent and inconsistency can be explained			
	С	Some inconsistency, reflecting genuine uncertainty around question	on		
	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact					
No significant impact found	А	Very large			
See Summary Table POQ3.17.S3	В	Substantial			
	С	Moderate			
	D	No difference			
4. Generalisability					
The study conducted in adults undergoing cardiac surgery.	Α	Evidence directly generalisable to target population			
	В	Evidence directly generalisable to target population with some cave	veats		
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied		
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to		
5. Applicability					
The study was conducted in the USA.	Α	Evidence directly applicable to Australian healthcare context			
	В	Evidence applicable to Australian healthcare context with few cave			
	С	Evidence probably applicable to Australian healthcare context with	n some caveats		
	D	Evidence not applicable to Australian healthcare context			

Other factors		
EVIDENCE STA	TEMENT	T MATRIX
Please summarise th	he develop	oment group's synthesis of the evidence relating to the key question, taking all the above factors into account.
Component	Rating	Description
Evidence base	С	One fair quality Level II study with moderate risk of bias
2. Consistency	NA	Not applicable (one study only)
3. Clinical impact	D	No statistically significant impact
4. Generalisability	Α	Evidence not directly generalisable to the target population but could be sensibly applied
5. Applicability	В	Evidence probably applicable to Australian healthcare context with some caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, the effect of the use of thromboelastography on coagulation status is uncertain.

Abbreviations: NA, not applicable.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I7.S3 Characteristics and results of studies examining the effect of point-of-care testing on coagulation status.

	Level of evidence	No. of trials /	Patient population /				Results			
Study	Quality	sample size	Surgical procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Shore-Lesserson (1999)	Level II	N=105	Adults undergoing cardiac surgery with a	Hospital in USA	TEG-guided transfusion algorithm compared with	Mean (SD) activated clotting time (baseline), seconds	165 (34)	170 (49)	P=0.55	
m fo	moderate to high risk for requiring a		standard laboratory-based transfusion therapy.	Mean (SD) activated clotting time (post-protamine), seconds	158 (93)	149 (20)	P=0.50			
			transfusion			Mean (SD) platelet count (baseline), X1000/µL	203 (66)	200 (78)	P=0.83	
					Mean (SD) platelet count (warming on CPB), X1000/μL	92 (79)	96 (79)	P=0.80		
						Mean (SD) platelet count (ICU), X1000/µL	111 (48)	120 (48)	P=0.34	
						Mean (SD) prothrombin time (baseline), seconds 13.0 (1.1)	13.0 (1.1)	12.9 (1.3)	P=0.67	
						Mean (SD) prothrombin time (post-protamine), seconds	18.1 (2.3)	21.3 (26)	P=0.38	
						Mean (SD) prothrombin time (ICU), seconds	16.1 (1.7)	15.7 (1.6)	P=0.22	
						Mean (SD) activated partial thromboplastin time (baseline), seconds	31.6 (6.9)	34.1 (13.1)	P=0.23	
						Mean (SD) activated partial thromboplastin time (post-protamine), seconds 52.2 (48.0)	52.2 (48.0)	43.0 (14)	P=0.19	
						Mean (SD) activated partial thromboplastin time (ICU), seconds	35.9 (6.1)	36.8 (10.2)	P=0.59	
						Mean (SD) fibrinogen concentration (baseline), mg/dL	409 (82)	416 (118)	P=0.73	
							Mean (SD) fibrinogen concentration (post-protamine), mg/dL	239 (86)	246 (86)	P=0.68
						Mean (SD) fibrinogen concentration (ICU), mg/dL	259 (95)	263 (118)	P=0.85	

Abbreviations: ICU, intensive care unit; SD, standard deviation; TEG, thromboelastography.

Key question(s): In patients undergoing surgery, what is the effect of point-of-care testing on le	nath	of hospital stay?	Evidence table ref*: POQ3.17.S5	
1. Evidence base	rigu	i di nospital stay:	1 0 20:17:00	
Level II evidence: Westbrook 2009 (fair quality; N=69)	А	One or more level I studies with a low risk of bias or several level II studies with a low risk of		
		One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias		
		One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias	
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias		
2. Consistency	1	<u> </u>		
	Α	All studies consistent		
	В	Most studies consistent and inconsistency can be explained		
	С	Some inconsistency, reflecting genuine uncertainty around question	on	
	D	Evidence is inconsistent		
	NA	Not applicable (one study only)		
3. Clinical impact				
Median (IQR) length of hospital stay (TEG vs clinician discretion), days: 9 (7, 3) vs. 8 (7, 12); P=NS (P-value not	Α	Very large		
reported)	В	Substantial		
	С	Moderate		
	D	Slight/Restricted		
4. Generalisability				
The study was conducted in patients undergoing cardiac surgery with the exception of one patient who underwent lung transplantation.	Α	Evidence directly generalisable to target population		
ung transplantation.	В	Evidence directly generalisable to target population with some cav	eats	
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied	
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to	
5. Applicability				
The study was conducted in Australia.	Α	Evidence directly applicable to Australian healthcare context		
	В	Evidence applicable to Australian healthcare context with few caveats		
	С	Evidence probably applicable to Australian healthcare context with	some caveats	
	D	Evidence not applicable to Australian healthcare context		

Other	factors
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EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	One Level II study with moderate risk of bias.
2. Consistency	NA	Not applicable (one study only)
3. Clinical impact	D	No statistically significant impact.
4. Generalisability	Α	Evidence not directly generalisable to the target population but could be sensibly applied
5. Applicability	Α	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, the effect of the use of thromboelastography on length of hospital stay is uncertain.

Abbreviations: IQR, interquartile range; NA, not applicable; NS, not significant; TEG, thromboelastography.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I7.S5 Characteristics and results of studies examining the effect of point-of-care testing on hospital length of stay.

0	Level of evidence	No. of trials /	Patient population / Surgical	0 111				Results	
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value
Westbrook (2009)	Level II Fair	N=69	Adults undergoing cardiac surgery with the exception of one patient who underwent lung transplantation	Hospital in Australia	TEG-guided transfusion algorithm vs. clinician directed administration with reference to laboratory coagulation tests	Median (IQR) length of hospital stay, days	9 (7, 3) *Extra day not due to bleeding	8 (7, 12)	NS (P-value NR)

Abbreviations: IQR, interquartile range; NR, not reported; NS, not significant; TEG, thromboelastography.

Key question(s):			Evidence table ref*:		
In patients undergoing surgery, what is the effect of point-of-care testing on IC	CU a	dmission and length of stay?	POQ3.I7.S6		
1. Evidence base					
Level II evidence: Westbrook 2009 (fair quality; N=69)	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias			
		One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	;		
2. Consistency	1				
	Α	All studies consistent			
	В	Most studies consistent and inconsistency can be explained			
	С	Some inconsistency, reflecting genuine uncertainty around question	on		
	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact					
Median (IQR) length of ICU stay (TEG vs clinician discretion), hours: 29.4 (14.3, 56.4) vs. 32.5 (22, 74.5); P=NS (P-value not reported)	Α	Very large			
(i -value not reported)	В	Substantial			
	С	Moderate			
	D	No difference			
4. Generalisability					
The study was conducted in patients undergoing cardiac surgery with the exception of one patient who underwent lung transplantation.	Α	Evidence directly generalisable to target population			
tung tanspantation.	В	Evidence directly generalisable to target population with some cav	veats		
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied		
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to		
5. Applicability					
The study was conducted in Australia.	Α	Evidence directly applicable to Australian healthcare context			
	В	Evidence applicable to Australian healthcare context with few caveats			
	С	Evidence probably applicable to Australian healthcare context with	n some caveats		
	D	Evidence not applicable to Australian healthcare context			

Other factors	
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EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	One Level II study with moderate risk of bias.
2. Consistency	NA	Not applicable (one study only)
3. Clinical impact	D	No statistically significant impact.
4. Generalisability	Α	Evidence not directly generalisable to the target population but could be sensibly applied
5. Applicability	Α	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, the effect of the use of thromboelastography on length of ICU stay is uncertain.

Abbreviations: ICU, intensive care unit; IQR, interquartile range; NA, not applicable; NS, not significant; TEG, thromboelastography.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.17.S6 Characteristics and results of studies examining the effect of point-of-care testing on length of ICU stay.

	Level of evidence	No. of trials /	Patient population / Surgical				Results		
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value
Westbrook (2009)	Level II Fair	N=69	Adults undergoing cardiac surgery with the exception of one patient who underwent lung transplantation	Hospital in Australia	TEG-guided transfusion algorithm vs. clinician directed administration with reference to laboratory coagulation tests	Median (IQR) length of ICU stay, hours	29.4 (14.3, 56.4)	32.5 (22, 74.5)	NS (P-value NR)

Abbreviations: ICU, intensive care unit; IQR, interquartile range; NR, not reported; NS, not significant; TEG, thromboelastography.

Recommendation(s) for point-of-care testing using thromboelastography

RECOMMENDATION	GRADE	RELE	VANT	
What recommendation(s) does the guideline development group draw from this evidence? Use action statements where possible.		EVIDENCE TABLE		
In adult patients undergoing cardiac surgery, the use of thromboelastography should be considered.	С	PO3.I7.P1, PO3.I7.P2		
		PC)3.17.P2	
IMPLEMENTATION OF RECOMMENDATION		•		
Please indicate yes or no to the following questions. Where the answer is yes please provide explanatory information about this. This information will be used to develop the implementation plan for the guidelines.				
Will this recommendation result in changes in usual care?		YES	NO	
Increased use of TEG and related devices.				
Are there any resource implications associated with implementing this recommendation?		YES	NO	
Capital investment, training and staffing (technicians).				
Will the implementation of this recommendation require changes in the way care is currently organised?		YES	NO	
Are the guideline development group aware of any barriers to the implementation of this recommendation		YES	NO	
Significant cost.				
What could help to facilitate implementation of the recommendation?		YES	NO	
Targeted funding for equipment.				

Intervention 8 – Administration of antifibrinolytics & DDAVP: Aprotinin

Key question(s):			Evidence table refa:			
In patients undergoing surgery, what is the effect of administration of aprotining	<u>n</u> on	transfusion incidence?	POQ3.I8a.P1			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)					
There is one pivotal Level I study (Henry 2007/good quality), four supportive level I studies (Henry 2009/good quality; Brown 2007; fair quality; Kagoma 2009/good quality; McIlroy 2009 /good quality) and one supportive	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias			
Level I/II studyc (Gurusamy 2009/good-fair quality). In addition, three additional RCTs (Later 2009/good quality;	В	One or two Level II studies with a low risk of bias or SR/several Lev	k of bias or SR/several Level III studies with a low risk of bias			
Nurözler 2008/fair quality, Colwell 2007/good quality) were identified that were published following the Henry 2007 literature search that were not included in any of the supportive Level I studies.	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Results consistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta- analyses described below. Additional RCT results consistent.	Α	All studies consistent				
Pivotal evidence – Henry 2007	В	Most studies consistent and inconsistency can be explained				
Moderate to substantial heterogeneity between studies. Appears to be due to differences in magnitude of effect rather than lack of effect in some studies. May also be due to different surgery types assessed.	С	Some inconsistency, reflecting genuine uncertainty around question	n			
Tathor than tack of shock in some statics. Thay they be due to direction sargery types assessed.		dence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>undetermined</u>)	nknow	<u>n</u> factor (not simply study quality or sample size) and thus the clinical impa	ct of the intervention could not be			
Pivotal evidence – Henry 2007	Α	Very large				
Pivotal evidence – Henry 2007 Any surgery – RR 0.66 (0.62, 0.71); 43.8% vs 64.3%; 96 RCTs (N=9949) Cardiac surgery – RR 0.66 (0.61, 0.72); 45.6% vs 66.1%; 76 RCTs (N=8793)	В	Substantial				
Orthopaedic surgery – RR 0.69 (0.56, 0.85); 23.1% vs 43.9%; 13 RCTs (N=771) Liver surgery – RR 0.58 (0.37, 0.90); 24.1% vs 43.3%; 2 RCTs (N=177)	С	Moderate				
Supportive evidence –see Summary Table POQ3.18a.P1	D	Slight/Restricted				
4. Generalisability (How well does the body of evidence match the population and clinical set	tings b	peing targeted by the Guideline?)				
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were included in the overall analysis; 76/96 studies in cardiac surgery.		Evidence directly generalisable to target population				
included in the overall analysis; 70/96 studies in cardiac surgery.	В	Evidence directly generalisable to target population with some cav	eats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to	judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of heal	Ith services/delivery of care and cultural factors?)				
Hospital setting. The pivotal review states that studies were conducted in a wide range of countries.	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few cave	ats			
	С	Evidence probably applicable to Australian healthcare context with	some caveats			
	D	Evidence not applicable to Australian healthcare context				
	_					

Other factors (Indicate here any other factors that you took into account when assessing the evidence base (for example, issues that might cause the group to downgrade or upgrade the recommendation)

In 2007, Bayer announced a worldwide suspension of aprotinin supply due to the results of the BART trial which suggested increased risk of mortality for aprotinin compared with the lysine analogues tranexamic acid and ε-aminocaproic acid.

The Henry (pivotal) review assessed quality and performed a subgroup analysis of transfusion incidence for all surgery types based on the rating (A,B or C) of treatment allocation. The analysis showed no substantial difference in the results between studies rated A. B or C.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I (good quality), four supportive level I studies, one supportive Level I/II study ^c and three additional RCTs.
2. Consistency	В	Significant heterogeneity in the pivotal level I study but mostly due to difference in magnitude of effect and not direction of effect. Differences may be due to different surgery types. Results of supportive level I studies and additional RCTs consistent with pivotal evidence.
3. Clinical impact	В	There were significant differences between intravenous aprotinin therapy and no therapy overall and for surgery subgroups. Substantial clinical impact.
4. Generalisability	В	The results are generalisable to an adult surgical population; the majority of studies were conducted in patients undergoing cardiac surgery.
5. Applicability	В	Overall there were a large number of studies conducted in a wide range of countries. Likely to be applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, intravenous aprotinin therapy reduces the incidence of allogeneic blood transfusion compared with no therapy.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI. confidence interval. RBC, red blood cell: SR, systematic review:

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission b Cochrane ratings defined as follows: Grade A, adequate allocation concealment; Grade B, uncertain allocation concealment; Grade C, inadequate allocation concealment.

^a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

^c A supportive level I/II study represents a systematic review which identified only one relevant RCT.

POQ3.l8a.P1 Characteristics and results of studies examining the effect of <u>aprotinin</u> on <u>transfusion incidence</u>.

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or AP (%) vs control (%)	Significance P-value	
ADULT POPULATION	/IV APROTININ								
Any surgery	_	1							_
Henry (2007)	Level I Good	96 RCTs N=9949	Adult patients undergoing any surgery	Hospital – planned surgery Various countries ^c	Aprotinin (IV) vs no aprotinin	Transfusion incidence (allogeneic blood)	RR 0.66 (0.62, 0.71)	Favours aprotinin <0.001	Substantial Phet<0.001 (l ² =68%)
Henry (2007)	Level I Good	16 RCTs N=1251	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Prime dose aprotinin (IV) vs no aprotinin d	Transfusion incidence (allogeneic blood)	RR 0.83 (0.71, 0.96)	Favours aprotinin 0.014	Substantial Phet<0.001 (I ² =75%)
		43 RCTs N=3073	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Low dose aprotinin (IV) vs no aprotinin e	Transfusion incidence (allogeneic blood)	RR 0.66 (0.59, 0.74)	Favours aprotinin <0.001	Substantial Phet<0.001 (I ² =53%)
		56 RCTs N=6569	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	High dose aprotinin (IV) vs no aprotinin f	Transfusion incidence (allogeneic blood)	RR 0.65 (0.60, 0.71)	Favours aprotinin <0.001	Substantial Phet<0.001 (I ² =66%)
Henry (2007)	Level I Good	76 RCTs N=8768	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin with transfusion protocol	Transfusion incidence (allogeneic blood)	RR 0.65 (0.60, 0.70)	Favours aprotinin <0.001	Substantial Phet<0.001 (I ² =70%)
		20 RCTs N=1182	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin without transfusion protocol	Transfusion incidence (allogeneic blood)	RR 0.73 (0.62, 0.86)	Favours aprotinin <0.001	Substantial Phet<0.001 (I ² =59%)
Henry (2007)	Level I Good Rating Ag	27 RCTs N=2113	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Transfusion incidence (allogeneic blood)	OR 0.65 (0.54, 0.78)	Favours aprotinin <0.001	Substantial Phet<0.001 (I ² =84%)
	Level I Good Rating Bg	57 RCTs N=6993	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Transfusion incidence (allogeneic blood)	OR 0.68 (0.63 0.73)	Favours aprotinin <0.001	Moderate Phet=0.75 (I ² =48%)
	Level I Good Rating C9	12 RCTs N=799	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Transfusion incidence (allogeneic blood)	OR 0.60 (0.49, 0.73)	Favours aprotinin <0.001	Moderate Phet=0.13 (I²=32%)

Study	Level of	No. of trials /	Patient population / Surgical procedure	Setting Location	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size included in analysis					Risk estimate (95% CI) or AP (%) vs control (%)	Significance P-value	
Cardiac surgery									
Henry (2007)	Level I Good	76 RCTs N=8793	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Transfusion incidence (allogeneic blood)	RR 0.66 (0.61, 0.72)	Favours aprotinin <0.001	Substantial Phet<0.001 (I ² =71%)
Henry (2007) Level I Good		15 RCTs N=1191	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Prime dose aprotinin (IV) vs no aprotinin d	Transfusion incidence (allogeneic blood)	RR 0.81 (0.69, 0.96)	Favours aprotinin 0.012	Substantial Phet<0.001 (l ² =78%)
	24 RCTs N=1995	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Low dose aprotinin (IV) vs no aprotinin e	Transfusion incidence (allogeneic blood)	RR 0.67 (0.58, 0.77)	Favours aprotinin <0.001	Substantial Phet<0.001 (I ² =62%)	
		55 RCTs N=6533	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	High dose aprotinin (IV) vs no aprotinin	Transfusion incidence (allogeneic blood)	RR 0.66 (0.60, 0.72)	Favours aprotinin <0.001	Substantial Phet<0.001 (I ² =67%)
Henry (2009)	Level I Good	81 RCTs N=9139 ^h	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries ^b	Aprotinin (IV) vs placebo	Transfusion incidence (allogeneic blood)	RR 0.66 (0.61, 0.72)	Favours aprotinin <0.05	NR
Brown (2007)	Level I Fair	49 RCTs N=4379	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	High dose aprotinin (IV) vs placeboi	Transfusion incidence (pRBCs)	RR 0.60 (0.53, 0.67)	Favours aprotinin <0.001	NR
		20 RCTs N=1645	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	<u>Low dose</u> aprotinin (IV) vs placeboi	Transfusion incidence (pRBCs)	RR 0.76 (0.66, 0.86)	Favours aprotinin <0.001	NR
McIlroy (2009)	Level I Good	10 RCTs N=856	Adult patients undergoing cardiac surgery receiving ASA	Hospital – planned surgery Various countries	Aprotinin (IV) vs placebo	Transfusion incidence (allogeneic blood)	OR 0.34 (0.25, 0.46)	Favours aprotinin <0.001	None Phet=0.75 (l ² =0%)
Later (2009)	Level II Good	1 RCT N=199	Adult patients undergoing low- and intermediate-risk cardiac surgery	Hospital – planned surgery The Netherlands	High-dose aprotinin (IV) vs placeboi	Transfusion incidence (pRBCs)	50% vs 70.9%	Favours aprotinin 0.004	NA
			THE INCHIGHANA		Transfusion incidence (blood products)	61.5% vs 78.6%	Favours aprotinin 0.009	NA	
Nurözler (2008) Level II Fair		N=51 pump coronary bypass who	Hospital – planned surgery	Low-dose aprotinin (IV) vs placebo ^k	Transfusion incidence (RBC)	68% vs 88%	Favours aprotinin 0.014	NA	
	have received clopidogrel within 5 days of surgery	Turkey		Transfusion incidence (blood products)	28% vs 53%	Favours aprotinin 0.002	NA		

Study	Level of	No. of trials /	Patient population / Surgical procedure	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size included in analysis		Location			Risk estimate (95% CI) or AP (%) vs control (%)	Significance P-value	
Orthopaedic surge	ery								
Henry (2007)	Level I Good	13 RCTs N=771	Adult patients undergoing orthopaedic surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Transfusion incidence (allogeneic blood)	RR 0.69 (0.56, 0.85)	Favours aprotinin <0.001	None Phet=0.23 (l ² =21%)
Kagoma (2009)	Level I Good	3 RCTs N=347	Adult patients undergoing orthopaedic surgery	Hospital – planned surgery Unknown	Aprotinin (IV) vs placebo	Transfusion incidence	RR 0.63 (0.50, 0.80)	Favours aprotinin <0.05	NR
Colwell (2007)	Level II Good	1 RCT N=352	Adults undergoing <u>unilateral</u> total hip arthroplasty	Hospital – planned surgery US/Canada	Aprotinin (IV) vs placebo	Transfusion incidence (whole blood or RBCs)	17% vs 32%	Favours aprotinin 0.0009	NA
		1 RCT N=352	Adults undergoing <u>unilateral</u> total hip arthroplasty	Hospital – planned surgery US/Canada	Aprotinin (IV) vs placebo	Transfusion incidence (allogeneic blood)	11% vs 22%	Favours aprotinin 0.006	NA
		1 RCT N=278	Adults undergoing <u>unilateral</u> total hip arthroplasty	Hospital – planned surgery US/Canada	Aprotinin (IV) vs placebo	Transfusion incidence (whole blood or RBCs without donation)	13% vs 24%	Favours aprotinin 0.02	NA
		1 RCT N=74	Adults undergoing <u>unilateral</u> total hip arthroplasty	Hospital – planned surgery US/Canada	Aprotinin (IV) vs placebo	Transfusion incidence (whole blood or RBCs with donation)	32% vs 62%	Favours aprotinin ND	NA
Liver surgery	II.		-1	•	1	1	1	1	•
Henry (2007)	Level I Good	2 RCTs N=177	Adult patients undergoing <u>liver</u> <u>surgery</u>	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Transfusion incidence (allogeneic blood)	RR 0.58 (0.37, 0.90)	Favours aprotinin 0.015	None Phet=0.31 (I ² =3%)
Gurusamy (2009)	Level I/II Good/Fair	1 RCT N=37	Adult patients undergoing <u>liver</u> resection	Hospital – planned surgery Unknown	Aprotinin (IV) vs placebo	Transfusion incidence (allogeneic blood)	RR 0.43 (0.21, 0.89)	Favours aprotinin 0.02	NA Phet=NA (I²=NA)
Other surgery								•	
Henry (2007)	Level I Good	2 RCTs N=62	Adult patients undergoing thoracic surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Transfusion incidence (allogeneic blood)	RR 0.28 (0.11, 0.74)	Favours aprotinin 0.011	None Phet=0.54 (I ² =0%)
Henry (2007)	Level I/II Good/Good	1 RCT N=60	Adult patients undergoing vascular surgery	Hospital – planned surgery Unknown	Aprotinin (IV) vs no aprotinin	Transfusion incidence (allogeneic blood)	RR 1.01 (0.72, 1.40)	No difference 0.98	None Phet=NA (I²=NA)

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or AP (%) vs control (%)	Significance P-value	
Henry (2007)	Level I/II Good/Poor	1 RCT N=56	Adult patients undergoing neurosurgery	Hospital – planned surgery Unknown	Aprotinin (IV) vs no aprotinin	Transfusion incidence (allogeneic blood)	RR 0.73 (0.40, 1.35)	No difference 0.32	NA Phet=NA (I ² =NA)
Henry (2007)	Level I/II Good/Poor	1 RCT N=30	Adult patients undergoing orthognathic surgery	Hospital – planned surgery Unknown	Aprotinin (IV) vs no aprotinin	Transfusion incidence (allogeneic blood)	RR 0.11 (0.02, 0.77)	Favours aprotinin 0.026	NA Phet=NA (I ² =NA)
PAEDIATRIC POPUL	ATION/IV APROTINI	N							
Orthopaedic surg	gery								
Tzortzopoulou (2008)	Level II Good	1 RCT N=43	Paediatric patients undergoing scoliosis surgery	Hospital – planned surgery Unknown	Aprotinin (IV) vs placebo	Transfusion incidence	RR 0.75 (0.44, 1.27)	No difference 0.28	NA
ADULT POPULATION	/TOPICAL APROTIN	IIN							
Cardiac surgery									
Abrishami (2009)	Level I Good	3 RCTs N=341	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Aprotinin (topical) vs placebo	Transfusion incidence (allogeneic RBC)	RR 0.72 (0.47, 1.08)	No difference 0.11	Substantial Phet=0.008 (I ² =60%)

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: ASA, acetylsalicylic acid; CI, confidence interval; het, heterogeneity; IV, intravenous; NA, not determined; NR, not reported; OR, odds ratio; RBC, red blood cell; RCT, randomised controlled trial; RR, risk ratio.

a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between aprotinin and no aprotinin.

d Prime' dose included regimens that added aprotinin to the pump prime solution of the cardiopulmonary bypass exclusively. 12/16 trials studied a 'prime' dose of 2 million KIU, 2/16 trials studied a 'prime' dose of 1 million KIU, 1/16 trials studied a 'prime' dose of 500,000 KIU and 1/16 trials studies a 'prime' dose of 25,000 KIU/kg.

^e Low-dose aprotinin was defined as any regimen that did not follow the 'full Hammersmith' regimen, including those studies that described their regimen as 'half Hammersmith'. For non-cardiac surgery trials, regimens were classified as low dose if the total dose was < 5 million KIU or 700 mg aprotinin.

f High-dose aprotinin was defined as any regimen that was described as the 'full Hammersmith' regimen. For non-cardiac surgery trials, regimens were classified as high-dose if the total dose was ≥ 5 million KIU or 700 mg aprotinin.

⁹ Cochrane ratings defined as follows: Grade A, adequate allocation concealment; Grade B, uncertain allocation concealment; Grade C, inadequate allocation concealment.

^hTotal number of trials available for analysis. The actual number of trials included in the analysis is not reported.

High dose (full-dose) aprotinin defined as a 2 million kallikrein-inhibiting units (KIU) IV loading dose, 2 million KIU pump-priming dose, and 0.5 million KIU IV/h maintenance dose.

Low dose (half-dose) aprotinin consisted of a 1 million KIU IV loading dose, 1 million KIU pump-priming dose, and 0.25 million KIU IV/h maintenance dose.

^{*}Low dose aprotinin consisted of 1 million KIU infused over 30 min followed by a continuous infusion of 0/5 million KIU/h until the end of surgery.

Key question(s):			Evidence table refa:			
In patients undergoing surgery, what is the effect of administration of aprotini	<u>n</u> on :	transfusion volume?	POQ3.I8a.P2			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	5)					
There is one pivotal Level I study (Henry 2007/good quality) study which included data from 35 RCTs which provides data on the transfusion volume in patients who received transfusion. There was one pivotal Level I study	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias			
(Henry 2007/good quality/61 RCTs) and five additional RCTs published following the Henry review (Later 2009/good quality; Nurözler 2008/fair quality; Colwell 2007/good quality; Apostolakis 2008/fair quality; Leijdekkers 2006/fair	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias			
quality) which provide data on transfusion volume in all patients (transfused or not transfused).	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Consistency of individual studies within meta-analyses described below. Additional RCTs consistent. Pivotal evidence – Henry 2007	Α	All studies consistent				
Moderate to substantial heterogeneity between studies. Heterogeneity could be due to differences in surgery types,	В	Most studies consistent and inconsistency can be explained				
degree of bleeding expected with different surgery types and transfusion triggers used in each study. Supportive evidence –see Summary Table POQ3.l8a.P1		Some inconsistency, reflecting genuine uncertainty around question	on			
		Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>undetermined</u>)	nknowi	n factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be			
Pivotal evidence – Henry 2007	Α	Very large				
Pivotal evidence – Henry 2007 Any surgery (transfused patients only) – WMD -0.96 units (-1.24, -0.68); 35 RCTs (N=3363) Supportive evidence –see Summary Table POQ3.l8a.P1	В	Substantial				
	С	Moderate				
	D	Slight/Restricted				
4. Generalisability (How well does the body of evidence match the population and clinical set	tings b	neing targeted by the Guideline?)				
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were included in the overall analysis.	Α	Evidence directly generalisable to target population				
included in the overall analysis.	В	Evidence directly generalisable to target population with some cav	eats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of heal					
Hospital setting. The pivotal review states that studies were conducted in a wide range of countries.	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few cave				
	С	Evidence probably applicable to Australian healthcare context with some caveats				
	D	Evidence not applicable to Australian healthcare context				

Other factors (Indicate here any other factors that you took into account when assessing the evidence base (for example, issues that might cause the group to downgrade or upgrade the recommendation)

In 2007, Bayer announced a worldwide suspension of aprotinin supply due to the results of the BART trial which suggested increased risk of mortality for aprotinin compared with the lysine analogues tranexamic acid and ε-aminocaproic acid.

The most relevant data comes from the Henry 2007 pivotal review which assessed transfusion volume in transfused patients only (ie, takes out effect of patients who received no transfusion). Heterogeneity was discussed by the CRG and it was concluded that it may be related to degree of bleeding, surgery type and transfusion triggers.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I (good quality) study which provides data on the transfusion volume in patients who received transfusion (includes 35 RCTs). There was one pivotal Level I (good quality) study and five additional RCTs which provide data on transfusion volume in all patients (transfused or not transfused).
2. Consistency	В	Most studies were reasonably consistent. Some differences which may be related to differences in surgery type, degree of bleeding and transfusion triggers.
3. Clinical impact	В	There was a significant difference in transfusion volume between intravenous aprotinin therapy and no therapy when only transfused patients were considered. There were significant differences between intravenous aprotinin therapy and no therapy overall and for surgery subgroups in transfused + non-transfused patients.
4. Generalisability	В	The results are generalisable to an adult surgical population.
5. Applicability	В	Overall there were a large number of studies conducted in a wide range of countries. Likely to be applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, intravenous aprotinin therapy reduces the volume of allogeneic blood transfusion compared with no therapy.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI, confidence interval, SR, systematic review;

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.l8a.P2 Characteristics and results of studies examining the effect of <u>aprotinin</u> on <u>transfusion volume</u>

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or Aprotinin (mean ± SD) vs control (mean ± SD)	Significance P-value	
ADULT POPULATION/	IV APROTININ								
Any surgery									
. , , , ,	Level I Good	61 RCTs N=6780	Adult patients undergoing any surgery (all patients)	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Transfusion volume (units; allogeneic blood)	WMD -1.07 (-1.31, -0.83)	Favours aprotinin <0.001	Substantial Phet<0.001 (I ² =90%)
		35 RCTs N=3363	Adult patients undergoing any surgery (transfused patients only)	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Transfusion volume (units; allogeneic blood)	WMD -0.96 (-1.24, -0.68)	Favours aprotinin <0.001	Substantial Phet<0.001 (I ² =60%)
Cardiac surgery	•					•		•	•
Later (2009)	Level II Good	1 RCT N=199	Adult patients undergoing low- or intermediate-risk cardiac surgery (all patients)	Hospital – planned surgery The Netherlands	High-dose aprotinin (IV) vs placebod	Transfusion volume (units; pRBCs)	MD -1.0 (-1.0, 0)	Favours aprotinin <0.001	NA
Nurözler (2008)	Level II Fair	1 RCT N=51	Adult patients undergoing off- pump coronary bypass who have received clopidogrel	Hospital – planned surgery Turkey	<u>Low-dose</u> aprotinin (IV) vs placebo ^e	Transfusion volume (units; pRBC)	1.7 ± 1.4 vs 2.9 ± 1.8	Favours aprotinin 0.014	NA
	within 5 days of surgery (all patients)	Tunoj		Transfusion incidence (units; platelets)	0.4 ± 0.6 vs 2.3 ± 1.2	Favours aprotinin 0.002	NA		
					Transfusion incidence (units; FFP)	0.6 ± 0.3 vs 1.4 ± 0.6	Favours aprotinin 0.008	NA	

Study	Level of	No. of trials /	Patient population / Surgical procedure	Setting Location	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size included in analysis					Risk estimate (95% CI) or Aprotinin (mean ± SD) vs control (mean ± SD)	Significance P-value	
Orthopaedic surge	ery	<u> </u>	<u>.</u>			•			
Colwell (2007)	Level II Good	1 RCT N=352	Adults undergoing <u>unilateral</u> total hip arthroplasty (all patients)	Hospital – planned surgery US/Canada	Aprotinin (IV) vs placebo	Transfusion volume (whole blood or RBCs)	0.27 vs 0.63	Favours aprotinin 0.0003	NA
		1 RCT N=352	Adults undergoing <u>unilateral</u> total hip arthroplasty (all patients)	Hospital – planned surgery US/Canada	Aprotinin (IV) vs placebo	Transfusion volume (allogeneic blood)	0.17 vs 0.42	Favours aprotinin 0.004	NA
		1 RCT N=278	Adults undergoing <u>unilateral</u> total hip arthroplasty (all patients)	Hospital – planned surgery US/Canada	Aprotinin (IV) vs placebo	Transfusion volume (whole blood or RBCs without donation)	0.21 vs 0.46	Favours aprotinin 0.0153	NA
			Adults undergoing <u>unilateral</u> total hip <u>arthroplasty (all patients)</u>	Hospital – planned surgery US/Canada	Aprotinin (IV) vs placebo	Transfusion volume (whole blood or RBCs with donation)	0.52 vs 1.21	Favours aprotinin ND	NA
Other surgery	1							I	
Apostolakis (2008)	Level II Fair			Hospital – planned surgery Greece	<u>Ultra-low-dose</u> aprotinin (IV) vs placebo ^f	Intraoperative transfusion volume (units; pRBCs)	0.17 ± 0.54 vs 0.17 ± 0.53	No difference 0.967	NA
					Postoperative transfusion volume (units; pRBCs)	0.00 ± 0.00 vs 0.03 ± 0.18	No difference 0.970	NA	
					Intraoperative transfusion volume (units; FFP)	0.21 ± 0.62 vs 0.20 ± 0.76	No difference 0.330	NA	
						Postoperative transfusion volume (units; FFP)	0.21 ± 0.62 vs 0.87 ± 1.53	Favours aprotinin 0.035	NA
Leijdekkers (2006)	Level II Fair	1 RCT N=35	Adult patients undergoing surgery for infra-renal abdominal aneurysm (all	Hospital – planned surgery The Netherlands	Aprotinin (IV) vs placebo	Mean total infusion (mL)	7845 ± 4888 vs 7835 ± 4776	No difference 0.99	NA
		patients)	THE NEUTERIANUS		Mean packed cells (units)	4.1 ± 3.1 vs 4.1 ± 2.9	No difference 0.95	NA	
						Mean FFP (units)	0.5 ±0.9 vs 0.3 ± 0.8	No difference 0.35	NA

Study	Level of evidence ^a Quality	No. of trials / sample size included in analysis	Patient population / Surgical procedure	Setting Location	Intervention	Outcome	Results		Heterogeneity ^b
							Risk estimate (95% CI) or Aprotinin (mean ± SD) vs control (mean ± SD)	Significance P-value	
PAEDIATRIC POPULA	TION/IV APROTINI	N							
Schouten (2009)	Level I Fair	3 RCTs N=250	Paediatric patients undergoing cardiac surgery	Hospital – planned surgery Unknown	Aprotinin (IV) vs no aprotinin	Transfusion volume (mL/kg; <u>pRBCs</u>)	WMD -4 (-7, -2)	Favours aprotinin NR	None Phet=NR (I²=0%)
	Level I Fair	2 RCTs N=228	Paediatric patients undergoing cardiac surgery	Hospital – planned surgery Unknown	Aprotinin (IV) vs no aprotinin	Transfusion volume (mL/kg; plasma)	WMD -5 (-8, -2)	Favours aprotinin NR	None Phet=NR (I²=0%)
Tzortzopoulou (2008)	Level I Good	2 RCTs N=87	Paediatric patients undergoing scoliosis surgery	Hospital – planned surgery Unknown	Aprotinin (IV) vs placebo	Transfusion volume (mL)	WMD -361 (-584, -139)	Favours aprotinin 0.0015	None Phet=0.80 (I ² =0%)
ADULT POPULATION/	TOPICAL APROTIN	VIN							
Cardiac surgery									
Abrishami (2009)	Level I Good	4 RCTs N=229	Adult patients undergoing primary on-pump cardiac surgery	Hospital – planned surgery Unknown	Aprotinin (topical) vs placebo	Transfusion volume (allogeneic RBC)	WMD -0.83 (-1.21, -0.44)	Favours aprotinin <0.001	None Phet=0.34 (I ² =11%)
Mehraien (2009)	Level II Good	1 RCT N=128	Adult patients undergoing first-time CABG (all patients)	Hospital – planned surgery Iran	Aprotinin (topical) vs placebo	Mean packed cells (units)	0.5 ± 0.7 vs 1.7 ± 1.0	Favours aprotinin 0.002	

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: CI, confidence interval; FFP, fresh frozen plasma; het, heterogeneity; IV, intravenous; kg, kilogram; mL, millilitre; ND, not determined (small sample size); NR, not reported; RBC, red blood cell; RCT, randomised controlled trial; SMD, standardised mean difference; WMD, weighted mean difference.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between aprotinin and no aprotinin.

d High dose aprotinin defined as a 2 million kallikrein-inhibiting units (KIU) IV loading dose, 2 million KIU pump-priming dose, and 0.5 million KIU IV/h maintenance dose.

^e Low dose aprotinin consisted of 1 million KIU infused over 30 min followed by a continuous infusion of 0/5 million KIU/h until the end of surgery.

f Ultra-low dose aprotinin defined as a test dose of 1mL at intubation, followed by 0.5 million IU over 15 min following intubation, and again following closure.

Key question(s):			Evidence table refa:			
In patients undergoing surgery, what is the effect of administration of aprotining	POQ3.I8a.P3					
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	5)					
There is one pivotal Level I study (Henry 2007/good quality) which included up to 79 RCTs depending on the specific blood loss outcome assessed, three supportive Level I studies (Brown 2007/fair quality; Kagoma 2009/good	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias				
quality; McIlroy 2009/good quality), one supportive Level I/II study (Gurusamy 2009/good-fair quality) and six additional RCTs (Grant 2008/fair quality; Later 2009/good quality; Nurözler 2008/fair quality; Colwell 2007/good	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias				
uality; Apostolakis 2008/fair quality; Leijdekkers 2006/fair quality).		One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Results consistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta- analyses described below. Additional RCT results consistent.	Α	All studies consistent				
Pivotal evidence – Henry 2007	В	Most studies consistent and inconsistency can be explained				
Moderate to substantial heterogeneity between studies. May be due to different surgery types assessed. Supportive evidence –see Summary Table POQ3.I8a.P3	С	Some inconsistency, reflecting genuine uncertainty around question				
Supportive evidence - see Summary Table 1 Octs. Ioa. 1 3	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>ur</u>	nknow	<u>n</u> factor (not simply study quality or sample size) and thus the clinical impa	ct of the intervention could not be			
Pivotal evidence – Henry 2007	Α	Very large				
Any surgery (total blood loss) – WMD -414 mL (-520, -309); 15 RCTs (N=1577) Cardiac surgery (total blood loss) – WMD -489 mL (-571, -407); 5 RCTs (N=1147)	В	Substantial				
Orthopaedic surgery (total blood loss) – WMD -399 mL (-563, -235); 10 RCTs (N=430) Liver surgery (total blood loss) – WMD -1200 mL (-2943, -543); 2 RCTs (N=137)	С	Moderate				
Other surgery/outcomes and supportive evidence – see Summary Table POQ3.l8a.P3	D	Slight/Restricted				
4. Generalisability (How well does the body of evidence match the population and clinical set	tings b	peing targeted by the Guideline?)				
e evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were luded in the overall analysis.	Α	Evidence directly generalisable to target population				
included in the overall analysis.	В	Evidence directly generalisable to target population with some cav	eats			
	С	Evidence not directly generalisable to the target population but could be sensibly applied				
	D	Evidence not directly generalisable to target population and hard to judge whether it is sensible to				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of heal	th services/delivery of care and cultural factors?)				
Hospital setting. The pivotal review states that overall, studies were conducted in a wide range of countries. It was not possible to determine the location of each of the individual studies from the review.	Α	Evidence directly applicable to Australian healthcare context				
not possible to determine the location of each of the maintaids statics from the review.	В	Evidence applicable to Australian healthcare context with few caveats				
		Evidence probably applicable to Australian healthcare context with some caveats				
	D	Evidence not applicable to Australian healthcare context				

In 2007, Bayer announced a worldwide suspension of aprotinin supply due to the results of the BART trial which suggested increased risk of mortality for aprotinin compared with the lysine analogues tranexamic acid and ε-aminocaproic acid.

A 400 mL difference in blood loss was considered to represent a moderate clinical impact by the CRG.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

	•	
Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (good quality), three supportive Level I studies, one supportive Level I/II study and six additional RCTs.
2. Consistency	В	Significant heterogeneity in the pivotal level I study but mostly due to difference in magnitude of effect and not direction of effect. Differences may be due to different surgery types. Results of supportive level I studies and additional RCTs consistent with pivotal evidence.
3. Clinical impact	С	There were significant (or near significant) differences in blood loss for all surgery types. The clinical impact was considered to be moderate.
4. Generalisability	В	The results are generalisable to an adult surgical population.
5. Applicability	В	Overall there were a large number of studies conducted in a wide range of countries. Likely to be applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, intravenous aprotinin therapy reduces blood loss compared with no therapy.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI, confidence interval, RBC, red blood cell: SR, systematic review;

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission b A Level I/II study is a systematic review which included only one RCT.

a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.l8a.P3 Characteristics and results of studies examining the effect of <u>aprotinin</u> on <u>blood loss</u>

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b								
	evidence ^a Quality	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or Aprotinin (mean ± SD) vs control (mean ± SD)	Significance P-value									
ADULT POPULATION	/IV APROTININ																
Any surgery					1												
Henry (2007)	Level I Good	15 RCTs N=1577	Adult patients undergoing any surgery	Hospital – planned surgery Various countries ^c	Aprotinin (IV) vs no aprotinin	Total blood loss (mL)	WMD -414 (-520, -309)	Favours aprotinin <0.001	Substantial Phet=0.003 (l ² =57%)								
		13 RCTs N=722	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Intraoperative blood loss (mL)	WMD -185 (-280, -90)	Favours aprotinin <0.001	Substantial Phet<0.001 (I ² =67%)								
		79 RCTs N=7414	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Postoperative blood loss (mL)	WMD -358 (-404, -313)	Favours aprotinin <0.001	Substantial Phet<0.001 (I ² =86%)								
Cardiac surgery																	
Henry (2007)	D7) Level I Good	5 RCTs N=1147	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Total blood loss (mL)	WMD -489 (-571, -407)	Favours aprotinin <0.001	None Phet=0.62 (I ² =0%)								
		5 RCTs N=360	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Intraoperative blood loss (mL)	WMD -140 (-244, -36)	Favours aprotinin 0.0086	Substantial Phet=0.01 (l ² =68%)								
											68 RCTs N=6948	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Postoperative blood loss (mL)	WMD -385 (-432, -339)	Favours aprotinin <0.001
Henry (2007)	Level I Good	15 RCTs N=1158	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Prime dose aprotinin (IV) vs no aprotinin d	Postoperative blood loss (mL)	WMD -343 (-458, -228)	Favours aprotinin <0.001	Substantial Phet<0.001 (I ² =88%)								
		21 RCTs N=1781	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Low dose aprotinin (IV) vs no aprotinin e	Postoperative blood loss (mL)	WMD -293 (-349, -238)	Favours aprotinin <0.001	Substantial Phet<0.001 (l ² =61%)								
		48 RCTs N=4819	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	High dose aprotinin (IV) vs no aprotinin	Postoperative blood loss (mL)	WMD -428 (-485, -371)	Favours aprotinin <0.001	Substantial Phet<0.001 (l ² =85%)								

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or Aprotinin (mean ± SD) vs control (mean ± SD)	Significance P-value	
Brown (2007)	Level I Fair	22 RCTs N=1760	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	High dose aprotinin (IV) vs placebog	Total blood loss (mL)	WMD -348 (-416, -281)	Favours aprotinin <0.001	NR
		6 RCTs N=515	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	Low dose aprotinin (IV) vs placeboh	Total blood loss (mL)	WMD -226 (-277, -175)	Favours aprotinin <0.001	NR
McIlroy(2009)	Level I Good	12 RCTs N=992	Adult patients <u>receiving ASA</u> undergoing cardiac surgery	Hospital – planned surgery Unknown	Aprotinin (IV) vs placebo	Postoperative chest tube blood loss (mL)	WMD -433 (-544, -321)	Favours aprotinin <0.001	Substantial Phet<0.001 (I ² =74%)
Grant (2008) Level II Fair			N=120 <u>pump coronary artery bypass</u> s	Hospital – planned surgery US	Aprotinin (IV) vs placebo	Intraoperative blood loss (mL)	867 ± 413^{j} and $870 \pm 383j^{j}$ vs 1252 ± 380	Favours aprotinin <0.02	NA
				03		Postoperative blood loss (mL/24 hrs)	415 ± 330 ^j and 427 ± 171 ^j vs 716 ± 336	Favours aprotinin <0.003	NA
Later (2009)	Level II Good	1 RCT N=199	Adult patients undergoing <u>low-or intermediate-risk</u> cardiac surgery	Hospital – planned surgery The Netherlands	High-dose aprotinin (IV) vs placebog	Mediastinal chest tube drain loss (mL)	MD -295 (-410, -185)	Favours aprotinin <0.001	NA
Nurözler (2008)	Level II Fair	1 RCT N=51	Adult patients undergoing off- pump coronary bypass who have received clopidogrel within 5 days of surgery (all patients)	Hospital – planned surgery Turkey	<u>Low-dose</u> aprotinin (IV) vs placebo ^k	Drainage (mL/24 hr)	423 ± 178 vs 748 ± 212	Favours aprotinin 0.005	NA
Orthopaedic surg	ery				•	•			
Henry (2007)	Level I Good	10 RCTs N=430	Adult patients undergoing orthopaedic surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Total blood loss (mL)	WMD -399 (-563, -235)	Favours aprotinin <0.001	Substantial Phet=0.01 (I ² =60%)
		5 RCTs N=201	Adult patients undergoing orthopaedic surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Intraoperative blood loss (mL)	WMD -151 (-318, 16)	No difference 0.076	Moderate Phet=0.16 (I ² =40%)
		7 RCTs N=318	Adult patients undergoing orthopaedic surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Postoperative blood loss (mL)	WMD -114 (-224, -3.5)	Favours aprotinin 0.043	Substantial Phet=0.005 (l²=68%)
Kagoma (2009)	Level I Good	4 RCTs N=230	Adults undergoing total knee or hip replacement	Hospital – planned surgery Unknown	Aprotinin (IV) vs no aprotinin	Total bleeding ^I (mL)	WMD -639 (-725, -536)	Favours aprotinin NR	NR

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or Aprotinin (mean ± SD) vs control (mean ± SD)	Significance P-value	
Colwell (2007)	Level II Good		Adults undergoing <u>unilateral</u> total hip arthroplasty (all	Hospital – planned surgery	Aprotinin (IV) vs placebo	Intraoperative blood loss (mL)	331 vs 385	Favours aprotinin 0.0217	NA
			patients)	US/Canada		0-6 hr drainage (mL)	96 vs 177	Favours aprotinin 0.0003	NA
		Tot		Total drainage (mL)	276 vs 390	Favours aprotinin 0.0141	NA		
						Total fluid loss (mL)	709 vs 957	Favours aprotinin 0.0002	NA
Liver surgery									
Henry (2007)	Level I Good	2 RCTs N=137	Adult patients undergoing <u>liver</u> <u>surgery</u>	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Intraoperative blood loss (mL)	WMD -1200 (-2943, -543)	Favours aprotinin 0.02	Substantial Phet=0.02 (l²=67%)
	Level I/II Fair	1 RCT N=24	Adult patients undergoing <u>liver</u> surgery	Hospital – planned surgery Unknown	Aprotinin (IV) vs no aprotinin	Postoperative blood loss (mL)	WMD -105 (-194, -16)	Favours aprotinin 0.021	NA
Gurusamy (2009)	Level I/II Good/Fair	1 RCT N=97	Adult patients undergoing liver resection	Hospital – planned surgery Unknown	Aprotinin (IV) vs placebo	Operative blood loss (mL)	WMD -436 (-874, 1.67)	No difference 0.051	NA
Other surgery		1	-	<u> </u>	•	1	1	•	•
Henry (2007)	Level I/II Good/Poor	1 RCT N=30	Adult patients undergoing orthognathic surgery	Hospital – planned surgery Unknown	Aprotinin (IV) vs no aprotinin	Postoperative blood loss (mL)	WMD -513 (-717, -309)	Favours aprotinin <0.001	NA
Henry (2007)	Level I/II Good/Fair	1 RCT N=24	Adult patients undergoing thoracic surgery	Hospital – planned surgery Unknown	Aprotinin (IV) vs no aprotinin	Intraoperative blood loss (mL)	WMD -532 (-863, -199)	Favours aprotinin 0.0016	NA
				Ulikilowii		Postoperative blood loss (mL)	WMD -441 (-786, -96)	Favours aprotinin 0.012	NA
Apostolakis (2008)	Level II Fair	1 RCT N=59	Adult patients undergoing thoracic surgery	Hospital – planned surgery Greece	Ultra-low-dose aprotinin (IV) vs placebom	Day 1 postoperative thoracic drainage (mL)	413 ± 199 vs 764 ± 214	Favours aprotinin <0.001	NA
						Day 2 postoperative thoracic drainage (mL)	248 ± 179 vs 455 ± 275	Favours aprotinin 0.001	NA
Henry (2007)	Level II Good/Good	1 RCT N=50	Adult patients undergoing vascular surgery	Hospital – planned surgery Unknown	Aprotinin (IV) vs no aprotinin	Postoperative blood loss (mL)	WMD -203 (-405, -1.07)	Favours aprotinin 0.049	NA

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or Aprotinin (mean ± SD) vs control (mean ± SD)	Significance P-value	
Leijdekkers (2006)	Level II Fair	1 RCT N=35	Adult patients undergoing surgery for infra-renal abdominal aneurysm	Hospital – planned surgery The Netherlands	Aprotinin (IV) vs placebo	Mean blood loss (mL)	2362 ± 1340 vs 2466 ± 1370	No difference 0.88	NA
PAEDIATRIC POPULA	TION/IV APROTINI	N							
Orthopaedic surge	ry								
Schouten (2009)	Level II Fair	1 RCT N=44	Paediatric patients undergoing scoliosis surgery	Hospital – planned surgery Unknown	Aprotinin (IV) vs no aprotinin	Blood loss (mL)	WMD -385 (-727, -42)	Favours aprotinin NR	NA
Tzortzopoulou (2008)	Level I Good	2 RCTs N=87	Paediatric patients undergoing scoliosis surgery	Hospital – planned surgery Unknown	Aprotinin <u>(IV)</u> vs placebo	Blood loss (mL)	WMD -450 (-726, -174)	Favours aprotinin 0.0014	None Phet=0.53 (I ² =0%)
ADULT POPULATION/	TOPICAL APROTIN	IN							
Cardiac surgery									
Abrishami (2009)	Level I Good	5 RCTs N=324	Adult patients undergoing primary on-pump cardiac surgery	Hospital – planned surgery Unknown	Aprotinin (topical) vs placebo	24-hr <u>postoperative</u> chest tube blood loss (mL)	WMD -204 (-276, -132)	Favours aprotinin <0.001	Substantial Phet=0.04 (I ² =60%)
Mehraien (2009)	Level II Good	1 RCT N=128	Adult patients undergoing first- time CABG (all patients)	Hospital – planned surgery Iran	Aprotinin (topical) vs placebo	24-hr postoperative chest tube blood loss (mL)	451 ± 218 vs 707 ± 269	Favours aprotinin 0.003	

Abbreviations: ASA, acetylsalicylic acid; CI, confidence interval; het, heterogeneity; IV, intravenous; mL, millilitres; NA, not applicable; NR, not reported; RCT, randomised controlled trial; SMD, standardised mean difference; WMD, weighted mean difference.

- ^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.
- b Heterogeneity defined as follows: (i) no significant heterogeneity if Pheto 0.1 and 12<25%; (ii) mild heterogeneity if 12<25%; moderate heterogeneity if 12 between 25-50%; substantial heterogeneity 12 > 50%.
- Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between aprotinin and no aprotinin.
- d Prime' dose included regimens that added aprotinin to the pump prime solution of the cardiopulmonary bypass exclusively. 12/16 trials studied a 'prime' dose of 2 million KIU, 2/16 trials studied a 'prime' dose of 500,000 KIU and 1/16 trials studies a 'prime' dose of 25,000 KIU/kg.
- c Low-dose aprotinin was defined as any regimen that did not follow the 'full Hammersmith' regimen, including those studies that described their regimen as 'half Hammersmith'. For non-cardiac surgery trials, regimens were classified as low dose if the total dose was < 5 million KIU or 700 mg aprotinin.
- f High-dose aprotinin was defined as any regimen that was described as the 'full Hammersmith' regimen. For non-cardiac surgery trials, regimens were classified as high-dose if the total dose was ≥ 5 million KIU or 700 mg aprotinin.
- 9 High dose (full-dose) aprotinin defined as a 2 million kallikrein-inhibiting units (KIU) IV loading dose, 2 million KIU pump-priming dose, and 0.5 million KIU IV/h maintenance dose.
- Low dose (half-dose) aprotinin consisted of a 1 million KIU IV loading dose, 1 million KIU pump-priming dose, and 0.25 million KIU IV/h maintenance dose.
- Patients with peak aprotinin levels > 271 KIU/mL.
- i Patients with peak aprotinin levels < 271 KIU/mL.
- Low dose aprotinin consisted of 1 million KIU infused over 30 min followed by a continuous infusion of 0/5 million KIU/h until the end of surgery.
- ¹ Total bleeding measured intraoperatively by weighing surgical sponges, posioperatively through drainage or perioperatively through the haemoglobin balance method which measures loss through comparison of pre- and postoperative haemoglobin concentrations (haematoma volumes as well as hidden or internal blood loss were excluded).
- m Ultra-low dose aprotinin defined as a test dose of 1mL at intubation, followed by 0.5 million IU over 15 min following intubation, and again following closure.

Key question(s):			Evidence table refa:			
In patients undergoing surgery, what is the effect of administration of aprotining	<u>n</u> on	mortality?	POQ3.I8a.P4			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	;)					
There is one pivotal Level I study (Henry 2007/good quality) which includes data from up to 37 RCTs (31 RCTs for cardiac surgery), two supportive level I studies (Henry 2009/good quality; Brown 2007/fair quality), one supportive	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias			
Level I/II study (Gurusamy 2009/good-fair quality) and five additional RCTs (Grant 2008/fair quality; Later	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias			
2009/good quality; Colwell 2007/good quality; Apostolakis 2008/fair quality; Leijdekkers 2006/fair quality) which were published after the Henry 2007 review and were not included in the supportive reviews.	С	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Results mostly consistent between pivotal and supportive meta-analyses. Consistency of individual studies within the pivotal meta-analysis described below. Some difference between high and low dose aprotinin in Brown 2007.	Α	All studies consistent				
Additional RCTs consistent.	В	Most studies consistent and inconsistency can be explained				
Pivotal evidence – Henry 2007 No heterogeneity between studies.	С	Some inconsistency, reflecting genuine uncertainty around question	on			
Supportive evidence - see Summary Table POQ3.l8a.P4	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>ur</u>	nknowi	<u>n</u> factor (not simply study quality or sample size) and thus the clinical impa	ct of the intervention could not be			
Pivotal evidence – Henry 2007 add percentages	Α	Very large				
Any surgery – RR 0.90 (0.67, 1.20); 2.4% vs 2.6%; 37 RCTs (N=6645) Cardiac surgery – RR 0.95 (0.70, 1.28); 2.5% vs 2.4%; 31 RCTs (N=6058)	В	Substantial				
Supportive evidence – Brown 2007 (for others see Summary Table POQ3.l8a.P4) Cardiac surgery (high dose) – RR 0.89 (0.65, 1.21); 43 RCTs (N=6175)	С	Moderate				
Cardiac surgery (light dose) – RR 0.67 (0.65, 1.21), 45 RCTs (N=0175) Cardiac surgery (low dose) – RR 1.37 (0.72, 2.59); 14 RCTs (N=786)	D	Underpowered				
4. Generalisability (How well does the body of evidence match the population and clinical set	tings t	peing targeted by the Guideline?)				
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were included in the overall analysis, although most were conducted in cardiac surgery.	Α	Evidence directly generalisable to target population				
included in the overall analysis, although most were conducted in cardiac surgery.	В	Evidence directly generalisable to target population with some cav	eats			
	С	Evidence not directly generalisable to the target population but cou	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to judge whether it is sensible to				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of heal	th services/delivery of care and cultural factors?)				
Hospital setting. The pivotal review states that studies were conducted in a wide range of countries.	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few cave	ats			
	С	Evidence probably applicable to Australian healthcare context with some caveats				
	D	Evidence not applicable to Australian healthcare context				

In 2007, Bayer announced a worldwide suspension of aprotinin supply due to the results of the BART trial which suggested increased risk of mortality for aprotinin compared with the lysine analogues tranexamic acid and ε-aminocaproic acid.

Mortality was underpowered in the individual studies but the Henry 2007 review included 6645 patients in total. Results from Brown 2007 shows a slight difference in direction of effect depending on dose (ie, high or low).

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (good quality), two supportive level I studies, one supportive Level I/II study ^b and five additional RCTs.
2. Consistency	С	There was no heterogeneity in the pivotal level I evidence. Most additional studies were consistent. There was some difference in the direction of effect due to dose in one of the supportive level I studies.
3. Clinical impact	D	While there is no significant difference in mortality between intravenous aprotinin therapy and no therapy, and the risk estimates suggest no increased risk, the findings are uncertain due to underpowering.
4. Generalisability	В	The results are generalisable to an cardiac adult surgical population; most studies were conducted in cardiac surgery.
5. Applicability	В	Overall there were a large number of studies conducted in a wide range of countries. Likely to be applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, the effect of intravenous aprotinin therapy on mortality, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: Cl. confidence interval. RBC, red blood cell: SR. systematic review:

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission b A Level I/II study is a systematic review which includes only one RCT.

^a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.l8a.P4 Characteristics and results of studies examining the effect of <u>aprotinin</u> on <u>mortality</u>

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size	Surgical procedure	Location			Risk estimate (95% CI) or AP (%) vs control (%)	Significance P-value	
ADULT POPULATION Any surgery	/IV APROTININ								
Henry (2007)	Level I Good	37 RCTs N=6645	Adult patients undergoing any surgery	Hospital – planned surgery Various countries ^c	Aprotinin (IV) vs no aprotinin	Mortality	RR 0.90 (0.67, 1.20)	No difference 0.47	None Phet=0.95 (l ² =0%)
Cardiac surgery	<u>'</u>	<u>'</u>		1	1			•	•
Henry (2007)	Level I Good	31 RCTs N=6058	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries ^c	Aprotinin (IV) vs no aprotinin	Mortality	RR 0.95 (0.70, 1.28)	No difference 0.72	None Phet=0.93 (I ² =0%)
Henry (2009)	Level I Good	32 RCTs N=6279	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries ^c	Aprotinin (IV) vs placebo	Mortality	RR 0.93 (0.69, 1.25)	No difference NR	NR
Brown (2007)	Level I Fair	43 RCTs N=6175	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	High dose aprotinin (IV) vs placebod	Mortality	RR 0.89 (0.65, 1.21)	No difference 0.46	NR
		14 RCTs N=1453	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	Low dose aprotinin (IV) vs placeboe	Mortality	RR 1.37 (0.72, 2.59)	No difference 0.34	NR
Grant (2008)	Level II Fair	1 RCT N=120	Adult patients undergoing off-pump coronary artery bypass surgery	Hospital – planned surgery US	Aprotinin (IV) vs placebo	1-year_mortality	5.1% vs 13.1%	No difference NS	NA
Later (2009)	Level II Good	1 RCT N=199	Adult patients undergoing low- to intermediate-risk cardiac surgery	Hospital – planned surgery The Netherlands	High dose aprotinin (IV) vs placebod	In-hospital mortality	2.1% vs 1.0%	No difference 0.61	NA
Orthopaedic surg	ery								
Colwell (2007)	Level II Good	1 RCT N=352	Adults undergoing <u>unilateral</u> total hip arthroplasty (all patients)	Hospital – planned surgery US/Canada	Aprotinin (IV) vs placebo	Mortality (up to 6 weeks)	0% vs 0.6%	No difference NS	NA
Liver surgery									
Gurusamy (2009)	Level I/II Good/Fair	1 RCT N=37	Adult patients undergoing liver resection	Hospital – planned surgery Unknown	Aprotinin (IV) vs placebo	Mortality	RR 1.18 (0.18, 7.48)	No difference 0.86	NA

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b	
	evidence ^a sample size Surgical procedure Location		Risk estimate (95% CI) or AP (%) vs control (%)	Significance P-value						
Other surgery	Other surgery									
Apostolakis (2008)	Level II Fair	1 RCT N=59	Adult patients undergoing thoracic surgery	Hospital – planned surgery Greece	Ultra-low-dose aprotinin (IV) vs placebof	In-hospital mortality	0% vs 0%	No difference NA	NA	
Leijdekkers (2006)	Level II Fair	1 RCT N=35	Adult patients undergoing surgery for infra-renal abdominal aneurysm	Hospital – planned surgery The Netherlands	Aprotinin (IV) vs placebo	In-hospital mortality	6.3% vs 5.3%	No difference 1.00	NA	

Abbreviations: CI, confidence interval; het, heterogeneity; IV, intravenous; NA, not applicable; NR, not reported; RCT, randomised controlled trial; RR, risk ratio.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I²<25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between aprotinin and no aprotinin.

d High dose (full-dose) aprotinin defined as a 2 million kallikrein-inhibiting units (KIU) IV loading dose, 2 million KIU pump-priming dose, and 0.5 million KIU IV/h maintenance dose.

Low dose (half-dose) aprotinin consisted of a 1 million KIU IV loading dose, 1 million KIU pump-priming dose, and 0.25 million KIU IV/h maintenance dose.

^{&#}x27;Ultra-low dose aprotinin defined as a test dose of 1mL at intubation, followed by 0.5 million IU over 15 min following intubation, and again following closure.

Key question(s): In patients undergoing surgery, what is the effect of administration of aprotining approximation of approxim	<u>1</u> on	morbidity (coronary artery graft occlusion)?	Evidence table refa: POQ3.I8a.P5 (CAGO)			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	5)					
There is one pivotal Level I study (Henry 2007/good quality) which includes 2 RCTs, and one additional RCT (Grant 2008/fair quality).	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias			
2000riali quality).	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias				
	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Pivotal evidence – Henry 2007 Significant heterogeneity between two included studies (l ² =56%).	Α	All studies consistent				
Supportive evidence – Grant 2008	В	Most studies consistent and inconsistency can be explained				
Additional RCT results consistent with one of the RCTs included in the Henry review.	С	Some inconsistency, reflecting genuine uncertainty around question				
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>undetermined</u>)	nknowi	<u>n</u> factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be			
Pivotal evidence – Henry 2007 Cardiac surgery – RR 0.76 (0.10, 5.67); 2 RCTs (N=728)	Α	Very large				
Supportive evidence – Grant 2008	В	Substantial				
Off-pump CABG surgery (saphenous vein grafts) – 3.8% vs 8.9%; 1 RCT (N=120)	С	Moderate				
	D	Underpowered/inconsistent				
4. Generalisability (How well does the body of evidence match the population and clinical set	tings b	peing targeted by the Guideline?)				
The evidence is generalisable to an adult population who are undergoing planned coronary artery bypass surgery.	Α	Evidence directly generalisable to target population				
	В	Evidence directly generalisable to target population with some cav	eats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to judge whether it is sensible to				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of heal					
Hospital setting. There was one large multinational trial and one small US trial included in the pivotal level I study. The additional RCT was conducted in the US. The evidence may be applicable to the Australian setting.	Α	Evidence directly applicable to Australian healthcare context				
The database for the solution of the objective may be approache to the ridstrainant setting.	В	Evidence applicable to Australian healthcare context with few cave				
	С	Evidence probably applicable to Australian healthcare context with	some caveats			
	D	Evidence not applicable to Australian healthcare context				

In 2007, Bayer announced a worldwide suspension of aprotinin supply due to the results of the BART trial which suggested increased risk of mortality for aprotinin compared with the lysine analogues tranexamic acid and ε-aminocaproic acid.

CAGO not specifically defined in Henry review. CRG concerned regarding definition of CAGO in individual studies. The results of the pivotal review showed substantial heterogeneity and wide confidence intervals; therefore, the effect of aprotinin therapy on CAGO was considered uncertain.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
Evidence base	Α	There is one pivotal Level I study (good quality) which includes 2 RCTs, and one additional RCT.
2. Consistency	С	The results of the two trials included in the pivotal level I study were inconsistent. Additional RCT results consistent with one small RCT included in the pivotal level I study.
Clinical impact	D	While there is no significant difference in coronary artery graft occlusion between intravenous aprotinin therapy and no therapy, the findings are uncertain due to inconsistency and underpowering.
4. Generalisability	А	The results are generalisable to an adult population undergoing coronary artery bypass graft.
5. Applicability	С	There was one large multinational trial and one small US trial included in the pivotal level I study. The additional RCT was conducted in the US. The evidence may be applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing coronary artery bypass surgery, the effect of intravenous aprotinin therapy on coronary artery graft occlusion, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI, confidence interval, RBC, red blood cell: SR, systematic review;

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.l8a.P5 (CAGO) Characteristics and results of studies examining the effect of aprotinin on morbidity (coronary artery graft occlusion)

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b		
	evidence ^a Quality	sample size	Surgical procedure				Risk estimate (95% CI)	Significance p-value			
ADULT POPULATION	DULT POPULATION/IV APROTININ										
Cardiac surgery	Cardiac surgery										
Henry (2007)	Level I Good	2 RCTs N=728	Adult patients undergoing cardiac surgery	Hospital – planned surgery Multinational and US	Aprotinin (IV) vs no aprotinin	Coronary artery graft occlusion	RR 0.76 (0.10, 5.67)	No difference 0.79	Substantial Phet=0.13 (l²=56%)		
Grant (2008)	Level II Fair	1 RCT N=120	Adult patients undergoing off-pump coronary artery bypass surgery	Hospital – planned surgery US	Aprotinin (IV) vs placebo	6-month acute occlusion	3.8% vs 8.9% (SVGs)	No difference NS	NA		

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: CI, confidence interval; DVT, deep vein thrombosis; het, heterogeneity; IV, intravenous; MI, myocardial infarction; NA, not applicable; NR, not reported; PE, pulmonary embolism; OR, odds ratio; RCT, randomised controlled trial; RD, risk difference; RR, risk ratio; SVG, saphenous vein graft.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

Key question(s):			Evidence table refa:		
In patients undergoing surgery, what is the effect of administration of aprotini	<u>n</u> on <u>l</u>	morbidity (myocardial infarction)?	POQ3.18a.P5 (MI)		
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	5)				
There is one pivotal Level I study (Henry 2007/good quality), which included data from 34 RCTs (31 RCTs for cardiac surgery), one supportive level I study (Brown 2007/fair quality) and four additional RCTs, three in cardiac	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias			
surgery (Grant 2008/fair quality; Later 2009/good quality/Nurôzler 2008/fair quality) and one in hip replacement surgery (Colwell 2007/good quality).	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias		
	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias		
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias			
2. Consistency (If only one study was available, rank this component as 'not applicable')					
Results consistent between Henry and Brown reviews. However, a slight difference in direction of the point estimate by dose was seen in the Brown review. Additional RCTs consistent.	Α	All studies consistent			
Pivotal evidence – Henry 2007	В	Most studies consistent and inconsistency can be explained			
No heterogeneity between studies.	С	Some inconsistency, reflecting genuine uncertainty around question	on		
	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>undetermined</u>)	nknowi	n factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be		
Pivotal evidence – Henry 2007	Α	Very large			
Any surgery – RR 0.92 (0.72, 1.18); 4.3% vs 4.6%; 34 RCTs (N=5758) Cardiac surgery – RR 0.95 (0.74, 1.22); 4.7% vs 4.7%; 31 RCTs (N=5279)	В	Substantial			
Supportive evidence – Brown 2007 (see also Supportive Table POQ3.18a.P5(MI)) Cardiac surgery (high dose) – RR 1.10 (0.83, 1.45); 31 RCTs (N=3315)	С	Moderate			
Cardiac surgery (low dose) – RR 1.10 (0.88, 1.49), 31 RC1s (N=1585) Cardiac surgery (low dose) – RR 0.94 (0.58, 1.54); 16 RCTs (N=1585)	D	No difference			
4. Generalisability (How well does the body of evidence match the population and clinical set	tings b	eing targeted by the Guideline?)			
The evidence is generalisable primarily to an adult population who are undergoing planned cardiac surgery: 31/34 studies included in the pivotal review were conducted in cardiac surgery, as were 3/4 RCTs. 1 recent RCT provides	Α	Evidence directly generalisable to target population			
data on hip replacement surgery.	В	Evidence directly generalisable to target population with some cav	eats		
	С	Evidence not directly generalisable to the target population but cou	uld be sensibly applied		
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to		
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of heal	th services/delivery of care and cultural factors?)			
Hospital setting. There are a large number of RCTs in the Henry review from a wide range of countries. It is unclear where these studies were conducted. The one additional recent RCT in hip replacement surgery was conducted in	Α	Evidence directly applicable to Australian healthcare context			
the US and Canada.	В	Evidence applicable to Australian healthcare context with few cave	eats		
	С	Evidence probably applicable to Australian healthcare context with	some caveats		
	D	Evidence not applicable to Australian healthcare context			

In 2007, Bayer announced a worldwide suspension of aprotinin supply due to the results of the BART trial which suggested increased risk of mortality for aprotinin compared with the lysine analogues tranexamic acid and ε-aminocaproic acid.

MI had a higher incidence than mortality (- 4% in Henry 2007), so the results presented in the Henry review for MI are more likely to be adequately powered than they were for mortality. Also had narrower confidence intervals than mortality. CRG concerned regarding the definition of MI in the individual included studies.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
Evidence base	А	There is one pivotal Level I study (good quality), one supportive level I study and four additional RCTs (three in cardiac surgery and one in hip replacement surgery).
2. Consistency	В	There was no heterogeneity in the pivotal level I evidence. Most additional studies were consistent. There was a slight difference in the direction of effect due to dose in the supportive level I studies.
3. Clinical impact	D	For cardiac surgery, there is no significant difference in the risk of myocardial infarction between intravenous aprotinin therapy and no therapy. For hip replacement surgery there was only RCT which was underpowered to detect a difference.
4. Generalisability	В	The results are generalisable primarily to cardiac surgery. There was also a recent RCT in hip replacement surgery.
5. Applicability	В	Studies were conducted in a wide range of countries. Likely to be applicable to the Australian setting. The single included RCT in hip replacement surgery was conducted in the US and Canada. Likely to be applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above

In adult patients undergoing cardiac surgery, intravenous aprotinin therapy does not appear to have an effect on the risk of myocardial infarction compared with no therapy.

In adult patients undergoing hip replacement surgery, the effect of intravenous aprotinin therapy on the risk of myocardial infarction, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI, confidence interval, RBC, red blood cell; SR, systematic review;

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.l8a.P5 (MI) Characteristics and results of studies examining the effect of <u>aprotinin</u> on <u>morbidity (myocardial infarction)</u>

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size	Surgical procedure				Risk estimate (95% CI) or AP (%) vs control (%)	Significance P-value	
ADULT POPULATION	/IV APROTININ								
Any surgery						1			
Henry (2007)	Level I Good	34 RCTs N=5758	Adult patients undergoing any surgery	Hospital – planned surgery Various countries ^c	Aprotinin (IV) vs no aprotinin	Myocardial infarction	RR 0.92 (0.72, 1.18)	No difference 0.50	None Phet=0.91 (l²=0%)
Cardiac surgery		•							
Henry (2007)	Level I Good	31 RCTs N=5279	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries ^c	Aprotinin (IV) vs no aprotinin	Myocardial infarction	RR 0.95 (0.74, 1.22)	No difference 0.69	None Phet=0.92 (l²=0%)
Henry (2009)	Level 1 Good	34 RCTs N=5441	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries ^c	Aprotinin (IV) vs placebo	Myocardial infarction	RR 0.94 (0.73, 1.21)	No difference NR	NR
Brown (2007)	Level I Fair	31 RCTs N=3315	Adult patients undergoing cardiac surgery	Hospital – planned surgery Countries not specified	High dose aprotinin (IV) vs placebod	Myocardial infarction	RR 1.10 (0.83, 1.45)	No difference 0.52	NR
		16 RCTs N=1585	Adult patients undergoing cardiac surgery	Hospital – planned surgery Countries not specified	Low dose aprotinin (IV) vs placeboe	Myocardial infarction	RR 0.94 (0.58, 1.54)	No difference 0.82	NR
Grant (2008)	Level II Fair	1 RCT N=120	Adult patients undergoing off-pump coronary artery bypass surgery	Hospital – planned surgery US	Aprotinin (IV) vs placebo	In-hospital myocardial infarction	1.7% vs 6.6%	No difference NS	NA
Later (2009)	Level II Good	1 RCT N=199	Adult patients undergoing low- to intermediate-risk cardiac surgery	Hospital – planned surgery The Netherlands	High dose aprotinin (IV) vs placebod	Perioperative myocardial infarction	1.0% vs 7.8%	Favours aprotinin 0.023	NA
Nurözler (2008)	Level II Fair	1 RCT N=51	Adult patients undergoing off-pump coronary bypass who have received clopidogrel within 5 days of surgery (all patients)	Hospital – planned surgery Turkey	Low-dose aprotinin (IV) vs placebo ^r	Myocardial infarction	0% vs 0%	No difference NA	NA

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results	Heterogeneity ^b	
	evidence ^a <i>Quality</i>	sample size	Surgical procedure				Risk estimate (95% CI) or AP (%) vs control (%)	Significance P-value	
Orthopaedic surge	ry								
Colwell (2007)	Level II Good	1 RCT N=352	Adults undergoing <u>unilateral</u> total hip arthroplasty (all patients)	Hospital – planned surgery US/Canada	Aprotinin (IV) vs placebo	Myocardial infarction	0.6% vs 0.6%	No difference NS	NA

Abbreviations: CI, confidence interval; het, heterogeneity; IV, intravenous; NR, not reported; RCT, randomised controlled trial; RR, risk ratio.

a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

^b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and |²<25%; (ii) mild heterogeneity if |² between 25-50%; substantial heterogeneity |² >50%.

c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between aprotinin and no aprotinin.

d High dose (full-dose) aprotinin defined as a 2 million kallikrein-inhibiting units (KIU) IV loading dose, 2 million KIU pump-priming dose, and 0.5 million KIU IV/h maintenance dose.

e Low dose (half-dose) aprotinin consisted of a 1 million KIU IV loading dose, 1 million KIU pump-priming dose, and 0.25 million KIU IV/h maintenance dose.

Low dose aprotinin consisted of 1 million KIU infused over 30 min followed by a continuous infusion of 0/5 million KIU/h until the end of surgery.

Key question(s):			Evidence table refa:	
In patients undergoing surgery, what is the effect of administration of aprotini	<u>n</u> on <u>i</u>	morbidity (renal failure/dysfunction)?	POQ3.I8a.P5 (renal)	
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	5)			
There is one pivotal Level I study (Henry 2007/good quality) which included data from 14 RCTs (11 RCTs for cardiac surgery), one supportive level I study (Brown 2007/fair quality) and three additional RCTs (Grant 2008/fair	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias	
quality; Later 2009/good quality; Colwell 2007/good quality).	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias	
	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias	
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias		
2. Consistency (If only one study was available, rank this component as 'not applicable')				
Results mostly consistent between the Henry and Brown reviews. However, significant renal dysfunction was seen for high dose aprotinin in Brown review. Most additional RCTs consistent, although Later 2009 showed less renal	Α	All studies consistent		
complications in aprotinin arm.	В	Most studies consistent and inconsistency can be explained		
Pivotal evidence – Henry 2007 No heterogeneity between studies.	С	Some inconsistency, reflecting genuine uncertainty around question	on	
la nota oganos, postación ocuación	D	Evidence is inconsistent		
	NA	Not applicable (one study only)		
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>undetermined</u>)	nknowi	n factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be	
Pivotal evidence – Henry 2007	Α	Very large		
Any surgery (renal failure/dysfunction) – RR 1.16 (0.79, 1.70); 3.0% vs 2.2%; 14 RCTs (N=3908) Cardiac surgery (renal failure/dysfunction) – RR 1.12 (0.74, 1.67); 2.9% vs 2.2%; 11 RCTS (N=3670)	В	Substantial		
Supportive evidence – Brown 2007 (see also Supportive Table POQ3.l8a.P5(renal) Cardiac surgery (renal failure; high dose) – RR 1.09 (0.68, 1.77); 27 RCTs (N=4681)	С	Moderate (renal dysfunction)		
Cardiac surgery (renal dalure, high dose) – RR 1.07 (0.06, 1.77), 27 RC1s (N=4001) Cardiac surgery (renal dysfunction; high dose) – RR 1.47 (1.12, 1.94); 19 RCTs (N=1778)	D	Slight (renal failure)		
4. Generalisability (How well does the body of evidence match the population and clinical set	tings b	eing targeted by the Guideline?)		
The evidence is generalisable to an adult population who are undergoing planned surgery. 11/14 studies included in the pivotal review were conducted in cardiac surgery.	Α	Evidence directly generalisable to target population		
une pivotai review were conducted in cardiac surgery.	В	Evidence directly generalisable to target population with some cav	eats	
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied	
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to	
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of heal	th services/delivery of care and cultural factors?)		
Hospital setting. Studies conducted in a wide range of countries.	Α	Evidence directly applicable to Australian healthcare context		
	В	Evidence applicable to Australian healthcare context with few cave	eats	
	С	Evidence probably applicable to Australian healthcare context with	some caveats	
	D	Evidence not applicable to Australian healthcare context		

In 2007, Bayer announced a worldwide suspension of aprotinin supply due to the results of the BART trial which suggested increased risk of mortality for aprotinin compared with the lysine analogues tranexamic acid and ε-aminocaproic acid.

CRG noted that eGFR calculation is not a good measure and only to be used in stable patients, not post-surgery. Grant used CT angiography with 100-150 mL contrast which may explain if baseline levels were high. Varying definitions between studies; some renal endpoints were soft.

Later 2009 RCT defined outcomes as follows: Renal failure as defined by Mangano (2006): required a postoperative serum creatinine of at least 2.0 mg/dL with an increase over the preoperative baseline level of at least 0.7 mg/dL; renal complication as defined by the RIFLE classification: risk of renal dysfunction defined as a 1.5 times increase in perioperative creatinine plasma concentration or a urine output < 0.5 mL/kg/h in 6 hours. Kidney injury was defined as a 2 times increase in perioperative creatinine plasma concentration or a urine output < 0.5 mL/kg/h in 12 hours.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating		Description
	Renal failure	Renal dysfunction	
1. Evidence base	А	А	There is one pivotal Level I (good quality) study, one supportive level I study and three additional RCTs.
2. Consistency	С	С	There is some inconsistency regarding dose in the supportive level I study and for renal dysfunction/complications which may be due to the definitions used in different studies.
3. Clinical impact	D		There was no significant difference between intravenous aprotinin therapy and no therapy for renal failure. There were inconsistent results on the basis of dose for renal dysfunction/complications (C for dysfunction, D for failure).
4. Generalisability	С	С	The results are generalisable to an adult surgical population but most studies were in cardiac surgery.
5. Applicability	В	1 1)	Overall there were a large number of studies conducted in a wide range of countries. Additional individual RCTs were conducted in the US, Canada and the Netherlands. Likely to be applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, intravenous aprotinin therapy does not appear to affect the risk of postoperative renal failure, compared with no therapy, but may impair postoperative renal function.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI, confidence interval, RBC, red blood cell: SR, systematic review;

^a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

POQ3.l8a.P5 (renal) Characteristics and results of studies examining the effect of <u>aprotinin</u> on <u>morbidity (renal failure/dysfunction)</u>

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	come Results		
	evidence ^a Quality	sample size	Surgical procedure	Location			Risk estimate (95% CI) or AP (%) vs control (%)	Significance P-value	
ADULT POPULATION	n/ IV aprotinin								
Any surgery									
Henry (2007)	Level I Good	14 RCTs N=3908	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Renal failure/dysfunction	RR 1.16 (0.79, 1.70)	No difference 0.46	None Phet=0.88 (I ² =0%)
Cardiac surgery		•							
Henry (2007)	Level I Good	11 RCTs N=3670	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries ^c	Aprotinin (IV) vs no aprotinin	Renal failure/dysfunction	RR 1.12 (0.74, 1.67)	No difference 0.60	None Phet=0.85 (I ² =0%)
Brown (2007)	Level I Fair	27 RCTs N=4681	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	High dose aprotinin (IV) vs placebod	Renal failure	RR 1.09 (0.68, 1.77)	No difference 0.71	NR
		7 RCTs N=786	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	Low dose aprotinin (IV) vs placebof	Renal failure	RR 1.86 (0.07, 49.26)	No difference 0.71	NR
Brown (2007)	Level I Fair	19 RCTs N=1778	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	High dose aprotinin (IV) vs placebod	Renal dysfunction ⁹	RR 1.47 (1.12, 1.94)	Favours no aprotinin 0.006	NR
		9 RCTs N=1041	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	Low dose aprotinin (IV) vs placebof	Renal dysfunction ^g	RR 1.01 (0.69, 1.49)	No difference 0.961	NR
Grant (2008)	Level II Fair	1 RCT N=120	Adult patients undergoing off-pump coronary artery bypass	Hospital – planned surgery US	Aprotinin (IV) vs placebo	Acute renal failure within 6 months ^h	3.4% vs 3.3%	No difference NS	NA
			surgery			Postoperative acute kidney injury	45.8 vs 24.6	Favours no aprotinin <0.03	NA
Later (2009)	Level II Good	1 RCT N=199	Adult patients undergoing low- to intermediate-risk cardiac	Hospital – planned surgery The Netherlands	High dose aprotinin (IV) vs placebod	Renal failure	3.1% vs 2.9%	No difference 1.0	NA
		surgery	The Netherlands		Renal complication ^k	10.4% vs 17.5%	Favours aprotinin 0.011	NA	

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results	Heterogeneity ^b	
	evidence ^a <i>Quality</i>	sample size	Surgical procedure	Location			Risk estimate (95% CI) or AP (%) vs control (%)	Significance P-value	
Orthopaedic surge	ery								
Colwell (2007)	Level II Good	1 RCT N=352	Adults undergoing <u>unilateral</u> total hip arthroplasty (all patients)	Hospital – planned surgery US/Canada	Aprotinin (IV) vs placebo	Renal failure (not defined)	1.1% vs 1.1%	No difference NS	NA

Abbreviations: CI, confidence interval; GFR, glomerular filtration rate; het, heterogeneity; IV, intravenous; NR, not reported; RCT, randomised controlled trial; RR, risk ratio.

- a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.
- b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and 12<25%; (ii) mild heterogeneity if 12 <25%; moderate heterogeneity if 12 between 25-50%; substantial heterogeneity 12 >50%.
- Estudies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between aprotinin and no aprotinin.
- d High dose (full-dose) aprotinin defined as a 2 million kallikrein-inhibiting units (KIU) IV loading dose, 2 million KIU pump-priming dose, and 0.5 million KIU IV/h maintenance dose.
- e Renal failure defined as a new onset of dialysis except in one study where it was defined as a ≥ 2 mg/dL creatinine level.
- Low dose (half-dose) aprotinin consisted of a 1 million KIU IV loading dose, 1 million KIU pump-priming dose, and 0.25 million KIU IV/h maintenance dose.
- ^g Renal dysfunction defined as a ≥ 0.5 mg/dL increase in creatinine.
- h Acute renal failure defined as postoperative eGFR < 75% of baseline and urine output is < 0.5 mL/kg/h for 6 hours.
- Postoperative kidney injury defined as postoperative eGFR < 75% of baseline.
- Renal failure as defined by Mangano (2006). Required a postoperative serum creatinine of at least 2.0 mg/dL with an increase over the preoperative baseline level of at least 0.7 mg/dL.
- kRenal complication as defined by the RIFLE classification². Risk of renal dysfunction defined as a 1.5 times increase in perioperative creatinine plasma concentration or a urine output < 0.5 mL/kg/h in 6 hours. Kidney injury was defined as a 2 times increase in perioperative creatinine plasma concentration or a urine output < 0.5 mL/kg/h in 12 hours, whilst renal failure was defined as a 3 times increase in perioperative creatinine plasma concentration or a urine output < 0.3 mL/kg/h in 24 hours.

 Renal complication defined as serum creatinine > 3.5 mg/dL or 309 umol/L.

¹ Mangano et al (2006) The risk associated with aprotinin in cardiac surgery. NEJM 354:353-365.

² Kuitunen et al (2006) Acute renal failure after cardiac surgery: evaluation of the RIFLE classification. Ann Thorac Surg 81: 542-546.

Key question(s): In patients undergoing surgery, what is the effect of administration of aprotini	morbidity (stroke)?	Evidence table refa: POQ3.18a.P5 (stroke)			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)				
There is one pivotal Level I study (Henry 2007/good quality) which includes data from 14 RCTs (9 in cardiac surgery), one supportive level I study (Brown 2007/fair quality) and two additional RCTs (Later 2009/good quality;	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias		
Nurözler 2008/fair quality) published after the Henry 2007 review.	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias		
	С	One or two Level III studies with a low risk of bias or Level I or II st	tudies with a moderate risk of bias		
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	5		
2. Consistency (If only one study was available, rank this component as 'not applicable')					
Results consistent between Henry and Brown reviews. Additional RCTs consistent. Pivotal evidence – Henry 2007	Α	All studies consistent			
No heterogeneity between studies.	В	Most studies consistent and inconsistency can be explained			
	С	Some inconsistency, reflecting genuine uncertainty around questi	ion		
	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u>	nknow	<u>n</u> factor (not simply study quality or sample size) and thus the clinical imp	act of the intervention could not be		
Pivotal evidence – Henry 2007 Any surgery – RR 0.78 (0.38, 1.62); 1.1% vs 1.7%; 14 RCTs (N=2158)	Α	Very large			
Cardiac surgery – RR 0.76 (0.30, 1.93); 1.3% vs 1.7%; 14 RCTs (N=2138)	В	Substantial			
Supportive evidence – Brown 2007 (see also Supportive Table POQ3.l8a.P5 (stroke)) Cardiac surgery (high dose) – RR 0.67 (0.30, 1.47); 22 RCTs (N=1737)	С	Moderate			
Cardiac surgery (low dose) – RR 0.47 (0.39, 1.47), 22 RC is (N=1737) Cardiac surgery (low dose) – RR 0.47 (0.09, 2.36); 10 RCTs (N=1049)	D	Underpowered			
4. Generalisability (How well does the body of evidence match the population and clinical se	ttings l	peing targeted by the Guideline?)			
The evidence is generalisable to an adult population who are undergoing planned surgery; 9/14 studies included in the Henry review were conducted in cardiac surgery.	Α	Evidence directly generalisable to target population			
line frem y review were conducted in cardiac surgery.	В	Evidence directly generalisable to target population with some car	veats		
	С	Evidence not directly generalisable to the target population but co	ould be sensibly applied		
	D	Evidence not directly generalisable to target population and hard	to judge whether it is sensible to		
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of hea	Ith services/delivery of care and cultural factors?)			
Hospital setting. Studies conducted in a wide range of countries.	Α	Evidence directly applicable to Australian healthcare context			
	В	Evidence applicable to Australian healthcare context with few cav	eats		
	С	Evidence probably applicable to Australian healthcare context wit	h some caveats		
	D	Evidence not applicable to Australian healthcare context			

E-aminocaproic acid is an antifibrinolytic agent. In 2007, Bayer announced a worldwide suspension of aprotinin supply due to the results of the BART trial which suggested increased risk of mortality for aprotinin compared with the lysine analogues tranexamic acid and ε-aminocaproic acid.

Studies underpowered for this outcome. Only one RCT showed a slight difference but this was based on a very small sample size (ie, only one patient in the aprotinin arm had a stroke).

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (good quality), one supportive level I study and two additional RCTs.
2. Consistency	Α	There was no heterogeneity in the pivotal level I study. Results of the supportive level I study and additional RCTs consistent with pivotal evidence.
3. Clinical impact	D	Results show no significant difference but studies likely to be underpowered to detect a difference in stroke.
4. Generalisability	В	The results are generalisable to an adult surgical population; more than half of studies were in cardiac surgery.
5. Applicability	В	Overall there were a large number of studies conducted in a wide range of countries. The additional RCTs were conducted in Turkey and the Netherlands. Likely to be applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intravenous aprotinin therapy on the risk of stroke, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI, confidence interval, RBC, red blood cell: SR, systematic review;

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.l8a.P5 (stroke) Characteristics and results of studies examining the effect of aprotinin on morbidity (stroke)

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a sample size Surgical procedure Quality				Risk estimate (95% CI) or Aprotinin (%) vs control (%)	Significance P-value			
ADULT POPULATION	/IV APROTININ								
Any surgery									
Henry (2007)	Level I Good	14 RCTs N=2158	Adult patients undergoing any surgery	Hospital – planned surgery Various countries ^c	Aprotinin (IV) vs no aprotinin	Stroke	RR 0.78 (0.38, 1.62)	No difference 0.51	None Phet=0.71 (l²=0%)
Cardiac surgery									
Henry (2007)	Level I Good	9 RCTs N=1163	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries ^c	Aprotinin (IV) vs no aprotinin	Stroke	RR 0.76 (0.30, 1.93)	No difference 0.57	None Phet=0.40 (2=4%)
Brown (2007)	Level I Fair	22 RCTs N=1737	Adult patients undergoing cardiac surgery	Hospital – planned surgery Countries not specified	High dose aprotinin (IV) vs placebod	Stroke	RR 0.67 (0.30, 1.47)	No difference 0.32	NR
	Level I Fair	10 RCTs N=1049	Adult patients undergoing cardiac surgery	Hospital – planned surgery Countries not specified	Low dose aprotinin (IV) vs placeboe	Stroke	RR 0.47 (0.09, 2.36)	No difference 0.36	NR
Later (2009)	Level II Good	1 RCT N=199	Adult patients undergoing low- to intermediate-risk cardiac surgery	Hospital – planned surgery The Netherlands	High dose aprotinin (IV) vs placebod	Stroke	1.0% vs 1.0%	No difference 1.0	NA
Nurözler (2008)	Level II Fair	1 RCT N=51	Adult patients undergoing off-pump coronary bypass who have received clopidogrel within 5 days of surgery (all patients)	Hospital – planned surgery Turkey	<u>Low-dose</u> aprotinin (IV) vs placebof	Stroke	4% vs 0%	No difference 0.317	NA

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: ASA, acetylsalicylic acid; CI, confidence interval; DVT, deep vein thrombosis; het, heterogeneity; IV, intravenous; MI, myocardial infarction; NR, not reported; PE, pulmonary embolism; OR, odds ratio; RCT, randomised controlled trial; RR, risk ratio

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I²<25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I²>50%.

Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between aprotinin and no aprotinin.

d High dose (full-dose) aprotinin defined as a 2 million kallikrein-inhibiting units (KIU) IV loading dose, 2 million KIU pump-priming dose, and 0.5 million KIU IV/h maintenance dose.

Low dose (half-dose) aprotinin consisted of a 1 million KIU IV loading dose, 1 million KIU pump-priming dose, and 0.25 million KIU IV/h maintenance dose.

Low dose aprotinin consisted of 1 million KIU infused over 30 min followed by a continuous infusion of 0/5 million KIU/h until the end of surgery.

Key question(s):			Evidence table refa:				
In patients undergoing surgery, what is the effect of administration of <u>aprotinin</u> on <u>morbidity (thrombosis)</u> ? POQ3.l8a.P5 (thrombosis)							
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)						
There is one pivotal Level I study (Henry 2007/good quality) which included up to 11 RCTs (depending on the specific outcome examined), three supportive level I studies (Kagoma 2009/good quality; Liu 2008/poor quality;	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias				
McIlroy 2009/good quality) and one additional RCT (Colwell 2007/good quality) published since the Henry review.	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias				
	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias					
2. Consistency (If only one study was available, rank this component as 'not applicable')							
Some inconsistency between results; may be due to different surgery types and specific thrombosis outcomes. Additional RCTs consistent.	Α	All studies consistent					
Pivotal evidence – Henry 2007	В	Most studies consistent and inconsistency can be explained					
No heterogeneity between studies.	С	Some inconsistency, reflecting genuine uncertainty around question	on				
	D	Evidence is inconsistent					
	NA	Not applicable (one study only)					
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u> determined)	nknow	<u>n</u> factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be				
Pivotal evidence – Henry 2007	Α	Very large					
Any surgery (DVT) – RR 0.79 (0.46, 1.34); 5.3% vs 5.4%; 11 RCTs (N=986) Cardiac surgery (DVT) – RR 2.52 (0.41, 15.45); 2.4% vs 1.0%; 2 RCTs (N=272)	В	Substantial					
Any surgery (PE) – RR 1.98 (0.38, 10.46); 3.1% vs 1.9%; 2 RCTs (N=175) Other outcomes and supportive evidence-see Supportive Table POQ3.18a.P5 (thrombosis)	С	Moderate					
Other outcomes and supportive evidence-see supportive rable POQs.iba.P3 (Infombosis)	D	Underpowered/inconsistent					
4. Generalisability (How well does the body of evidence match the population and clinical se	ttings b	peing targeted by the Guideline?)					
The evidence is generalisable to an adult population who are undergoing planned surgery; 2/11 studies included in the Henry review were conducted in cardiac surgery for DVT and 3/7 studies conducted in cardiac surgery for other	Α	Evidence directly generalisable to target population					
thrombosis.	В	Evidence directly generalisable to target population with some cav	eats				
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied				
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of heal	th services/delivery of care and cultural factors?)					
Hospital setting. Studies conducted in a wide range of countries.	Α	Evidence directly applicable to Australian healthcare context					
	В	Evidence applicable to Australian healthcare context with few cave	eats				
	С	Evidence probably applicable to Australian healthcare context with	some caveats				
	D	Evidence not applicable to Australian healthcare context					

In 2007, Bayer announced a worldwide suspension of aprotinin supply due to the results of the BART trial which suggested increased risk of mortality for aprotinin compared with the lysine analogues tranexamic acid and ε-aminocaproic acid.

CRG noted that variations in definition and how measured may make a difference. Included studies underpowered to detect these outcomes.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (good quality), three supportive level I studies and one additional RCT.
2. Consistency	С	There was no heterogeneity in the pivotal level I study for each thrombosis outcome (DVT, PE, other thrombosis). Results of the supportive level I studies and additional RCT showed some inconsistency, possibly due to different surgeries and definitions of thrombosis outcomes.
3. Clinical impact	D	There was no significant difference in any results but some of the risk estimates were large. Likely to be underpowered for thrombosis outcomes.
4. Generalisability	В	The results are generalisable to an adult surgical population.
5. Applicability	В	Overall there were a reasonable number of studies conducted in a range of countries. The additional RCT was conducted in the US/Canada. Likely to be applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intravenous aprotinin therapy on the risk of venous thromboembolism, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI. confidence interval. RBC, red blood cell: SR, systematic review:

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.l8a.P5 (thrombosis) Characteristics and results of studies examining the effect of <u>aprotinin</u> on <u>morbidity (thrombosis)</u>

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size	Surgical procedure				Risk estimate (95% CI)	Significance P-value	
ADULT POPULATION	I/IV APROTININ								
Any surgery									
Henry (2007)	Level I Good	11 RCTs N=986	Adult patients undergoing any surgery	Hospital – planned surgery Various countries ^c	Aprotinin (IV) vs no aprotinin	Deep vein thrombosis	RR 0.79 (0.46, 1.34)	No difference 0.38	None Phet=0.80 (I ² =0%)
Henry (2007)	Level I Good	2 RCTs N=175	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Pulmonary embolism	RR 1.98 (0.38, 10.46)	No difference 0.42	None Phet=0.95 (I ² =0%)
Henry (2007)	Level I Good	7 RCTs N=583	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Other thrombosis (not MI, stroke, DVT or PE)	RR 0.73 (0.25, 2.15)	No difference 0.57	None Phet=0.64 (I ² =0%)
Cardiac surgery									
Henry (2007)	Level I Good	2 RCTs N=272	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries ^c	Aprotinin (IV) vs no aprotinin	Deep vein thrombosis	RR 2.52 (0.41, 15.45)	No difference 0.32	None Phet=0.71 (I ² =0%)
Henry (2007)	Level I Good	3 RCTs N=370	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Other thrombosis (not MI, stroke, DVT or PE)	RR 0.62 (0.11, 3.36)	No difference 0.58	None Phet=0.50 (I ² =0%)
McIlroy (2009)	Level I Good	3 RCTs N=174	Adult patients <u>receiving</u> <u>aspirin</u> undergoing cardiac surgery	Hospital – planned surgery Countries not specified	Aprotinin (IV) vs placebo	Thrombotic complication (includes DVT, stroke, MI or PE)	OR 0.51 (0.21, 1.20)	No difference 0.12	None Phet=0.76 (I2=0%)
Orthopaedic surg	jery								
Kagoma (2009)	Level I Good	3 trials N=97	Adult patients undergoing orthopaedic surgery	Hospital – planned surgery Countries not specified	Aprotinin (IV) vs placebo	Venous thromboembolism (including DVT and PE)	RD -0.04 (-0.09, 0.02)	No difference NR	NR
Colwell (2007)	Level II Good	1 RCT N=352	Adults undergoing <u>unilateral</u> total hip arthroplasty	Hospital – planned surgery US/Canada	Aprotinin (IV) vs placebo	Deep vein thrombosis	1.1% vs 1.7%	No difference NS	NA
				US/Cariaua		Pulmonary embolism	1.1% vs 1.1%	No difference NS	NA

Study	Level of	No. of trials /	Patient population /		Outcome	ome Results			
	evidence ^a sample size Surgical procedure <i>Quality</i>			Risk estimate (95% CI)	Significance P-value				
Liver surgery									•
Liu (2008)	Level I Poor	2 RCTs N=200	Adult patients undergoing orthotopic liver transplantation	Hospital – planned surgery Countries not specified	Aprotinin (IV) vs no aprotinin	Thromboembolic events (not defined)	OR 0.38 (0.09, 1.64)	No difference >0.05	None Phet=0.88

Abbreviations: CI, confidence interval; DVT, deep vein thrombosis; het, heterogeneity; IV, intravenous; MI, myocardial infarction; NA, not applicable; NR, not reported; PE, pulmonary embolism; OR, odds ratio; RCT, randomised controlled trial; RD, risk difference; RR, risk ratio.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and 12<25%; (ii) mild heterogeneity if 12 <25%; moderate heterogeneity if 12 between 25-50%; substantial heterogeneity 12 >50%.

c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between aprotinin and no aprotinin.

Key question(s):			Evidence table refa:
In patients undergoing surgery, what is the effect of administration of aprotining	<u>on (</u>	quality of life?	POQ3.I8a.P6
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies,)		
No studies of any level were identified which assessed the effect of tranexamic acid on quality of life.	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias
	В	One or two Level II studies with a low risk of bias or SR/several Lev	rel III studies with a low risk of bias
	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	
2. Consistency (If only one study was available, rank this component as 'not applicable')			
NA	Α	All studies consistent	
	В	Most studies consistent and inconsistency can be explained	
	С	Some inconsistency, reflecting genuine uncertainty around question	on
	D	Evidence is inconsistent	
	NA	Not applicable (one study only)	
3. Clinical impact (Indicate in the space below if the study results varied according to some undetermined)	knowi	n factor (not simply study quality or sample size) and thus the clinical impa	ct of the intervention could not be
NA NA	Α	Very large	
	В	Substantial	
	С	Moderate	
	D	Slight/Restricted	
4. Generalisability (How well does the body of evidence match the population and clinical sett	tings b		
NA	Α	Evidence directly generalisable to target population	
	В	Evidence directly generalisable to target population with some cav	eats
	С	Evidence not directly generalisable to the target population but cou	ıld be sensibly applied
	D	Evidence not directly generalisable to target population and hard to	judge whether it is sensible to
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of heal	·	
NA	Α	Evidence directly applicable to Australian healthcare context	
	В	Evidence applicable to Australian healthcare context with few cave	
	С	Evidence probably applicable to Australian healthcare context with	some caveats
	D	Evidence not applicable to Australian healthcare context	

In 2007, Bayer announced a worldwide suspension of aprotinin supply due to the results of the BART trial which suggested increased risk of mortality for aprotinin compared with the lysine analogues tranexamic acid and ε-aminocaproic acid.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	NA	
2. Consistency	NA	
3. Clinical impact	NA	
4. Generalisability	NA	
5. Applicability	NA	

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intravenous aprotinin therapy on quality of life, compared with no therapy, is unknown.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI. confidence interval. RBC, red blood cell: SR, systematic review:

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^a Interventions: 11 = ANH, 12 = intraoperative cell salvage, 13 = perioperative ANH and intraoperative cell salvage, 14 = postoperative cell salvage, 15 = deliberate induced hypotension, 16 = prevention of hypothermia, 17 = point-of-care testing for coagulation status and haemoglobin, 18 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, 19 = appropriate patient positioning, 110 = PAD

Key question(s):			Evidence table refa:			
In patients undergoing surgery, what is the effect of administration of aprotini	<u>n</u> on	re-operation for bleeding?	POQ3.I8a.S2			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)					
There is one pivotal Level I study (Henry 2007/good quality) which included data from 36 RCTs (33 RCTs for cardiac surgery), three supportive level I studies (Henry 2009/good quality; Brown 2007/fair quality; McIlroy	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias				
2009/good quality) and four additional RCTs (Later 2009/good quality; Nurözler 2008/fair quality; Apostolakis 2008/fair quality; Leijdekkers 2006/fair quality) published following the Henry review.	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Results consistent between pivotal and supportive studies. Additional RCTs consistent. Pivotal evidence – Henry 2007	Α	All studies consistent				
No heterogeneity between studies.	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around question	on			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u>	nknow	n factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be			
Pivotal evidence – Henry 2007 (re-operation for bleeding)	Α	Very large				
Any surgery – RR 0.48 (0.35, 0.68); 1.9% vs 4.7%; 36 RCTs (N=4715) Cardiac surgery – RR 0.49 (0.34, 0.70); 1.9% vs 4.5%; 33 RCTs (N=4534)	В	Substantial (cardiac)				
Supportive evidence – Brown 2007 (return to operating room) – see also Supportive Table POQ3.l8a.S2 Cardiac surgery (high dose) – RR 0.47 (0.32, 0.69); 49 RCTs (N=3912)	С	Moderate				
Cardiac surgery (low dose) – RR 0.69 (0.41, 1.18); 20 RCTs (N=1623)	D	Underpowered (non-cardiac)				
4. Generalisability (How well does the body of evidence match the population and clinical se	ttings Ł	peing targeted by the Guideline?)				
The evidence is generalisable to an adult population who are undergoing planned surgery; 33/36 studies included in Henry 2007 conducted in cardiac surgery.	Α	Evidence directly generalisable to target population				
Treffly 2007 Conducted in Cardiac Surgery.	В	Evidence directly generalisable to target population with some cav	eats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of heal	th services/delivery of care and cultural factors?)				
Hospital setting. Studies conducted in a wide range of countries.	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few cave	eats			
	С	Evidence probably applicable to Australian healthcare context with	some caveats			
	D	Evidence not applicable to Australian healthcare context				

In 2007, Bayer announced a worldwide suspension of aprotinin supply due to the results of the BART trial which suggested increased risk of mortality for aprotinin compared with the lysine analogues tranexamic acid and ε-aminocaproic acid.

Later 2009 study presented results for re-operation due to surgical bleeding and re-operation due to non-surgical bleeding. When these are combined to include re-operation due to any bleeding, there is no difference between arms. Differences due to does in Brown 2007 study. Low dose category may be underpowered, thus more uncertainty.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Component Rating		Description
	Cardiac	Non-cardiac	
Evidence base	Α	А	There is one pivotal Level I (good quality) study, three supportive level I studies and four additional RCTs.
2. Consistency	В	1 1)	There was no heterogeneity in the pivotal level I study. There was some inconsistency for reoperation (not defined) due to dose in one of the supportive level I studies. The results for cardiac surgery were largely consistent.
3. Clinical impact	В		For cardiac surgery, there were significantly less re-operations due to bleeding in patients receiving aprotinin therapy compared with no therapy. For non-cardiac surgery, the results were likely underpowered.
4. Generalisability	В	В	The results are generalisable to an adult surgical population; most studies in cardiac surgery.
5. Applicability	В	В	Overall there were a large number of studies conducted in a wide range of countries. Additional RCTs conducted in the Netherlands, Turkey and Greece. Likely to be applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, intravenous aprotinin therapy reduces the risk of reoperation for bleeding compared with no therapy.

In adult patients undergoing noncardiac surgery, the effect of intravenous aprotinin therapy on reoperation for bleeding, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI, confidence interval, RBC, red blood cell: SR, systematic review;

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.l8a.S2 Characteristics and results of studies examining the effect of <u>aprotinin</u> on <u>reoperation for bleeding</u>

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size	Surgical procedure	Location			Risk estimate (95% CI) or Aprotinin (%) vs control (%)	Significance P-value	
ADULT POPULATION	n/ IV aprotinin								
Any surgery									
Henry (2007)	Level I Good	36 trials N=4715	Adult patients undergoing any surgery	Hospital – planned surgery Various countries ^c	Aprotinin (IV) vs placebo	Reoperation for bleeding	RR 0.48 (0.35, 0.68)	Favours aprotinin <0.001	None Phet=0.51 (l²=0%)
Cardiac surgery									
Henry (2007)	Level I Good	33 RCTs N=4534	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries ^c	Aprotinin (IV) vs placebo	Reoperation for bleeding	RR 0.49 (0.34, 0.70)	Favours aprotinin <0.001	None Phet=0.41 (l²=4%)
Henry (2009)	Level I Fair	NR ^d	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs placebo	Reoperation for bleeding	RR 0.48 (0.34, 0.67)	Favours aprotinin NR	NR
Brown (2007)	Level I Fair	40 RCTs N=3912	Adult patients undergoing cardiac surgery	Hospital – planned surgery Countries not specified	High dose aprotinin (IV) vs placeboe	Return to operating room (reason/s not defined)	RR 0.47 (0.32, 0.69)	Favours aprotinin <0.001	NR
		20 RCTs N=1623	Adult patients undergoing cardiac surgery	Hospital – planned surgery Countries not specified	Low dose aprotinin (IV) vs placebof	Return to operating room (reason/s not defined)	RR 0.69 (0.41, 1.18)	No difference 0.176	NR
McIlroy (2009)	Level I Good	4 RCTs N=198	Adult patients <u>receiving</u> <u>ASA</u> undergoing cardiac surgery	Hospital – planned surgery Countries not specified	Aprotinin (IV) vs placebo	Reoperation for bleeding	OR 0.42 (0.13, 1.36)	No difference 0.15	None Phet=0.61 (I ² =0%)
Later (2009)	Level II Good	1 RCT N=199	Adult patients undergoing low- to intermediate-risk cardiac surgery	Hospital – planned surgery The Netherlands	High dose aprotinin (IV) vs placeboe	Reoperation for <u>any</u> reason	5.2% vs 13.6%	No difference 0.054	NA
				The incinentalius		Reoperation <u>due to</u> <u>surgical bleeding</u>	4.2% vs 2.9%	No difference 0.71	NA
						Reoperation <u>due to non-</u> <u>surgical bleeding</u>	0% vs 3.9%	No difference 0.12	NA

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size	Surgical procedure	Location			Risk estimate (95% CI) or Aprotinin (%) vs control (%)	Significance P-value	
Nurözler (2008)	Level II Fair	1 RCT N=51	Adult patients undergoing off-pump coronary bypass who have received clopidogrel within 5 days of surgery (all patients)	Hospital – planned surgery Turkey	Low-dose aprotinin (IV) vs placebo ^g	Reoperation for bleeding	0% vs 7.7%	No difference 0.157	NA
Other surgery	_	•							•
Apostolakis (2008)	Level II Fair	1 RCT N=59	Adult population undergoing thoracic surgery	Hospital – planned surgery Greece	Ultra-low-dose aprotinin vs placebo ^h	Reoperation for bleeding	0% vs 0%	No difference NA	NA
Leijdekkers (2006)	Level II Fair	1 RCT N=35	Adult patients undergoing surgery for infra-renal abdominal aneurysm	Hospital – planned surgery The Netherlands	Aprotinin (IV) vs placebo	Reoperation for bleeding	6.3% vs 10.5%	No difference 0.65	NA

Abbreviations: ASA, acetylsalicylic acid; CI, confidence interval; het, heterogeneity; IV, intravenous; NR, not reported; OR, odds ratio; RCT, randomised controlled trial; RR, risk ratio.

a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

bHeterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I²<25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between aprotinin and no aprotinin.

^d Not specifically reported in the text of the publication.

e High dose (full-dose) aprotinin defined as a 2 million kallikrein-inhibiting units (KIU) IV loading dose, 2 million KIU pump-priming dose, and 0.5 million KIU IV/h maintenance dose.

Low dose (half-dose) aprotinin consisted of a 1 million KIU IV loading dose, 1 million KIU pump-priming dose, and 0.25 million KIU IV/h maintenance dose.

⁹ Low dose aprotinin consisted of 1 million KIU infused over 30 min followed by a continuous infusion of 0/5 million KIU/h until the end of surgery.

h Ultra-low dose aprotinin defined as a test dose of 1mL at intubation, followed by 0.5 million IU over 15 min following intubation, and again following closure.

Key question(s):			Evidence table refa:			
In patients undergoing surgery, what is the effect of administration of aprotinii	<u>n</u> on	hospital length of stay?	POQ3.I8a.S5			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	5)					
There is one pivotal Level I study (Henry 2007/good quality) which includes data from 21 RCTs (13 for cardiac surgery) and two additional RCTs (Later 2009/good quality; Nurözler 2008/fair quality) published after the Henry	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias				
review.	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias				
	С	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Pivotal evidence – Henry 2007	Α	All studies consistent				
Mild-moderate heterogeneity between studies.	В	Most studies consistent and inconsistency can be explained				
Additional RCTs consistent.	С	Some inconsistency, reflecting genuine uncertainty around question	on			
		Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>ur</u>	nknow	n factor (not simply study quality or sample size) and thus the clinical impa	ct of the intervention could not be			
Pivotal evidence – Henry 2007	Α	Very large				
Any surgery – WMD -0.01 (-0.50, 0.48); 21 RCTs (N=1570) Cardiac surgery – WMD -0.10 (-0.64, 0.44); 13 RCTs (N=1412)	В	Substantial				
Supportive evidence – see Supportive Table POQ3.l8a.S5	С	Moderate				
	D	No difference				
4. Generalisability (How well does the body of evidence match the population and clinical set	tings Ł	peing targeted by the Guideline?)				
The evidence is generalisable to an adult population who are undergoing planned surgery; 13/21 studies included in the Henry review conducted in cardiac surgery.	Α	Evidence directly generalisable to target population				
and many review conducted in cardiac surgery.	В	Evidence directly generalisable to target population with some cav	eats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of heal	th services/delivery of care and cultural factors?)				
Hospital setting. Studies conducted in a wide range of countries.	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few caveats				
	С	Evidence probably applicable to Australian healthcare context with	some caveats			
	D	Evidence not applicable to Australian healthcare context				

In 2007, Bayer announced a worldwide suspension of aprotinin supply due to the results of the BART trial which suggested increased risk of mortality for aprotinin compared with the lysine analogues tranexamic acid and ε-aminocaproic acid.

Differences of 1% of a day not considered by the CRG to be clinically important.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I (good quality) study and two additional RCTs.
2. Consistency	В	There was some heterogeneity in the pivotal level I (good quality) study. The results of the additional RCTs were consistent.
3. Clinical impact	D	There was no significant difference in hospital length of stay for intravenous aprotinin therapy compared with no therapy.
4. Generalisability	В	The results are generalisable to an adult surgical population.
5. Applicability	В	Studies were conducted in a wide range of countries. The additional RCTs were conducted in the Netherlands and Turkey. Likely to be applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, intravenous aprotinin therapy has no effect on hospital length of stay compared with no therapy.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI, confidence interval, RBC, red blood cell; SR, systematic review;

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

a Interventions: 11 = ANH, 12 = intraoperative cell salvage, 13 = perioperative ANH and intraoperative cell salvage, 14 = postoperative cell salvage, 15 = deliberate induced hypotension, 16 = prevention of hypothermia, 17 = point-of-care testing for coagulation status and haemoglobin, 18 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, 19 = appropriate patient positioning, 110 = PAD

POQ3.l8a.S5 Characteristics and results of studies examining the effect of aprotinin on hospital length of stay

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size	Surgical procedure	Location			Risk estimate (95% CI) Or Aprotinin (Mean ± SD) vs control (mean ± SD)	Significance P-value	
ADULT POPULATION	/IV APROTININ								
Any surgery									
Henry (2007)	Level I Good	21 trials N=1570	Adult patients undergoing any surgery	Hospital – planned surgery Various countries ^c	Aprotinin (IV) vs no aprotinin	Hospital length of stay (days)	WMD -0.01 (-0.50, 0.48)	No difference 0.96	Mild Phet=0.19 (l ² =23%)
Cardiac surgery									·
Henry (2007)	Level I Good	13 RCTs N=1412	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Aprotinin (IV) vs no aprotinin	Hospital length of stay (days)	WMD -0.10 (-0.64, 0.44)	No difference 0.73	Moderate Phet=0.12 (l ² =33%)
Later (2009)	Level II Good	1 RCT N=199	Adult patients undergoing low- to intermediate-risk cardiac surgery	Hospital – planned surgery The Netherlands	High-dose aprotinin (IV) vs placebod	Hospital length of stay (days)	7.8 ± 6.7 vs 8.5 ± 7.4	No difference 0.49	NA
Nurözler (2008)	Level II Fair	1 RCT N=51	Adult patients undergoing off-pump coronary bypass who have received clopidogrel within 5 days of surgery	Hospital – planned surgery Turkey	Low-dose aprotinin (IV) vs placebo ^o	Hospital length of stay (days)	5.3 ± 1.6 vs 5.5 ± 1.4	No difference 0.660	NA

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: ASA, acetylsalicylic acid; CI, confidence interval; het, heterogeneity; IV, intravenous; NR, not reported; OR, odds ratio; RCT, randomised controlled trial; RR, risk ratio.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between aprotinin and no aprotinin.

d High dose (full-dose) aprotinin defined as a 2 million kallikrein-inhibiting units (KIU) IV loading dose, 2 million KIU pump-priming dose, and 0.5 million KIU IV/h maintenance dose.

ELOW dose aprotinin consisted of 1 million KIU infused over 30 min followed by a continuous infusion of 0/5 million KIU/h until the end of surgery.

Recommendation(s) for administration of aprotinin

RECOMMENDATION	GRADE	RELE	VANT	
What recommendation(s) does the guideline development group draw from this evidence? Use action statements where possible.		EVIDENCE TAB		
No recommendation made because the drug has been withdrawn due to concerns that it is less safe than alternative therapies.				
IMPLEMENTATION OF RECOMMENDATION	•	1		
Please indicate yes or no to the following questions. Where the answer is yes please provide explanatory information about this. This information will be used to develop the implementation plan for the guidelines.				
Will this recommendation result in changes in usual care?		YES	NO	
Are there any resource implications associated with implementing this recommendation?		YES	NO	
Will the implementation of this recommendation require changes in the way care is currently organised?		YES	NO	
Are the guideline development group aware of any barriers to the implementation of this recommendation		YES	NO	
What could help to facilitate implementation of the recommendation?		YES	NO	
		•		

Intervention 8 - Administration of antifibrinolytics & DDAVP: Tranexamic acid

Key question(s): In patients undergoing surgery, what is the effect of administration of <u>tranexar</u>	nic a	acid on transfusion incidence?	Evidence table refa: POQ3.I8b.P1				
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	.)						
There is one pivotal Level I (Henry 2007/good quality) which includes data from up to 51 RCTs, two supportive level	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias				
l studies (Brown 2007/fair quality; Kagoma 2009/good quality), two supportive level I/II studies which include one RCT each (Kongnyuy 2009/good-good quality; McIlroy 2009/good-poor quality) and nine additional RCTs (Jimenez	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias				
2007/good quality; Later 2009/good quality; Mehr-Aein 2007/good quality; Taghaddomi 2009/fair quality; Alvarez	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias				
2008/fair quality; Elwatidy 2008/fair quality; Sadeghi 2007/good quality; Wong 2008/good quality; Choi 2009/fair quality).	D	Level IV studies or Level I to III studies/SRs with a high risk of bias					
2. Consistency (If only one study was available, rank this component as 'not applicable')							
Results mostly consistent between pivotal and supportive meta-analyses and RCTs. Consistency of individual	Α	All studies consistent					
studies within meta-analyses described below. Pivotal evidence – Henry 2007	В	Most studies consistent and inconsistency can be explained					
Moderate to substantial heterogeneity (see note) in main analyses and most subgroup analyses. Differences may be due to different surgery types, and transfusion of different blood products.	С	Some inconsistency, reflecting genuine uncertainty around question	ninty around question				
be due to different surgery types, and transmission of different blood products.	D	Evidence is inconsistent					
	NA	Not applicable (one study only)					
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>ur</u>	know	n factor (not simply study quality or sample size) and thus the clinical impa	ct of the intervention could not be				
Pivotal evidence – Henry 2007	Α	Very large					
Any surgery – 27.0% vs 43.8%; RR 0.61 (0.54, 0.70); 51 RCTs (N=3751) Cardiac surgery – 27.8% vs 40.8%; RR 0.69 (0.60, 0.79); 28 RCTs (N=2443)	В	Substantial					
Orthopaedic surgery – 26.7% vs 52.2%; RR 0.44 (0.33, 0.60); 20 RCTs (N=953) Liver surgery – 19.6% vs 36.5%; RR 0.16 (0.00, 32.47); 2 RCTs (N=296)	С	Moderate					
Supportive evidence – see Summary Table POQ3.l8b.P1	D	Slight/Restricted					
4. Generalisability (How well does the body of evidence match the population and clinical set	tings Ł	peing targeted by the Guideline?)					
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were included in the overall analysis. There were also analyses of patients undergoing cardiac surgery who had received	Α	Evidence directly generalisable to target population					
ASA within 7 days prior to surgery.	В	Evidence directly generalisable to target population with some cav	eats				
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied				
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of heal	th services/delivery of care and cultural factors?)					
Hospital setting.	Α	Evidence directly applicable to Australian healthcare context					
The pivotal review included studies from a wide range of countries. Included RCTs were from a number of different countries including several from the Middle East and Asia.	В	Evidence applicable to Australian healthcare context with few cave	ats				
Š	С	Evidence probably applicable to Australian healthcare context with	some caveats				
	D	Evidence not applicable to Australian healthcare context					

The Advisory Committee on Prescription Medicines (ACPM) has recently recommended approval of tranexamic acid injection "for the reduction of peri- and postoperative blood loss and of the need for blood transfusion in adult patients undergoing cardiac surgery or total knee or hip arthroplasty; and paediatric patients undergoing cardiac surgery". Tranexamic acid tablets are approved in Australia for a number of indications including haemostatic, hereditary angioedema, short-term treatment of traumatic hyphaema, patients with coagulopathies undergoing minor surgery, and menorrhagia.

The Henry (pivotal) review assessed quality and performed a subgroup analysis of transfusion incidence for all surgery types based on the rating (A,B or C) of treatment allocation^b. The analysis showed no substantial difference in the results between studies rated A. B or C.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	А	There is one pivotal Level I (good quality), two supportive level I studies, two supportive level I/II studies and nine additional RCTs.
2. Consistency		Significant heterogeneity in the pivotal level I study but mostly due to difference in magnitude of effect and not direction of effect. Differences may be due to different surgery types. Results of supportive level I studies and additional RCTs consistent with pivotal evidence.
3. Clinical impact	В	There were significant differences between intravenous tranexamic acid therapy and no therapy overall and for surgery subgroups.
4. Generalisability	В	The results are generalisable to an adult surgical population undergoing cardiac, major joint and spinal surgery.
5. Applicability	В	Overall there were a large number of studies conducted in a wide range of countries. Likely to be applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery and major orthopaedic surgery, intravenous tranexamic acid therapy reduces the incidence of allogeneic blood transfusion compared with no therapy.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50% Abbreviations: ASA, acetylsalicylic acid; RCT, randomised controlled trial; RR, risk ratio; SR, systematic review.

a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission b Cochrane ratings defined as follows: Grade A. adequate allocation concealment: Grade B. uncertain allocation concealment: Grade C. inadequate allocation concealment.

POQ3.l8b.P1 Characteristics and results of studies examining the effect of <u>tranexamic acid</u> on <u>transfusion incidence</u>.

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size proce included in analysis	procedure	Location			Risk estimate (95% CI) or TXA (%) vs control (%)	Significance P-value	
ADULT POPULATION	ON/IV TRANEXAMIC	ACID							
Any surgery									
Henry (2007)	Level I Good	51 RCTs N=3751	Adult patients undergoing any surgery	Hospital – planned surgery Various countries ^c	Tranexamic acid (IV) vs no tranexamic acid	Transfusion incidence (allogeneic blood)	RR 0.61 (0.54, 0.70)	Favours tranexamic acid <0.001	Substantial Phet<0.001 (l ² =50%)
Henry (2007)	Level I Good	45 RCTs N=3191	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid with transfusion protocol	Transfusion incidence (allogeneic blood)	RR 0.57 (0.49, 0.66)	Favours tranexamic acid <0.001	Moderate Phet=0.001 (I ² =44%)
		6 RCTs N=560	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid without transfusion protocol	Transfusion incidence (allogeneic blood)	RR 0.82 (0.63, 1.07)	No difference 0.15	Substantial Phet=0.02 (l ² =63%)
Henry (2007)	Level I Good Rating A ^d	21 RCTs N=1610	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Transfusion incidence (allogeneic blood)	RR 0.60 (0.49, 0.72)	Favours tranexamic acid <0.001	Moderate Phet=0.02 (1 ² =42%)
	Level I Good Rating Bd	20 RCTs N=1254	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Transfusion incidence (allogeneic blood)	RR 0.55 (0.42, 0.73)	Favours tranexamic acid <0.001	Substantial Phet<0.001 (l²=62%)
	Level I Good Rating Cd	10 RCTs N=927	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Transfusion incidence (allogeneic blood)	RR 0.69 (0.56, 0.86)	Favours tranexamic acid 0.0012	Moderate Phet=0.09 (l²=40%)

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or TXA (%) vs control (%)	Significance P-value	
Cardiac surgery									
Henry (2007)	Level I Good	28 RCTs N=2443	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Transfusion incidence (allogeneic blood)	RR 0.69 (0.60, 0.79)	Favours tranexamic acid <0.001	Moderate Phet=0.03 (l²=37%)
		16 RCTs N=926	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Tranexamic acid total dose < 2.0 q (IV) vs no tranexamic acid	Transfusion incidence (allogeneic blood)	RR 0.72 (0.59, 0.88)	Favours tranexamic acid 0.0013	Moderate Phet=0.05 (I ² =40%)
		13 RCTs N=1571	N=1571 cardiac surgery	Hospital – planned surgery Various countries	Tranexamic acid total dose 2.0 – 10.0 g (IV) vs no tranexamic acid	Transfusion incidence (allogeneic blood)	RR 0.67 (0.55, 0.83)	Favours tranexamic acid <0.001	Moderate Phet=0.09 (I ² =37%)
Henry (2009)	Level I Good	N RCTs NR N=NR	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Transfusion incidence (allogeneic blood)	RR 0.70 (0.61, 0.80)	Favours tranexamic acid NR	NR
Brown (2007)	Level I Fair	22 RCTs N=2429	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs placebo	Transfusion incidence (pRBCs)	RR 0.75 (0.60, 0.92)	Favours tranexamic acid 0.007	NR
McIlroy (2009)	Level I/II Good/Poor	1 RCT N=79	Adult patients receiving aspirin undergoing CABG surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs placebo	Transfusion incidence (allogeneic blood products)	RR 0.97 (0.32, 2.90)	No difference 0.95	NA
Jimenez (2007)	Level II Good		N=50 <u>cardiopulmonary bypass</u> su	Hospital - planned surgery Spain		Transfusion incidence (RBC and plasma/0-4 hr)	4.2% vs 7.6%	No difference 0.39	NA
				Gpa		Transfusion incidence (RBC and plasma/chest tube withdrawal)	37.5% vs 73.1%	Favours tranexamic acid 0.01	NA
						Transfusion incidence (plasma/chest tube withdrawal)	4.2% vs 30.8%	Favours tranexamic acid 0.02	NA
	Level II Good	1 RCT N=202	Adults undergoing <u>first-time</u> , <u>non-complex cardiac surgery</u> <u>with CPB</u>	Hospital - planned surgery The Netherlands	Tranexamic acid (IV) vs placebo	Transfusion incidence (pRBC)	57.6% vs 70.9%	No difference 0.057	NA
				realistanta		Transfusion incidence (blood products)	69.7% vs 78.6%	No difference 0.15	NA
Mehr-Aein (2007)	Level II Good	1 RCT N=66	Adults undergoing off-pump CABG surgery	Hospital - planned surgery	Tranexamic acid (IV) vs placebo	Transfusion incidence (whole blood or pRBC)	15.2% vs 24.2%	No difference 0.07	NA

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or TXA (%) vs control (%)	Significance P-value	
				Iran		Transfusion incidence (FFP)	0% vs 18.2%	No difference 0.05	NA
						Transfusion incidence (platelets)	0% vs 0%	No difference NA	NA
				Transfusion incidence (any blood products)	15.2% vs 36.4%	No difference 0.09	NA		
	Adults undergoing off-pump CABG surgery	Hospital - planned surgery	Tranexamic acid (IV) vs placebo	Transfusion incidence (pRBC/intraoperative)	0% vs 6%	No difference 0.24	NA		
		Iran		Transfusion incidence (pRBC/0-4 hr)	0% vs 30%	Favours tranexamic acid <0.001	NA		
					Transfusion incidence (pRBC/4-24 hr)	16.0% vs 18.0%	No difference 1.00	NA	
					Transfusion incidence (FFP/0-4 hr)	4.0% vs 4.0%	No difference 1.00	NA	
						Transfusion incidence (FFP/4-24 hr)	0% vs 0%	No difference NA	NA
						Transfusion incidence (RBC or FFP/up to 24 hr)	16.0% vs 54.0%	Favours tranexamic acid <0.001	NA
Orthopaedic surge	ery								
Henry (2007)	Level I Good	20 RCTs N=953	Adult patients undergoing orthopaedic surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Transfusion incidence (allogeneic blood)	RR 0.44 (0.33, 0.60)	Favours tranexamic acid <0.001	Substantial Phet<0.001 (l²=65%)
Kagoma (2009)	Level I Good	18 RCTs N=943	Adult patients undergoing hip and knee replacement surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs placebo	Transfusion incidence	RR 0.47 (0.40, 0.55)	Favours tranexamic acid NR	NR
Alvarez (2008)	Level II Fair	1 RCT N=95	Adult patients undergoing total knee arthroplasty	Hospital – planned surgery Spain	Tranexamic acid (IV) vs placebo	Transfusion incidence (allogeneic and autologous blood)	2.2% vs 12.2%	No difference 0.11	NA
				•		Transfusion incidence (recovered blood)	4.3% vs 73.5%	Favours tranexamic acid <0.001	NA

Study Level of			Patient population / Surgical procedure	Setting Location	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size included in analysis					Risk estimate (95% CI) or TXA (%) vs control (%)	Significance P-value	
Elwatidy (2008)	Level II Fair	1 RCT N=64	Adults <u>or children</u> undergoing spine surgery	Hospital – planned surgery Saudi Arabia	Tranexamic acid (IV) vs placebo	Transfusion incidence	12.5% vs 37.5%	Favours tranexamic acid 0.021	NA
Sadeghi (2007) Level II 1 RCT Good N=67	N=67 <u>surgery</u> s	Hospital – planned surgery Iran	Tranexamic acid (IV) vs placebo	Transfusion incidence (whole blood or pRBC)	37.5% vs 57.1%	Favours tranexamic acid 0.04	NA		
					Transfusion incidence (FFP)	3.1% vs 0%	No difference >0.05	NA	
				Transfusion incidence (platelets)	0% vs 0%	No difference NA	NA		
						Transfusion incidence (any blood products)	37.5% vs 57.1%	Favours tranexamic acid 0.04	NA
Wong (2008)	Level II Good	1 RCT N=147	Adults undergoing spinal fusion surgery	Hospital – planned surgery	urgery (IV) vs placebo	Transfusion incidence (pRBC/perioperative)	31% vs 40%	No difference 0.25	NA
				Canada		Transfusion incidence (AWB/perioperative)	32% vs 36%	No difference 0.65	NA
						Transfusion incidence (cell saver/perioperative)	45% vs 63%	Favours tranexamic acid 0.026	NA
						Transfusion incidence (FFP/perioperative)	7% vs 12%	No difference 0.27	NA
				Transfusion incidence (platelets/perioperative)	3% vs 3%	No difference 0.99	NA		

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size included in analysis	included in	Location			Risk estimate (95% CI) or TXA (%) vs control (%)	Significance P-value	
Wong (2008)	Level II Good	1 RCT N=147	Adults undergoing spinal fusion surgery	Hospital – planned surgery Canada	Tranexamic acid (IV) vs placebo	Transfusion incidence (pRBC/intraoperative)	19% vs 23%	No difference 0.57	NA
				Canaua		Transfusion incidence (AWB/intraoperative)	25% vs 28%	No difference 0.61	NA
						Transfusion incidence (<u>cell</u> saver/intraoperative)	45% vs 62%	Favours tranexamic acid 0.039	NA
					Transfusion incidence (FFP/intraoperative)	5% vs 9%	No difference 0.36	NA	
				Transfusion incidence (platelets/intraoperative)	3% vs 3%	No difference 0.99	NA		
Wong (2008) Level II Good		1 RCT N=147	Adults undergoing spinal fusion surgery	Hospital – planned surgery Canada	Tranexamic acid (IV) vs placebo	Transfusion incidence (pRBC/postoperative)	15% vs 28%	No difference 0.051	NA
			Sanaaa		Transfusion incidence (AWB/postoperative)	13% vs 13%	No difference 0.97	NA	
						Transfusion incidence (cell saver/postoperative)	3% vs 4%	No difference 0.66	NA
						Transfusion incidence (FFP/postoperative)	0% vs 0%	No difference NA	NA
						Transfusion incidence (platelets/postoperative)	0% vs 0%	No difference NA	NA
Liver surgery		1		1	1	1	•	1	1
Henry (2007)	Level I Good	2 RCTs N=296	Adult patients undergoing <u>liver</u> surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Transfusion incidence (allogeneic blood)	RR 0.16 (0.00, 32.47)	No difference 0.50	Substantial Phet<0.001 (I ² =93%)
Other surgery									
Henry (2007)	Level I/II Good/Fair	1 RCT N=59	Adult patients undergoing vascular surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Transfusion incidence (allogeneic blood)	RR 0.56 (0.33, 0.96)	Favours tranexamic acid 0.035	NA

Study	Level of	No. of trials /	Patient population / Surgical procedure	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size included in analysis		Location			Risk estimate (95% CI) or TXA (%) vs control (%)	Significance P-value	
Kongnyuy (2009)	Level I/II Good/Good	1 RCT N=100	Adult patients undergoing myomectomy	Hospital – planned surgery Turkey	Tranexamic acid (IV) vs placebo	Transfusion incidence	RR 1.71 (0.68, 4.30)	No difference 0.25	NA
Choi (2009)	Level II Fair	1 RCT N=61	Adult patients undergoing orthognathic surgery	Hospital – planned surgery China (Hong Kong)	Tranexamic acid (IV) vs placebo	Transfusion incidence	12.5% vs 24.1%	No difference 0.32	NA
PAEDIATRIC POPULA	TION/IV APROTINI	N							
Orthopaedic surg	ery								
Tzortzopoulou (2008)	Level II Good	2 RCT N=84	Paediatric patients undergoing scoliosis surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs placebo	Transfusion incidence	RR 0.84 (0.56, 1.27)	No difference 0.41	None Phet=0.94 (I ² =0%)
ADULT POPULATION	TOPICAL TRANEX	AMIC ACID							
Cardiac surgery									
Abrishami (2009)	Level I Good	2 RCTs N=233	Adult patients undergoing on- pump cardiac surgery	Hospital – planned surgery Unknown	Tranexamic acid (topical) vs placebo	Transfusion incidence (allogeneic RBC)	RR 0.98 (0.75, 1.27)	No difference 0.88	None Phet=0.69 (I ² =0%)
ADULT POPULATION	ORAL TRANEXAMI	IC ACID							
Cardiac surgery									
Gurusamy (2009)	Level I/II Poor	1 RCT N=214	Adults patients undergoing liver resection	Hospital – planned surgery China	Tranexamic acid (oral) vs no tranexamic acid	Transfusion incidence (allogeneic blood)	RR 0.03 (0.00, 0.46)	Favours tranexamic acid 0.012	NA

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: ASA, acetylsalicylic acid; CI, confidence interval; het, heterogeneity; IV, intravenous; NA, not applicable; NR, not reported; OR, odds ratio; RBC, red blood cell; RCT, randomised controlled trial; RR, risk ratio.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level II.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between aprotinin and no aprotinin.

d Cochrane ratings defined as follows: Grade A, adequate allocation concealment; Grade B, uncertain allocation concealment; Grade C, inadequate allocation concealment.

Key question(s):		Evidence table refa:				
In patients undergoing surgery, what is the effect of administration of tranexal	mic a	acid on transfusion volume?	POQ3.I8b.P2			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	5)					
There is one pivotal Level I (Henry 2007/good quality) study which includes data from 11 RCTs and four additional level II studies (Taghaddomi 2009/fair quality; Alvarez 2008/fair quality; Elwatidy 2008/fair quality; Wong 2008/good	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias			
quality) which provide data on the transfusion volume in patients who received transfusion. There was one pivotal Level I (Henry 2007/good quality) study which includes data from 14 RCTs and five additional Level II studies (Later	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias			
2009/good quality; Maddali 2007/good quality; Mehr-Aein 2007/good quality; Elwatidy 2008/fair quality; Sadeghi 2007/good quality) which provide data on transfusion volume in all patients (transfused or not).	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
2007/good quality, milor provide data of italistasion volume in air patients (italistased of net).	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	3			
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Results somewhat inconsistent between pivotal meta-analysis and additional RCTs. Consistency of individual studies within meta-analysis described below. Results either favour tranexamic acid or show no difference. Potential	Α	All studies consistent				
causes for differences between studies include different denominators used (all patients or transfused patients),	В	Most studies consistent and inconsistency can be explained				
different surgery types and different blood products transfused. Pivotal evidence – Henry 2007	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
Substantial heterogeneity (see note) in main analyses.	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>undetermined</u>)	nknow	<u>n</u> factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be			
Pivotal evidence – Henry 2007	Α	Very large				
Pivotal evidence – Henry 2007 Any surgery (units; all patients) – WMD -1.12 (-1.59, -0.64); 14 RCTs (N=965) Any surgery (units; transfused patients – WMD -0.51 (-1.06, 0.04); 11 RCTs (N=429)	В	Substantial				
Supportive evidence – see Summary Table POQ3.18b.P2	С	Moderate				
	D	Slight/Restricted				
4. Generalisability (How well does the body of evidence match the population and clinical set	tings b	peing targeted by the Guideline?)				
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were included in the overall analysis. There were also analyses of patients undergoing cardiac surgery who had received	Α	Evidence directly generalisable to target population				
ASA within 7 days prior to surgery.	В	Evidence directly generalisable to target population with some cave	veats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of hear	Ith services/delivery of care and cultural factors?)				
Hospital setting. The pivotal reviews included studies from a wide range of countries.	Α	Evidence directly applicable to Australian healthcare context				
Included RCTs were from a number of different countries including several from the Middle East and Asia.	В	Evidence applicable to Australian healthcare context with few caveats				
	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
	D	Evidence not applicable to Australian healthcare context				

The Advisory Committee on Prescription Medicines (ACPM) has recently recommended approval of tranexamic acid injection "for the reduction of peri- and postoperative blood loss and of the need for blood transfusion in adult patients undergoing cardiac surgery or total knee or hip arthroplasty; and paediatric patients undergoing cardiac surgery". Tranexamic acid tablets are approved in Australia for a number of indications including haemostatic, hereditary angioedema, short-term treatment of traumatic hyphaema, patients with coagulopathies undergoing minor surgery, and menorrhagia.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I (good quality) study and four additional level II studies which provide data on the transfusion volume in patients who received transfusion. There was one pivotal Level I (good quality) study and five additional Level II studies which provide data on transfusion volume in all patients (transfused or not).
2. Consistency	В	Some inconsistency, likely due to different denominators (all vs transfused patients), surgery type and blood products.
3. Clinical impact	С	There was generally a slight to moderate reduction in transfusion volume associated with tranexamic acid therapy compared with no therapy
4. Generalisability	С	The results are generalisable to an adult surgical population; the majority of evidence is in cardiac and orthopaedic surgery.
5. Applicability	В	There were a reasonable number of studies conducted in different countries. The additional RCTs were conducted in the Netherlands, Oman, Iran, Saudi Arabia, Spain and Canada. Likely to be applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, intravenous tranexamic acid therapy may reduce the volume of allogeneic blood transfusion compared with no therapy.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <5%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: ASA, acetylsalicylic acid: RCT, randomised controlled trial: SR, systematic review: WMD, weighted mean difference.

Primary outcomes; P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.l8b.P2 Characteristics and results of studies examining the effect of <u>tranexamic acid</u> on <u>transfusion volume</u>

Study	Level of	No. of trials /	Patient population / Surgical	Setting Location	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size included in analysis	procedure				Risk estimate (95% CI) or TXA (mean) vs control (mean)	Significance P-value	
ADULT POPULATION/	IV TRANEXAMIC A	ACID							
Any surgery									
Henry (2007)	Level I Good	14 RCTs N=965	Adult patients undergoing any surgery (all patients)	Hospital – planned surgery Various countries ^c	Tranexamic acid (IV) vs no tranexamic acid	Transfusion volume (units; allogeneic blood)	WMD -1.12 (-1.59, -0.64)	Favours tranexamic acid <0.001	Substantial Phet<0.001 (l²=73%)
	Level I Good	11 RCTs N=429	Adult patients undergoing any surgery (transfused patients only)	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Transfusion volume (units; allogeneic blood)	WMD -0.51 (-1.06, 0.04)	No difference 0.071	Substantial Phet<0.001 (l²=74%)
Cardiac surgery					•				
Later (2009)	Level II Good	1 RCT N=202	Adult patients undergoing <u>first-time</u> , <u>non-complex cardiac</u> <u>surgery with CPB (all patients)</u>	Hospital – planned surgery The Netherlands	Tranexamic acid (IV) vs placebo	Transfusion volume (units; <u>pRBC</u>)	Comparison of medians: 1.0 vs 2.0	Favours tranexamic acid 0.038	NA
Maddali (2007)	Level II 1 RCT Good N=222	N=222 CABG surgery (all patients)	Hospital – planned surgery Oman	d Tranexamic acid (IV) vs placebo	Transfusion volume (mL; total pRBC)	609 vs 952	Favours tranexamic acid 0.001	NA	
						Transfusion volume (units; total FFP)	0.72 vs 1.6	Favours tranexamic acid <0.01	NA
					Transfusion volume (units; total platelets)	0.7 vs 0.8	No difference NS	NA	
Mehr-Aein (2007)	Level II Good	1 RCT N=66	Adults undergoing primary off- pump CABG surgery (all patients)	Hospital – planned surgery Iran	Tranexamic acid (IV) vs placebo	Transfusion volume (units; whole blood or pRBC)	0.46 vs 0.94	Favours tranexamic acid 0.001	NA

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or TXA (mean) vs control (mean)	Significance P-value	
Taghaddomi (2009)	Level II Fair	1 RCT N=100	Adult patients undergoing off- pump CABG surgery (transfused patients only)	Hospital – planned surgery Iran	Tranexamic acid (IV) vs placebo	Transfusion volume (units; pRBC/intraoperative)	0 vs 1	No difference 0.36	NA
						Transfusion volume (units; pRBC/0-4 postoperative)	0 vs 1.3	Favours tranexamic acid <0.001	NA
						Transfusion volume (units; pRBC/4-24 postoperative)	1 vs 1	No difference 0.5	NA
						Transfusion volume (units; FFP/0-4 postoperative)	3 vs 2.5	No difference 0.8	NA
						Transfusion volume (units; FFP/4-24 postoperative)	0 vs 0	No difference NA	NA
						Transfusion volume (units; FFP/4-24 postoperative)	1 vs 1.1	No difference NR	NA
Orthopaedic surge	ery								
Alvarez (2008)	Level II Fair	1 RCT N=95	Adult patients undergoing total knee arthroplasty (transfused patients only)	Hospital – planned surgery Spain	Tranexamic acid (IV) vs placebo	Transfusion volume (total RBC; units)	1 vs 1.8	NR	NA
				Эран		Transfusion volume (allogeneic RBC; units)	1 vs NR (8 units in unspecified number of patients)	NR	NA
						Transfusion volume (autologous RBC; units)	0 vs NR (3 units in unspecified number of patients)	NR	NA
Elwatidy (2008)	Level II Fair	1 RCT N=64	Adult or paediatric patients undergoing spine surgery (all patients)	Hospital – planned surgery Saudi Arabia	Tranexamic acid (IV) vs placebo	Transfusion volume (mL)	94 vs 531	Favours tranexamic acid 0.008	NA
			Adult or paediatric patients undergoing spine surgery (transfused patients only)			Transfusion volume (units)	1.5 vs 2.8 ^d	NA	NA
Sadeghi (2007)	Level II Good	1 RCT N=67	Adult patients undergoing hip fracture surgery (all patients)	Hospital – planned surgery Iran	Tranexamic acid (IV) vs placebo	Transfusion volume (whole blood or pRBC; units)	1.25 vs 1.95	Favours tranexamic acid 0.001	NA

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity⁵
	evidence ^a Quality	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or TXA (mean) vs control (mean)	Significance P-value	
Wong (2008)	Level II Good	1 RCT N=147	Adult patients undergoing spinal fusion surgery (transfused patients)e	Hospital – planned surgery Canada	Tranexamic acid (IV) vs placebo	Transfusion volume (mL; pRBC/perioperative)	266 vs 406	No difference 0.16	NA
						Transfusion volume (mL; AWB/perioperative)	222 vs 315	No difference 0.30	NA
						Transfusion volume (mL; cell-saver/perioperative)	218 vs 334	No difference 0.083	NA
						Transfusion volume (mL; pRBC/intraoperative)	169 vs 208	No difference 0.61	NA
						Transfusion volume (mL; AWB/intraoperative)	150 vs 249	No difference 0.24	NA
						Transfusion volume (mL; cell-saver/intraoperative)	210 vs 323	No difference 0.086	NA
						Transfusion volume (mL; pRBC/postoperative)	97 vs 198	No difference 0.057	NA
						Transfusion volume (mL; AWB/postoperative)	72 vs 66	No difference 0.85	NA
						Transfusion volume (mL; cell-saver/postoperative)	8 vs 11	No difference 0.73	NA
PAEDIATRIC POPULA	TION/IV TRANEXA	MIC ACID							
Cardiac surgery									
Schouten (2009)	Level I Good	NR N=460	Paediatric patients undergoing cardiac surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs no tranexamic acid	Transfusion volume (mL/kg; <u>pRBC</u>)	WMD -7 (-10, -5)	Favours tranexamic acid NR	None Phet=NR (I²=6%)
		NR N=419	Paediatric patients undergoing cardiac surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs no tranexamic acid	Transfusion volume (mL/kg; <u>plasma</u>)	WMD -7 (-9, -4)	Favours tranexamic acid NR	None Phet=NR (I²=0%)

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size included in analysis	procedure	Location		Transfusion volume (mL/kg; <u>thrombo</u>)	Risk estimate (95% CI) or TXA (mean) vs control (mean)	Significance P-value	
		NR N=370	Paediatric patients undergoing cardiac surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs no tranexamic acid		WMD -5 (-7, -3)	Favours tranexamic acid NR	None Phet=NR (I²=0%)
Orthopaedic surge	ery	•	-1	•	1	1	1	'	•
Schouten (2009)	Level I Good	2 RCTs N=84	Paediatric patients undergoing scoliosis surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs no tranexamic acid	Transfusion volume (mL; <u>pRBC</u>)	WMD -349 (-620, -77)	Favours tranexamic acid NR	None Phet=NR (I²=0%)
						Transfusion volume (mL; plasma)	WMD -15 (-127, -98)	Favours tranexamic acid NR	None Phet=NR (I²=24%)
Tzortzopoulou (2008)	Level I Good	2 RCT N=84	Paediatric patients undergoing scoliosis surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs placebo	Transfusion volume (mL)	WMD -395 (-688, -103)	Favours ε- aminocaproic acid 0.0081	None Phet=0.51 (I ² =0%)
ADULT POPULATION	TOPICAL APROTIN	IIN							
Abrishami (2009)	Levell	3 RCTs N=229	Adult patients undergoing on- pump cardiac surgery	Hospital – planned surgery Unknown	Tranexamic acid (topical) vs no tranexamic acid	Transfusion volume (units)	WMD -1.58 (-2.26, -0.90)	Favours tranexamic acid <0.001	None Phet=0.29 (I ² =20%)
Fawzy (2009)	Level II Good	1 RCT N=38	Adult patients undergoing primary elective CABG surgery	Hospital – planned surgery Saudi Arabia	Tranexamic acid (topical) vs placebo	Transfusion volume (units; pRBC/postoperative)	Comparison of medians: 1.0 vs 1.0	No difference 0.82	NA
			Sa			Transfusion volume (units; FFP/postoperative)	Comparison of medians: 0 vs 2.0	No difference 0.42	NA
						Transfusion volume (units; plasma/postoperative)	Comparison of medians: 0 vs 2.0	Favours tranexamic acid 0.03	NA

Note: Studies/analyses providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies/analyses provide supportive evidence.

Abbreviations: CI, confidence interval; FFP, fresh frozen plasma; het, heterogeneity; IV, intravenous; kg, kilogram; mL, millilitre; NA, not applicable; NR, not reported; RBC, red blood cell; RCT, randomised controlled trial; SMD, standardised mean difference; WMD, weighted mean difference.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level II.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I²<25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between aprotinin and no aprotinin.

d One patient received 14 units of blood. If this person is excluded the mean number of units transfused per transfused patient is 1.8.

^e Not specifically stated but appears to be based on transfused patients only.

Key question(s):			Evidence table refa:
In patients undergoing surgery, what is the effect of administration of tranexa	mic a	acid on <u>blood loss</u> ?	POQ3.I8b.P3
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)		
There is one pivotal Level I study (Henry 2007/good quality) which includes data from up to 23 RCTs, two supportive level I studies (Brown 2007/fair quality; Kagoma 2009/good quality), two supportive Level I/II studies which included	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias
data from one RCT each (McIlroy 2009/good-fair quality; Kongnyuy 2009/good-good quality) and 13 additional RCTs published since the pivotal review (Jimenez 2007/good quality; Later 2009/good quality; Maddali 2007/good	В	One or two Level II studies with a low risk of bias or SR/several Lev	/el III studies with a low risk of bias
quality; Mehr-Aein 2007/good quality; Taghaddomi 2009/fair quality; Alvarez 2008/fair quality; Elwatidy 2008/good quality; Sadeghi 2007/good quality; Wong 2008/good quality; Chen 2008/fair quality; Choi 2009/fair quality; Mayur	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias
2007/poor quality; Sekhavat 2009/poor quality).	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	1
2. Consistency (If only one study was available, rank this component as 'not applicable')			
Results consistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta- analyses described below.	Α	All studies consistent	
Pivotal evidence – Henry 2007	В	Most studies consistent and inconsistency can be explained	
Substantial significant heterogeneity (see note) in main analyses (postoperative) and subgroup analyses. Differences likely due to different surgeries and measurement and timing of blood loss.	С	Some inconsistency, reflecting genuine uncertainty around question	on
Supportive evidence	D	Evidence is inconsistent	
Most results showed significantly less blood loss with tranexamic acid compared with no tranexamic acid.	NA	Not applicable (one study only)	
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u>	nknow	<u>n</u> factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be
Pivotal evidence – Henry 2007	Α	Very large	
Any surgery (mL; total blood loss) – WMD -444 (-572, -315); 17 RCTs (N=955) Cardiac surgery (mL; total blood loss) – WMD -440 (-607, -273); 3 RCTs (N=245)	В	Substantial	
Orthopaedic surgery (mL; total blood loss) – WMD -440 (-591, -288); 14 RCTs (N=690) Supportive evidence – See Summary Table POQ3.l8b.P3	С	Moderate	
Supportive evidence – See Summary Table POQS.lob.PS	D	Slight/Restricted	
4. Generalisability (How well does the body of evidence match the population and clinical se	ttings b	being targeted by the Guideline?)	
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were included in the overall analysis. There were also analyses of patients undergoing cardiac surgery who had received	Α	Evidence directly generalisable to target population	
ASA within 7 days prior to surgery.	В	Evidence directly generalisable to target population with some cav	eats
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of heal	Ith services/delivery of care and cultural factors?)	
Hospital setting. The pivotal reviews included studies from a wide range of countries.	Α	Evidence directly applicable to Australian healthcare context	
Included RCTs were from a number of different countries including several from the Middle East and Asia.	В	Evidence applicable to Australian healthcare context with few cave	ats
	С	Evidence probably applicable to Australian healthcare context with	ı some caveats
	D	Evidence not applicable to Australian healthcare context	

The Advisory Committee on Prescription Medicines (ACPM) has recently recommended approval of tranexamic acid injection "for the reduction of peri- and postoperative blood loss and of the need for blood transfusion in adult patients undergoing cardiac surgery or total knee or hip arthroplasty; and paediatric patients undergoing cardiac surgery". Tranexamic acid tablets are approved in Australia for a number of indications including haemostatic, hereditary angioedema, short-term treatment of traumatic hyphaema, patients with coagulopathies undergoing minor surgery, and menorrhagia.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (good quality), two supportive level I studies, two supportive Level I/II study and 13 additional RCTs.
2. Consistency	В	Some inconsistency, likely due to different surgery types and timing of outcome measurement. Inconsistency related to magnitude of effect rather than direction of effect.
3. Clinical impact	В	There was generally a substantial reduction in blood loss associated with TXA.
4. Generalisability	С	The results are generalisable to a general surgical population.
5. Applicability	В	Overall there were a reasonable number of studies conducted in a range of countries. Individual additional RCTs were conducted in Spain, The Netherlands, Oman, Iran, Saudi Arabia, Canada, Turkey, Hong Kong and India. Likely to be applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, intravenous tranexamic acid therapy reduces blood loss compared with no therapy.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: ASA, acetylsalicylic acid; RCT, randomised controlled trial; SR, systematic review; TXA, tranexamic acid; WMD, weighted mean difference.

a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

POQ3.l8b.P3 Characteristics and results of studies examining the effect of <u>tranexamic acid</u> on <u>blood loss</u>

Study	Level of	No. of trials /	Patient population / Surgical procedure	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size included in analysis		Location			Risk estimate (95% CI) or TXA (mean) vs control (mean)	Significance P-value	
ADULT POPULATIO	ON/ IV TRANEXAMIC A	ACID							
Any surgery									
Henry (2007)	Level I Good	10 RCTs N=553	Adult patients undergoing any surgery	Hospital – planned surgery Various countries ^c	Tranexamic acid (IV) vs no tranexamic acid	Intraoperative blood loss (mL)	WMD -55 (-105, -4.5)	Favours tranexamic acid 0.033	None Phet=0.26 (I ² =20%)
		23 RCTs N=1423	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	vs no tranexamic loss (mL)	WMD -248 (-313, -183)	Favours tranexamic acid <0.001	Substantial Phet<0.001 (I ² =76%)	
		18 RCTs N=955	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Total blood loss (mL)	WMD -444 (-572, -315)	Favours tranexamic acid <0.001	Substantial Phet<0.001 (I ² =72%)
Cardiac surgery	′				•				
Henry (2007)	Level I Good	3 RCTs N=144	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Intraoperative blood loss (mL)	WMD -287 (-482, -93)	Favours tranexamic acid 0.0038	None Phet=0.66 (I ² =0%)
		17 RCTs N=1130	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Postoperative blood loss (mL)	WMD -263 (-319, -207)	Favours tranexamic acid <0.001	Moderate Phet=0.01 (I ² =48%)
		9 RCTs N=302	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Postoperative blood loss (mL; total dose < 2.0 g)	WMD -252 (-352, -151)	Favours tranexamic acid <0.001	Moderate Phet=0.07 (1 ² =45%)
		8 RCTs N=828	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Postoperative blood loss (mL; total dose 2.0-10.0 g)	WMD -272 (-341, -205)	Favours tranexamic acid <0.001	Substantial Phet=0.03 (I ² =54%)
		3 RCTs N=245	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Total blood loss (mL)	WMD -440 (-607, -273)	Favours tranexamic acid <0.001	None Phet=0.82 (l ² =0%)
Brown (2007)	Level I Fair	11 RCTs N=1100	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	Tranexamic acid_(IV) vs placebo	Total blood loss (mL)	WMD -285 (-394, -175)	Favours tranexamic acid <0.001	NR

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity⁵
	evidence ^a <i>Quality</i>	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or TXA (mean) vs control (mean)	Significance P-value	
McIlroy(2009)	Level I/II Good/Fair	1 RCT N=79	Adult patients receiving ASA undergoing cardiac surgery	Hospital – planned surgery Unknown	Lysine analogues (IV) vs placebo	Postoperative chest tube blood loss (mL)	WMD -189 (-287, -91)	Favours lysine analogues <0.001	Substantial Phet=0.05 I2=67%
Jimenez (2007)	Level II Good	1 RCT N=50	Adults undergoing cardiopulmonary bypass surgery	Hospital - planned surgery Spain	Tranexamic acid (IV) vs placebo	24-hour blood loss (mL)	464 vs 1037	Favours tranexamic acid <0.01	NA
						Total blood loss (mL)	835 vs 1466	Favours tranexamic acid <0.01	NA
Later (2009)	Level II Good	1 RCT N=202	Adults undergoing fi <u>rst-time</u> , non-complex cardiac surgery with CPB	Hospital - planned surgery The Netherlands	Tranexamic acid (IV) vs placebo	Total mediastinal chest tube blood loss (mL)	760 vs 860	Favours tranexamic acid 0.034	NA
Maddali (2007)	Level II Good	1 RCT N=222	Adults undergoing primary CABG surgery	Hospital – planned surgery Oman	Tranexamic acid (IV) vs placebo	Total drainage (mL)	633 vs 981	Favours tranexamic acid 0.001	NA
Mehr-Aein (2007)	Level II Good	1 RCT N=66	Adults undergoing off-pump CABG surgery	Hospital - planned surgery Iran	Tranexamic acid (IV) vs placebo	Postoperative blood loss <u>0-2 hr</u> (mL)	90 vs 180	Favours tranexamic acid <0.001	NA
						Postoperative blood loss 2-6 hr (mL)	190 vs 290	Favours tranexamic acid 0.001	NA
						Total postoperative blood loss (mL)	320 vs 480	Favours tranexamic acid 0.001	NA

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or TXA (mean) vs control (mean)	Significance P-value	
Taghaddomi (2009)	Level II Fair	1 RCT N=100	Adults undergoing off-pump CABG surgery	Hospital - planned surgery Iran	Tranexamic acid (IV) vs placebo	Intraoperative bleeding (mL)	467 vs 531	No difference 0.62	NA
				II dii		Postoperative bleeding (mL; 0-4 hr)	87 vs 210	Favours tranexamic acid 0.005	NA
						Postoperative bleeding (mL; 4-24 hr)	462 vs 570	No difference 0.07	NA
						Total bleeding (mL; within 24 hr)	471 vs 844	Favours tranexamic acid <0.001	NA
Orthopaedic surg	jery		<u> </u>	l	<u> </u>	<u>I</u>	l		
Henry (2007)	Level I Good	7 RCTs N=409	Adult patients undergoing orthopaedic surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Intraoperative blood loss (mL)	WMD -30 (-69, 10)	No difference 0.14	None Phet=0.69 (I ² =0%)
		6 RCTs N=293	Adult patients undergoing orthopaedic surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Postoperative blood loss (mL)	WMD -210 (-384, -35)	Favours tranexamic acid 0.019	Substantial Phet<0.001 (l²=91%)
		14 RCT N=690	Adult patients undergoing orthopaedic surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Total blood loss (mL)	WMD -440 (-591, -288)	Favours tranexamic acid <0.001	Substantial Phet<0.001 (l ² =78%)
Kagoma (2009)	Level I Good	15 RCTs N=778	Adults undergoing total knee or hip replacement	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs no tranexamic acid	Total bleeding ^d (mL)	WMD -393 (-442, -345)	Favours tranexamic acid NR	NR
Alvarez (2008)	Level II Fair	1 RCT N=95	Adult patients undergoing total knee arthroplasty	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs placebo	Chest-tube blood loss (mL; 0-6 hr postoperative)	159 vs 534	Favours tranexamic acid <0.001	NA
						Chest-tube blood loss (mL; <u>6 hr – 4 day</u> postoperative)	132 vs 132	No difference 0.98	NA
						Total chest-tube blood loss (mL)	170 vs 551	Favours tranexamic acid <0.001	NA

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or TXA (mean) vs control (mean)	Significance P-value	
Elwatidy (2008)	Level II Fair	1 RCT N=64	Adult and paediatric patients undergoing spine surgery	Hospital – planned surgery Saudi Arabia	Tranexamic acid (IV) vs placebo	Intraoperative blood loss (mL)	311 vs 585	Favours tranexamic acid 0.03	NA
						Wound drain blood loss (mL)	98 vs 215	Favours tranexamic acid 0.004	NA
						Total blood loss (mL)	406 vs 800	Favours tranexamic acid 0.007	NA
Sadeghi (2007)	Level II Good	1 RCT N=67	Adults undergoing hip fracture surgery	Hospital – planned surgery Iran	Tranexamic acid (IV) vs placebo	Perioperative blood loss (mL)	652 vs 1108	Favours tranexamic acid 0.003	NA
						Postoperative blood loss 1 hr (mL)	111 vs 139	No difference 0.39	NA
						Postoperative blood loss 2 hr (mL)	192 vs 246	No difference 0.28	NA
						Postoperative blood loss 5 hr (mL)	255 vs 323	No difference 0.31	NA
						Postoperative blood loss 12 hr (mL)	296 vs 375	No difference 0.20	NA
						Postoperative blood loss 24 hr (mL)	300 vs 390	No difference 0.11	NA
						Total blood loss (mL)	960 vs 1484	Favours tranexamic acid 0.001	NA

Study	Level of	No. of trials / sample size	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	included in analysis	procedure	Location			Risk estimate (95% CI) or TXA (mean) vs control (mean)	Significance P-value	
Wong (2008)	Level II Good	1 RCT N=147	Adults undergoing spinal fusion surgery	Hospital – planned surgery Canada	Tranexamic acid (IV) vs placebo	Perioperative blood loss (estimated; mL)	1592 vs 2138	Favours tranexamic acid 0.026	NA
						Perioperative blood loss (calculated; mL)	3079 vs 4363	Favours tranexamic acid 0.017	NA
						Perioperative RBC loss (calculated; mL)	1078 vs 1527	Favours tranexamic acid 0.017	NA
						Intraoperative blood loss (estimated; mL)	1203 vs 1600	Favours tranexamic acid 0.044	NA
						Postoperative blood loss (estimated; mL)	536 vs 737	Favours tranexamic acid 0.039	NA
Liver surgery									
Henry (2007)	Level I/II Good/Poor	1 RCT N=20	Adult patients undergoing orthotopic liver transplant	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs placebo	Blood loss (mL)	MD -6552 (-14330, 1226)	No difference 0.099	NA
Other surgery									
Kongnyuy (2009)	Level I/II Good/Good	1 RCT N=100	Adult patients undergoing myomectomy	Hospital – planned surgery Turkey	Tranexamic acid (IV) vs placebo	Transection blood loss (mL)	MD -243 (-460, -26)	Favours tranexamic acid 0.028	NA
Chen (2008)	Level II Fair	1 RCT N=55	Adult patients undergoing head and neck surgery	Hospital – planned surgery Taiwan	Tranexamic acid (IV) vs placebo	Perioperative bleeding (mL)	87 vs 116	No difference 0.392	NA
				. S. Pull		Drainage amount (mL)	50 vs 89	Favours tranexamic acid 0.04	NA

Study	Level of	No. of trials /	Patient population / Surgical procedure	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or TXA (mean) vs control (mean)	Significance P-value	
Choi (2009)	Level II Fair	1 RCT N=44	Adult patients undergoing anterior mandibular surgery	Hospital – planned surgery Hong Kong	Tranexamic acid (IV) vs placebo	Intraoperative or postoperative blood loss (mL)	277 vs 416	No difference NS	NA
		1 RCT N=61	Adult patients undergoing maxillary surgery	Hospital – planned surgery Hong Kong	Tranexamic acid (IV) vs placebo	Intraoperative or postoperative blood loss (mL)	428 vs 644	Favours tranexamic acid <0.05	NA
		1 RCT N=41	Adult patients undergoing ramus surgery	Hospital – planned surgery Hong Kong	Tranexamic acid (IV) vs placebo	Intraoperative or postoperative blood loss (mL)	287 vs 329	No difference NS	NA
		1 RCT N=61	Adult patients undergoing <u>any</u> orthognathic surgery	Hospital – planned surgery Hong Kong	Tranexamic acid (IV) vs placebo	Intraoperative or postoperative blood loss (mL)	879 vs 1257	Favours tranexamic acid	NA
Mayur (2007)	Level II Poor	1 RCT N=100	Adult patients undergoing caesarean section	Hospital – planned surgery India	Tranexamic acid (IV) vs placebo	Post-partum haemorrhage (<u>placental delivery to</u> <u>end of surgery</u> ; mL)	299 vs 340	No difference 0.056	NA
						Post-partum haemorrhage (end of surgery to 2 hr post- partum; mL)	76 vs 133	Favours tranexamic acid 0.001	NA
						Post-partum haemorrhage (placental delivery to 2 hr post-partum; mL)	375 vs 473	Favours tranexamic acid 0.003	NA
Sekhavat (2009)	Level II Poor	1 RCT N=90	Adult patients undergoing caesarean section	Hospital – planned surgery Iran	Tranexamic acid (IV) vs placebo	Blood loss up to 2 hr postoperative (mL)	28 vs 37	Favours tranexamic acid <0.001	NA
PAEDIATRIC POPULA	ATION/IV TRANEXA	MIC ACID							
Cardiac surgery Schouten (2009)	Level I Good	NR N=542	Paediatric patients undergoing scoliosis surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs no tranexamic acid	Blood loss (mL/kg)	WMD -11 (-13, -8)	Favours tranexamic acid NR	Moderate Phet=NR (l²=31)

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or TXA (mean) vs control (mean)	Significance P-value	
Orthopaedic surge	ery	•							•
Schouten (2009)	Level I Good	2 RCTs N=84	Paediatric patients undergoing scoliosis surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs no tranexamic acid	Blood loss (mL)	WMD -682 (-1149, -214)	Favours tranexamic acid NR	Unclear Phet=NR (l²=24)
Tzortzopoulou (2008)	Level I Good	2 RCTs N=84	Paediatric patients undergoing scoliosis surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs placebo	Blood loss (mL)	WMD -682 (-1149, -214)	Favours tranexamic acid 0.0042	Mild Phet=0.25 (l²=24)
ADULT POPULATION/	TOPICAL TRANEX	AMIC ACID							
Cardiac surgery									
Abrishami (2009)	Level I Good	4 RCTs N=269	Adult patients undergoing on- pump cardiac surgery	Hospital – planned surgery Unknown	Tranexamic acid (topical) vs placebo	24-hr postoperative chest-tube loss (mL)	WMD -250 (-465, -35)	Favours tranexamic acid 0.02	Substantial Phet<0.001 (l²=95%)
Fawzy (2009)	Level II Good	1 RCT N=38	Adult patients undergoing primary isolated CABG surgery	Hospital – planned surgery Unknown	Tranexamic acid (topical) vs placebo	24-hr chest tube blood loss (mL)	626 vs 1040	Favours tranexamic acid 0.04	NA
						Total chest-tube blood loss (mL)	656 vs 1056	<i>Unclear</i> NR	NA
Jabalameli (2006)	Level II Poor	1 RCT N=56	Adult patients undergoing endoscopic sinus surgery	Hospital – planned surgery Iran	Tranexamic acid (topical) vs placebo	Intraoperative blood loss (mL)	174 vs 229	Favours tranexamic acid <0.05	NA
Other surgery									
Athanasiadis (2007)	Level II Fair	1 RCT N=30	Adult patients undergoing endoscopic sinus surgery	Hospital – planned surgery Australia	Tranexamic acid 100 mg (topical) vs placebo	Bleeding grading scalese at 0, 2, 4, 6, 8 and 10 mins	NR	Favours tranexamic acid <0.05	NA
					Tranexamic acid 1 g (topical) vs placebo	Bleeding grading scalese at 0, 2, 4, 6, 8 and 10 mins	NR	Favours tranexamic acid <0.05	NA

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidencea sample size included in analysis N/ORALTRANEXAMIC ACID			Risk estimate (95% CI) or TXA (mean) vs control (mean)	Significance P-value				
ADULT POPULATION	ORAL TRANEXAM	IC ACID							
Liver surgery									
Gurusamy (2009)	Level I/II Fair	1 RCT N=214	Adult patients undergoing liver surgery	Hospital – planned surgery China	Tranexamic acid (oral) vs placebo	Transection blood loss (mL)	MD -260 (-435, -85)	Favours tranexamic acid 0.0036	NA
						Operative blood loss (mL)	MD -300 (-502, -98)	Favours tranexamic acid 0.0036	NA

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: ASA, acetylsalicylic acid; CI, confidence interval; het, heterogeneity; IV, intravenous; mL, millilitres; NA, not applicable; NR, not reported; RCT, randomised controlled trial; SMD, standardised mean difference; WMD, weighted mean difference.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level II.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between tranexamic acid and no tranexamic acid.

^d Total bleeding measured intraoperatively by weighing surgical sponges, postoperatively through drainage or perioperatively through the haemoglobin balance method which measures loss through comparison of pre- and postoperative haemoglobin concentrations (haematoma volumes as well as hidden or internal blood loss were excluded).

^e Wormald grading scale and Boezaart grading scale.

In patients undergoing surgery, what is the effect of administration of tranexamic acid on mortality? 1. Evidence base (number of studies, level of evidence and risk of bias in the included studies) There is one pital terel study (Brown 2007/hiar quality) and four additional RCTs published since the photol review (Immerz 2007/spood quality). Letter 2009/spood quality. Mehr Aein 2007/spood quality. All xerial and part 2007/spood quality. Mehr Aein 2007/spood quality. All xerial and part 2007/spood quality. Mehr Aein 2007/spood quality. All xerial and part 2007/spood quality. All xerial 2007	Key question(s):			Evidence table refa:			
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The Advisory Committee on Prescription Medicines (ACPM) has recently recommended approval of tranexamic acid injection "for the reduction of peri- and postoperative blood loss and of the need for blood transfusion in adult patients undergoing cardiac surgery or total knee or hip arthroplasty; and paediatric patients undergoing cardiac surgery". Tranexamic acid tablets are approved in Australia for a number of indications including haemostatic, hereditary angioedema, short-term treatment of traumatic hyphaema, patients with coagulopathies undergoing minor surgery, and menorrhagia.

Generalisability made a C as most studies were conducted in cardiac surgery but the evidence statement has been applied to the whole surgical population.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (good quality), one supportive Level I study and four additional RCTs.
2. Consistency	Α	Results are consistent. There was no heterogeneity in the pivotal study.
3. Clinical impact	D	While there is no significant difference in mortality between intravenous tranexamic acid therapy and no therapy, and the risk estimates suggest no increased risk, the findings are uncertain due to underpowering.
4. Generalisability	С	The results are generalisable to a surgical population; however, most studies were conducted in cardiac surgery.
5. Applicability	В	There were a moderate number of studies conducted in a range of countries. The additional RCTs were conducted in Spain, The Netherlands and Iran. Likely to be applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intravenous tranexamic acid therapy on mortality compared with no therapy is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: RCT, randomised controlled trial; RR, risk ratio; SR, systematic review.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.l8b.P4 Characteristics and results of studies examining the effect of <u>tranexamic acid</u> on <u>mortality</u>

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size	Surgical procedure	Location			Risk estimate (95% CI) or TXA (%) vs control (%)	Significance P-value	
Adult Population Any surgery	IV TRANEXAMIC A	ACID							
Henry (2007)	Level I Good	16 RCTs N=1684	Adult patients undergoing any surgery	Hospital – planned surgery Various countries ^c	Tranexamic acid (IV) vs no tranexamic acid	Mortality	RR 0.60 (0.32, 1.12)	No difference 0.11	None Phet=0.84 (I ² =0%)
Cardiac surgery		•	1	1	1	•	<u>'</u>	•	'
Henry (2007)	Level I Good	11 RCTs N=1390	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Mortality	RR 0.55 (0.24, 1.25)	No difference 0.15	None Phet=0.73 (l²=0%)
Henry (2009)	Level I Good	11 RCTs N=1390	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Mortality	RR 0.55 (0.24, 1.25)	No difference NR	NR
Brown (2007)	Level I Fair	18 RCTs N=2229	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs placebo	Mortality	RR 0.67 (0.33, 1.37)	No difference 0.276	NR
Jimenez (2007)	Level II Good	1 RCT N=50	Adult patients undergoing cardiopulmonary bypass surgery	Hospital – planned surgery Spain	Tranexamic acid (IV) vs placebo	In-hospital mortality	0% vs 0%	No difference NA	NA
Later (2009)	Level II Good	1 RCT N=202	Adult patients undergoing first-time, non-complex cardiac surgery with CPB	Hospital – planned surgery The Netherlands	Tranexamic acid (IV) vs placebo	In-hospital mortality	1% vs 1%	No difference 1.00	NA
Mehr-Aein (2007)	Level II Good	1 RCT N=66	Adult patients undergoing primary CABG surgery	Hospital – planned surgery Iran	Tranexamic acid (IV) vs placebo	In-hospital mortality	0% vs 0%	No difference NA	NA
Orthopaedic surg	ery			•					
Sadeghi (2007)	Level II Good	1 RCT N=67	Adult patients undergoing hip fracture surgery	Hospital – planned surgery Iran	Tranexamic acid (IV) vs placebo	In-hospital mortality	0% vs 3%	No difference 1.00	NA
ADULT POPULATION	TOPICAL TRANEX	AMIC ACID							
Cardiac surgery	_								
Fawzy (2009)	Level II Good	1 RCT N=28	Adult patients undergoing primary isolated CABG surgery	Hospital – planned surgery Saudi Arabia	Tranexamic acid (topical) vs placebo	In-hospital mortality	0% vs 0%	No difference NA	NA

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
Adult population/	evidence ^a <i>Quality</i> ORAL TRANEXAMO	sample size	Surgical procedure	Location			Risk estimate (95% CI) or TXA (%) vs control (%)	Significance P-value	
Cardiac surgery									
Gurusamy (2009)	Level I/II Poor	1 RCT N=214	Adults patients undergoing liver resection	Hospital – planned surgery China	Tranexamic acid (oral) vs no tranexamic acid	Mortality	0% vs 0%	No difference NA	NA

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: CI, confidence interval; het, heterogeneity; IV, intravenous; NA, not applicable; NR, not reported; RCT, randomised controlled trial; RR, risk ratio.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level II.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and |2<25%; (ii) mild heterogeneity if |2<25%; moderate heterogeneity if |2 between 25-50%; substantial heterogeneity |2 >50%.

c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between tranexamic acid and no tranexamic acid.

Key question(s):			Evidence table refa:			
In patients undergoing surgery, what is the effect of administration of tranexa	mic a	acid on morbidity (myocardial infarction)?	POQ3.I8b.P5 (MI)			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studie	s)					
There is one pivotal Level I study (Henry 2007/good quality), one supportive Level I study (Brown 2007/fair quality)	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias			
and four additional RCTs (Later 2009/good quality; Mehr-Aein 2007/good quality; Taghaddomi 2009/fair quality; Wong 2008/good quality).	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias			
Trong 2000, good quality)	С	One or two Level III studies with a low risk of bias or Level I or II st				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Results consistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta-	Α	All studies consistent				
analyses described below. Pivotal evidence – Henry 2007	В	Most studies consistent and inconsistency can be explained				
No significant heterogeneity (see note) in main analyses	С	Some inconsistency, reflecting genuine uncertainty around question	on			
Supportive evidence – nearly all studies showed no difference between tranexamic acid and no tranexamic acid.	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u> determined)	nknow	<u>n</u> factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be			
Pivotal evidence – Henry 2007	Α	Very large				
Any surgery – 1.7% vs 1.9%; RR 0.96 (0.48, 1.90); 12 RCTs (N=1344) Cardiac surgery – 1.5% vs 1.9%; RR 0.91 (0.44, 1.88); 9 RCTs (N=1048)	В	Substantial				
Supportive evidence – Brown 2007 (see also Summary Table POQ3.l8b.P5 (MI)	С	Moderate				
Cardiac surgery – no absolute risks reported; RR 0.94 (0.51, 1.74) (N=2219)	D	No difference				
4. Generalisability (How well does the body of evidence match the population and clinical se	ttings l	peing targeted by the Guideline?)				
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were	Α	Evidence directly generalisable to target population				
included in the overall analysis of the pivotal review but 9/12 included RCTs in cardiac surgery.	В	Evidence directly generalisable to target population with some cav	reats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to			
		apply				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of hea					
Hospital setting.	Α	Evidence directly applicable to Australian healthcare context				
The pivotal review included studies from a wide range of countries. Included RCTs were from a number of different countries including The Netherlands, Iran and Canada.	В	Evidence applicable to Australian healthcare context with few caveats				
misidaded 1013 were from a number of different countries including the neutronalids, it all and callada.	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
	D	Evidence not applicable to Australian healthcare context				

The Advisory Committee on Prescription Medicines (ACPM) has recently recommended approval of tranexamic acid injection "for the reduction of peri- and postoperative blood loss and of the need for blood transfusion in adult patients undergoing cardiac surgery or total knee or hip arthroplasty; and paediatric patients undergoing cardiac surgery". Tranexamic acid tablets are approved in Australia for a number of indications including haemostatic, hereditary angioedema, short-term treatment of traumatic hyphaema, patients with coagulopathies undergoing minor surgery, and menorrhagia.

CRG concerned regarding the definition of MI in the individual included studies.

Generalisability made a C as most studies were conducted in cardiac surgery but the evidence statement is being applied to the whole surgical population.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (good quality), one supportive Level I study and four additional RCTs.
2. Consistency	Α	There was no heterogeneity in the pivotal level I evidence. Additional studies were consistent.
3. Clinical impact	D	There was no significant difference in the risk of myocardial infarction between intravenous tranexamic acid therapy and no therapy.
4. Generalisability	С	The results are generalisable to a surgical population; however, most studies were conducted in cardiac surgery.
5. Applicability	В	A reasonable number of studies were conducted in a range of countries. The additional RCTs were conducted in The Netherlands, Iran and Canada. Likely to be applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, intravenous tranexamic acid therapy does not appear to have an effect on risk of myocardial infarction compared with no therapy.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI, confidence interval, RBC, red blood cell: RCT, randomised controlled trial; RR, relative risk; SR, systematic review.

a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

POQ3.l8b.P5 (MI) Characteristics and results of studies examining the effect of <u>tranexamic acid</u> on <u>morbidity (myocardial infarction)</u>

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		
	evidence ^a <i>Quality</i>	sample size	Surgical procedure				Risk estimate (95% CI) or TXA (%) vs control (%)	Significance P-value	
ADULT POPULATION	IV TRANEXAMIC A	ACID							
Any surgery									
Henry (2007)	Level I Good	12 RCTs N=1344	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries ^c	Tranexamic acid (IV) vs no tranexamic acid	Myocardial infarction	RR 0.96 (0.48, 1.90)	No difference 0.91	None Phet=0.96 (I ² =0%)
Cardiac surgery		•							
Henry (2007)	Level I Good	9 RCTs N=1048	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Myocardial infarction	RR 0.91 (0.44, 1.88)	No difference 0.79	None Phet=0.91 (l²=0%)
Henry (2009)	Level I Good	10 RCTs N=1148	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs placebo	Myocardial infarction	RR 0.86 (0.43, 1.75)	No difference NR	NR
Brown (2007)	Level I Fair	16 RCTs N=2219	Adult patients undergoing cardiac surgery	Hospital – planned surgery Countries not specified	Tranexamic acid (IV) vs placebo	Myocardial infarction	RR 0.94 (0.51, 1.74)	No difference 0.85	NR
Later (2009)	Level II Good	1 RCT N=202	Adult patients undergoing first-time, non-complex cardiac surgery with CPB	Hospital – planned surgery The Netherlands	Tranexamic acid (IV) vs placebo	Perioperative myocardial infarction	0% vs 8%	Favours tranexamic acid 0.007	NA
Mehr-Aein (2007)	Level II Good	1 RCT N=66	Adult patients undergoing primary CABG surgery	Hospital – planned surgery Iran	Tranexamic acid (IV) vs placebo	Myocardial infarction	0% vs 0%	No difference NA	NA
Taghaddomi (2009)	Level II Fair	1 RCT N=100	Adult patients off-pump CABG surgery	Hospital – planned surgery Iran	Tranexamic acid (IV) vs placebo	Myocardial infarction	0% vs 0%	No difference NA	NA
						Myocardial ischaemia	0% vs 0%	No difference NA	NA

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size	Surgical procedure				Risk estimate (95% CI) or TXA (%) vs control (%)	Significance P-value	
Orthopaedic sur	gery								
Wong (2008)	Level II Good N/TOPICAL TRANEX	1 RCT N=147	Adult patients undergoing spinal fusion surgery	Hospital – planned surgery Canada	Tranexamic acid (IV) vs placebo	Myocardial infarction	1% (asymptomatic only) vs 0%	No difference NA	NA
Cardiac surgery		AMIC ACID							
Fawzy (2009)	Level II Good	1 RCT N=38	Adult patients undergoing primary isolated CABG surgery	Hospital – planned surgery Saudi Arabia	Tranexamic acid (topical) vs placebo	In-hospital myocardial infarction	0% vs 0%	No difference NA	NA

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: CABG, coronary artery bypass graft; CI, confidence interval; het, heterogeneity; IV, intravenous; NA, not applicable; NR, not reported; RCT, randomised controlled trial; RR, risk ratio.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level II.

^b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between tranexamic acid and no tranexamic acid.

In patients undergoing surgery, what is the effect of administration of <u>tranexamic acid</u> on <u>morbidity (renal)?</u> 1. Evidence base (number of studies, level of evidence and risk of bias in the included studies) There is one photal twel I study (Henry 2007/good quality) which includes data from 4 RCTs, one supportive Level study (Henry 2007/good quality) and two additional RCTs published since the photal review (Later 2009/good quality). A One or more level I studies with a low risk of bias or several level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or Level I or II studies with a low risk of bias or Level I or III studies wit	enal)				
There is one pivotal Level I study (Henry 2007/good quality) which includes data from 4 RCTs, one supportive Level I study (Brown 2007/fair quality) and two additional RCTs published since the pivotal review (Later 2009/good quality). ### A					
Study (Brown 2007/fpair quality) and two additional RCTs published since the pivotal review (Later 2009/good quality). C					
pualify. Mehr-Aein 2007/good quality). B One or two Level II studies with a low risk of bias or SR/several Level II studies with a low risk of bias or SR/several Level II studies with a low risk of bias or SR/several Level I or II studies with a moderate risk D Level IV studies or Level I to III studies/SRs with a high risk of bias 2. Consistency (If only one study was available, rank this component as 'not applicable') Results inconsistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta-analyses described below. Protal evidence – Henry 2007 No significant heterogeneity (see note) in analysis. Supportive evidence While all results not stalistically significant, point estimates from Brown meta-analyses for renal failure and renal deformation of them yesult. 3. Clinical impact (Indicate in the space below if the study results varied according to some unknown factor (not simply study quality or sample size) and thus the clinical impact of the intervention could relaterationard. Protal evidence – Henry 2007 Cardiac surgery (renal failure) – no absolute risks reported: RR 1.43 (0.30, 6.85): 3 RCTs (N-840) Supportive evidence – Brown 2007 Cardiac surgery (renal disfurce) – no absolute risks reported: RR 1.43 (0.30, 6.85): 3 RCTs (N-840) Supportive evidence of Summary Table PO03.8b.P5 (renal) 4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?) The evidence is generalisable to an adult population who are undergoing cardiac surgery. A Evidence directly generalisable to target population	of bias				
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2. Consistency (if only one study was available, rank this component as 'not applicable') Results inconsistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta-analyses described below. Pivotal evidence – Henry 2007 Not applicable (inconsistent) (see note) in analysis. Supportive evidence While all results not statistically significant, point estimates from Brown meta-analyses for renal failure and renal dysfunction were in the opposite direction to Henry 2007 point estimate for both outcomes combined. 2 included RCTs were consistent with Henry result. 3. Clinical impact (Indicate in the space below if the study results varied according to some unknown factor (not simply study quality or sample size) and thus the clinical impact of the intervention could redetermined? Privotal evidence – Henry 2007 Cardiac surgery (renal failure) – no absolute risks reported: RR 1.43 (0.30, 6.85): 3 RCTs (N=80) Supportive evidence – Brown 2007 Cardiac surgery (renal failure) – no absolute risks reported: RR 2.02 (0.73, 5.60): 4 RCTs (N=684) See also Summary Table POQ3.18b.P5 (renal) 4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?) The evidence is generalisable to an adult population who are undergoing cardiac surgery. A Il studies consistent B All studies consistent B Most studies consistent and inconsistency can be explained C Some inconsistent B Most studies consistent and inconsistency an	k of bias				
Results inconsistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta-analyses described below. Privotal evidence – Henry 2007 No significant heterogeneity (see note) in analysis. Supportive evidence While all results not statistically significant, point estimates from Brown meta-analyses for renal failure and renal dysfunction were in the opposite direction to Henry 2007 point estimate for both outcomes combined. 2 included RCTs were consistent with Henry result. 3. Clinical impact (Indicate in the space below if the study results varied according to some unknown factor (not simply study quality or sample size) and thus the clinical impact of the intervention could relative intervention of the intervention could relative intervention of the interv					
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Privotal evidence – Henry 2007 No significant heterogeneity (see note) in analysis. Supportive evidence While all results not statistically significant, point estimates from Brown meta-analyses for renal failure and renal dysfunction were in the opposite direction to Henry 2007 point estimate for both outcomes combined. 2 included RCTs were consistent with Henry result. 3. Clinical impact (Indicate in the space below if the study results varied according to some unknown factor (not simply study quality or sample size) and thus the clinical impact of the intervention could relatermined. Privotal evidence – Henry 2007 Cardiac surgery (renal failure) – no absolute risks reported; RR 1.43 (0.30, 6.85); 3 RCTs (N=840) See also Summary Table POQ3.18b.P5 (renal) 4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?) The evidence is generalisable to an adult population who are undergoing cardiac surgery. A Some inconsistent and inconsistency can be explained C Some inconsistency, reflecting genuine uncertainty around question Evidence is inconsistent NA Not applicable (one study only) Evidence is inconsistent NA Very large Substantial C Moderate C Moderate D Inconsistent 4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?) A Evidence directly generalisable to target population					
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Supportive evidence – Brown 2007 Cardiac surgery (renal failure) – no absolute risks reported; RR 1.43 (0.30, 6.85); 3 RCTs (N=840) Cardiac surgery (renal dysfunction) - no absolute risks reported; RR 2.02 (0.73, 5.60); 4 RCTs (N=684) See also Summary Table POQ3.l8b.P5 (renal) 4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?) The evidence is generalisable to an adult population who are undergoing cardiac surgery. A Evidence directly generalisable to target population					
Cardiac surgery (renal dysfunction) - no absolute risks reported; RR 2.02 (0.73, 5.60): 4 RCTs (N=684) See also Summary Table POQ3.l8b.P5 (renal) D Inconsistent 4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?) The evidence is generalisable to an adult population who are undergoing cardiac surgery. A Evidence directly generalisable to target population					
See also Summary Table POQ3.l8b.P5 (renal) 4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?) The evidence is generalisable to an adult population who are undergoing cardiac surgery. A Evidence directly generalisable to target population					
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The evidence is generalisable to an adult population who are undergoing cardiac surgery. A Evidence directly generalisable to target population					
B Evidence directly generalisable to target population with some caveats					
C Evidence not directly generalisable to the target population but could be sensibly applied					
D Evidence not directly generalisable to target population and hard to judge whether it is sensi	sible to				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of health services/delivery of care and cultural factors?)					
Hospital setting. A Evidence directly applicable to Australian healthcare context The pivotal review included studies from a number of countries.					
Included RCTs were from The Netherlands and Iran. B Evidence applicable to Australian healthcare context with few caveats					
C Evidence probably applicable to Australian healthcare context with some caveats					
D Evidence not applicable to Australian healthcare context					

The Advisory Committee on Prescription Medicines (ACPM) has recently recommended approval of tranexamic acid injection "for the reduction of peri- and postoperative blood loss and of the need for blood transfusion in adult patients undergoing cardiac surgery or total knee or hip arthroplasty; and paediatric patients undergoing cardiac surgery". Tranexamic acid tablets are approved in Australia for a number of indications including haemostatic, hereditary angioedema, short-term treatment of traumatic hyphaema, patients with coagulopathies undergoing minor surgery, and menorrhagia.

Brown defined renal outcomes as follows: (i) renal failure defined as a ≥ 0.5 mg/dL increase in creatinine.

Later 2009 RCT defined outcomes as follows: (i) renal failure as defined by Mangano (2006): required a postoperative serum creatinine of at least 2.0 mg/dL with an increase over the preoperative baseline level of at least 0.7 mg/dL; (ii) renal complication as defined by the RIFLE classification: risk of renal dysfunction defined as a 1.5 times increase in perioperative creatinine plasma concentration or a urine output < 0.5 mL/kg/h in 6 hours. Kidney injury was defined as a 2 times increase in perioperative creatinine plasma concentration or a urine output < 0.5 mL/kg/h in 24 hours.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (good quality), one supportive Level I study and two additional RCTs.
2. Consistency	D	Different direction of point estimate for pivotal and supportive Level I studies.
3. Clinical impact	D	There was no difference in risk of renal failure/dysfunction in the primary study but there was potentially an increased risk in the supportive study; thus the results are not consistent.
4. Generalisability	В	The results are generalisable to a cardiac surgical population.
5. Applicability	В	A small number of studies were conducted in a range of countries. The additional RCTs were conducted in The Netherlands and Iran. Likely to be applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, the effect of intravenous tranexamic acid therapy on risk of renal failure or dysfunction, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: RCT, randomised controlled trial: RR, risk ratio: SR, systematic review,

a Interventions: 11 = ANH, 12 = intraoperative cell salvage, 13 = perioperative ANH and intraoperative cell salvage, 14 = postoperative cell salvage, 15 = deliberate induced hypotension, 16 = prevention of hypothermia, 17 = point-of-care testing for coagulation status and haemoglobin, 18 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, 19 = appropriate patient positioning, 110 = PAD

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

POQ3.l8c.P5 (renal) Characteristics and results of studies examining the effect of <u>tranexamic acid</u> on <u>morbidity (renal failure/dysfunction)</u>

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results	3		ilts	
	evidence ^a Quality	sample size	Surgical procedure				Risk estimate (95% CI) or TXA (%) vs control (%)	Significance P-value			
ADULT POPULATION	/IV TRANEXAMIC A	ACID									
Cardiac surgery											
Henry (2007)	Level I Good	4 RCTs N=400	Adult patients undergoing any surgery	Hospital – planned surgery Unknown ^c	Tranexamic acid (IV) vs no tranexamic acid	Renal failure/dysfunction	RR 0.73 (0.16, 3.32)	No difference 0.68	None P <i>het</i> =0.69 (I ² =0%)		
Brown (2007)	Level I Fair	3 RCTs N=840	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs placebo	Renal failure ^d	RR 1.43 (0.30, 6.85)	No difference 0.66	NR		
		4 RCTs N=684	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs placebo	Renal dysfunction ^e	RR 2.02 (0.73, 5.60)	No difference 0.18	NR		
Later (2009)	Level II Good	1 RCT N=202	Adult patients undergoing first-time, non-complex cardiac surgery with CPB	Hospital – planned surgery The Netherlands	Tranexamic acid (IV) vs placebo	Renal failure ^f	3% vs 3%	No difference 1.00	NA		
			Surado Surgery With Of D	The Netherlands		Renal complication ^g	8% vs 18%	No difference 0.059	NA		
Mehr-Aein (2007)	Level II Good	1 RCT N=66	Adult patients undergoing primary CABG	Hospital – planned surgery Iran	Tranexamic acid (IV) vs placebo	Renal dysfunction (creatinine > 2 mg/dL)	0% vs 3%	No difference >0.05	NA		

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: CABG, coronary artery bypass graft; CI, confidence interval; DVT, deep vein thrombosis; *het*, heterogeneity; IV, intravenous; MI, myocardial infarction; NA, not applicable; NR, not reported; PE, pulmonary embolism; OR, odds ratio; RCT, randomised controlled trial; RD, risk difference; RR, risk ratio.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level II.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between tranexamic acid and no tranexamic acid.

d Renal failure defined as a new onset of dialysis except in one study where it was defined as a ≥ 2 mg/dL creatinine level.

e Renal dysfunction defined as a ≥ 0.5 mg/dL increase in creatinine.

Fenal failure as defined by Mangano (2006)3. Required a postoperative serum creatinine of at least 2.0 mg/dL with an increase over the preoperative baseline level of at least 0.7 mg/dL.

⁹ Renal complication as defined by the RIFLE classification ⁴. Risk of renal dysfunction defined as a 1.5 times increase in perioperative creatinine plasma concentration or a urine output < 0.5 mL/kg/h in 6 hours. Kidney injury was defined as a 2 times increase in perioperative creatinine plasma concentration or a urine output < 0.5 mL/kg/h in 12 hours, whilst renal failure was defined as a 3 times increase in perioperative creatinine plasma concentration or a urine output < 0.5 mL/kg/h in 24 hours.

³ Mangano et al (2006) The risk associated with aprotinin in cardiac surgery. NEJM 354:353-365.

⁴ Kuitunen et al (2006) Acute renal failure after cardiac surgery: evaluation of the RIFLE classification. Ann Thorac Surg 81: 542-546.

Key question(s):			Evidence table refa:		
In patients undergoing surgery, what is the effect of administration of tranexal	mic a	<u>acid</u> on <u>morbidity (stroke)</u> ?	POQ3.I8b.P5 (stroke)		
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	5)				
There is one pivotal Level I study (Henry 2007/good quality) which includes data from 7 RCTs, one supportive Level	Α	One or more level I studies with a low risk of bias or several level	I studies with a low risk of bias		
I study (Brown 2007/fair quality) and one additional RCT (Later 2009/good quality).	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias		
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias		
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias			
2. Consistency (If only one study was available, rank this component as 'not applicable')					
Results consistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta-	Α	All studies consistent			
analyses described below. Pivotal evidence – Henry 2007	В	Most studies consistent and inconsistency can be explained			
No significant heterogeneity (see note) in main analyses.	С	Some inconsistency, reflecting genuine uncertainty around questi	on		
Supportive evidence	D	Evidence is inconsistent			
Similar results between systematic reviews and single additional RCT.	NA	Not applicable (one study only)			
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>un</u> determined)	<u>nknow.</u>	n factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be		
Pivotal evidence – Henry 2007	Α	Very large			
Any surgery – 1.4% vs 1.1%; RR 1.25 (0.47, 3.31); 7 RCTs (N=937)	В	Substantial			
Cardiac surgery – 1.3% vs 0.8%; RR 1.52 (0.52, 4.41); 5 RCTs (N=841) Supportive evidence – Brown 2007 (see also Summary Table POQ3.l8b.P5 (stroke)	С	Moderate			
Cardiac surgery – no absolute risks reported; RR 1.31 (0.59, 2.93); 15 RCTs (N=2098)	D	Underpowered			
4. Generalisability (How well does the body of evidence match the population and clinical set	tings b	neing targeted by the Guideline?)			
The evidence is generalisable to an adult population who are undergoing planned surgery. Most evidence in cardiac	Α	Evidence directly generalisable to target population			
surgery.	В	Evidence directly generalisable to target population with some cav	reats		
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied		
	D	Evidence not directly generalisable to target population and hard to judge whether it is sensible to apply			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of heal	th services/delivery of care and cultural factors?)			
Hospital setting.	Α	Evidence directly applicable to Australian healthcare context			
The pivotal review included studies from a number of countries. The additional included RCT was from The Netherlands.	В	Evidence applicable to Australian healthcare context with few cave	eats		
The additional included NOT was from the Netherlands.	С	Evidence probably applicable to Australian healthcare context with some caveats			
	D	Evidence not applicable to Australian healthcare context			

The Advisory Committee on Prescription Medicines (ACPM) has recently recommended approval of tranexamic acid injection "for the reduction of peri- and postoperative blood loss and of the need for blood transfusion in adult patients undergoing cardiac surgery or total knee or hip arthroplasty; and paediatric patients undergoing cardiac surgery". Tranexamic acid tablets are approved in Australia for a number of indications including haemostatic, hereditary angioedema, short-term treatment of traumatic hyphaema, patients with coagulopathies undergoing minor surgery, and menorrhagia.

Generalisability made a C as most studies in cardiac surgery but evidence statement being applied to the whole surgical population.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (good quality), one supportive Level I study and one additional RCT.
2. Consistency	Α	There was no heterogeneity in the pivotal study. The supportive study was consistent. The additional RCT was underpowered to detect a difference in stroke between tranexamic acid therapy and no therapy.
3. Clinical impact	D	Results show a slightly increased risk with no significant difference (9/711 versus 5/634 in cardiac surgery) but studies likely underpowered to detect a difference in stroke.
4. Generalisability	С	The results are generalisable to a surgical population; however, most studies conducted in cardiac surgery.
5. Applicability	В	Studies were conducted in a range of countries. The additional RCT was conducted in The Netherlands. Likely to be applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intravenous tranexamic acid therapy on risk of stroke, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: RCT, randomised controlled trial: RR, risk ratio: SR, systematic review.

^a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

POQ3.l8b.P5(stroke) Characteristics and results of studies examining the effect of <u>tranexamic acid</u> on <u>morbidity (stroke)</u>

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results	Heterogeneity ^b	
	evidence ^a sample size Surgical procedure <i>Quality</i>		Risk estimate (95% CI) or TXA (%) vs control (%)	Significance P-value					
ADULT POPULATIO	N/ IV TRANEXAMIC A	CID							
Any surgery									
Henry (2007)	Level I Good	7 RCTs N=937	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Stroke	RR 1.25 (0.47, 3.31)	No difference 0.65	None Phet=0.79 (l ² =0%)
Cardiac surgery				•					
Henry (2007)	Level I Good	5 RCTs N=841	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Stroke	RR 1.52 (0.52, 4.41)	No difference 0.44	None Phet=0.78 (I ² =0%)
Brown (2007)	Level I Fair	15 RCTs N=2098	Adult patients undergoing cardiac surgery	Hospital – planned surgery Countries not specified	Tranexamic acid (IV) vs placebo	Stroke	RR 1.31 (0.59, 2.93)	No difference 0.51	NR
Later (2009)	Level II Good	1 RCT N=202	Adult patients undergoing first-time, non-complex cardiac surgery with CPB	Hospital – planned surgery The Netherlands	Tranexamic acid (IV) vs placebo	Stroke	1% vs 1%	No difference 1.00	NR

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: ASA, acetylsalicylic acid; CI, confidence interval; CPB, cardiopulmonary bypass; DVT, deep vein thrombosis; het, heterogeneity; IV, intravenous; MI, myocardial infarction; NR, not reported; PE, pulmonary embolism; OR, odds ratio; RCT, randomised controlled trial; RR, risk ratio; TIA, transient ischaemic attack.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level II.

^b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I²<25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

^c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between tranexamic acid and no tranexamic acid.

Key question(s):			Evidence table refa:		
In patients undergoing surgery, what is the effect of administration of tranexam	nic a	cid on morbidity (thrombosis)?	POQ3.I8b.P5 (thromb)		
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies)					
	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias			
supportive Level I study (Kagoma 2009/good quality), one supportive Level I/II study which included data from one RCT (McIlroy 2009/good-fair quality) and eight additional RCTs published since the pivotal review (Taghaddomi	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias		
209/fair quality; Alvarez 2008/fair quality; Elwatidy 2008/fair quality; Wong 2008/fair quality; Chen 2008/fair quality	С	One or two Level III studies with a low risk of bias or Level I or II st			
Choi 2009/fair quality; Mayur 2007/poor quality; Sekhavat 2009/poor quality).	D	Level IV studies or Level I to III studies/SRs with a high risk of bias			
2. Consistency (If only one study was available, rank this component as 'not applicable')					
	Α	All studies consistent			
analyses described below. Pivotal evidence – Henry 2007	В	Most studies consistent and inconsistency can be explained			
No significant heterogeneity (see note) in analyses of any surgery or cardiac surgery for DVT or PE.	С	Some inconsistency, reflecting genuine uncertainty around question	on		
Supportive evidence	D	Evidence is inconsistent			
All studies showed no difference between tranexamic and no tranexamic acid.	NA	Not applicable (one study only)			
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>unk</u> determined)	knowr	n factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be		
	Α	Very large			
Any surgery (DVT) – 1.9% vs 2.9%; RR 0.77 (0.37, 1.61); 10 RCTs (N=681)	В	Substantial			
Any surgery (PE) – 0.4% vs 1.3%; RR 0.55 (0.17, 1.76); 7 RCTs (N=568) Cardiac surgery (DVT) – 0% vs 1.0%; RR 0.37 (0.04, 3.47); 2 RCTs (N=291)	С	Moderate			
Cardiac surgery (PE) – 0% vs 0.7%; RR 0.33 (0.04, 3.15); 2 RCTs (N=289)	D	Underpowered			
4. Generalisability (How well does the body of evidence match the population and clinical setting	ings b	eing targeted by the Guideline?)			
	Α	Evidence directly generalisable to target population			
included in the overall analysis of the pivotal review; a number of studies were in orthopaedic surgery.	В	Evidence directly generalisable to target population with some cav	reats		
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied		
	D	Evidence not directly generalisable to target population and hard to judge whether it is sensible to apply			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	f heali	th services/delivery of care and cultural factors?)			
' '	Α	Evidence directly applicable to Australian healthcare context			
The pivotal review included studies from a number of countries. Included RCTs were from a number of different countries including several from the Middle East and Asia.	В	Evidence applicable to Australian healthcare context with few caveats			
included NC13 were from a number of different countries including several from the whole East alid Asia.	С	Evidence probably applicable to Australian healthcare context with some caveats			
	D	Evidence not applicable to Australian healthcare context			

The Advisory Committee on Prescription Medicines (ACPM) has recently recommended approval of tranexamic acid injection "for the reduction of peri- and postoperative blood loss and of the need for blood transfusion in adult patients undergoing cardiac surgery or total knee or hip arthroplasty; and paediatric patients undergoing cardiac surgery". Tranexamic acid tablets are approved in Australia for a number of indications including haemostatic, hereditary angioedema, short-term treatment of traumatic hyphaema, patients with coagulopathies undergoing minor surgery, and menorrhagia.

The thrombosis outcome includes deep vein thrombosis, pulmonary embolism and outcomes that have been specified as thrombosis but not further defined in included studies.

Generalisability made a C as being applied to the whole surgical population.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (good quality), one supportive Level I study , one supportive Level I/II study and eight additional RCTs.
2. Consistency	Α	There was no heterogeneity in the pivotal study. Other studies showed consistent results.
3. Clinical impact	D	There was no significant difference in any results but some of the risk estimates were large. Likely to be underpowered for thrombosis outcomes.
4. Generalisability	С	The results are generalisable to a surgical population; however, the majority of studies included in the pivotal and supportive Level I evidence were conducted in orthopaedic surgery.
5. Applicability	В	A moderate number of studies were conducted in a range of countries. The additional RCTs were conducted in Iran, Spain, Saudi Arabia, Canada, Taiwan, China and India. Likely to be applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intravenous tranexamic acid therapy on risk of thrombosis, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: DVT, deep vein thrombosis; PR, pulmonary embolism; RCT, randomised controlled trial; RR, risk ratio; SR, systematic review.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.l8c.P5 (thrombosis) Characteristics and results of studies examining the effect of <u>tranexamic acid</u> on <u>morbidity (thrombosis)</u>

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results	Heterogeneity ^b	
	evidence ^a sample size Surgical procedure Quality			Risk estimate (95% CI) or TXA (%) vs control (%)	Significance P-value				
ADULT POPULATION	N/ IV TRANEXAMIC A	ACID							
Any surgery									
Henry (2007)	Level I Good	10 RCTs N=681	Adult patients undergoing any surgery	Hospital – planned surgery Unknown ^c	Tranexamic acid (IV) vs no tranexamic acid	DVT	RR 0.77 (0.37, 1.61)	No difference 0.49	None P <i>het</i> =0.81 (I ² =0%)
Henry (2007)	Level I Good	7 RCTs N=568	Adult patients undergoing any surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs no tranexamic acid	PE	RR 0.55 (0.17, 1.76)	No difference 0.31	None P <i>het</i> =0.93 (I ² =0%)
Henry (2007)	Level I Good	2 RCTs N=114	Adult patients undergoing any surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs no tranexamic acid	Other thrombosis	RR 2.10 (0.49, 8.99)	No difference 0.32	None P <i>het</i> =0.80 (I ² =0%)
Cardiac surgery	•	•							
Henry (2007)	Level I Good	2 RCTs N=291	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs no tranexamic acid	DVT	RR 0.37 (0.04, 3.47)	No difference 0.38	None P <i>het</i> =0.95 (I ² =0%)
Henry (2007)	Level I Good	2 RCTs N=289	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs no tranexamic acid	PE	RR 0.33 (0.04, 3.15)	No difference 0.34	None P <i>het</i> =0.98 (I ² =0%)
McIlroy (2009)	Level I/II Good/Fair	1 RCT N=79	Adult patients receiving ASA undergoing cardiac surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs placebo	Thrombotic complications	OR 0.32 (0.01, 8.02)	No difference 0.49	NA
Taghaddomi (2009)	Level II Fair	1 RCT N=100	Adult patients undergoing off-pump CABG surgery	Hospital – planned surgery Iran	Tranexamic acid (IV) vs placebo	Thrombosis	0% vs 0%	No difference NA	NA
Orthopaedic surg	gery								
Kagoma (2009)	Level I Good	10 RCTs N=459	Adult patients undergoing total hip or knee replacement surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs placebo	VTE	RD -0.01 (-0.04, 0.02)	No difference NR	NR
Alvarez (2008)	Level II Fair	1 RCT N=95	Adult patients undergoing total knee arthroplasty	Hospital – planned surgery Spain	Tranexamic acid (IV) vs placebo	Thrombosis	0% vs 0%	No difference NA	NA

	Patient population /	Setting	Intervention	Outcome	Results	Results			
	evidence ^a sample size Surgical procedure Ouality		Risk estimate (95% CI) or TXA (%) vs control (%)	Significance P-value					
Elwatidy (2008)	Level II Fair	1 RCT N=64	Adult and paediatric patients undergoing spine surgery	Hospital – planned surgery Saudi Arabia	Tranexamic acid (IV) vs placebo	Thrombosis	0% vs 0%	No difference NA	NA
Wong (2008)	Level II Good	1 RCT N=147	Adult patients undergoing spinal fusion surgery	Hospital – planned surgery Canada	Tranexamic acid (IV) vs placebo	Thrombosis	0% vs 1%	No difference 1.00	NA
Other surgery	•	•			•			•	•
Chen (2008)	Level II Fair	1 RCT N=55	Adult patients undergoing head and neck surgery	Hospital – planned surgery Taiwan	Tranexamic acid (IV) vs placebo	DVT	0% vs 0%	No difference NA	NA
Choi (2009)	Level II Fair	1 RCT N=61	Adult patients undergoing orthognathic surgery	Hospital – planned surgery China	Tranexamic acid (IV) vs placebo	Thrombosis	0% vs 0%	No difference NA	NA
Mayur (2007)	Level II Poor	1 RCT N=100	Adult patients undergoing caesarean section	Hospital – planned surgery India	Tranexamic acid (IV) vs placebo	Thrombosis	0% vs 0%	No difference NA	NA
Sekhavat (2009)	Level II Poor	1 RCT N=90	Adult patients undergoing caesarean section	Hospital – planned surgery Iran	Tranexamic acid (IV) vs placebo	Thrombosis	0% vs 0%	No difference NA	NA

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: ASA, acetylsalicylic acid; CI, confidence interval; DVT, deep vein thrombosis; *het*, heterogeneity; IV, intravenous; MI, myocardial infarction; NA, not applicable; NR, not reported; PE, pulmonary embolism; OR, odds ratio; RCT, randomised controlled trial; RD, risk difference; RR, risk ratio.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level II.

^b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

Key question(s):			Evidence table refa:
In patients undergoing surgery, what is the effect of administration of tranexar	nic a	cid on quality of life?	POQ3.I8b.P6
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies,)		
No studies of any level were identified which assessed the effect of tranexamic acid on quality of life.	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias
	В	One or two Level II studies with a low risk of bias or SR/several Lev	rel III studies with a low risk of bias
	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	
2. Consistency (If only one study was available, rank this component as 'not applicable')			
NA	Α	All studies consistent	
	В	Most studies consistent and inconsistency can be explained	
	С	Some inconsistency, reflecting genuine uncertainty around question	on
	D	Evidence is inconsistent	
	NA	Not applicable (one study only)	
3. Clinical impact (Indicate in the space below if the study results varied according to some undetermined)	knowi	n factor (not simply study quality or sample size) and thus the clinical impa	ct of the intervention could not be
NA	Α	Very large	
	В	Substantial	
	С	Moderate	
	D	Slight/Restricted	
4. Generalisability (How well does the body of evidence match the population and clinical sett	tings b		
NA	Α	Evidence directly generalisable to target population	
	В	Evidence directly generalisable to target population with some cav	eats
	С	Evidence not directly generalisable to the target population but cou	ıld be sensibly applied
	D	Evidence not directly generalisable to target population and hard to	judge whether it is sensible to
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of heal	·	
NA	Α	Evidence directly applicable to Australian healthcare context	
	В	Evidence applicable to Australian healthcare context with few cave	
	С	Evidence probably applicable to Australian healthcare context with	some caveats
	D	Evidence not applicable to Australian healthcare context	

The Advisory Committee on Prescription Medicines (ACPM) has recently recommended approval of tranexamic acid injection "for the reduction of peri- and postoperative blood loss and of the need for blood transfusion in adult patients undergoing cardiac surgery or total knee or hip arthroplasty; and paediatric patients undergoing cardiac surgery". Tranexamic acid tablets are approved in Australia for a number of indications including haemostatic, hereditary angioedema, short-term treatment of traumatic hyphaema, patients with coagulopathies undergoing minor surgery, and menorrhagia.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	NA	
2. Consistency	NA	
3. Clinical impact	NA	
4. Generalisability	NA	
5. Applicability	NA	

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intravenous tranexamic acid therapy on quality of life, compared with no therapy, is unknown.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI. confidence interval. RBC, red blood cell: SR, systematic review:

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^a Interventions: 11 = ANH, 12 = intraoperative cell salvage, 13 = perioperative ANH and intraoperative cell salvage, 14 = postoperative cell salvage, 15 = deliberate induced hypotension, 16 = prevention of hypothermia, 17 = point-of-care testing for coagulation status and haemoglobin, 18 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, 19 = appropriate patient positioning, 110 = PAD

Key question(s):			Evidence table refa:			
In patients undergoing surgery, what is the effect of administration of tranexa	mic a	acid on re-operation for bleeding?	POQ3.l8b.S2			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)					
There is one pivotal Level I study (Henry 2007/good quality) which includes data from up to 18 RCTs, two suppo		One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias			
Level I studies (Henry 2009/good quality; Brown 2007/fair quality), one supportive Level I/II study which includes data from one RCT (McIlroy 2009/goog-fair quality) and three additional RCTs published since the pivotal review	В	One or two Level II studies with a low risk of bias or SR/several Level	vel III studies with a low risk of bias			
ater 2009/good quality; Maddali 2007/good quality; Mehr-Aein 2007/good quality).	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	;			
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Results consistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta-	Α	All studies consistent				
analyses described below. Pivotal evidence – Henry 2007	В	Most studies consistent and inconsistency can be explained				
No significant heterogeneity (see note) in main analyses	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
Supportive evidence	D	Evidence is inconsistent				
Results similar for Brown meta-analysis and additional RCTs.	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u> determined)	<u>ınknow</u>	n factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be			
Pivotal evidence – Henry 2007	Α	Very large				
Any surgery – 2.9% vs 5.0%; RR 0.67 (0.41, 1.09); 18 RCTs (N=1598) Cardiac surgery – 2.7% vs 4.9%; RR 0.65 (0.39, 1.08); 17 RCTs (N=1540)	В	Substantial				
Supportive evidence – Brown 2007 (see also Summary table POQ3.18b.S2)	С	Moderate				
Cardiac surgery – no absolute risks reported; RR 0.70 (0.44, 1.11); 21 RCTs (N=2255)	D	Underpowered				
4. Generalisability (How well does the body of evidence match the population and clinical se	ettings b	peing targeted by the Guideline?)				
The evidence is generalisable to an adult population who are undergoing planned surgery. Nearly all included	Α	Evidence directly generalisable to target population				
evidence was in cardiac surgery.	В	Evidence directly generalisable to target population with some cav	veats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to judge whether it is sensible to apply				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of hear	Ith services/delivery of care and cultural factors?)				
Hospital setting.	Α	Evidence directly applicable to Australian healthcare context				
The pivotal review included studies from a wide range of countries. Included RCTs were from a number of different countries including The Netherlands, Oman and Iran.	В	Evidence applicable to Australian healthcare context with few caveats				
included KC15 were from a number of different countries including the ineffections, Offidit did fidit.	С	Evidence probably applicable to Australian healthcare context with some caveats				
	D	Evidence not applicable to Australian healthcare context				

The Advisory Committee on Prescription Medicines (ACPM) has recently recommended approval of tranexamic acid injection "for the reduction of peri- and postoperative blood loss and of the need for blood transfusion in adult patients undergoing cardiac surgery or total knee or hip arthroplasty; and paediatric patients undergoing cardiac surgery". Tranexamic acid tablets are approved in Australia for a number of indications including haemostatic, hereditary angioedema, short-term treatment of traumatic hyphaema, patients with coagulopathies undergoing minor surgery, and menorrhagia.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (good quality), one supportive Level I study, one supportive Level I/II study and three additional RCTs.
2. Consistency	Α	There was no heterogeneity in the pivotal study and the results of the supportive and additional studies were consistent.
3. Clinical impact	D	There was no difference in risk of re-operation; however, the included studies may be underpowered to detect a difference.
4. Generalisability	С	The results are generalisable to a cardiac surgical population.
5. Applicability	В	There were a reasonable number of studies conducted in a range of countries. The additional RCTs were conducted in The Netherlands, Oman and Iran. Likely to be applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, the effect of intravenous tranexamic acid therapy on risk of reoperation due to bleeding, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: RCT. randomised controlled trial: RR. risk ratio: SR. systematic review.

^a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

POQ3.l8b.S2 Characteristics and results of studies examining the effect of <u>tranexamic acid</u> on <u>re-operation for bleeding</u>

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results	Heterogeneity ^b	
ADULT POPULATION.	evidence ^a <i>Quality</i> IV TRANEXAMIC A	sample size	Surgical procedure	Location			Risk estimate (95% CI) or TXA (%) vs control (%)	Significance P-value	
Any surgery									
Henry (2007)	Level I Good	18 RCTs N=1598	Adult patients undergoing any surgery	Hospital – planned surgery Various countries ^c	Tranexamic acid (IV) vs no tranexamic acid	Re-operation for bleeding	RR 0.67 (0.41, 1.09)	No difference 0.11	None Phet=0.92 (I ² =0%)
Cardiac surgery	•	•						•	
Henry (2007)	Level I Good	17 trials N=1540	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Re-operation for bleeding	RR 0.65 (0.39, 1.08)	No difference 0.097	None Phet=0.90 (I ² =0%)
Henry (2009)	Level I Good	NR NR	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs placebo	Re-operation for bleeding	RR 0.67 (0.41, 1.12)	No difference NR	NR
Brown (2007)	Level I Fair	21 RCTs N=2255	Adult patients undergoing cardiac surgery	Hospital – planned surgery Countries not specified	Tranexamic acid (IV) vs placebo	Return to operating room	RR 0.70 (0.44, 1.11)	No difference 0.125	NR
McIlroy(2009)	Level I/II Good/Fair	1 RCT N=79	Adult patients <u>receiving</u> <u>ASA</u> undergoing cardiac surgery	Hospital – planned surgery Unknown	Tranexamic acid (IV) vs placebo	Surgical re-exploration	OR 0.30 (0.01, 8.02)	No difference NR	NA
Later (2009)	Level II Good	1 RCT N=202	Adult patients undergoing first-time, non-complex cardiac surgery with CPB	Hospital – planned surgery The Netherlands	Tranexamic acid (IV) vs placebo	Re-operation for <u>any</u> reason	14% vs 14%	No difference 1.00	NA
				, no nomenana		Re-operation for <u>surgical</u> <u>bleeding</u>	3% vs 3%	No difference 1.00	NA
						Re-operation for non- surgical bleeding	2% vs 4%	No difference 0.68	NA
Maddali (2007)	Level II Good	1 RCT N=222	Adult patients undergoing primary CABG surgery	Hospital – planned surgery Oman	Tranexamic acid (IV) vs placebo	Re-operation for bleeding	3% vs 3%	No difference NS	NA
Mehr-Aein (2007)	Level II Good	1 RCT N=66	Adult patients undergoing primary CABG surgery	Hospital – planned surgery Iran	Tranexamic acid (IV) vs placebo	Re-exploration for bleeding	0% vs 3%	No difference >0.05	NA

Study		Patient population / Setting	Setting	Intervention	Outcome	Results	Heterogeneity ^b		
Adult population/	evidence ^a <i>Quality</i> TOPICAL TRANEXA	sample size	Surgical procedure	Location			Risk estimate (95% CI) or TXA (%) vs control (%)	Significance P-value	
Cardiac surgery									
Fawzy (2009)	Level II Good	1 RCT N=38	Adult patients undergoing primary isolated CABG surgery	Hospital – planned surgery Saudi Arabia	Tranexamic acid (topical) vs placebo	Re-operation for bleeding	5% vs 0%	No difference 1.00	NA

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: ASA, acetylsalicylic acid; CABG, coronary artery bypass graft; CI, confidence interval; CPB, cardiac pulmonary bypass; het, heterogeneity; IV, intravenous; NA, not applicable; NR, not reported; OR, odds ratio; RCT, randomised controlled trial; RR, risk ratio.

^a Where only one RCT is available in a systematic review, the level of evidence has been downgraded to Level II.

^b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between tranexamic acid and no tranexamic acid.

Key question(s): In patients undergoing surgery, what is the effect of administration of tranexamic acid on hospital.length.of.stay ? Description of POQ3.l8b.S5 1. Evidence base (number of studies, level of evidence and risk of bias in the included studies) There is one pivotal Level I study (Henry 2007/good quality): Including data from up to 4 RCTs and eight additional RCTs published since the pivotal review (Jimenez 2007/good quality; Mehr-Aein 2007/good quality: Elwatidy 2008/fair quality: Sadeghi 2007/good quality: Wong 2008/good quality: Wong 2008/good quality: Chen 2008/fair quality: Chen 2008/fair quality: Chen 2009/fair quality: Wong 2008/good quality: Chen 2008/fair quality: Wong 2008/good quality: Wong 2008/good quality: Chen 2008/fair quality: Studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or Level I on or two Level III studies with a low risk of bias or Level I on or two Level III studies with a low risk of bias or Level I on or two Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias
There is one pivotal Level I study (Henry 2007/good quality) including data from up to 4 RCTs and eight additional RCTs published since the pivotal review (Jimenez 2007/good quality; Later 2009/good quality; Mehr-Aein 2007/good quality; Elwatidy 2008/fair quality; Sadeghi 2007/good quality; Wong 2008/good quality; Chen 2008/fair quality; Choi 2009/fair quality). A One or more level I studies with a low risk of bias or several level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias or Level I or II studies with a high risk of bias or Level I or II studies with a moderate risk of bias or Level I or II studies with a high risk of bias or Level I or II studies with a high risk of bias or Level I or II studies with a high risk of bias or Level I or II studies with a high risk of bias or Level I or II studies with a high risk of bias or Level I or II studies with a high risk of bias or Level I or II studies with a high risk of bias or Level I or II studies with a low risk of bias or Level I or II studies with a low risk of bias or Level I or II studies with a low risk of bias or Level I or II studies with a low risk of bias or Level I or II studies with a low risk of bias or Level I or II studies with a low risk of bias or Level I or II studies with a low risk of bias or Level I or II studies with a low risk of bias or Level I or II studies with a low risk of bias or Level I or II studies with a low risk of bias or Level I or II studies with a low risk of bias or Level I or II studies with a low risk of bias or Level I or II o
RCTs published since the pivotal review (Jimenez 2007/good quality; Later 2009/good quality; Mehr-Aein 2007/good quality; Elwatidy 2008/fair quality; Sadeghi 2007/good quality; Wong 2008/good quality; Chen 2008/fair quality; Choi 2009/fair quality). B One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias or Level IV studies or Level I to III studies/SRs with a high risk of bias 2. Consistency (If only one study was available, rank this component as 'not applicable') Some inconsistency between SR and individual RCTs. Consistency of individual studies within meta-analyses described below. Pivotal evidence – Henry 2007
2007/good quality; Elwatidy 2008/fair quality; Sadeghi 2007/good quality; Wong 2008/good quality; Chen 2008/fair quality; Choi 2009/fair quality; Choi 2008/fair quality; Choi 2008/fair quality; Choi 2008/fair quality; Choi 2008/fair quality; Sadeghi 2007/good quality; Choi 2008/fair quality; C
quality; Choi 2009/fair quality). C One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias D Level IV studies or Level I to III studies/SRs with a high risk of bias 2. Consistency (If only one study was available, rank this component as 'not applicable') Some inconsistency between SR and individual RCTs. Consistency of individual studies within meta-analyses described below. Pivotal evidence – Henry 2007 C One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias A All studies consistent B Most studies consistent and inconsistency can be explained
2. Consistency (If only one study was available, rank this component as 'not applicable') Some inconsistency between SR and individual RCTs. Consistency of individual studies within meta-analyses described below. Pivotal evidence – Henry 2007 All studies consistent B Most studies consistent and inconsistency can be explained
Some inconsistency between SR and individual RCTs. Consistency of individual studies within meta-analyses described below. Pivotal evidence – Henry 2007 All studies consistent B Most studies consistent and inconsistency can be explained
described below. Pivotal evidence – Henry 2007 B Most studies consistent and inconsistency can be explained
Pivotal evidence – Henry 2007
i ivolai evidence – neni y 2007
No significant heterogeneity (see note) in analyses. C Some inconsistency, reflecting genuine uncertainty around question
Supportive evidence D Evidence is inconsistent
No difference between tranexamic acid and no tranexamic acid in most supportive studies although in some cases the length of stay is slightly longer with tranexamic acid. NA Not applicable (one study only)
3. Clinical impact (Indicate in the space below if the study results varied according to some unknown factor (not simply study quality or sample size) and thus the clinical impact of the intervention could not be determined)
Pivotal evidence – Henry 2007 A Very large
Any surgery (days) – WMD -0.30 (-0.71, 0.10); 4 RCTs (N=176) Cardiac surgery (days) – WMD -0.23 (-0.67, 0.21); 2 RCTs (N=116) B Substantial
Supportive evidence – see Summary Table POQ3.l8b.S5
D No difference
4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?)
The evidence is generalisable to an adult population who are undergoing planned surgery. There is evidence A Evidence directly generalisable to target population
available for a number of different surgery types. B Evidence directly generalisable to target population with some caveats
C Evidence not directly generalisable to the target population but could be sensibly applied
D Evidence not directly generalisable to target population and hard to judge whether it is sensible to apply
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of health services/delivery of care and cultural factors?)
Hospital setting. A Evidence directly applicable to Australian healthcare context
The pivotal review included studies from a wide range of countries. Included RCTs were from a number of different countries including several from the Middle East and Asia. B Evidence applicable to Australian healthcare context with few caveats
C Evidence probably applicable to Australian healthcare context with some caveats
D Evidence not applicable to Australian healthcare context

The Advisory Committee on Prescription Medicines (ACPM) has recently recommended approval of tranexamic acid injection "for the reduction of peri- and postoperative blood loss and of the need for blood transfusion in adult patients undergoing cardiac surgery or total knee or hip arthroplasty; and paediatric patients undergoing cardiac surgery". Tranexamic acid tablets are approved in Australia for a number of indications including haemostatic, hereditary angioedema, short-term treatment of traumatic hyphaema, patients with coagulopathies undergoing minor surgery, and menorrhagia.

Consistency changed from A to D as some results went slightly in the opposite direction.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (good quality) and eight additional RCTs.
2. Consistency	D	There was some inconsistency between studies with most studies showing no difference (although the direct was slightly different in some) and one study showing a significant difference in favour of tranexamic acid.
3. Clinical impact	D	There was no difference in length of hospital stay.
4. Generalisability	С	The results are generalisable to a surgical population.
5. Applicability	В	There were studies from a number of countries including Spain, The Netherlands, Iran, Saudi Arabia, Canada, Taiwan and Hong Kong. Likely to be applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, intravenous tranexamic acid therapy does not appear to affect hospital length of stay compared with no therapy.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: RCT, randomised controlled trial; SR, systematic review; WMD, weighted mean difference.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.l8b.S5 Characteristics and results of studies examining the effect of <u>tranexamic acid</u> on <u>hospital length of stay</u>

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size	Surgical procedure	Location			Risk estimate (95% CI) or TXA (mean) vs control (mean)	Significance P-value	
ADULT POPULATION	IV TRANEXAMIC A	CID							
Any surgery									
Henry (2007)	Level I Good	4 RCTs N=176	Adult patients undergoing any surgery	Hospital – planned surgery Various countries ^c	Tranexamic acid (IV) vs no tranexamic acid	Hospital length of stay (days)	WMD -0.30 (-0.71, 0.10)	No difference 0.14	None Phet=0.66 (I ² =0%)
Cardiac surgery									
Henry (2007)	Level I Good	2 RCTs N=116	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	Tranexamic acid (IV) vs no tranexamic acid	Hospital length of stay (days)	WMD -0.23 (-0.67, 0.21)	No difference 0.31	None Phet=0.64 (l ² =0%)
Jimenez (2007)	Level II Good	1 RCT N=50	Adult patients undergoing CPB surgery	Hospital – planned surgery Spain	Tranexamic acid (IV) vs placebo	Hospital length of stay (days)	4.5 vs 4	No difference 0.34	NA
Later (2009)	Level II Good	1 RCT N=202	Adult patients undergoing first-time, non-complex cardiac surgery with CPB	Hospital – planned surgery The Netherlands	Tranexamic acid (IV) vs placebo	Hospital length of stay (days)	9.4 vs 8.5	No difference 0.43	NA
Mehr-Aein (2007)	Level II Good	1 RCT N=66	Adult patients undergoing primary CABG surgery	Hospital – planned surgery Iran	Tranexamic acid (IV) vs placebo	Hospital length of stay (days)	4.8 vs 4.8	No difference 0.09	NA
Orthopaedic surge	ery	П		•	1		1	-1	1
Elwatidy (2008)	Level II Fair	1 RCT N=64	Adult and paediatric patients undergoing spine surgery	Hospital – planned surgery Saudi Arabia	Tranexamic acid (IV) vs placebo	Hospital length of stay (days)	8.5 vs 10.7	No difference 0.21	NA
Sadeghi (2007)	Level II Good	1 RCT N=67	Adult patients undergoing hip fracture surgery	Hospital – planned surgery Iran	Tranexamic acid (IV) vs placebo	Hospital length of stay (days)	4.3 vs 5.8	Favours tranexamic acid <0.05	NA
Wong (2008)	Level II Good	1 RCT N=147	Adult patients undergoing spinal fusion surgery	Hospital – planned surgery Canada	Tranexamic acid (IV) vs placebo	Hospital length of stay (days)	9.2 vs 8.5	No difference 0.38	NA

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	ea sample size Surgical procedure Location		Risk estimate (95% CI) or TXA (mean) vs control (mean)	Significance P-value				
Other surgery	•	•						-	1
Chen (2008)	Level II Fair	1 RCT N=55	Adults undergoing <u>head and</u> neck surgery	Hospital – planned surgery Taiwan	Tranexamic acid (IV) vs placebo	Hospital length of stay (days)	4.8 vs 5.3	No difference 0.087	NA
Choi (2009)	Level II Fair	RCT N=61	Adults undergoing orthognathic surgery	Hospital – planned surgery Hong Kong	Tranexamic acid (IV) vs placebo	Hospital length of stay (days)	7.2 vs 7.5	No difference 0.32	NA
ADULT POPULATION	TOPICAL TRANEX	AMIC ACID							
Cardiac surgery									
Fawzy (2009)	Level II Good	1 RCT N=38	Adult patients undergoing primary isolated CABG surgery	Hospital – planned surgery Saudi Arabia	Tranexamic acid (topical) vs placebo	Hospital length of stay (days)	7.5 vs 7.8	No difference 0.68	NA
ADULT POPULATION	ORAL TRANEXAM	C ACID							
Liver surgery									
Gurusamy (2009)	Level I/II Good/Fair	1 RCT N=214	Adult patients undergoing liver resection	Hospital – planned surgery China	Tranexamic acid (oral) vs placebo	Hospital length of stay (days)	8 vs 9	No difference 0.34	NA

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: ASA, acetylsalicylic acid; CABG, coronary artery bypass graft; CI, confidence interval; CPB, cardiopulmonary bypass; het, heterogeneity; IV, intravenous; NA, not applicable; NR, not reported; OR, odds ratio; RCT, randomised controlled trial; RR, risk ratio.

^a Where only one RCT is available in a systematic review, the level of evidence has been downgraded to Level II.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between tranexamic acid and no tranexamic acid.

Recommendation(s) for administration of tranexamic acid

RECOMMENDATION What recommendation(s) does the guideline development group draw from this evidence? Use action statements where possible.	GRADE	RELEVANT EVIDENCE TABLI
In adult patients undergoing cardiac surgery, the use of intravenous tranexamic acid is recommended.	А	PO3.l8b.P1, PO3.l8b.P2, PO3.l8b.P3, PO3.l8b.P5
In adult patients undergoing non-cardiac surgery, if substantial blood loss is anticipated, the use of intravenous tranexamic acid is recommended.	В	PO3.l8b.P1, PO3.l8b.P2, PO3.l8b.P3, PO3.l8b.P5
IMPLEMENTATION OF RECOMMENDATION Please indicate yes or no to the following questions. Where the answer is yes please provide explanatory information about this. This information will be used to develop the implementation plan for the guidelines.		
Will this recommendation result in changes in usual care? Increased use of tranexamic acid.		YES NO
Are there any resource implications associated with implementing this recommendation?		YES NO
Drug cost (albeit modest).		
Will the implementation of this recommendation require changes in the way care is currently organised?		YES NO
Are the guideline development group aware of any barriers to the implementation of this recommendation		YES NO
Potential resistance.		
What could help to facilitate implementation of the recommendation?		YES NO
Education; promotion of PO guideline.		

Intervention 8 – Administration of antifibrinolytics & DDAVP: ε-aminocaproic acid

Key question(s): In patients undergoing surgery, what is the effect of administration of ε -amino	capr	oic acid on transfusion incidence?	Evidence table refa: POQ3.18c.P1			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)					
There is one pivotal Level I study (Henry 2007/good quality) which includes data from up to 14 RCTs and two	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias				
supportive Level I studies (Brown 2007/fair quality; Kagoma 2009/good quality).	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	S			
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Results consistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta-	Α	All studies consistent				
analyses described below. Pivotal evidence – Henry 2007	В	Most studies consistent and inconsistency can be explained				
Moderate to substantial heterogeneity between studies. May be due to different surgery types assessed. Only one	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
study available for liver surgery.	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u> determined)	nknow	<u>n</u> factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be			
Pivotal evidence – Henry 2007	Α	Very large				
Any surgery - 33.3% vs 45.1%; RR 0.75 (0.58, 0.96); 14 RCTs (N=801)	В	Substantial (cardiac)				
Cardiac surgery - 26.2% vs 39.8%; RR 0.65 (0.47, 0.91); 10 RCTs (N=597) Orthopaedic surgery - 33.9% vs 36.5%; RR 0.96 (0.61, 1.50); 3 RCTs (N=122)	С	Moderate				
Liver surgery – 85.7% vs 92.5%; RR 0.93 (0.80, 1.08); 1 RCT (N=82) Supportive evidence –see Summary Table POQ3.l8c.P1	D	Underpowered (non-cardiac)				
4. Generalisability (How well does the body of evidence match the population and clinical sea	ttinas l	being targeted by the Guideline?)				
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were	A	Evidence directly generalisable to target (cardiac) population				
included in the overall analysis although studies were predominantly in cardiac surgery. There were also subgroup analyses of patients who had undergone surgery with/without a transfusion protocol.	В	Evidence directly generalisable to target population with some cav	/eats			
analyses of patients who had undergone surgery withwithout a transfusion protocol.	С	Evidence not directly generalisable to the target (non-cardiac) pop	pulation but could be sensibly applied			
	D	Evidence not directly generalisable to target population and hard tapply	o judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of hea	Ith services/delivery of care and cultural factors?)				
Hospital setting.	Α	Evidence directly applicable to Australian healthcare context				
The pivotal review states that studies were conducted in a wide range of countries.	В	Evidence applicable to Australian healthcare context with few caveats				
	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
	D	Evidence not applicable to Australian healthcare context				
		·				

E-aminocaproic acid is a synthetic derivative of the amino acid lysine. It is not currently TGA-approved for use in Australia.

The Henry (pivotal) review assessed quality and performed a subgroup analysis of transfusion incidence for all surgery types based on the rating (A,B or C) of treatment allocation. The analysis showed no substantial difference in the results between studies rated A, B or C, with the exception that potentially poorer studies showed less effect.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Cardiac	Other	Description
1. Evidence base	Α	Α	There is one pivotal Level I study (good quality) and two supportive Level I studies.
2. Consistency	В	В	Most studies were reasonably consistent with differences mostly related to magnitude of effect rather than direction of effect. Differences may be related to surgery type.
3. Clinical impact	В		There was a significant difference for cardiac surgery only. There was no difference for non-cardiac surgery (predominantly orthopaedic surgery) but includes few studies so may be underpowered to detect a difference.
4. Generalisability	Α	С	The results are generalisable to an adult surgical population; in particular those undergoing cardiac surgery
5. Applicability	В	В	Studies were conducted in a range of countries. Likely to be applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, intravenous ε-aminocaproic acid therapy reduces the incidence of allogeneic blood transfusion compared with no therapy. In adult patients undergoing non-cardiac surgery in which substantial blood loss is anticipated, the effect of intravenous ε-aminocaproic acid therapy on the incidence of allogeneic transfusion, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI, confidence interval, RBC, red blood cell: SR, systematic review;

^a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission by Cochrane ratings defined as follows: Grade A, adequate allocation concealment: Grade B, uncertain allocation concealment: Grade C, inadequate allocation concealment.

POQ3.l8c.P1 Characteristics and results of studies examining the effect of ε-aminocaproic acid on transfusion incidence.

Study Level of		No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size included in analysis	procedure	Location			Risk estimate (95% CI)	Significance P-value	
ADULT POPULATIO	N/ IV E -AMINOCAPR	OIC ACID							
Any surgery									
Henry (2007)	Level I Good	14 RCTs N=801	Adult patients undergoing any surgery	Hospital – planned surgery Various countries ^c	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Transfusion incidence (allogeneic blood)	RR 0.75 (0.58, 0.96)	Favours ε- aminocaproic acid 0.023	Moderate Phet=0.03 (I ² =47%)
Henry (2007)	Level I Good	13 RCTs N=771	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	E-aminocaproic acid (IV) vs no ε- aminocaproic acid with transfusion protocol	Transfusion incidence (allogeneic blood)	RR 0.73 (0.56, 0.95)	Favours ε- aminocaproic acid 0.019	Substantial Phet=0.02 (l²=52%)
	Level I/II Good/Fair	1 RCT N=30	Adult patients undergoing any surgery (orthopaedic surgery only)	Hospital – planned surgery Various countries	E-aminocaproic acid (IV) vs no ε- aminocaproic acid without transfusion protocol	Transfusion incidence (allogeneic blood)	RR 1.33 (0.36, 4.97)	No difference 0.67	NA
Henry (2007)	Level I Good Rating Ad	3 RCTs N=218	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Transfusion incidence (allogeneic blood)	RR 0.68 (0.44, 1.04)	No difference 0.076	Moderate Phet=0.25 (I ² =29%)
	Level I Good Rating Bd	9 RCTs N=455	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Transfusion incidence (allogeneic blood)	RR 0.68 (0.46 1.03)	No difference 0.068	Moderate Phet=0.13 (I ² =36%)
	Level I Good Rating Cd	2 RCTs N=128	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Transfusion incidence (allogeneic blood)	RR 0.93 (0.81, 1.08)	No difference 0.35	None Phet=0.72 (I ² =0%)
Cardiac surgery									
Henry (2007)	Level I Good	10 RCTs N=597	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Transfusion incidence (allogeneic blood)	RR 0.65 (0.47, 0.91)	Favours ε- aminocaproic acid 0.011	Moderate Phet=0.11 (I ² =38%)
Brown (2007)	Level I Fair	10 RCTs N=628	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	E-aminocaproic acid (IV) vs placebo	Transfusion incidence (pRBCs)	RR 0.63 (0.44, 0.90)	Favours ε- aminocaproic acid 0.010	NR

,	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size included in analysis	procedure	Location			Risk estimate (95% CI)	Significance P-value	
Orthopaedic surg	ery								
Henry (2007)	Level I Good	3 RCTs N=122	Adult patients undergoing orthopaedic surgery	Hospital – planned surgery Various countries	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Transfusion incidence (allogeneic blood)	RR 0.96 (0.61, 1.50)	No difference 0.85	None Phet=0.64 (l²=0%)
Kagoma (2009)	Level I Good	3 RCTs N=180	Adult patients undergoing orthopaedic surgery	Hospital – planned surgery Unknown	E-aminocaproic acid (IV) vs placebo	Transfusion incidence	RR 0.64 (0.21, 1.93)	No difference NR	NR
Liver surgery		•			•				
Henry (2007)	Level I/II Good/Fair	1 RCT N=82	Adult patients undergoing <u>liver</u> surgery	Hospital – planned surgery Various countries	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Transfusion incidence (allogeneic blood)	RR 0.93 (0.80, 1.08)	No difference 0.33	NA
PAEDIATRIC POPULA	TION/IV E-AMINOC	APROIC ACID							
Orthopaedic surg	ery								
Tzortzopoulou (2008)	Level I/II Good/Good	1 RCT N=36	Paediatric patients undergoing scoliosis surgery	Hospital – planned surgery Unknown	E-aminocaproic acid (IV) vs placebo	Transfusion incidence	RR 1.04 (0.69, 1.57)	No difference 0.84	NA

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: CI, confidence interval; *het*, heterogeneity; IV, intravenous; NA, not applicable; NR, not reported; pRBC, packed red blood cell; RCT, randomised controlled trial; RR, risk ratio.

a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between ε-aminocaproic acid and no ε-aminocaproic acid.

d Cochrane ratings defined as follows: Grade A, adequate allocation concealment; Grade B, uncertain allocation concealment; Grade C, inadequate allocation concealment.

Key question(s):			Evidence table refa:			
In patients undergoing surgery, what is the effect of administration of ε-amino	ocapr	oic acid on transfusion volume?	POQ3.I8c.P2			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studie.	s)					
There is one pivotal Level I study (Henry 2007/good quality) which includes data from up to 4 RCTs and one additional RCT (Berenholtz 2009/good quality) in major spine surgery (conducted in the US).		One or more level I studies with a low risk of bias or several level	Il studies with a low risk of bias			
additional No.1 (Berefilloliz 2007/g000 quality) in major spille surgery (conducted in the 0.5).	В	One or two Level II studies with a low risk of bias or SR/several Level	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Results somewhat inconsistent. Likely to be due to differences in reporting of results in pivotal review and supportive studies (ie, includes all patients or transfused patients only; different blood products assessed; different transfusion	e A	All studies consistent				
units included such as units or mL).	В	Most studies consistent and inconsistency can be explained				
Pivotal evidence – Henry 2007 Significant heterogeneity in analysis of transfusion volume for all patients but not for transfusion volume including	С	Some inconsistency, reflecting genuine uncertainty around question				
only transfused patients.	D	Evidence is inconsistent				
Supportive evidence –Mix of results showing effect favouring ε-ACA and no difference.	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u> determined)	inknow	<u>n</u> factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be			
Pivotal evidence – Henry 2007	Α	Very large				
Any surgery (all patients; units) – WMD -1.77 (-2.59, -0.95); 4 RCTs (N=198) Any surgery (transfused patients; units) – WMD 0.22 (-0.34, 0.79); 3 RCTs (N=119)	В	Substantial				
Supportive evidence – (see Summary Table POQ3.l8d.P2)	С	Moderate				
	D	Inconsistent				
4. Generalisability (How well does the body of evidence match the population and clinical se	ttings b					
The evidence is generalisable to an adult population who are undergoing planned surgery. The majority of studies were in cardiac or orthopaedic surgery.	Α	Evidence directly generalisable to target population				
note in caldade of orthopacode sangery.	В	Evidence directly generalisable to target population with some caveats				
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of hear	Ith services/delivery of care and cultural factors?)				
Hospital setting. The pivotal review states that studies were conducted in a range of countries. The additional RCT was conducted in	Α	Evidence directly applicable to Australian healthcare context				
the US.	В	Evidence applicable to Australian healthcare context with few caveats				
	С	Evidence probably applicable to Australian healthcare context with some caveats				
	D	Evidence not applicable to Australian healthcare context				

E-aminocaproic acid is a synthetic derivative of the amino acid lysine. It is not currently listed for use in Australia.

Generalisability rated C as most evidence in a restricted surgical population but applied to general surgical population.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (good quality) and one additional RCT.
2. Consistency	С	There are some inconsistencies depending on the denominator used (all patients vs transfused patients), surgery type and blood products.
3. Clinical impact	D	There was generally significantly less transfusion when all patients were included in the analysis; however, there was no significant difference when only transfused patients were considered.
4. Generalisability	С	The results are generalisable to a surgical population; in particular those undergoing cardiac and orthopaedic surgery.
5. Applicability	С	Studies were conducted in a number of countries. The additional RCT was conducted in the US. Possibly applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intravenous ε-aminocaproic acid therapy on volume of allogeneic blood transfusion compared with no therapy is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: ACA, aminocaproic acid: CI, confidence interval, RBC, red blood cell: SR, systematic review:

Primary outcomes: P1 = transfusion incidence. P2 = transfusion volume, P3 = blood loss. P4 = mortality, P5 = morbidity. P6 = quality of life

^a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.l8c.P2 Characteristics and results of studies examining the effect of ε-aminocaproic acid on transfusion volume

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size included in analysis	procedure	Location			Risk estimate (95% CI)	Significance P-value	
ADULT POPULATION/	IV E-AMINOCAPRO	DIC ACID							
Any surgery									
Henry (2007)	Level I Good	4 RCTs N=198	Adult patients undergoing any surgery (all patients)	Hospital – planned surgery Various countries ^c	ε -aminocaproic acid (IV) vs no ε- aminocaproic acid	Transfusion volume (units; allogeneic blood)	WMD -1.77 (-2.59, -0.95)	Favours ε- aminocaproic acid <0.001	Substantial Phet=0.02 (I ² =69%)
		3 RCTs N=119	Adult patients undergoing any surgery (transfused patients only)	Hospital – planned surgery Various countries	ε -aminocaproic acid (IV) vs no ε- aminocaproic acid	Transfusion volume (units; allogeneic blood)	WMD 0.22 (-0.34, 0.79)	No difference 0.44	None Phet=0.76 (I ² =0%)
Orthopaedic surge	ery		<u>. </u>						
Berenholtz (2009)	holtz (2009) Level II Good 1 RCT N=182 Adult patients undergoing major spine surgery (all patients) US Hospita surgery US	0 ,	urgery (IV) vs placebo to	Transfusion volume (units; total allogeneic RBC)	MD -1.00 (-2.47, 0.47) ^d	No difference 0.18 ^d	NA		
		US		Transfusion volume (units; postoperative RBC)	MD -0.80 (-1.48, -0.12) ^d	Favours ε- aminocaproic acid 0.02 ^d	NA		
						Transfusion volume (units; total FFP)	MD -0.70 (-2.17, 0.77) ^d	No difference 0.35 ^d	NA
						Transfusion volume (units; total platelets)	MD 0.00 (-1.17, 1.17) ^d	No difference	NA
						Transfusion volume (units; total all blood products)	MD -2.60 (-6.38, 1.18) ^d	No difference 0.18 ^d	NA
PAEDIATRIC POPULA	TION/IV E-AMINOC	APROIC ACID						1	
Cardiac surgery									
Schouten (2009)	Level I Good	3 RCTs N=410	Paediatric patients undergoing cardiac surgery	Hospital – planned surgery Unknown	ε -aminocaproic acid (IV) vs no ε- aminocaproic acid	Transfusion volume (mL/kg; <u>plasma</u>)	WMD -3 (-5, -1)	Favours ε- aminocaproic acid NR	None Phet=NR (I ² =20%)
Orthopaedic surge	ery	•			•			•	
Tzortzopoulou (2008)	Level I/II Good/Good	1 RCT N=87	Paediatric patients undergoing scoliosis surgery	Hospital – planned surgery Unknown	ε -aminocaproic acid (IV) vs placebo	Transfusion volume (mL)	WMD -245 (-481, -8.97)	Favours ε- aminocaproic acid 0.042	NA

Note: Studies/analyses providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies/analyses provide supportive evidence.

Abbreviations: CI, confidence interval; FFP, fresh frozen plasma; het, heterogeneity; IV, intravenous; MD, mean difference; NA, not applicable; NR, not reported; RBC, red blood cell; RCT, randomised controlled trial; SMD, standardised mean difference; US, United States of America; WMD; weighted mean difference.

a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

Appendix D: Evidence matrixes – Intervention 8	(Administration of	e-aminocaproic acid

c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between ε-aminocaproic acid and no εaminocaproic acid. d Post-hoc calculation for this summary.

Key question(s):			Evidence table refa:
In patients undergoing surgery, what is the effect of administration of ε-amino	capr	oic acid on blood loss?	POQ3.I8c.P3
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)		
There is one pivotal Level I study (Henry 2007/good quality) including data from up to 12 RCTs, three supportive Level I studies (Brown 2007/fair quality; Kagoma 2009/good quality; McIlroy 2009/good quality [which combined	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias
data for TXA and ACA) and two additional RCTs (Gharebaghian 2006/fair quality; Berenholtz 2009/good quality) published since the pivotal review.	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	
2. Consistency (If only one study was available, rank this component as 'not applicable')			
Results consistent between pivotal and supportive meta-analyses and RCTs. Consistency of individual studies within meta-analyses described below.	Α	All studies consistent	
Pivotal evidence – Henry 2007	В	Most studies consistent and inconsistency can be explained (card	iac)
Moderate to substantial heterogeneity (see note) in main analyses. No heterogeneity in analyses by surgery type. Some inconsistency between non-cardiac surgery types.	С	Some inconsistency, reflecting genuine uncertainty around question	on (orthopaedic)
Some medialition of between non earline surgery types.	D	Evidence is inconsistent	
	NA	Not applicable (one study only)	
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u>	nknow	rn factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be
Pivotal evidence – Henry 2007	Α	Very large	
Any surgery (intraoperative blood loss) – WMD -142 (-285, 0.92); 4 RCTs (N=171) Any surgery (postoperative blood loss) – WMD -202 (-274, -131); 12 RCTs (N=940)	В	Substantial (cardiac)	
Cardiac surgery (intraoperative blood loss) – WMD -214 (-310, -117); 2 RCTs (N=79) Cardiac surgery (postoperative blood loss) – WMD -196 (-272, -121); 11 RCTs (N=894)	С	Moderate (orthopaedic)	
Orthopaedic surgery (postoperative blood loss) – WMD -176 (-272, -121), 11 RC1s (N=674)	D	Slight/Restricted	
4. Generalisability (How well does the body of evidence match the population and clinical se	ttings I	being targeted by the Guideline?)	
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were included in the overall analysis although studies were predominantly in cardiac and orthopaedic surgery. There was	Α	Evidence directly generalisable to target population (cardiac)	
also evidence relating to adults undergoing cardiac surgery who had received aspirin.	В	Evidence directly generalisable to target population with some cav	eats (orthopaedic)
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of hea	Ith services/delivery of care and cultural factors?)	
Hospital setting. The pivotal review states that studies were conducted in a wide range of countries.	Α	Evidence directly applicable to Australian healthcare context	
Additional RCTs were conducted in Iran and the US.	В	Evidence applicable to Australian healthcare context with few cave	eats (cardiac surgery)
	С	Evidence probably applicable to Australian healthcare context with	n some caveats (orthopaedic surgery)
	D	Evidence not applicable to Australian healthcare context	
			·

E-aminocaproic acid is a synthetic derivative of the amino acid lysine. It is not currently TGA-approved for use in Australia.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Cardiac	Ortho	Description
1. Evidence base	Α	Α	There is one pivotal Level I study (good quality), three supportive Level I studies and two additional RCTs.
2. Consistency	В	С	Results were generally consistent. Substantial heterogeneity in any surgery analysis may be due to surgery types.
3. Clinical impact	В	С	There was generally significantly less blood loss with ACA, particularly postoperatively. This was strongest in cardiac surgery.
4. Generalisability	Α	В	Results generalisable to an adult surgical population; in particular those undergoing cardiac and orthopaedic surgery.
5. Applicability	В	С	Studies were conducted in a number of countries. The additional RCTs were conducted in the US and Iran. Possibly applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, intravenous ε-aminocaproic acid therapy reduces blood loss compared with no therapy.

In adult patients undergoing major orthopaedic surgery, intravenous ε-aminocaproic acid therapy may reduce blood loss compared with no therapy.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%.

Abbreviations: ACA, ϵ -aminocaproic acid; CI, confidence interval, RBC, red blood cell; RCT, randomised controlled trial; TXA, tranexamic acid.

a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

POQ3.l8c.P3 Characteristics and results of studies examining the effect of ε-aminocaproic acid on blood loss

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or ε-ACA (mean) vs control (mean)	Significance P-value	
ADULT POPULATIO	N/IV E-AMINOCAPR	OIC ACID							
Any surgery									
Henry (2007)	Level I Good	4 RCTs N=171	Adult patients undergoing any surgery	Hospital – planned surgery Various countries ^c	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Intraoperative blood loss (mL)	WMD -142 (-285, 0.92)	No difference 0.051	Moderate Phet=0.19 (I ² =37%)
		12 RCTs N=940	Adult patients undergoing any surgery	Hospital – planned surgery Various countries	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Postoperative blood loss (mL)	WMD -202 (-274, -131)	Favours ε- aminocaproic acid <0.001	Substantial Phet<0.001 (I ² =89%)
Cardiac surgery								·	
Henry (2007)	Level I Good	2 RCTs N=79	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Intraoperative blood loss (mL)	WMD -214 (-310, -117)	Favours ε- aminocaproic acid <0.001	None Phet=0.73 (I ² =0%)
		11 RCTs N=894	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Postoperative blood loss (mL)	WMD -196 (-272, -121)	Favours ε- aminocaproic acid <0.001	Substantial Phet<0.001 (I ² =89%)
Brown (2007)	Level I Fair	3 RCTs N=144	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	E-aminocaproic acid (IV) vs placebo	Total blood loss (mL)	WMD -240 (-341, -140)	Favours ε- aminocaproic acid <0.001	NR
McIlroy(2009)	Level I Good	3 RCTs N=259	Adult patients receiving ASA undergoing cardiac surgery	Hospital – planned surgery Unknown	Lysine analoguesh (IV) vs placebo	Postoperative chest tube blood loss (mL)	WMD -189 (-287, -91)	Favours lysine analogues <0.001	Substantial Phet=0.05 (l²=67%)

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results	Heterogeneity⁵				
	evidence ^a Quality	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or ε-ACA (mean) vs control (mean)	Significance P-value				
Gharebaghian (2006)	Level II Fair	1 RCT N=60	Adult patients undergoing major CABG surgery	Hospital – planned surgery Iran	E-aminocaproic acid (IV) pre-incision regimend vs placebo	Chest tube blood loss at <u>6 hrs</u> (mL)	~ 300 vs ~ 600	Favours ε- aminocaproic acid <0.05	NA			
						Chest tube blood loss at 12 hrs (mL)	~500 vs ~650	No difference >0.05	NA			
						Chest tube blood loss at removal (mL)	~1000 vs ~2000	Favours ε- aminocaproic acid <0.05	NA			
					Chest tube blood loss at <u>6 hrs</u> (mL)	~ 300 vs ~ 600	Favours ε- aminocaproic acid <0.05	NA				
						Chest tube blood loss at 12 hrs (mL)	~500 vs ~650	No difference >0.05	NA			
									Chest tube blood loss at removal (mL)	~800 vs ~2000	Favours ε- aminocaproic acid <0.05	NA
Orthopaedic surge	ery											
Henry (2007)	Level I Good	2RCTs N=92	Adult patients undergoing orthopaedic surgery	Hospital – planned surgery Various countries	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Total blood loss (mL)	WMD -300 (-523, -77)	Favours ε- aminocaproic acid 0.0084	None Phet=0.39 (I ² =0%)			
						Intraoperative blood loss (mL)	WMD 10.9 (-260, 282)	No difference 0.94	None Phet=0.26 (l ² =22%)			
	Level I/II Good/Fair	1 RCT N=46	Adult patients undergoing orthopaedic surgery	Hospital – planned surgery Various countries	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Postoperative blood loss (mL)	WMD -276 (-449, -103)	Favours ε- aminocaproic acid 0.0017	NA			
Kagoma (2009)	Level I Good	3 RCTs N=141	Adults undergoing total knee or hip replacement	Hospital – planned surgery Unknown	E-aminocaproic acid (IV) vs no ε-aminocaproic acid	Total bleeding ^f (mL)	WMD -331 (-544, -118)	Favours ε- aminocaproic acid P<0.05	NR			
Berenholtz (2009)	Level II Good	1 RCT N=182	Adult patients undergoing major spine surgery	Hospital – planned surgery US	E-aminocaproic acid (IV) vs placebo	Intraoperative blood loss (mL)	MD -335 (-990, 320) ⁹	No difference 0.32 g	NA			
						Post-surgical to POD 1 blood loss (mL)	MD -430 (-1121, 261) ^g	No difference 0.22 g	NA			

Study	Level of No. of trials /		Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size included in analysis	procedure	Location			Risk estimate (95% CI) or ε-ACA (mean) vs control (mean)	Significance P-value	
PAEDIATRIC POPULA	TION/IV E-AMINOC	APROIC ACID							
Orthopaedic surge	ery								
Schouten (2009)	Level I/II Good/Good	1 RCT ⁱ N=36	Paediatric patients undergoing scoliosis surgery	Hospital – planned surgery Unknown	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Blood loss (mL/kg/day)	WMD -59 (-262, 144)	No difference NR	NA
Tzortzopoulou (2008)	Level I/II Good/Good	1 RCT [†] N=36	Paediatric patients undergoing scoliosis surgery	Hospital – planned surgery Unknown	E-aminocaproic acid (IV) vs placebo	Total blood loss (mL)	WMD -325 (-587, -63)	Favours ε- aminocaproic acid 0.015	NA
ADULT POPULATION/	TOPICAL E-AMINO	CAPROIC ACID							
Other surgery									
Athanasiadis (2007)	Level II Fair	1 RCT N=20	Adult patients undergoing endoscopic sinus surgery	Hospital – planned surgery Australia	E-aminocaproic acid (topical) vs placebo	Bleeding grading scales at 0, 2, 4, 6, 8 and 10 mins	NR	No difference NR	NA

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: ASA, acetylsalicylic acid; CI, confidence interval; het, heterogeneity; IV, intravenous; mL, millilitres; NA, not applicable; NR, not reported; RCT, randomised controlled trial; SMD, standardised mean difference; WMD, weighted mean difference.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Pheto 0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

The pre-incision group received 150 mg/kg ε-ACA over 10 mins as pre-incision bolus, followed by 15 mg/kg/hr as post-incision infusion and normal saline for post-heparin and 15 mg/kg/hr ε-ACA as 3 min following heparin to the end of CPB infusion.

e The post-heparin group received normal saline in pre-incision and post-incision and post-incision and ε-ACA comprising 150 mg/kg over 10 mins after heparin injection followed by 15 mg/kg/hr ε-ACA from 3 mins following heparin injection to the end of CPB.

^rTotal bleeding measured intraoperatively by weighing surgical sponges, postoperatively through drainage or perioperatively through the haemoglobin balance method which measures blood loss through comparison or pre- and postoperative haemoglobin concentrations (haematoma volumes as well as hidden or internal blood loss were excluded).

^g Calculated post-hoc for this Guideline.

h Includes ε-aminocaproic acid (2 RCTs) and tranexamic acid (1 RCT).

¹ These two reviews included the same study (Florentino 2004).⁵

Wormald grading scale and Boezaart grading scale.

⁵ Florentino-Pineda et al (2004) The effect of amicar on perioperative blood loss in idiopathic scoliosis: the results of a prospective, randomized double-blind study. Spine 29: 233-238.

Key question(s):			Evidence table refa:	
In patients undergoing surgery, what is the effect of administration of ε-amino	ocapr	oic acid on mortality?	POQ3.I8c.P4	
1. Evidence base (number of studies, level of evidence and risk of bias in the included studie	s)			
There is one pivotal Level I study (Henry 2007/good quality) which includes data from up to 5 RCTs, one supportive Level I study (Brown 2007/fair quality) and one additional RCT (Berenholtz 2009/good quality) in major spine	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias	
surgery conducted in the US.	В	One or two Level II studies with a low risk of bias or SR/several Lev	rel III studies with a low risk of bias	
	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias	
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias		
2. Consistency (If only one study was available, rank this component as 'not applicable')				
Results consistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta- analyses described below.	Α	All studies consistent in finding no significant difference		
Pivotal evidence – Henry 2007	В	Most studies consistent and inconsistency can be explained		
No significant heterogeneity (see note) in main analysis and cardiac surgery analysis.	С	Some inconsistency, reflecting genuine uncertainty around question	on	
	D	Evidence is inconsistent		
	NA	Not applicable (one study only)		
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u>	nknow	<u>n</u> factor (not simply study quality or sample size) and thus the clinical impa	ct of the intervention could not be	
Pivotal evidence – Henry 2007	Α	Very large		
Any surgery - 2.6% vs 1.9%; RR 1.17 (0.47, 2.93); 5 RCTs (N=714) Cardiac surgery - 2.0% vs 0.9%; RR 1.65 (0.50, 5.43); 4 RCTs (N=632)	В	Substantial		
Supportive evidence – Brown 2007	С	Moderate		
Cardiac surgery – absolute risk not reported; RR 1.82 (0.55, 5.98); 6 RCTs (N=735) See also Summary Table POQ3.l8c.P4	D	Underpowered		
4. Generalisability (How well does the body of evidence match the population and clinical se	ttings b	peing targeted by the Guideline?)		
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were included in the overall analysis although studies were predominantly in cardiac surgery.	Α	Evidence directly generalisable to target population		
included in the overall aliasysis although studies were predominantly in cardiac surgery.	В	Evidence directly generalisable to target population with some cav	eats	
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied	
	D	Evidence not directly generalisable to target population and hard to	judge whether it is sensible to	
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of hear	th services/delivery of care and cultural factors?)		
Hospital setting. The pivotal review states that studies were conducted in a range of countries. An additional RCT was conducted in	Α	Evidence directly applicable to Australian healthcare context		
the US.	В	Evidence applicable to Australian healthcare context with few cave	ats	
	С	Evidence probably applicable to Australian healthcare context with	some caveats	
	D	Evidence not applicable to Australian healthcare context		

E-aminocaproic acid is a synthetic derivative of the amino acid lysine. It is not currently TGA-approved for use in Australia.

Generalisability rated C as most evidence in a restricted surgical population but applied to general surgical population.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (good quality), one supportive Level I study and one additional RCT.
2. Consistency	Α	There was no heterogeneity in the pivotal Level I study. All studies showed no significant difference.
3. Clinical impact	D	Results show a slightly increased risk with no significant difference (7/346 versus 3/326 in cardiac surgery) but studies likely underpowered to detect a difference in mortality.
4. Generalisability	С	The results are generalisable to an adult surgical population; in particular those undergoing cardiac surgery
5. Applicability	В	Studies were conducted in a range of countries. The additional RCT was conducted in the US. Likely to be applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery, the effect of intravenous ε-aminocaproic acid therapy on mortality compared with no therapy is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI, confidence interval, RBC, red blood cell; SR, systematic review;

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.l8c.P4 Characteristics and results of studies examining the effect of ε-aminocaproic acid on mortality

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size	Surgical procedure	Location			Risk estimate (95% CI)	Significance P-value	
ADULT POPULATION/	IV E-AMINOCAPRO	DIC ACID							
Henry (2007)	Level I Good	5 RCTs N=714	Adult patients undergoing any surgery	Hospital – planned surgery Various countries ^c	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Mortality	RR 1.17 (0.47, 2.93)	No difference 0.73	None Phet=0.78 (I ² =0%)
Cardiac surgery									
Henry (2007)	Level I Good	4 RCTs N=632	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Mortality	RR 1.65 (0.50, 5.43)	No difference 0.41	None Phet=0.81 (I ² =0%)
Brown (2007)	Level I Fair	6 RCTs N=735	Adult patients undergoing cardiac surgery	Hospital – planned surgery Unknown	E-aminocaproic acid (IV) vs placebo	Mortality	RR 1.82 (0.55, 5.98)	No difference 0.32	NR
Orthopaedic surge	ery				•				•
Berenholtz (2009)	Level II Good	1 RCT N=182	Adults undergoing <u>major</u> spine surgery	Hospital – planned surgery US	E-aminocaproic acid (IV) vs placebo	In-hospital mortality	RR 0.30 (0.01, 8.08) ^d	No difference 0.50 d	NA

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: CI, confidence interval; het, heterogeneity; IV, intravenous; NA, not applicable; NR, not reported; RCT, randomised controlled trial; RR, risk ratio.

a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I²<25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between ε-aminocaproic acid and no ε-aminocaproic acid.

d Post-hoc analysis conducted for this guideline.

	OQ3.I8c.P5 (MI)			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies)				
There is one pivotal Level I study (Henry 2007/good quality) which includes data from up to 4 RCTs, one supportive A One or more level I studies with a low risk of bias or several level II studies.	dies with a low risk of bias			
Level I study (Brown 2007/fair quality) and one additional RCT conducted in the US. B One or two Level II studies with a low risk of bias or SR/several Level III studies.	I studies with a low risk of bias			
C One or two Level III studies with a low risk of bias or Level I or II studies	s with a moderate risk of bias			
D Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')				
Results generally consistent between pivotal and supportive meta-analyses, however there is a slight difference in A All studies consistent				
direction of effect. Consistency of individual studies within meta-analyses described below. Pivotal evidence – Henry 2007 B Most studies consistent and inconsistency can be explained				
No significant heterogeneity (see note) in cardiac surgery analysis. C Some inconsistency, reflecting genuine uncertainty around question				
D Evidence is inconsistent	Evidence is inconsistent			
NA Not applicable (one study only)	Not applicable (one study only)			
3. Clinical impact (Indicate in the space below if the study results varied according to some unknown factor (not simply study quality or sample size) and thus the clinical impact of the determined)	the intervention could not be			
Pivotal evidence – Henry 2007 A Very large				
Cardiac surgery - 3.5% vs 4.3%; RR 0.89 (0.37, 2.18); 4 RCTs (N=632) Supportive evidence - Brown 2007 B Substantial				
Cardiac surgery – absolute risk not reported; RR 1.14 (0.50, 2.60); 8 RCTs (N=839)				
D Underpowered				
4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?)				
The evidence is generalisable to an adult population who are undergoing planned surgery. Only cardiac surgery A Evidence directly generalisable to target population				
was included in the systematic reviews. There was one RCT in orthopaedic surgery. B Evidence directly generalisable to target population with some caveats				
C Evidence not directly generalisable to the target population but could be	e sensibly applied			
D Evidence not directly generalisable to target population and hard to judg apply	lge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of health services/delivery of care and cultural factors?)				
Hospital setting. A Evidence directly applicable to Australian healthcare context				
The pivotal review states that studies were conducted in a wide range of countries. B Evidence applicable to Australian healthcare context with few caveats	Evidence applicable to Australian healthcare context with few caveats			
C Evidence probably applicable to Australian healthcare context with some	ne caveats			
D Evidence not applicable to Australian healthcare context				

E-aminocaproic acid is a synthetic derivative of the amino acid lysine. It is not currently TGA-approved for use in Australia.

Generalisability rated C as most evidence in a restricted surgical population but applied to general surgical population.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
Evidence base	Α	There is one pivotal Level I study (good quality), one supportive Level I study and one additional RCT.
2. Consistency	В	No heterogeneity in the pivotal study. Small difference in direction of effect between the pivotal and supportive Level I studies. There were no events in the additional RCT.
3. Clinical impact	D	There was no significant difference in the analyses; however, the studies are likely to be underpowered to detect a difference.
4. Generalisability	С	The results are generalisable to an adult surgical population; in particular those undergoing cardiac surgery
5. Applicability	В	Studies were conducted in a range of countries. The additional RCT was conducted in the US. Likely to be applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intravenous ε-aminocaproic acid therapy on the risk of myocardial infarction, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI, confidence interval, RBC, red blood cell; SR, systematic review;

a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

POQ3.l8c.P5 (MI) Characteristics and results of studies examining the effect of ε-aminocaproic acid on morbidity (myocardial infarction)

1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		No. of trials /			Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size	Surgical procedure				Risk estimate (95% CI)	Significance P-value	
ADULT POPULATION	IV E-AMINOCAPRO	DIC ACID							
Cardiac surgery									
Henry (2007)	Level I Good	4 RCTs N=632	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries ^c	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Myocardial infarction	RR 0.89 (0.37, 2.18)	No difference 0.80	None Phet=0.33 (I ² =12%)
Brown (2007)	Level I Fair	8 RCTs N=839	Adult patients undergoing cardiac surgery	Hospital – planned surgery Countries not specified	E-aminocaproic acid (IV) vs placebo	Myocardial infarction	RR 1.14 (0.50, 2.60)	No difference 0.76	NR
Orthopaedic surge	ery								
Berenholtz (2009)	Level II Good	1 RCT N=182	Adult patients undergoing major spine surgery	Hospital – planned surgery US	E-aminocaproic acid (IV) vs placebo	Myocardial infarction	NAd	No difference NA	NA

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: CI, confidence interval; het, heterogeneity; IV, intravenous; NA, not applicable; NR, not reported; RCT, randomised controlled trial; RR, risk ratio.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

^b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between ε-aminocaproic acid and no ε-aminocaproic acid.

^d There were no myocardial infarctions in either treatment group.

Key question(s): In patients undergoing surgery, what is the effect of administration of ε-amino	capr	oic acid on morbidity (stroke)?	Evidence table refa: POQ3.I8c.P5 (stroke)			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)					
There is one pivotal Level I study (Henry 2007/good quality) which includes data from 3 RCTs, one supportive Level I study (Brown 2007/fair quality) and one additional RCT (Berenholtz 2009/good quality) in major spine surgery,	Α	One or more level I studies with a low risk of bias or several level	I studies with a low risk of bias			
conducted in the US.	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Results consistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta- analyses described below.	Α	All studies consistent				
Pivotal evidence – Henry 2007	В	Most studies consistent and inconsistency can be explained				
No significant heterogeneity (see note) in main analysis.	С	Some inconsistency, reflecting genuine uncertainty around question				
	D	Evidence is inconsistent				
	NA	A Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some undetermined)	nknow	<u>n</u> factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be			
Pivotal evidence – Henry 2007 Cardiac surgery – 0.6% vs 0.9%; RR 0.59 (0.10, 3.44); 3 RCTs (N=541)	А	Very large				
Supportive evidence – Brown 2007	В	Substantial				
Cardiac surgery – absolute risk not reported; RR 0.60 (0.13, 2.81); 8 RCTs (N=833)	С	Moderate				
	D	Underpowered				
4. Generalisability (How well does the body of evidence match the population and clinical set	ttings t	peing targeted by the Guideline?)				
The evidence is generalisable to an adult population who are undergoing planned surgery. Only cardiac surgery was included in the systematic reviews. There was one RCT in orthopaedic surgery.	Α	Evidence directly generalisable to target population				
was included in the systematic reviews. There was the NCT in orthopaedic surgery.	В	Evidence directly generalisable to target population with some cave	reats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of hear	Ith services/delivery of care and cultural factors?)				
Hospital setting.	Α	Evidence directly applicable to Australian healthcare context				
The pivotal review states that studies were conducted in a wide range of countries.	В	Evidence applicable to Australian healthcare context with few caveats				
	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
	D	Evidence not applicable to Australian healthcare context				

E-aminocaproic acid is a synthetic derivative of the amino acid lysine. It is not currently TGA-approved for use in Australia.

Generalisability rated C as most evidence in a restricted surgical population but applied to general surgical population.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (good quality), one supportive Level I study and one additional RCT.
2. Consistency	Α	There was no heterogeneity in the pivotal Level I study. Similar results were seen in the supportive Level I study and the additional RCT.
3. Clinical impact	D	There was no significant difference in the analyses; however, the studies are likely to be underpowered to detect a difference in stroke.
4. Generalisability	С	The results are generalisable to an adult surgical population; in particular those undergoing cardiac surgery
5. Applicability	В	Studies were conducted in a range of countries. The additional RCT was conducted in the US. Likely to be applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intravenous ε-aminocaproic acid therapy on risk of stroke, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI, confidence interval, RBC, red blood cell; SR, systematic review;

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.l8a.P5 (stroke) Characteristics and results of studies examining the effect of ε-aminocaproic acid on morbidity (stroke)

evid	Level of	No. of trials /	Patient population / Surgical procedure	Setting	Intervention	Outcome	Results	Heterogeneity ^b	
	evidence ^a Quality	sample size					Risk estimate (95% CI)	Significance P-value	
ADULT POPULATION	IV E-AMINOCAPRO	DIC ACID							
Cardiac surgery									
Henry (2007)	Level I Good	3 RCTs N=541	Adult patients undergoing cardiac surgery	Hospital – planned surgery Various countries ^c	E-aminocaproic acid (IV) vs no ε-aminocaproic acid	Stroke	RR 0.59 (0.10, 3.44)	No difference 0.55	None Phet=0.47 (I ² =0%)
Brown (2007)	Level I Fair	8 RCTs N=833	Adult patients undergoing cardiac surgery	Hospital – planned surgery Countries not specified	E-aminocaproic acid (IV) vs placebo	Stroke	RR 0.60 (0.13, 2.81)	No difference 0.52	NR
Orthopaedic surge	ery								
Berenholtz (2009)	Level II Good	1 RCT N=182	Adults undergoing major spine surgery	Hospital – planned surgery US	E-aminocaproic acid (IV) vs placebo	Cerebral infarction/TIA	RR 0.30 (0.01, 8.08) ^d	No difference 0.50 ^d	NA

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: ASA, acetylsalicylic acid; CI, confidence interval; DVT, deep vein thrombosis; het, heterogeneity; IV, intravenous; MI, myocardial infarction; NR, not reported; PE, pulmonary embolism; OR, odds ratio; RCT, randomised controlled trial; RR, risk ratio; TIA, transient ischaemic attack.

a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I²<25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

^c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between ε-aminocaproic acid and no ε-aminocaproic acid.

^d Post-hoc analysis conducted for this guideline.

In patients undergoing surgery, what is the effect of administration of e-aminocaproic acid on morbidity (thrombosis)? 1. Evidence base (number of studies, level of evidence and risk of bias in the included studies with a low risk of bias or several level II studies with a low risk of bias or several level II studies with a low risk of bias or several level II studies with a low risk of bias or several level II studies with a low risk of bias or several level II studies with a low risk of bias or several level II studies with a low risk of bias or several level II studies with a low risk of bias or several level II studies with a low risk of bias or several level II studies with a low risk of bias or several level II studies with a low risk of bias or several level II studies with a low risk of bias or several level III studies with a low risk of bias or several level II studies with a lo	Key question(s):			Evidence table refa:		
There were one photal Level III study (Herry 2007) and entirely which included data from only one syndle pool graphity (Arr on the photal evaluation one small dar usually of Crif febre surgery, one supported. Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or		capr	oic acid on morbidity (thrombosis)?	POQ3.I8c.P5 (thromb)		
Public Note for findingselds supply and make small find quality with in land quality which in land data from three RCTs and one additional RCT (Berenholtz 2009/space quality) in major spine surgery conducted in the US. C One or two Level III studies with a low risk of bias or Level I or III studies with a moderate risk of bias or Level I or III studies with a high risk of bias C One or two Level III studies with a high risk of bias D Level IV studies or Level I or III studies with a high risk of bias A All studies consistent	1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	5)				
Regional 2009/good qualityly which included data from three RCTs and one additional RCT (Berembutz 2009/god quality) in major spine surgery conducted in the US. C	There was one pivotal Level I/II study (Henry 2007/good quality) which included data from only one small good	Α	One or more level I studies with a low risk of bias or several level	Il studies with a low risk of bias		
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		D	Evidence not applicable to Australian healthcare context			

E-aminocaproic acid is a synthetic derivative of the amino acid lysine. It is not currently TGA-approved for use in Australia.

Generalisability rated C as most evidence in a restricted surgical population but applied to general surgical population.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	В	There was one pivotal Level I study (good quality) which included data from only one small good quality RCT, one supportive Level I study (which included three RCTs) and one additional RCT.
2. Consistency	Α	Results similar suggesting potentially no difference or less risk with ε-aminocaproic acid.
3. Clinical impact	D	There was no significant difference in the analyses; however, studies likely to be underpowered to detect a difference.
4. Generalisability	С	The results are generalisable to an adult surgical population; in particular those undergoing orthopaedic surgery
5. Applicability	С	A small number of studies were conducted in a range of countries. The additional RCT was conducted in the US. Possibly applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intravenous ε-aminocaproic acid therapy on risk of venous thromboembolism, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI, confidence interval, RBC, red blood cell: SR, systematic review;

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.l8c.P5 (thrombosis) Characteristics and results of studies examining the effect of ε-aminocaproic acid on morbidity (thrombosis)

Study Level of evidence ^a Quality		No. of trials /	Patient population /	Setting	Intervention	Outcome	Results	Heterogeneity ^b	
		3.7				Risk estimate (95% CI)	Significance P-value		
ADULT POPULATION/	IV E-AMINOCAPRO	DIC ACID							
Orthopaedic surge	ery								
. , , , , , , , , , , , , , , , , , , ,	Level I/II Good/Good	1 RCT N=46	Adult patients undergoing orthopaedic surgery	Hospital – planned surgery Unknown	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Deep vein thrombosis	RR 1.09 (0.25, 4.85)	No difference 0.91	NA
						Pulmonary embolism	RR 0.36 (0.02, 8.46)	No difference 0.53	NA
Kagoma (2009)	Level I Good	3 RCTs N=180	Adults undergoing orthopaedic surgery	Hospital – planned surgery Unknown	E-aminocaproic acid (IV) vs placebo	VTE complications	RD 0.00 (-0.07, 0.07)	No difference NR	NR
Berenholtz (2009)	Level II Good	1 RCT N=182		Hospital – planned surgery US	E-aminocaproic acid (IV) vs placebo	Deep vein thrombosis	RR 0.20 (0.01, 4.11) ^c	No difference 0.30°	NA
						Pulmonary embolism	RR 0.33 (0.04, 3.15) ^c	No difference 0.34 °	NA
						Any thrombotic complication	RR 0.33 (0.07, 1.61) ^c	No difference 0.17 °	NA
Liver surgery			•	•	•	•	•		•
Henry (2007)	Level I/II Good/Fair	1 RCT N=82	Adult patients undergoing liver transplant	Hospital – planned surgery Unknown	E-aminocaproic acid (IV) vs placebo	Other thrombosis (not MI, stroke, DVT or PE)	RR 0.95 (0.14, 6.44)	No difference 0.96	NA

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: CI, confidence interval; DVT, deep vein thrombosis; het, heterogeneity; IV, intravenous; MI, myocardial infarction; NA, not applicable; NR, not reported; PE, pulmonary embolism; OR, odds ratio; RCT, randomised controlled trial; RD, risk difference; RR, risk ratio.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I²<25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

^c Post-hoc analysis for this guideline.

Key question(s):			Evidence table refa:
In patients undergoing surgery, what is the effect of administration of $\underline{\epsilon}$ -amino	capro	<u>pic acid</u> on <u>quality of life</u> ?	POQ3.I8c.P6
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies,)		
No studies of any level were identified which assessed the effect of ε-aminocaproic acid on quality of life.	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias
	В	One or two Level II studies with a low risk of bias or SR/several Lev	rel III studies with a low risk of bias
	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	
2. Consistency (If only one study was available, rank this component as 'not applicable')			
NA	Α	All studies consistent	
	В	Most studies consistent and inconsistency can be explained	
	С	Some inconsistency, reflecting genuine uncertainty around question	on
	D	Evidence is inconsistent	
	NA	Not applicable (one study only)	
3. Clinical impact (Indicate in the space below if the study results varied according to some undetermined)	knowi	n factor (not simply study quality or sample size) and thus the clinical impa	ct of the intervention could not be
NA NA	Α	Very large	
	В	Substantial	
	С	Moderate	
	D	Slight/Restricted	
4. Generalisability (How well does the body of evidence match the population and clinical sett	tings b		
NA	Α	Evidence directly generalisable to target population	
	В	Evidence directly generalisable to target population with some cav	eats
	С	Evidence not directly generalisable to the target population but cou	ıld be sensibly applied
	D	Evidence not directly generalisable to target population and hard to	judge whether it is sensible to
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of heal	·	
NA	A	Evidence directly applicable to Australian healthcare context	
	В	Evidence applicable to Australian healthcare context with few cave	
	С	Evidence probably applicable to Australian healthcare context with	some caveats
	D	Evidence not applicable to Australian healthcare context	

E-aminocaproic acid is a synthetic derivative of the amino acid lysine. It is not currently listed for use in Australia.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
Evidence base	NA	
2. Consistency	NA	
3. Clinical impact	NA	
4. Generalisability	NA	
5. Applicability	NA	

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intravenous ε-aminocaproic acid therapy on quality of life, compared with no therapy, is unknown.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI, confidence interval, RBC, red blood cell: SR, systematic review;

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Key question(s):			Evidence table refa:			
In patients undergoing surgery, what is the effect of administration of ε -amino	capr	oic acid on re-operation for bleeding?	POQ3.l8c.S2			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)					
There is one pivotal Level I study (Henry 2007/good quality) including data from 5 RCTs, one supportive Level I	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias			
study (Brown 2007/fair quality), one supportive Level I/II study which includes data from one poor quality RCT (McIlroy 2009/good quality) and one additional RCT in major spine surgery (Berenholtz 2009/good quality) published	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias			
since the pivotal review and conducted in the US.	С	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bia				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	S			
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Results consistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta-	Α	All studies consistent				
lyses described below. otal evidence – Henry 2007		Most studies consistent and inconsistency can be explained				
votal evidence – Henry 2007 o significant heterogeneity (see note) in main analysis.	С	Some inconsistency, reflecting genuine uncertainty around quest	ion			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>un</u> determined)	<u>nknow</u>	<u>n</u> factor (not simply study quality or sample size) and thus the clinical imp	act of the intervention could not be			
Pivotal evidence – Henry 2007	Α	Very large				
Cardiac surgery - 0.8% vs 3.3%; RR 0.35 (0.11, 1.17); 5 RCTs (N=740) Supportive evidence - Brown 2007	В	Substantial (potential)				
Cardiac surgery (return to operating room) – absolute risk not reported; RR 0.51 (0.15, 1.82); 9 RCTs (N=851)	С	Moderate				
, , , , , , , , , , , , , , , , , , ,	D	Slight/Restricted				
4. Generalisability (How well does the body of evidence match the population and clinical set	tings i	being targeted by the Guideline?)				
The evidence is generalisable to an adult population who are undergoing planned surgery. Evidence was	Α	Evidence directly generalisable to target population				
predominantly in cardiac surgery although there was one RCT in orthopaedic surgery.	В	Evidence directly generalisable to target population with some car	veats			
	С	Evidence not directly generalisable to the target population but co	ould be sensibly applied			
	D	Evidence not directly generalisable to target population and hard	to judge whether it is sensible to			
		apply				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of hea					
Hospital setting.	Α	Evidence directly applicable to Australian healthcare context				
The pivotal review states that studies were conducted in a range of countries.	В	Evidence applicable to Australian healthcare context with few caveats				
	С	Evidence probably applicable to Australian healthcare context wit	h some caveats			
	D	Evidence not applicable to Australian healthcare context				

E-aminocaproic acid is a synthetic derivative of the amino acid lysine. It is not currently TGA-approved for use in Australia.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
Evidence base	А	There is one pivotal Level I study (good quality), one supportive Level I studies, one supportive Level I/II study (including 1 RCT only) and one additional RCT (major spine surgery).
2. Consistency	Α	There was no heterogeneity in the pivotal study and the results of the supportive and additional studies were consistent.
3. Clinical impact	В	There was no significant difference in the analyses but potentially substantial if trend upheld by greater powering.
4. Generalisability	Α	The results are generalisable to an adult cardiac surgical population.
5. Applicability	В	Studies were conducted in a range of countries. Likely to be applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, the effect of intravenous ε-aminocaproic acid therapy on risk of reoperation for bleeding, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI, confidence interval, Pot, potential; RBC, red blood cell; SR, systematic review;

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

^b Potentially substantial if results upheld in studies with greater power

POQ3.l8c.S2 Characteristics and results of studies examining the effect of ε-aminocaproic acid on re-operation for bleeding

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results	Heterogeneity ^b	
	evidence ^a sample size Surgical procedure Location			Risk estimate (95% CI)	Significance P-value				
ADULT POPULATION/	IV E-AMINOCAPRO	DIC ACID							
Cardiac surgery									
Henry (2007)	Level I Good	5 RCTs N=662	Adult patients undergoing any surgery	Hospital – planned surgery Various countries ^c	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Reoperation for bleeding	RR 0.35 (0.11, 1.17)	No difference 0.087	None Phet=0.78 (l²=0%)
Brown (2007)	Level I Fair	9 RCTs N=851	Adult patients undergoing cardiac surgery	Hospital – planned surgery Countries not specified	E-aminocaproic acid (IV) vs placebo	Return to operating room	RR 0.51 (0.15, 1.82)	No difference 0.30	NR
McIlroy(2009)	Level I/II Good/Poor	1 RCT N=30	Adult patients <u>receiving</u> <u>ASA</u> undergoing cardiac surgery	Hospital – planned surgery Unknown	E-aminocaproic acid (IV) vs placebo	Surgical re-exploration	OR 0.31 (0.01, 8.28)	No difference NR	NA
Orthopaedic surge	ery								
Berenholtz (2009)	Level II Good	1 RCT N=182	Adult patients undergoing major spine surgery	Hospital – planned surgery US	E-aminocaproic acid (IV) vs placebo	Reoperation for bleeding	RR 0.20 (0.01, 4.11) ^d	No difference 0.30 d	NA

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: ASA, acetylsalicylic acid; CI, confidence interval; het, heterogeneity; IV, intravenous; NA, not applicable; NR, not reported; OR, odds ratio; RCT, randomised controlled trial; RR, risk ratio.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and 12<25%; (ii) mild heterogeneity if 12 <25%; moderate heterogeneity if 12 between 25-50%; substantial heterogeneity 12 >50%.

c Studies conducted in a wide range of countries. Eight of the 211 studies included in the review were conducted in Australia; however, it is unknown how many of these were related specifically to the comparison between.

^d Post-hoc analysis conducted for this guideline.

Key question(s):		ssis acid on beauthal langth of star 2	Evidence table refa: POQ3.18c.S5				
In patients undergoing surgery, what is the effect of administration of ε-amino	<u>)capi</u>	oic acid on nospital length of stay?	F 0Q3.106.33				
1. Evidence base (number of studies, level of evidence and risk of bias in the included studie	s)						
There is one pivotal Level I/II study (Henry 2007/good quality) which contains one RCT (good quality) and one	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias				
additional RCT (Berenholtz 2009/good quality), both in orthopaedic surgery.	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias				
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias				
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	;				
2. Consistency (If only one study was available, rank this component as 'not applicable')							
Only one RCT included in Henry review (in orthopaedic surgery) and one additional RCT (in major spine surgery).	Α	All studies consistent					
Results conflicting; both show no significant difference but post estimates in different directions.	В	Most studies consistent and inconsistency can be explained					
	С	Some inconsistency, reflecting genuine uncertainty around questi	cy, reflecting genuine uncertainty around question				
	D	Evidence is inconsistent					
	NA	Not applicable (one study only)					
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u> determined)	<u>nknow</u>	<u>n</u> factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be				
Pivotal evidence – Henry 2007	Α	Very large					
Orthopaedic surgery (days) - MD 2.90 (-0.96, 6.76); 1 RCT (N=46) Supportive evidence - Berenholtz 2009	В	Substantial					
Major spine surgery (days) – MD -1.00 (-2.94, 0.94); 1 RCT (N=182)	С	Moderate					
	D	Inconsistent					
4. Generalisability (How well does the body of evidence match the population and clinical se	ttings l	being targeted by the Guideline?)					
The evidence is generalisable to an adult population who are undergoing planned surgery. Both included studies	Α	Evidence directly generalisable to target population					
were in orthopaedic surgery.	В	Evidence directly generalisable to target population with some cav	/eats				
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied				
	D	Evidence not directly generalisable to target population and hard tapply	o judge whether it is sensible to				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of hea	Ith services/delivery of care and cultural factors?)					
Hospital setting.	Α	Evidence directly applicable to Australian healthcare context					
The location of the single RCT from the Henry review is unknown. The Berenholtz RCT was conducted in the US.	В	Evidence applicable to Australian healthcare context with few cave	eats				
	С	Evidence probably applicable to Australian healthcare context with	n some caveats				
	D	Evidence not applicable to Australian healthcare context					

E-aminocaproic acid is a synthetic derivative of the amino acid lysine. It is not currently TGA-approved for use in Australia.

Generalisability rated C as most evidence in a restricted surgical population but applied to general surgical population.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	В	There is one pivotal Level I/II study (good quality) which contains one RCT (good quality) and one additional RCT, both in orthopaedic surgery.
2. Consistency	D	Conflicting direction of the point estimates between the two RCTs.
3. Clinical impact	D	There was no significant difference in the analyses; however, the results were conflicting.
4. Generalisability	С	The results are generalisable to an adult population undergoing orthopaedic/spine surgery.
5. Applicability	С	One RCT was conducted in the US, the other is unknown. Possibly applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intravenous ε-aminocaproic acid therapy on length of hospital stay, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI. confidence interval. RBC, red blood cell: SR, systematic review:

Primary outcomes; P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.l8c.S5 Characteristics and results of studies examining the effect of ε-aminocaproic acid on hospital length of stay

Study	, I	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size	Surgical procedure	Location			Risk estimate (95% CI)	Significance P-value	
ADULT POPULATION/	IV E-AMINOCAPRO	DIC ACID							
Orthopaedic surge	ery								
Henry (2007)	Level I/II Good/Good	1 RCT N=46	Adult patients undergoing orthopaedic surgery	Hospital – planned surgery Unknown	E-aminocaproic acid (IV) vs no ε- aminocaproic acid	Hospital length of stay (days)	MD 2.90 (-0.96, 6.76)	No difference 0.14	NA
Berenholtz (2009)	Level II Good	1 RCT N=182	Adult patients undergoing major spine surgery	Hospital – planned surgery US	E-aminocaproic acid (IV) vs placebo	Hospital length of stay (days)	MD -1.00 (-2.94, 0.94)°	No difference 0.31°	NA

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: CI, confidence interval; het, heterogeneity; IV, intravenous; NR, not reported; RCT, randomised controlled trial; RR, risk ratio.

a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and |2<25%; (ii) mild heterogeneity if |2<25%; moderate heterogeneity if |2 between 25-50%; substantial heterogeneity |2 >50%.

^c Post-hoc analysis conducted for this Guideline.

Recommendation(s) for administration of ε-aminocaproic acid

RECOMMENDATION	GRADE	RELE	VANT
What recommendation(s) does the guideline development group draw from this evidence? Use action statements where possible.		EVIDENO	CE TABLE
In adult patients undergoing cardiac surgery, the use of intravenous ε-aminocaproic acid is recommended.	С	PO3.	l8c.P1,
		PO3.	l8c.P3
IMPLEMENTATION OF RECOMMENDATION			
Please indicate yes or no to the following questions. Where the answer is yes please provide explanatory information about this. This information will be used to develop the implementation plan for the guidelines.			
Will this recommendation result in changes in usual care?		YES	NO
Increased use of ε-aminocaproic acid.		•	
Are there any resource implications associated with implementing this recommendation?		YES	NO
Drug cost (albeit modest).			
Will the implementation of this recommendation require changes in the way care is currently organised?		YES	NO
Are the guideline development group aware of any barriers to the implementation of this recommendation		YES	NO
Potential resistance.			
What could help to facilitate implementation of the recommendation?		YES	NO
Education; promotion of PO guideline.			

Intervention 8 – Administration of antifibrinolytics & DDAVP: Desmopressin

			ı		
Key question(s):			Evidence table refa:		
In patients undergoing surgery, what is the effect of administration of desmor	oress	in on transfusion incidence?	POQ3.l8d.P1		
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)				
There is one pivotal Level I study (Crescenzi 2008/ fair quality) which includes data from up to 21 RCTs, one	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias		
supportive Level I study (Carless 2008/ good quality) and one supportive Level I/II study (Gurusamy 2009/good-fair quality) in a specific surgery type (liver resection).	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias		
quality) in a specific surgery type (liver resection).	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias		
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	i		
2. Consistency (If only one study was available, rank this component as 'not applicable')					
Results consistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta-	Α	All studies consistent			
analyses described below. Pivotal evidence – Crescenzi 2008	В	Most studies consistent and inconsistency can be explained			
No significant heterogeneity (see note) in main analyses and most subgroup analyses. Moderate heterogeneity in	С	Some inconsistency, reflecting genuine uncertainty around questi	on		
analyses of cardiac surgery subgroup.	D	Evidence is inconsistent			
Supportive evidence – Carless 2008 No significant heterogeneity in most analyses except moderate heterogeneity in some subgroups (see attached table).	NA	Not applicable (one study only)			
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u> determined)	nknow	n factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be		
Pivotal evidence – Crescenzi 2008	Α	Very large			
Any surgery (all blood products) – 55.1% vs 57.9%; OR 0.88 (0.70, 1.10); 21 RCTs (N=1429) Cardiac surgery (all blood products) – 54.9% vs 57.9%; OR 0.87 (0.68, 1.11); 16 RCTs (N=1213)	В	Substantial			
Cardiac surgery (platelets) – 9.6% vs 9.1%; OR 0.64 (0.41, 1.01); 11 RCTs (N=769)	С	Moderate			
Non-cardiac surgery – 56.6% vs 57.9%; OR 0.93 (0.48, 1.79); 5 RCTs (N=216) Supportive evidence – Carless 2008 (see Summary Table POQ3.l8d.P1) especially surgery type subgroups	D	Slight/Restricted (primary CABG; complex surgery and non-cardia only])	c surgery; cardiac surgery [platelets		
4. Generalisability (How well does the body of evidence match the population and clinical se	ttings b	peing targeted by the Guideline?)			
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were	Α	Evidence directly generalisable to target population			
included in the overall analysis although studies were predominantly in cardiac surgery (16/21 RCTs). There were also analyses of patients undergoing cardiac surgery who had or had not received ASA within 7 days prior to	В	Evidence directly generalisable to target population with some cav	veats		
surgery.	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied (non-cardiac)		
	D	Evidence not directly generalisable to target population and hard tapply	o judge whether it is sensible to		
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of heal	th services/delivery of care and cultural factors?)			
Hospital setting.	Α	Evidence directly applicable to Australian healthcare context			
The pivotal review does not state which countries the RCTs were conducted in but the supportive study (which includes most of the studies included in the pivotal review) states that studies were conducted in US, Canada,	В	Evidence applicable to Australian healthcare context with few cave	eats		
Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK.	С	Evidence probably applicable to Australian healthcare context witl	n some caveats		
	D	Evidence not applicable to Australian healthcare context			

Desmopressin injection currently listed by TGA for the following indication: "patients undergoing cardiac surgery with cardiopulmonary bypass for prosthetic valve replacement or aortocoronary bypass grafting, especially when it is complicated by platelet function defects sufficient to prolong bleeding time despite relatively normal platelet cover. Desmopressin acetate offers no benefit as routine therapy in patients having an uncomplicated (simple) cardiopulmonary bypass procedure".

The Crescenzi (pivotal) review includes more studies than the Carless (supportive) review due to the more restricted inclusion criteria used in Carless (ie, elective or non-urgent surgery). The Crescenzi review assesses transfusion of whole blood products (including RBCs, FFP and platelets) while the Carless review assesses only transfusion of RBCs. Both the Crescenzi and Carless reviews include any surgery type, while the earlier reviews assessed only cardiac surgery. The Crescenzi review has been rated as *fair* quality due to the lack of formal quality assessment and lack of investigation of heterogeneity. The Carless review assessed quality and performed a subgroup analysis of transfusion incidence for all surgery types based on the rating (A,B or C) of treatment allocation. The analysis showed no substantial difference in the results between studies rated A, B or C, suggesting study quality may not have greatly influenced the findings. All studies included in the Carless review were included in the Crescenzi review. The Carless review performed extensive subgroup analyses for this outcome (shown in Table POQ3.l8d.P1).

There was significant discussion amongst the CRG around different surgical subgroups so separate ratings have been given for each.

EVIDENCE STATEMENT MATRIX

Component	Primary CABG	Complex cardiac	Cardiac (PLT)	Noncardiac	Description
Evidence base	А	А	А	А	There is one pivotal Level I study (fair quality), one supportive Level I study and one supportive Level I/II study in a specific surgery type.
2. Consistency	В	В	В	В	Most studies were reasonably consistent. Some heterogeneity in cardiac subgroup analyses
3. Clinical impact	D	D	D	D	There was no significant difference in the majority of surgery types. Large potential risk estimate for platelets in cardiac surgery but not significant, possibly due to underpowering.
4. Generalisability	А	А	А	С	The results are generalisable to an adult surgical population; in particular those undergoing cardiac surgery
5. Applicability	В	В	В	В	Studies were conducted in a wide range of countries. Likely to be applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

In adult patients undergoing primary coronary artery bypass surgery, intravenous desmopressin therapy reduces the incidence of allogeneic blood transfusion compared with no therapy.

In adult patients undergoing complex cardiac surgery, intravenous desmopressin therapy does not reduce the incidence of allogeneic blood transfusion compared with no therapy.

In adult patients undergoing cardiac surgery, intravenous desmopressin therapy may reduce the incidence of platelet transfusion compared with no therapy.

In adult patients undergoing noncardiac surgery in which substantial blood loss is anticipated, intravenous desmopressin therapy does not appear to reduce the incidence of allogeneic blood transfusion compared with no therapy.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity if I² >50%. Abbreviations: CI, confidence interval, Pot, potential; RBC, red blood cell; SR, systematic review.

^a Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

OQ3.l8d.P1 Characteristics and results of studies examining the effect of <u>desmopressin</u> on <u>transfusion incidence</u>.

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size	procedure	Location			Risk estimate (95% CI)	Significance P-value	
ADULT POPULATION	IV DESMOPRESSI	N							
Any surgery									
Crescenzi (2008)	Level I Fair	21 RCTs N=1429	Adult patients undergoing any surgery	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Transfusion incidence (blood products) ^c	OR 0.88 (0.70, 1.10)	No difference P=0.26	None Phet=0.19 (I ² =21.4%)
Carless (2008)	Level I Good	17 RCTs N=1308	Adult patients undergoing any surgery	Hospital – elective or non-urgent surgery Various ^d	Desmopressin (IV) vs no desmopressin	Transfusion incidence (RBCs)	RR 0.96 (0.87, 1.06)	No difference P=0.42	None Phet=0.19 (I ² =22%)
Carless (2008)	Level I Good	10 RCTs N=736	Adult patients undergoing any surgery with transfusion protocol	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion incidence (RBCs)	RR 0.90 (0.77, 1.04)	No difference P=0.16	None Phet=0.25 (I ² =21%)
		7 RCTs N=572	Adult patients undergoing any surgery without transfusion protocol	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion incidence (RBCs)	RR 1.03 (0.93, 1.14)	No difference P=0.60	None Phet=0.40 (l²=4%)
Carless (2008)	Level I Good	8 RCTs N=635	Adult patients undergoing any surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion incidence (RBCs) <u>using</u> <u>autologous</u> <u>techniques</u>	RR 1.00 (0.84, 1.19)	No difference P=0.97	None Phet=0.31 (I ² =15%)
		9 RCTs N=673	Adult patients undergoing any surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion incidence (RBCs) not using autologous techniques	RR 0.91 (0.78, 1.07)	No difference P=0.25	Moderate Phet=0.04 (I ² =50%)
Carless (2008)	Level I Good Rating Ae	2 RCTs N=190	Adult patients undergoing any surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion incidence (RBCs)	RR 0.97 (0.75, 1.24)	No difference P=0.80	None Phet=0.50 (l ² =0%)
	Level I Good Rating Be	10 RCTs N=746	Adult patients undergoing any surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion incidence (RBCs)	RR 0.88 (0.75, 1.03)	No difference P=0.12	Substantial Phet=0.04 (I ² =50%)
	Level I Good Rating Ce	5 RCTs N=372	Adult patients undergoing any surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion incidence (RBCs)	RR 1.11 (0.94, 1.33)	No difference P=0.22	None Phet=0.75 (l²=0%)
Cardiac surgery						_			
Crescenzi (2008)	Level I Fair	16 trials N=1213	Adult patients undergoing cardiac surgery	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Transfusion incidence (blood products) ^a	OR 0.87 (0.68, 1.11)	No difference P=0.26	Moderate Phet=0.07 (l ² =37.0%)

Study	Level of	No. of trials /	Patient population / Surgical	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size	procedure	Location			Risk estimate (95% CI)	Significance P-value	
		11 RCTs N=769	Adult patients undergoing cardiac surgery	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Transfusion incidence (platelets)	OR 0.64 (0.41, 1.01)	No difference P=0.06	None Phet=0.22 (l ² =23.1%)
Carless (2008)	Level I Good	14 RCTs N=1137	Adult patients undergoing cardiac surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion incidence (RBCs)	RR 0.95 (0.84, 1.07)	No difference P=0.39	Moderate Phet=0.11 (I ² =33%)
Carless (2008)	Level I Good	8 RCTs N=527	Adult patients undergoing primary CABG	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion incidence (RBCs)	RR 0.85 (0.73, 0.99)	Favours DDAVP P=0.038	None Phet=0.43 (I ² =0%)
		6 RCTs N=610	Adult patients undergoing <u>CABG + valve ±</u> <u>combination/redo surgery</u>	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion incidence (RBCs)	RR 1.03 (0.88, 1.19)	No difference P=0.75	Moderate Phet=0.14 (I ² =40%)
Carless (2008)	Level I Good	5 RCTs N=340	Adult patients undergoing cardiac surgery who have had ASA within 7 days prior	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion incidence (RBCs)	RR 0.89 (0.64, 1.23)	No difference P=0.49	Moderate Phet=0.12 (I²=40%)
		4 RCTs N=286	Adult patients undergoing cardiac surgery who have had no ASA within 7 days prior	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion incidence (RBCs)	RR 0.79 (0.62, 1.01)	No difference P=0.056	None Phet=0.36 (I ² =7%)
Non-cardiac surge	ery								
Crescenzi (2008)	Level I Fair	5 RCTs N=216	Adults patients undergoing surgery other than cardiac	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Transfusion incidence (blood products) ^c	OR 0.93 (0.48, 1.79)	No difference P=0.83	None Phet=0.81 (I ² =0%)
Carless (2008)	Level I Good	3 RCTs N=171	Adults patients undergoing surgery other than cardiac	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion incidence (RBCs)	RR 1.01 (0.81, 1.26)	No difference P=0.91	None Phet=0.59 (I ² =0%)
Liver surgery	•	•	•		•	•	•	•	•
Gurusamy (2009)	Level I/II Good/Fair	1 RCT N=59	Adult patients undergoing liver resection	Hospital – planned surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion incidence (allogenic blood)	RR 0.58 (0.15, 2.21)	No difference 0.42	NA Phet=NA (I ² =NA)

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: ASA, acetylsalicylic acid; CABG, coronary artery bypass graft; CI, confidence interval; FFP, fresh frozen plasma; het, heterogeneity; IV, intravenous; OR, odds ratio; RBC, red blood cell; RCT, randomised controlled trial; RR, risk ratio.

a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I²<25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

^c Blood products include RBCs, FFP and platelets.

d US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland, UK.

º Cochrane ratings defined as follows: Grade A, adequate allocation concealment; Grade B, uncertain allocation concealment; Grade C, inadequate allocation concealment.

Key question(s): In patients undergoing surgery, what is the effect of administration of desmog	ress	in on transfusion volume?	Evidence table refa: POQ3.8d.P2
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)		
There is one pivotal Level I study (Crescenzi 2008/fair quality) which includes data from up to 11 RCTs, one supportive Level I study (Carless 2008/good quality) and one supportive level I/II study (Gurusamy 2009/good-fair quality) which includes data from one RCT in liver resection surgery.	A B	One or more level I studies with a low risk of bias or several level I One or two Level II studies with a low risk of bias or SR/several Level II studies with a low risk or SR/several Level II studies with a low risk or SR/several	vel III studies with a low risk of bias
	C D	One or two Level III studies with a low risk of bias or Level I or II st Level IV studies or Level I to III studies/SRs with a high risk of bias	
2 Consistancy (If only one study was qualished work this component as (not applicable))	D	Level IV Studies of Level I to III Studies/SRS with a high risk of blas	
2. Consistency (If only one study was available, rank this component as 'not applicable') Results consistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta-analyses described below. Pivotal evidence – Crescenzi 2008: Substantial heterogeneity (see note) in all analyses. May be due to inclusion of studies measuring transfusion in units and mL. Supportive evidence – Carless 2008: Moderate heterogeneity in analysis including all surgery and all patients. No	A B C	All studies consistent Most studies consistent and inconsistency can be explained Some inconsistency, reflecting genuine uncertainty around question Evidence is inconsistent	on
supportive evidence – Carless 2008: Moderate neterogeneity in analysis including all surgery and all patients. N significant heterogeneity seen in analysis of all surgery including transfused patients only. Moderate to substantial heterogeneity seen in most subgroup analyses which included all patients (ie, those who required transfusion and those who didn't).		Not applicable (one study only)	
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u>	nknow	\underline{n} factor and thus the clinical impact of the intervention could not be determ	nined)
Pivotal evidence – Crescenzi 2008 (based on SMD - not easy to interpret differences clinically) Any surgery (all patients) – SMD -0.29 (-0.52, -0.06); 34 RCTs (N=2065) Cardiac surgery (all patients) – SMD -0.22 (-0.52, 0.08); 23 RCTs (N=1607) Non-cardiac surgery (all patients) – SMD -0.45 (-0.77, -0.13); 11 RCTs (N=458) Supportive evidence – Carless 2008 (see Summary Table POQ3.l8d.P2 for all results) Any surgery (all patients) – WMD -0.30 (-0.60, -0.01); 14 RCTs (N=885) Any surgery (transfused patients) – WMD -0.49 (-0.94, -0.04); 5 RCTs (N=211)	A B C D	Very large Substantial Moderate Slight/Restricted	
4. Generalisability (How well does the body of evidence match the population and clinical set	ttings b	neing targeted by the Guideline?)	
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were to be included in the overall analysis although the majority of studies were in patients undergoing cardiac surgery (23/34 RCTs). There were also separate analyses of patients undergoing orthopaedic surgery and vascular surgery although these included few studies (2 RCTs each).		Evidence directly generalisable to target population Evidence directly generalisable to target population with some cave in the second directly generalisable to the target population but contained by the second directly generalisable to target population and hard the second directly generalisable to target population and hard the second directly generalisable to target population and hard the second directly generalisable to target population and hard the second directly generalisable to target population and hard the second directly generalisable to target population but contained the second directly generalisable to target population and hard the second directly generalisable to target population but contained the second directly generalisable to target population and hard the second directly generalisable to target population and hard the second directly generalisable to target population and hard the second directly generalisable to target population and hard the second directly generalisable to target population and hard the second directly generalisable to target population and hard the second directly generalisable to target population and hard the second directly generalisable to target population and hard the second directly generalisable to target population and the second directly generalisable to target population directly generalisable to target population dire	uld be sensibly applied
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of heal	th services/delivery of care and cultural factors?)	
Hospital setting. The pivotal review does not state which countries the RCTs were conducted in but the supportive study (which includes most of the studies included in the pivotal review) states that studies were conducted in US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK.	A B C	Evidence directly applicable to Australian healthcare context Evidence applicable to Australian healthcare context with few cave Evidence probably applicable to Australian healthcare context with	
	D	Evidence not applicable to Australian healthcare context	

Desmopressin injection currently listed by TGA for the following indication: "patients undergoing cardiac surgery with cardiopulmonary bypass for prosthetic valve replacement or aortocoronary bypass grafting, especially when it is complicated by platelet function defects sufficient to prolong bleeding time despite relatively normal platelet cover. Desmopressin acetate offers no benefit as routine therapy in patients having an uncomplicated (simple) cardiopulmonary bypass procedure".

Generalisability rated C as most studies in a specific surgical type but the evidence statement is applied to the general population.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (fair quality) and one supportive Level I study and one supportive level I/II study.
2. Consistency	С	There was some heterogeneity, particularly in the pivotal study. May be due to inclusion of different volume measures.
3. Clinical impact	В	There was a significant difference in a number of the main analyses and no difference in others.
4. Generalisability	С	The results are generalisable to an adult surgical population; in particular those undergoing cardiac surgery
5. Applicability	В	Studies were conducted in a wide range of countries. Likely to be applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, intravenous desmopressin therapy may reduce the volume of transfusion compared with no therapy.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I² <25%; (ii) mild heterogeneity if Phet<0.1 and I² <25%; moderate heterogeneity if P>0.1 and I² between 25-50%; substantial heterogeneity if P<0.1 and I² >50% Abbreviations: CI, confidence interval, RBC, red blood cell; SR, systematic review.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

a Interventions: 1 = ANH, 2 = intraoperative cell salvage, 3 = perioperative ANH and intraoperative cell salvage, 4 = postoperative cell salvage, 5 = deliberate induced hypotension, 6 = prevention of hypothermia, 7 = point-of-care testing for coagulation status and haemoglobin, 8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, 9 = appropriate patient positioning, 10 = PAD

^b Gurusamy included one study only which related to liver resection. This is described where appropriate in the Evidence Statement Summary Tables.

^c Publication dated 2004 but includes update to March 2008

POQ3.8d.P2 Characteristics and results of studies examining the effect of <u>desmopressin</u> on <u>transfusion volume</u>

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size	Surgical procedure				Risk estimate (95% CI)	Significance P-value	
ADULT POPULATION	/IV DESMOPRESSI	N							
Any surgery									
Crescenzi (2008)	Level I Fair	34 RCTs N=2065	Adult patients undergoing any surgery	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Transfusion volume ^c (blood products) Includes all patients	SMD -0.29 (-0.52, -0.06)	Favours DDAVP 0.01	Substantial Phet<0.001 (I ² =84.5%)
Carless (2008)	Level I Good	14 RCTs N=885	Adult patients undergoing any surgery	Hospital – elective or non-urgent surgery Various ^d	Desmopressin (IV) vs no desmopressin	Transfusion volume in units (RBCs) Includes all patients	WMD -0.30 (-0.60, -0.01)	Favours DDAVP 0.042	Moderate Phet=0.07 (I ² =39%)
		5 RCTs N=211	Adult patients undergoing any surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion volume in units (RBCs) Includes transfused patients only	WMD -0.49 (-0.94, -0.04)	Favours DDAVP 0.033	None Phet=0.49 (I ² =0%)
Carless (2008)	rless (2008) Level I Good	4 RCTs N=151	Adult patients undergoing any surgery in whom autologous techniques were used	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion volume in units (RBCs) Includes all patients	WMD -0.47 (-1.15, 0.20)	No difference 0.17	Substantial Phet=0.08 (I ² =56%)
		10 RCTs N=734	Adult patients undergoing any surgery in whom <u>no</u> autologous techniques were <u>used</u>	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion volume in units (RBCs) Includes all patients	WMD -0.22 (-0.55, 0.10)	No difference 0.18	Moderate Phet=0.19 (I ² =28%)
Cardiac surgery	1	•					•	•	
Crescenzi (2008)	Level I Fair	23 trials N=1607	Adult patients undergoing cardiac surgery	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Transfusion volume (blood products) Includes all patients	SMD -0.22 (-0.52, 0.08)	No difference 0.14	Substantial Phet<0.001 (l ² =87.8%)
Carless (2008)	Level I Good	10 RCTs N=621	Adult patients undergoing cardiac surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion volume in units (RBCs) Includes all patients	WMD -0.39 (-0.77, -0.01)	Favours DDAVP 0.047	Substantial Phet=0.03 (I ² =52%)
Non-cardiac surg	ery								•
Crescenzi (2008)	Level I Fair	11 RCTs N=458	Adults patients undergoing surgery other than cardiac	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Transfusion volume (blood products) Includes all patients	SMD -0.45 (-0.77, -0.13)	Favours DDAVP 0.006	Substantial Phet=0.003 (l²=62.4%)

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a <i>Quality</i>	sample size	Surgical procedure				Risk estimate (95% CI)	Significance P-value	
Orthopaedic surg	ery				•			•	
Carless (2008)	Level I Good	2 RCTs N=129	Adults patients undergoing orthopaedic surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion volume in units (RBCs) Includes all patients	WMD -0.15 (-0.64, 0.33)	No difference 0.54	None Phet=0.43 (I ² =0%)
Vascular surgery	•	•							
Carless (2008)	Level I Good	2 RCTs N=135	Adults patients undergoing vascular surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Transfusion volume in units (RBCs) Includes all patients	WMD 0.06 (-0.89, 1.02)	No difference 0.90	None Phet=0.40 (I ² =0%)
Liver surgery									
Gurusamy (2009)	Level I/II Good/Fair	1 RCT N=59	Adult patients undergoing liver resection	Hospital – planned surgery Unknown	Desmopressin (IV) vs placebo	Transfusion volume in units (RBCs) Includes all patients	SMD -0.31 (-0.82, 0.21)	No difference 0.24	NA Phet=NA (I2=NA)

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: CI, confidence interval; FFP, fresh frozen plasma; RBC, red blood cell; RCT, randomised controlled trial; SMD, standardised mean difference; WMD, weighted mean difference.

a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

^c Blood products include RBCs, FFP and platelets. Due to differences in the way this outcome was reported in individual RCTs, the analysis has been performed using the SMD.

d US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland, UK.

Key question(s):			Evidence table refa:						
In patients undergoing surgery, what is the effect of administration of desmo	<u>in</u> on <u>blood loss</u> ?	POQ3.8d.P3							
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	es)								
There is one pivotal Level I study (Crescenzi 2008/fair quality) which includes data from up to 11 RCTs, one	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias							
supportive Level I study (Carless 2008/good quality) and one supportive level I/II study (Gurusamy 2009/good-fair quality) which includes data from one RCT in liver resection surgery.	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias						
quality) initial initial acts and in the rest in invertes extended and invertes extended and in the rest in invertes extended and in the rest in	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias						
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	i						
2. Consistency (If only one study was available, rank this component as 'not applicable')									
Results consistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta-	А	All studies consistent							
analyses described below. Pivotal evidence – Crescenzi 2008	В	Most studies consistent and inconsistency can be explained							
Substantial heterogeneity (see note) in the analysis of all surgery types and cardiac surgery. No heterogeneity in	С	Some inconsistency, reflecting genuine uncertainty around questi	on						
analysis of non-cardiac surgery.	D	Evidence is inconsistent							
Supportive evidence – Carless 2008 (see Summary Table POQ3.l8d.P3) Moderate to substantial heterogeneity seen in most analyses and sub-analyses.	NA	Not applicable (one study only)							
3. Clinical impact (Indicate in the space below if the study results varied according to some	3. Clinical impact (Indicate in the space below if the study results varied according to some unknown factor and thus the clinical impact of the intervention could not be determined)								
Pivotal evidence – Crescenzi 2008 (based on SMD - not easy to interpret differences clinically)	Α	Very large							
Any surgery – SMD -0.20 (-0.34, -0.06); 40 RCTs (N=2445)	В	Substantial							
Cardiac surgery - SMD -0.23 (-0.40, -0.05); 29 RCTs (N=1928) Non-cardiac surgery - SMD -0.10 (-0.28, 0.07); 11 RCTs (N=517)	С	Moderate							
Supportive evidence – Carless 2008 (see Summary Table POQ3.l8d.P3 for all results)	D	Slight/Restricted							
Any surgery – WMD -92.98 (-149.86, -36.11); 18 RCTs (N=1201)									
Cardiac surgery – WMD -96.58 (-163.04, -30.12); 16 RCTs (N=1107)									
4. Generalisability (How well does the body of evidence match the population and clinical se	ettings b	peing targeted by the Guideline?)							
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were	Α	Evidence directly generalisable to target population							
included in the overall analysis although studies were predominantly in cardiac surgery (29/40 RCTs). There were also separate analyses of patients undergoing cardiac surgery who did or did not receive ASA.	В	Evidence directly generalisable to target population with some cav							
J	С	Evidence not directly generalisable to the target population but co	3 11						
	D	Evidence not directly generalisable to target population and hard t apply	o judge whether it is sensible to						
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of hear	Ith services/delivery of care and cultural factors?)							
Hospital setting.	Α	Evidence directly applicable to Australian healthcare context							
The pivotal review does not state which countries the RCTs were conducted in but the supportive study (which includes most of the studies included in the pivotal review) states that studies were conducted in US, Canada,	В	Evidence applicable to Australian healthcare context with few caveats							
Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK.	С	Evidence probably applicable to Australian healthcare context with some caveats							
	D	Evidence not applicable to Australian healthcare context							

Desmopressin injection currently listed by TGA for the following indication: "patients undergoing cardiac surgery with cardiopulmonary bypass for prosthetic valve replacement or aortocoronary bypass grafting, especially when it is complicated by platelet function defects sufficient to prolong bleeding time despite relatively normal platelet cover. Desmopressin acetate offers no benefit as routine therapy in patients having an uncomplicated (simple) cardiopulmonary bypass procedure".

In cardiac surgery, the difference was statistically significant but not clinically important.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Cardiac	Description
1. Evidence base	Α	There is one pivotal Level I study (fair quality) and one supportive Level I study and one supportive level I/II study.
2. Consistency	С	There was some heterogeneity, particularly in the pivotal study. May be due to different surgery types and blood loss measures used in different studies.
3. Clinical impact	D	There was a significant difference in most of the main analyses and no difference in some subgroup analyses
4. Generalisability	В	The results are generalisable to an adult surgical population; in particular those undergoing cardiac surgery
5. Applicability	В	Studies were conducted in a wide range of countries. Likely to be applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, intravenous desmopressin therapy reduces blood loss compared with no therapy.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I² <25%; (ii) mild heterogeneity if Phet<0.1 and I² <25%; moderate heterogeneity if P>0.1 and I² between 25-50%; substantial heterogeneity if P<0.1 and I² >50%. Abbreviations: CI, confidence interval, RBC, red blood cell: SR, systematic review.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

a Interventions: 1 = ANH, 2 = intraoperative cell salvage, 3 = perioperative ANH and intraoperative cell salvage, 4 = postoperative cell salvage, 5 = deliberate induced hypotension, 6 = prevention of hypothermia, 7 = point-of-care testing for coagulation status and haemoglobin, 8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, 9 = appropriate patient positioning, 10 = PAD

POQ3.8d.P3 Characteristics and results of studies examining the effect of <u>desmopressin</u> on <u>blood loss</u>

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size	Surgical procedure				Risk estimate (95% CI)	Significance P-value	
ADULT POPULATION	IC DESMOPRESSI	N							
Any surgery									
Crescenzi (2008)	Level I Fair	40 RCTs N=2445	Adult patients undergoing any surgery	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Blood loss (mL)	SMD -0.20 (-0.34, -0.06)	Favours DDAVP 0.004	Substantial Phet<0.001 (l ² =63.7%)
Carless (2008)	Level I Good	7 RCTs N=493	Adult patients undergoing any surgery	Hospital – elective or non-urgent surgery Various ^c	Desmopressin (IV) vs no desmopressin	Intraoperative blood loss (mL)	WMD -90.07 (-199.56, 19.42)	No difference 0.11	Moderate Phet=0.17 (I ² =34%)
		12 RCTs N=787	Adult patients undergoing any surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	0-24 hours postoperative blood loss (mL)	WMD –100.41 (-176.48, -24.34)	Favours DDAVP 0.0097	Substantial Phet=0.004 (l²=59%)
		18 RCTs N=1201	Adult patients undergoing any surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Postoperative blood loss (mL)	WMD -92.98 (-149.86, -36.11)	Favours DDAVP 0.0014	Substantial Phet=0.001 (l²=58%)
		10 RCTs N=669	Adult patients undergoing any surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Intraoperative + postoperative blood loss(mL)	WMD -241.78 (-387.55, -96.01)	Favours DDAVP 0.0012	Substantial Phet=0.002 (l²=66%)
Cardiac surgery									
Crescenzi (2008)	Level I Fair	29 trials N=1928	Adult patients undergoing cardiac surgery	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Blood loss (mL)	SMD -0.23 (-0.40, -0.05)	Favours DDAVP 0.01	Substantial Phet<0.001 (l ² =71.0%)

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size	Surgical procedure				Risk estimate (95% CI)	Significance P-value	
Carless (2008)	Level I Good	3 RCTs N=229	Adult patients undergoing cardiac surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Intraoperative blood loss (mL)	WMD -119.79 (-314.57, 75.00)	No difference 0.23	Substantial Phet=0.06 (I ² =65%)
		1 RCT N=59	Adult patients undergoing cardiac surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	0-6 hours postoperative blood loss (mL)	WMD -98.00 (-304.99, 108.99)	No difference 0.35	NA Phet=NA (l²=NA)
		3 RCTs N=233	Adult patients undergoing cardiac surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	0-12 hours postoperative blood loss (mL)	WMD -114.05 (-269.46, 41.36)	No difference 0.15	Substantial Phet=0.004 (12=82%)
		2 RCTs N=122	Adult patients undergoing cardiac surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	0-16 hours postoperative blood loss (mL)	WMD –18.01 (-113.34, 77.32)	No difference 0.71	None Phet=0.42 (l²=0%)
		10 RCTs N=693	Adult patients undergoing cardiac surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	0-24 hours postoperative blood loss (mL)	WMD -107.46 (-207.12, -7.80)	Favours DDAVP 0.035	Substantial Phet=0.002 (l²=65%)
		16 RCTs N=1107	Adult patients undergoing cardiac surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Postoperative blood loss (mL)	WMD -96.58 (-163.04, -30.12)	Favours DDAVP 0.0044	Substantial Phet<0.001 (l ² =62%)
		7 RCTs N=496	Adult patients undergoing cardiac surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Intraoperative + postoperative blood loss (mL)	WMD -237.92 (-413.43, -62.40)	Favours DDAVP 0.0079	Substantial Phet<0.001 (l ² =74%)
Carless (2008)	Level I Good	10 RCTs N=633	Adult patients undergoing cardiac surgery with ASA use	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Postoperative blood loss (mL)	WMD -109.57 (-200.11, -19.03)	Favours DDAVP 0.018	Substantial Phet=0.01 (I ² =60%)
		3 RCTs N=221	Adult patients undergoing cardiac surgery without ASA use	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Postoperative blood loss (mL)	WMD -112.69 (-227.59, 2.22)	No difference 0.055	Moderate Phet=0.16 (I ² =45%)

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	3			Risk estimate (95% CI)	Significance P-value			
Carless (2008)	Level I Good	3 RCTs N=198	Adult patients undergoing cardiac surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Postoperative blood loss after <u>CPB < 80 mins</u> (mL)	WMD -41.22 (-157.25, 74.80)	No difference 0.49	Substantial Phet=0.07 (I ² =62%)
		5 RCTs N=330	Adult patients undergoing cardiac surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Postoperative blood loss after <u>CPB 80 – 100 mins</u> (mL)	WMD -104.18 (-184.75, -23.61)	Favours DDAVP 0.011	Moderate Phet=0.21 (I ² =31%)
		3 RCTs N=129	Adult patients undergoing cardiac surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Postoperative blood loss after <u>CPB 101 – 120</u> <u>mins</u> (mL)	WMD 53.08 (-156.33, 262.50)	No difference 0.62	Moderate Phet=0.15 (I ² =47%)
		2 RCTs N=196	Adult patients undergoing cardiac surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Postoperative blood loss after <u>CPB 121 – 140</u> <u>mins</u> (mL)	WMD -46.53 (-366.29, 273.23)	No difference 0.78	Substantial Phet=0.01 (I ² =84%)
		2 RCTs N=171	Adult patients undergoing cardiac surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Postoperative blood loss after CPB > 140 mins (mL)	WMD -344.74 (-478.50, -210.97)	Favours DDAVP <0.001	None Phet=0.42 (l²=0%)
		15 RCTs N=1024	Adult patients undergoing cardiac surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Postoperative blood loss after CPB of any duration (mL)	WMD -93.34 (-162.24, -24.44)	Favours DDAVP 0.0079	Substantial Phet<0.001 (P=64%)
Non-cardiac surge	ery	•					<u> </u>	•	
Crescenzi (2008)	Level I Fair	11 RCTs N=517	Adults patients undergoing surgery other than cardiac	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Blood loss (mL)	SMD -0.10 (-0.28, 0.07)	No difference 0.25	None Phet=0.45 (I ² =0%)
Liver surgery									
Gurusamy (2009)	Level I/II Good/Fair	1 RCT N=97	Adult patients undergoing liver resection	Hospital – planned surgery Various	Desmopressin (IV) vs no desmopressin	Operative blood loss (mL)	MD 32.50 (-695.69, 760.69)	No difference 0.93	NA Phet=NA (I²=NA)

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: ASA, acetylsalicylic acid; CI, confidence interval; CPB, cardiopulmonary bypass; MD, mean difference; NA, not applicable; RBC, red blood cell; RCT, randomised controlled trial; SMD, standardised mean difference; WMD, weighted mean difference

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (iii) mild heterogeneity if I²<25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

^cUS, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland, UK.

Key question(s):		Evidence table refa:				
In patients undergoing surgery, what is the effect of administration of desmo	<u>in</u> on <u>mortality</u> ?	POQ3.l8d.P4				
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	es)					
There is one pivotal Level I study (Crescenzi 2008/fair quality) which includes data from up to 8 RCTs and one supportive Level I study (Carless 2008/good quality).		One or more level I studies with a low risk of bias or several level	I studies with a low risk of bias			
supportive Lever's study (Carress 2000/good quality).	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Results consistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta- analyses described below.	Α	All studies consistent				
Pivotal evidence – Crescenzi 2008	В	Most studies consistent and inconsistency can be explained				
No heterogeneity in analyses of any surgery or cardiac surgery (non-cardiac surgery analysis includes data from only 1 RCT)	С	Some inconsistency, reflecting genuine uncertainty around question	on			
Supportive evidence – Carless 2008	D	Evidence is inconsistent				
No heterogeneity in analyses of any surgery.	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some undetermined)	unknov	<u>rn</u> factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be			
Pivotal evidence – Crescenzi 2008	А	Very large				
Any surgery - 1.2% vs 0.9%; OR 1.25 (0.51, 3.04); 8 RCTs (N=673) Cardiac surgery - 1.0% vs 1.1%; OR 1.00 (0.38, 2.62); 7 RCTs (N=582)	В	Substantial				
Non-cardiac surgery – 2.1% vs 0%; OR 5.84 (0.27, 125.19); 1 RCT (N=91)	С	Moderate				
Supportive evidence – Carless 2008 Any surgery – 2.4% vs 1.3%; RR 1.72 (0.68, 4.33); 8 RCTs (N=774)	D	No difference				
4. Generalisability (How well does the body of evidence match the population and clinical se	ettings	being targeted by the Guideline?)				
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were included in the overall analysis although studies were predominantly in cardiac surgery (7/8 RCTs). There were als	Α	Evidence directly generalisable to target population				
separate analyses of patients undergoing cardiac surgery or non-cardiac surgery. However, the non-cardiac	В	Evidence directly generalisable to target population with some cav	eats			
surgery analysis includes data from only one RCT in vascular surgery.	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to judge whether it is sensible to				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of hea	Ith services/delivery of care and cultural factors?)				
Hospital setting. The pivotal review does not state which countries the RCTs were conducted in but the supportive study (which	Α	Evidence directly applicable to Australian healthcare context				
includes most of the studies included in the pivotal review) states that studies were conducted in US, Canada,	В	Evidence applicable to Australian healthcare context with few caveats				
Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK.	С	Evidence probably applicable to Australian healthcare context with some caveats				
	D	Evidence not applicable to Australian healthcare context				

Desmopressin injection currently listed by TGA for the following indication: "patients undergoing cardiac surgery with cardiopulmonary bypass for prosthetic valve replacement or aortocoronary bypass grafting, especially when it is complicated by platelet function defects sufficient to prolong bleeding time despite relatively normal platelet cover. Desmopressin acetate offers no benefit as routine therapy in patients having an uncomplicated (simple) cardiopulmonary bypass procedure".

Generalisability rated C as most studies in a specific surgical type but the evidence statement is applied to the general population.

The CRG noted that the point estimate indicated a potentially increased risk of mortality, although the evidence is underpowered to show a difference (difference not statistically significant).

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (fair quality) and one supportive Level I study.
2. Consistency	Α	There was no heterogeneity between studies
3. Clinical impact	D	There was no significant difference in all analyses. The studies are likely underpowered to detect a difference in mortality.
4. Generalisability	С	The results are generalisable to an adult surgical population; in particular those undergoing cardiac surgery
5. Applicability	В	Studies were conducted in a range of countries. Likely to be applicable to the Australian setting.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intravenous desmopressin therapy on mortality, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I² <25%; moderate heterogeneity if P>0.1 and I² between 25-50%; substantial heterogeneity if P<0.1 and I² >50%. Abbreviations: CI, confidence interval, Pot, potential; RBC, red blood cell; SR, systematic review.

^a Interventions: 1 = ANH, 2 = intraoperative cell salvage, 3 = perioperative ANH and intraoperative cell salvage, 4 = postoperative cell salvage, 5 = deliberate induced hypotension, 6 = prevention of hypothermia, 7 = point-of-care testing for coagulation status and haemoglobin, 8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, 9 = appropriate patient positioning, 10 = PAD

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

POQ3.8d.P4 Characteristics and results of studies examining the effect of <u>desmopressin</u> on <u>mortality</u>

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size	Surgical procedure				Risk estimate (95% CI)	Significance P-value	
ADULT POPULATION	IV DESMOPRESSI	N							
Any surgery									
Crescenzi (2008)	Level I Fair	8 RCTs N=673	Adult patients undergoing any surgery	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Mortality	OR 1.25 (0.51, 3.04)	No difference 0.63	None Phet=0.76 (I ² =0%)
Carless (2008)	Level I Good	8 RCTs N=774	Adult patients undergoing any surgery	Hospital – elective or non-urgent surgery Various ^c	Desmopressin (IV) vs no desmopressin	Mortality	RR 1.72 (0.68, 4.33)	No difference 0.25	None Phet=0.80 (I ² =0%)
Cardiac surgery	•								
Crescenzi (2008)	Level I Fair	7 trials N=582	Adult patients undergoing cardiac surgery	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Mortality	OR 1.00 (0.38, 2.62)	No difference 1.00	None Phet=0.81 (I ² =0%)
Non-cardiac surge	ery							•	
Crescenzi (2008)	Level I/II Fair/ Unknown	1 RCT N=91	Adults patients undergoing surgery other than cardiac	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Mortality	OR 5.84 (0.27, 125.19)	No difference 0.26	NA Phet=NA (I ² =NA)

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: CI, confidence interval; NA, not applicable; OR, odds ratio; RCT, randomised controlled trial; RR, risk ratio.

a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I²<25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

^c US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland, UK.

Key question(s):			Evidence table refa:			
In patients undergoing surgery, what is the effect of administration of desmor	ress	in on morbidity (hypotension)?	POQ3.I8d.P5			
			(hypotension)			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)					
There is one pivotal Level I study (Crescenzi 2008/fair quality) including data from up to 13 RCTs and one	Α	One or more level I studies with a low rik of bias or several level II	studies with a low risk of bias			
supportive Level I study (Carless 2008/good quality).	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Results consistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta-	Α	All studies consistent				
analyses described below. Pivotal evidence – Crescenzi 2008	В	Most studies consistent and inconsistency can be explained				
No heterogeneity in analyses of any surgery, cardiac surgery or non-cardiac surgery	С	Some inconsistency, reflecting genuine uncertainty around question	on			
Supportive evidence – Carless 2008	D	Evidence is inconsistent				
No heterogeneity in analyses of any surgery.	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u> determined)	<u>nknow</u>	n factor (not simply study quality or sample size) and thus the clinical impa	nct of the intervention could not be			
Pivotal evidence – Crescenzi 2008	Α	Very large				
Any surgery - 8.2% vs 2.1%; OR 4.84 (2.31, 10.13); 7 RCTs (N=320) Cardiac surgery - 5.2% vs 0.3%; OR 8.92 (2.54, 31.37); 5 RCTs (N=221)	В	Substantial				
Non-cardiac surgery = 3.2% vs 0.3%; OR 8.92 (2.34, 31.37); 5 RCTs (N=221)	С	Moderate				
Supportive evidence – Carless 2008	D	Slight/Restricted				
Any surgery – 37.0% vs 9.9%; RR 2.81 (1.50, 5.27); 5 RCTs (N=183)						
4. Generalisability (How well does the body of evidence match the population and clinical se	ttings l	being targeted by the Guideline?)				
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were	Α	Evidence directly generalisable to target population				
included in the overall analysis although studies were predominantly in cardiac surgery (5/7 RCTs). There were also separate analyses of patients undergoing cardiac surgery or non-cardiac surgery.	В	Evidence directly generalisable to target population with some cav	eats			
populate unarjood of patients undergoing salado salgor) of non-salado salgor).	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard tapply	o judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of hea	Ith services/delivery of care and cultural factors?)				
Hospital setting.	Α	Evidence directly applicable to Australian healthcare context				
The pivotal review does not state which countries the RCTs were conducted in but the supportive study (which	В	Evidence applicable to Australian healthcare context with few cave	eats			
includes most of the studies included in the pivotal review) states that studies were conducted in US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK.	С	Evidence probably applicable to Australian healthcare context with some caveats				
Spain, Shoush, Somany, Turkey, Islam, Shina, Hormay, Finland and Orc.		Evidence not applicable to Australian healthcare context				

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Desmopressin injection currently listed by TGA for the following indication: "patients undergoing cardiac surgery with cardiopulmonary bypass for prosthetic valve replacement or aortocoronary bypass grafting, especially when it is complicated by platelet function defects sufficient to prolong bleeding time despite relatively normal platelet cover. Desmopressin acetate offers no benefit as routine therapy in patients having an uncomplicated (simple) cardiopulmonary bypass procedure".

Generalisability rated C as most studies in a specific surgical type but the evidence statement is applied to the general population.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (fair quality) and one supportive Level I study.
2. Consistency	Α	There was no heterogeneity in the Level I studies and the two Level I studies showed consistent results.
3. Clinical impact	D	There was a significant risk of hypotension for desmopressin therapy compared with no therapy but the authors note that this is transient and mild.
4. Generalisability	С	The results are generalisable to an adult surgical population; in particular those undergoing cardiac surgery
5. Applicability	В	Studies were conducted in a range of countries. Likely to be applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, intravenous desmopressin therapy increases the risk of mild and transient hypotension compared with no therapy.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I² <25%; (ii) mild heterogeneity if Phet<0.1 and I² <25%; moderate heterogeneity if P>0.1 and I² between 25-50%; substantial heterogeneity if P<0.1 and I² >50%. Abbreviations: CI, confidence interval, RBC, red blood cell; SR, systematic review.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

a Interventions: 1 = ANH, 2 = intraoperative cell salvage, 3 = perioperative ANH and intraoperative cell salvage, 4 = postoperative cell salvage, 5 = deliberate induced hypotension, 6 = prevention of hypothermia, 7 = point-of-care testing for coagulation status and haemoglobin, 8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, 9 = appropriate patient positioning, 10 = PAD

POQ3.l8d.P5 (hypotension) Characteristics and results of studies examining the effect of <u>desmopressin</u> on <u>morbidity (hypotension)</u>

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence ^a Quality	sample size	Surgical procedure				Risk estimate (95% CI)	Significance P-value	
ADULT POPULATION/	IV DESMOPRESSIN	ı							
Any surgery									
Crescenzi (2008)	Level I Fair	7 RCTs N=320	Adult patients undergoing any surgery	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Hypotension	OR 4.84 (2.31, 10.13)	Favours no desmopressin <0.001	None Phet=0.85 (l²=0%)
Carless (2008)	Level I Good	5 RCTs N=183	Adult patients undergoing any surgery	Hospital – elective or non-urgent surgery Various ^c	Desmopressin (IV) vs no desmopressin	Hypotension during infusion requiring treatment (fluids and/or vasoactive drugs)	RR 2.81 (1.50, 5.27)	Favours no desmopressin 0.0013	None Phet=0.50 (I ² =0%)
Cardiac surgery									
Crescenzi (2008)	Level I Fair	5 RCTs N=221	Adult patients undergoing cardiac surgery	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Hypotension	OR 8.92 (2.54, 31.37)	Favours no desmopressin <0.001	None Phet=0.94 (I ² =0%)
Non-cardiac surge	ry								
Crescenzi (2008)	Level I Fair	2 RCTs N=99	Adults patients undergoing surgery other than cardiac	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Hypotension	OR 3.04 (1.18, 7.87)	Favours no desmopressin 0.02	None Phet=0.64 (l²=0%)

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: CI, confidence interval; NA, not applicable; OR, odds ratio; RCT, randomised controlled trial; RR, risk ratio.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and |2<25%; (ii) mild heterogeneity if |2 <25%; moderate heterogeneity if |2 between 25-50%; substantial heterogeneity |2 >50%.

^c US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland, UK.

Key question(s): In patients undergoing surgery, what is the effect of administration of desmor	Evidence table refa: POQ3.I8d.P5 (MI)					
1. Evidence base (number of studies, level of evidence and risk of bias in the included studie	s)					
There is one pivotal Level I study (Crescenzi 2008/fair quality) which includes data from up to 13 RCTs and one	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias			
supportive Level I study (Carless 2008/good quality).	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Results consistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta-	Α	All studies consistent				
analyses described below. Pivotal evidence – Crescenzi 2008	В	Most studies consistent and inconsistency can be explained				
No heterogeneity in analyses of any surgery, cardiac surgery or non-cardiac surgery. Difference in direction of effec	t C	Some inconsistency, reflecting genuine uncertainty around question	on			
for different surgery types.	D	Evidence is inconsistent				
Supportive evidence – Carless 2008 No heterogeneity in analyses of any surgery.	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u> determined) Pivotal evidence – Crescenzi 2008	inknow A	n factor (not simply study quality or sample size) and thus the clinical impa	nct of the intervention could not be			
Any surgery – 3.8% vs 2.9%; OR 1.27 (0.73, 2.20); 13RCTs (N=916)	В	Substantial				
Cardiac surgery – 4.3% vs 3.1%; OR 1.36 (0.75, 2.48); 11 RCTs (N=775)	C	Moderate				
Non-cardiac surgery – 1.8% vs 2.2%; OR 0.84 (0.20,3.53); 2 RCTs (N=141) Supportive evidence – Carless 2008 Any surgery – 6.3% vs 4.1%; RR 1.38 (0.77, 2.50); 9 RCTs (N=731)	D	Underpowered				
4. Generalisability (How well does the body of evidence match the population and clinical se	ttings l	being targeted by the Guideline?)				
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were	Α	Evidence directly generalisable to target population				
included in the overall analysis although studies were predominantly in cardiac surgery (11/13 RCTs). There were also separate analyses of patients undergoing cardiac surgery or non-cardiac surgery.	В	Evidence directly generalisable to target population with some cav	eats			
and soparate analysis of patients analysing saratas sargery or non-saratas sargery.	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard tapply	o judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of hea	lth services/delivery of care and cultural factors?)				
Hospital setting.	Α	Evidence directly applicable to Australian healthcare context				
The pivotal review does not state which countries the RCTs were conducted in but the supportive study (which includes most of the studies included in the pivotal review) states that studies were conducted in US, Canada,	В	Evidence applicable to Australian healthcare context with few cave	eats			
Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK.	С	Evidence probably applicable to Australian healthcare context with	some caveats			
	D	Evidence not applicable to Australian healthcare context				

Desmopressin injection currently listed by TGA for the following indication: "patients undergoing cardiac surgery with cardiopulmonary bypass for prosthetic valve replacement or aortocoronary bypass grafting, especially when it is complicated by platelet function defects sufficient to prolong bleeding time despite relatively normal platelet cover. Desmopressin acetate offers no benefit as routine therapy in patients having an uncomplicated (simple) cardiopulmonary bypass procedure".

Generalisability rated C as most studies in a specific surgical type but the evidence statement is applied to the general population.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
Evidence base	Α	There is one pivotal Level I study (fair quality) and one supportive Level I study.
2. Consistency	В	There was no heterogeneity in the two included Level I studies. There was a difference in the direction of effect for cardiac surgery versus non-cardiac surgery.
3. Clinical impact	D	Results show an slightly increased risk with no significant difference (28/441 versus 19/435 in any surgery) but studies likely underpowered to detect a difference in myocardial infarction.
4. Generalisability	С	The results are generalisable to an adult surgical population; in particular those undergoing cardiac surgery
5. Applicability	В	Studies were conducted in a range of countries. Likely to be applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intravenous desmopressin therapy on risk of myocardial infarction, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I² <25%; (ii) mild heterogeneity if Phet<0.1 and I² <25%; moderate heterogeneity if P>0.1 and I² between 25-50%; substantial heterogeneity if P<0.1 and I² >50%.

a Interventions: 1 = ANH, 2 = intraoperative cell salvage, 3 = perioperative ANH and intraoperative cell salvage, 4 = postoperative cell salvage, 5 = deliberate induced hypotension, 6 = prevention of hypothermia, 7 = point-of-care testing for coagulation status and haemoglobin, 8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, 9 = appropriate patient positioning, 10 = PAD

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

POQ3.l8d.P5 (MI) Characteristics and results of studies examining the effect of <u>desmopressin</u> on <u>morbidity (myocardial infarction)</u>

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence Quality	sample size	Surgical procedure				Risk estimate (95% CI)	Significance P-value	
ADULT POPULATION	IV DESMOPRESSI	N							
Any surgery									
Crescenzi (2008)	Level I Fair	13 RCTs N=916	Adult patients undergoing any surgery	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Myocardial infarction	OR 1.27 (0.73, 2.20)	No difference 0.40	None Phet=0.88 (I ² =0%)
Carless (2008)	Level I Good	9 RCTs N=731	Adult patients undergoing any surgery	Hospital – elective or non-urgent surgery Various ^c	Desmopressin (IV) vs no desmopressin	Myocardial infarction	RR 1.38 (0.77, 2.50)	No difference 0.28	None Phet=0.87 (l ² =0%)
Cardiac surgery	•								
Crescenzi (2008)	Level I Fair	11 RCTs N=775	Adult patients undergoing cardiac surgery	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Myocardial infarction	OR 1.36 (0.75, 2.48)	No difference 0.31	None Phet=0.86 (l²=0%)
Non-cardiac surg	ery							•	•
Crescenzi (2008)	Level I Fair	2 RCTs N=141	Adults patients undergoing surgery other than cardiac	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Myocardial infarction	OR 0.84 (0.20,3.53)	No difference 0.81	None Phet=0.35 (I ² =0%)

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: CI, confidence interval; NA, not applicable; OR, odds ratio; RCT, randomised controlled trial; RR, risk ratio.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and |2<25%; (ii) mild heterogeneity if |2 <25%; moderate heterogeneity if |2 between 25-50%; substantial heterogeneity |2 >50%.

^c US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland, UK.

Key question(s): In patients undergoing surgery, what is the effect of administration of desmop	oress	in on morbidity (stroke)?	Evidence table refa: POQ3.I8d.P5 (stroke)			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)					
There is one supportive Level I study (Carless 2008/good quality) which includes data from seven RCTs.	А	One or more level I studies with a low risk of bias or several level II studies with a low risk of				
		One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias			
	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Supportive evidence – Carless 2008 No heterogeneity in analyses of any surgery.	Α	All studies consistent				
no neterogeneity in analyses of any surgery.	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around question				
	D	Evidence is inconsistent				
	NA	A Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some undetermined)	nknow	<u>n</u> factor (not simply study quality or sample size) and thus the clinical impa	act of the intervention could not be			
Supportive evidence – Carless 2008	Α	Very large				
Any surgery – 4.3% vs 1.1%; RR 2.40 (0.68, 8.43); 7 RCTs (N=591)	В	Substantial (potential)				
	С	Moderate				
	D	Slight/Restricted				
4. Generalisability (How well does the body of evidence match the population and clinical set	ttings b	peing targeted by the Guideline?)				
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were included in the overall analysis although studies were predominantly in cardiac surgery.	Α	Evidence directly generalisable to target population				
included in the overall analysis although studies were predominantly in cardiac surgery.	В	Evidence directly generalisable to target population with some cav	eats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to judge whether it is sensible to				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of heal					
Hospital setting. The Carless 2008 study states that studies were conducted in US, Canada, Spain, Sweden, Germany, Turkey,	Α	Evidence directly applicable to Australian healthcare context				
Israel, China, Norway, Finland and UK.	В	Evidence applicable to Australian healthcare context with few caveats				
	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
	D	Evidence not applicable to Australian healthcare context				

Desmopressin injection currently listed by TGA for the following indication: "patients undergoing cardiac surgery with cardiopulmonary bypass for prosthetic valve replacement or aortocoronary bypass grafting, especially when it is complicated by platelet function defects sufficient to prolong bleeding time despite relatively normal platelet cover. Desmopressin acetate offers no benefit as routine therapy in patients having an uncomplicated (simple) cardiopulmonary bypass procedure".

Generalisability rated C as most studies in a specific surgical type but the evidence statement is applied to the general population.

The CRG noted that the point estimate indicated a potentially increased risk of stroke, although the evidence is underpowered to show a difference (difference not statistically significant).

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one supportive Level I study (good quality) which includes data from seven RCTs.
Consistency A There was no heterogeneity between studies		There was no heterogeneity between studies
3. Clinical impact	D	There was no significant difference in the analysis. The studies are likely underpowered to detect a difference in risk of stroke.
4. Generalisability	С	The results are generalisable to an adult surgical population; in particular those undergoing cardiac surgery
5. Applicability	В	Studies were conducted in a range of countries. Likely to be applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intravenous desmopressin therapy on risk of stroke, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I² <25%; moderate heterogeneity if P>0.1 and I² between 25-50%; substantial heterogeneity if P<0.1 and I² >50%. Abbreviations: CI, confidence interval, Pot, potential; SR, systematic review.

a Interventions: 1 = ANH, 2 = intraoperative cell salvage, 3 = perioperative ANH and intraoperative cell salvage, 4 = postoperative cell salvage, 5 = deliberate induced hypotension, 6 = prevention of hypothermia, 7 = point-of-care testing for coagulation status and haemoglobin, 8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, 9 = appropriate patient positioning, 10 = PAD

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

POQ3.l8d.P5 (stroke) Characteristics and results of studies examining the effect of <u>desmopressin</u> on <u>morbidity</u> (stroke)

Study	Level of	No. of trials /	Patient population /	Setting	Intervention Outcome		Results	Heterogeneity ^b	
	evidence ^a Quality	sample size	Surgical procedure				Risk estimate (95% CI)	Significance P-value	
Adult population/IV desmopressin									
Any surgery									
Carless (2008)	Level I Good	7 RCTs N=591	Adult patients undergoing any surgery	Hospital – elective or non-urgent surgery Various ^c	Desmopressin (IV) vs no desmopressin	Stroke	RR 2.40 (0.68, 8.43)	No difference 0.17	None Phet=0.90 (I ² =0%)

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: CI, confidence interval; NA, not applicable; OR, odds ratio; RCT, randomised controlled trial; RR, risk ratio.

a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review. b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

CUS, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland, UK.

Key question(s):			Evidence table refa:		
In patients undergoing surgery, what is the effect of administration of desmop	POQ3.I8d.P5				
			(thrombosis)		
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	c)		(till officeolo)		
There is one pivotal Level I study (Crescenzi 2008/fair quality) which includes data from up to 14 RCTs and one		O			
supportive Level I study (Carless 2008/good quality).	A	One or more level I studies with a low risk of bias or several level II			
	В	One or two Level II studies with a low risk of bias or SR/several Lev			
	С	One or two Level III studies with a low risk of bias or Level I or II stu	idies with a moderate risk of bias		
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias			
2. Consistency (If only one study was available, rank this component as 'not applicable')					
Results consistent between pivotal and supportive meta-analyses. Consistency of individual studies within meta-	Α	All studies consistent			
analyses described below. Pivotal evidence – Crescenzi 2008	В	Most studies consistent and inconsistency can be explained			
No heterogeneity in analyses of any surgery or cardiac surgery. Moderate heterogeneity in analysis of non-cardiac	С	Some inconsistency, reflecting genuine uncertainty around questic	n		
surgery (3 RCTs only)	D	Evidence is inconsistent			
Supportive evidence – Carless 2008	NA	Not applicable (one study only)			
No heterogeneity in analyses of any surgery.					
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>un</u> determined)	<u>nknow.</u>	<u>n</u> factor (not simply study quality or sample size) and thus the clinical impa	ct of the intervention could not be		
Pivotal evidence – Crescenzi 2008	Α	Very large			
Any surgery – 2.9% vs 2.5%; OR 1.20 (0.68, 2.09); 14 RCTs (N=1151)	В	Substantial			
Cardiac surgery – 2.5% vs 2.0%; OR 1.27 (0.64, 2.50); 11 RCTs (N=931) Non-cardiac surgery – 4.4% vs 4.1%; OR 1.06 (0.39,2.84); 3 RCTs (N=220)	С	Moderate (potential)			
Supportive evidence – Carless 2008	D	Slight/restricted			
Any surgery – 3.9% vs 3.0%; RR 1.46 (0.64, 3.35); 7 RCTs (N=591)					
4. Generalisability (How well does the body of evidence match the population and clinical set	ttings Ł	eing targeted by the Guideline?)			
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were	Α	Evidence directly generalisable to target population			
included in the overall analysis although studies were predominantly in cardiac surgery (11/14 RCTs). There were also separate analyses of patients undergoing cardiac surgery or non-cardiac surgery.	В	Evidence directly generalisable to target population with some cave	eats		
also separate analyses of patients undergoing cardiac surgery of non-cardiac surgery.	С	Evidence not directly generalisable to the target population but cou	ıld be sensibly applied		
		Evidence not directly generalisable to target population and hard to apply	judge whether it is sensible to		
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of heal	th services/delivery of care and cultural factors?)			
Hospital setting.	Α	Evidence directly applicable to Australian healthcare context			
The pivotal review does not state which countries the RCTs were conducted in but the supportive study (which	В	Evidence applicable to Australian healthcare context with few cave	ats		
includes most of the studies included in the pivotal review) states that studies were conducted in US, Čanada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK.	С	Evidence probably applicable to Australian healthcare context with some caveats			
Spani, Strady, Tandy, Island, Olina, Hornay, Filliana and Oli.	D	Evidence not applicable to Australian healthcare context			
		1			

Desmopressin injection currently listed by TGA for the following indication: "patients undergoing cardiac surgery with cardiopulmonary bypass for prosthetic valve replacement or aortocoronary bypass grafting, especially when it is complicated by platelet function defects sufficient to prolong bleeding time despite relatively normal platelet cover. Desmopressin acetate offers no benefit as routine therapy in patients having an uncomplicated (simple) cardiopulmonary bypass procedure".

Generalisability rated C as most studies in a specific surgical type but the evidence statement is applied to the general population.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description				
1. Evidence base	А	There is one pivotal Level I study (fair quality) and one supportive Level I study.				
2. Consistency A There		was no heterogeneity in the Level I studies and consistent results between the two studies.				
3. Clinical impact	D	Results show a slightly increased risk with no significant difference (14/361 versus 10/330 in any surgery) but studies likely underpowered to detect a difference in thrombosis.				
4. Generalisability	С	The results are generalisable to an adult surgical population; in particular those undergoing cardiac surgery				
5. Applicability	В	Studies were conducted in a range of countries. Likely to be applicable to the Australian setting				

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of intravenous desmopressin therapy on risk of thrombosis, compared with no therapy, is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I² >50%; (ii) mild heterogeneity if Phet<0.1 and I² >50%. Abbreviations: CI, confidence interval, SR, systematic review.

a Interventions: 1 = ANH, 2 = intraoperative cell salvage, 3 = perioperative ANH and intraoperative cell salvage, 4 = postoperative cell salvage, 5 = deliberate induced hypotension, 6 = prevention of hypothermia, 7 = point-of-care testing for coagulation status and haemoglobin, 8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, 9 = appropriate patient positioning, 10 = PAD

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

POQ3.l8d.P5 (thrombosis) Characteristics and results of studies examining the effect of <u>desmopressin</u> on <u>morbidity (thrombosis)</u>

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneityb
	evidence ^a sample size Surgical procedure Quality					Risk estimate (95% CI)	Significance P-value		
ADULT POPULATION	IV DESMOPRESSI	N							
Any surgery									
Crescenzi (2008)	Level I Fair	14 RCTs N=1151	Adult patients undergoing any surgery	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Thromboses (other than myocardial infarction)	OR 1.20 (0.68, 2.09)	No difference 0.53	None Phet=0.82 (I ² =0%)
Carless (2008)	Level I Good	7 RCTs N=591	Adult patients undergoing any surgery	Hospital – elective or non-urgent surgery Various ^c	Desmopressin (IV) vs no desmopressin	Any thrombosis	RR 1.46 (0.64, 3.35)	No difference 0.37	None Phet=0.78 (I ² =0%)
Cardiac surgery	•								
Crescenzi (2008)	Level I Fair	11 RCTs N=931	Adult patients undergoing cardiac surgery	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Thromboses (other than myocardial infarction)	OR 1.27 (0.64, 2.50)	No difference 0.49	None Phet=0.86 (I ² =0%)
Non-cardiac surg	ery								
Crescenzi (2008)	Level I Fair	3 RCTs N=220	Adults patients undergoing surgery other than cardiac	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Thromboses (other than myocardial infarction)	OR 1.06 (0.39,2.84)	No difference 0.92	Moderate Phet=0.24 (I ² =30.2%)

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: CI, confidence interval; NA, not applicable; OR, odds ratio; RCT, randomised controlled trial; RR, risk ratio.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I²<25%; (ii) mild heterogeneity if I² <25%; moderate heterogeneity if I² between 25-50%; substantial heterogeneity I² >50%.

^cUS, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland, UK.

In patients undergoing surgery, what is the effect of administration of desmorps:ssin on quality of life? 1. Evidence base (number of studies, level of evidence and risk of bias in the included studies) 1. Evidence base (number of studies, level of evidence and risk of bias in the included studies) 1. Evidence base (number of studies, level of evidence and risk of bias in the included studies) 2. Consistency (if unity one study was available, rank this component as not applicable) 1. Evidence I studies with a low risk of bias or several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias	Key question(s):			Evidence table refa:				
No studies of any level were identified which assessed the effect of desrropressin on quality of Ille A One or more level I studies with a low risk of bias or several level III studies with a low risk of bias or SR/several Level III studies with	In patients undergoing surgery, what is the effect of administration of desmop	ressi	in on quality of life?	POQ3.I8d.P6				
No studies of any level were identified which assessed the effect of desrropressin on quality of Ille A One or more level I studies with a low risk of bias or several level III studies with a low risk of bias or SR/several Level III studies with	1 Evidence has a forumber of studies level of evidence and rick of bias in the included studies)							
B One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias or Level III studies with a low risk of bias or Level I or III studies with a moderate risk of bias or Level II or III studies/SRs with a high risk of bias or Level I or III studies/SRs with a high risk of bias or Level II or III studies/SRs with a high risk of bias or Level I or III studies/SRs with a high risk of bias or Level I or III studies/SRs with a high risk of bias or Level I or III studies/SRs with a high risk of bias or Level I or III studies/SRs with a high risk of bias or Level I or III studies/SRs with a high risk of bias or Level I or III studies/SRs with a high risk of bias or Level I or III studies/SRs with a high risk of bias or Level I or III studies/SRs with a high risk of bias or Level I or III studies/SRs with a high risk of bias or Level I or III studies/SRs with a high risk of bias or Level I or III studies/SRs with a high risk of bias or Level I or III studies with a low risk of bias or Level I or III studies with a low risk of bias or Level I or III studies with a low risk of bias or Level I or III studies with a low risk of bias or Level I or III studies with a high risk of bias or Level I or III studies with a high risk of bias or Level I or III studies with a high risk of bias or Level I or III studies with a high risk of bias or Level I or III studies with a high risk of bias or Level I or III studies with a high risk of bias or Level III studies with a high risk of bias or Level III studies with a high risk of bias or Level III studies with a high risk of bias or Level III studies with a high risk of bias or Level III studies with a high risk of bias or Level III studies with a high risk of bias or Level III studies with a high risk of bias or Level III studies with a high risk of bias or Level III studies with a high risk of bias or Level III studies with a hi	,		One or more level I studies with a low risk of higs or several level I	I studies with a low risk of hias				
Consistency (it only one study was available, rank this component as not applicable) Pack	states of any local note admined miner accessed the shock of desireprocessing in quality of more							
Consistency (it any one study was available, rank this component as 'not applicable') Na		В	One of two Level II studies with a low risk of bias of SR/several Lev	ver ill studies with a low risk of dias				
2. Consistency (if only one study was available, rank this component as 'not applicable') NA All studies consistent B Most studies consistent and inconsistency can be explained C Some inconsistency, reflecting genuine uncertainty around question D Evidence is inconsistent NA Not applicable (one study only) 3. Clinical impact (Indicate in the space below if the study results varied according to some unknown factor (not simply study quality or sample size) and thus the clinical impact of the intervention could not be determined. NA Very large B Substantial C Moderate D Slight/Restricted 4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?) NA Very large A Very large B Substantial C Moderate D Slight/Restricted 4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?) NA Evidence directly generalisable to target population with some caveats C Evidence not directly generalisable to target population with some caveats 5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of health service/sidelivery of care and cultural factors?) NA Evidence applicable to Australian healthcare context with few caveats C Evidence probably applicable to Australian healthcare context with some caveats		С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias				
A All studies consistent B Most studies consistent and inconsistency can be explained C Some inconsistency, reflecting genuine uncertainty around question D Evidence is inconsistent NA Not applicable (one study only) 3. Clinical impact (Indicate in the space below if the study results varied according to some <u>unknown</u> factor (not simply study quality or sample size) and thus the clinical impact of the intervention could not be adecordinated. NA Very large B Substantial C		D	Level IV studies or Level I to III studies/SRs with a high risk of bias					
A All studies consistent B Most studies consistent and inconsistency can be explained C Some inconsistency, reflecting genuine uncertainty around question D Evidence is inconsistent NA Not applicable (one study only) 3. Clinical impact (Indicate in the space below if the study results varied according to some <u>unknown</u> factor (not simply study quality or sample size) and thus the clinical impact of the intervention could not be adecordinated. NA Very large B Substantial C	2. Consistency (If only one study was available, rank this component as 'not applicable')							
C Some inconsistency, reflecting genuine uncertainty around question D Evidence is inconsistent NA Not applicable (one study only) 3. Clinical impact (indicate in the space below if the study results varied according to some unknown factor (not simply study quality or sample size) and thus the clinical impact of the intervention could not be advantaged by the study on the intervention could not be determined. A Very large B Substantial C Moderate D Slight/Restricted 4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?) A Evidence directly generalisable to target population with some caveats C Evidence not directly generalisable to the target population but could be sensibly applied D Evidence not directly generalisable to target population and hard to judge whether it is sensible to be sensible to directly generalisable to target population and hard to judge whether it is sensible to be sensible to a Evidence of directly generalisable to target population and hard to judge whether it is sensible to be sensible to a Evidence of directly generalisable to target population and hard to judge whether it is sensible to be sensible to a Evidence of directly applicable to Australian healthcare context with few caveats E Evidence applicable to Australian healthcare context with few caveats C Evidence probably applicable to Australian healthcare context with some caveats	NA	Α	All studies consistent					
D Evidence is inconsistent NA Not applicable (one study only) 3. Clinical impact (Indicate in the space below if the study results varied according to some unknown factor (not simply study quality or sample size) and thus the clinical impact of the intervention could not be determined. A Very large B Substantial C Moderate D Slight/Restricted 4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?) NA Evidence directly generalisable to target population with some caveats C Evidence not directly generalisable to the target population but could be sensibly applied D Evidence not directly generalisable to target population and hard to judge whether it is sensible to surget population and hard to judge whether it is sensible to surget population and hard to judge whether it is sensible to surget population and hard to judge whether it is sensible to surget population and hard to judge whether it is sensible to surget population and hard to judge whether it is sensible to surget population and hard to judge whether it is sensible to surget population and hard to judge whether it is sensible to surget population and hard to judge whether it is sensible to account directly applicable to Australian healthcare context with few caveats E vidence directly applicable to Australian healthcare context with few caveats C Evidence probably applicable to Australian healthcare context with some caveats		В	Most studies consistent and inconsistency can be explained					
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EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	NA	
2. Consistency	NA	
3. Clinical impact	NA	
4. Generalisability	NA	
5. Applicability	NA	

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of desmopressin therapy on quality of life, compared with no therapy, is unknown.

Abbreviations: NA, not applicable.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^a Interventions: 1 = ANH, 2 = intraoperative cell salvage, 3 = perioperative ANH and intraoperative cell salvage, 4 = postoperative cell salvage, 5 = deliberate induced hypotension, 6 = prevention of hypothermia, 7 = point-of-care testing for coagulation status and haemoglobin, 8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, 9 = appropriate patient positioning, 10 = PAD

In patients undergoing surgery, what is the effect of administration of desmopressin on reoperation for bleeding? 1. Evidence base (number of studies, level of evidence and risk of bias in the included studies) There is one pivotal Level I study (Crescenzi 2008/fair quality) including data from up to 15 RCTs, and one supportive Level I study (Carless 2008/good quality). A One or more level I studies with a low risk of bias or several level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias or SR/several Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias or Level I or II studies with a low risk of bias or Level I or II studies with a high risk of bias or Level I or II studies with a low risk of bias or Level I or II studies with a high risk of bias or Level I or II studies with a low risk of bias or Level I or II studies	Key question(s):			Evidence table refa:					
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Cardiac surgery – 2.8% vs 4.7%; OR 0.63 (0.36, 1.08); 14 RCTs (N=1136) Non-cardiac surgery – 2.6% vs 2.5%; OR 1.00 (0.18,5.51); 1 RCT (N=50) Supportive evidence – Carless 2008 Any surgery – 1.8% vs 3.5%; RR 0.69 (0.26, 1.83); 9 RCTs (N=693) 4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?) The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were included in the overall analysis although studies were predominantly in cardiac surgery (14/15 RCTs). There were also separate analyses of patients undergoing cardiac surgery or non-cardiac surgery. Evidence directly generalisable to target population with some caveats C Evidence not directly generalisable to target population but could be sensibly applied Evidence not directly generalisable to target population and hard to judge whether it is sensible to apply 5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of health services/delivery of care and cultural factors?) Hospital setting. The pivotal review does not state which countries the RCTs were conducted in but the supportive study (which includes most of the studies included in the pivotal review) states that studies were conducted in US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK.		Α	Very large						
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Supportive evidence – Carless 2008 Any surgery – 1.8% vs 3.5%; RR 0.69 (0.26, 1.83); 9 RCTs (N=693) 4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?) The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were included in the overall analysis although studies were predominantly in cardiac surgery (14/15 RCTs). There were also separate analyses of patients undergoing cardiac surgery or non-cardiac surgery. Evidence directly generalisable to target population with some caveats C Evidence not directly generalisable to target population but could be sensibly applied D Evidence not directly generalisable to target population and hard to judge whether it is sensible to apply 5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of health services/delivery of care and cultural factors?) Hospital setting. The pivotal review does not state which countries the RCTs were conducted in but the supportive study (which includes most of the studies included in the pivotal review) states that studies were conducted in US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK.		С	Moderate						
4. Generalisability (How well does the body of evidence match the population and clinical settings being targeted by the Guideline?) The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were also separate analysis although studies were predominantly in cardiac surgery (14/15 RCTs). There were also separate analyses of patients undergoing cardiac surgery or non-cardiac surgery. B Evidence directly generalisable to target population with some caveats C Evidence not directly generalisable to the target population but could be sensibly applied D Evidence not directly generalisable to target population and hard to judge whether it is sensible to apply 5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of health services/delivery of care and cultural factors?) Hospital setting. The pivotal review does not state which countries the RCTs were conducted in but the supportive study (which includes most of the studies included in the pivotal review) states that studies were conducted in US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK. A Evidence directly generalisable to target population with some caveats Evidence and cultural factors?) A Evidence directly generalisable to target population and hard to judge whether it is sensible to apply Evidence not directly generalisable to Australian healthcare context A Evidence directly applicable to Australian healthcare context B Evidence directly generalisable to Australian healthcare context Evidence directly generalisable to Australian healthcare context Evidence applicable to Australian healthcare context with some caveats C Evidence probably applicable to Australian healthcare context with some caveats		D	Slight/Restricted						
The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were included in the overall analysis although studies were predominantly in cardiac surgery (14/15 RCTs). There were also separate analyses of patients undergoing cardiac surgery or non-cardiac surgery. B Evidence directly generalisable to target population with some caveats C Evidence not directly generalisable to target population but could be sensibly applied D Evidence not directly generalisable to target population and hard to judge whether it is sensible to apply 5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of health services/delivery of care and cultural factors?) Hospital setting. The pivotal review does not state which countries the RCTs were conducted in but the supportive study (which includes most of the studies included in the pivotal review) states that studies were conducted in US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK. A Evidence directly generalisable to target population with some caveats Evidence directly generalisable to target population and hard to judge whether it is sensible to apply Evidence not directly generalisable to target population and hard to judge whether it is sensible to apply Evidence not directly generalisable to target population with some caveats Evidence and cultural factors?) A Evidence directly applicable to Australian healthcare context B Evidence applicable to Australian healthcare context Evidence applicable to Australian healthcare context with few caveats C Evidence probably applicable to Australian healthcare context with some caveats	Any surgery – 1.8% vs 3.5%; RR 0.69 (0.26, 1.83); 9 RCTs (N=693)								
Included in the overall analysis although studies were predominantly in cardiac surgery (14/15 RCTs). There were also separate analyses of patients undergoing cardiac surgery or non-cardiac surgery. B Evidence directly generalisable to target population with some caveats C Evidence not directly generalisable to target population but could be sensibly applied D Evidence not directly generalisable to target population and hard to judge whether it is sensible to apply 5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of health services/delivery of care and cultural factors?) Hospital setting. The pivotal review does not state which countries the RCTs were conducted in but the supportive study (which includes most of the studies included in the pivotal review) states that studies were conducted in US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK. B Evidence applicable to Australian healthcare context with some caveats C Evidence probably applicable to Australian healthcare context with some caveats	4. Generalisability (How well does the body of evidence match the population and clinical se	ttings Ł	peing targeted by the Guideline?)						
also separate analyses of patients undergoing cardiac surgery or non-cardiac surgery. C Evidence not directly generalisable to the target population but could be sensibly applied Evidence not directly generalisable to target population and hard to judge whether it is sensible to apply 5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of health services/delivery of care and cultural factors?) Hospital setting. The pivotal review does not state which countries the RCTs were conducted in but the supportive study (which includes most of the studies included in the pivotal review) states that studies were conducted in US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK. E vidence directly applicable to Australian healthcare context with few caveats C Evidence probably applicable to Australian healthcare context with some caveats	The evidence is generalisable to an adult population who are undergoing planned surgery. All surgery types were	Α	Evidence directly generalisable to target population						
C Evidence not directly generalisable to the target population but could be sensibly applied D Evidence not directly generalisable to target population and hard to judge whether it is sensible to apply 5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of health services/delivery of care and cultural factors?) Hospital setting. The pivotal review does not state which countries the RCTs were conducted in but the supportive study (which includes most of the studies included in the pivotal review) states that studies were conducted in US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK. C Evidence not directly generalisable to target population but could be sensibly applied Evidence and cultural factors?) A Evidence directly applicable to Australian healthcare context B Evidence applicable to Australian healthcare context with few caveats C Evidence probably applicable to Australian healthcare context with some caveats		В	Evidence directly generalisable to target population with some cav	eats					
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of health services/delivery of care and cultural factors?) Hospital setting. The pivotal review does not state which countries the RCTs were conducted in but the supportive study (which includes most of the studies included in the pivotal review) states that studies were conducted in US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK. A Evidence directly applicable to Australian healthcare context with few caveats C Evidence probably applicable to Australian healthcare context with some caveats	and copulate analyses of patients analogous great and sargery or non-saratas sargery	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied					
Hospital setting. The pivotal review does not state which countries the RCTs were conducted in but the supportive study (which includes most of the studies included in the pivotal review) states that studies were conducted in US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK. A Evidence directly applicable to Australian healthcare context B Evidence applicable to Australian healthcare context with few caveats C Evidence probably applicable to Australian healthcare context with some caveats		D							
The pivotal review does not state which countries the RCTs were conducted in but the supportive study (which includes most of the studies included in the pivotal review) states that studies were conducted in US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK. B Evidence applicable to Australian healthcare context with few caveats C Evidence probably applicable to Australian healthcare context with some caveats	5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of heal	th services/delivery of care and cultural factors?)						
includes most of the studies included in the pivotal review) states that studies were conducted in US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK. C Evidence probably applicable to Australian healthcare context with some caveats		Α	Evidence directly applicable to Australian healthcare context						
Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK. Evidence probably applicable to Australian healthcare context with some caveats		В	Evidence applicable to Australian healthcare context with few cave	eats					
		С	Evidence probably applicable to Australian healthcare context with some caveats						
		D	Evidence not applicable to Australian healthcare context						

Desmopressin injection currently listed by TGA for the following indication: "patients undergoing cardiac surgery with cardiopulmonary bypass for prosthetic valve replacement or aortocoronary bypass grafting, especially when it is complicated by platelet function defects sufficient to prolong bleeding time despite relatively normal platelet cover. Desmopressin acetate offers no benefit as routine therapy in patients having an uncomplicated (simple) cardiopulmonary bypass procedure".

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	Α	There is one pivotal Level I study (fair quality) and one supportive Level I study.
2. Consistency	Α	There was no heterogeneity in the Level I studies and consistent results between the two studies.
3. Clinical impact	В	There was no significant difference in all analyses; however, there was a slight trend towards favouring DDAVP in cardiac surgery. The included studies may not be suffiently powered to detect a difference.
4. Generalisability	А	The results are generalisable to an adult surgical population undergoing cardiac surgery
5. Applicability	В	Studies were conducted in a range of countries. Likely to be applicable to the Australian setting

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing cardiac surgery, the effect of intravenous desmopressin therapy on risk of reoperation due to bleeding compared with no therapy is uncertain.

Note: Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and I² <25%; moderate heterogeneity if P>0.1 and I² between 25-50%; substantial heterogeneity if P<0.1 and I² >50%. Abbreviations: CI, confidence interval, Pot, potential; RBC, red blood cell; SR, systematic review.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

a Interventions: 1 = ANH, 2 = intraoperative cell salvage, 3 = perioperative ANH and intraoperative cell salvage, 4 = postoperative cell salvage, 5 = deliberate induced hypotension, 6 = prevention of hypothermia, 7 = point-of-care testing for coagulation status and haemoglobin, 8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, 9 = appropriate patient positioning, 10 = PAD

POQ3.l8d.S2 Characteristics and results of studies examining the effect of desmopressin on reoperation for bleeding

Study	Level of	No. of trials /	Patient population /	Setting	Intervention	Outcome	Results		Heterogeneity ^b
	evidence sample size Surgical procedure Quality Surgical procedure			Risk estimate (95% CI)	Significance P-value				
ADULT POPULATION	IV DESMOPRESSI	N							
Any surgery									
Crescenzi (2008)	Level I Fair	15 RCTs N=1186	Adult patients undergoing any surgery	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Surgical revision for bleeding	OR 0.65 (0.39, 1.09)	No difference 0.11	None Phet=0.50 (I ² =0%)
Carless (2008)	Level I Good	9 RCTs N=693	Adult patients undergoing any surgery	Hospital – elective or non-urgent surgery Various	Desmopressin (IV) vs no desmopressin	Reoperation for bleeding	RR 0.69 (0.26, 1.83)	No difference 0.45	None Phet=0.39 (l ² =6%)
Cardiac surgery	•								
Crescenzi (2008)	Level I Fair	14 RCTs N=1136	Adult patients undergoing cardiac surgery	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Surgical revision for bleeding	OR 0.63 (0.36, 1.08)	No difference 0.09	None Phet=0.44 (l²=0.6%)
Non-cardiac surge	ery								
Crescenzi (2008)	Level I/II Fair/ Unknown	1 RCT N=50	Adults patients undergoing surgery other than cardiac	Hospital – any surgery Countries not specified	Desmopressin (IV) vs placebo	Surgical revision for bleeding	OR 1.00 (0.18,5.51)	No difference 1.00	NA Phet=NA (I ² =NA)

Note: Studies providing pivotal evidence for the Evidence Statement Form are shown in shading. Unshaded studies provide supportive evidence.

Abbreviations: CI, confidence interval; NA, not applicable; OR, odds ratio; RCT, randomised controlled trial; RR, risk ratio.

^a Where only one RCT is available in a systematic review, the level of evidence will be downgraded to Level I/II. The quality of the included level II study will be rated based on the quality assessment of the systematic review.

b Heterogeneity defined as follows: (i) no significant heterogeneity if Phet>0.1 and 12<25%; (ii) mild heterogeneity if 12 <25%; moderate heterogeneity if 12 between 25-50%; substantial heterogeneity 12 >50%.

^c US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland, UK.

Recommendation(s) for administration of desmopressin

RECOMMENDATION	GRADE	RELE	VANT
What recommendation(s) does the guideline development group draw from this evidence? Use action statements where possible.	EVIDENC	E TABLE	
No recommendation made due to safety concerns.			
IMPLEMENTATION OF DECOMMENDATION			
IMPLEMENTATION OF RECOMMENDATION Please indicate yes or no to the following questions. Where the answer is yes please provide explanatory information about this. This information will be used to develop the implementation plan for the guidelines.			
Will this recommendation result in changes in usual care?		YES	NO
Are there any resource implications associated with implementing this recommendation?		YES	NO
Will the implementation of this recommendation require changes in the way care is currently organised?		YES	NO
Are the guideline development group aware of any barriers to the implementation of this recommendation		YES	NO
What could help to facilitate implementation of the recommendation?		YES	NO

Intervention 9 – Appropriate patient positioning

Key question(s): In patients undergoing surgery, what is the effect of <u>patient positioning</u> on <u>tran</u>	nsfusi	ion incidence?	Evidence table ref*: POQ3.19.P1			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)					
Four level II studies:	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias			
Park 2000 (N=40); good quality. Pace et al. 2008 (N=101), Ong et al. 2003 (N=60), Widman et al. 2001 (N=74); all fair quality.	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias			
Pace et al. 2008 (N=101), Ong et al. 2003 (N=60), Widman et al. 2001 (N=74); all fall quality.	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
None of the studies observed a significant effect of alternate patient positioning on the incidence of	Α	All studies consistent				
transfusion during surgery.	В	Most studies consistent and inconsistency can be explained				
Two studies examined the use of the lateral position compared to supine position during hip	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
arthroplasty. However, both studies failed to detect a significant effect of patient positioning on	D	Evidence is inconsistent				
transfusion incidence.	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>ur</u>	<u>nknown</u>	n factor (not simply study quality or sample size) and thus the clinical impac	ct of the intervention could not be			
There were no significant effects observed in any of the studies.	Α	Very large				
The examination of different surgical procedures and the use of different patient positions in each	В	Substantial				
study makes it difficult to assess the clinical impact of the patient positions examined.	С	Moderate				
	D					
4. Generalisability (How well does the body of evidence match the population and clinical set	tings be	eing targeted by the Guideline?)				
The four studies identified examined patients undergoing hip arthroplasty, knee surgery and lumbar spinal surgery. As such the evidence is likely generalisable to patients undergoing such surgical	Α	Evidence directly generalisable to target population				
procedures.	В	Evidence directly generalisable to target population with some caveats				
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of healti	h services/delivery of care and cultural factors?)				
There were two studies conducted in the UK and one in Sweden. The evidence from these studies	Α	Evidence directly applicable to Australian healthcare context				
are likely applicable in the Australian context. The fourth study, examining lumbar spinal surgery, was conducted in South Korea. Differences in the	В	Evidence applicable to Australian healthcare context with few cave	eats			
healthcare system between South Korea and Australia may limit the applicability of the evidence in	С	Evidence probably applicable to Australian healthcare context with	n some caveats			
the Australian context.	D	Evidence not applicable to Australian healthcare context				

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	В	1 good quality RCT and 3 fair quality RCT.
2. Consistency	В	All four studies examined patients undergoing orthopaedic surgery. None of the studies observed a significant effect of patient positioning on transfusion incidence.
3. Clinical impact	D	No significant effect was observed
4. Generalisability	В	The four studies identified examined patients undergoing hip arthroplasty, knee surgery and lumbar spinal surgery. As such the evidence is likely generalisable to patients undergoing such surgical procedures.
5. Applicability	В	There were two studies conducted in the UK and one in Sweden. The evidence from these studies are likely applicable in the Australian context. One study was conducted in South Korea, differences in their healthcare system may limit the applicability of the evidence in the Australian context.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing orthopaedic surgery, the effect of patient positioning on the incidence of allogeneic blood transfusion is uncertain.

POQ3.I9.P1

Characteristics and results of studies examining the effect of <u>patient positioning</u> on <u>transfusion incidence</u>

	Level of evidence	No. of trials /	Patient population /			Outcome	Results				Nata
Study	Quality Sample size Surgical procedure Setting Intervention / Comparator	Intervention / Comparator	Outcome	Intervention		Comparator	p-value	Notes			
Pace et al. (2008)	Level II Fair	N=101	Patients undergoing hip arthroplasty.	Hospital in UK.	Lateral position / Supine position	Transfusion incidence n/N (%)	5/51	(9.8)	8/50 (16)	P=0.65	-
Ong et al.	Level II	N=60 unilateral total knee	Transfusion incidence	Intervention A	Intervention B						
(2003)	Fair			UK.	Comparator:	n/N (%)	7/20 (35)	7/20 (35)	11/20 (55)	P=0.3	-
Widman et al. (2001)	Level II Fair	N=74	Patients undergoing hip replacement surgery.	Hospital in Sweden	Lateral position / Supine position	Transfusion incidence n/N (%)	17/3) (57)	30/44 (68)	P=0.336	-
Park 2000	Level II Good	N=40	ASA class I and II patients undergoing posterior lumbar spinal surgery.	Hospital in South Korea.	Narrow pad width on support / Wide pad width on spinal support	Transfusion incidence n/N (%)	5/20	(25)	1/20 (5)	NS	-

Abbreviations: ASA, American Society of Anaesthesiologists; NS, not statistically significant.

Key question(s): In patients undergoing surgery, what is the effect of patient positioning on tra	Evidence table ref*: POQ3.19.P2					
1. Evidence base (number of studies, level of evidence and risk of bias in the included studie	s)					
Three level II studies:	Α	One or more level I studies with a low risk of bias or several level	Il studies with a low risk of bias			
Park 2000 (N=40); good quality. Ong et al. 2003 (N=60), Widman et al. 2001 (N=74); both fair quality.	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias			
ong et al. 2000 (14-00), waithair et al. 2001 (14-71), bott hair quality.	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	i			
2. Consistency (If only one study was available, rank this component as 'not applicable')						
None of the studies observed a significant effect of alternate patient positioning on the incidence of	Α	All studies consistent				
transfusion during surgery.	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around questi	on			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>u</u>	nknown	n factor (not simply study quality or sample size) and thus the clinical impa	ct of the intervention could not be			
There were no significant effects observed in any of the studies.	А	Very large				
The examination of different surgical procedures and the use of different patient positions in each	В	Substantial				
study make it difficult to assess the clinical impact of the patient positions examined.	С	Moderate				
	D	No difference				
4. Generalisability (How well does the body of evidence match the population and clinical se	ttings be	eing targeted by the Guideline?)				
The three studies identified examined patients undergoing hip arthroplasty, knee surgery and lumbar spinal surgery. As such the evidence is likely generalisable to patients undergoing such surgical	Α	Evidence directly generalisable to target population				
procedures.	В	Evidence directly generalisable to target population with some cave	reats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of healt					
One study was conducted in the UK and one in Sweden. The evidence from these studies are likely applicable in the Australian context.	Α	Evidence directly applicable to Australian healthcare context				
The third study, examining lumbar spinal surgery, was conducted in South Korea. Differences in the	В	Evidence applicable to Australian healthcare context with few cave				
healthcare system between South Korea and Australia may limit the applicability of the evidence in	С	Evidence probably applicable to Australian healthcare context with	ı some caveats			
the Australian context.	D	Evidence not applicable to Australian healthcare context				

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	В	1 good quality RCT and 2 fair quality RCT.
2. Consistency	Α	None of the studies observed a significant effect.
3. Clinical impact	D	No significant effect was observed
4. Generalisability	В	The three studies identified examined patients undergoing hip arthroplasty, knee surgery and lumbar spinal surgery. As such the evidence is likely generalisable to patients undergoing such surgical procedures.
5. Applicability	С	One study was conducted in the UK and one in Sweden. The evidence from these studies are likely applicable in the Australian context. One study was conducted in South Korea; differences in their healthcare system may limit the applicability of the evidence in the Australian context.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing orthopaedic surgery, the effect of patient positioning on the volume of allogeneic blood transfusion is uncertain.

POQ3.19.P2

Characteristics and results of studies examining the effect of <u>patient positioning</u> on <u>transfusion volume</u>

Study	Level of evidence	No. of trials /	Patient population /	2			Results				N.A.
	Quality	sample size	Surgical procedure	Setting	Intervention / Comparator	Outcome	Intervention		Comparator	p-value	Notes
Ong et al.	Level II		Patients undergoing primary unilateral total knee	Hospital in	Intervention A: Leg elevated with knee flexed Intervention B:	Blood transfusion dose	Intervention A	Intervention B		P=0.3	
(2003)	Fair	N=60	replacement for osteoarthritis.	UK	Leg elevated with knee extended Comparator: Knee extended and level with bed	Median (range)	0 (0, 2)	0 (0, 2)	2 (0, 3.5)		-
Widman et al. (2001)	Level II Fair	N=74	Patients undergoing hip replacement surgery.	Hospital in Sweden	Lateral position / Supine position	Blood transfusion dose Mean (SD)	on 321mL (341)		407mL (362)	P=0.307	-
Park 2000	Level II Good	N=40	ASA class I and II patients undergoing posterior lumbar spinal surgery.	Hospital in South Korea	Narrow pad width on support / Wide pad width on spinal support	Blood transfusion dose Mean (SD)	2.2 Uni	ts (NR)	2 Units (NR)	NS	-

Abbreviations: ASA, American Society of Anaesthesiologists; NR, not reported; NS, not statistically significant; SD, standard deviation.

Key question(s): In patients undergoing surgery, what is the effect of <u>patient positioning</u> on <u>block</u>	Evidence table ref*: POQ3.19.P3				
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	.)				
Five level II studies:	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias		
Park 2000 (N=40); good quality. Ko et al. 2008 (N=60), Pace et al. 2008 (N=101), Widman et al. 2001 (N=74); all fair quality.	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias		
	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias		
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias			
2. Consistency (If only one study was available, rank this component as 'not applicable')					
Patient positioning had a significant effect on blood loss in four out of the five studies.	Α	All studies consistent			
Two studies examined the effect of lateral versus supine position during hip arthroplasty, however, a	В	Most studies consistent and inconsistency can be explained			
significant effect was only observed in one of the studies.	С	Some inconsistency, reflecting genuine uncertainty around question	on		
	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>ur</u>	known	factor (not simply study quality or sample size) and thus the clinical impac	t of the intervention could not be		
The effect of patient position on the volume of blood loss was 442mL during spinal surgery, 125.7mL	Α	Very large			
during endoscopic sinus surgery, and between 27mL to 215mL during hip arthroplasty,	В	Substantial			
The examination of different surgical procedures and the use of different patient positions in each	С	Moderate			
study make it difficult to synthesise a single effect estimate for the clinical impact of the patient positioning.	D	Slight/Restricted			
4. Generalisability (How well does the body of evidence match the population and clinical set	tings be	eing targeted by the Guideline?)			
The five studies identified examined patients undergoing endoscopic sinus surgery, hip arthroplasty and	Α	Evidence directly generalisable to target population			
lumbar spinal surgery. As such the evidence is likely generalisable to patients undergoing such surgical procedures.	В	Evidence directly generalisable to target population with some cav	eats		
	С	Evidence not directly generalisable to the target population but con	uld be sensibly applied		
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to		
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of healti				
Three of the studies were conducted in the UK and Sweden. The evidence from these studies are	Α	Evidence directly applicable to Australian healthcare context			
likely applicable in the Australian context. One study was conducted in South Korea, while another was conducted in Taiwan. Differences in the	В	Evidence applicable to Australian healthcare context with few caveats			
healthcare system of these countries with Australia may limit the applicability of the evidence in the	С	Evidence probably applicable to Australian healthcare context with	some caveats		
Australian context.	D	Evidence not applicable to Australian healthcare context			

The study by Ko et al. examined blood loss during endoscopic sinus surgery. In this study, the importance of blood loss is related more to the obstruction of surgical field rather than to issues relating to blood transfusion requirements. Consequently, less emphasis has been placed on the findings from this study.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	1 good quality RCT and 3 fair quality RCT.
2. Consistency	С	3 of the 4 studies found a significant effect of patient positioning on blood loss.
3. Clinical impact	D	The effect of patient position on the volume of blood loss varied between 27mL to 442mL.
4. Generalisability	В	The five studies identified examined patients undergoing endoscopic sinus surgery, hip arthroplasty and lumbar spinal surgery. As such the evidence is likely generalisable to patients undergoing such surgical procedures.
5. Applicability	В	Two studies were conducted in the UK and Sweden. The evidence from these studies are likely applicable in the Australian context. One study was conducted in South Korea, while another was conducted in Taiwan. Differences in the healthcare system of these countries with Australia may limit the applicability of the evidence in the Australian context.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing certain types of surgery, the head-up and lateral patient positions are associated with reduced blood loss.

POQ3.19.P3

Characteristics and results of studies examining the effect of <u>patient positioning</u> on <u>blood loss</u>

Charles	Level of	No of trials /	Patient population /	Callian	Intervention / Comparator	0.1	Re		Notes	
Study	evidence <i>Quality</i>	sample size	Surgical procedure	Setting	Intervention / Comparator	Outcome	Intervention	Comparator	p-value	Notes
Ko et al. (2008)	Level II Fair	N=60	Patients undergoing Endoscopic sinus surgery.	Hospital in Taiwan.	Reverse Trendelenburg position / Supine position	Blood loss (mL) Mean (SD)	126.0 (85.8)	251.7 (139.1)	P<0.001	Difference in blood loss: 125.7mL
Pace et al. (2008)	Level II Fair	N=101	Patients undergoing hip arthroplasty.	Hospital in UK.	Lateral position / Supine position	Blood loss (mL) Mean (95% CI)	1129 (989, 1310)	1156 (954, 1265)	P=0.41	Difference in blood loss: 27mL
Widman et al.	Level II	N=74	Patients undergoing hip	hip Hospital in Lateral position / Suping position	Lateral position / Cuping position	Blood loss (mL)	Intraoperative: 508 (316)	723 (316)	P=0.001	Difference in blood loss: 215mL
(2001)	Fair	ir N=74	replacement surgery.	Sweden	Lateral position / Supine position	Mean (SD)	After 24 hr: 1273 (407)	1374 (458)	P=0.043	Difference in blood loss: 101mL
Park 2000	Level II Good	N=40	ASA class I and II patients undergoing posterior lumbar spinal surgery.	Hospital in South Korea.	Narrow pad width on support / Wide pad width on spinal support	Blood loss (mL) Mean (SD)	878 (521)	436 (159)	P<0.05	Difference in blood loss: 442mL

Abbreviations: ASA, American Society of Anaesthesiologists; CI, confidence interval; SD, standard deviation.

Key question(s): In patients undergoing surgery, what is the effect of patient positioning on mo	Evidence table ref*: POQ3.19.P4			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)			
There were no level I or II studies that reported data on mortality.	One or more level I studies with a low risk of bias or several level	a low risk of bias or several level II studies with a low risk of bias		
	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias	
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias	
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	;	
2. Consistency (If only one study was available, rank this component as 'not applicable')				
NA	Α	All studies consistent		
	В	Most studies consistent and inconsistency can be explained		
	С	Some inconsistency, reflecting genuine uncertainty around questi	on	
	D	Evidence is inconsistent		
	NA	Not applicable (one study only)		
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>undetermined</u>)	nknowr	n factor (not simply study quality or sample size) and thus the clinical impac	ct of the intervention could not be	
NA NA	Α	Very large		
	В	Substantial		
	С	Moderate		
	D	Slight/Restricted		
4. Generalisability (How well does the body of evidence match the population and clinical set	ttings b	eing targeted by the Guideline?)		
NA	Α	Evidence directly generalisable to target population		
	В	Evidence directly generalisable to target population with some cave	veats	
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied	
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to	
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of heali			
NA	Α	Evidence directly applicable to Australian healthcare context		
	В	Evidence applicable to Australian healthcare context with few cave		
	С	Evidence probably applicable to Australian healthcare context with	n some caveats	
	D	Evidence not applicable to Australian healthcare context		

Other factors (Inc	dicate here	any other factors that you took into account when assessing the evidence base (for example, issues that might cause the group to downgrade or upgrade the recommendation)
EVIDENCE STA	TEMEN	Γ MATRIX
Please summarise t	the develop	oment group's synthesis of the evidence relating to the key question, taking all the above factors into account.
Component	Rating	Description
1. Evidence base	NA	There were no level I or II studies that reported data on mortality.
2. Consistency	NA	
3. Clinical impact	NA	
4. Generalisability	NA	
5. Applicability	NA	
DRAFT EVIDEN	CE STA	TEMENT
Based on the body	of evidence	e above.
In adult patie	nts unde	rgoing surgery in which substantial blood loss is anticipated, the effect of patient positioning on mortality is unknown.

Key question(s):	Evidence table ref*:							
In patients undergoing surgery, what is the effect of <u>patient positioning</u> on <u>mod</u>	POQ3.I9.P5							
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies)								
Three level II studies:								
DeSio et al. 2008 (N=75); good quality. Pace et al. 2008 (N=101), Ong et al. 2003 (N=60); both fair quality	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias					
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias					
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias						
2. Consistency (If only one study was available, rank this component as 'not applicable')								
None of the studies observed a significant effect of patient positioning on morbidity outcomes	Α	All studies consistent						
Two studies examined the effect of patient positioning on the incidence of DVT; none of the studies	В	Most studies consistent and inconsistency can be explained						
observed a significant effect.	С	Some inconsistency, reflecting genuine uncertainty around question						
		Evidence is inconsistent						
	NA	Not applicable (one study only)						
3. Clinical impact (Indicate in the space below if the study results varied according to some undetermined)	<u>ıknown</u>	factor (not simply study quality or sample size) and thus the clinical impac	ct of the intervention could not be					
There were no significant effects observed in any of the studies.	Α	Very large						
The examination of different surgical procedures and the use of different patient positions in each	В	Substantial						
study make it difficult to assess the clinical impact of the patient positions examined.	С	Moderate						
	D	No difference						
4. Generalisability (How well does the body of evidence match the population and clinical sett	tings be	eing targeted by the Guideline?)						
The three studies identified examined patients undergoing nephrolithotomy, hip arthroplasty and knee	Α	Evidence directly generalisable to target population						
surgery. As such the evidence is likely generalisable to patients undergoing these surgical procedures.	В	Evidence directly generalisable to target population with some cave	reats					
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied					
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to					
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of healti	h services/delivery of care and cultural factors?)						
The studies were conducted in the UK or Italy, as such the evidence from these studies are likely	Α	Evidence directly applicable to Australian healthcare context						
applicable in the Australian context.	В	Evidence applicable to Australian healthcare context with few cave	eats					
	С	Evidence probably applicable to Australian healthcare context with	n some caveats					
	D	Evidence not applicable to Australian healthcare context						

п	a.,			
и	()thar tactors	/l		e (for example, issues that might cause the group to downgrade or upgrade the recommendation)
ш	L JILLIEL LAC LOLS	Undicate here any other factors that	I VALLIAAK INTA ACCALINI WAAN ASSASSINA INA AVIAANCA NASA I	A ITOL AYAMDIA. ISSUAS INAL MIANT CAUSA INA ALCOLO IN ALCOLO IN ALCOLO IN ALCOLO IN ALCOLO IN ALCOLO IN ALCOLO
п	Othici luctors	(indicate ricie diry other ractors that	you took into account which assessing the evidence base (e from example, 1334e3 that might eduse the group to downgrade or appraise the recommendation,

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	1 good quality RCT and 2 fair quality RCT.
2. Consistency	Α	None of the studies observed a significant effect of patient positioning on morbidity outcomes
3. Clinical impact	D	There were no significant effects observed in any of the studies.
4. Generalisability	В	The three studies identified examined patients undergoing nephrolithotomy, hip arthroplasty and knee surgery. As such the evidence is likely generalisable to patients undergoing these surgical procedures.
5. Applicability	В	The studies were conducted in the UK or Italy, as such the evidence from these studies are likely applicable in the Australian context.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of patient positioning on morbidity is uncertain.

POQ3.19.P5

Characteristics and results of studies examining the effect of patient positioning on morbidity

Chindre	Level of	No. of trials /	Patient population /	Catting	Intervention / Comments	Comparator Outcome Results					Notes				
Study	evidence <i>Quality</i>	sample size	Surgical procedure	Setting	Intervention / Comparator	ervention / Comparator Outcome		ention	Comparator	p-value	Notes				
DeSio et al.	Level II	N=75	Patients undergoing	Medical	Modified supine position /	Major complications ^a n/N (%)	1/39	(2.6)	0/36 (0)	P=0.2	-				
(2008)	Good	N=75	nephrolithotomy.	Institutions in Italy. Prone position Minor complications b n/N (%)	Italy. Prone position Minor complications b 7/39 (18)		(18)	5/36 (14)	P=0.16	-					
Pace et al.	Level II	N 101	Patients undergoing hip		Hospital in Lateral position / Supine		n		Incidence of D ^o n/N (%)		1/51	(1.9)	0/50 (0)	NS	-
(2008)	Fair	N=101	arthroplasty.		position	Wound infection n/N (%)	0/51 (0)		2/50 (4)	NS	-				
					Intervention A: Leg elevated with knee flexed Intervention B:	Leg elevated with knee	Incidence of DVT	Intervention A	Intervention B	0/20 (0)	NR	_			
Ong et al.	Level II		Patients undergoing primary	Hospital in		n/N (%)	1/20 (5)	1/20 (5)	0/20 (0)	NIX	_				
(2003)	Fair	N=60	unilateral total knee replacement for osteoarthritis.	UK.	Leg elevated with knee extended	Knee sweeling (cm)	Intervention A	Intervention B	3.8						
					Comparator: Knee extended and level with bed	Mean (range)	3.4 (1.0, 7.0)	3.3 (1.5, 8.0)	(1.5, 8.0)	P=0.6	-				

Abbreviations: DVT, deep vein thrombosis; NR, not reported; NS, not significant; SD, standard deviation.

^a Major complications include septicaemia, haemorrhaging requiring transfusion, thoracic or abdominal organ injury, acute pancreatitis.

^b Minor complications include fever, insignificant bleeding, urinary tract infection, colic.

Key question(s):			Evidence table ref*:					
In patients undergoing surgery, what is the effect of patient positioning on qua	POQ3.I9.P6							
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies)								
There were no studies that reported data on quality of life.	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias					
	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias					
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias					
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	;					
2. Consistency (If only one study was available, rank this component as 'not applicable')								
NA	Α	All studies consistent						
	В	Most studies consistent and inconsistency can be explained						
	С	Some inconsistency, reflecting genuine uncertainty around questi	on					
	D	Evidence is inconsistent						
	NA	Not applicable						
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>ur</u>	nknown	factor (not simply study quality or sample size) and thus the clinical impac	ct of the intervention could not be					
NA	Α	Very large						
	В	Substantial						
	С	Moderate						
	D	Slight/Restricted						
4. Generalisability (How well does the body of evidence match the population and clinical set	tings be	eing targeted by the Guideline?)						
NA	Α	Evidence directly generalisable to target population						
	В	Evidence directly generalisable to target population with some call	veats					
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied					
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to					
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of healt	·						
NA	Α	Evidence directly applicable to Australian healthcare context						
	В	Evidence applicable to Australian healthcare context with few cave	eats					
	С	Evidence probably applicable to Australian healthcare context with	n some caveats					
	D	Evidence not applicable to Australian healthcare context						

Other factors (Indi	Other factors (Indicate here any other factors that you took into account when assessing the evidence base (for example, issues that might cause the group to downgrade or upgrade the recommendation)							
EVIDENCE STAT	ΓEMENΙ	MATRIX						
Please summarise th	ne develop	ment group's synthesis of the evidence relating to the key question, taking all the above factors into account.						
Component	Rating	Description						
1. Evidence base	NA	There were no studies that reported data on quality of life.						
2. Consistency	NA							
3. Clinical impact	NA							
4. Generalisability	NA							
5. Applicability	NA							
DRAFT EVIDENC	CE STAT	TEMENT						
Based on the body o	f evidence	e above.						
In adult patier	its under	going surgery in which substantial blood loss is anticipated, the effect of patient positioning on quality of life is unknown.						

Recommendation(s) for appropriate patient positioning

RECOMMENDATION	GRADE	RELE	VANT			
What recommendation(s) does the guideline development group draw from this evidence? Use action statements where possible.						
No recommendation made.						
IMPLEMENTATION OF RECOMMENDATION	I					
Please indicate yes or no to the following questions. Where the answer is yes please provide explanatory information about this. This information will be used to develop the implementation plan for the guidelines.						
Will this recommendation result in changes in usual care?		YES	NO			
Are there any resource implications associated with implementing this recommendation?		YES	NO			
Will the implementation of this recommendation require changes in the way care is currently organised?		YES	NO			
Are the guideline development group aware of any barriers to the implementation of this recommendation		YES	NO			
What could help to facilitate implementation of the recommendation?		YES	NO			
		<u> </u>				

Intervention 10 – Preoperative autologous donation

Key question(s): In patients undergoing surgery, what is the effect of <u>PAD</u> on <u>transfusion incic</u>	(allogeneic blood)?	Evidence table ref*: POQ3.I10.P1a		
1. Evidence base				
1 level SR (Henry 2001) ^a and 2 Level II RCTs (Bouchard 2008; Hashimoto 2007)	А	One or more level I studies with a low risk of bias or several level	I studies with a low risk of bias	
Pivotal evidence – Henry 2001 (Level I; good quality); 11 RCTs (assessed RBCs only), all fair quality; N=1423, Most up-to-date search; includes largest number of studies	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias	
Supportive evidence – Bouchard 2008 (Level II; fair quality); N=48, Adult patients undergoing cardiac	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias	
surgery Supportive evidence – Hashimoto 2007 (Level II; poor quality); N=79, Adult patients undergoing liver graft procurement.	D	Level IV studies or Level I to III studies/SRs with a high risk of bias		
2. Consistency	•			
There is significant overall heterogeneity between the trials in Henry 2001 (Phet=0.00052). The results are	Α	All studies consistent		
consistent between the subgroups in Henry 2001. The results from Bouchard (2008) were not significant. No patients in Hashimoto 2007 were transfused with allogeneic blood.	В	Most studies consistent and inconsistency can be explained		
	С	Some inconsistency, reflecting genuine uncertainty around question	on	
	D	Evidence is inconsistent		
	NA	Not applicable (one study only)		
3. Clinical impact				
Pivotal evidence – Henry 2001 Any surgery – RR 0.36 (0.25, 0.51); 11 trials (N=1423)	А	Very large		
Cancer surgery - RR 0.49 (0.38, 0.63); 5 trials (N=950)	В	Substantial		
Orthopaedic surgery – RR 0.21 (0.11, 0.43); 5 (N=425) Maxillofacial surgery – RR 0.02 (0.00, 0.28); 1 trial (N=48)	С	Moderate		
Supportive evidence – Bouchard 2008 ^b and Hashimoto 2007 ^c (see Summary Table POQ3.I10.P1)	D	Slight/Restricted		
4. Generalisability				
The evidence is generalisable to an adult population who are undergoing elective surgery. Surgical operations	Α	Evidence directly generalisable to target population		
assessed include cancer surgery, orthopaedic surgery, maxillofacial surgery, cardiac surgery, and liver graft procurement.	В	Evidence directly generalisable to target population with some cav		
production.	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied	
	D	Evidence not directly generalisable to target population and hard t	o judge whether it is sensible to	
5. Applicability				
All the studies were conducted in a hospital setting. The RCTs included in Henry 2001 were conducted in Germany, Greece, Japan, Sweden, the Netherlands, and the USA. Bouchard 2008 and Hashimoto 2007 were	Α	Evidence directly applicable to Australian healthcare context		
conducted in Canadian and Japanese hospitals respectively.	В	Evidence applicable to Australian healthcare context with few cave	eats	
	С	Evidence probably applicable to Australian healthcare context with	n some caveats	
	D	Evidence not applicable to Australian healthcare context		

Other factors

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	Thirteen level II studies with a moderate risk of bias
2. Consistency	В	Some inconsistency, which is mainly in orthopaedic surgery. Overall direction of effect consistent.
3. Clinical impact	Α	The reduction in transfusion incidence associated with PAD is very large
4. Generalisability	В	Evidence directly generalisable to target population with some caveats
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, preoperative autologous donation reduces the incidence of allogeneic RBC transfusion.

Abbreviations: PAD, Preoperative autologous donation; RCT, randomised controlled trial; RR, relative risk; SR, systematic review.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hosp

^{* &}lt;u>Interventions</u>: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

b Results from Bouchard 2008 – RR (people transfused with allogeneic blood products): 0.41 (0.15, 1.15); RR (people transfused with allogeneic blood): 0.06 (0.00, 1.02).

Results from Hashimoto 2007 - RR 0 (0,0).

Key question(s): In patients undergoing surgery, what is the effect of <u>PAD</u> on <u>transfusion incid</u>	ence	(allogeneic and/or autologous blood)?	Evidence table ref*: POQ3.I10.P1b	
1. Evidence base				
1 level I SR (Henry 2001) ^a and 2 Level II RCTs (Bouchard 2008; Hashimoto 2007)	Α	One or more level I studies with a low risk of bias or several level	II studies with a low risk of bias	
Pivotal evidence – Henry 2001 (Level I; good quality); 11 RCTs (assessed RBCs only); N=1423 Most up-to-date search; includes largest number of studies Supportive evidence – Bouchard 2008 (Level II; fair quality); N=48	В	One or two Level II studies with a low risk of bias or SR/several Le	vel III studies with a low risk of bias	
Adult patients undergoing cardiac surgery	С	One or two Level III studies with a low risk of bias or Level I or II st	rudies with a moderate risk of bias	
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias	3	
2. Consistency				
There is significant overall heterogeneity between the trials in Henry 2001 (Phet<0.). The results are consistent between the subgroups in Henry 2001. The results from Bouchard (2008) were not significant.	Α	All studies consistent		
between the subgroups in Henry 2001. The results from Bouchard (2008) were not significant.	В	Most studies consistent and inconsistency can be explained		
	С	Some inconsistency, reflecting genuine uncertainty around question		
	D	Evidence is inconsistent		
	NA	Not applicable (one study only)		
3. Clinical impact				
Pivotal evidence – Henry 2001 Any surgery – RR 1.33 (1.10, 1.61); 9 trials (N=1232)	Α	Very large		
Cancer surgery – RR 1.38 (1.20, 1.58); 5 trials (N=950)	В	Substantial		
Orthopaedic surgery – RR 1.78 (0.61, 5.20); 3 trials (N=234) Maxillofacial surgery – RR 0 (0, 0); 1 trial (N=48)	С	Moderate		
Supportive evidence – Bouchard 2008 (see Summary Table POQ3.I10.P1)	D	Slight/Restricted		
4. Generalisability				
The evidence is generalisable to an adult population who are undergoing elective surgery. Surgical operations assessed include cancer surgery, orthopaedic surgery, maxillofacial surgery, and cardiac surgery.	Α	Evidence directly generalisable to target population		
assessed include cancer surgery, orthopaedic surgery, maximulacial surgery, and caldiac surgery.	В	Evidence directly generalisable to target population with some case	/eats	
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied	
	D	Evidence not directly generalisable to target population and hard	to judge whether it is sensible to	
5. Applicability				
All the studies were conducted in a hospital setting. The RCTs included in Henry 2001 were conducted in Germany, Greece, Japan, Sweden, the Netherlands, and the USA. Bouchard 2008 was conducted in a Canadian	Α	Evidence directly applicable to Australian healthcare context		
hospital.	В	Evidence applicable to Australian healthcare context with few cav		
	С	Evidence probably applicable to Australian healthcare context with some caveats		
	D	Evidence not applicable to Australian healthcare context		

Other factors

The CRG considered the conclusions from the Henry (2001) review, which outlines the potential for harm, offsetting benefits from preoperative autologous donation.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description			
1. Evidence base	С	hirteen level II studies with a low risk of bias			
2. Consistency	С	Some inconsistency, reflecting genuine uncertainty around question			
3. Clinical impact	В	Substantial clinical impact			
4. Generalisability	В	Evidence directly generalisable to target population with some caveats			
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats			

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss, preoperative autologous donation increases the overall incidence of blood transfusion.

Abbreviations: PAD, Preoperative autologous donation; RCT, randomised controlled trial; RR, relative risk; SR, systematic review.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission

^a Publication dated 2001 but includes update to January 2004

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I10.P1 Characteristics and results of studies examining the effect of PAD on transfusion incidence.

Study	Level of evidence Quality	No. of trials / sample size	Patient population / Surgical procedure	Setting	Intervention	Outcome	Results			Notes
							Intervention	Comparator	p-value	
Henry (2001)	Level I Good	11 trials N=1423	Adult patients undergoing any elective surgery	Studies conducted in developed countries	PAD vs standard care	Patients transfused with allogeneic blood	149/716 (21%)	375/707 (53%)	P<0.05	Phet<0.01
		9 trials N=1232				Patients transfused with allogeneic and/or autologous	496/620 (80%)	343/612 (56%)	P<0.05	Phet<0.01
Cancer surgery										
Henry (2001)	Level I Good	5 trials N=950	Adult patients undergoing any elective surgery	Studies conducted in developed countries	PAD vs standard care	Patients transfused with allogeneic blood	128/467 (27%)	280/483 (58%)	P<0.05	Phet=0.15
		5 trials N=950				Patients transfused with allogeneic and/or autologous	363/467 (78%)	260/483 (54%)	P<0.05	Phet=0.13
Orthopaedic sur	gery	•		•	•	•	•	•	•	'
Henry (2001)	Level I Good	5 trials N=425	Adult patients undergoing any elective surgery	Studies conducted in developed countries	PAD vs standard care	Patients transfused with allogeneic blood	21/221 (10%)	75/204 (37%)	P<0.05	Phet=0.07
		3 trials N=234				Patients transfused with allogeneic and/or autologous	105/125 (84%)	43/109 (39%)	P>0.05	Phet<0.01
Maxillofacial sur	gery					-				
Henry (2001)	Level I Good	1 trial N=48	Adult patients undergoing any elective surgery	Studies conducted in developed countries	PAD vs standard care	Patients transfused with allogeneic blood	0/28 (0%)	20/20 (100%)	P<0.05	
		1 trial N=48				Patients transfused with allogeneic and/or autologous	28/28 (100%)	20/20 (100%)	NA	
Cardiac surgery		•		•	•	•	•	•	•	•
Bouchard (2008)	Level II Fair	N=48	Adults undergoing elective cardiac surgery	Canadian hospital	Two units of 350 mL each (or 6mL/kg when the patent's weight was below 60 kg). Blood was reinfused postoperatively	Patients transfused with autologous blood	6/25 (24%)	NA	NA	
						Patients transfused with allogeneic blood products	4/25 (16%)	9/23 (39%)	P=0.09	
						Patients transfused with allogeneic blood	0/25 (0%)	7/23 (30%)	P=0.05	

	Level of evidence	No. of trials /	Patient population / Surgical procedure	Setting	Intervention	Outcome		Results		Notes
Study	Quality	sample size					Intervention	Comparator	p-value	
Liver resection	'	•							•	1
Hashimoto (2007)	Level II Poor	N=79	Adults undergoing liver graft procurement	Japanese hospital	Blood volume equal to approximately 0.7% of the patient's body weight was collected before the liver transection. The collected blood was reinfused into the patient after the graft procurement.	Patients transfused with allogeneic blood	0/40 (0%)	0/39 (0%)	Not estimable	
Studies with a t	ransfusion proto	ocol								
Henry (2001)	Level I Good	7 trials N=1206	Adult patients undergoing any elective surgery	Studies conducted in developed countries	PAD vs standard care	Patients transfused with allogeneic blood	138/595 (23%)	299/611 (49%)	P<0.05	Phet=0.18
		5 trials N=1015				Patients transfused with allogeneic and/or autologous	384/499 (77%)	267/516 (52%)	P<0.05	Phet<0.01
Studies without	t a transfusion pi	rotocol								
Henry (2001)	Level I Good	4 trials N=217	Adult patients undergoing any elective surgery	Studies conducted in developed countries	PAD vs standard care	Patients transfused with allogeneic blood	11/121 (9%)	76/96 (79%)	P<0.05	Phet=0.08
		4 trials N=217				Patients transfused with allogeneic and/or autologous	112/121 (93%)	76/96 (79%)	P>0.05	Phet<0.001

Abbreviations: PAD, preoperative autologous donation.

Key question(s):			Evidence table ref*:		
In patients undergoing surgery, what is the effect of PAD on transfusion volur	<u>ne</u> ?		POQ3.I10.P2		
1. Evidence base					
1 Level II RCT (Bouchard 2008); Fair quality; N=48	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias			
It is unclear whether allocation to treatment groups was concealed from those responsible for recruiting subjects. The treatment arms had similar demographic characteristics. Neither the patient nor the surgeon was blinded to the group assignment; however, the ICU intensivist, nurses, and residents were blinded. A transfusion protocol	В	One or two Level II studies with a low risk of bias or SR/several Lev	R/several Level III studies with a low risk of bias		
was uses. All analyses were conducted ITT. PAD was not completed in 2 patients (8%) because of worsened	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias		
angina pectoris. NB: Level I evidence (Henry 2001) did not report this outcome. ^a	D	Level IV studies or Level I to III studies/SRs with a high risk of bias			
2. Consistency (If only one study was available, rank this component as 'not applicable')					
NA	Α	All studies consistent			
	В	Most studies consistent and inconsistency can be explained			
	С	Some inconsistency, reflecting genuine uncertainty around question	on		
	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact (Indicate in the space below if the study results varied according to some undetermined)	nknowi	n factor (not simply study quality or sample size) and thus the clinical impa	ct of the intervention could not be		
Bouchard 2008 (N=48)	Α	Very large			
Autologous blood (mean [SD]: PAD vs. control), units: 2 (1.2) vs. 0 (0) Allogeneic blood (mean [SD]: PAD vs. control), units: 0 (0) vs. 2 (1.2)	В	Substantial			
Fresh frozen plasma (mean difference [SD]), units: 4.0 (0) vs. 2.8 (1) Platelets (mean difference [SD]), units: 4.3 (2.9) vs. 6 (0)	С	Moderate			
Cryoprecipitate (mean difference [SD]), units: 4.3 (2.9) vs. 6 (0)	D	Slight/Restricted			
4. Generalisability (How well does the body of evidence match the population and clinical set	tings b	eing targeted by the Guideline?)			
The study was conducted in adults undergoing elective cardiac surgery.	Α	Evidence directly generalisable to target population			
	В	Evidence directly generalisable to target population with some cav			
	С	Evidence not directly generalisable to the target population but cou	7 11		
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to		
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms	of heal	th services/delivery of care and cultural factors?)			
The study was conducted in a Canadian hospital.	Α	Evidence directly applicable to Australian healthcare context			
	В	Evidence applicable to Australian healthcare context with few cave			
	С	Evidence probably applicable to Australian healthcare context with	some caveats		
	D	Evidence not applicable to Australian healthcare context			

Other factors (Indicate here any other factors that you took into account when assessing the evidence base (for example, issues that might cause the group to downgrade or upgrade the recommendation)

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	One level II study with moderate risk of bias
2. Consistency	NA	Not applicable (one study only)
3. Clinical impact	В	Not statistically significant
4. Generalisability	С	Evidence not directly generalisable to the target population but could be sensibly applied
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, preoperative autologous donation may reduce the volume of allogeneic blood transfusion.

In adult patients undergoing surgery in which substantial blood loss is anticipated, preoperative autologous donation does not appear to have an effect on the overall volume of blood transfusion.

Abbreviations: ICU, intensive care unit: ITT, intention-to-treat: NA, not applicable: PAD, preoperative autologous donation; RCT, randomised controlled trial: SD, standard deviation.

Primary outcomes: P1 = transfusion incidence. P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission are a transfusion volume was one the primary outcomes in Henry 2001; however, none of the RCTs provided sufficient detail on this outcome for meta-analysis.

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I10.P2 Characteristics and results of studies examining the effect of PAD on transfusion volume.

0	Level of evidence	No. of trials /	Patient population / Surgical procedure	Setting	Intervention	Outcome		Results		Notes
Study	Quality	sample size			Intervention	Outcome	Intervention	Comparator	p-value	
Henry (2001)	Level I Good	11 trials N=1423	Adult patients undergoing any elective surgery	Studies conducted in developed countries	PAD vs standard care	Mean (SD), units	NR	NR	NR	None of the included RCTs provided sufficient evidence to conduct a meta-analysis.
Autologous bloo	d									
Bouchard (2008)	Level II Fair	N=48	Adults undergoing elective cardiac surgery	Canadian hospital	Two units of 350 mL each (or 6mL/kg when the patent's weight was below 60 kg) was withdrawn preoperatively. Blood was reinfused postoperatively	Mean (SD), units	2 (1.2)	0 (0)	Not estimable	
Allogeneic blood	1									
Bouchard (2008)	Level II Fair	N=48	Adults undergoing elective cardiac surgery	Canadian hospital	Two units of 350 mL each (or 6mL/kg when the patent's weight was below 60 kg) was withdrawn preoperatively. Blood was reinfused postoperatively	Mean (SD), units	0 (0)	2 (1.2)	Not estimable	
Fresh frozen plas	sma	1	1	1	•	-1	1	1	'	
Bouchard (2008)	Level II Fair	N=48	Adults undergoing elective cardiac surgery	Canadian hospital	Two units of 350 mL each (or 6mL/kg when the patent's weight was below 60 kg) was withdrawn preoperatively. Blood was reinfused postoperatively	Mean (SD), units	4 (0)	2.8 (1)	Not estimable	
Platelets	•	•	•	•	•	-	-	-	-	
Bouchard (2008)	Level II Fair	N=48	Adults undergoing elective cardiac surgery	Canadian hospital	Two units of 350 mL each (or 6mL/kg when the patent's weight was below 60 kg) was withdrawn preoperatively. Blood was reinfused postoperatively	Mean (SD), units	4.3 (2.9)	6 (0)	Not estimable	

Cryoprecipitate	Cryoprecipitate									
Bouchard (2008)	Level II Fair	N=48	Adults undergoing elective cardiac surgery	Canadian hospital	Two units of 350 mL each (or 6mL/kg when the patent's weight was below 60 kg) was withdrawn preoperatively. Blood was reinfused postoperatively	Mean (SD), units	0 (0)	10	Not estimable	

Abbreviations: NR, not reported; PAD, preoperative autologous donation; RCT, randomised controlled trial; SD, standard deviation.

Key question(s):			Evidence table ref*:			
In patients undergoing surgery, what is the effect of <u>PAD</u> on <u>blood loss</u> ?			POQ3.I10.P3			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)					
Bouchard 2008 (Level II; fair quality); N=48 Allocation concealment not reported. Neither the patient nor the surgeon was blinded to the group assignment;	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias				
however, the ICU intensivist, nurses, and residents were blinded. A transfusion protocol was uses. Analyses conducted ITT.	В	One or two Level II studies with a low risk of bias or SR/several Lev	rel III studies with a low risk of bias			
Hashimoto 2007 (Level II; poor quality); N=79	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias			
Allocation concealment not reported. The study was not blinded. No transfusion protocol was reported. The study was not conducted ITT ^a .	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
In both studies there was no significant difference in operative blood loss; however there was a significant difference in the blood loss during transection between PAD and control in Hashimoto 2007.	Α	All studies consistent				
unierence in the blood loss during transection between FAD and control in Hashimoto 2007.	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around question	on			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>undetermined</u>)	nknowi	n factor (not simply study quality or sample size) and thus the clinical impa	ct of the intervention could not be			
Bouchard 2008	Α	Very large				
Operative blood loss – mean difference -34 (-171, 102); P=0.62 Postoperative blood loss – mean difference 27 (-302, 355); P=0.88	В	Substantial				
Hashimoto 2007 Operative blood loss – mean difference -37 (-101, 27); P=0.25	С	Moderate				
Transection blood loss – mean difference -90 (-172, -8); P=0.031	D	Slight/Restricted				
4. Generalisability (How well does the body of evidence match the population and clinical set	tings b	neing targeted by the Guideline?)				
Bouchard 2008 was conducted in patients undergoing cardiac surgery. Hashimoto 2007 was conducted in patients undergoing liver resection.	Α	Evidence directly generalisable to target population				
patients divergeing five resection.	В	Evidence directly generalisable to target population with some cav	eats			
	С	Evidence not directly generalisable to the target population but cou	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to judge whether it is sensible to				
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of heal					
Bouchard 2008 was conducted in a Canadian hospital and Hashimoto 2007 was conducted in a Japanese hospital.	A	Evidence directly applicable to Australian healthcare context				
Подрам	В	Evidence applicable to Australian healthcare context with few caveats				
	С	Evidence probably applicable to Australian healthcare context with some caveats				
	D	Evidence not applicable to Australian healthcare context				

Other factors (Indicate here any other factors that you took into account when assessing the evidence base (for example, issues that might cause the group to downgrade or upgrade the recommendation)

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	Two level II studies of fair and poor quality and small size (N=48 and N=79)
2. Consistency	С	Some inconsistency, reflecting genuine uncertainty around question (refer to clinical impact)
3. Clinical impact	D	No statistically significant difference in operative blood loss. A small, but significant, difference in transection blood loss.
4. Generalisability	С	Evidence not directly generalisable to the target population but could be sensibly applied. One cardiac study and one liver graft procurement
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats. Studies conducted in Canada and Japan.

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of preoperative autologous donation on blood loss is uncertain.

Abbreviations: ICU, intensive care unit; ITT, intention to treat; PAD, preoperative autologous donation.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and one patient in the control group was excluded from analysis after randomisation because the operation was stopped due to an asthmatic attack.

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I10.P3 Characteristics and results of studies examining the effect of PAD on outcome blood loss.

Study evidence	Level of	No. of trials /	Patient population / Surgical procedure	Setting	Intervention C	Outcome		Results		Notes		
	Quality	sample size					Intervention	Comparator	p-value			
Bouchard (2008)	Level II Fair	N=48	Adults undergoing elective cardiac surgery	Canadian hospital	Two units of 350 mL each (or 6mL/kg when the patent's weight was below 60 kg). Blood was reinfused	Operative blood loss Mean difference (95% CI), mL	-34 (-171, 102)		P=0.62			
	postoperatively	Postoperative blood loss Mean difference (95% CI), mL	27 (-302, 355)		P=0.88							
Hashimoto (2007)	Level II Poor	N=79	Adults undergoing liver graft procurement	Japanese hospital	Blood volume equal to approximately 0.7% of the patient's body weight was collected before the liver transection. The collected	approximately 0.7% of the patient's body weight was collected before the liver	approximately 0.7% of the patient's body weight was collected before the liver transection. The collected	Operative blood loss Mean difference (95% CI), mL	-37 (-101, 27)		P=0.25	
				patient after the graft procurement.		Transection blood loss Mean difference (95% CI), mL	-90 (-172, -8)		P=0.031			

Abbreviations: CI, confidence interval; PAD, preoperative autologous donation.

Key question(s):			Evidence table ref*:			
In patients undergoing surgery, what is the effect of <u>PAD</u> on <u>mortality</u> ?			POQ3.I10.P4			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)					
1 Level II evidence RCT (Hashimoto 2007). Poor quality. (N=79; 40 PAD, 39 control) It is unclear whether allocation was concealed from those in charge of recruiting subjects. Baseline characteristics	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias				
were similar between treatment groups. The patients and surgeons were not blinded to randomisation results. No transfusion protocol was reported. One patient in the control group was excluded from analysis after	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias				
randomisation because the operation was stopped due to an asthmatic attack Henry 2001 (level I; good quality) reported "insufficient evidence" for an association between PAD and mortality	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias			
but did not report any more detail.		Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
	Α	All studies consistent				
	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around question	on			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact (Indicate in the space below if the study results varied according to some <u>ur</u>	nknowi	n factor (not simply study quality or sample size) and thus the clinical impa	ct of the intervention could not be			
The mortality rate was 0% in both treatment arms.	Α	Very large				
	В	Substantial				
	С	Moderate				
	D	No difference				
4. Generalisability (How well does the body of evidence match the population and clinical set	tings b					
The study population was patients undergoing liver graft procurement.	Α	Evidence directly generalisable to target population				
	В	Evidence directly generalisable to target population with some cav	eats			
	С	Evidence not directly generalisable to the target population but cou	ıld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to	judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of heal					
The study was conducted in a Japanese university hospital.	Α	Evidence directly applicable to Australian healthcare context				
	В	Evidence applicable to Australian healthcare context with few cave				
	С	Evidence probably applicable to Australian healthcare context with	some caveats			
	D	Evidence not applicable to Australian healthcare context				

Other factors (Indicate here any other factors that you took into account when assessing the evidence base (for example, issues that might cause the group to downgrade or upgrade the recommendation)

Included studies were underpowered to detect a mortality difference.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	D	One level II study with a high risk of bias
2. Consistency	NA	Not applicable (one study only)
3. Clinical impact	D	Not statistically significant
4. Generalisability	С	Evidence not directly generalisable to target population and hard to judge whether it is sensible to
5. Applicability	С	Evidence probably applicable to Australian healthcare context with some caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of preoperative autologous donation on mortality is uncertain.

Abbreviations: PAD, preoperative autologous donation; RCT, randomised controlled trial.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ÅNH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I10.P4 Characteristics and results of studies examining the effect of PAD on mortality.

Charles	Level of evidence Quality	No. of trials / sample size	Patient population / Surgical procedure	Setting	Intervention	Outcome	Results			Notes
Study							Intervention	Comparator	p-value	
Henry (2001)	Level I Good	11 trials N=1423	Adult patients undergoing any elective surgery	Studies conducted in developed countries	PAD vs standard care	Mortality	NR	NR	NR	The authors found "insufficient evidence" for any association between PAD and mortality.
Hashimoto (2007)	Level I Good	N=79	Adult population undergoing liver resection	Japanese hospital	PAD: Blood volume equal to approximately 0.7% of the patient's body weight was collected before the liver transection. The collected blood was reinfused into the patient after the graft procurement.	Mortality	0/40 (0%)	0/39 (0%)	not estimable	

Abbreviations: NR, not reported; PAD, preoperative autologous donation.

Key question(s):			Evidence table ref*:			
In patients undergoing surgery, what is the effect of <u>PAD</u> on <u>morbidity</u> ?			POQ3.I10.P5			
1. Evidence base (number of studies, level of evidence and risk of bias in the included studies	s)					
Evidence for rate of infection: 1 Level I SR (Henry 2001); good quality; N=621 Evidence for rate of thrombosis: 1 level I SR (Henry 2001); good quality; N=250	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias				
Evidence for rate of stroke, DVP, and pulmonary embolus: 1 level I SR (Henry 2001); good quality; N=NR Evidence for rate of bile leak: 1 level II RCT (Hashimoto 2007); poor quality; N=79	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias				
Evidence for rate of intra-abdominal bleeding: 1 level II RCT (Hashimoto 2007); poor quality; N=79	С	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias				
		Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency (If only one study was available, rank this component as 'not applicable')						
Infection: one trial reported in Henry 2001 found that the infection rate for patients with PAD was significantly lower than patients without PAD (RR 0.44; 95% CI: 0.20, 0.98). The other two trials reporting infection as an	Α	All studies consistent				
outcome found no significant difference between PAD and no PAD. The level of heterogeneity was not significant	В	Most studies consistent and inconsistency can be explained				
(Phet=0.07). <u>Thrombosis:</u> all three trials found no significant difference. The level of heterogeneity between the trials was not	С	Some inconsistency, reflecting genuine uncertainty around questic	Some inconsistency, reflecting genuine uncertainty around question			
ignificant (Phet=0.53)	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact						
<u>Infection</u> – RR 0.70 (0.34, 1.43); 3 trials (N=621) <u>Thrombosis</u> – RR 0.82 (0.21, 3.13); 3 trials (N=250)	Α	Very large				
Bile leak – RR 0.33 (0.01, 7.75); 1 trial (N=79)	В	Substantial				
Intra-abdominal bleeding – RR 0.33 (0.01, 7.75); 1 trial (N=79)	С	Moderate				
	D	No difference				
4. Generalisability (How well does the body of evidence match the population and clinical set	tings b					
The evidence is generalisable to an adult population who are undergoing elective surgery. Surgical operations assessed include cancer surgery, orthopaedic surgery, maxillofacial surgery, and liver graft procurement.	Α	Evidence directly generalisable to target population				
accessed include surror cargory, or inopacture surgery, marinolastic cargory, and inter-grant procedures in	В	Evidence directly generalisable to target population with some cav	eats			
	С	Evidence not directly generalisable to the target population but cou	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to			
5. Applicability (Is the body of evidence relevant to the Australian healthcare context in terms of	of heal					
All the studies were conducted in a hospital setting. The RCTs included in Henry 2001 were conducted in Germany, Greece, Japan, Sweden, the Netherlands, and the USA. Hashimoto 2007 was conducted in a	A	Evidence directly applicable to Australian healthcare context				
Japanese hospital.	В	Evidence applicable to Australian healthcare context with few cave				
	С	Evidence probably applicable to Australian healthcare context with some caveats				
	D	Evidence not applicable to Australian healthcare context				

Other factors (Indicate here any other factors that you took into account when assessing the evidence base (for example, issues that might cause the group to downgrade or upgrade the recommendation)

The CRG noted that the studies used varying definitions and sources of infection. Studies were of poor methodological quality and were underpowered for morbidity outcomes.

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description					
1. Evidence base	1. Evidence base C Level II studies with a moderate risk of bias						
2. Consistency C Some inconsistency, reflecting genuine uncertainty around question							
3. Clinical impact	D	Not statistically significant					
4. Generalisability	В	Evidence directly generalisable to target population with some caveats					
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats					

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect on preoperative autologous donation on morbidity is uncertain.

Abbreviations: DVP, deep vein thrombosis; NR, not reported; PAD, preoperative autologous donation; NR, not reported; RCT, randomised controlled trials; RR, relative risk; SR, systematic review.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I10.P5 Characteristics and results of studies examining the effect of PAD on morbidity.

Study	Level of evidence	No. of trials /	Patient population / Surgical				Results			Notes
	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Henry (2001)	Level I Good	3 trials N=621	Adult patients undergoing any elective surgery	Studies conducted in developed countries	PAD vs standard care	Infection	74/309 (24%)	81/312 (26%)	P>0.05	Phet=0.07
		3 trials N=250	Adult patients undergoing any elective surgery	Studies conducted in developed countries	PAD vs standard care	Thrombosis	6/140 (4%)	3/110 (3%)	P>0.05	Phet=0.53
		NR	Adult patients undergoing any elective surgery	Studies conducted in developed countries	PAD vs standard care	Other adverse events	NR	NR	NR	Insufficient data for stroke, DVP, and pulmonary embolus
Hashimoto (2007)	Level II Poor		0 0 0	Japanese hospital	Blood volume equal to approximately 0.7% of the patient's body weight was	Bile leak	0/40 (0%)	1/39 (3%)	P=0.49	
				collected before the liver	Intra-abdominal bleeding	0/40 (0%)	1/39 (3%)	P=0.49		

Abbreviations: DVP, deep vein thrombosis; NR, not reported; PAD, preoperative autologous donation.

Key question(s):		Evidence table ref*:			
In patients undergoing surgery, what is the effect of administration of PAD on	POQ3.I10.P6				
1. Evidence base					
No evidence found	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias		
	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias		
	С	One or two Level III studies with a low risk of bias or Level I or II stu	udies with a moderate risk of bias		
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias			
2. Consistency					
	Α	All studies consistent			
	B Most studies consistent and inconsistency can be explained				
C Some inconsistency, reflecting genuine uncertainty around question					
	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact					
	Α	Very large			
	В	Substantial			
	С	Moderate			
	D	Slight/Restricted			
4. Generalisability					
	Α	Evidence directly generalisable to target population			
	В	Evidence directly generalisable to target population with some cav	eats		
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied		
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to		
5. Applicability					
	Α	Evidence directly applicable to Australian healthcare context			
	В	Evidence applicable to Australian healthcare context with few cave			
	С	Evidence probably applicable to Australian healthcare context with	n some caveats		
	D	Evidence not applicable to Australian healthcare context			

Other factors		
EVIDENCE STA	TEMENT	MATRIX
Please summarise ti	he develop	oment group's synthesis of the evidence relating to the key question, taking all the above factors into account.
Component	Rating	Description
1. Evidence base	NA	
2. Consistency	NA	
3. Clinical impact	NA	
4. Generalisability	NA	
5. Applicability	NA	
DRAFT EVIDEN	CE STAT	rement
Based on the body of	of evidence	e above.
In adult pa	itients un	dergoing surgery in which substantial blood loss is anticipated, the effect of PAD on quality of life is unknown.

Abbreviations: PAD, preoperative autologous donation.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

Key question(s):			Evidence table ref*:			
In patients undergoing surgery, what is the effect of <u>PAD</u> on change in <u>haemo</u>	alob	oin concentration	POQ3.I10.S1			
1. Evidence base						
Henry 2001 (Level I; good quality); 5 RCTs (assessed RBCs only); N=534	Α	One or more level I studies with a low risk of bias or several level I	I studies with a low risk of bias			
Most up-to-date search; includes largest number of studies (only reports preoperative haemoglobin concentration) Bouchard 2008 (Level II; fair quality); N=48	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias			
Adult patients undergoing cardiac surgery. Reports pre- and postoperative haemoglobin concentration.						
Hashimoto 2007 (Level II; poor quality); N=79 Adult patients undergoing liver graft procurement. (only reports preoperative haemoglobin concentration)	С	One or two Level III studies with a low risk of bias or Level I or II stu				
NB: the timeframe between PAD and surgery was not reported in either Henry 2001, Bouchard 2008, or Hashimoto 2007.	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency						
Preoperative haemoglobin concentration	Α	All studies consistent				
There is a significant level of heterogeneity between the trials in Henry 2001 (Phet=0.004). The results from Henry 2001 do not agree with the results from Bouchard 2008 and Hashimoto 2007.	В	Most studies consistent and inconsistency can be explained				
Postoperative haemoglobin concentration Only one of the studies (Bouchard 2008) reported postoperative haemoglobin concentration as an outcome.	С	Some inconsistency, reflecting genuine uncertainty around question	nuine uncertainty around question			
Only one of the studies (bouchard 2000) reported postoperative fractiographic concentration as an outcome.	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact						
Henry 2001 – mean difference in preoperative concentration between PAD and control, -1.16 (-1.60, -0.73)	Α	Very large				
Bouchard 2008 – There was no significant difference between PAD and control at any time point. Mean difference: Preoperative, -0.60 (-1.36, 0.16); 24 hours postoperative, -0.40 (-1.11, 0.31); 5 days	В	Substantial				
postoperative, -0.50 (-1.18, 0.18) Healthingto 2007 Median (IOD) propagative becomed table consentration, DAD vs. control 13 0 (11.0 to 15.7)	С	Moderate				
Hashimoto 2007 – Median (IQR) preoperative haemoglobin concentration, PAD vs. control: 13.0 (11.0 to 15.7) vs. 13.6 (11.6 to 15.9)	D	Slight/Restricted				
4. Generalisability						
The evidence for difference in preoperative haemoglobin concentration between PAD and control is generalisable to an adult population who are undergoing elective surgery. Surgical operations assessed include cancer surgery,	Α	Evidence directly generalisable to target population				
orthopaedic surgery, maxillofacial surgery, cardiac surgery, and liver graft procurement.	В	Evidence directly generalisable to target population with some cav	eats			
The evidence for difference in postoperative haemoglobin concentration between PAD and control comes from one trial of patients undergoing cardiac surgery.	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
		Evidence not directly generalisable to target population and hard to judge whether it is sensible to				
5. Applicability						
All the studies were conducted in a hospital setting. The RCTs included in Henry 2001 were conducted in Germany, Greece, Japan, Sweden, the Netherlands, and the USA. Bouchard 2008 and Hashimoto 2007 were	Α	Evidence directly applicable to Australian healthcare context				
conducted in Canadian and Japanese hospitals respectively.	В	Evidence applicable to Australian healthcare context with few caveats				
	С	idence probably applicable to Australian healthcare context with some caveats				
	D	Evidence not applicable to Australian healthcare context				

Other factors		
EVIDENCE CTA	TENNENIT	MATRIV
EVIDENCE STA	IEWENI	MATRIX
Please summarise th	he develop	oment group's synthesis of the evidence relating to the key question, taking all the above factors into account.
Component	Rating	Description
Evidence base	С	Seven level II studies with a moderate risk of bias
2. Consistency	В	There is a significant degree of heterogeneity regarding

DRAFT EVIDENCE STATEMENT

В

3. Clinical impact

Generalisability
 Applicability

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, preoperative autologous donation reduces preoperative haemoglobin concentration.

PAD is associated with a moderate decrease in preoperative haemoglobin concentration compared with control

Evidence not directly generalisable to the target population but could be sensibly applied

Evidence applicable to Australian healthcare context with few caveats

Abbreviations: IQR, interquartile range; PAD, preoperative autologous donation; RCT, randomised controlled trial.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{* &}lt;u>Interventions</u>: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I10.S1 Characteristics and results of studies examining the effect of PAD on haemoglobin concentration.

	Level of	evidence No. of trials / Patient population / Surgical Setting						Notes		
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Henry (2001)	Level I Good	5 trials (2 in orthopaedic surgery and 3 in surgery for cancer) N=534	Adult patients undergoing any elective surgery	Studies conducted in developed countries	PAD vs standard care	Preoperative Hb concentration. Mean difference (95% CI), g/dL	-1.16 (-1.60, -0.73)		P<0.05	Phet=0.004
Bouchard (2008)	Level II Fair	N=48	Adults undergoing elective cardiac surgery	Canadian hospital	Two units of 350 mL each (or 6mL/kg when the patent's weight was below 60 kg). Blood was reinfused postoperatively	Preoperative Hb concentration. Mean difference (95% CI), g/dL	-0.60 (-1.36, 0.16)		P=0.12	
				Hb conce 24 hc surge Mean (95% Hb conce days surge Mean Mean Mean Mean Mean Mean Mean Mea	Hb concentration 24 hours after surgery. Mean difference (95% CI), g/dL	-0.40 (-1.11, 0.31)		P=0.27		
						Hb concentration 5 days after surgery. Mean difference (95% CI), g/dL	-0.50 (-1.18, 0.18)		P=0.15	
Hashimoto (2007)	Level II Poor	N=79	Adults undergoing liver graft procurement	Japanese hospital	Blood volume equal to approximately 0.7% of the patient's body weight was collected before the liver transection. The collected blood was reinfused into the patient after the graft procurement.	Preoperative Hb concentration. Median (IQR), g/dL	13.0 (11.0 to 15.7)	13.6 (11.6 to 15.9)	P=0.455	

Abbreviations: CI, confidence interval; IQR, interquartile range; Hb, haemoglobin; PAD, preoperative autologous donation.

Key question(s):	aulation status?	Evidence table ref*: POQ3.I10.S3				
In patients undergoing surgery, what is the effect of administration of <u>PAD</u> on 1. Evidence base	cual	guiation status?	1 0 2 3 .11 0 . 3 3			
Two Level II studies: Bouchard 2009 (fair quality; N=48); Hashimoto 2007 (poor quality; N=79)	Τ	One or many level letydics with a levy rick of his or several level l	U akudiaa wikh a law wiak af hiaa			
Bouchard 2009 reports prothrombin time, and fibrinogen concentration	A	One or more level I studies with a low risk of bias or several level II studies with a low risk of bias				
Hashimoto 2007 reports prothrombin time and INR	В	One or two Level II studies with a low risk of bias or SR/several Level III studies with a low risk of bias				
	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias				
2. Consistency						
Both studies found no significant impact on prothrombin time or INR. Bouchard 2009 is the only study that reports fibringen concentration.	Α	All studies consistent				
ilbilingen concentiation.	В	Most studies consistent and inconsistency can be explained				
	С	Some inconsistency, reflecting genuine uncertainty around question	วท			
	D	Evidence is inconsistent				
	NA	Not applicable (one study only)				
3. Clinical impact						
See Summary Table POQ3.I10.S3.	А	Very large				
	В	Substantial				
	С	Moderate				
	D	No difference				
4. Generalisability						
Bouchard 2008 was conducted in patients undergoing cardiac surgery. Hashimoto 2007 was conducted in patients undergoing liver resection.	Α	Evidence directly generalisable to target population				
patients undergoing liver resection.	В	Evidence directly generalisable to target population with some cav	eats			
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied			
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to			
5. Applicability						
Bouchard 2008 was conducted in a Canadian hospital and Hashimoto 2007 was conducted in a Japanese hospital.	Α	Evidence directly applicable to Australian healthcare context				
поэрни.	В	Evidence applicable to Australian healthcare context with few cave	ats			
	С	Evidence probably applicable to Australian healthcare context with	some caveats			
	D	Evidence not applicable to Australian healthcare context				

Other factors

EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description					
Evidence base	1. Evidence base C Two Level II studies with moderate risk of bias						
2. Consistency A All studies consistent							
3. Clinical impact D No statistically significant impact on prothrombin time							
4. Generalisability	С	Evidence not directly generalisable to the target population but could be sensibly applied					
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats					

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, preoperative autologous donation does not appear to have an effect on prothrombin time.

Abbreviations: INR, international normalised ratio; PAD, preoperative autologous donation.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I10.S3 Characteristics and results of studies examining the effect of PAD on coagulation status.

	Level of evidence	No. of trials /	Patient population / Surgical	Setting				Notes			
Study	Quality	sample size	procedure		Intervention	Outcome	Intervention	Comparator	p-value		
Bouchard (2008)	ouchard (2008) Level II Fair Adults undergoing elective cardiac surgery Canadian hospital Two units of 350 mL each (or 6mL/kg when the patent's weight was below 60 kg). Blood was reinfused postoperatively	Preoperative prothrombin time (mean [SD]), seconds	9.7 (2.8)	9.4 (1.1)	P=0.62						
		reinfused postoperatively	Prothrombin time 30 minutes after surgery (mean [SD]), seconds	13.2 (3.9)	13.5 (2.2)	P=0.74					
							Prothrombin time 24 hours after surgery (mean [SD]), seconds	10.3 (1.3)	10.9 (1.7)	P=0.17	
							Preoperative fibrinogen concentration (mean [SD]), g/L	4.3 (1.5)	3.1 (0.9)	P=0.0007	
							Fibrinogen concentration 30 minutes after surgery (mean [SD]), g/L	3.0 (0.9)	2.6 (0.7)	P=0.08	
						Fibrinogen concentration 24 hours after surgery (mean [SD]), g/L	6.2 (1.3)	5.1 (1.2)	P=0.002		
Hashimoto (2007)	Poor procurement approximately 0.7% of the patient's body weight was collected before the	Prothrombin time 24 hours after surgery (median [IQR]), seconds	12.3 (9.6 to 15.9)	12.5 (10.5 to 15.0)	P=0.280						
					liver transection. The collected blood was reinfused into the patient after the graft procurement.	Preoperative INR (median [IQR])	1.11 (0.95 to 1.34)	1.10 (0.91 to 1.31)	P=0.350		
						INR 24 hours postoperative (median [IQR])	1.76 (1.30 to 2.37)	1.77 (1.29 to 2.32)	P=0.456		

Abbreviations: IQR, interquartile range; PAD, preoperative autologous donation; SD, standard deviation.

Key question(s): In patients undergoing surgery, what is the effect of PAD on length of hospita	l stay	?	Evidence table ref*: POQ3.I10.S5					
1. Evidence base								
Bouchard 2008 (Level II; fair quality); N=48								
Allocation concealment not reported. Neither the patient nor the surgeon was blinded to the group assignment; however, the ICU intensivist, nurses, and residents were blinded. A transfusion protocol was uses. Analyses conducted ITT.	В	One or two Level II studies with a low risk of bias or SR/several Lev	vel III studies with a low risk of bias					
Hashimoto 2007 (Level II; poor quality); N=79	С	One or two Level III studies with a low risk of bias or Level I or II st	udies with a moderate risk of bias					
Allocation concealment not reported. The study was not blinded. No transfusion protocol was reported. The study was not conducted ITT ^a .	D	Level IV studies or Level I to III studies/SRs with a high risk of bias						
2. Consistency								
Both studies found no significant effect of administration of PAD on length of hospital stay.	Α	All studies consistent						
	В	Most studies consistent and inconsistency can be explained						
	С	Some inconsistency, reflecting genuine uncertainty around question	on					
	D	Evidence is inconsistent						
	NA	Not applicable (one study only)						
3. Clinical impact								
Bouchard 2008 Mean difference (95% CI), days: 0.00 (-0.51, 0.51); P=1.00	Α	Very large						
Hashimoto 2007	В	Substantial						
PAD vs. control (Median [IQR]), days: 14 (10 to 36) vs. 14 (11 to 46); P=0.476	С	Moderate						
	D	No difference						
4. Generalisability								
Bouchard 2008 was conducted in patients undergoing cardiac surgery. Hashimoto 2007 was conducted in patients undergoing liver resection.	Α	Evidence directly generalisable to target population						
patients undergoing liver resection.	В	Evidence directly generalisable to target population with some cav	eats					
	С	Evidence not directly generalisable to the target population but co	uld be sensibly applied					
	D	Evidence not directly generalisable to target population and hard to	o judge whether it is sensible to					
5. Applicability								
Bouchard 2008 was conducted in a Canadian hospital and Hashimoto 2007 was conducted in a Japanese hospital.	Α	Evidence directly applicable to Australian healthcare context						
поэрна.	В	Evidence applicable to Australian healthcare context with few caveats						
	С	Evidence probably applicable to Australian healthcare context with some caveats						
	D	Evidence not applicable to Australian healthcare context						

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EVIDENCE STATEMENT MATRIX

Please summarise the development group's synthesis of the evidence relating to the key question, taking all the above factors into account.

Component	Rating	Description
1. Evidence base	С	Two level II studies with a moderate risk of bias
2. Consistency	Α	Both studies showed consistent results
3. Clinical impact	D	Not statistically significant
4. Generalisability	С	Evidence not directly generalisable to the target population but could be sensibly applied
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats

DRAFT EVIDENCE STATEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of preoperative autologous donation on length of hospital stay is uncertain.

Abbreviations: CI, confidence interval; ICU, intensive care unit; IQR, interguartile range; ITT, intention-to-treat; PAD, preoperative autologous donation.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = mortality, P5 = morbidity, P6 = quality of life

Secondary outcomes: S1 = change in haemoglobin, S2 = reoperation for bleeding, S3 = correction/prevention of DIC and coagulopathy, S4 = cost, S5 = hospital length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hospital readmission and length of stay, S6 = ICU admission and length of stay, S7 = hospital readmission and length of stay, S7 = hospital readmissi

^{* &}lt;u>Interventions</u>: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I10.S5 Characteristics and results of studies examining the effect of PAD on hospital length of stay.

	CHICK	ence No. of trials / Pa	Patient population / Surgical procedure	Setting Intervention O			Results			Notes
Study					Outcome	Intervention	Comparator	p-value		
Bouchard (2008)	Level II Fair	N=48	Adults undergoing elective cardiac surgery	Canadian hospital	Two units of 350 mL each (or 6mL/kg when the patent's weight was below 60 kg). Blood was reinfused postoperatively	Mean difference (95% SD), days	0.00 (-0.51, 0.51)		P=1.00	
Hashimoto (2007)	Level II Poor	N=79	Adults undergoing liver graft procurement	Japanese hospital	Blood volume equal to approximately 0.7% of the patient's body weight was collected before the liver transection. The collected blood was reinfused into the patient after the graft procurement.	Median (IQR), days	14 (10 to 36)	14 (11 to 46)	P=0.476	

Abbreviations: IQR, interquartile range; PAD, preoperative autologous donation; SD, standard deviation.

Key question(s):		Evidence table ref*:			
In patients undergoing surgery, what is the effect of PAD on ICU admission a	ngth of stay?	POQ3.I10.S6			
1. Evidence base					
Bouchard 2008 (Level II; fair); N=48 Allocation concealment not reported. Neither the patient nor the surgeon was blinded to the group assignment;	Α	One or more level I studies with a low risk of bias or several level II studies with a low risk of			
however, the ICU intensivist, nurses, and residents were blinded. A transfusion protocol was uses. Analyses conducted ITT.	В	One or two Level II studies with a low risk of bias or SR/several Lev	rel III studies with a low risk of bias		
	С	One or two Level III studies with a low risk of bias or Level I or II studies with a moderate risk of bias			
	D	Level IV studies or Level I to III studies/SRs with a high risk of bias			
2. Consistency					
	Α	All studies consistent			
	В	Most studies consistent and inconsistency can be explained			
	С	Some inconsistency, reflecting genuine uncertainty around question	on		
	D	Evidence is inconsistent			
	NA	Not applicable (one study only)			
3. Clinical impact					
Mean difference (95% CI): 0.00 (-3.34, 0.34)	Α	Very large			
	В	Substantial			
		Moderate			
	D	No difference			
4. Generalisability					
Study was conducted in adults undergoing elective cardiac surgery.	Α	Evidence directly generalisable to target population			
	В	Evidence directly generalisable to target population with some cav	eats		
	С	Evidence not directly generalisable to the target population but cou	ıld be sensibly applied		
	D	Evidence not directly generalisable to target population and hard to	judge whether it is sensible to		
5. Applicability					
Study was conducted in a Canadian hospital.	Α	Evidence directly applicable to Australian healthcare context			
	В	Evidence applicable to Australian healthcare context with few cave			
	С	Evidence probably applicable to Australian healthcare context with	some caveats		
	D	Evidence not applicable to Australian healthcare context			

OIL 6 I		
Other factors		
EVIDENCE STA	TEMENT	T MATRIX
Discourse	l	
Please summarise ti	ne aeveiop	pment group's synthesis of the evidence relating to the key question, taking all the above factors into account.
Component	Rating	Description
1. Evidence base	С	One level II study with a moderate risk of bias
2. Consistency	NA	Not applicable (one study only)
3. Clinical impact	D	Not statistically significant
4. Generalisability	С	Evidence not directly generalisable to the target population but could be sensibly applied
5. Applicability	В	Evidence applicable to Australian healthcare context with few caveats
DRAFT EVIDEN	CE STA	TEMENT

Based on the body of evidence above.

In adult patients undergoing surgery in which substantial blood loss is anticipated, the effect of preoperative autologous donation on length of ICU stay is uncertain.

Abbreviations: CI, confidence interval; ICU, intensive care unit; ITT, intention-to-treat; PAD, preoperative autologous donation.

Primary outcomes: P1 = transfusion incidence, P2 = transfusion volume, P3 = blood loss, P4 = morbidity, P5 = morbidity, P6 = quality of life

^{*} Interventions: I1 = ANH, I2 = intraoperative cell salvage, I3 = perioperative ANH and intraoperative cell salvage, I4 = postoperative cell salvage, I5 = deliberate induced hypotension, I6 = prevention of hypothermia, I7 = point-of-care testing for coagulation status and haemoglobin, I8 = administration of antifibrinolytics (tranexamic acid, EACA, aprotinin) and DDAVP, I9 = appropriate patient positioning, I10 = PAD

POQ3.I10.S6 Characteristics and results of studies examining the effect of PAD on outcome ICU admission and length of stay.

	Level of evidence	No. of trials /	Patient population / Surgical				Results			Notes
Study	Quality	sample size	procedure	Setting	Intervention	Outcome	Intervention	Comparator	p-value	
Bouchard (2008)	Level II Fair	N=48	Adults undergoing elective cardiac surgery	Canadian hospital	Two units of 350 mL each (or 6mL/kg when the patent's weight was below 60 kg). Blood was reinfused postoperatively	Length of ICU stay, days Mean difference (95% CI)	0.00 (-0.34, 0.34)		P=1.00	

Abbreviations: CI, confidence interval; ICU, intensive care unit; PAD, preoperative autologous donation.

Recommendation(s) for preoperative autologous donation

RECOMMENDATION	GRADE	RELE	VANT
What recommendation(s) does the guideline development group draw from this evidence? Use action statements where possible.		EVIDENC	E TABLE
The routine use of preoperative autologous donation is not recommended because, although it reduces the risk	С	PC	3.I10.P1
of allogeneic RBC transfusion, it increases the risk of receiving any RBC transfusion (allogeneic and autologous).			
IMPLEMENTATION OF RECOMMENDATION		<u> </u>	
Please indicate yes or no to the following questions. Where the answer is yes please provide explanatory information about this.			
The information will be used to develop the implementation plan for the guidelines.			
Will this recommendation result in changes in usual care?		YES	NO
Routine use of PAD should reduce.			
Are there any resource implications associated with implementing this recommendation?		YES	NO
Potential cost savings; frees up ARCBS time.			
Will the implementation of this recommendation require changes in the way care is currently organised?		YES	NO
Are the guideline development group aware of any barriers to the implementation of this recommendation		YES	NO
Existence of MBS item number.			_
What could help to facilitate implementation of the recommendation?			
Successful lobbying to remove MBS item number.			

Appendix E: Quality analyses

Intervention 1 – Acute normovolemic haemodilution

Level I evidence

Citation	Bryson GL, Laupacis A, and Wells GA. (1998) Does acute normovolemic hemodilution reduce
	perioperative allogeneic transfusion? A meta-analysis. Anesthesia and Analgesia 86:9-15.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results if the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	
Overall	Good
assessment	

Citation	Carless P, Moxey A, O'Connell D, and Henry D. (2004) Autologous transfusion techniques: A
	systematic review of their efficacy. Transfusion Medicine
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
N	D. Was a quality assessment of included studies undertaken?
N	E. Were the characteristics and results if the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
N	Were the sources of heterogeneity explored?
Comments	The SR includes both RCTs and observational studies, however the results are evaluated separately
	by study type.
Overall	Fair
assessment	

Citation	Gurusamy KS, Li J, Sharma D, and Davidson BR. (2009) Cardiopulmonary interventions to decrease blood loss and blood transfusion requirements for liver resection. Cochrane
	Database of Systematic Reviews: Reviews 2009. Issue. 4
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results if the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	
Overall	Good
assessment	

Citation	Laupacis A and Fergusson D. (1998) The efficacy of technologies to minimise perioperative allogeneic transfusion (Structured abstract). Kluwer. Academic Publishers.17-36.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
N	E. Were the characteristics and results if the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
N	Were the sources of heterogeneity explored?
Comments	Baseline characteristics and use of transfusion protocol not reported.
Overall	Fair
assessment	

Citation	Segal JB, Blasco-Colmenares E, Norris EJ, and Guallar E. (2004) Preoperative acute
	normovolemic hemodilution: A meta-analysis. Transfusion 44:632-644.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
N	D. Was a quality assessment of included studies undertaken?
N	E. Were the characteristics and results if the individual studies appropriately summarised?
N	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	Authors did not analyse data by surgery type or use of transfusion protocol.
Overall	Fair
assessment	

Level II evidence

Citation	Akhlagh SH, Chohedri AH, Bazojoo A, and Nemati MH. (2007) A comparison of total amount of blood needed in patients taking autologous or homologous blood transfusion in coronary artery bypass grafting: A clinical randomized case-control trial. Pakistan Journal of Medical Sciences 23:542-545.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
NR	D. Were all randomised patients included in the analysis?
N	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Although the study reports the total number of patients in the study it does not specify the numbers randomised to treatment arms. No transfusion protocol was reported.
Overall	Poor
assessment	

Citation	Bennett J, Haynes S, Torella F, Grainger H, and McCollum C. (2006) Acute normovolemic hemodilution in moderate blood loss surgery: A randomized controlled trial. Transfusion 46:1097-1103.
NR	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Neither the anaesthetist nor the surgical team could be blinded.
Overall	Fair
assessment	

Citation	Casati V, Speziali G, D'Alessandro C, Cianchi C, Antonietta Grasso M, Spagnolo S, and Sandrelli L. (2002) Intraoperative low-volume acute normovolemic hemodilution in adult open-heart surgery. Anesthesiology 97:367-373.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	
Overall	Poor
assessment	

Citation	Casati V, Benussi S, Sandrelli L, Grasso MA, Spagnolo S, and D'Angelo A. (2004) Intraoperative Moderate Acute Norvolemic Hemodilution Associated with a Comprehensive Blood-Sparing Protocol in Off-Pump Coronary Surgery. Anesthesia and Analgesia 98:1217-1223.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	
Overall	Fair
assessment	

Citation	Hohn L, Schweizer A, Licker M, and Morel DR. (2002) Absence of beneficial effect of acute normovolemic hemodilution combined with aprotinin on allogeneic blood transfusion requirements in cardiac surgery. Anesthesiology 96:276-282.
NR	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	
Overall	Poor
assessment	

Citation	Friesen RH, Perryman KM, Weigers KR, Mitchell MB, and Friesen RM. (2006) A trial of fresh autologous whole blood to treat dilutional coagulopathy following cardiopulmonary bypass
	in infants. Paediatric Anaesthesia 16:429-435.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	
Overall	Fair
assessment	

Citation	Jarnagin WR, Gonen M, Maithel SK, Fong Y, DAngelica MI, Dematteo RP, Grant F, Wuest D, Kundu K, Blumgart LH, and Fischer M. (2008) A prospective randomized trial of acute normovolemic hemodilution compared to standard intraoperative management in patients undergoing major hepatic resection. Annals of Surgery 248:360-368.
NR	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
Υ	F. Were any subgroup analyses carried out?
Comments	
Overall	Fair
assessment	

Citation	Juelsgaard P, Moller MB, and Larsen UT. (2002) Preoperative acute normovolaemic hemodilution (ANH) in combination with hypotensive epidural anaesthesia (HEA) during knee arthroplasty surgery. No effect on transfusion rate. A randomized controlled trial [ISRCTN87597684]. BMC Anesthesiology 2.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	
Overall	Fair
assessment	

Citation	Lim YJ, Kim CS, Bahk JH, Ham BM, and Do SH. (2003) Clinical trial of esmolol-induced controlled hypotension with or without acute normovolemic hemodilution in spinal surgery. Acta Anaesthesiologica Scandinavica 47:74-78.
NR	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
NA	F. Were any subgroup analyses carried out?
Comments	
Overall	Fair
assessment	

Citation	Matot I, Scheinin O, Jurim O, and Eid A. (2002) Effectiveness of acute normovolemic hemodilution to minimize allogeneic blood transfusion in major liver resections. Anesthesiology 97:794-800.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	
Overall	Fair
assessment	

Citation	Obasi C, Arendt J, and Antoszewski Z. (2006) An assessment of the efficacy of preoperative controlled haemodilution in the perioperative management of patients including the elderly. Chirurgia Polska 8:111-124.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
N	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Very little methodological detail. Assessed multiple surgery types without evaluating heterogeneity. No transfusion protocol.
Overall	Poor
assessment	

Citation	Sanders G, Mellor N, Rickards K, Rushton A, Christie I, Nicholl J, Copplestone A, and Hosie K. (2004) Prospective randomized controlled trial of acute normovolaemic haemodilution in major gastrointestinal surgery. British Journal of Anaesthesia 93:775-781.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
Υ	F. Were any subgroup analyses carried out?
Comments	
Overall	Fair
assessment	

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Citation	Saricaoglu F, Akinci SB, and Aypar U. (2005) The effect of acute normovolemic hemodilution and acute hypervolemic hemodilution on coagulation and allogeneic transfusion. Saudi Medical Journal 26:792-798.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
NA	F. Were any subgroup analyses carried out?
Comments	
Overall	Good
assessment	

Citation	Wolowczyk L, Nevin M, Smith FCT, Baird RN, and Lamont PM. (2003) Haemodilutional effect of standard fluid management limits the effectiveness of acute normovolaemic haemodilution in AAA surgery - Results of a pilot trial. European Journal of Vascular and Endovascular 26:405-411.
NR	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
NA	F. Were any subgroup analyses carried out?
Comments	
Overall	Fair
assessment	

Intervention 2 – Intraoperative cell salvage

Level I evidence

Citation	Carless P, Moxey A, O'Connell D, and Henry D. (2004) Autologous transfusion techniques: A
	systematic review of their efficacy. Transfusion Medicine 14:123-144.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
N	E. Were the characteristics and results if the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	Number of participants in studies not reported
Overall	Fair
assessment	

Citation	Carless PA, Henry DA, Moxey AJ, O'connell DL, Brown T, and Fergusson DA. (2006) Cell
	salvage for minimising perioperative allogeneic blood transfusion. Cochrane database of
	reviews (Online)CD001888.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results if the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	Includes intra- and postoperative data
Overall	Good
assessment	

Citation	Davies L, Brown TJ, Haynes S, Payne K, Elliott RA, and McCollum C. (2006) Cost- effectiveness of cell salvage and alternative methods of minimising perioperative allogeneic blood transfusion: A systematic review and economic model. Health Technology Assessment 10:1-114.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results if the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	
Overall	Good
assessment	

Citation	Huet C, Salmi R, Fergusson D, Koopman-Van Gemert AWMM, Rubens F, and Laupacis A. (1999) A meta-analysis of the effectiveness of cell salvage to minimize perioperative allogeneic blood transfusion in cardiac and orthopedic surgery. Anesthesia and Analgesia 89:861-869.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results if the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	
Overall	Good
assessment	

Citation	Takagi H, Sekino S, Kato T, Matsuno Y, and Umemoto T. (2007) Intraoperative autotransfusion in abdominal aortic aneurysm surgery: meta-analysis of randomized controlled trials (Structured abstract). Archives of Surgery 142:1098-1101.
Υ	A. Was a clinical question clearly defined?
N	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
	E. Were the characteristics and results if the individual studies appropriately summarised?
	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	Specific search strategy not described,
Overall	Fair
assessment	

Citation	Bowley DM, Barker P, and Boffard KD. (2006) Intraoperative blood salvage in penetrating
	abdominal trauma: A randomised, controlled trial. World Journal of Surgery 30:1074-1080.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	It is unclear whether the analysis was ITT
Overall	Fair
assessment	

Citation	Damgaard S and Steinbruchel DA. (2006) Autotransfusion with cell saver for off-pump coronary artery bypass surgery: A randomized trial. Scandinavian Cardiovascular Journal 40:194-198.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	The surgical and anesthetic team were blinded during the operation, but not after. The ICU staff were blinded.
Overall	Good
assessment	

Citation	Goel P, Pannu H, Mohan D, and Arora R. (2007) Efficacy of cell saver in reducing homologous blood transfusions during OPCAB surgery: A prospective randomized trial. Transfusion Medicine 17:285-289.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
NA	F. Were any subgroup analyses carried out?
Comments	
Overall	Fair
assessment	

Citation	Mercer KG, Spark JI, Berridge DC, Kent PJ, and Scott DJA. (2004) Randomized clinical trial of intraoperative autotransfusion in surgery for abdominal aortic aneurysm. British Journal of Surgery 91:1443-1448.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
NA	F. Were any subgroup analyses carried out?
Comments	
Overall	Good
assessment	

Citation	Murphy GJ, Rogers CS, Lansdowne WB, Channon I, Alwair H, Cohen A, Caputo M, and Angelini GD. (2005) Safety, efficacy, and cost of intraoperative cell salvage and autotransfusion after off-pump coronary artery bypass surgery: A randomized trial. Journal of Thoracic and Cardiovascular 130:20-28.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	
Overall	Fair
assessment	

Citation	Niranjan G, Asimakopoulos G, Karagounis A, Cockerill G, Thompson M, and Chandrasekaran V. (2006) Effects of cell saver autologous blood transfusion on blood loss and homologous blood transfusion requirements in patients undergoing cardiac surgery on- versus off-cardiopulmonary bypass: a randomised trial. European Journal of Cardio-thoracic Surgery 30:271-277.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
Υ	F. Were any subgroup analyses carried out?
Comments	
Overall	Good
assessment	

Citation	Selo-Ojeme DO and Feyi-Waboso PA. (2007) Salvage autotransfusion versus homologous blood transfusion for ruptured ectopic pregnancy. International Journal of Gynecology and
	Obstetrics 96:108-111.
NR	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
NA	F. Were any subgroup analyses carried out?
Comments	No transfusion protocol was used.
Overall	Fair
assessment	

Citation	Wiefferink A, Weerwind PW, van Heerde W, Teerenstra S, Noyez L, de Pauw BE, and Brouwer RM. (2007) Autotransfusion management during and after cardiopulmonary bypass alters fibrin degradation and transfusion requirements. The Journal of Extra-corporeal Technology 39:66-70.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
N	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
N	E. Were the statistical methods appropriate?
NA	F. Were any subgroup analyses carried out?
Comments	No transfusion protocol was reported, low
Overall	Fair
assessment	

Citation	Zhang XL, Qian BH, and Luo QF. (2004) Effects of blood transfusion modes during perioperative period on prognosis of patients with scoliosis. Chinese Journal of Clinical Rehabilitation 8:7308-7310.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
N	E. Were the statistical methods appropriate?
NA	F. Were any subgroup analyses carried out?
Comments	No transfusion protocol was used; insufficient detail provided for many of the outcomes; surgical procedure not described; although paper states that there was no difference in baseline characteristics, the baseline values themselves were not reported.
Overall	Poor
assessment	

Intervention 3 – Perioperative acute normovolemic haemodilution combined with intraoperative cell salvage

Level II evidence

Citation	Haynes SL, Torella F, Wong JCL, Dalrymple K, James M, and McCollum CN. (2002) Economic evaluation of a randomized clinical trial of haemodilution with cell salvage in aortic surgery. British Journal of Surgery 89:731-736.
NA	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
NA	B. Was the study double-blinded?
NA	C. Were patient characteristics and demographics similar between treatment arms at baseline?
NA	D. Were all randomised patients included in the analysis?
NA	E. Were the statistical methods appropriate?
NA	F. Were any subgroup analyses carried out?
Comments	Authors did not conduct subgroup analysis by type of aortic surgery.
Overall	Fair (See Wong et al [2002])
assessment	

Citation	McGill N, O'Shaughnessy D, Pickering R, Herbertson M, and Gill R. (2002) Mechanical methods of reducing blood transfusion in cardiac surgery: Randomised controlled trial.
	British Medical Journal 324:1299-1302.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
N	E. Were the statistical methods appropriate?
Υ	F. Were any subgroup analyses carried out?
Comments	
Overall	Fair
assessment	

Citation	Wong JC, Torella F, Haynes SL, Dalrymple K, Mortimer AJ, McCollum CN, and ATIS I. (2002) Autologous versus allogeneic transfusion in aortic surgery: a multicenter randomized clinical trial. Annals of surgery 235:145-151.
NR	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
Υ	F. Were any subgroup analyses carried out?
Comments	A transfusion protocol was used
Overall	Fair
assessment	

Intervention 4 – Postoperative cell salvage

Citation	Carless P, Moxey A, O'Connell D, and Henry D. (2004) Autologous transfusion techniques: A
	systematic review of their efficacy. Transfusion Medicine 14:123-144.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
N	E. Were the characteristics and results if the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	Number of participants in studies not reported
Overall	Fair
assessment	

Citation	Carless PA, Henry DA, Moxey AJ, O'connell DL, Brown T, and Fergusson DA. (2006) Cell salvage for minimising perioperative allogeneic blood transfusion. Cochrane database of reviews (Online)CD001888.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results if the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	Includes intra- and postoperative data
Overall	Good
assessment	

Citation	Davies L, Brown TJ, Haynes S, Payne K, Elliott RA, and McCollum C. (2006) Cost- effectiveness of cell salvage and alternative methods of minimising perioperative allogeneic blood transfusion: A systematic review and economic model. Health Technology Assessment 10:1-114.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results if the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	
Overall	Good
assessment	

Citation	Duffy G and Neal KR. (1996) Differences in postoperative infection rates between patients receiving autologous and allogeneic blood transfusion: a meta-analysis of published randomized and nonrandomized studies (Structured abstract). Transfusion Medicine 6:325-328.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
N	D. Was a quality assessment of included studies undertaken?
N	E. Were the characteristics and results if the individual studies appropriately summarised?
NA	Were the methods for pooling the data appropriate?
NA	Were the sources of heterogeneity explored?
Comments	Limited trial info provided
Overall	Fair
assessment	

Citation	Huet C, Salmi R, Fergusson D, Koopman-Van Gemert AWMM, Rubens F, and Laupacis A. (1999) A meta-analysis of the effectiveness of cell salvage to minimize perioperative allogeneic blood transfusion in cardiac and orthopedic surgery. Anesthesia and Analgesia 89:861-869.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
N	E. Were the characteristics and results if the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
N	Were the sources of heterogeneity explored?
Comments	
Overall	Fair
assessment	

Citation	Amin A, Watson A, Mangwani J, Nawabi D, Ahluwalia R, and Loeffler M. (2008) A prospective randomised controlled trial of autologous retransfusion in total knee replacement. Journal of
	Bone and Joint - Series B 90:451-454.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	
Overall	Fair
assessment	

Citation	Cheng SC, Hung TS, and Tse PY. (2005) Investigation of the use of drained blood reinfusion after total knee arthroplasty: a prospective randomised controlled study. Journal of
	Orthopaedic Surgery (Hong Kong) 13:120-124.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Near the end of each operation, the corresponding envelope for each patient was opened, and the surgeon was informed at the time of drain insertion to achieve a single-blind effect. The control group had a larger proportion of patients with a pre-morbid condition (65% vs 54%) and a larger proportion of males compared with the reinfusion group (35% vs 23%). However, these differences were not significant.
Overall	Fair
assessment	

Citation	Zacharopoulos A, Apostolopoulos A, and Kyriakidis A. (2007) The effectiveness of reinfusion after total knee replacement. A prospective randomised controlled study. International Orthopaedics 31:303-308.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
N	C. Were patient characteristics and demographics similar between treatment arms at baseline?
NR	D. Were all randomised patients included in the analysis?
N	E. Were the statistical methods appropriate?
NA	F. Were any subgroup analyses carried out?
Comments	The study did not report use of a transfusion protocol. Clinical outcomes and patient demographics were insufficiently reported. It is unclear whether all patients were included in the analysis. The authors report results as "average" without clarifying whether they are referring to mean or median.
Overall	Poor
assessment	

Intervention 5 – Deliberate induced hypotension

Level I evidence

Citation	Paul JE, Ling E, Lalonde C, Thabane L. Deliberate hypotension in orthopedic surgery reduces blood loss and transfusion requirements: A meta-analysis of randomized controlled trials. Can J Anesth 2007;54(10):799-810.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results if the individual studies appropriately summarised?
Υ	F. Were the methods for pooling the data appropriate?
Υ	G. Were the sources of heterogeneity explored?
Comments	
Overall	Good
assessment	

Citation	Boldt J, Weber A, Mailer K, Papsdorf M, Schuster P. Acute normovolaemic haemodilution vs controlled hypotension for reducing the use of allogeneic blood in patients undergoing
	radical prostatectomy. Br J Anaesth 1999;82(2):170-174.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
Υ	F. Were any subgroup analyses carried out?
Comments	
Overall	Good.
assessment	

Citation	Elsharnouby NM, Elsharnouby MM. Magnesium sulphate as a technique of hypotensive
	anaesthesia. Br J Anaesth 2006;96(6):727-731.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
Υ	F. Were any subgroup analyses carried out?
Comments	The statistical analyses were appropriate and presented well. However, it does not provide an assessment/discussion of possible limitations or bias in this study.
Overall	Good
assessment	

Citation	Fredin H, Gustafson C, Rosberg B. Hypotensive anesthesia, thromboprophylaxis and postoperative thromboembolism in total hip arthroplasty. Acta Anaesthesiol Scand 1984;28(5):503-507.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	However, it is lacking details regarding the randomisation and blinding procedure. Statistical analyses performed were appropriate and well presented. Discussion did not address the presence of possible biases in the study.
Overall	Fair
assessment	

Citation	Jacobi KE, Bohm BE, Rickauer AJ, Jacobi C. Moderate controlled hypotension with sodium nitroprusside does not improve surgical conditions or decrease blood loss in endoscopic sinus surgery. J Clin Anesth 2000;12(3):202-207.
NR	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
NR	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
Υ	F. Were any subgroup analyses carried out?
Comments	This study did not include a description of the blinding methods employed, if any. Modest sample size (n=32).
Overall	Fair
assessment	

Citation	Karakaya D, Ustun E, Tur A, Baris S, Sarihasan B, Sahinoglu H, Guldogus F. Acute normovolemic hemodilution and nitroglycerin-induced hypotension: Comparative effects on tissue oxygenation and allogeneic blood transfusion requirement in total hip arthroplasty. J Clin Anesth 1999;11(5):368-374.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
NR	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
N	E. Were the statistical methods appropriate?
Υ	F. Were any subgroup analyses carried out?
Comments	This RCT did not describe the blinding method employed, if any. Due to the small sample size (n=10 per group), non-parametric methods should have been used.
Overall	Fair
assessment	

Citation	Kop EC, Spauwen PHM, Kouwenberg PPGM, Heymans FJM, van Beem HBH. Influence of controlled hypotension versus normotension on amount of blood loss during breast reduction. J Plast Reconstr Aesthetic Surg 2009;62(2):200-205.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	The discussion included an assessment of possible limitations such as the in the measurement of intraoperative blood loss.
Overall	Good
assessment	

Citation	O'Connor PJ, Hanson J, Finucane BT. Induced hypotension with epidural/general anesthesia reduces transfusion in radical prostate surgery. Can J Anesth 2006;53(9):873-880.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
Υ	F. Were any subgroup analyses carried out?
Comments	Anaesthesiologists were not blinded, however, only 23% of the transfusions were decided by the anaesthesiologist. The remaining 77% of transfusions were initiated postoperatively by non-study personnel.
Overall	Good.
assessment	

Citation	Piper SN, Suttner SW, Maleck WH, Kumle B, Haisch G, Boldt J. Effects of sodium nitroprusside-induced controlled hypotension on pancreatic function assessed by pancreatitis-associated protein in patients undergoing radical prostatectomy. Eur J
V	Anaesthesiol 2002;19(8):609-613.
Y	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
NR	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
Υ	F. Were any subgroup analyses carried out?
Comments	This study did not include a description of the blinding methods employed, if any. Also, as the primary aim of this study was not blood loss/transfusion, the authors did not discuss these outcomes in detail.
Overall	Good.
assessment	

Citation	Sood S, Jayalaxmi TS, Vijayaraghavan S, Nundy S. Use of sodium nitroprusside induced hypotensive anaesthesia for reducing blood loss in patients undergoing lienorenal shunts for portal hypertension. Br J 1987;74(11):1036-1038.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
NR	B. Was the study double-blinded?
N	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	The study provides no assessment/discussion of possible limitations or bias. The final study sample size of 18 patients is small, however statistical significance was achieved in the analysis.
Overall	Fair
assessment	

Citation	Suttner SW, Piper SN, Lang K, Huttner I, Kumle B, Boldt J. Cerebral effects and blood sparing efficiency of sodium nitroprusside-induced hypotension alone and in combination with acute normovolaemic haemodilution. Br J Anaesth 2001;87(5):699-705.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
Υ	F. Were any subgroup analyses carried out?
Comments	This RCT included a modest number of subjects, although initial power calculations suggest that the study was sufficiently powered. Information as to the method of randomising and blinding was not specified. The analyses performed were appropriate and the results presented clearly.
Overall	Good
assessment	

Intervention 6 – Prevention of hypothermia

Level I evidence

Citation	Mahoney CB, Odom J. Maintaining intraoperative normothermia: a meta-analysis of
	outcomes with costs (Structured abstract). AANA J 1999;67:155-164.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
N	C. Were the inclusion criteria appropriate and applied in an unbiased way?
N	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results of the individual studies appropriately summarised?
N	F. Were the methods for pooling the data appropriate?
N	G. Were the sources of heterogeneity explored?
Comments	The inclusion of non-randomised trials (3/18) and the lack of information on the allocation method
	and blinding of individual studies diminish the quality of this study.
Overall	Poor
assessment	

Citation	Rajagopalan S, Mascha E, Na J, Sessler Dl. The effects of mild perioperative hypothermia on
	blood loss and transfusion requirement. Anesthesiology 2008;108(1):71-77.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
N	E. Were the characteristics and results of the individual studies appropriately summarised?
Υ	F. Were the methods for pooling the data appropriate?
Υ	G. Were the sources of heterogeneity explored?
Comments	Detailed characteristics of included studies were absent; quality scores were assigned for each study. The statistical analyses were well conducted and clearly presented. The presence of publication bias and study effect were also examined.
Overall	Good
assessment	

Citation	Scott EM, Buckland R. A systematic review of intraoperative warming to prevent postoperative complications (Structured abstract). AORN Journal 2006;83:1090-1104.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results of the individual studies appropriately summarised?
N	F. Were the methods for pooling the data appropriate?
N	G. Were the sources of heterogeneity explored?
Comments	This review provides clear description of the randomisation, inclusion and exclusion criteria, and quality assessment of the included studies. However, the pooled estimates for morbid cardiac events were derived from just two studies, while the need for blood transfusion was derived from three studies. No assessment of heterogeneity or publication bias was performed.
Overall	Fair
assessment	

Citation	Jeong SM, Hahm KD, Jeong YB, Yang HS, Choi IC. Warming of intravenous fluids prevents hypothermia during off-pump coronary artery bypass graft surgery. Journal of cardiothoracic and vascular anesthesia 2008;22:67-70.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
N	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	The authors recognise that the small sample size was likely underpowered to detect changes in clinical data, as the study was designed to be powered to detect a change in patient temperature. Investigators were not blinded the treatment group of the subjects, this may have led to bias.
Overall	Poor.
assessment	

Citation	Kim YS, Lee JY, Yang SC, Song JH, Koh HS, Park WK. Comparative Study of the Influence of Room-Temperature and Warmed Fluid Irrigation on Body Temperature in Arthroscopic Shoulder Surgery. Arthroscopy J Arthroscopic Relat Surg 2009;25(1):24-29.
NR	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
Υ	F. Were any subgroup analyses carried out?
Comments	This RCT had clearly defined research questions and methods. It did not clearly describing the blinding or randomisation methods used. Based on the results of previous studies, their power calculation indicated that they had over 80% power.
Overall	Fair
assessment	

Citation	Melling AC, Ali B, Scott EM, Leaper DJ. Effects of preoperative warming on the incidence of wound infection after clean surgery: A randomised controlled trial. Lancet 2001;358(9285):876-
	880.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
NR	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
Υ	F. Were any subgroup analyses carried out?
Comments	Prospective power calculations indicated that the sample size provided 90% power to detect a 5% change in infection rates. Statistical analyses performed were appropriate, with multivariate analysis used to identify possible risk factors.
Overall	Good
assessment	

Citation	Yau TM, Carson S, Weisel RD, Ivanov J, Sun Z, Yu R, Glynn MF, Teasdale SJ. The effect of warm heart surgery on postoperative bleeding. The Journal of thoracic and cardiovascular surgery 1992;103:1155-1162.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
N	E. Were the statistical methods appropriate?
Υ	F. Were any subgroup analyses carried out?
Comments	Antifibrinolytic therapy was concurrently used by some patients in this study, and was not controlled for in this study; however, the use of such therapy had no bearing on the assignment to treatment group and as such would have had a non-differential effect, if any. The authors recognised that their sample size may have been underpowered to detect differences between treatment groups.
Overall	Fair
assessment	

Citation	Zhao J, Luo AL, Xu L, Huang YG. Forced-air warming and fluid warming minimize core hypothermia during abdominal surgery. Chinese medical sciences journal / Chinese Academy
	of Medical Sciences 2005;20:261-264.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	This RCT did not provide a description of the blinding or randomisation methods employed.
Overall	Fair
assessment	

Intervention 7 – Point-of-care testing using thromboelastography

Citation	Ak, K., Isbir, SC., et al., Thromboblastography-based algorithm reduces blood product use
	after elective CABG: a prospective randomised study. J Card Surg 2009;24:404-410.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
NA	F. Were any subgroup analyses carried out?
Comments	Anaesthesiologist performing transfusion was blinded to the patient's group assignment.
Overall	Fair
assessment	

Citation	Avidan M.S., Alcock E.L. et al. Comparison of structured use of routine laboratory tests or near-patient assessment with clinical judgement in the management of bleeding after cardiac surgery. British Journal of Anaesthesia. 2004; 2:176-86.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
NA	F. Were any subgroup analyses carried out?
Comments	
Overall	Fair
assessment	

Citation	Royston D. and von Kier S. Reduced haemostatic factor transfusion using heparinise- modified thromboelastography during cardiopulmonary bypass. British Journal of
	Anaesthesia. 2001; 4:575-8.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Uncertain	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	
Overall	Poor
assessment	

Citation	Shore-Lesserson L., Manspeizer H.E. et al. Thromboelastography-guided transfusion algorithm reduces transfusions in complex cardiac surgery. Anesthesia and analgesia. 1999; 88:312-9.
Unclear	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	All staff appointed with caring for the patient directly were blinded.
Overall	Fair
assessment	

Citation	Westbrook AJ., Olsen J. et al. Protocol based on thrombolestaograph (TEG) out-performs physician preference using laboratory coagulation tests to guide blood replacement during
	and after cardiac surgery: a pilot study. Heart, Lung and Circulation.2009;18:277-288.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
NA	F. Were any subgroup analyses carried out?
Comments	Surgeons were blinded to the method of haemostasis.
Overall	Fair
assessment	

Citation	Avidan M.S., Alcock E.L. et al. Comparison of structured use of routine laboratory tests or near-patient assessment with clinical judgement in the management of bleeding after cardiac
	surgery. British Journal of Anaesthesia. 2004; 2:176-86.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
NA	F. Were any subgroup analyses carried out?
Comments	
Overall	Fair
assessment	

Citation	Spalding, GJ., Hartrumpf, M. et al. Cost reduction of peri operative coagulation management in cardiac surgery: value of "bedside" thrombelastography (ROTEM). Eur J Cardiothorac Surg 2007;31:1052-1057.
Υ	A. How were subjects selected for the 'new' intervention?
Υ	B. How were subjects selected for the comparison or control group?
Υ	C. Does the study adequately control for demographic characteristics, clinical features and other potential confounding variables in the study design or analysis?
N	D. Was the measurement of outcomes unbiased (ie, blinded to treatment group and comparable across groups?)
N	E. Was follow-up long enough for outcomes to occur?
Υ	F. Was follow-up complete and were there exclusions from analysis?
Comments	
Overall	Fair
assessment	

Intervention 8 – Administration of antifibrinolytics & DDAVP

Citation	Abrishami A, Chung F, Wong J (2009) Topical application of antifibrinolytic drugs for on-
	pump cardiac surgery: a systematic review and meta-analysis. Can J Anesth 56: 202-212.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results of the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	
Overall	Good
assessment	

Citation	Brown JR, Birkmeyer NJO, O'Connor GT (2007) Meta-analysis comparing the effectiveness
	and adverse outcomes of antifibrinolytic agents in cardiac surgery. Circulation 115: 2801-
	2813.
N	A. Was a clinical question clearly defined?
N	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
N	E. Were the characteristics and results of the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	Medline search only conducted. Given the funnel plot indicated potential publication bias it may have been wise to expand the search to identify additional studies. No reporting of results of individual included studies. A number of data extraction errors were identified when checking one of the subgroup analyses.
Overall	Fair
assessment	

Citation	Carless PA, Stokes BJ, Moxey AJ, Henry DA (2004) Desmopressin use for minimising perioperative allogenic blood transfusion. Cochrane Database of Systematic Reviews. Issue 1. Article No.: CD001884. DOI: 10.1002/14651858.CD001884.pub2.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results of the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	Included studies generally considered to be of poor methodological quality.
Overall	Good
assessment	

Citation	Crescenzi G, Landoni G, Biondi-Zoccai G et al (2008) Desmopressin reduces transfusion needs after surgery: a meta-analysis of randomized clinical trials. Anesthesiology 109: 1063-1076.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
N	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results of the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
N	Were the sources of heterogeneity explored?
Comments	Reasonable literature search and selection and extraction of data. No quality assessment undertaken so limited quality information available for individual studies. No exploration of reasons for heterogeneity carried out.
Overall	Fair
assessment	

Citation	Gurusamy KS, Sharma D, Davidson BR (2009) Pharmacological interventions to decrease blood loss and blood transfusion requirements for liver resection. Cochrane Database of Systematic Reviews. 009,Issue 4.Art.No.: CD008085. DOI:10.1002/14651858. CD008085.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results of the individual studies appropriately summarised?
NA	Were the methods for pooling the data appropriate?
NA	Were the sources of heterogeneity explored?
Comments	Comprehensive literature search carried out. Quality assessment undertaken. Single study only available for each comparison. Authors note all studies at high risk of bias.
Overall assessment	Good

Citation	Henry DA, Carless PA, Moxey AJ et al (2007) Antifibrinolytic use for minimising perioperative allogenic blood transfusion. Cochrane Database of Systematic Reviews 2007, Issue 4. Art. No.: CD001886. DOI: 10.1002/14651858.CD001886.pub2.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results of the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	Comprehensive literature search carried out. Quality assessment undertaken. Subgroup analyses performed on <i>a priori</i> categories including surgery, transfusion protocol, dose and trial quality.
Overall	Good
assessment	

Citation	Henry DA, Carless PA, Fergusson D, et al (2009) The safety of aprotinin and lysine-derived antifibrinolytic drugs in cardiac surgery: a meta-analysis. CMAJ 180(2): 183-193.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
N	E. Were the characteristics and results of the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	Comprehensive literature search carried out. Some missing data on individual studies but the majority of these were provided in the Henry 2007 review.
Overall	Good
assessment	

Citation	Kagoma YK, Crowther MA, Douketis J et al (2009) Use of antifibrinolytic therapy to reduce transfusion in patients undergoing orthopaedic surgery: a systematic review of randomized
	trials. Thrombosis Research 123: 687-696.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results of the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	Included studies generally considered to be of good methodological quality.
Overall	Good
assessment	

Citation	Systematic review: Kongnyuy EJ, Wiysonge CS (2009) Interventions to reduce haemorrhage during myomectomy for fibroids. Cochrane Database of Systematic Reviews 2009, Issue 3. Art. No.: CD005335. DOI: 10.1002/14651858.CD005335.pub3. Single included RCT: Caglar GS, Tasci Y, Kayikcioglu F et al (2008) Intravenous tranexamic acid use in myomectomy: a prospective randomised double-blind placebo controlled study. European Journal of Obstetrics, Gynecology and Reproductive Biology 137(): 227-231.
Systematic revi	
Y	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results of the individual studies appropriately summarised?
NA	Were the methods for pooling the data appropriate?
NA	Were the sources of heterogeneity explored?
Comments	Cochrane review; extensive literature search; quality assessment of included studies; appropriate analysis.
Overall assessment	Good
RCT	
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Computer generated randomisation code; treatments in sequentially numbered identical containers; patients, surgeons and anaesthetists blinded to treatment allocation; full follow-up of patients.
Overall	Good
assessment	

Citation	Liu C-M, Chen J, Wang X-H (2008) Requirements for liver transfusion and postoperative outcomes in orthotopic liver transplantation: a meta-analysis on aprotinin. World J Gatroenterol 14(9): 1425-1429.
Υ	A. Was a clinical question clearly defined?
N	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
N	D. Was a quality assessment of included studies undertaken?
N	E. Were the characteristics and results of the individual studies appropriately summarised?
N	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	Inadequate search; included one non-RCT and one active-controlled RCT; no quality assessment; no details on individual studies provided.
Overall	Poor
assessment	

Citation	McIlroy DR, Myles PS, Phillips LE, Smith JA (2009) Antifibrinolytics in cardiac surgical patients receiving aspirin: a systematic review and meta-analysis.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results of the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	Comprehensive literature search carried out. Quality assessment undertaken. Appropriate analysis methods used. Subgroup and sensitivity analyses undertaken.
Overall	Good.
assessment	

Citation	Schouten ES, van de Pol A, Schouten ANJ et al (2009) The effect of aprotinin, tranexamic acid and aminocaproic acid on blood loss and use of blood products in major pediatric surgery: a meta-analysis. Pediatr Cri Care Med 10(2): 182-190.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
N	E. Were the characteristics and results of the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	Comprehensive literature search carried out. Quality assessment undertaken. Not all individual results and pooled results provided. Meta-regression analysis carried out using potential confounders for the cardiac studies due to heterogeneity (age, weight and time on cardiopulmonary bypass).
Overall	Fair.
assessment	

Citation	Tzortzopoulou A, Cepeda MS, Schumann R et al (2008) Antifibrinolytic agents for reducing blood loss in scoliosis surgery in children. Cochrane Database of Systematic Reviews 2008, Issue 3. Art. No.: CD006883. DOI: 10.1002/14651858.pub2.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results of the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
N	Were the sources of heterogeneity explored?
Comments	Comprehensive literature search carried out. Quality assessment undertaken. 4/6 studies considered to have low risk of bias and 2/6 considered to have moderate risk of bias.
Overall	Good
assessment	

Citation	Alvarez JC, Santiveri FX, Ramos I et al (2008) Tranexamic acid reduces blood transfusion in total knee arthroplasty even when a blood conservation program is applied. Transfusion 48: 519-525.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Computer-generated stratified randomisation method used; sealed envelopes. Double-blind. 15 patients excluded from analysis following randomisation, more in treatment than control group.
Overall	Fair
assessment	

Citation	Apostolakis E, Panagopoulos N, Koletsis EN, Crockett J, Stamou-Kouki H, Sourgiadaki E, Filos K, Dougenis D (2008) Influence of ultra low dose aprotinin on thoracic surgical operations: a prospective randomized trial. Journal of Cardiothoracic Surgery 3:14.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Randomisation; single-blind – treating anaesthetist aware of treatment allocation so potential for bias; small trial.
Overall	Fair
assessment	

Citation	Athanasiadis T, Beule AG, Wormald PJ (2007) Effects of topical antifibrinolytics in
	endoscopic sinus surgery: a pilot randomized controlled trial. Am J Rhinol 21: 737-742.
Unclear	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	The method of randomisation was not reported. The anaesthetist prepared treatments so unclear if
	this could have resulted in unblinding. Outcome rating scales used not validated.
Overall	Fair
assessment	

Citation	Berenholtz SM, Pham JC, Garrett-Mayer E et al (2009) effect of epsilon aminocaproic acid on red-cell transfusion requirements in major spinal surgery. Spine 34(19): 2096-2103.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Computer-generated, stratified randomisation method used. Double-blind. All patients included in analysis. Study was underpowered to detect a 1-unit difference in total blood transfusion.
Overall	Good.
assessment	

Citation	Chen CC, Wang CC, Wang, CP et al (2008) Prospective, randomized, controlled trial of tranexamic acid in patients who undergo head and neck surgery. Otolaryngology – Head and Neck Surgery 138: 762-767.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Computer-generated stratified randomisation method used. Double-blind. 7 patients (8%) excluded from analysis following randomisation.
Overall	Fair
assessment	

Citation	Choi WS, Irwin MG, Samman N (2009) The effect of tranexamic acid on blood loss during
	orthognathic surgery: a randomized controlled trial. J Oral Maxillofac Sug 67: 125-133.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Computer-generated stratified randomisation method used/sealed envelopes. Double-blind. 12
	patients (16%) excluded from analysis following randomisation.
Overall	Fair
assessment	

Citation	Colwell Jr CW, Chelly JE, Murkin JM, Stevens D, O'Keefe TJ, Hall R, Parvizi J (2007) Randomized study of aprotinin effect on transfusions and blood loss in primary THA. Clinical
	Orthopaedics and Related Research 465: 189-195.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Randomised; double-blind; all patients included in analysis.
Overall	Good
assessment	

Citation	Elwatidy S, Jamjoon Z, Elgamal E et al (2008) Efficacy and safety of prophylactic large dose of tranexamic acid in spine surgery: a prospective, randomized, double-blind, placebo-controlled study. Spine 33(24): 2577-2580.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Randomised using odd/even numbers so could be easily worked out; all subjects included in
	analysis.
Overall	Fair
assessment	

Citation	Fawzy H, Elmistekawy E, Bonneau D et al (2009) Can local application of tranexamic acid reduce post-coronary bypass surgery blood loss? A randomized controlled trial. Journal of Cardiothoracic Surgery 4:25.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Randomisation using random number tables; double-blind; full follow-up of patients
Overall	Good
assessment	

Citation	Gharabaghian M, Eghtesadi-Araghi P (2006) The efficacy of epsilon-aminocaproic acid and its timing in reducing blood loss in major cardiac coronary bypass surgery: a randomized double-blinded placebo-controlled study. International journal of Pharmacology 2(1): 131-135.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Study was described as randomised but no details of method and concealment of allocation was provided. Not stated whether all patients included in the analysis although this was likely.
Overall	Fair.
assessment	

Citation	Grant MC, Kon Z, Joshi A, Christenson E, Kallam S, Burris N, Gu J, Poston RS (2008) Is aprotinin safe to use in a cohort at increased risk for thrombotic events: results from a randomized, prospective trial in off-pump coronary artery bypass. Ann Thorac Surg 86: 815-822.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
N	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Computer-generated randomisation; double-blind; no baseline details reported; some patients not included in analysis.
Overall assessment	Fair

Citation	Jabalami M, Zakeri K (2006) Evaluation of topical tranexamic acid on intraoperative bleeding in endoscopic sinus surgery. Iran J Med Sci 31(4): 221-223.
NR	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
NR	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
NR	D. Were all randomised patients included in the analysis?
NR	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Poor reporting of randomisation, blinding, outcome assessment and follow-up
Overall	Poor
assessment	

Citation	Jimenez JJ, Iribarren JL, Lorente L et al (2007) Tranexamic acid attenuates inflammatory response in cardiopulmonary bypass surgery through blockade of fibrinolysis: a case
	control study followed by a randomized controlled trial. Critical care: 11 R117.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Randomisation using random number tables; double-blind; full follow-up of patients
Overall	Good
assessment	

Citation	Later AFL, Maas JJ, Engbers FHM et al (2009) Tranexamic acid and aprotinin in low- and intermediate risk cardiac surgery: a non-sponsored, double-blind, randomised placebo-controlled trial. European Journal of Cardiothoracic Surgery 36: 322-329.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Randomisation; double-blind; 9% of patients not included in analysis but reasonably large trial.
Overall	Good
assessment	

Citation	Leijdekkers VJ, Vahl AC, Mackaay AJC, Huijgens PC, Rauwerda JA (2006) Aprotinin does not diminish blood loss in elective operations for infrarenal abdominal aneurysms: a randomized, double-blind controlled trial. Ann Vasc Surg 20: 322-329.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Randomised; described as double-blind but no details given; all patients included in analysis but very small trial.
Overall	Fair
assessment	

Citation	Maddali MM, Rajakumar MC (2007) Tranexamic acid and primary coronary artery bypass surgery: a prospective study. Asian Cardiovascular and Thoracic Annals 15: 313-319.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Randomisation; double-blind; all patients included in analysis.
Overall	Good
assessment	

Citation	Mayur G, Purvi P, Ashoo G, Panjak D (2007) Efficacy of tranexamic acid in decreasing blood
	loss during and after cesarean section: a randomized case controlled prospective study. The
	Journal of Obstetrics and Gynecology of India 57(3): 227-230.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Unsecure method of randomisation, open-label.
Overall	Fair
assessment	

Citation	Mehr-Aein A, Sadeghi M, Madani-civi M (2007) Does tranexamic acid reduce blood loss in off-pump coronary artery bypass? Asian cardiovascular and Thoracic Annals 15: 285-289.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Randomised, double-blind, all patients included in analysis.
Overall	Good
assessment	

Citation	Mehraien A, Ghafari A, Mohammadi SS (2009) Effect of topical aprotinin on early postoperative bleeding and ICU stay after coronary artery bypass graft surgeries. Pakistan Journal of Biological Sciences 12(10): 813-816.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Randomised; double-blind; all patients included in analysis.
Overall	Good
assessment	

Citation	Nurözler F, Kutlu T, Küçük G (2008) Aprotinin for patients exposed to clopidogrel before off- pump coronary bypass. Asian Cardiovascular and Thoracic Annals 16: 483-487.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Randomised; double-blind; all patients included in analysis; small study.
Overall	Fair
assessment	

Citation	Sadegi M, Mehr-Aein A (2007) Does a single bolus dose of tranexamic acid reduce blood loss and transfusion requirements during hip fracture surgery? A prospective randomised double-blind study in 67 patients. Acta Medica Iranica 45(6): 437-442.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Randomised, double-blind, all patients included in analysis.
Overall	Good
assessment	

Citation	Sekhavat L, Tabatabah A, Dalili M et al (2009) Efficacy of tranexamic acid in reducing blood
	loss after cesarean section. The Journal of maternal-Fetal and Neonatal Medicine 22(1): 72-
	75.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
NA	F. Were any subgroup analyses carried out?
Comments	Unsecure method of randomisation, open-label.
Overall	Poor
assessment	

Citation	Taghaddomi RJ, Mirzaee A, Attar AS et al (2009) Tranexamic acid reduces blood loss in off- pump coronary artery bypass surgery. Journal of Cardiothoracic and Vascular Anaesthesia 23(3): 312-315.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Randomised, double-blind, 7.4% of randomised patients not included in the analysis.
Overall	Fair
assessment	

Citation	Wong J, El Beheiry H, Rampersaud YR et al (2008) Tranexamic acid reduces perioperative blood loss in adult patients having spinal fusion surgery. Anesth Analg 107: 1479-1486.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Randomised, double-blind, 2.4% of randomised patients not included in the analysis.
Overall	Good
assessment	

Intervention 9 – Appropriate patient positioning

Citation	De Sio M, Autorino R, Quarto G, Calabro F, Damiano R, Giugliano F, Mordente S, D'Armiento M. Modified Supine versus Prone Position in Percutaneous Nephrolithotomy for Renal Stones Treatable with a Single Percutaneous Access: A Prospective Randomized Trial. Eur Urol 2008;54(1):196-203.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
Υ	F. Were any subgroup analyses carried out?
Comments	Not possible for surgeons to be blinded of intervention.
Overall assessment	Good.

Citation	Ko MT, Chuang KC, Su CY. Multiple analyses of factors related to intraoperative blood loss and the role of reverse Trendelenburg position in endoscopic sinus surgery. Laryngoscope 2008;118(9):1687-1691.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Not possible for surgeons to be blinded of intervention.
Overall	Fair
assessment	

Citation	Ong SM, Taylor GJSC. Can knee position save blood following total knee replacement? Knee 2003;10(1):81-85.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
NR	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	
Overall	Fair
assessment	

Citation	Pace A, Yousef A. The effect of patient position on blood loss in primary cemented total hip arthroplasty. Arch Orthop Trauma Surg 2008;128(10):1209-1212.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Not possible to blind surgeon from patients' treatment group.
Overall	Fair
assessment	

Citation	Park CK. The effect of patient positioning on intraabdominal pressure and blood loss in spinal surgery. Anesth Analg 2000;91(3):552-557.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
Υ	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
Υ	F. Were any subgroup analyses carried out?
Comments	This study was well described. Appropriate statistical tests, such as the use of non-parametric tests were used, in light of the modest sample size (n=40).
Overall	Good
assessment	

Citation	Widman J, Isacson J. Lateral position reduces blood loss in hip replacement surgery: A prospective randomized study of 74 patients. Int Orthop 2001;25(4):226-227.
Υ	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
N	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	Not possible for the surgeon to be blinded to the treatment group of the patient.
Overall	Fair.
assessment	

Intervention 10 – Preoperative autologous donation

Level I evidence

Citation	Carless P, Moxey A, O'Connell D, and Henry D. (2004) Autologous transfusion techniques: A
	systematic review of their efficacy. Transfusion Medicine 14:123-144.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
N	E. Were the characteristics and results if the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	
Overall	Good
assessment	

Davies L, Brown TJ, Haynes S, Payne K, Elliott RA, and McCollum C. (2006) Cost- effectiveness of cell salvage and alternative methods of minimising perioperative allogeneic blood transfusion: a systematic review and economic model (Provisional abstract). Health Technology Assessment 10:1-228.
A. Was a clinical question clearly defined?
B. Was an adequate search strategy used?
C. Were the inclusion criteria appropriate and applied in an unbiased way?
D. Was a quality assessment of included studies undertaken?
E. Were the characteristics and results if the individual studies appropriately summarised?
Were the methods for pooling the data appropriate?
Were the sources of heterogeneity explored?
Good

Citation	Duffy G and Neal KR. (1996) Differences in postoperative infection rates between patients receiving autologous and allogeneic blood transfusion: a meta-analysis of published randomized and nonrandomized studies (Structured abstract). Transfusion Medicine 6:325-328.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
N	D. Was a quality assessment of included studies undertaken?
N	E. Were the characteristics and results if the individual studies appropriately summarised?
NA	Were the methods for pooling the data appropriate?
NA	Were the sources of heterogeneity explored?
Comments	Limited trial information provided
Overall	Fair
assessment	

Citation	Forgie MA, Wells PS, Laupacis A, and Fergusson D. (1998) Preoperative autologous donation decreases allogeneic transfusion but increases exposure to all red blood cell transfusion:
	Results of a meta- analysis. Archives of Internal Medicine 158:610-616.
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results if the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	
Overall	Good
assessment	

Citation	Gurusamy KS, Li J, Sharma D, and Davidson BR. (2009) Cardiopulmonary interventions to decrease blood loss and blood transfusion requirements for liver resection. Cochrane
	Database of Systematic Reviews: Reviews 2009. Issue. 4
Υ	A. Was a clinical question clearly defined?
Υ	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
Υ	D. Was a quality assessment of included studies undertaken?
Υ	E. Were the characteristics and results if the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
Υ	Were the sources of heterogeneity explored?
Comments	
Overall	Good
assessment	

Citation	Henry DA, Carless PA, Moxey AJ, O'Connell D, Forgie MA, Wells P, and Fergusson DA. (2001)						
	Preoperative autologous donation for minimising perioperative allogeneic blood transfusion.						
	Cochrane Database of Systematic Reviews 2001; Issue 4						
Υ	A. Was a clinical question clearly defined?						
Υ	B. Was an adequate search strategy used?						
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?						
Υ	D. Was a quality assessment of included studies undertaken?						
Υ	E. Were the characteristics and results if the individual studies appropriately summarised?						
Υ	Were the methods for pooling the data appropriate?						
Υ	Were the sources of heterogeneity explored?						
Comments							
Overall	Good						
assessment							

Citation	Laupacis A and Fergusson D. (1998) The efficacy of technologies to minimise perioperative					
	allogeneic transfusion (Structured abstract). Kluwer. Academic Publishers.17-36.					
Υ	A. Was a clinical question clearly defined?					
Υ	B. Was an adequate search strategy used?					
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?					
Υ	D. Was a quality assessment of included studies undertaken?					
N	E. Were the characteristics and results if the individual studies appropriately summarised?					
Υ	Were the methods for pooling the data appropriate?					
N	Were the sources of heterogeneity explored?					
Comments	Baseline characteristics not reported					
Overall	Fair					
assessment						

Citation	Vamvakas EC. (2002) Meta-analysis of randomized controlled trials investigating the risk of postoperative infection in association with white blood cell-containing allogeneic blood transfusion: The effects of the type of transfused red blood cell product and surgical setting. Transfusion Medicine Reviews 16:304-314.
Υ	A. Was a clinical question clearly defined?
NR	B. Was an adequate search strategy used?
Υ	C. Were the inclusion criteria appropriate and applied in an unbiased way?
N	D. Was a quality assessment of included studies undertaken?
N	E. Were the characteristics and results if the individual studies appropriately summarised?
Υ	Were the methods for pooling the data appropriate?
N	Were the sources of heterogeneity explored?
Comments	No reporting of transfusion protocol, allocation concealment, blinding, ITT analysis.
Overall	Poor
assessment	

Citation	Bouchard D, Marcheix B, Al Shamary S, Vanden Eynden F, Demers P, Robitaille D, Pellerin M, Perrault LP, and Carrier M. (2008) Preoperative autologous blood donation reduces the need for allogeneic blood products: A prospective randomized study. Canadian Journal of Surgery 51:422-427.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
Υ	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
N	F. Were any subgroup analyses carried out?
Comments	
Overall	Fair
assessment	

Citation	Hashimoto T, Kokudo N, Orii R, Seyama Y, Sano K, Imamura H, Sugawara Y, Hasegawa K, and Makuuchi M. (2007) Intraoperative blood salvage during liver resection: A randomized
	controlled trial. Annals of Surgery 245:686-691.
N	A. Was the allocation to treatment groups concealed from those responsible for recruiting subjects?
N	B. Was the study double-blinded?
Υ	C. Were patient characteristics and demographics similar between treatment arms at baseline?
N	D. Were all randomised patients included in the analysis?
Υ	E. Were the statistical methods appropriate?
NA	F. Were any subgroup analyses carried out?
Comments	No transfusion protocol was used
Overall	Poor
assessment	

Appendix F: Evidence summaries

Intervention 1 – Acute normovolemic haemodilution

Level I evidence

Citation						
Bryson GL, Laupa	acis A, and Wells GA. (eta-analysis. Anesthesi			emic hemodilu	ution reduce perior	perative allogeneic
Affiliation/Sourc	e of funds					
None declared						
Study design		Level of evidence			Location/setting	
SR Search conducted August 1996		I			NA	
Intervention			Comparator			
Intraoperative ANH: only those trials in which whole blood was withdrawn on the day of surgery and replaced with a crystalloid or colloid solution were considered to represent ANH.			Control group			
Population chara	acteristics					
Any patients unde	ergoing a surgical proce	edure.				
Length of follow	-up		Outcon	nes measure	d	
NR			Proportion of patients transfused with at least 1 unit of allogeneic blood in the perioperative period, volume of allogeneic blood transfused, volume of blood withdrawn during haemodilution, discharge haematocrit.			
INTERNAL VALI	DITY					
Allocation	Results	Blinding analy	Blinding analysis		measurement	Follow-up (ITT)
All trials were randomised (pseudorandom trials were excluded). The median Jadad score was 1 of a possible 5. Only five trials, all scoring 2, exceeded the median.	SR did not report baseline characteristics.			Fifteen trials did not specify a transfusion protocol or used a protocol that set different transfusion thresholds in the ANH and control groups.		NR
Overall quality a	ı ssessment (descripti [,]	ve)		l		<u>I</u>
Good						
RESULTS						

Outcome	Intervention group	Comparator group	Statistical significance
Mean volume of blood reserved, mL 24 trials (n=629; 629 ANH)	936		NA
Mean perioperative blood loss, mL 24 trials (n=1218; 629 ANH, 589 control)	1268	1348	NR
Patients exposed to at least 1 unit of allogeneic blood 16 trials (n=615; 308 ANH, 307 control)			OR (95% CI): 0.31 (0.15, 0.62) P<0.05 (Phet=0.013) ANH improves outcome
Patients exposed to at least 1 unit of allogeneic blood (cardiac surgery) 6 trials (n=266; 128 ANH, 138 control)			OR (95% CI): 0.51 (0.26, 0.99) P<0.05 (P <i>het</i> =0.945) ANH improves outcome
Patients exposed to at least 1 unit of allogeneic blood (orthopaedic surgery) 3 trials (n=174; 88 ANH, 86 control)			OR (95% CI): 1.00 (0.01, 4.47) P>0.05 (P <i>het</i> =0.032) ANH worsens outcome
Patients exposed to at least 1 unit of allogeneic blood (miscellaneous surgery) 7 trials (n=175; 92 ANH, 83 control)			OR (95% CI): 0.05 (0.01, 0.18) P<0.05 (P <i>het</i> =0.226) ANH improves outcome
Patients exposed to at least 1 unit of allogeneic blood (<1000 mL of blood withdrawn preoperatively in ANH group)			OR (95% CI): 0.43 (0.18, 1.02) P>0.05 (Phet=NR) No significant difference
Patients exposed to at least 1 unit of allogeneic blood (≥1000 mL of blood withdrawn preoperatively in ANH group)			OR (95% CI): 0.16 (0.04, 0.65) P<0.05 (Phet=NR) ANH improves outcome
Patients exposed to at least 1 unit of allogeneic blood (transfusion protocol)			OR (95% CI): 0.64 (0.31, 1.31) P>0.05 (Phet=NR) No significant difference
Patients exposed to at least 1 unit of allogeneic blood (no transfusion protocol)			OR (95% CI): 0.12 (0.04, 0.37) P<0.05 (P <i>het</i> =NR) ANH improves outcome
Units of allogeneic blood transfused 13 trials (n=568; 299 ANH, 269 control)			WMD (95% CI): -2.22 (-3.57, -0.86) P<0.05 (P <i>het</i> <0.001) ANH improves outcome

Units of allogeneic blood transfused (cardiac surgery) 6 trials (n=365; 197 ANH, 168 control)	WMD (95% CI): -2.83 (-5.34, -0.31) P<0.05 (Phet<0.001) ANH improves outcome	e
Units of allogeneic blood transfused (orthopaedic surgery) 2 trials (n=60; 30 ANH, 30 control)	WMD (95% CI): -1.54 (-4.41, 1.32) P>0.05 (Phet<0.001) No significant difference	e
Units of allogeneic blood transfused (miscellaneous surgery) 5 trials (n=143; 72 ANH, 71 control)	WMD (95% CI): -2.26 (-3.71, -0.80) P<0.05 (Phet=0.004) ANH improves outcome	e
Units of allogeneic blood transfused (<1000 mL of blood withdrawn preoperatively in ANH group)	WMD (95% CI): -2.30 (-3.79, -0.81) P<0.05 (Phet=NR) ANH improves outcome	e
Units of allogeneic blood transfused (≥1000 mL of blood withdrawn preoperatively in ANH group)	WMD (95% CI): -1.69 (-3.42, -0.03) P<0.05 (Phet=NR) ANH improves outcome	e
Units of allogeneic blood transfused (transfusion protocol)	WMD (95% CI): 0.25 (- 0.10) P>0.05 (Phet=NR) No significant difference	
Units of allogeneic blood transfused (no transfusion protocol)	WMD (95% CI): -3.01 (-3.47, -2.55) P<0.05 (Phet=NR) AMH improves outcom	ıe
Difference in perioperative blood loss, mL 13 trials (n=500; 245 ANH, 255 control)	WMD (95% CI): -117 (-58) P>0.05 (Phet<0.001) No significant difference	
Difference in perioperative blood loss (cardiac surgery), mL 7 trials (n=350; 169 ANH, 181 control)	WMD (95% CI): -233 (-5) P<0.05 (Phet<0.001) ANH had less perioper blood	
Difference in perioperative blood loss (orthopaedic surgery), mL 1 trial (n=31; 16 ANH, 15 control)	WMD (95% CI): 33 (-57 578) P>0.05 (Phet=NA) No significant difference	

Difference in perioperative		WMD (95% CI): -97 (-339,						
blood loss (miscellaneous		145)						
surgery), mL		P>0.05 (Phet=0.013)						
5 trials (n =119; 60 ANH, 59 control)		No significant difference						
Mortality		OR (95% CI): 0.67 (0.14,						
6 trials (n=170)		3.20)						
Myocardial infarction		OR (95% CI): 1.00 (0.06,						
2 trials (n=40)		17.07)						
DVP		OR (95% CI): 0.67 (0.14,						
2 trials (n=123)		3.20)						
Clinical importance	Clinical relevar	Clinical relevance						
EXTERNAL VALIDITY	•							
Generalisability								
Patients considered similar to guideline target populat	tion							
Applicability	Applicability							
All the studies included in this review were conducted in countries with well developed healthcare systems (not specifically Aus/NZ).								
Comments								

Abbreviations: ANH, acute normovolemic haemodilution; NA, not applicable; NR, not reported; SR, systematic review.

Carless P, Moxey A, O'Connell D, and Henry D. (2004) Autologous transfusion techniques: A systematic review of their efficacy. Transfusion Medicine 14:123-144.

Affiliation/Source of funds

The research was supported by a grant obtained from the National Health and Medical Research Council of Australia and a special purpose grant from the Hunter Area Pathology Service, Australia.

Study design	Level of evidence	Location/setting
Systematic review of RCTs and observational studies with meta-analysis	1	NA
Search conducted July 2002		

Intervention	Comparator			
Autologous transfusion techniques: preoperative Autologous blood deposit (PAD), ANH, and cell salvage. NOTE: This form only contains RCT info relevant for ANH.	Comparator: no Autologous transfusion technique (active versus active comparisons were excluded). Sample size n=591			
Sample size n=704				

Population characteristics

Patients older than 18 years undergoing any type of surgery. The mean age of participants in ANH trials was 56 years. The trials included more than twice as many males as females (2.3:1). Twelve trials involved cardiac surgery, seven involved orthopaedic surgery, and 11 involved various other operative procedures (eg, urological, thoracic, or vascular).

Length of follow-up	Outcomes measured				
NA	Mortality, re-operation, infection, wound complication, thrombosis, non-fatal MI, rate of allogeneic red blood cell transfusion, and volume of allogeneic blood transfused.				

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Allocation concealment and the method of randomisation were judged by the SR authors to be inadequate in 100 and 92% of trials respectively (kappa=0.78- 1.0).	SR did not discuss similarity between preoperative data and baseline characteristics for the intervention groups.	The majority (96%) of the included RCTs assessing ANH were unblended.	Not detected	NR

Overall quality assessment (descriptive)

Fair

Outcome	Intervention group	Comparator group	Statistical significance		
Mortality 8 trials (n=NR)	NR	NR	RR (95% CI): 1.16 (0.19, 7.15) Phet=NR RR (95% CI): 4.94 (0.61, 40.19) Phet=NR		
Morbidity: infection 2 trials (n=NR)	NR	NR			
Morbidity: thrombosis 3 trials (n=NR)	NR	NR	RR (95% CI): 0.44 (0.21, 0.93) Phet=NR		
Morbidity: non-fatal MI 3 trials (n=NR)	NR	NR	RR (95% CI): 3.43 (0.15, 79.74) Phet=NR		
Re-operation 7 trials (n=NR)	NR	NR	RR (95% CI):1.59 (0.20, 12.53) Phet=NR		
Rate of allogeneic blood transfusion: all studies 25 trials (n=1081; 567 ANH, 514 control)	273/567 (48%)	357/514 (69%)	RR (95% CI): (Phet<0.00001)		
Rate of allogeneic blood transfusion: cardiac surgery 10 trials (n=NR)	NR	NR	RR (95% CI): 0.77 (0.57, 1.04) RR (95% CI): 0.79 (0.60, 1.06)		
Rate of allogeneic blood transfusion: orthopaedic surgery 6 trials (n=NR)	NR	NR			
Rate of allogeneic blood transfusion: miscellaneous surgery 9 trials (n=NR)	NR	NR	RR (95% CI): 0.42 (0.24, 0.74)		
Rate of allogeneic blood transfusion: transfusion protocol used 16 trials (n=NR)	NR	NR	RR (95% CI): 0.81 (0.62, 1.00)		
Rate of allogeneic blood transfusion: transfusion protocol not used/reported 9 trials (n=NR)	NR	NR	RR (95% CI): 0.53 (0.36, 0.76)		
Difference in units of allogeneic blood transfused 17 trials	NR	NR	WMD (95% CI) Overall: -1.9 (-1.1, -2.7) Studies with a transfusion protocol: -1.0 (-1.7, -0.4) Studies without a transfusion protocol: -3.0 (-4.9, -1.1)		
Hospital length of stay, d 3 trials (N=96)	NR	NR	WMD (95% CI): 0.21 (-1.26, 1.68)		

Clinical importance	Clinical relevance				
EXTERNAL VALIDITY					
Generalisability					
Patients considered similar to guideline target population					
Applicability					
All the studies included in this review were conducted in countries with well developed healthcare systems (not specifically Aus/NZ).					
Comments					

Abbreviations: ANH, acute normovolemic haemodilution; MI, myocardial infarction; PAD, preoperative autologous donation; NR, not reported; RCT, randomised controlled trial; RR, risk ratio; WMD, weighted mean difference.

Citation								
Gurusamy KS, Li								e blood loss and blood
transfusion requir		er resection	. Cochrane Data	base of S	Systematic Re	views 2009;	lssue.	4
Affiliation/Source	e of funds							
None declared						Г		
Study design		Le	evel of evidence	9		Location/s	settino	9
SR of RCTs		1				NA		
Search conducted	Search conducted November 2008							
Intervention				Compa	arator			
Cardiopulmonary interventions to decrease blood loss and blood transfusion requirements for liver resection: low central venous pressure, PAD, ANH, ANH with controlled hypotension, and hypoventilation. NOTE: This form only contains info relevant for ANH. ¹ n=115								
Population chara	acteristics			l				
Patients undergoi normal or cirrhotic								or minor liver resections, resection.
Length of follow	-up			Outco	mes measure	d		
NR					erative mortalit ncy, operating			orbidity, transfusion od loss.
INTERNAL VALII	DITY							
Allocation	Results		Blinding an	alysis	Treatment/bias	measureme	nt	Follow-up (ITT)
All three trials had randomised allocation but unclear allocation concealment.	The authors stated that it unclear whe the three tria free of base imbalance.	was ther any of als were	None of the trials blinded outcomes.				Analysis performed with ITT.	
Overall quality as	ssessment (d	lescriptive))		1			1
Good								
RESULTS								
Outcome		Intervent	on group	Coi	mparator grou	ıp	Stat	istical significance
Perioperative mortality 2 trials (n=150; 73 ANH, 77 control)					3.32 P>0	(95% CI): 0.35 (0.04,) .05 (P <i>het</i> =1.00) significant difference		
Perioperative morbidity: bile leak 1 trial (n=78; 39 ANH, 39 control) RR (95% CI): 1.5 (0.27, 8.49) P>0.05 (Phet=NA))			

Limited: the SR only included p	apers assessing ANH for live	r resection, not otl	ner surgery types	5.
Generalisability				
EXTERNAL VALIDITY				
Outcome	Clinical importance		Clinical releva	ince
Length of hospital stay, days 1 trial (n=130; 63 ANH, 67 control)				Mean difference (95% CI): 0.0 (-2.66, 2.66) P>0.05 (Phet=NA) No significant difference
Operative blood loss, mL 2 trials (n=98; 49 ANH, 49 control)				Mean difference (95% CI): 1.53 (-102.37, 105.44) P>0.05 (Phet=0.83) ANH improves outcome
Operating time in minutes 2 trials (n=208; 102 ANH, 106 control)				Mean difference (95% CI): -28.86 (-57.37, -0.35) P<0.05 (Phet=0.90) ANH improves outcome
Red cell transfusion 2 trials (n=150; 73 ANH, 77 control)				Mean difference (95% CI): -0.09 (-0.48, 0.29) P>0.05 (Phet<0.00001) No significant difference
Number requiring allogeneic blood transfusion 3 trials (n=233; 115 ANH, 118 control)				RR (95% CI): 0.41 (0.25, 0.66) P<0.05 (P <i>het</i> =0.70) ANH improves outcome
Perioperative morbidity: chest infection 1 trial (n=78; 39 ANH, 39 control)				RR (95% CI): 1.50 (0.27, 8.49) P>0.05 (Phet=NA)
Perioperative morbidity: wound infection 2 trials (n=208; 102 ANH, 106 control)				RR (95% CI): 0.84 (0.34, 2.03) P>0.05 (Phet=0.18)
Perioperative morbidity: intra- abdominal collection requiring drainage 1 trial (n=130; 63 ANH, 67 control)				RR (95% CI): 1.26 (0.061, 2.60) P>0.05 (Phet=NA)
Perioperative morbidity: intra- abdominal infection 1 trial (n=78; 39 ANH, 39 control)				RR (95% CI): 0.33 (0.04, 3.07) P>0.05 (Phet=NA)
Perioperative morbidity: intra- abdominal bleeding 2 trials (n=208; 102 ANH, 106 control)				RR (95% CI): 1.87 (0.4, 8.67) P>0.05 (Phet=0.39)

Applicability

Of the three RCTs included in this SR, one was conducted in America, one in Israel, and the other was conducted in China. These countries have some differences to Aus/NZ health systems.

Comments

¹ The SR assesses both haemodilution versus control and haemodilution with bovine haemoglobin (HBOC-201) versus haemodilution with hydroxyl ethyl starch. This extraction form only includes data for haemodilution versus control.

Abbreviations: ANH; acute normovolemic haemodilution; M-H, Mantel-Haenszel estimate; NA, not applicable; NR, not reported; PAD, preoperative autologous donation; RCT, randomised controlled trial; RR, risk ratio; SR, systematic review.

Citation								
Laupacis A and F (Structured abstra				ologies to	minimise per	ioperative allog	gene	ic transfusion
Affiliation/Source	e of funds							
NR								
Study design	Study design Level of evidence Location/setting							J
SR		1				NA		
Search conducted March 1997								
Intervention Comparator								
Technologies to minimise perioperative allogeneic transfusion: aprotinin, desmopressin, tranexamic acid, epsilon aminocaproic acid, erythropoietin, PAD, ANH. NB: this form only reports results for ANH.								
Population chara	acteristics							
Adult patients und surgery.	dergoing elect	tive surgery	. Types of surge	ry include	d cardiac, col	orectal, liver re	esect	ion and orthopaedic
Length of follow	-up			Outcon	nes measure	d		
NR Proportion of patients receiving at least one unit of allogeneic packed red blood cells, perioperative MI, re-operations because of bleeding.								
INTERNAL VALI	DITY			II.				
Allocation	Results		Blinding anal	ysis	Treatment/ bias	measurement	t	Follow-up (ITT)
NR	Baseline characterist	tics NR	NR		Use of trans	sfusion protocol NR		NR
Overall quality a	ssessment (descriptive)					
Fair								
RESULTS								
Outcome		Interventi	on group	Con	nparator grou	ıp :	Stati	stical significance
allogeneic blood 0.62) "The likelihood in cardiac and miscellaneous p				likelihood was reduced rdiac and ellaneous procedures ot in orthopaedic				
Patients transfused with allogeneic blood: <1000 mL blood withdrawn OR (95% CI): 0.43 (0.18, 1.02)								
Patients transfused with allogeneic blood: >1000 mL blood withdrawn OR (95% CI): 0.16 (0.04, 0.65)								

Outcome	Clinical importance	Clinical relevance				
Patients transfused with allogeneic blood	1	1				
Patients transfused with allogeneic blood: <1000 mL blood withdrawn	4	1				
Patients transfused with allogeneic blood: >1000 mL blood withdrawn	1	1				
EXTERNAL VALIDITY						
Generalisability						
The SR is generalisable for ele	The SR is generalisable for elective, non urgent surgery.					
Applicability						
The studies were mostly from countries with similar health-care systems to Australia						
Comments	Comments					

Abbreviations: ANH, acute normovolemic haemodilution; MI, myocardial infarction; NA, not applicable; NR, not reported; OR, odds ratio; PAD, preoperative autologous deposit; SR, systematic review.

Citation						
Segal JB, Blasco- analysis. Transfus	Colmenares E, Norris ion 44:632-644.	EJ, and Guallar E.	(2004) Pr	eoperative ac	ute normovolemic	hemodilution: A meta-
Affiliation/Source	e of funds					
None declared						
Study design		Level of evidence	9		Location/setting	9
SR		1			NA	
Search conducted	October 2002					
Intervention			Compa	rator		
ANH			Compai	rison group tha	at did not receive A	ANH.
Population chara	ecteristics		I			
Patients undergoi	ng any surgery type					
Length of follow-	·up		Outcon	nes measure	d	
NR I			Number of subjects who received allogeneic blood transfusion in the perioperative period, average amount of allogeneic blood received per patient in each study group, average volume of blood loss perioperatively, adverse events.			
INTERNAL VALID	DITY		l		·	
Allocation	Results	Blinding analy	ysis	Treatment/r bias	measurement	Follow-up (ITT)
Of the 42 included studies, only 12 studies reported that the outcomes assessor was masked as to treatment assignment. Although all studies were randomised only 14 of the studies provided any description of the randomisation procedure.	SR did not report baseline characteristics	The patients w masked to trea assignment in four studies.	ıtment	the studies r threshold at		NR
Overall quality as	ssessment (descripti	ve)				
Fair						

Outcome	Intervention group	Comparator group	Statistical significance
Proportion receiving allogeneic blood perioperatively (ANH vs standard care) 25 trials (n=1409; 703 ANH, 706 standard care)	mior volution group	oomparator group	RR (95% CI): 0.96 (0.90, 1.01) P>0.05 (Phet=0.98) No significant difference
Proportion receiving allogeneic blood perioperatively (trials with concurrent use of other blood conservation methods in both arms) 10 trials (n=NR)			RR (95% CI): 0.98 (0.91, 1.07) P>0.05 (Phet=NA) No significant difference
Volume of allogeneic blood transfused intraoperatively (ANH vs standard care), mL ¹ Number of trials not reported			WMD (95% CI): -303 (-555, - 55) p<0.05 (P <i>het</i> <0.001) ANH improves outcome
Volume of allogeneic blood transfused intra- and postoperatively (ANH vs standard care), mL ² 13 trials (n=735; 406 ANH, 329 standard care)			WMD (95% CI): -201 (-309, - 92) P<0.05 (Phet<0.001) ANH improves outcome
Volume of intraoperative blood loss (ANH vs standard care), mL Number of trials not reported			Pooled average difference (95% CI): 15 (-27, 58) P>0.05 (Phet=0.26) No significant difference
Volume of intra- and postoperative blood loss (ANH vs standard care), mL 20 trials (n=1138; 565 ANH, 573 standard care)			WMD (95% CI): -91 (-157, - 25) P<0.05 (Phet<0.0001)
Mortality 17 trials (n=1191)	6/607 (1%)	10/584 (2%)	NR
Morbidity	Myocardial infarction: 7/502 (1%) Cardiac ischemia: 8/140 (6%) Left ventricular dysfunction: 2/133 (2%) Venous thromboembolism: 3/180 (2%) Cerebral infarction: 3/323 (1%) Hypotension during haemodilution: 2/234 (1%) Transfusion reaction: 0/131	Myocardial infarction: 9/480 (2%) Cardiac ischemia: 9/137 (7%) Left ventricular dysfunction: 7/110 (6%) Venous thromboembolism: 2/180 (1%) Cerebral infarction: 2/343 (1%) Hypotension during haemodilution: 0/243 (0%) Transfusion reaction: 0/153	NR

Outcome	Clinical importance	Clinical relevance					
Outcome	Clinical importance	Clinical relevance					
EXTERNAL VALIDITY	EXTERNAL VALIDITY						
Generalisability							
Patients considered similar to	guideline target population						
Applicability							
Study performed in UK which has many similarities with the Aus/NZ healthcare systems.							
Comments							
NB: in this review units of blood were transformed to millilitres by assuming that 1 unit of whole blood contained 450 mL of blood and that 1 unit of RBCs had a volume of 300 mL.							

¹The one trial that was an extreme outlier in the volume of blood transfused (Lilleaasen 1977) was removed from these analyses. The two studies that used the largest volume haemodilution (mean of 1500 mL of blood withdrawn) reduced the volume of intraoperative allogeneic transfusion most extremely (weighted mean difference, -720 mL; 95% CI: -475, -982).

²Trials using higher Hct levels to trigger transfusion demonstrated greater savings with ANH (P<0.001), as did older trials (P=0.04). There is also little difference between the large-volume and lower-volume studies (P=0.6).

Abbreviations: ANH, acute normovolemic haemodilution; NA, not applicable; NR, not reported; RR, risk ratio; SR, systematic review; WMD,

Abbreviations: ANH, acute normovolemic haemodilution; NA, not applicable; NR, not reported; RR, risk ratio; SR, systematic review; WMD, weighted mean difference.

Level II evidence

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C	It 2	١tı	n	n
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Akhlagh SH, Chohedri AH, Bazojoo A, and Nemati MH. (2007) A comparison of total amount of blood needed in patients taking autologous or homologous blood transfusion in coronary artery bypass grafting: A clinical randomized case-control trial. Pakistan Journal of Medical Sciences 23:542-545.

Affiliation/Source of funds

None declared

Study design	Level of evidence	Location/setting
RCT	II	Iran

Intervention	Comparator
ANH and retransfusion of autologous blood after separating the patient from the cardiopulmonary machine.	Control: no withdrawal of blood and only allogeneic blood transfused.
n=30? ¹	n=30? ¹

Population characteristics

Patients undergoing coronary artery bypass grafting (on-pump). Eighty percent of patients were male.

Length of follow-up	Outcomes measured	
24 hours		

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
It is unclear whether allocation was concealed from those responsible for recruiting subjects.	The baseline characteristics of the treatment arms were matched.	All data were registered by an independent investigator	No transfusion protocol was reported.	There was no reported loss to follow-up

Overall quality assessment (descriptive)

Poor

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance
Mean (SD) 24 h post- operational haematocrit concentration, %	36.5 (1.5)	37 (2)	P=0.21
Mean total volume of blood transfused, mL	1815 ²	2010	P=0.024
Clinical importance		Clinical relevance	

EXTERNAL VALIDITY

Generalisability

The study population is consistent with the guideline population.

Applicability

The study was conducted in Iran, which may limit its applicability to the Australian context.

Comments

¹Although the paper reports the total study population (n=60), it does not specify the number of patients randomised to each arm. The values of 30 patients for each arm is based on the assumption that the participants were evenly split between active and control treatments. ²Of the 1815 mL blood needed in the ANH group, 870 mL (47.9%) was homologous and 945 mL (52.1%) was autologous.

Citation						
Bennett J, Haynes	s S, Torella F, Grainger I				volemic hemodilut	ion in moderate blood
	ndomized controlled trial	. Transfusion 46:	1097-110	3.		
Affiliation/Sourc	e of funds					
NR						
Study design	L	evel of evidence	Э		Location/setting	9
RCT	ll l				UK hospital	
Intervention			Compa	rator		
surgery, aiming to target of 110 g pe mL bags containin crystalloids were in normovolaemia. A 6 hours of collecti if a transfusion trig	blood was collected immoreduce haemoglobin corr. ANH blood was colled and 63 mL of citrate-based infused simultaneously to all autologous blood was on, starting on wound clapger was reached. If furthogeneic blood was administration	encentration to a ected into 450 d anticoagulant; o maintain returned within osure or sooner her transfusion	Allogen reached	eic blood was d. The trigger f	or both intervention	nsfusion trigger was in and comparator vel of less than 80 g per
Population chara	acteristics					
hip replacement,	· · · · · · · · · · · · · · · · · · ·		IH and 8 i	n standard tra	nsfusion) and 1 hi	nderwent primary total p resurfacing procedure.
Length of follow	· · · · · · · · · · · · · · · · · · ·			nes measured		
Unclear (at least (until hospital discharge)		Proportion of patients requiring allogeneic transfusion and the volume of allogeneic blood transfused, postoperative complications (infective complications and hospital inpatient stay)			
INTERNAL VALI	DITY		•			
Allocation	Results	Blinding anal	ysis	Treatment/r bias	neasurement	Follow-up (ITT)
Randomised allocation. Unclear whether allocation to treatment groups was concealed from those responsible for recruiting subjects.	Twenty patients in each group were on anti-PLT, or anticoagulant drugs at the time of surgery; 29 in standard transfusion and 29 in ANH were on NSAIDs. There were no differences in preoperative Hb or PLT count.	Neither the anaesthetist no surgical team of be blinded but patients were the allocated treatment.	could	transfusion a received allo violation of the (2 units each to reach any	ese 5 patients cluded, the transfused units vs 25 have been	All analysis follows ITT.
Overall quality a	ssessment (descriptive	e)				
Fair						
RESULTS						

Comparator group

Intervention group

Outcome

Statistical significance

Mortality	1/78 (1.3%)	0/77 (0%)	P=0.50
Patients with at least one significant postoperative complication.	14/78 (18%)	30/77 (38%)	P=0.006
Morbidity	Cardiovascular event: 1/78 (1%) Postoperative infection: 7/78 (9%) Wound (non-infective): 2/78 (3%) Bleeding: 0/78 (0%) Venous thromboembolism: 2/78 (3%) Urinary retention: 3/78 (4%) Transfusion reaction: 0/78 (0%)	Cardiovascular event: 4/77 (5%) Postoperative infection: 17/77 (22%) Wound (non-infective): 0/77 (0%) Bleeding: 1/77 (1%) Venous thromboembolism: 1/77 (1%) Urinary retention: 3/77 (4%) Transfusion reaction: 1/77 (1%)	P-value Cardiovascular event: 0.21 Postoperative infection: 0.03 Wound (non-infective): 0.30 Bleeding: 0.49 Venous thromboembolism: 0.58 Urinary retention: 0.99 Transfusion reaction: 0.49
Patients requiring allogeneic transfusion	15 /78(19%)	22/77 (29%)	P=0.18
Units of allogeneic blood transfused	33	63	P=0.1
Median (IQR) postoperative hospital stay	7 (6, 9)	8 (6, 11)	P=0.03
Median (IQR) intraoperative blood loss (mL)	692 (452, 1019)	641 (477, 1007)	P=0.82
Median (IQR) total blood loss (mL)	1182 (840, 1646)	1210 (816, 1545)	P=0.82
Patients who needed to seek medical attention after discharge (either via their GP or local hospital)	29/78 (37%)	43/77 (56%)	P=0.02
EXTERNAL VALIDITY			
Generalisability			
Patients considered similar to	guideline target population		
Applicability			
Study performed in UK which	has many similarities with the Au	us/NZ healthcare systems.	
Comments			

Abbreviations: ANH, acute normovolemic haemodilution; Hb, haemoglobin; IQR, inter-quartile range; NR, not reported; NSAID, non-steroidal anti-inflammatory drug; OR, odds ratio; PLT, platelet; RCT, randomised controlled trial.

Casati V, Speziali G, D'Alessandro C, Cianchi C, Antonietta Grasso M, Spagnolo S, and Sandrelli L. (2002) Intraoperative low-volume acute normovolemic hemodilution in adult open-heart surgery. Anesthesiology 97:367-373.

Affiliation/Source of funds

NR

Study design	Level of evidence	Location/setting
RCT	II	Italy, hospital

Intervention	Comparator	
Low volume ANH: 5-8 mL/kg of blood w systemic heparinisation and replaced wi solutions. n=103	Standard care: no had n=101	emodilution

Population characteristics

Patients undergoing on-CPB cardiac surgery.

Preoperative exclusion criteria: age < 18 years, LVEF < 30%, preoperative haematocrit < 36% or haemoglobin less than 12 g/dl, history of haematologic diseases, chronic renal insufficiency (plasma creatinine > 2 mg/dl), and history of hepatic diseases.

Length of follow-up	Outcomes measured
Samples for evaluation of haemoglobin, haematocrit, platelet count, prothrombin time, activated partial thromboplastin time, creatinine, creatine phosphokinase, and creatine phosphokinase myocardial band isoenzyme were performed before the induction of anaesthesia (time 1), on arrival in ICU (time 2), 24 h after the arrival in ICU (time 3), 48 h after surgery (time 4), and at discharge (time 5). Blood loss was recorded during the first 24 h.	Transfusion frequency, amount of allogeneic blood transfused, amount of postoperative bleeding, postoperative complications, and postoperative haematochemical data.

INTERNAL VALIDITY

	T		T .	
Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Allocation was randomised using a computer-generated random number sequence. It is unclear whether allocation to treatment groups was concealed from those responsible for recruiting subjects.	Treatment groups had similar patient demographics, haematochemical levels. The intervention/control split for the entire set of cardiac operations assessed was close to even. Only the amounts of colloids and crystalloids infused before CPB, greater in ANH group, were significantly different.	Neither the patients nor the surgical staff/assessors were blinded to treatment allocation.	Not detected.	Study used per protocol analysis

589

(descriptive)		
Intervention group	Comparator group	Statistical significance ¹
4/103 (3.9%)	4/101 (4%)	P=0.98
Myocardial infarction: 2/103 (2%) renal failure: 3/103 (2.9%) minor neurological complications: 7/103 (6.9%) stroke: 2/103 (2%) pulmonary embolism: 0/103 (0%)	Myocardial infarction: 1/101 (1%) renal failure: 4/101 (4%) minor neurological complications: 8/101 (8%) stroke: 1/101 (1%) pulmonary embolism: 1/101 (1%)	P-value Myocardial infarction: P=0.58 renal failure: 0.68 minor neurological complications: 0.86 stroke: 0.58 pulmonary embolism: 0.49
35/103 (34%)	36/101 (36%)	P=0.88
123/32	126/34	P=0.47
158 (106, 305)	172 (117.5, 265)	P=0.93
374 (255, 704)	412 (313, 552)	P=0.94
7 (6, 9)	7 (6, 8.25)	P=0.54
1 (1, 1)	1 (1, 2)	P=0.49
	Intervention group 4/103 (3.9%) Myocardial infarction: 2/103 (2%) renal failure: 3/103 (2.9%) minor neurological complications: 7/103 (6.9%) stroke: 2/103 (2%) pulmonary embolism: 0/103 (0%) 35/103 (34%) 123/32 158 (106, 305) 374 (255, 704) 7 (6, 9)	Intervention group 4/103 (3.9%) Myocardial infarction: 2/103 (2%) renal failure: 3/103 (2.9%) minor neurological complications: 7/103 (6.9%) stroke: 2/103 (2%) pulmonary embolism: 0/103 (0%) 35/103 (34%) 123/32 126/34 158 (106, 305) 7 (6, 9) Comparator group 4/101 (4%) Myocardial infarction: 1/101 (1%) renal failure: 4/101 (4%) minor neurological complications: 8/101 (8%) stroke: 1/101 (1%) pulmonary embolism: 1/101 (1%) 126/34 172 (117.5, 265) 412 (313, 552) 7 (6, 9) 7 (6, 8.25)

EXTERNAL VALIDITY

Generalisability

Patients considered similar to guideline target population.

Applicability

RCT performed in Italy; however the results should be applicable to the Australian setting.

Comments

Two patients (1 per group) did not complete the study: they died during the first 24 h postoperatively after cardiogenic shock refractory to maximal pharmacologic support and intra-aortic counter pulsation. They were excluded by statistical analysis. 202 patients entered in the statistical analysis.

Abbreviations: FFP, fresh frozen plasma; ICU, intensive care unit; IQR, interquartile range; LVEF, left ventricular ejection fraction; NR, not reported; PLTC, platelet concentrate; PRBC, packed red blood cells; RCT, randomised controlled trial.

¹The authors of the study conducted all statistical tests with per protocol analysis.

²Not including the two patients who died 24 h postoperatively.

Casati V, Benussi S, Sandrelli L, Grasso MA, Spagnolo S, and D'Angelo A. (2004) Intraoperative Moderate Acute Norvolemic Hemodilution Associated with a Comprehensive Blood-Sparing Protocol in Off-Pump Coronary Surgery. Anesthesia and Analgesia 98:1217-1223.

Affiliation/Source of funds

NR

Study design	Level of evidence	Location/setting
RCT	II	Italy / Hospital

Intervention Comparator

ANH + tranexamic acid: intraoperative tranexamic acid as an IV bolus of 1 g 20 min before sternotomy, followed by continuous infusion of 400 mg/h until the end of surgery. The blood shed from the surgical field was collected in a cardiotomy reservoir and, in case of intraoperative bleeding more than 250 mL, reinfused after washing and concentration in a cell salvage circuit.

n=50

ANH protocol: The whole-blood volume targeted for removal was 17% \pm 2% of the circulating volume, as calculated from a body-surface area nomogram. The blood was drawn after the induction of anesthesia and before systemic heparinization, and was collected into sterile bags containing citrate phosphate dextrose by using a blood mixer and balance system. During blood withdrawl, 4% succinylated gelatine in 0.9% NaCl was infused at a 1:1 ratio. Irrespective of heamatocrit values, reinfusion of the harvested Autologous blood was started after protamine administration and on-demand reinfusion of the shed blood.

Tranexamic acid control: Intraoperative tranexamic acid as an IV bolus of 1 g 20 min before sternotomy, followed by continuous infusion of 400 mg/h until the end of surgery. The blood shed from the surgical field was collected in a cardiotomy reservoir and, in case of intraoperative bleeding more than 250 mL, reinfused after washing and concentration in a cell salvage circuit.

n=50

Population characteristics

Patients undergoing OPCAB (baseline hematocrit > 34%).

Length of follow-up	Outcomes measured
NR	Mortality, perioperative complications (respiratory failure, myocardial infarction, acute renal failure, venous thromboembolism, neurological complications), postoperative bleeding, transfusion frequency of allogeneic blood, units of allogeneic blood transfused, ICU stay, postoperative hospital stay, intubation time.

INTERNAL VALIDITY				
Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Allocation randomised, however it is unclear whether allocation to treatment groups was concealed from those responsible for recruiting subjects.	Treatment groups had similar patient demographics and haematochemical levels.	Surgical staff/assessors were not blinded to treatment group. It is unclear whether the participants were blinded to treatment allocation.	Not detected	There was no loss to follow-up. All analyses were performed ITT

Overall quality assessment (descriptive)

Fair

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance
Mortality	0/50 (0%)	1/50 (2%)	-
Morbidity	Respiratory failure: 1/50 (2%) atrial fibrillation: 5/50 (10%) major ventricular arrhythmia: 1/50 (2%) myocardial infarction: 1/50 (2%) creatinine double the baseline: 1/50 (2%) minor neurological complications: 2/50 (4%)	Respiratory failure: 1/50 (2%) atrial fibrillation: 6/50 (12%) major ventricular arrhythmia: 1/50 (2%) myocardial infarction: 1/50 (2%) creatinine double the baseline: 2/50 (4%) minor neurological complications: 1/50 (2%)	P-value Respiratory failure: 1.00 atrial fibrillation: 0.75 major ventricular arrhythmia: 1.00 myocardial infarction: 1.00 creatinine double the baseline: 0.57 minor neurological complications: 0.57
Mean (IQR) total postoperative bleeding (mL)	375 (248, 475)	350 (300, 443)	NS
Mean (IQR) bleeding 0 to 4 h after surgery (mL)	160 (110, 235)	150 (100, 220)	NS
Patients transfused with allogeneic PRBC	2/50 (4%)	10/50 (20%)	P=0.028
Units of allogeneic PRBC transfused	5	24	P<0.001
Total number of patients transfused with allogeneic blood (including PRBC, FFP, and PLTC)	2/50 (4%)	10/50 (20%)	P=0.028
Mean (IQR) ICU stay (d)	1 (1, 1)	1 (1, 1)	P=1
Mean (IQR) postoperative hospital stay (d)	6 (6, 7)	6 (6, 7)	NR
Mean (IQR) intubation time (min)	252 (151, 186)	244 (165, 182)	NR

592

EXTERNAL VALIDITY
Generalisability
Patients considered similar to guideline target population.
Applicability
RCT performed in Italy; however the results should be applicable to the Australian setting.
Comments

Abbreviations: OPCAB, Off-pump coronary artery bypass

Friesen RH, Perryman KM, Weigers KR, Mitchell MB, and Friesen RM. (2006) A trial of fresh autologous whole blood to treat dilutional coagulopathy following cardiopulmonary bypass in infants. Paediatric Anaesthesia 16:429-435.

Affiliation/Source of funds

The research was supported by a grant from the General Clinical Research Centers Program, National Center for Research Resources, NIH.

Study design	Level of evidence	Location/setting
RCT	II	USA / hospital

Intervention	Comparator
Whole blood in the amount of 15 mL/kg was withdrawn from the patient through the central venous catheter. Isovolemia was maintained by infusion of 1 mL of 5% albumin solution for each mL of blood withdrawn. The autologous blood was retransfused postoperatively. n=16	Patients in the control group did not have withdrawal of autologous whole blood or infusion of albumin. n=16

Population characteristics

Infants undergoing non-complex open cardiac surgery.

Length of follow-up	Outcomes measured
After 2 hours in the ICU.	Primary outcome: Coagulation status (measured by PC, PA, PT, aPTT, and fibrinogen concentration.
	Secondary outcomes: activation of fibrinolysis (not reported in this form) ¹ , measured by tissue plasminogen activator, plasminogen activator inhibitor, and d-dimer; haematocrit; 24 h postoperative blood loss, measured as the sanguinous output through the mediastinal drainage tube; and the transfusion of homologous blood components during the intraoperative and 24 h postoperative periods.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Allocation was concealed from those responsible for recruiting subjects.	The intervention groups were similar in patient demographics, type of operation, and baseline coagulation tests.	The study was not blinded.	Transfusion protocol was used.	There was no loss to follow-up

Overall quality assessment (descriptive)

Fair

Outcome	Intervention group	Comparator group	Statistical significance
Mean (SD) haematocrit, %	T1: 32 (3) T2: 32 (8) T3: 33 (7)	T1: 32 (4) T2: 34 (6) T3: 34 (6)	From T2 to T4, the treatment group had a greater improvement in haematocrit (P=0.009).
	T4:35 (8) ΔT2 – T3: +1 (2) ΔT2 – T4: +3 (4)	T4: 34 (5) ΔT2 – T3: +1 (1) ΔT2 – T4: 0 (3)	,
Mean (SD) PC, 10 ⁹ /L	T1: 353 (92) T2: 126 (49) T3: 161 (55) T4:207 (53) ΔT2 – T3: +36 (22) ΔT2 – T4: +82 (43)	T1: 335 (92) T2: 140 (47) T3: 158 (57) T4: 217 (59) ΔT2 – T3: +18 (17) ΔT2 – T4: +70 (42)	From T2 to T3, the treatment group had greater improvement in PC (P=0.018)
Mean (SD) PA, s	T1: 205 (62) T2: 222 (71) T3: 144 (58) T4: 112 (23) ΔT2 – T3: -78 (53) ΔT2 – T4: -109 (67)	T1: 189 (54) T2: 210 (70) T3: 159 (72) T4: 113 (32) ΔT2 – T3: -49 (77) ΔT2 – T4: -97 (64)	NR
Mean (SD) PT, s	T1: 13.4 (0.9) T2: 20.4 (4.3) T3: 18.1 (3.1) T4: 15.9 (2.1) ΔT2 – T3: -2.3 (1.9) ΔT2 – T4: -4.5 (3.2)	T1: 14.1 (1.1) T2: 19.9 (3.8) T3: 18.9 (3.6) T4:16.8 (2.0) ΔT2 – T3: -0.9 (1.2) ΔT2 – T4: -3.0 (2.7)	From T2 to T3, the treatment group had greater improvement in PT (P=0.015)
Mean (SD) aPTT, s	T1: 35.9 (9.3) T2: 46.7 (14.2) T3: 42.2 (14.1) T4: 37.8 (13.2) ΔT2 – T3: -4.4 (7.7) ΔT2 – T4: -8.9 (11.0)	T1: 36.9 (8.7) T2: 44.1 (12.6) T3: 43.7 (13.1) T4: 41.9 (17.2) ΔT2 – T3: -0.4 (9.6) ΔT2 – T4: -2.3 (16.7)	NR
Mean (SD) fibrinogen concentration, mg/dL	T1: 235 (63) T2: 109 (37) T3: 132 (44) T4: 152 (51) ΔT2 – T3: +14 (9) ΔT2 – T4: +35 (18)	T1: 215 (55) T2: 129 (38) T3: 128 (32) T4: 146 (36) ΔT2 – T3: -1 (16) ΔT2 – T4: +17 (20)	The treatment group had a greater improvement in fibrinogen from T2 to T3 (P=0.003), and T2 to T4 (P=0.019)
Postoperative 24 h blood loss.			Was less in the treatment group when measured as mL per 24 h (P=0.036), but not significantly less when measured as mL/kg.24h (P=0.16).

Transfusion misc.		One subject in the treatment group and five subjects in the control group received postoperative FFP or platelet transfusion (P=0.06).
Clinical importance	Clinical relevance	
•		
EXTERNAL VALIDITY		
Generalisability		
The study population was infants, and therefore the study is n	ot generalisable to an adult popu	lation.
Applicability		
The study was conducted in the USA, and the procedures are	likely to be comparable to those	used in Australia.
Comments		

Abbreviations: aPTT, activated partial thromboplastin time; CPB, cardiopulmonary bypass; ICU, intensive-care unit; PA, platelet aggregation; PC, platelet count; PT, prothrombin time; T1, baseline; T2, following conclusion of CPB and modified ultrafiltration; T3, 20 min after T2; T4, after 2 hours in the ICU

¹Tests of fibrinolysis underwent similar changes in both groups.

Hohn L, Schweizer A, Licker M, and Morel DR. (2002) Absence of beneficial effect of acute normovolemic hemodilution combined with aprotinin on allogeneic blood transfusion requirements in cardiac surgery. Anesthesiology 96:276-282.

Affiliation/Source of funds

The current study was initiated while the authors were in discussion with Biosafe Corporation about the opportunity to test a new blood separator device to evaluate the efficiency of thrombocytapheresis during cardiac surgery.

Study design	Level of evidence	Location/setting
RCT	II	Switzerland / university hospital

Intervention	Comparator
ANH: ANH was added to the control procedure described. Blood was withdrawn from the mean haematocrit of 43% to 28%. n=39 (although 3 participants were lost to follow up and not included in analysis)	Control: filling of extracorporeal circuit with saline isotonic fluid only, intraoperative blood salvage and reinfusion of shed mediastinal blood, integral reinfusion of blood contained in the extracorporeal circuit at the end of surgery, administration of intravenous aprotinin, and external heating at the end of CPD. n=41

Population characteristics

Patients undergoing on-CPB cardiac surgery

Length of follow-up	Outcomes measured
Five days after hospital discharge.	

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
The study was randomised however it is not clear whether allocation was concealed from those responsible for recruiting subjects.	The treatment groups had similar preoperative demographics, except that more patients were taking diuretics in the ANH group than in the control group (P=0.001).	The study was not blinded	A transfusion protocol was used.	Three ANH patients (8%) were withdrawn because of postoperative surgical lesion bleeding that needed reoperation for haemostasis (P=0.098 compared with the control group). These patients were not included in further analysis.

Overall quality assessment (descriptive)

Poor

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance
Mean ± SD (range) blood volume collected during ANH, mL	1099 ± 333 (430 to 1900)		
Patients receiving allogeneic blood	12/39 (31%)	12/41 (29%)	P=0.88

Median (range) number of allogeneic blood units per transfused patient	2 (1 to 5)	2 (1 to 3)	P=0.219
Mean (SD) baseline haematocrit concentration, %	43.3 (3.9)	43.2 (2.4)	P=0.89
Mean (SD) immediate postoperative haematocrit concentration, %	25 (3.5)	25.7 (3.3)	P=0.36
In-hospital mortality	0/39 (0%)	2/41 (5%)	P=0.31
Mean (SD) postoperative length of hospital stay, d	13.1 (3.7)	13.4 (8.3)	P=0.83
Mean (SD) length of ICU stay, d	3.1 (1.3)	3.0 (1.3)	P=0.73
Reoperation for bleeding	3/39 (8%)	0/41 (0%)	P=0.18
Mean (SD) surgical time, min	245 (65)	271 (80)	P=0.11
Clinical importance		Clinical relevance	·
EXTERNAL VALIDITY			
Generalisability			
Patients studied were similar to the target population of the guidelines.			
Applicability			
Study was conducted in Switzerland, but is likely to be applicable to the Australian context.			

Comments

Jarnagin WR, Gonen M, Maithel SK, Fong Y, DAngelica MI, Dematteo RP, Grant F, Wuest D, Kundu K, Blumgart LH, and Fischer M. (2008) A prospective randomized trial of acute normovolemic hemodilution compared to standard intraoperative management in patients undergoing major hepatic resection. Annals of Surgery 248:360-368.

Affiliation/Source of funds

Authors received financial support from Robert Wittes, MD, Physician-in-Chief, Memorial Hospital

Study design	Level of evidence	Location/setting
RCT	II	USA / hospital

Intervention	Comparator
ANH: blood was withdrawn to a target Hgb of 8.0 g/dL, with a maximum of 3 L of blood removed. Euvolemia was maintained by replacing half of the removed blood volume with 5% albumin and the other half with crystalloid. n=63	Standard anaesthetic management n=67

Population characteristics

Patients undergoing major hepatic resection (3 or more liver segments) for any diagnosis, with or without any other planned procedures.

Length of follow-up	Outcomes measured	
-	Proportion of patients who required transfusion of allogeneic red blood cell products, impact of ANH on FFP transfusion, intraoperative management, postoperative complications, operating time, and length of stay.	

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
It is not clear whether allocation was concealed from those responsible for recruiting subjects.	The groups were well matched for demographic and preoperative variables. The only difference noted was the proportion of patients with comorbid medical conditions, which was higher in the ANH group.	The study was not blinded.	A transfusion protocol was used.	ITT analysis was performed.

Overall quality assessment (descriptive)

Fair

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance
Median (range) volume of blood removed during ANH, mL	2250 (800 to 3000)		

			T =	
Median (range) estimated blood loss, mL	800 (100 to 3200)	700 (100 to 4000)	P=0.42	
Median (range) length of surgery, min	255 (135 to 546)	288 (140 to 535)	P=0.35	
Patients undergoing any allogeneic transfusion	14/63 (22.2%)	23/67 (34%)	P=0.13	
Patients transfused with allogeneic RBC (total)	8/63 (12.7%)	17/67 (25.4%)	P=0.08	
Patients transfused with allogeneic RBC (intraoperative)	1/63 (1.6%)	7/67 (10.4%)	P=0.07	
Patients transfused with FFP	11/63 (17.5%)	19/67 (28.4%)	P=0.15	
Mean (SD) units of allogeneic RBC transfused (for those who received any allogeneic RBC transfusion)	3.5 (1.3)	2.1 (0.5)	P=0.6	
Mean (SD) units of any allogeneic transfusion (RBC or FFP; for those who received any RBC or FFP transfusion)	5.6 (1.7)	6.9 (2.7)	P=0.72	
Overall morbidity	28/63 (44%)	22/67 (33%)	P=0.17	
Grade ≥ 3 morbidity	19/63 (30%)	19/67 (28%)	P=0.82	
Median (range) length of hospital stay, d	7 (5 to 50)	7 (4 to 26)	P=0.33	
Clinical importance		Clinical relevance	Clinical relevance	
EXTERNAL VALIDITY				
Generalisability				
Patients studied were similar to the target population of the guidelines.				
Applicability				
The study was conducted in USA, but is likely to be applicable to the Australian context.				
Comments				

Abbreviations: FFP, fresh frozen plasma

Juelsgaard P, Moller MB, and Larsen UT. (2002) Preoperative acute normovolaemic hemodilution (ANH) in combination with hypotensive epidural anaesthesia (HEA) during knee arthroplasty surgery. No effect on transfusion rate. A randomized controlled trial [ISRCTN87597684]. BMC Anesthesiology 2.

Affiliation/Source of funds

None declared

Study design	Level of evidence	Location/setting
RCT	II	Denmark, hospital

Intervention	Comparator
ANH: 20% of the total blood volume was drawn before anaesthesia. This volume was simultaneously replaced with an equal volume of Hydroxy Ethyl Starch 6%. Blood re-transfusion was completed within 6 h. n=14	No ANH. n=14

Population characteristics

Patients undergoing total knee arthroplasty surgery.

Length of follow-up	Outcomes measured
	Amount of allogeneic blood transfused, blood loss

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Patients were randomised, however it is unclear whether the allocation was blinded from those responsible for recruiting subjects.	The participants in the ANH group were older than control: 75.8 (6.4) years vs 70.1 (9.6) years; P=0.008 The control group had a higher mean arterial pressure (mm Hg) compared with ANH: 118 (19) vs 103 (12); P=0.02	The patients were blinded to treatment allocation; however, it is unclear whether the allocation was blinded to the people conducting the trial.	Not detected	There was no loss to follow up.

Overall quality assessment (descriptive)

Fair

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance
Patients transfused with PRBCs	7/14 (50%)	6/14 (43%)	
Mean volume of PRBCs transfused (mL)	386	343	0.85
Mean (SD) intraoperative blood loss (mL)	131 (78)	111 (56)	0.45

601

Mean (SD) total blood loss (mL)	1306 (300)	1026 (294)	0.02		
Clinical importance		Clinical relevance			
EXTERNAL VALIDITY	EXTERNAL VALIDITY				
Generalisability					
Patients studied were similar to the target population of the guidelines.					
Applicability					
The study was conducted in Denmark; however it should be applicable to the Australian context.					
Comments					

Lim YJ, Kim CS, Bahk JH, Ham BM, and Do SH. (2003) Clinical trial of esmolol-induced controlled hypotension with or without acute normovolemic hemodilution in spinal surgery. Acta Anaesthesiologica Scandinavica 47:74-78.

Affiliation/Source of funds

None declared

Intervention

Study design	Level of evidence	Location/setting
RCT	II	South Korea / university hospital

Combined ANH and esmolol-induced controlled hypotension (E-ANH group): for ANH autologous blood was withdrawn from the radial artery aiming for 28% haematocrit. To maintain normovolemia, the first 500 mL of blood drawn was simultaneously replaced with an

of blood drawn was simultaneously replaced with an equal amount of 6% hydroxyethyl starch, and the blood thereafter was replaced with three times that volume of Lactated Ringer's solution.

n=15

Comparator

esmolol-induced controlled hypotention alone (esmolol group): controlled hypotension was performed during the period of bony decompression and instrumental fusion of the spine. Following an initial bolus injection of esmolol 500 μ /kg, esmolol was continuously infused at a rate of 0-300 μ g/kg/min to bring the MAP n=15

Population characteristics

Patients undergoing spinal surgery

Length of follow-up	Outcomes measured
1 week	

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
It was not clear whether allocation was concealed from those responsible for recruiting subjects.	There were no significant differences in demographic data between the two groups.	The study was not blinded.	A transfusion protocol was used.	There was no loss to follow-up.

Overall quality assessment (descriptive)

Fair

RESULTS

Outcome	Intervention group (E-ANH)	Comparator group (esmolol)	Statistical significance
Mean (SE) volume withdrawn during ANH, mL ¹	717 (50)	-	-
Mean (SE) volume of intraoperative bleeding, mL	1600 (160)	1500 (180)	P>0.05
Mean (SE) volume of postoperative bleeding, mL	600 (96)	883 (122)	P>0.05

Mean (SE) concentration of haemoglobin one week postoperative, g%	11.3 (0.3)	11.3 (0.2)	P>0.05		
Patients transfused with PRBCs	10/15 (67%)	15/15 (100%)	P=0.04		
Mean (SE) units of PRBCs transfused	2.2 (0.6)	4.3 (0.4)	P=0.0052		
Morbidity			All patients were evaluated 1 week after the operation, and there were no postoperative complications (thromboembolism, neurologic sequelae or wound infection) in either group.		
Clinical importance		Clinical relevance			
EXTERNAL VALIDITY					
Generalisability	Generalisability				
Patients studied were similar to the target population of the guidelines.					
Applicability					
The study was conducted in South Korea; however it should be applicable to the Australian context.					
Comments					
_					

¹All autologous blood was returned to the patients postoperatively. Abbreviations: MAP, mean arterial pressure

Matot I, Scheinin O, Jurim O, and Eid A. (2002) Effectiveness of acute normovolemic hemodilution to minimize allogeneic blood transfusion in major liver resections. Anesthesiology 97:794-800.

Affiliation/Source of funds

None declared

Study design	Level of evidence	Location/setting
RCT	II	Israel, medical centre

Intervention	Comparator
ANH: Preoperatively, blood was transfused from the patients into standard citrate-phosphate-dextrose blood storage bags, and was simultaneously replaced by colloid solutions. The volume of blood collected was 2,020 ± 412 mL. n=39	No ANH n=39

Population characteristics

Patients undergoing liver resection.

Length of follow-up	Outcomes measured
NR	

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Allocation was randomised; however, it is unclear whether allocation to treatment groups was concealed from those responsible for recruiting subjects.	The treatment arms were similar in baseline characteristics.	Although the surgical staff were not blinded, the investigator who verified the electrocardiogram for possible ischemic episodes was blinded to group assignment. It is unclear the participants were blinded to treatment assignment.	Not detected.	There was no loss to follow up.

Overall quality assessment (descriptive)

Fair

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance
Mortality	0/39 (0%)	0/39 (0%)	Not estimable
Morbidity (adverse cardiac, renal, or neurological outcomes	0/39 (0%)	0/39 (0%)	Not estimable
Patients receiving PRBCs	4/39 (10%)	14/39 (36%)	P=0.014
Haematocrit (%) (before vs after)	40.8 ± 2.7 vs 23.5 ± 1.2 (P<0.05)	41.6 ± 3.2 vs 40.9 ± 2.8 (P>0.05)	

Mean surgical blood loss (mL)	1442 ± 1827	1528 ± 1822		
Clinical importance		Clinical relevance		
EXTERNAL VALIDITY				
Generalisability				
This trial was conducted on a specific patient population (people undergoing liver resection), however it is likely to be generalisable to patients undergoing other elective surgical procedures with moderate blood loss.				
Applicability				
The study was performed in Israel, which may limit its applicability in the Australian context.				
Comments				

Obasi C, Arendt J, and Antoszewski Z. (2006) An assessment of the efficacy of preoperative controlled haemodilution in the perioperative management of patients including the elderly. Chirurgia Polska 8:111-124.

Affiliation/Source of funds

None declared

Study design	Level of evidence	Location/setting
RCT		Poland / hospital

Intervention	Compa	arator	
ANH:	Contro	:	
n=31	n=31		

Population characteristics

Patients undergoing surgical procedures: endoprosthesis of hip joint (13% ANH vs 10% control); anastomis of the femur for fracture (23% ANH vs 29% control); leg amputation (16% ANH vs 19% control); plastic perineal surgery (13% ANH vs 13% control); gastrointestinal anastomosis (6% in both groups).

Length of follow-up	Outcomes measured

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
It is unclear whether allocation was concealed from those responsible for recruiting subjects. Randomisation method was not disclosed.	The percentage of patient > 70 years was 52% in the ANH group and 39% in the control group. Other baseline characteristics were similar between the groups.	The study was not blinded.	There was no transfusion protocol.	There was no loss to follow up.

Overall quality assessment (descriptive)

Poor

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance
Mean (SD) preoperative concentration of haemoglobin, mmol/L	8.37 (0.43)	8.37 (0.63)	NS
Mean (SD) concentration of haemoglobin immediately postoperative, mmol/L	6.45 (0.52)	6.46 (0.56)	NS
Mean (SD) concentration of haemoglobin 6 hours postoperative, mmol/L	7.20 (0.53)	6.48 (0.56)	P<0.005

Clinical importance	Clinical relevance		
EXTERNAL VALIDITY			
Generalisability			
Patients studied were similar to the target population of the guidelines.			
Applicability			
The study was conducted in Poland; which may limit its applicability to the Australian context.			
Comments			

Sanders G, Mellor N, Rickards K, Rushton A, Christie I, Nicholl J, Copplestone A, and Hosie K. (2004) Prospective randomized controlled trial of acute normovolaemic haemodilution in major gastrointestinal surgery. British Journal of Anaesthesia 93:775-781.

Affiliation/Source of funds

NR

Study design	Level of evidence	Location/setting
RCT		UK, hospital

Intervention	Comparator
ANH: Maximum 3 units of blood withdrawn and transfused into blood bags containing citrate-phosphate-dextrose (anticoagulant). Warmed cell-free fluid was administered during blood withdrawal to maintain normovolaemia. At the end of the operation, all the autologous blood was re-transfused. n=78	No ANH n=82

Population characteristics

Patients undergoing major gastrointestinal surgery (colorectal, gastric, or pancreatic). These operations were considered high risk (>40%) for allogeneic transfusion.

Length of follow-up	Outcomes measured
NR (at least 3 days)	Proportion of patients transfused with allogeneic red blood cells, number of units transfused in the first 3 days after surgery, time taken to venesect, perioperative complications, hospital length of stay.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Patients were randomised to treatment arm, however it is unclear whether allocation was concealed from the people responsible for recruiting participants.	Treatment groups were similar in baseline characteristics. There was a statistically significant difference between the groups, in both preoperative and postoperative temperature (P<0.01); however, the difference in median temperatures was 0.1 and 0.3°C, respectively.	The patients were blinded; however, the surgical team were not blinded.	None detected. Transfusion protocol was not overruled for any of the patients.	There was no loss to follow-up

Overall quality assessment (descriptive)

Fair

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance

Mortality	2/78 (3%)	1/82 (1%)	NR
Morbidity	Pyrexia: 0/78 (0%)	Pyrexia: 3/82 (4%)	Pyrexia: P=0.21
	UTI: 8/78 (10%)	UTI: 7/82 (9%)	UTI: P=0.71
	RTI: 2/78 (3%)	RTI: 1/82 (1%)	RTI: P=0.54
	Wound infection: 3/78 (4%)	Wound infection: 6/82 (7%)	Wound infection: P=0.35
	Deep infection: 1/78 (1%)	Deep infection: 0/78 (0%)	Deep infection: P=0.48
	Septicaemia: 1/78 (1%)	Septicaemia: 1/82 (1%)	Septicaemia: P=0.97
	DVT: 2/78 (3%)	DVT: 2/82 (2%)	DVT: P=0.96
	PE: 0/78 (0%)	PE: 2/82 (2%)	PE: P=0.31
	Anastomotic leak: 0/78 (0%)	Anastomotic leak: 3/82 (4%)	Anastomotic leak: P=0.21
Patients receiving allogeneic blood	22/78 (28%)	25/82 (30%)	NS
Units of allogeneic blood transfused.	90	93	NS
Median (range) blood loss (mL)	750-1000 (100-4500)	750-1000 (100-4368)	
Median (range) length of hospital stay, d	8 (5 to 110)	10 (5 to 92)	NS
Clinical importance		Clinical relevance	
EVEDNAL VALIDITY			

EXTERNAL VALIDITY

Generalisability

The study population was people undergoing major gastrointestinal surgery; however, the study is likely to be somewhat generalisable for surgical procedures with a high likelihood for transfusion.

Applicability

Study performed in Canada which has many similarities with the Aus/NZ healthcare systems.

Comments

Abbreviations: DVT, deep vein thrombosis; PE, pulmonary embolism; RTI, respiratory tract infection; UTI, urinary tract infection.

Citation								
Saricaoglu F, Aki hemodilution on o							nd acı	ute hypervolemic
Affiliation/Source	e of funds							
None declared								
Study design Level of evide		evel of evidence	9		Location/s	etting]	
RCT		II				Turkey / Un	iversi	ty hospital
Intervention			Comparators					
ANH: Autologous blood 15 mL/kg was withdrawr replaced by ~15 mL/kg 6% HES n=10		drawn and	· · ·			out removal of any		
Population char	acteristics							
Patients undergo		polastv.						
Length of follow		<u> </u>		Outcor	nes measure	d		
24 h postoperativ						-		
INTERNAL VALI								
Allocation	Results	Blinding ana		ysis	sis Treatment/measurement bias		nt	Follow-up (ITT)
Allocation was concealed from those responsible for recruiting subjects.	The 3 grou similar rega gender, he weight, dur operation, intraoperat loss, posto drainage, a amount of intraoperat crystalloid (p>0.05)	arding age, ight, ration of ive blood perative and the ive	The study was blinded.	not	A transfusio used.	n protocol wa	IS	There was no loss to follow-up.
Overall quality a	ssessment	(descriptive))					
Fair								
RESULTS								
Outcome Intervention group		on group	Con	nparator grou	р	Stati	stical significance	
Median (95% CI) volume of blood withdrawn during ANH, mL		1065 (975,	1170)					
Median (95% CI) intraoperative blo mL		740 (600, 8	•): 650 (500, 85 trol: 695 (510,	•	P=0.	275
		105 (95, 12	, 125)		HHD: 102.5 (95, 125)		P=0.	795

Control: 105 (95, 125)

operation, min

Patients transfused with allogeneic RBCs	2/10 (20%)	HDD: 4/10 (40%) Control: 10/10 (100%)	
Total units of allogeneic RBCs transfused	3 (one patient required 1 unit and the other required 2)	HDD: 5 (three patients required one unit and one patient required 2 units) Control: 13 (7 patients required one unit and 3 patients required 2 units)	P(ANH+HDD vs control)<0.05
Median (95% CI) preoperative haematocrit concentration, %	39.2 (34.6, 46.0)	HHD: 41.1 (37, 45.3) Control: 43.2 (35.8, 45.8)	P=0.5
Median (95% CI) postoperative haematocrit concentration, %	32.7 (26.5, 38.6)	HHD: 29.1 (26.5, 38.6) Control: 32.3 (26.5, 38.6)	P=0.398
Median (95% CI) 24 h postoperative haematocrit concentration, %	32.7 (30.1, 40.1)	HHD: 34.9 (30.2, 36.7) Control: 32.9 (30, 36.5)	P=0.89
Mean (95% CI) preoperative platelet count, 1000/mm ³	280 (132, 367)	HDD: 286 (240, 387) Control: 285 (240, 387)	P=0.98
Mean (95% CI) postoperative platelet count, 1000/mm ³	258 (123, 354)	HDD: 204 (167, 300) Control: 241 (175, 310)	P=0.96
Mean (95% CI) 24 h postoperative platelet count, 1000/mm ³	283 (138, 356)	HDD: 195 (163, 300) Control: 283 (190, 356)	P=0.010
Mean (95% CI) preoperative INR	1.1 (0.92, 1.3)	HDD: 1.15 (0.95, 1.4) Control: 1.15 (0.92, 1.14)	P=0.6
Mean (95% CI) postoperative INR	1.2 (1.1, 2.3)	HDD: 1.4 (1.2, 1.5) Control: 1.35 (1.2, 1.5)	P=0.052
Mean (95% CI) 24 h postoperative INR	1.2 (1.1, 1.87)	HDD: 1.2 (1.1, 1.3) Control: 1.2 (1.1, 1.3)	P=0.68
Mean (95% CI) preoperative aPTT, seconds	27.6 (26.4, 35.9)	HDD: 28.5 (26.8, 32.1) Control: 27.6 (26.4, 32.1)	P=0.4
Mean (95% CI) postoperative aPTT, seconds	26.75 (23.8, 32.3)	HDD: 33.8 (30.1, 35.6) Control: 27.5 (24.7, 34.2)	P=0.01 P(ANH v HDD)<0.008
Mean (95% CI) 24 h postoperative aPTT, seconds	26.5 (24.7, 30.1)	HDD: 30.1 (24.7, 34.2) Control: 24.2 (24.2, 34.7)	P=0.182
Clinical importance		Clinical relevance	
EXTERNAL VALIDITY			
Generalisability			

Patients studied were similar to the target population of the guidelines.

Applicability

The study was conducted in Turkey; which may limit its applicability to the Australian context.

Comments

Abbreviations: HES, hydroxyethyl starch; HHD, hypervolemic haemodilution; INR, international normalised ratio; aPTT, activated partial thromboplastin time

Wolowczyk L, Nevin M, Smith FCT, Baird RN, and Lamont PM. (2003) Haemodilutional effect of standard fluid management limits the effectiveness of acute normovolaemic haemodilution in AAA surgery - Results of a pilot trial. European Journal of Vascular and Endovascular 26:405-411.

Affiliation/Source of funds

None declared.

Study design	Level of evidence	Location/setting
RCT	II	UK / hospital

Intervention	Comparator
ANH and cell salvage: 15 g/kg of blood was withdrawn and replaced with a similar volume of 6% HAES	Standard care (including cell salvage) n=18
n=18 (although two patients were not included in the	

Population characteristics

Patients undergoing abdominal aortic aneurysm repair.

Length of follow-up	Outcomes measured
Seven days postoperative	

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
It is unclear whether allocation was concealed from those responsible for recruiting subjects.	Both groups were well matched. The incidence of ischemic heart disease in the form of previous myocardial infarction and/or angina was similer: 5/16 ANH and 7/18 control patients. One patient randomised to the ANH group had a successful coronary artery bypass graft prior to AAA repair. All AAAs were infrarenal, but supra-renal aortic clamping was necessary in 2 ANH and 3 control patients.	The study was not blinded.	A transfusion protocol was used.	The study was not performed ITT. Two patients randomised into the ANH group exhibited signs of poor cardiac reserve soon after induction of general anaesthesia, which precluded the ANH. Those two patients were excluded from further analysis.

Overall quality assessment (descriptive)

Fair

RESULTS	Interventing and	Commonator	Chatiatical along!
Outcome	Intervention group	Comparator group	Statistical significance
Median (range) amount of blood withdrawn during ANH, g	890 (670, 1620)	-	-
Median (IQR) intraoperative blood loss, mL	1780 (930, 5000)	1700 (750, 2600)	P=0.55
Patients with blood loss below 1000 mL	4/16 (25%)	5/18 (28%)	P=1.0
Median (IQR) volume of RBCs concentrate recovered by intraoperative cell salvage and retransfused	590 (200, 1410)	540 (210, 740)	P=0.60
Patients transfused with banked autologous blood intraoperatively	7/16 (44%)	7/18 (39%)	P=1
Patients transfused with banked autologous blood postoperatively	5/16 (31%)	10/18 (56%)	P=0.73
Total patients transfused with banked autologous blood(intraoperatively and postoperatively)	10/16 (63%)	13/18 (72%)	P=0.99
Median (IQR) units of banked autologous blood transfused intraoperatively	0 (0 to 4)	0 (0 to 2)	P=0.51
Median (IQR) units of banked autologous blood transfused postoperatively	0 (0 to 2)	1 (0 to 2)	P=0.33
Median (IQR) units of banked autologous blood transfused (intraoperatively and postoperatively)	2 (0 to 5)	2.5 (0 to 5)	P=0.68
Median (range) Haemoglobin concentration, g/dL	Preoperative: 14.2 (12.1 to 16.5) Post-ANH: 9.4 (7.0 to 12.1) At aortic clamping: 9.2 (6.8 to 10.6) At clamp release: 7.7 (6.6 to 9.3) Post-op 1-2 h: 10.8 (8.8 to 13.3) Post-op day 1: 10.4 (8.3 to 12.4) Post-op day 2: 10.6 (8.2 to 13.3) Post-op day 7: 11.5 (10.2 to	Preoperative: 13.8 (12.1 to 15.6) Post-ANH: NA At aortic clamping: 11.3 (7.2 to 14.5) At clamp release: 9.1 (5.1 to 11.9) Post-op 1-2 h:10.3 (8.1 to 12.7) Post-op day 1: 10.4 (8.2 to 12.8) Post-op day 2: 9.7 (8.5 to 13.7) Post-op day 7: 10.7 (9.1 to	P-value Preoperative: 0.57 Post-ANH: NA At aortic clamping: 0.001 At clamp release: 0.004 Post-op 1-2 h: 0.68 Post-op day 1: 0.68 Post-op day 2: 0.60 Post-op day 7: 0.021

Clinical importance	Clinical relevance		
EXTERNAL VALIDITY			
Generalisability			
The study population was people undergoing abdominal aortic repair; however, the study is likely to be somewhat generalisable for surgical procedures with moderate blood loss.			
Applicability			
The study is likely to be applicable to the Australian context.			
Comments			

Abbreviations: HAES, hydroxyethyl starch

Intervention 2 – Intraoperative cell salvage

Level I evidence

LEVEL I EVIDENCE						
Citation						
	A, O'Connell D, and Hen ion Medicine 14:123-144.		ologous ti	ransfusion tec	hniques: A system	natic review of their
Affiliation/Source	e of funds					
Research supported by a grant obtained from the National Health and Medical Research Council of Australia and a special purpose grant from the Hunter Area Pathology Service, Australia						
Study design	Le	evel of evidence)		Location/setting	g
Systematic review observational studentallysis	dies with meta-			NA		
Search conducted	1 July 2002					
Intervention			Compa	rator		
Autologous transfusion techniques: preoperative Autologous blood deposit (PAD), ANH, and cell salvage (CS). NOTE: This form only contains RCT info relevant for			Comparator: No Autologous transfusion technique (active versus active comparisons were excluded) Sample size (control for perioperative cell salvage) N=1052			
intraoperative cell salvage.						
	operative cell salvage) N=	=10/3				
Population chara						101 1 1
orthopaedic surge	n 18 years undergoing ar ery, and four involved vas were studies (1.8:1).					
Length of follow	-up		Outcon	nes measure	d	
NA			thrombo	osis, non-fatal	n, infection, wound MI, rate of alloger me of allogeneic b	neic red blood cell
INTERNAL VALI	DITY					
Allocation	Results	Blinding analy	<i>y</i> sis	Treatment/i bias	measurement	Follow-up (ITT)
Both the allocation concealment and the method of randomisation were judged by the authors to be inadequate in every case.	SR did not discuss similarity between preoperative data and baseline characteristics for the intervention groups.	The majority of trials were unbl (96%).		agreement l SR raters w	assessment, the petween the two ere moderate to a=0.65 to 1.0)	NR
	ssessment (descriptive))				
Fair						

RESULTS			
Outcome	Intervention group	Comparator group	Statistical significance
Number of subjects exposed to allogeneic RBC transfusion (perioperative CS) 26 trials (N=1939; 973 cell salvage, 966 control)			RR (95% CI): 0.58 (0.47, 0.73) Phet<0.00001
Number of subjects exposed to allogeneic RBC transfusion (perioperative CS): transfusion protocol used 23 trials (N=NR)			RR (95% CI): 0.62 (0.50, 0.78) Phet=NR
Number of subjects exposed to allogeneic RBC transfusion (perioperative CS): no transfusion protocol used/reported 3 trials (N=NR)			RR (95% CI): 0.14 (0.00, 4.48) Phet=NR
Number of subjects exposed to allogeneic RBC transfusion (perioperative CS): cell washing used 13 trials (N=NR)			RR (95% CI): 0.52 (0.38, 0.71) Phet=NR
Number of subjects exposed to allogeneic RBC transfusion (perioperative CS): cell washing not used/reported 12 trials (N=NR)			RR (95% CI): 0.74 (0.58, 0.94) P <i>het</i> =NR
Number of subjects exposed to allogeneic RBC transfusion (perioperative CS): cardiac surgery 12 trials (N=NR)			RR (95% CI): 0.82 (0.70, 0.95) Phet=NR
Number of subjects exposed to allogeneic RBC transfusion (perioperative CS): orthopaedic surgery 11 trials (N=NR)			RR (95% CI): 0.35 (0.24, 0.52) Phet=NR
Number of subjects exposed to allogeneic RBC transfusion (perioperative CS): miscellaneous surgery 3 trials (N=NR)			RR (95% CI): 0.55 (0.13, 2.36) Phet=NR
Number of subjects exposed to allogeneic RBC transfusion (intraoperative CS) 5 trials (N=NR)			RR (95% CI): 0.61 (0.39, 0.95) Phet=NR

Mean (95% CI) units of allogeneic blood transfused (intra- and postoperative) 17 trials (N=NR)	RR (95% CI): 0.91 (0.51, 1.31) Phet<0.00001
Length of hospital stay (intra- and postoperative), days 5 trials (N=NR)	WMD (95% CI): -1.28 (-2.65, 0.08) P <i>het</i> =NR
Mortality (intra- and postoperative) 11 trials (N=NR)	RR (95% CI): 1.53 (0.65, 3.61) Phet=0.66
Infection (intra- and postoperative) 9 trials (N=NR)	RR (95% CI): 0.75 (0.41, 1.37) Phet=0.37
Wound complications (intra- and postoperative) 7 trials (N=NR)	RR (95% CI): 0.88 (0.42, 1.81) Phet=NR
Thrombosis (intra- and postoperative) 6 trials (N=NR)	RR (95% CI): 1.46 (0.56, 3.83) Phet=NR
Non-fatal MI (intra- and postoperative) 5 trials (N=NR)	RR (95% CI): 0.58 (0.28, 1.19) Phet=NR
Re-operation (intra- and postoperative) 8 trials (N=NR)	RR (95%CI): 1.08 (0.47, 2.48) Phet=NR
Clinical importance	Clinical relevance

EXTERNAL VALIDITY

Generalisability

Patients considered similar to guideline target population

Applicability

All the studies included in this review were conducted in countries with well developed healthcare systems (not specifically Aus/NZ).

Comments

According to the authors, 5 trials assessed the use of intraoperative CS. The authors do not identify these papers nor do they state the number of participants in the trials.

Carless PA, Henry DA, Moxey AJ, O'connell DL, Brown T, and Fergusson DA. (2006) Cell salvage for minimising perioperative allogeneic blood transfusion. Cochrane database of reviews; Issue 4.

Affiliation/Source of funds

None declared

Study design	Level of evidence	Location/setting
SR	1	NA
Jan 2004		

Intervention	Comparator
Cell salvage. Studies with a combination of a comparisons were included if both the intervence control groups were equally exposed to the a treatment (ie, active plus cell salvage versus comparisons). n (perioperative cell salvage): 1952 n (intraoperative cell salvage): 282 n (postoperative cell salvage): 1448 n (intra- + postoperative cell salvage): 142 The authors found 51 studies.	ntion and n (perioperative cell salvage): 1905

Population characteristics

Adults (over 18 years) undergoing elective, non-urgent surgery. Surgery types found in the search include cardiac (23 studies), orthopaedic (23 studies), and vascular (5 studies) surgery. 33 of the trials studied cell salvage during the postoperative period, 10 studied intraoperative cell salvage, and seven studied both intraoperative and postoperative cell salvage. One trial failed to describe the timing of cell salvage. Twenty trials studied cell salvage systems that reinfused washed salvaged blood, and 29 trials studied cell salvage systems that reinfused unwashed filtered salvaged blood. One trial studied both washed and unwashed cell salvage (4-arm trial) and provided two comparisons of cell salvage. 38 trials reported the use of transfusion protocols.

Length of follow-up	Outcomes measured
NA	Number of patients transfused with allogeneic and/or autologous blood, amounts of allogeneic and/or autologous blood transfused, re-operation for bleeding, postoperative complications, mortality, and length of hospital stay.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
According to the authors, the description of the method to conceal allocation of either inadequate or unclear for all of the studies.	-	Based on the Schulz criteria, blinding was reported in only one of the trials.	Nine of the 51 studies did not report a transfusion protocol.	-

Overall quality assessment (descriptive)			
Good			
RESULTS			
Outcome	Intervention group	Comparator group	Statistical significance
Rate of allogeneic transfusion (intraoperative cell salvage; aggregated analysis¹) 7 trials (N=564; 282 cell salvage, 282 control)			RR (95% CI): 0.53 (0.35, 0.80) P=0.0027 (Phet=0.00015)
Rate of allogeneic transfusion (intraoperative cell salvage; active vs control) 5 trials (N=382; 191 cell salvage, 191 control)	76/191 (40%)	113/191 (59%)	RR (95% CI): 0.61 (0.39, 0.95) P=0.029 (Phet=0.01)
Clinical importance	•	Clinical relevance	
EXTERNAL VALIDITY			
Generalisability			

The SR is generalisable for elective, non urgent surgery.

Applicability

The studies were mostly from countries with similar health-care systems to Australia

Comments

The systematic review includes trials assessing the intraoperative, postoperative, and both intra- and postoperative. There is more data but most of it combines intra- and postoperative data.

¹ Includes both studies where an active treatment is compared with a control intervention and those studies where both the intervention and control arms also received an additional active intervention (c.f. active versus control, where the controls are untreated). Abbreviations: RR, relative risk; ARR, absolute risk reduction; RRR, relative risk reduction

Davies L, Brown TJ, Haynes S, Payne K, Elliott RA, and McCollum C. (2006) Cost-effectiveness of cell salvage and alternative methods of minimising perioperative allogeneic blood transfusion: A systematic review and economic model. Health Technology Assessment 10:1-114.

Affiliation/Source of funds

One author received sponsorship from haemonetics and AstraTech to attend the International Society of Blood Transfusion (ISBT) VIIIth European Congress. The author has also given invited lectures for AstraTech Ltd and Unomedical with honoraria and expenses paid.

Study design	Level of evidence	Location/setting
SR with economic analysis (SR search is an update of Carless 2003: a Cochrane review that had been updated in 2006. The study includes a meta-analysis combining the results of Carless 2003 and the search update.		NA
Search conducted Jan 2004.		

Intervention	Comparator
Transfusion strategies to minimise perioperative allogeneic blood transfusion: cell salvage, PAD, PAD plus EPO, EPO, ANH, cell salvage plus ANH, AFs, FSs, restrictive transfusion thresholds or protocols.	No cell salvage or allogeneic blood.
NB: This form only includes information relevant for perioperative cell salvage.	
Specific characteristics of the 1 included RCT (Zhao 2003)	
Non-washed shed mediastinal blood retransfused postoperatively after CABG; mean 280 mL autologous blood retransfused/	

Population characteristics

For inclusion, the SRs had to only include adults undergoing elective, non-urgent surgery.

Specific characteristics of participants in the 1 included RCT (Zhao 2003)

Patients undergoing CABG surgery

Mean (SD) age (study vs control): 59.2 (8.2) vs 59.5 (8)

Sex (M/F): 27/3 vs 26/4

Length of follow-up	Outcomes measured
NA	Proportion/number of patients transfused with allogeneic and/or autologous blood; the volume of allogeneic and/or autologous blood transfused; reoperation for bleeding; adverse transfusion reactions; preoperative morbidity and Hb levels; postoperative complications; length of hospital stay; mortality.

INTERNAL VALII	INTERNAL VALIDITY ¹			
Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Method of randomisation not described and allocation concealment unclear.	Zhao 2003 had adequate baseline comparability.	Did not have participant blinding and it is unclear whether study had allocation concealment.	The study had a well defined transfusion protocol.	Unclear intention to treat (however there was no loss to follow-up).

Overall quality assessment (descriptive)

Good

RESULTS

1,125215		
Intervention group	Comparator group	Statistical significance
74/191 (41%)	113/191 (59%)	RR (95% CI): 0.61 (0.39, 0.95) Phet=0.01
<u> </u>	Clinical relevance	<u> </u>
		74/191 (41%) 113/191 (59%)

EXTERNAL VALIDITY

Generalisability

The review includes all surgery types and performs subgroup analyses by surgery type. Therefore the results are likely to be generalisable for other elective, non-emergency operations.

Applicability

Low applicability to the question - With the exception of transfusion frequency, all of the outcomes assessed combined data from intra- and postoperative cell salvage.

Comments

The updated lit search included 2 RCTs (1 as abstract only): Naumenko 2003 and Zhao 2003. Zhao 2003 investigated intraoperative cell salvage (described in this form), and Naumenko investigated postoperative cell salvage (described in I4).

¹Refers only to the one intraoperative cell salvage RCT included in the systematic update (Zhao 2003)

²Does not include studies where both the intervention and control arms received an additional active intervention.

³Combines data from 5 trials using intraoperative cell salvage, 18 trials of postoperative cell salvage and 5 trials assessing combined intra-+ postoperative cell salvage.

⁴Includes both studies where an active treatment is compared with a control intervention and those studies where both the intervention and control arms also received an additional active intervention. Includes cell salvage in both the intra- and postoperative periods.

Abbreviations: CABG, Coronary Artery Bypass Graft; MI, myocardial infarction

Huet C, Salmi R, Fergusson D, Koopman-Van Gemert AWMM, Rubens F, and Laupacis A. (1999) A meta-analysis of the effectiveness of cell salvage to minimize perioperative allogeneic blood transfusion in cardiac and orthopedic surgery. Anesthesia and Analgesia 89:861-869.

Affiliation/Source of funds

Coordinating Centre has been funded by Janssen Ortho Inc, Canada. One of the authors is the recipient of the First Fellowship from the International Society of Technology Assessment in Health Care, funded by the PPP Medical Trust.

Study design	Level of evidence	Location/setting
SR of RCTS with Meta-analysis	1	NA
Search conducted 1997		

Intervention	Comparator
Intraoperative cell salvage	Control

Population characteristics

Patients who underwent cardiac or orthopaedic surgery (two articles dealing with vascular surgery were not considered). The one study for intraoperative cell salvage included was in patients undergoing hip surgery.

Length of follow-up	Outcomes measured
NA	Proportion of patients receiving at least one unit of allogeneic packed red blood cells

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Three of the included trials were pseudorandomised.		Unclear: "all of the 28 trials included in this study scored between zero and three on the Jadad scale. Because it is difficult to "blind" the operative team to the presence or absence of cell salvage, the Jadad score would rarely be expected to be greater than 3"	The SR did not report on whether or not a transfusion protocol was used in the RCTs	NR

Overall quality assessment (descriptive)

Good

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance
Patients transfused with allogeneic blood	6/20 (30%)	18/20 (90%)	RR (95% CI): 0.33 (0.17, 0.66)
1 trial (N=40; 20 cell salvage, 20 control)			P<0.05 (P <i>het</i> =NA)

Mean units of allogeneic blood transfused	0.7	2.7	NR			
1 trial (N=40; 20 cell salvage, 20 control)						
Clinical importance		Clinical relevance				
·						
EXTERNAL VALIDITY						
Generalisability						
The review includes all surgery types and performs subgroup analyses by surgery type. Therefore the results are likely to be generalisable for other elective, non-emergency operations.						
Applicability						
Low applicability to the question - all of the outcomes assessed combined data from intra- and postoperative cell salvage.						
Comments						
1 trial for intraoperative cell salvage included						

¹Includes both studies where an active treatment is compared with a control intervention and those studies where both the intervention and control arms also received an additional active intervention (c.f. active versus control, where the controls are untreated).

Citation							
Takagi H, Sekino aneurysm surgery							abdominal aortic Surgery 142:1098-1101.
Affiliation/Source	e of funds						
None declared							
Study design		Lo	evel of evidence	9		Location/setti	ng
SR		1				NA	
November 2005							
Intervention				Compa	rator		
Intraoperative auto	otransfusion			Control			
Population chara	acteristics						
Patients undergoi	ng elective ir	frarenal abo	lominal aortic and	eurysm sı	ırgery.		
Length of follow-	-up			Outcon	nes measure	d	
NR			Incidend	ce of allogene	ic blood transfus	ion	
INTERNAL VALIDITY							
Allocation	Results	Results Blinding anal		ysis	is Treatment/measurement Follow-up bias		Follow-up (ITT)
It is not clear whether any of the trials concealed allocation from those responsible for recruiting subjects.		r baseline were not blinde		ed in			
Overall quality as	ssessment (descriptive)				
Good							
RESULTS							
Outcome		Intervention	on group	Com	parator grou	p St	atistical significance
Number of patient received allogene transfusion 4 trials (N=292)	lumber of patients who eceived allogeneic ansfusion 67/147 (45.6)		109/145 (75.1)		0.9	R (95% CI): 0.63 (0.41, 95) 0.05 (P <i>het</i> =0.02)	
Clinical importance			Clinical relevance				
EXTERNAL VALI	DITY						
Generalisability							
The study populat	ions are simi	lar to the gu	ideline populatio	n.			

Applicability
The results are applicable to the Australian context.
Comments

Level II evidence

Citation									
Bowley DM, Bark controlled trial. W	er P, and Bot orld Journal (fard KD. (20 of Surgery 30	06) Intraoperativ 0:1074-1080.	e blood s	alvage in pene	etrating abdor	minal	trauma: A randomised,	
Affiliation/Sourc	e of funds								
NR									
Study design		Le	evel of evidence	9		Location/se	etting	J	
RCT		II				Johannesbu Africa), hosp		Republic of South	
Intervention				Compa	rator				
Intraoperative blo allogeneic and wa N=21	ashed autolog		on of both	Allogen medical N=23		sfusion at the	discr	etion of the attending	
Population chara									
Patients with pen- arrival and in who						d hypotension	eithe	er pre-hospital or on	
Length of follow	-up			Outcon	nes measure	d			
24 hours post-injury				Volume of allogeneic blood transfused, mortality					
INTERNAL VALI	DITY								
Allocation	Results		Blinding analy	ysis Treatment/measuremer bias		measuremen	t	Follow-up (ITT)	
Patients were randomised and allocation was concealed	Score, Pen Injury Seve	not blinded. stics in re-hospital n known), time, rauma ıry Severity		n was	Transfusion protocol NR.			It is unclear whether the analyses were conducted ITT.	
Overall quality a	ssessment ((descriptive)						
Fair									
RESULTS									
Outcome		Intervention	on group	Con	omparator group St		Stati	Statistical significance	
Mean volume (un allogeneic blood t		6.47 (5.14)		11.1	1.17 (6.06) P:		P=0.008		
Mortality		14/21 (67%	<u> </u>	15/2	3 (65%)		P=NS		
Mean length of ho (days)	ospital stay	15.7 (9.17)		14.6 (6.8)			P=0.	79	

Clinical importance

EXTERNAL VALIDITY

Generalisability

The study assessed the use of cell salvage in traumatic surgery. Therefore, the study is not generalisable for elective, non-emergency surgery.

Applicability

The study was conducted in Johannesburg, which may limit the applicability of the study.

Comments

The mean volume of salvaged blood retransfused in the cell salvage group: 1493 mL ± 617.43 mL.

Citation						
	teinbruchel DA. (2006 Scandinavian Cardiova			saver for off-po	ump coronary arte	ry bypass surgery: A
Affiliation/Source	of funds					
One of the authors	was funded by a rese	arch grant from H:	S Copenl	nagen Hospita	al Corporation	
Study design		Level of evidence	9		Location/settin	g
RCT		l			Denmark, hospit	tal
Intervention			Compa	rator		
The continuously heparinised suction and reservoir belonging to the cell saver were used for all patients in both groups. The suctioned blood from patients in the cell saver group was processed and autotransfused before the patient was transferred to the ICU. N=30		Control group had their suctioned blood discharged. N=30				
Population charac						
	or elective or sub-acut	e coronary bypass				
Length of follow-u	ıb			nes measure		
			Proportion of patients receiving transfusion with allogeneic blood components, average number of units transfused per patient during the admission, intraoperative and postoperative bleeding, changes in haemoglobin and haematocrit, and registration of complications and costs.			
INTERNAL VALIDI	ITY	1		T		T
Allocation	Results	Blinding anal	ysis	Treatment/ bias	measurement	Follow-up (ITT)
were	The intervention groups had similar baseline characteristics. Cell saver group patients received median 1.26 mL of autologous cell saver blood.	Both treatment control groups underwent blood salvage. Randomisation blood retransfudischarge did noccur until after operation. The the surgical and anaesthetic teatwere blinded do the operation, after. However ICU and ward personnel were informed about procedure had performed.	n to usion or not er the refore ad am luring but not r, the e not t which	A transfusioused.	n protocol was	Intraoperatively, one patient in the control group was excluded (before randomisation). Due to logistic or technical problems the cell saver was not used for five patients in the cell saver group. According to intention to treat principles, they were kept in the study analysis.
Overall quality as:	sessment (descriptiv	re)				1
Good						

RESULTS			
Outcome	Intervention group	Comparator group	Statistical significance
Patients receiving allogeneic blood components	17/30 (57%)	21/29 (72%)	P=0.28
Median (IQR) volume (units) of allogeneic blood components transfused per patient	1 (0 to 2)	2 (0 to 7)	P=0.06
Median (IQR) units of allogeneic RBCs transfused per patient	1 (0 to 2)	2 (0 to 5)	P=0.07
Median (IQR) units of FFP transfused (ICU)	0 (0 to 0) (range: 0 to 4)	0 (0,0) (range: 0 to 22)	P=0.40
Median (IQR) units of FFP transfused (ward)	0	0 (0, 0) (range: 0 to 1)	P=0.31
Median (IQR) units of pooled platelets transfused	0 (0 to 0) (range:0 to 1)	0 (0 to 0) (range: 0 to 1)	P=NR
Median (IQR) haemoglobin, intensive care (lowest), mmole/L	5.9 (5.3 to 6.6)	5.8 (5.2 to 6.7)	P=0.97
Median (IQR) haematocrit, intensive care (lowest), %	29 (27 to 33)	29 (25 to 33)	P=0.69
Median (IQR) haemoglobin, ward (lowest), mmole/L	6.4 (5.9 to 6.8)	6.6 (5.8 to 7.1)	P=0.58
Median (IQR) haemoglobin, at discharge, %	7.1 (6.5 to 7.4)	7.2 (6.5 to 8.1)	P=0.25
Median (IQR) duration of operation, min	165 (135 to 186)	150 (135 to 188)	P=0.39
Median (IQR) net blood less, mL	300 (193 to 403)	610 (450 to 928)	P<0.001
Mortality	0/30 (0%)	2/30 (7%)	P=0.24

Morbidity	Stroke: 0/30 (0%)	Stroke: 1/30 (3%)	P-value
	MI: 0/30 (0%)	MI: 1/30 (3%)	Stroke: NS
	Reoperation for bleeding: 1/30 (3%)	Reoperation for bleeding: 3/30 (10%)	MI: NS Reoperation for bleeding:
	Pneumonia: 2/30 (7%)	Pneumonia: 3/30 (10%)	0.35
	GI bleeding: 0/30 (0%)	GI bleeding: 3/30 (10%)	Pneumonia: NS
	Deep sterna wound infection: 0/30 (0%) Leg wound infection: 0/30 (0%) Dialysis: 1/30 (3%) Ventilator > 24 hours: 0/30 (0%) Low cardiac output syndrome: 0/30 (0%) Atrial arrhythmia: 14/30 (47%) Ventricular arrhythmia: 0/30 (0%)	Deep sterna wound infection: 1/30 (3%) Leg wound infection: 1/30 (3%) Dialysis: 2/30 (7%) Ventilator > 24 hours: 3/30 (10%) Low cardiac output syndrome: 6/30 (20%) Atrial arrhythmia: 20/30 (67%) Ventricular arrhythmia: 3/30 (10%)	GI bleeding: 0.11 Deep sterna wound infection: NS Leg wound infection: NS Dialysis: NS Ventilator > 24 hours: 0.11 Low cardiac output syndrome: 0.01 Atrial arrhythmia: 0.12 Ventricular arrhythmia: 0.11 Inotropic infusion: 0.38
	Inotropic infusion: 6/30 (20%)	Inotropic infusion: 9/30 (30%)	
Median (IQR) length of hospital admission, days	7 (6 to 8)	7 (6 to 9)	NS
Intensive care unit > 24 hours	1 (3%)	6 (21%)	P=0.05
Clinical importance	•	Clinical relevance	
EXTERNAL VALIDITY			
Generalisability			
The study assesses coronal	ry artery bypass, but is still somew	hat generalisable other elective	surgeries.
Applicability			
Although the trial was condu	ucted in Denmark, it is likely to be	applicable to the Australian conte	ext.
Comments			

Abbreviations: EF, ejection fraction; CPB, cardiopulmonary bypass; OPCAB, off-pump coronary artery bypass grafting

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Citation								
Cool D. Dannu H.	Mohan D. ar	ad Arora D	(2007) Efficacy of	f coll cav	or in roducing	homologous	blood	transfusions during
OPCAB surgery:							טטטוע	i italisiusions uuring
Affiliation/Source	e of funds							
None declared								
Study design		L	evel of evidence	;		Location/s	ettino]
RCT		II				India / Hos	pital	
Intervention				Compa	rator			
Intraoperative cell salvage and autotransfusion of washed shed blood and transfusion of allogeneic blood if required. N=25 (although one patient was excluded from analysis)			Control N=25	only transfus	ed with allog	eneic	homologous blood.	
Population char	acteristics		·					
Patients undergo	ing off-pump	coronary art	ery bypass grafti	ng.				
Length of follow	-up			Outcor	nes measure	d		
NR			Transfusion frequency and volume of transfusion, haematocrit concentration, haemoglobin concentration, morbidity, mortality					
INTERNAL VALI	INTERNAL VALIDITY							
Allocation	Results		Blinding analy	Blinding analysis Treatment bias		measureme	nt	Follow-up (ITT)
Allocation was concealed from those responsible for recruiting subjects.	The interve groups had demograph characteris	d similar blinded blinded		not	A transfusion protocol was		as	One patient randomised to the cell saver group was excluded as he crashed on opening the left anterior descending artery and the grafting was completed on pump.
Overall quality a	ssessment ((descriptive	<u>)</u>		l			
Fair								
RESULTS								
Outcome		Interventi	on group	Comparator grou		р	Statistical significance	
Mean (SD) volum autotransfused, n		714.8 (317	7.5)	NA			NA	
Mean haematocrit 34.6 (4.6) concentration of the autotransfused blood, %			NA			NA		
Patients requiring blood transfusion		20/24 (839	%)	25/25 (100%)			0.05	
Mean (SD) units of allogeneic blood to		1.5 (1.1)		2.4 (1.3)			P=0.02	

Mean (SD) postoperative haemoglobin concentration, g/dL	10.9 (1.5)	9.6 (0.9)	P=0.0007			
Mean (SD) decrease in haemoglobin from preoperative to immediate postoperative, g/dL	1.8 (1.2)	2.7 (1.6)	P=0.02			
Mortality	0/24 (0%)	0/25 (0%)	NS			
Morbidity			There was no re-exploration or deep sterna wound infection in either group.			
Clinical importance		Clinical relevance	Clinical relevance			
EXTERNAL VALIDITY						
Generalisability						
Patients considered similar to	guideline target population	on.				
Applicability						
The study was conducted in India, which may limit its applicability to the Australian context.						
Comments						

STUDY DETAILS

Citation

Mercer KG, Spark JI, Berridge DC, Kent PJ, and Scott DJA. (2004) Randomized clinical trial of intraoperative autotransfusion in surgery for abdominal aortic aneurysm. British Journal of Surgery 91:1443-1448.

Affiliation/Source of funds

Authors received support from The Sir Jules Thorne Charitable Trust

Study design	Level of evidence	Location/setting
RCT		UK / university hospital

Intervention	Comparator
Intraoperative cell salvage. Processed blood was returned to the patient as soon as haemostasis ha achieved. N=40	d been Control: allogeneic blood transfusion only N=41

Population characteristics

Patients undergoing surgery for abdominal aortic aneurysm.

Length of follow-up	Outcomes measured
NR (at least until hospital discharge)	Operative blood loss, patients transfused with allogeneic blood, morbidity, mortality, length of hospital stay.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Allocation was concealed from those responsible for recruiting subjects (using sealed envelopes)	The treatment groups had similar demographic characteristics.	Patients were blinded to the transfusion group allocation. Members of the operating surgical team were responsible for the continuing care of patients, decision to use blood transfusion and investigation of postoperative complications. They were independent of the research team, but were not blinded.	A transfusion protocol was used.	All analyses were conducted ITT. Four patients in the treatment arm did not receive autologous transfusion because less than 500 mL of shed blood was collected.

Overall quality assessment (descriptive)

Good

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance
Median (IQR) operative blood loss, mL	1950 (775 to 285)	1270 (775 to 2850)	P=0.140

Median (IQR) volume of salvaged blood retransfused, mL	650 (500 to 1125)	-	-		
Patients transfused with allogeneic blood	21/40 (53%)	31/41 (76%)	P=0.038		
Median (IQR) units of allogeneic blood transfused	1 (0 to 3)	3 (1 to 5)	P=0.012		
Median (IQR) units of allogeneic and/or autologous blood transfused	3 (2 to 6)	3 (1 to 5)	P=0.783		
Morbidity	SIRS: 9/40 (23%) Infection: 5/40 (13%) ¹ Sepsis: 4/40 (10%)	SIRS: 20/41 (49%) Infection: 14/41 (34%) ² Sepsis: 8/41 (20%)	P-value SIRS: 0.020 Infection: 0.035 Sepsis: 0.349		
Mortality	1/40 (3%)3	1/41 (2%)	P=1.000		
Median (IQR) length of hospital stay, d	12 (8 to 19)	13 (10 to 19)	P=0.385		
Clinical importance		Clinical relevance			
EXTERNAL VALIDITY					
Generalisability					
The study population is similar	r to the guideline population				
Applicability					
The results are applicable to the Australian context.					
Comments					

¹Including four chest infections and one line infection. ²Including twelve chest infections, one graft infection, and one blood infection.

³The patient died within 30 days of surgery owing to postoperative myocardial infarction and

⁴The patient died in hospital 37 days after surgery with pneumonia, MRSA septicaemia and acute renal failure Abbreviations: MRSA, methicillin-resistant Staphylococcus aureus; SIRS, systemic inflammatory response syndrome

Murphy GJ, Rogers CS, Lansdowne WB, Channon I, Alwair H, Cohen A, Caputo M, and Angelini GD. (2005) Safety, efficacy, and cost of intraoperative cell salvage and autotransfusion after off-pump coronary artery bypass surgery: A randomized trial. Journal of Thoracic and Cardiovascular Surgery 130:20-28.

Affiliation/Source of funds

None declared

Study design	Level of evidence	Location/setting
RCT	II	UK / hospital

Intervention	Comparator
Intraoperative cell salvage, with Autotransfusion of washed, salvaged red blood cells at the completion of the operative procedure. Salvaged washed RBCs were autotransfused at the time of skin closure.	In the control group, all blood spilled, from skin incision to skin closure, was aspirated with a high-pressure sucker and discarded. N=31
N=30	

Population characteristics

Patients scheduled for non-emergency first-time CABG (off-pump).

Lengt	h of follow-up	Outcomes measured
NR		Transfusion frequency, mortality, ICU length of stay, hospital length of stay, morbidity, change in haemoglobin levels.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Patients were randomised to treatment arms and allocation was concealed from those in charge of recruiting subjects.	The study groups were similar in terms of baseline characteristics	The study was not blinded.	Transfusion protocol utilised.	There was no loss to follow-up.

Overall quality assessment (descriptive)

Fair

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance
Number of patients autotransfused, n (%)	20/30 (67%)		
Median (IQR) volume salvaged (range) / median (IQR) volume autotransfused, mL	747 (607 to 978) / 236 (206 to 342)		
Number of patients receiving any allogeneic blood product, n (%)	5/30 (17%)	11/31 (36%)	OR (95% CI): 0.36 (0.11, 1.22) P=0.095)

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Number of patients receiving homologous blood transfusion, n (%)	4/30 (13%)	7/31 (23%)	OR (95% CI): 0.53 (0.14, 2.03) P=0.35
Number of patients receiving platelet transfusion, n (%)	2/30 (7%)	6/31 (19%)	OR (95% CI): 0.30 (0.06, 1.61) P=0.26
Number of patients receiving clotting factor transfusion, n (%)	0/30 (0%)	1/31 (3%)	P>0.99
Mortality	0/30 (0%)	0/31 (0%)	P=NS
Median (IQR) length of hospital stay, d	6.0 (5.0, 8.3)	6.0 (5.0, 8.0)	OR (95% CI): 1.08 (0.65, 1.80) P=0.73
Median (IQR) length of ICU stay, d	1.0 (1.0, 1.0)	1.0 (1.0, 1.0)	OR (95% CI): 1.14 (0.67, 1.96) P=0.50
Readmission to ICU, n (%)	1/30 (3)	1/31 (3)	OR (95% CI): 1.0 (0.06, 17.32) P=0.98)
Morbidity, n (%)	Arrhythmia: 6/30 (20%)	Arrhythmia: 7/31 (23%)	OR (95% CI) Arrhythmia: 0.86 (0.25, 2.92); P=0.81
	Pulmonary complications: 0/30 (0%)	Pulmonary complications: 4/31 (13)	Pulmonary complications: NA; P=0.11
	Stroke: 0/30 (0%)	Stroke: 0/31 (0%)	Stroke: NA; P=1.0
	Infective complications: 2/30 (7%)	Infective complications: 1/31 (3%)	Infective complications: NA; P=0.41
	Renal complications: 0/30 (0%)	Renal complications: 2/31 (7%)	Renal complications: NA; P=0.49
	Myocardial infarction: 2/30 (7%)	Myocardial infarction: 0/31 (0%)	Myocardial infarction: NA; P=0.24
Mean (SE) haemoglobin, g/dL	After protamine: 11.14 (0.21) 1 h: 10.55 (0.21) 24 h: 11.71 (0.21)	After protamine: 11.25 (0.21) 1 h: 10.40 (0.20) 24 h: 10.69 (0.20)	Difference/ratio (95% CI) After protamine: 0.11 (-0.47, 0.70) P=0.71 1 h: -0.15 (-0.74, 0.43)
			P=0.60 24 h: -1.02 (-1.60, 0.44) P=0.0007
Mean (SE) haematocrit, L/L			Difference/ratio (95% CI) After protamine: -0.001
	After protamine: 0.345 (0.006)	After protamine: 0.344 (0.006)	(-0.019, 0.017) P=0.91
	1 h: 0.312 (0.006)	1 h: 0.305 (0.006)	1 h: -0.007 (-0.024, 0.011) P=0.46
	24 h: 0.350 (0.006)	24 h: 0.319 (0.006)	24 h: -0.031 (-0.049, -0.013) P=0.0008
Mean (SE) platelet count,	1 h: 192.8 (0.028)	1 h: 189.7 (0.026)	
X109/L	24 h: 225.4 (0.027)	24h: 218.2 (0.026)	

Mean (SE) prothrombin ratio	After protamine: 1.27 (0.012)	After protamine: 1.27 (0.012)	
	1 h: 1.19 (0.012)	1 h: 1.19 (0.011)	
	24 h: 1.15 (0.012)	24 h: 1.15 (0.012)	
APTT ratio	After protamine: 1.17 (0.024)	After protamine: 1.14 (0.022)	
	1 h: 1.08 (0.022)	1 h: 1.13 (0.022)	
	24 h: 1.08 (0.022)	24 h: 1.11 (0.022)	
Fibrinogen concentration,	After protamine: 2.59 (0.036)	After protamine:2.68 (0.033)	
g/L	1 h: 2.21 (0.034)	1 h: 2.34 (0.033)	
	24 h: 4.92 (0.035)	24 h:5.04 (0.034)	
Clinical importance		Clinical relevance	

EXTERNAL VALIDITY

Generalisability

The patient population is limited to people undergoing CABG; however, the results are still somewhat generalisable to all elective non-emergency surgery with moderate blood loss.

Applicability

The study was conducted in the UK, however it is still applicable to the Australian context.

Comments

Niranjan G, Asimakopoulos G, Karagounis A, Cockerill G, Thompson M, and Chandrasekaran V. (2006) Effects of cell saver autologous blood transfusion on blood loss and homologous blood transfusion requirements in patients undergoing cardiac surgery on- versus off-cardiopulmonary bypass: a randomised trial. European Journal of Cardio-thoracic Surgery 30:271-277

Affiliation/Source of funds

Funding from the British Heart Foundation.

Study design	Level of evidence	Location/setting
RCT	I	UK, hospital

Intervention	Comparator
Intraoperative cell salvage, with Autotransfusion of washed, salvaged red blood cells at the conclusion of the procedure.	Control: All los
N=20 (on-pump CPB) and 20 (off-pump CPB)	N=20 (on-pun

Control: All lost blood from the skin incision to closure was suctioned with a high pressure sucker into a waste container. N=20 (on-pump CPB) and 20 (off-pump CPB)

Population characteristics

Patients undergoing first-time isolated CABG.

Length of follow-up	Outcomes measured
Patient followed up until discharge from hospital.	Change in haemoglobin levels, total amount of homologous blood transfusion (HBT), length of ICU stay, postoperative complications

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
The allocation was randomised and was concealed from those responsible for recruiting subjects.	The treatment groups had similar baseline characteristics.	The study was not blinded.	A transfusion protocol was utilised.	There was one death during the study in the on-CPD without cell saver group that was sudden on the predischarge day and attributed to an arrhythmia with no obvious cause of death found at postmortem. The patient was included in the postoperative analysis.

Overall quality assessment (descriptive)

Good

RESULTS			
Outcome	Intervention group	Comparator group	Statistical significance
Mean (SD) volume of homologous blood transfusion	On-CPB: 179 (214) Off-CPB: 141 (183) Combined: 159 (196)	On-CPB: 230 (240) Off-CPB: 595 (438) Combined: 413 (394)	Off-CPB was significantly higher than the other groups (P<0.005)
			Significantly lower for patients receiving cell saver (combined on-CPB and off-CPB) (p < 0.001)
Mean (SD) 24 h postoperative blood loss, mL	On-CPB: 842 (276) Off-CPB: 869 (286)	On-CPB: 1023 (291) Off-CPB: 903 (315)	P>0.05
Mean (SD) change in Hb levels from preoperative to postoperative day 1, g/dl	On-CPB: 4.95 (1.1) Off-CPB: 4.95 (1.5)	On-CPB: 4.4 (0.9) Off-CPB: 5.0 (1.6)	P>0.05
Morbidity: atrial fibrillation, n (%)	On-CPB: 7 (35) Off-CPB: 3 (25)	On-CPB: 5 (25) Off-CPB: 4 (20)	P>0.05
Morbidity: Pulmonary complications, n (%)	On-CPB: 4 (20) Off-CPB: 2 (10)	On-CPB: 3 (15) Off-CPB: 1 (5)	P>0.05
Morbidity: Renal complications, n (%)	On-CPB: 2 (10) Off-CPB: 1 (5)	On-CPB: 1 (5) Off-CPB: 0 (0)	P>0.05
Morbidity: CVA, n (%)	On-CPB: 0 (0) Off-CPB: 1 (5)	On-CPB: 1 (5) Off-CPB: 0 (0)	P>0.05
Mean (SD) length of hospital stay, d	On-CPB: 8.1 (2) Off-CPB: 7.2 (2.3)	On-CPB: 8.3 (3.1) Off-CPB: 7.4 (2.1)	P>0.05
Mean (SD) length of ICU stay, h	On-CPB: 22.1 (9.2) Off-CPB: 23 (8.4)	On-CPB: 23 (8.9) Off-CPB: 21.7 (5.8)	P>0.05
Prothrombin time			There was a significant rise on the first postoperative day from preoperative levels (P<0.0005) in all groups with no significant difference between groups. At the fifth postoperative day the PT was still elevated compared to preoperative levels in all groups with no difference between groups.
Partial thromboplastin time (ratio)			Showed a significant increase on the first postoperative day in all patient groups (P<0.001) with no significant difference between groups and was still significantly raised on the fifth postoperative day but again there was no difference between groups.

Clinical importance	Clinical relevance	
EXTERNAL VALIDITY		
Generalisability		
The study population is restricted to patients undergoing CABG; however, it is somewhat generalisable to other elective, non-emergency surgeries with moderate blood loss.		
Applicability		
The study was conducted in the UK and is likely to be applicable to the Australian context.		
Comments		

Abbreviations: CPB, cardiopulmonary bypass; CVA, cardiovascular accident

Citation								
Selo-Ojeme DO a ectopic pregnanc							transfusion for ruptured	
Affiliation/Source	e of funds							
None declared								
Study design	Level of evidence			e Location/setting			ing	
RCT	II					Nigeria / Unive	ersity hospital	
Intervention				Compa	rator			
Intraoperative cell salvage with transfusion of filtered autologous blood. N=56			of filtered	Control N=56	: allogeneic bl	ood transfusion	only.	
Population char	acteristics							
Women with a dia	agnosis of rup	otured ectop	ic pregnancy.					
Length of follow	-up			Outcor	nes measure	d		
Follow-up up until hospital discharge.				Patients transfused with ≥ 1000 mL of blood, haematocrit concentration, morbidity, length of hospital stay				
INTERNAL VALI	DITY							
Allocation	Results Blinding an		Blinding anal	ysis Treatment/measubias		measurement	Follow-up (ITT)	
It is not clear whether allocation was concealed from those responsible for recruiting subjects.			The study was blinded.			on protocol was	There was no loss to follow-up.	
Overall quality a	ssessment ((descriptive)				- 1	
Fair								
RESULTS								
Outcome		Interventi	on group	n group Com		ip S	tatistical significance	
Patients transfused with ≥ 34/56 (60° 1000 mL of blood		34/56 (60%	6) 1-		11/56 (20%)		RR (95% CI): 6.41 (2.75, 15.24)	
Mean postoperati haematocrit concentration, %	crit [*]			26%			<0.01	
Duration of surgery					lo	The duration of surgery was nger in the autologous roup, but not significantly."		

Morbidity			RR (95% CI)
	Postoperative fever: 20/56 (36%) Wound infection: 3/56 (5%)	Postoperative fever: 21/56 (38%) Wound infection: 4/56 (7%)	Postoperative fever: 0.95 (0.43, 2.01) Wound infection: 0.73 (0.17, 3.19)
Mortality			"There were no deaths from ectopic pregnancies during the study period."
Patients who had a length of hospital stay of more than 7 days.	8/56 (14%)	6/56 (11%)	RR (95% CI): 1.37 (0.44, 4.31)
Clinical importance		Clinical relevance	
EVEDNIAL VALIDITY	-		-

Generalisability

The study is of women diagnosed with ruptured ectopic pregnancy and may not be generalisable to men, or people undergoing other surgery types.

Applicability

The study was conducted in Nigeria, which may limit its applicability in high-resource countries such as Australia.

Comments

Wiefferink A, Weerwind PW, van Heerde W, Teerenstra S, Noyez L, de Pauw BE, and Brouwer RM. (2007) Autotransfusion management during and after cardiopulmonary bypass alters fibrin degradation and transfusion requirements. The Journal of extra-corporeal technology 39:66-70.

Affiliation/Source of funds

None declared

Study design	Level of evidence	Location/setting
RCT	II	The Netherlands / hospital

Intervention	Comparator
Intraoperative cell salvage: The mediastinal and residual CPB blood was processed by a continuous autotransfusion system before reinfusion. N=15	Control group: did not undergo cell salvage N=15

Population characteristics

Patients undergoing CABG with CPB

Length of follow-up	Outcomes measured
20 hours after arrival at ICU.	Number of patients transfused, length of operation

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Allocation was concealed.	The patient demographics were similar between the two groups.	The ICU staff were blinded to the group, however the surgical staff were not.	No transfusion protocol was used.	There was no loss to follow-up.

Overall quality assessment (descriptive)

Fair

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance
Patients who received at least one unit of packed blood cells postoperatively	8/15 (54%)	10/15 (67%)	P>0.05
Patients who received at least two units of packed blood cells postoperatively	2/15 (13%)	7/15 (47%)	P<0.05
Mean (SD) bypass time, min	98 (25)	86 (21)	P>0.05
Clinical importance		Clinical relevance	

EXTERNAL VALIDITY

Generalisability

The study population was people undergoing CPB; however, the study is likely to be somewhat generalisable for surgical procedures with moderate blood loss.

Applicability
The study is likely to be applicable to the Australian context.
Comments

Abbreviations: CABG, coronary artery bypass graft; CPB, cardiopulmonary bypass

Citation								
Zhang XL, Qian E patients with scol						g perioperative _l	period on prognosis of	
Affiliation/Source	e of funds							
None declared								
Study design		L	evel of evidence	е		Location/sett	ing	
RCT	II					China / hospit	al	
Intervention				Compa	rator			
Intraoperative cell salvage and retransfusion of washed autologous blood. N=36			on of washed	Control N=12	: allogeneic tra	ansfusion only		
Population char	acteristics							
Patients undergoi	ing operation	for scoliosis	i.					
Length of follow	-up			Outcor	nes measure	d		
				Perioperative bleeding, patients transfused with allogeneic blood, mortality, allergic reaction				
INTERNAL VALI	DITY							
Allocation	Results		Blinding anal	ysis Treatment/measuremen bias		measurement	Follow-up (ITT)	
It is unclear whether allocation was concealed from those responsible for recruiting subjects.		reatment arms imilar baseline icteristics The study was blinded.		not	not No transfusion protocol was reported.		There was no loss to follow-up.	
Overall quality a	ssessment ((descriptive)					
Poor								
RESULTS								
Outcome		Interventi	on group	Con	Comparator group		Statistical significance	
Perioperative bleeding						qı	There was no difference in uantity of bleeding between two groups."	
Patients transfuse allogeneic blood	ed with	11/36 (31%	11/36 (31%)		12/12(100%)		<0.01	
Allergic reaction		0/36 (0%)		3/12 (25%)		N	NR	
Mortality	Mortality		0/36 (0%)		0/12 (0%)			

Clinical importance	Clinical relevance					
EXTERNAL VALIDITY						
Generalisability						
The patient population were people undergoing surgery for scoliosis and may not be generalisable to other surgery types.						
Applicability						
Study was conducted in China, which may limit its applicability to the Australian context.						
Comments						

Intervention 3 – Perioperative acute normovolemic haemodilution combined with intraoperative cell salvage

Level II evidence

Citation								
Haynes SL, Torel clinical trial of hae							luation of a randomized 6.	
Affiliation/Sourc	e of funds							
Funded by the NF	HS Executive	North West	Research and D	evelopen	nent Directorat	e		
Study design	Level of evidence)		Location/settin	ng	
RCT and econom	ic analysis	II				UK / university h	nospital	
Intervention				Compa	rator			
ANH and intraope n=74	erative cell sa	lvage		Standar n=71	rd transfusion	practice (allogene	eic transfusion)	
Population chara	acteristics			•				
Patients undergoi	ng aortic sur	gery. 34 und	erwent aortobife	moral byp	ass and 111 ι	underwent aortic	aneurysm repair.	
Length of follow	-up			Outcon	nes measured	d		
			Operative blood loss, units of allogeneic blood transfused, operative time, length of hospital stay, length of ICU stay, morbidity, mortality, cost.					
INTERNAL VALI	DITY							
Allocation	Results	Blinding analy		ysis	sis Treatment/measurement Follobias		Follow-up (ITT)	
It is not clear whether allocation was concealed from those responsible for recruiting subjects.	compare th	tudy did not are the baseline acteristics of the rention arms. The study was blinded.			A transfusion used.	n protocol was		
Overall quality a	ssessment (descriptive)					
Fair			•					
RESULTS								
Outcome		Intervention	on group	Comparator group		p Sta	Statistical significance	
Median (IQR) ope time, min	erative	195 (162 to	o 238)			P =	0.86	
Median (range) length of ICU stay, days		1 (0		0 to 25) P=0).89		
Mean cost of treatment¹ £5384		£585	59	NS				
Clinical importance			Clin	ical relevance	9			
EXTERNAL VAL	IDITY			ı				

Generalisability

The study population is similar to the guideline population.

Applicability

The clinical outcomes are applicable to the Australian context, however the cost outcomes are not.

Comments

¹The majority of the total cost was due to intensive care and ward stays: transfusion accounted for only 6 and 7% of the total in control and ANH + cell salvage groups respectively.

Citation							
	ghnessy D, Pickering R, diac surgery: Randomis					educing blood	
Affiliation/Source	e of funds						
The study was su	pported by a grant from	the local blood tra	nsfusion	service.			
Study design		Level of evidence	9		Location/setting	g	
RCT		II			UK, hospital		
Intervention			Compa	rator			
ANH + intraopera	tive cell salvage (n=86)		1. 2.	•	ve cell salvage (n ical blood conserv	= 84) ration, control (n = 84)	
Population chara	acteristics						
Patients undergoi	ng cardiac surgery.						
Length of follow-	-up		Outcon	nes measure	d		
NR				Transfusion frequency and dose, perioperative complications, change in haemoglobin, length of hospital stay, time in intensive care, operative time			
INTERNAL VALID	OITY		l .				
Allocation	Results	Blinding anal	ysis	Treatment/r bias	measurement	Follow-up (ITT)	
Allocation was randomised and allocation was adequately concealed from those responsible for recruiting subjects.	The three treatment arms had similar baseline characteristics. Parsonnet scores, which reflect a number of patient variables and allow preoperative risk stratification, were also similar across the groups.	The patients, intensive care and trial asses were blinded to allocation. The surgical staff c be blinded. Note: because intensive care were blinded to allocation to grand no protocc violations occur can be assume the reduction in allogeneic red cell transfusion related to the experimental of the treatmental care and the reduction in the treatmental care and the treatmental care and the treatmental care and trial care and transfusion related to the experimental care and trial care	sors ould not the staff oup, oll ired, it ed that n blood n is	Not detected	d.	Two patients who were randomised could not be included because cell salvage or blood harvest machines were not available. Another two patients were excluded because of inappropriate perioperative transfusion. These four patients were not included in the analysis because of insufficient data. They were replaced with other patients in the trial who were randomised by the independent observer so that their next allocation was concealed.	
Overall quality as	ssessment (descriptiv	e)				1	
Fair	<u> </u>						

RESULTS				
Outcome	Intervention group	Comparator group	Statistical significance	
Number of patients transfused with any allogeneic blood product	33/86 (38%)	Cell salvage: 32/84 (38%) Control: 47/84 (56%)	Combined treatment vs cell salvage: OR (95% CI): 1.05 (0.56, 1.98) Wald test P-value = 0.872	
Number of patients transfused with allogeneic whole blood transfusion	29/86 (34%)	Cell salvage: 26/84 (31%) Control: 43/84 (51%)	Combined treatment vs cell salvage: OR (95% CI): 1.18 (0.62, 2.24) Wald test p value = 0.622	
Mean (SD) units of allogeneic blood transfused per patient ¹	0.63 (1.22)	Cell salvage: 0.68 (1.55) Control: 1.07 (1.56)	Kruskal-Wallis p value = 0.015	
Number of patients transfused with FFP	13/86 (15%)	Cell salvage: 14/84 (17%) Control: 13/84 (15%)	Combined treatment vs cell salvage: OR (95% CI): 0.91 (0.40, 2.11) Wald test P-value = 0.831	
Mean (SD) units of FFP transfused	0.43 (1.12)	Cell salvage: 0.57 (1.47) Control: 0.49 (1.25)	Kruskal-Wallis P-value = 0.952	
Number of patients transfused with platelets	15/86 (17%)	Cell salvage: 11/84 (13%) Control: 15/84 (18%)	Combined treatment vs cell salvage: OR (95% CI): 1.46 (0.62, 3.47): Wald test P-value: 0.386	
Mean (SD) units of platelets transfused	0.31 (0.81)	Cell salvage: 0.20 (0.62) Control: 0.29 (0.67)	Kruskal-Wallis P-value = 0.601	

Morbidity (perioperative complications)	No perioperative complications: 46/86 (53%) Inotropes required after 24 hours: 11/86 (13%) Surgical bleeding: 2/86 (2%) Cerebrovascular accident: 1/86 (1%) Arrhythmias: 20/86 (23%) Renal failure: 2/86 (2%) Proven infection: 7/86 (8%) Myocardial infarction: 4/86 (5%)	Cell salvage No perioperative complications: 46/84 (55%) Inotropes required after 24 hours: 12/84 (14%) Surgical bleeding: 2/84 (2%) Cerebrovascular accident: 1/84 (1%) Arrhythmias: 17/84 (20%) Renal failure: 1/84 (1%) Proven infection: 11/84 (13%) Myocardial infarction: 5/84 (6%) Control No perioperative complications: 42/84 (50%) Inotropes required after 24 hours: 9/84 (11%) Surgical bleeding: 3/84 (4%) Cerebrovascular accident: 2/84 (2%) Arrhythmias: 27/84 (32%) Renal failure: 0/84 (0%) Proven infection: 7/84 (8%) Myocardial infarction: 10/84 (12%)	NR
Median (IQR) haemoglobin concentration (g/l)	Before operation: 145 (138, 150) On admission to ICU: 108 (99, 116) Day 1 after operation: 105 (96, 113) Day 3 after operation: 108 (100, 119)	Cell salvage Before operation: 145 (136, 150) On admission to ICU: 105 (98, 116) Day 1 after operation: 104 (95, 115) Day 3 after operation: 105 (98, 115) Control Before operation: 142 (135, 150) On admission to ICU: 100 (91, 107) Day 1 after operation: 100 (94, 109) Day 3 after operation: 106 (98, 112)	NR
Median (IQR) length of hospital stay (h)	170.4 (147.1, 221.6)	cell salvage: 160.7 (145.5, 198.8) control: 168.9 (140.3, 219.3)	Kruskal-Wallis p-value: 0.724

Median (IQR) time in ICU (h)	23.3 (22.5, 25.0)	Cell salvage: 22.7 (22.0, 24.6)	Kruskal-Wallis p-value: 0.249
Mean (IQR) operative time (minutes)	154 (131, 174)	control: 22.9 (21.8, 24.5) Cell salvage: 160 (140, 184)	NR
Clinical importance		control: 160 (135, 196) Clinical relevance	

Generalisability

This trial was conducted on a specific patient population (people undergoing cardiac surgery), however it is likely to be generalisable to patients undergoing other elective surgical procedures with moderate blood loss.

Applicability

The study was performed in UK and is likely to be applicable to the Australian context

Comments

The combination of acute perioperative normovolaemic haemodilution and intraoperative cell salvage did not show any additional benefit over intraoperative cell salvage alone (?: check)

¹Nine patients needed a markedly higher amount of transfused blood (≥ 3 units). These patients were returned to the operating theatre for re-exploration of the mediastinum. A surgical cause of bleeding was found in seven of these patients (three in the control group and two each in the cell salvage and combined treatment groups).

Wong JC, Torella F, Haynes SL, Dalrymple K, Mortimer AJ, McCollum CN, and ATIS I. (2002) Autologous versus allogeneic transfusion in aortic surgery: a multicenter randomized clinical trial. Annals of surgery 235:145-151.

Affiliation/Source of funds

Funded by the NHS Executive Research and Development

Study design	Level of evidence	Location/setting
RCT	II	UK / hospital

Intervention	Comparator
ANH and ICS: Before skin incision, sufficient blood was taken to reduce the haemoglobin concentration to 11 g/dL. Blood volume was replaced simultaneously with crystalloids, maintaining a steady central venous pressure during blood collection. ANH blood, containing fresh platelets and clotting factors, was retained for reinfusion at wound closure when haemostasis was secure. Blood lost during the procedure was salvaged using one of three centrifugal cell salvage devices with comparable efficacy in red cell recovery under standard conditions. All autologous blood was reinfused within 6 hours of withdrawal.	Allogeneic transfusion: Patients did not receive ANH or ICS. Patients received allogeneic blood transfusion when required. N=71

Population characteristics

Patients undergoing aortic surgery, including 111 (59 ANH + cell salvage; 52 control) patients with aneurysms and 34 (15 ANH + cell salvage; 19 control) with occlusive disease.

Length of follow-up	Outcomes measured
Patients were assessed at wound closure, 2 hours, 1, 2, and 7 days after surgery.	Clinical signs, complications, transfusion requirements, and laboratory assay results.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Subjects were randomised; however, it is unclear whether allocation was censored for those responsible for recruiting subjects.	The baseline characteristics were mostly similar between the treatment arms. However, "allogeneic" patients had slightly higher mean preoperative haemoglobin concentrations (14.03 vs 13.57 g/dL; P=0.053) and ANH+cell salvage patients were slightly older (72 vs 69 years; P=0.04).	The trial was single- blind. However, the decision to give allogeneic transfusion was made by a rigid protocol and was made by a physician independent from the research team.	A transfusion protocol was used. Members of the research team attended all operations and recorded all data independently from the clinical team.	All analyses conducted ITT.

Overall quality assessment	(descriptive)		
RESULTS			
Outcome	Intervention group	Comparator group	Statistical significance
Mean (SD) volume of blood withdrawn during ANH, units (450 mL)	1.66 (0.71)	The process of the pr	
Median (IQR) blood loss during surgery, mL	921 (661 to 1374)	1000 (688 to 1734)	P=0.37
Median (IQR) packed red cell volume reinfused after cell salvage, mL	415 (225 to 543) ¹	-	-
Patients requiring transfusion of allogeneic whole blood during surgery	32/74 (43%)	40/71 (56%)	P=0.12
Median (IQR) units of allogeneic blood transfused (for all patients) during surgery	0 (0 to 2)	2 (0 to 4)	P=0.008
Total units of allogeneic blood transfused	117	251	
Patients requiring transfusion of allogeneic whole blood within 24 hours of surgery	19/74 (26%)	50/71 (70%)	P<0.001
Median (IQR) units of allogeneic blood transfused (for all aneurysm patients) during surgery)	0 (0 to 2)	2 (0 to 4)	P=0.002
Total units of allogeneic blood transfused (aneurysm patients)	102	201	
Median (IQR) units of allogeneic blood transfused (for all occlusive disease patients) during surgery)	0 (0 to 2)	0 (0 to 2)	0.87
Total units of allogeneic blood transfused (occlusive disease patients)	15	50	
Mortality	13/74 (18%)	11/71 (15%)	P=0.91
Morbidity: postoperative infection	16/74 (22%)	19/71 (27%)	P=0.6
Morbidity: transfusion reaction (minor)	0/0 (0%)	1/71 (1%)	NR
Morbidity: cardiac events	13/74 (18%)	8/71 (11%)	P=0.4
Morbidity: haemorrhagic complications	5/74 (7%) ²	8/71 (11%)³	NR
Reoperation	10/74 (14%)4	7/71 (10%)5	NR

Median (IQR) hospital stay, days	10 (8 to 13)	9 (7 to 12)	P=0.17	
Clinical importance		Clinical relevance		
EXTERNAL VALIDITY				
Generalisability				
The patient population (adults	undergoing cardiac surgery) is s	similar to the guideline population	٦.	
Applicability				
The study was conducted in UK, and is applicable to the Australian context.				
Comments				

¹Equivalent to more than one unit of allogeneic blood because the haematocrit of reinfused red cells was approximately 65%.

Abbreviations: ICS, intraoperative cell salvage

²Two of the patients required a laparotomy (one for massive bleeding from the proximal aortic anastomosis, one for upper gastrointestinal haemorrhage).

³Three patients had intraoperative bleeding and a further five required reoperation for intra-abdominal bleeding.

⁴Five thromboembolectomies, two laparotomies for haemorrhage, two laparotomies for bowel obstruction, one groin resuturing.

 $^{{}^5\}text{Five}$ required reoperation for intra-abdominal bleeding, and two thromboembolectomies.

Intervention 4 – Postoperative cell salvage

Level I evidence

Citation					
Carless P, Moxey A, O'Connell D, and Henry D. (2004) Autologous transfusion techniques: A systematic review of their efficacy. Transfusion Medicine 14:123-144.					
Affiliation/Source of funds					
The research was supported by a grar special purpose grant from the Hunter				dical Research Co	uncil of Australia and a
Study design	Level of evidence	е		Location/setting]
Systematic review of RCTs and observational studies with meta-analysis	I			NA	
Search conducted July 2002					
Intervention		Compa	rator		
Autologous transfusion techniques: preoperative Autologous blood deposit (PAD), ANH, and cell salvage. NOTE: This form only contains RCT info relevant for ANH. Comparator: no Autologous transfusion technique (active versus active comparisons were excluded). Sample size n=591					
Sample size n=704					
Population characteristics					
Patients older than 18 years undergoin trials included more than twice as man orthopaedic surgery, and 11 involved was a surgery.	y males as females	(2.3:1). Tv	velve trials inv	olved cardiac surg	ery, seven involved
Length of follow-up		Outcon	nes measure	d	
NA		thrombo	osis, non-fatal	n, infection, wound MI, rate of allogen me of allogeneic bl	eic red blood cell
INTERNAL VALIDITY					
Allocation Results	Blinding anal	ysis	Treatment/r bias	measurement	Follow-up (ITT)
Allocation concealment and the method of randomisation were judged by the SR authors to be inadequate in 100 and 92% of trials respectively (kappa=0.78-1.0). SR did not discuss similarity between preoperative data and baseline characteristics for the intervention groups. The majority (96%) of the included RCTs assessing ANH were unblended. NR Not detected NR Not detected NR Not detected NR Overall quality assessment (descriptive)			NR		
Overall quality assessment (descrip	tive)				

Fair			
RESULTS			
Outcome	Intervention group	Comparator group	Statistical significance
Mortality 8 trials (n=NR)	NR	NR	RR (95% CI): 1.16 (0.19, 7.15) Phet=NR
Morbidity: infection 2 trials (n=NR)	NR	NR	RR (95% CI): 4.94 (0.61, 40.19) Phet=NR
Morbidity: thrombosis 3 trials (n=NR)	NR	NR	RR (95% CI): 0.44 (0.21, 0.93) Phet=NR
Morbidity: non-fatal MI 3 trials (n=NR)	NR	NR	RR (95% CI): 3.43 (0.15, 79.74) Phet=NR
Re-operation 7 trials (n=NR)	NR	NR	RR (95% CI):1.59 (0.20, 12.53) Phet=NR
Rate of allogeneic blood transfusion: all studies 25 trials (n=1081; 567 ANH, 514 control)	273/567 (48%)	357/514 (69%)	RR (95% CI): (Phet<0.00001)
Rate of allogeneic blood transfusion: cardiac surgery 10 trials (n=NR)	NR	NR	RR (95% CI): 0.77 (0.57, 1.04)
Rate of allogeneic blood transfusion: orthopaedic surgery 6 trials (n=NR)	NR	NR	RR (95% CI): 0.79 (0.60, 1.06)
Rate of allogeneic blood transfusion: miscellaneous surgery 9 trials (n=NR)	NR	NR	RR (95% CI): 0.42 (0.24, 0.74)
Rate of allogeneic blood transfusion: transfusion protocol used 16 trials (n=NR)	NR	NR	RR (95% CI): 0.81 (0.62, 1.00)
Rate of allogeneic blood transfusion: transfusion protocol not used/reported 9 trials (n=NR)	NR	NR	RR (95% CI): 0.53 (0.36, 0.76)
Difference in units of allogeneic blood transfused 17 trials	NR	NR	WMD (95% CI) Overall: -1.9 (-1.1, -2.7) Studies with a transfusion protocol: -1.0 (-1.7, -0.4) Studies without a transfusio protocol: -3.0 (-4.9, -1.1)

Hospital length of stay, d 3 trials (N=96)	NR	NR	WMD (95% CI): 0.21 (-1.26, 1.68)	
Clinical importance	nical importance Clinical relevance			
EXTERNAL VALIDITY				
Generalisability	Generalisability			
Patients considered similar to	guideline target population			
Applicability				
All the studies included in this review were conducted in countries with well developed healthcare systems (not specifically Aus/NZ).				
Comments				

Abbreviations: ANH, acute normovolemic haemodilution; MI, myocardial infarction; PAD, preoperative autologous donation; NR, not reported; RCT, randomised controlled trial; RR, risk ratio; WMD, weighted mean difference.

Carless PA, Henry DA, Moxey AJ, O'connell DL, Brown T, and Fergusson DA. (2006) Cell salvage for minimising perioperative allogeneic blood transfusion. Cochrane database of reviews; Issue 4. Affiliation/Source of funds None declared Study design Level of evidence Location/setting SR Search conducted Jan 2004

Intervention	Comparator	·
Cell salvage. Studies with a combination of comparisons were included if both the intercontrol groups were equally exposed to the treatment (ie, active plus cell salvage versus comparisons). N (perioperative cell salvage) = 1952 The authors found 51 studies.	vention and N=1905 active	

Population characteristics

Adults (over 18 years) undergoing elective, non-urgent surgery. Surgery types found in the search include cardiac (23 studies), orthopaedic (23 studies), and vascular (5 studies) surgery. 33 of the trials studied cell salvage during the postoperative period, 10 studied intraoperative cell salvage, and seven studied both intraoperative and postoperative cell salvage. One trial failed to describe the timing of cell salvage. Twenty trials studied cell salvage systems that reinfused washed salvaged blood, and 29 trials studied cell salvage systems that reinfused unwashed filtered salvaged blood. One trial studied both washed and unwashed cell salvage (4-arm trial) and provided two comparisons of cell salvage.

Length of follow-up	Outcomes measured	
NA	Number of patients transfused with allogeneic and/or autologous blood, amounts of allogeneic and/or autologous blood transfused, re-operation for bleeding, postoperative complications, mortality, and length of hospital stay.	

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
According to the authors, the description of the method to conceal allocation of either inadequate or unclear for all of the studies.		Based on the Schulz criteria, blinding was reported in only one of the trials.	Nine of the 51 studies did not report a transfusion protocol.	

Overall quality assessment (descriptive)

Good

RESULTS					
Outcome	Intervention group	Comparator group	Statistical significance		
Rate of allogeneic transfusion (postoperative cell salvage) 16 trials (n=NR)			RR (95% CI): 0.58 (0.43, 0.79) P <i>het</i> =NR		
Clinical importance		Clinical relevance	Clinical relevance		
EXTERNAL VALIDITY					
Generalisability					
The SR is generalisable for e	elective, non urgent surgery.				
Applicability					
The studies were mostly from countries with similar health-care systems to Australia					
Comments					
The systematic review includes trials assessing the intraoperative, postoperative, and both intra- and postoperative. There is more data but most of it combines intra- and postoperative data.					

Abbreviations: RR, relative risk; ARR, absolute risk reduction; RRR, relative risk reduction

Davies L, Brown TJ, Haynes S, Payne K, Elliott RA, and McCollum C. (2006) Cost-effectiveness of cell salvage and alternative methods of minimising perioperative allogeneic blood transfusion: A systematic review and economic model. Health Technology Assessment 10:1-114.

Affiliation/Source of funds

One author received sponsorship from haemonetics and AstraTech to attend the International Society of Blood Transfusion (ISBT) VIIIth European Congress. The author has also given invited lectures for AstraTech Ltd and Unomedical with honoraria and expenses paid.

Study design	Level of evidence	Location/setting
SR with economic analysis (SR search is an update of Carless 2003: a Cochrane review that had been updated in 2006. The study includes a meta-analysis combining the results of Carless 2003 and the search update.		NA
Search conducted Jan 2004		

Intervention	Comparator
Transfusion strategies to minimise perioperative allogeneic blood transfusion: cell salvage, PAD, PAD plus EPO, EPO, ANH, cell salvage plus ANH, AFs, FSs, restrictive transfusion thresholds or protocols.	No cell salvage or allogeneic blood.
NB: This form only includes information relevant for postoperative cell salvage.	
Specific characteristics of the 1 included RCT (Naumenko 2003)	
Drainage discharge collected for 8 hours postoperatively and reinfused. Erythrocytes reinfused postoperatively after washing.	
NB: the authors of the SR were not able obtain a full version of Naumenko 2003, and therefore had to rely on information provided in the abstract.	

Population characteristics

For inclusion, the SRs had to only include adults undergoing elective, non-urgent surgery.

Specific characteristics of the 1 included RCT (Naumenko 2003)

Patients undergoing CABG surgery

Length of follow-up	Outcomes measured	
NA	Proportion/number of patients transfused with allogeneic and/or autologous blood; the volume of allogeneic and/or autologous blood transfused; reoperation for bleeding; adverse transfusion reactions; preoperative morbidity and Hb levels; postoperative complications; length of hospital stay; mortality.	

INTERNAL VALIDITY ¹				
Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Method of randomisation not described and allocation concealment unclear.	Naumenko 2003 had unclear baseline comparability: 'no significant difference between groups was detected at any stage of the study'	Did not have participant blinding and it is unclear whether study had allocation concealment.	The abstract of Naumenko 2003 does not describe the use of a transfusion protocol.	Unclear intention to treat (however there was no loss to follow-up).

Overall quality assessment (descriptive)

Good

RESULTS

REGOETO			
Outcome Intervention group		Comparator group	Statistical significance
Number of patients transfused with allogeneic blood (postoperative cell salvage; active versus control ²) 287/738 (39%) 287/738 (39%)		473/724 (65%)	RR (95% CI): 0.60 (0.45, 0.79) P<0.05
Clinical importance		Clinical relevance	

EXTERNAL VALIDITY

Generalisability

The review includes all surgery types and performs subgroup analyses by surgery type. Therefore the results are likely to be generalisable for other elective, non-emergency operations.

Applicability

Transfusion frequency outcome: Good applicability

Other outcomes - poor applicability to the question - With the exception of transfusion frequency, all of the outcomes assessed combined data from intra- and postoperative cell salvage.

Comments

The updated lit search included 2 RCTs (1 as abstract only): Naumenko 2003 and Zhao 2003. Zhao 2003 investigated intraoperative cell salvage (described in this form), and Naumenko 2003 investigated postoperative cell salvage (described in 14).

The SR includes a meta-analysis using data from the above two trials, as well as the data from Carless 2003 (a Cochrane review that had been updated in 2006. Except for transfusion frequency, all of the outcomes meta-analysed combined trials using intra- and postoperative cell salvage.

¹Refers only to the one intraoperative cell salvage RCT included in the systematic update (Zhao 2003)

Citation Duffy G and Neal KR. (1996) Differences in postoperative infection rates between patients receiving autologous and allogeneic blood transfusion: a meta-analysis of published randomized and nonrandomized studies (Structured abstract). Transfusion Medicine 6:325-328. Affiliation/Source of funds None declared Study design Level of evidence Location/setting SR of RCTs and retrospective studies NA Search date NR Intervention Comparator Autologous transfusion (including PAD or cell salvage) Allogeneic blood transfusion only Population characteristics Length of follow-up Outcomes measured Patients undergoing any surgical operation. Infections. INTERNAL VALIDITY Allocation Results Blinding analysis Treatment/measurement Follow-up (ITT) bias NR Baseline NR SR did not define whether a NR characteristics NR transfusion protocol was used Overall quality assessment (descriptive) Fair **RESULTS** Outcome Intervention group Comparator group Statistical significance OR (95% CI): 3.2 (0.4, 29.0) Infection (1 trial) 1/35 (3%) 3/35 (9%) Clinical importance Clinical relevance **EXTERNAL VALIDITY** Generalisability Limited generalisability (due to uncertainty regarding the small sample size, and lack of info regarding potential sources of bias). **Applicability** The SR does not report the location of the Newman 1995 trial. Comments NB: only one postoperative cell salvage RCT included (Newman 1995)

Citation Huet C, Salmi R, Fergusson D, Koopman-Van Gemert AWMM, Rubens F, and Laupacis A. (1999) A meta-analysis of the effectiveness of cell salvage to minimize perioperative allogeneic blood transfusion in cardiac and orthopedic surgery. Anesthesia and Analgesia 89:861-869. Affiliation/Source of funds Coordinating Centre has been funded by Janssen Ortho Inc, Canada. One of the authors is the recipient of the First Fellowship from the International Society of Technology Assessment in Health Care, funded by the PPP Medical Trust. Study design Level of evidence Location/setting SR of RCTS with Meta-analysis NA Search conducted December 1997 Intervention Comparator Perioperative cell salvage Population characteristics Patients who underwent cardiac or orthopaedic surgery (two articles dealing with vascular surgery were not considered). Length of follow-up Outcomes measured NA Proportion of patients receiving at least one unit of allogeneic packed red blood cells INTERNAL VALIDITY Allocation Results Blinding analysis Treatment/measurement Follow-up (ITT) bias Three of the Unclear: "all of the 28 The SR did not report on NR included trials trials included in this whether or not a transfusion were pseudostudy scored between protocol was used in the randomised. zero and three on the **RCTs** Jadad scale. Because it is difficult to "blind" the operative team to the presence or absence of cell salvage, the Jadad score would rarely be expected to be greater than 3" Overall quality assessment (descriptive) Fair **RESULTS** Outcome Intervention group Comparator group Statistical significance RR: 0.84 (95% CI: 0.77, Patients transfused with allogeneic blood (active 0.93)versus: CABG surgery using P<0.05 (Phet=NR) washed salvage (active vs

control analysis) 6 trials (n=482; 246 cell salvage, 236 control) Clinical importance Clinical relevance

EXTERNAL VALIDITY

Generalisability

The review includes all surgery types and performs subgroup analyses by surgery type. Therefore the results are likely to be generalisable for other elective, non-emergency operations.

Applicability

Low applicability to the question - all of the outcomes assessed combined data from intra- and postoperative cell salvage.

Comments

27 references, representing 28 RCTs, were included in the meta-analysis.

NB: The authors did not separately analyse intra- and postoperative cell salvage; however, all of the cardiac surgery trials used postoperative cell salvage. See section I2 for the perioperative cell salvage values for orthopaedic surgery, and other outcomes not assessed in a specific postoperative population.

¹ Includes both studies where an active treatment is compared with a control intervention and those studies where both the intervention and control arms also received an additional active intervention (c.f. active versus control, where the controls are untreated).

Level II evidence

Level II evid	ence							
Citation								
Amin A, Watson autologous retrai								mised controlled trial of
Affiliation/Source	ce of funds							
None declared								
Study design		L	evel of evidence	9		Location/s	ettinç]
RCT		11				UK / Hospi	tal	
Intervention		'		Compa	rator			
Postoperative cell salvage: The tourniquet was released before closure and one deep drain was inserted within the joint space. The drain was connected either to the Bellovac ABT Autotransfusion system. The shed blood was returned to the patient after collecting up to 500 mL and no later than six hours after surgery. A maximum of 1200 mL was retransfused. N=92		erted within either to the e shed blood up to 500 mL	Control: The drain was connected to a standard vacuum chamber, and the collected blood was discarded. N=86					
Population char	acteristics							
Patients undergo knee.	ing unilateral	TKA. All pat	ients aged over 5	55 years v	with osteoarthr	itis and/or in	ıflamn	natory arthritis of the
Length of follow	/-up			Outcor	nes measured	t		
NR				Transfusion (%), transfusion (vol), haemoglobin concentration, morbidity, length of hospital stay				
INTERNAL VAL	IDITY							
Allocation	Results		Blinding analy	ysis	Treatment/r bias	measureme	nt	Follow-up (ITT)
Allocation was concealed	groups had	and control I similar nographics.	Not blinded A transfus used.		A transfusion used.	n protocol wa	as	There were eight patients who were not retransfused. ¹ These patients were included in the study based on an 'intention to treat' principle.
Overall quality a	assessment ((descriptive)					
Fair								
RESULTS								
Outcome		Intervention group		Comparator group		p	Stat	istical significance
Mean (SD) chang haemoglobin, g/o	ĬL	2.2 (0.7)		2.6 (0.8)			P=0.	354
Patients transfus allogeneic blood		12/92 (13%	3%) 13		6 (15%)		P=0.	439
Total units of bloom	bd	22		26	26		NR	

638 (86 to 1470)

659 (100 to 1900)

Median (IQR) drainage

volume, mL

P=0.468

Median (IQR) of autologous blood retransferred, mL	481 (200 to 1110)	NA	NA		
Morbidity	Morbidity Wound infection: 3/92 (3%) DVT: 1/92 (1%)		NS		
	Persistent wound drainage (no infection): 2/92 (2%)	Persistent wound drainage (no infection): 1/86 (1%)			
	Other infections: 2/92 (2%)	Other infections: 2/86 (2%)			
	Returns to operating theatre: 1/92 (1%)	Returns to operating theatre: 0/86 (0%)			
Median (IQR) length of hospital stay, days	6.6 (3 to 14)	7.0 (3 to 16)	P=0.54		
Clinical importance		Clinical relevance			
EXTERNAL VALIDITY		l			
Generalisability					
The study population is genera	alisable for elective surgery with	moderate blood loss			
Applicability					
Study is applicable to the Australian context.					
Comments	Comments				

¹Five patients were not retransfused because of low drainage volumes (< 100 mL) and three patients who were not retransfused because of technical difficulties such as problems with the tubing and filter system.

Cheng SC, Hung TS, and Tse PY. (2005) Investigation of the use of drained blood reinfusion after total knee arthroplasty: a prospective randomised controlled study. Journal of orthopaedic surgery (Hong Kong) 13:120-124.

Affiliation/Source of funds

Sponsored by the Tung Wah Group of Hospitals Research Fund.

Study design	Level of evidence	Location/setting	
RCT	II	Hong Kong / Hospital	

		· · ·
Intervention	Comparator	
Postoperative cell salvage: patients in the group had their blood reinfused from dra of surgery. N=26	Control: Shed blood v N=34	vas not reinfused.

Population characteristics

Patients undergoing knee arthroplasty

Length of follow-up	Outcomes measured
3 days post-surgery	Patients requiring allogeneic transfusion, units of allogeneic transfusion, mean haemoglobin level, total operative blood loss, reinfusion volume, febrile complications.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Subjects were randomised into intervention groups. Allocation was concealed from those responsible for recruiting subjects.	The control group had a larger proportion of patients with a premorbid condition (65% vs 54%) and a larger proportion of males compared with the reinfusion group (35% vs 23%). However, these differences were not significant.	Near the end of each operation, the corresponding envelope for each patient was opened, and the surgeon was informed at the time of drain insertion to achieve a single-blind effect.	Transfusion protocol implemented.	There was no loss to follow-up.

Overall quality assessment (descriptive)

Fair

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance
Median (IQR) total operative blood loss, mL	273 (100 to 600)	280 (100 to 800)	P=0.84
Median (IQR) reinfusion volume, mL	425.2 (180 to 620)	NA	NA
Patients requiring allogeneic transfusion	4/26 (15%)	13/34 (38%)	P=0.05

Median (IQR) units of allogeneic transfusion	0.15 (0 to 1)	0.46 (0 to 4)	P=0.033
Median haemoglobin level (IQR), g/L	Immediately postoperative: 101 (84 to 128) Day 3: 98 (77 to 130)	Immediately postoperative: 104 (87 to 137) Day 3: 101 (77 to 130)	P-value Immediately postoperative: 0.332 Day 3: 0.401
Febrile complications	2/26 (8%)	1/34 (3%)	P=0.403
Clinical importance		Clinical relevance	

Generalisability

The patients in the study were specifically undergoing total knee arthroplasty; however the study is generalisable to other elective, non-emergency surgery types associated with moderate blood loss.

Applicability

The study was conducted in Hong Kong; however, it is likely to be applicable to the Australian context.

Comments

Citation								
Zacharopoulos A, prospective rando						reinfusion afte	er tot	al knee replacement. A
Affiliation/Source	e of funds							
None declared								
Study design		Le	evel of evidence	9		Location/se	tting	
RCT		II				Greece, hosp	pital	
Intervention				Compa	rator			
Postoperative cell blood within 6 hou			f washed	standar		d unit was give nage system w		raoperatively, and a sed.
N=30				N=30				
Population chara								
		replacemen	t. There were 47	ı	•		mear	age of 69.7 years.
Length of follow	•				nes measure			
15 days postoperative			Intra- and postoperative blood loss, volume of unwashed blood salvage returned, number of autologous blood transfusions, perioperative haemoglobin values, operation time, length of hospital stay.					
INTERNAL VALI	DITY							
Allocation	Results		Blinding analy	ysis	Treatment/r bias	neasurement	t	Follow-up (ITT)
Allocation concealment was not reported	Patient den were not re		The study was not blinded.		A transfusion protocol was not used.		5	It is unclear whether all patients randomised were included in the analyses.
Overall quality as	ssessment ((descriptive)						
Poor								
RESULTS								
Outcome		Intervention	on group	Con	parator grou	р :	Stati	stical significance
"Average" (range of blood reinfused		808 (300 to 1750)						
Mean volume of homologous blood transfused, units	d	0.3).3				NR	
Patients requiring postoperative hon blood transfusion		5/30 (16.69) (16.6%)		0 (33.3%)		NR	
Difference in haer and haematocrit	moglobin						NS	

Clinical importance	Clinical relevance		
EXTERNAL VALIDITY			
Generalisability			
The patients were restricted to those undergoing total knee replacement; however, the study is generalisable for elective, non-emergency surgery associated with moderate blood loss.			
Applicability			
The study was conducted in a hospital situated in a small town of 8000 inhabitants. This may limit the applicability to larger hospitals.			
Comments			

¹The authors report some results as "average" without explicitly stating whether they are referring to mean or median.

Intervention 5 - Deliberate induced hypotension

Level I evidence

Citation

Paul JE, Ling E, Lalonde C, Thabane L. Deliberate hypotension in orthopedic surgery reduces blood loss and transfusion requirements: A meta-analysis of randomized controlled trials. Canadian Journal of Anaesthesia 2007;54(10):799-810.

Affiliation/Source of funds

 $Department\ of\ An esthesia,\ McMaster\ University,\ Hamilton,\ Ontario,\ Canada.$

Funding: Hamilton Health Sciences, Department of Anesthesia Academic Fund.

Study design	Level of evidence	Location/setting
Systematic review including 17 RCTs that investigated the effects of deliberate hypotension on blood loss and transfusion requirements in patients undergoing orthopaedic surgery.	Level I	Hospital

Intervention	Comparator
Deliberate hypotension by any method	No deliberate hypotension

Population characteristics

Patients undergoing orthopaedic surgery

Length of follow-up	Outcomes measured
NR	Intraoperative blood loss, transfusion requirements, duration of surgery

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomised allocation was reported in all studies. Method of randomisation not reported.	Baseline characteristics of intervention and control groups not reported. A random effects model was use in the mateanalyses as the test of heterogeneity was significant (p<0.05).	One was a double blinded study. Six were single blinded studies. Ten were open- labelled studies.	Egger's test for bias was non- significant (p=0.955), suggesting that there was no publication bias. Data extraction for each study was performed independently by the three authors. Consensus between reviewers was considered good with a kappa score of 0.87.	No lost to follow-up in all studies

Overall quality assessment (descriptive)

Good. This review clearly defined the research question, scope, search terms and inclusion/exclusion criteria. The search strategy employed appeared robust, and the methods for analysis were appropriate. The summary, as well as a quality rating, for each included study was provided. This review provided pooled data, for each of the specified outcomes, through meta-analysis of the data from included studies.

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance
Blood loss	NR	NR	-286 mL (95%CI : -447, -127)
Transfusion requirements	NR	NR	-667 mL (95%CI : -963, -370)

Surgery duration	NR	NR		-1.9 min (95%CI : -7.2, 3.5)
Outcome	Clinical importance		Clinical relevance	
Blood loss	1: reduced blood loss		1	
Transfusion requirements	1: reduced requirements		1	
Surgery duration	4		2	

Generalisability

This systematic review focus on patients undergoing orthopaedic surgery, which may not share clinical characteristics with the general surgical patient population.

Applicability

The studies in this review were mostly conducted in developed countries (mostly European), comparable to Australia. The surgeries performed (and the possible benefits) are likely applicable in Australia.

Comments

This review suggests that induced hypotension may decrease blood loss, transfusion requirement and surgery duration.

Abbreviations: CI, confidence intervals; NR, not reported; RCTs, randomised controlled trials.

675

Level II evidence

Citation

Boldt J, Weber A, Mailer K, Papsdorf M, Schuster P. Acute normovolaemic haemodilution vs controlled hypotension for reducing the use of allogeneic blood in patients undergoing radical prostatectomy. British Journal of Anaesthesia1999;82(2):170-174.

Affiliation/Source of funds

Department of Anaesthesiology and Intensive Care Medicine and the Clinic of Urology, Klinikum der Stadt Ludwigshafen, Bremserstr, Germany.

Funding: NR

Study design	Level of evidence	Location/setting
RCT	Level II	Hospital

Intervention		Comparator	
Controlled hypotension (MAP: 50mm Henitroprusside	g) using sodium	No induced hypotens	ion
Controlled hypotension with haemodilut scope of Q3:Intervention 5)	ion (beyond the		

Population characteristics

40 patients, under the age of 75 years undergoing retropubic radical prostatectomy with bilateral pelvic lymphadenectomy. Exclusion criteria: ASA class IV, myocardial infarction within 6 months, documented coronary artery disease or carotid artery stenosis, abnormal coagulation, liver dysfunction, medication with aspirin, renal insufficiency.

Length of follow-up	Outcomes measured
Up to the first postoperative day in the ward	Blood loss, transfusion dose and frequency, coagulation status

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomisation using blinded envelopes	No significant differences in patient clinical characteristics were observed between the intervention and control group	Blinded envelopes used. Postoperative administration of blood/blood products provided by an independent physician blinded to patient's study group.	All patients operated on by one of two surgical teams. Postoperative administration of blood/blood products provided by an independent physician blinded to patient's study group.	No death or lost to follow-up

Overall quality assessment (descriptive)

Good. This RCT provides clear description for the research question and methods. The analyses performed were appropriate and the results presented clearly.

RESULTS			
Outcome	Intervention group	Comparator group	Statistical significance
Blood loss (mL)	1260 (SD: 570)	1920 (SD: 590)	P<0.05
Transfusion frequency	25%	60%	P<0.05
Total transfusion dose in group (units)	14	28	P<0.05
Coagulation status	NR	NR	No significant difference
Cost per patient (blood, volume replacement, intervention)	US\$82.81	US\$139.99	P<0.05

Outcome	Clinical importance	Clinical relevance
Blood loss	1: reduced blood loss	1
Transfusion frequency	1: reduced frequency	1
Transfusion dose	1: reduced requirements	1
Coagulation status	4	1
Cost per patient	NA	NA

Generalisability

This RCT was conducted on a specific patient population (males with ASA status I-III, undergoing retropubic radical prostatectomy with bilateral pelvic lymphadenectomy, in addition to several other exclusion criteria). Consequently, the findings of this study may not be applicable to other surgical patient populations.

Applicability

This trial was conducted in Germany, with a developed healthcare system comparable to that in Australia. As such the findings of the study are applicable.

Comments

This study shows that induced hypotension reduces blood loss, costs, transfusion dose and frequency.

Abbreviations: ASA, American Society of Anaesthesiologists physical status; MAP, mean arterial pressure; NA, not applicable; NR, not reported; RCT, randomised clinical trial; SD, standard deviation.

Elsharnouby NM, Elsharnouby MM. Magnesium sulphate as a technique of hypotensive anaesthesia. British Journal of Anaesthesia 2006;96(6):727-731.

Affiliation/Source of funds

Faculty of Medicine, Ain-shams University, Cairo, Egypt.

Funding: NR

Study design	Level of evidence	Location/setting
Double-blinded RCT	Level II	Hospital

Intervention	Comparator	
Magnesium sulphate induced hypotension	No induced hypotensi	on

Population characteristics

60 patients undergoing functional endoscopic sinuses surgery.

Length of follow-up	Outcomes measured
NR, includes postoperative monitoring	Blood loss, surgery duration

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Randomised, computerized random allocation.	No significant differences in patient clinical characteristics were observed between the intervention and control group	Double –blinded. Anaesthetist was unaware of the patients' treatment group.	Intraoperative bleeding was evaluated by the same surgeon every 15 minutes	No subjects excluded

Overall quality assessment (descriptive)

Good. This RCT provided a clear description of its research scope and methods. The statistical analyses were appropriate and presented well. However, it does not provide an assessment/discussion of possible limitations or bias in this study.

RESULTS

Outcome	Intervention group	Comparator	group	Statistical significance
Blood loss (mL)	165 (SD: 19)	257 (SD: 21)		P<0.05
Surgery duration (minutes)	68 (SD: 15)	88 (SD: 10)		P<0.001
Outcome	Clinical importance		Clinical relevance	
Blood loss	1: reduced blood loss		1	
Surgery duration	1: reduced duration		2	

EXTERNAL VALIDITY

Generalisability

This RCT was conducted on a specific patient population (patients undergoing functional endoscopic sinuses surgery). Consequently, the findings of this study may not be applicable to other surgical patient populations.

Applicability

This trial was conducted in Egypt, which likely has comparable healthcare facilities to Australia. As endoscopic sinuses surgeries are performed in Australia, this study is likely applicable.

Comments

This study shows that magnesium sulphate induced hypotension reduces intraoperative blood loss and surgery duration.

Abbreviations: NR, not reported; RCT, randomised clinical trial; SD, standard deviation.

Fredin H, Gustafson C, Rosberg B. Hypotensive anesthesia, thromboprophylaxis and postoperative thromboembolism in total hip arthroplasty. Acta Anaesthesiologica Scandinavica 1984;28(5):503-507.

Affiliation/Source of funds

Department of Anaesthesiology and Orthopaedic Surgery. Malmo General Hospital, Sweden.

Funding: Swedish National Association against Heart and Chest Diseases and Herman Jarnhardt's Foundation.

Study design	Level of evidence	Location/setting
RCT	Level II	Hospital

		I
Intervention	Comparator	
Induced hypotension using sodium nitro blood pressure: 70-80mm Hg) with low-dihydroergotamine (HDHE)	No induced hypotens	ion with HDHE

Population characteristics

57 patients undergoing total hip arthroplasty.

Exclusion criteria: Patients with cardiovascular, pulmonary, renal, hepatic or thyroid diseases.

Length of follow-up	Outcomes measured	
10-14 days after surgery	Blood loss, transfusion dose, incidence of deep vein thrombosis (DVT) and pulmonary embolism (PE).	

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomised. Method NR	No significant differences in patient clinical characteristics were observed between the intervention and control group	Patient blinding NR. Diagnosis of DVT and PE were re-evaluated by an independent specialist, who were blinded to the patients' treatment status	NR	8 subjects with revision arthroplasties were excluded from study and not included in analysis

Overall quality assessment (descriptive)

Fair. This RCT provides clear description for the research question and scope. However, it is lacking details regarding the randomisation and blinding procedure. Statistical analyses performed were appropriate and well presented. Discussion did not assess possible biases in the study.

RESULTS			
Outcome	Intervention group	Comparator group	Statistical significance
Intraoperative blood loss (mL)	620 (SD: 240)	1070 (SD: 630)	P<0.001
Total blood loss (mL)	1170 (SD: 395)	1700 (SD: 860)	P<0.01
Intraoperative transfusion dose (mL)	580 (SD: 390)	1210 (SD: 620)	P<0.01
Total transfusion dose (mL)	920 (SD: 580)	1540 (SD: 1050)	P<0.01
Incidence of DVT	11/24	10/26	Not significant
Incidence of PE	6/26	1/28	Not significant

Outcome	Clinical importance	Clinical relevance
Peroperative blood loss	1: reduced blood loss	1
Total blood loss	1: reduced blood loss	1
Peroperative transfusion dose	1: reduced transfusion dose	1
Total transfusion dose	1: reduced transfusion dose	1
Incidence of DVT	4	1
Incidence of PE	4	1

Generalisability

This RCT was conducted on a specific patient population (patients undergoing Hip Arthroplasty, in addition to several other exclusion criteria). Consequently, their clinical characteristics may differ from a general surgery patient population.

Applicability

This trial was conducted in Sweden, with a similarly developed healthcare system as Australia. However, it is important to note that as this study was conducted in 1983, changes to patient management or clinical practice may affect the applicability of this study.

Comments

This study shows that induce hypotension reduces blood loss, transfusion dose.

Abbreviations: DVT, deep vein thrombosis; HDHE, low-dose heparin and dihydroergotamine; NR, not reported; PE, pulmonary embolism; RCT, randomised clinical trial; SD, standard deviation.

Jacobi KE, Bohm BE, Rickauer AJ, Jacobi C. Moderate controlled hypotension with sodium nitroprusside does not improve surgical conditions or decrease blood loss in endoscopic sinus surgery. Journal of Clinical Anaesthesia 2000;12(3):202-207.

Affiliation/Source of funds

University of Erlangen-Nuremberg, Erlangen, Germany.

Study design	Level of evidence	Location/setting
RCT	Level II	Hospital

Intervention	Comparator
Moderate hypotension (MAP: 65–75mmHg) using sodium nitroprusside	No induced hypotension

Population characteristics

32 patients undergoing endoscopic sinus surgery (ASA class I and II)

Length of follow-up	Outcomes measured
3 hours post-operation	Blood loss

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomised via computer generated random numbers.	No significant differences in patient clinical characteristics, duration of anaesthesia, intraoperative treatment were observed between the intervention and control group	NR	Hypotension group received same treatment as control group, plus sodium nitroprusside.	No lost to follow-up

Overall quality assessment (descriptive)

Fair. This study did not include a description of the blinding methods employed, if any.

RESULTS				
Outcome	Intervention group	Comparator g	roup	Statistical significance
Blood loss (mL)	278 ± 110 245 ± 132			P>0.05
Outcome	Clinical importance		Clinical rele	evance
Blood loss	4		1	

EXTERNAL VALIDITY

Generalisability

This RCT was conducted on a specific patient population (patients undergoing endoscopic sinus surgery, ASA class I and II). Consequently, the findings of this study may not be applicable to other surgical patient populations.

Applicability

This trial was conducted in Germany, with a similarly developed healthcare system as Australia. As such the findings of the study may be relevant in the Australian context.

Comments

The authors suggest that moderate hypotension may not be effective in reducing blood loss, while other studies suggest that profound hypotension (MAP=50mmHg) can provide a reduction in blood loss during surgery.

Abbreviations: ASA, American Society of Anaesthesiologists physical status; MAP, mean arterial pressure; NR, not reported; RCT, randomised clinical trial; SD, standard deviation.

Perioperative Question 3

Appendix F: Evidence summaries – Intervention 5 (Deliberate induced hypotension)

Karakaya D, Ustun E, Tur A, Baris S, Sarihasan B, Sahinoglu H, Guldogus F. Acute normovolemic hemodilution and nitroglycerin-induced hypotension: Comparative effects on tissue oxygenation and allogeneic blood transfusion requirement in total hip arthroplasty. Journal of Clinical Anesthesia 1999;11(5):368-374.

Affiliation/Source of funds

Ondokuz Mayis University, Kurupelit-SAMSUN, Turkey.

Funding: NR

Study design	Level of evidence		Location/setting
RCT	Level II		Hospital
1 1 11		0 1	

Intervention	Comparator
Nitroglycerine induced hypotension (60–65 mmHg)	No induced hypotension

Population characteristics

20 ASA class I and II patients undergoing primary total hip arthroplasty, performed via the posterior approach in the lateral decubitus position.

Length of follow-up	Outcomes measured
Up to fifth postoperative day	Surgery duration, transfusion requirements, haemoglobin level.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Randomised. Method NR.	No significant differences in patient clinical characteristics were observed between the intervention and control group	NR	All surgeries performed by a single surgeon. Subjects with bleeding disorders not included in study.	No lost to follow-up.

Overall quality assessment (descriptive)

Fair. This RCT did not describe the blinding method employed, if any. Due to the small sample size in each group, non-parametric methods should have been used.

RESULTS				
Outcome	Intervention group	Intervention group Comparator gro		Statistical significance
Surgery duration (minutes)	171.0 (SD: 26.6)	163.5 (SD: 24	4.9)	P>0.05
Blood transfusion (units)	2.3 (SD: 0.8)	2.7 (SD: 1.1)		P>0.05
Haemoglobin (g/dL) 5 minutes after intubation End of operation Fifth postoperative day	11.6 (SD: 0.4) 9.2 (SD: 0.19) 10.2 (SD: 0.3)	11.9 (SD: 0.8 9.7 (SD: 0.2) 10.3 (SD: 0.5	,	P>0.05
Outcome	Clinical importance	Clinical importance		elevance
Surgery duration	4	4		
Blood transfusion	4	4		
Haemoglobin	4		1	

Generalisability

This RCT was conducted on a specific patient population (ASA status I and II, undergoing hip arthroplasty, in addition to several other exclusion criteria). Consequently, the findings of this study may not be generalisable to other surgical patient populations.

Applicability

This trial was conducted in Turkey. Possible differences in the healthcare system, in addition to the small sample size make it difficult to assess the applicability in the Australian context.

Comments

Abbreviations: ASA, American Society of Anaesthesiologists physical status; NR, not reported; RCT, randomised clinical trial.

Kop EC, Spauwen PHM, Kouwenberg PPGM, Heymans FJM, van Beem HBH. Influence of controlled hypotension versus normotension on amount of blood loss during breast reduction. Journal of Plastic, Reconstructive and Aesthetic Surgery 2009;62(2):200-205.

Affiliation/Source of funds

University Medical Centre, Nijmegen, Netherlands. Slingeland Hospital, Doetinchem, Netherlands. Funding: Article was not supported by any funds.

Study design	Level of evidence		Location/setting
Double-blinded RCT	Level II		Hospital
Intervention		Comparator	

intervention	Comparator
Controlled hypotension using nitroprusside	Normotension
(MAP: 50mmHg)	

Population characteristics

85 Patients (<60 years, ASA I and II) undergoing bilateral breast reduction surgery.

Other Exclusion criteria: diabetes, hypertension, coagulation disturbances, kidney or liver dysfunction.

Length of follow-up	Outcomes measured
6 weeks post-operation	Blood loss

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomised. Random selection of envelope assigning patient to treatment group	No significant differences in patient clinical characteristics were observed between the intervention and control group	Double-blinded. Only anaesthesiologist knew the patients' treatment group.	Surgeon, who was unaware of patient's treatment group, determined when perfusor was stopped. Anaesthesiologist was not involved in data analysis.	34 patients did not meet selection criteria. Excluded from analysis.

Overall quality assessment (descriptive)

Good. This RCT provided a clear description of the research scope and study methods. The discussion included an assessment of possible limitations such as the in the measurement of intraoperative blood loss.

RESULTS				
Outcome	Intervention group	Comparator gr	oup	Statistical significance
Blood loss (mL)	316 (Range: 133–560) 598 (Range (250		0–1335)	P<0.001
Surgery duration (minutes)	56.4 (Range: 41-73)	62.7 (Range: 48	i–78)	P=0.013
Outcome	Clinical importance		Clinical re	elevance
Blood loss	1: Reduced blood loss		1	
Surgery duration	1: Reduced surgery duration		2	

Generalisability

This RCT was conducted on a specific patient population (females with ASA status I-II under the age of 60, undergoing bilateral breast reduction, in addition to several other exclusion criteria). Consequently, the findings of this study may not be applicable to other surgical patient populations.

Applicability

This trial was conducted in the Netherlands, which is similarly well developed like Australia. As this surgical procedure is performed in Australia, the findings of this study is likely applicable.

Comments

This study shows that induced hypotension reduces blood loss and surgery duration during breast reduction surgery

Abbreviations: ASA, American Society of Anaesthesiologists physical status; MAP, mean arterial pressure; NR, not reported; RCT, randomised clinical trial.

O'Connor PJ, Hanson J, Finucane BT. Induced hypotension with epidural/general anesthesia reduces transfusion in radical prostate surgery. Canadian Journal of Anesthesia 2006;53(9):873-880.

Affiliation/Source of funds

University of Alberta, and the Division of Health, Population and Information, Cross Cancer Institute, Edmonton, Alberta, Canada.

Study design	Level of evidence	Location/setting
Prospective, randomised, single-blind trial	Level II	University of Alberta Hospital, Canada

titai			
Intervention		Comparator	
Combined epidural and anaesthesia to control mean arterial pressure (MAP) at 50-60mmHg.		General anaesthesia	alone, no control of MAP.

Population characteristics

99 patients, ASA status I-III, with adenocarcinoma of the prostate to undergo radical retropubic prostatectomy.

Length of follow-up	Outcomes measured
Patients monitored up till discharge.	Primary outcome: Blood loss, transfusion frequency and dose.
	Secondary outcome: Operating time, hospital length of stay, occurrence of serious adverse events.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Randomisation using a computer-generated table of random numbers, patients were block randomised (block size=10) using blinded study envelopes which were opened prior to surgery.	No significant differences in patient clinical characteristics were observed between the intervention and control group	Single-blinded.	The lack of blinding of the anaesthesiologists involved. The use of rigid automatic laboratory-based transfusion trigger could introduce potential bias due to the effects of haemodilution. However, the trigger rates in the intervention and control group were comparable.	NR

Overall quality assessment (descriptive)

Good. This RCT provides clear description for the randomisation, inclusion and exclusion criteria, methods, outcomes measured, and employed suitable statistical methods for analysis. The discussion was comprehensive and assessed possible limitations and biases. Anaesthesiologists were not blinded, however, only 23% of the transfusions were decided by the anaesthesiologist. The remaining 77% of transfusions were initiated postoperatively by non-study personnel.

RESULTS			
Outcome	Intervention group	Comparator group	Statistical significance
Blood loss (mL)	955 (SD: 517)	1477 (SD: 823)	P<0.001
Transfusion frequency	2/49 (4%)	9/50 (18%)	P=0.028
Total transfusion dose in group (units)	3	24	NR
Operating time (minutes)	107 (SD:36)	122 (SD:32)	P=0.038

Hospital length of stay (>5 days)	24/49 (49%)	34/5	50 (68%)	P=0.055	
Serious adverse events	0	0		NA	
Outcome	Clinical important	Clinical importance		ance	
Blood loss	1: reduced loss	1: reduced loss		1	
Transfusion requirements	1: reduced requirer	1: reduced requirements		1	
Mean transfusion dose	NA	NA			
Operating time	2: reduced duration	2: reduced duration			
Hospital length of stay	4	4			
Serious adverse events	NA	NA			

Generalisability

This RCT was conducted on a specific patient population (males with ASA status I-III, with adenocarcinoma of the prostate, undergoing radical retropubic prostatectomy and bilateral pelvic lymphadenectomy). Consequently, the findings of this study may not be applicable to other surgical patient populations (e.g. females, patients with ASA status III-V).

Applicability

This trial was conducted in Canada, with a similarly developed healthcare system as Australia. As such the findings of the study may be relevant in the Australian context.

Comments

This study shows that induce hypotension reduces blood loss, transfusion frequency and operating time.

Abbreviations: ASA, American Society of Anaesthesiologists physical status; MAP, mean arterial pressure; NA, not applicable; NR, not reported; RCT, randomised clinical trial; SD, standard deviation.

Piper SN, Suttner SW, Maleck WH, Kumle B, Haisch G, Boldt J. Effects of sodium nitroprusside-induced controlled hypotension on pancreatic function assessed by pancreatitis-associated protein in patients undergoing radical prostatectomy. European Journal of Anaesthesiology 2002;19(8):609-613.

Affiliation/Source of funds

Department of Anaesthesiology and Intensive Care Medicine, Klinikum Ludwigshafen, Germany. Funding: NR

Study design	Level of evidence	Location/setting
RCT	Level II	Hospital

Intervention		Comparator	
Controlled hypotension (MAP: 50mm Hg nitroprusside) using sodium	No induced hypotens	ion

Population characteristics

30 patients undergoing elective radical prostatectomy (ASA class II and III only)

Length of follow-up	Outcomes measured
24 hours post-operation	Blood loss, blood transfusion dose, duration of surgery

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomised using a random list.	No significant differences in patient clinical characteristics were observed between the intervention and control group	NR	Routine intraoperative care used for both study groups.	No lost to follow-up

Overall quality assessment (descriptive)

Fair. This study did not include a description of the blinding methods employed, if any. Also, as the primary aim of this study was not blood loss/transfusion, the authors did not discuss these outcomes in detail.

RESULTS				
Outcome	Intervention group	Comparato	r group	Statistical significance
Blood loss (mL)	843 (SD: 233)	1526 (SD: 4	09)	P<0.05
Transfusion frequency	0 patients	4 patients		P<0.05
Total transfusion dose in group (units)	0	10		P<0.05
Surgery duration (minutes)	154 (SD: 20.6)	164 (SD: 20	.6)	P>0.05
Haemoglobin concentration	Higher in intervention group than cont		rol group	P<0.05
Outcome	Clinical importance		Clinical re	levance
Blood loss	1:reduced blood loss		1	
Transfusion frequency	1: reduced frequency		2	
Transfusion dose	1: reduced requirements		2	
Surgery duration	4		2	

Haemoglobin concentration	1:Increased haemoglobin concentration	1
EXTERNAL VALIDITY		
Generalisability		
This RCT was conducted on a specific patient population (males with ASA status II-III, undergoing radical prostatectomy, in addition to several other exclusion criteria). Consequently, the findings of this study may not be applicable to other surgical patient populations.		
Applicability		
This trial was conducted in Germany, with a similarly developed healthcare system as Australia. As such the findings of the study may be relevant in the Australian context.		
Comments		

Abbreviations: ASA, American Society of Anaesthesiologists physical status; MAP, mean arterial pressure; NR, not reported; RCT, randomised clinical trial; SD, standard deviation.

This study shows that induced hypotension reduces blood loss and blood transfusion.

Sood S, Jayalaxmi TS, Vijayaraghavan S, Nundy S. Use of sodium nitroprusside induced hypotensive anaesthesia for reducing blood loss in patients undergoing lienorenal shunts for portal hypertension. British Journal of Surgery 1987;74(11):1036-1038.

Affiliation/Source of funds

All India Institute of Medical Sciences, Ansari Nagar, New Delhi, India.

Funding: NR

Study design	Level of evidence		Location/setting
RCT	Level II		Hospital

Intervention	Comparator
Sodium nitroprusside induced hypotensive anaesthesia (systolic blood pressure < 95mmHg)	No induced hypotension

Population characteristics

18 patients undergoing elective, proximal, lienorenal shunts for portal hypertension.

Length of follow-up	Outcomes measured
48 hours after surgery	Blood loss, transfusion dose.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomised using table of random numbers.	Control group had more males and more subjects with extrahepatic obstruction.	NR	Preoperative investigations, surgical technique and postoperative management were undertaken by the same surgical and anaesthetic teams for both treatment groups.	2 patients were excluded, not considered in analysis.

Overall quality assessment (descriptive)

Fair. This RCT did not provide a clear description for the blinding methods used, if any. The study also provides no assessment/discussion of possible limitations or bias. The final study sample size of 18 patients is small, however statistical significance was achieved in the analysis.

RESULTS					
Outcome Intervention group Comparator gr		group	Statistical significance		
Blood loss (mL)	517 (SD: 220) 1286 (SD: 523)		3)	P<0.01	
Transfusion requirement (units)	0.88 (SD: 0.9) 3.0 (SD: 1.2)			P<0.01	
Outcome	Clinical importance		Clinical relevance		
Blood loss	1: reduced blood loss		1		
Transfusion requirement	1: reduced requirements		1		

EXTERNAL VALIDITY

Generalisability

This RCT was conducted on a specific patient population (patients undergoing elective, proximal, lienorenal shunts for portal hypertension). In addition, the small sample size makes it difficult for findings of this study to be extended to the general surgical population.

Applicability

This trial was conducted in India, in a large well-established medical institution. As such the level and quality of healthcare is likely comparable to that in Australia. As such the findings of this study are likely applicable.

Comments

This study shows that sodium nitroprusside induced hypotension blood loss and transfusion requirements.

Abbreviations: NR, not reported; RCT, randomised clinical trial; SD, standard deviation.

Suttner SW, Piper SN, Lang K, Huttner I, Kumle B, Boldt J. Cerebral effects and blood sparing efficiency of sodium nitroprusside-induced hypotension alone and in combination with acute normovolaemic haemodilution. British Journal of Anaesthesia 2001;87(5):699-705.

Affiliation/Source of funds

Department of Anaesthesiology and Intensive Care Medicine, Klinikum der Stadt Ludwigshafen, Germany. Funding: NR

Study design	Level of evidence	Location/setting	
RCT	Level II	Hospital	

Intervention	Comparator
Controlled hypotension (MAP: 50mm Hg) using sodium nitroprusside	No induced hypotension
Controlled hypotension with haemodilution (beyond the scope of Q3:Intervention 5)	

Population characteristics

28 patients, undergoing elective radical prostatectomy.

Exclusion criteria: ASA class greater than III, myocardial infarction within 6 months, documented coronary artery disease or carotid artery stenosis, abnormal coagulation, liver dysfunction, medication with aspirin, renal insufficiency.

Length of follow-up	Outcomes measured		
Up to discharge from post anaesthesia care unit	Blood loss, transfusion dose and frequency.		

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomised, method not specified.	No significant differences in patient clinical characteristics were observed between the intervention and control group	NR	Anaesthesiologist was not involved in data analysis.	6 subjects failed to meet inclusion criteria, excluded from analysis

Overall quality assessment (descriptive)

Good. This RCT included a modest number of subjects, although initial power calculations suggest that the study was sufficiently powered. Information as to the method of randomising and blinding was not specified. The analyses performed were appropriate and the results presented clearly.

RESULTS					
Outcome	Intervention group Compara		or group	Statistical significance	
Blood loss (mL)	788 (SD: 193) 1335 (SD		460)	P<0.05	
Transfusion frequency	1/14 (7%)	7/14 (50%)		P<0.05	
Total transfusion dose in group (units)	3	17		P<0.05	
Outcome	Clinical importance	Clinical rel		evance	
Blood loss	1: reduced blood loss	1			
Transfusion frequency	1: reduced frequency	1			
Transfusion dose	1: reduced requirements		1		
EXTERNAL VALIDITY					

Generalisability

This RCT was conducted on a specific patient population (males with ASA status I-III, undergoing radical prostatectomy, in addition to several other exclusion criteria). Consequently, the findings of this study may not be applicable to other surgical patient populations.

Applicability

This trial was conducted in Germany, with a comparable level of healthcare as in Australia. As such the findings of the study are likely applicable.

Comments

This study shows that induced hypotension reduces blood loss, transfusion dose and frequency.

Abbreviations: ASA, American Society of Anaesthesiologists physical status; MAP, mean arterial pressure; NR, not reported; RCT, randomised clinical trial; SD, standard deviation.

Intervention 6 - Prevention of hypothermia

Level I evidence

abstract). AANA J 1	Maintaining intraoporative no					
		ormothe	ermia: a meta	a-analysis of out	comes with costs (Struc	ctured
Affiliation/Source	of funds					
	Management, University of Minne	esota.				
Funding: NR					-	
Study design		Level	of evidence)	Location/setting	
Meta-analysis of 18 studies to determine the difference in patients' outcomes between normothermic and mildly hypothermic patients undergoing surgery.			Level I Hospitals		Hospitals	
Intervention			Comparate	or		
Maintenance of normothermia			Mild hypothermia			
Population charac	teristics					
Studies which include	de patients undergoing any surg	ery. Pa	tient populati	on does not inclu	ude cases of extreme h	ypothermia.
Length of follow-u	p		Outcomes m	neasured		
NR. Appears variab information on lengt	le, with only some studies provic h of stay.		Transfusion dose and frequency, mortality rate, myocardial infarction, cost, length of stay in hospital and ICU			
INTERNAL VALIDI	ΤΥ					
Allocation	Results		linding nalysis	Treatment/ measurement bias		Follow-up (ITT)
3 of the 18 included studies were not randomised. Method NR.	The authors state that there were no significant differences in the general characteristics between the intervention and control group, in any of the included studies.	addressed by statistical adjustment during meta-analysis, where information was available. Publication bias of the processor addressed by statistical adjustment during meta-analysis, where information was available.		NR		

Overall quality assessment (descriptive)

Poor. The literature search conducted in this meta-analysis is acceptable. However, the inclusion of non-randomised trials (3/18) and the lack of information on the allocation method and blinding of individual studies diminish the quality of this study. The meta-analysis explored possible sources of bias and provided useful estimates for several outcomes of interest.

RESULTS					
Outcome	Intervention group	Comparator group	Statistical significance		
Units of red blood cells transfused	0.117 (SD: 0.0247)	1.167 (SD: 0.0867)	P<0.05		
Units of plasma transfused	0.3 (0.09)	1.4 (0.2)	P<0.05		
Units of platelets transfused	0.2 (0.01)	0.9 (0.06)	P<0.05		
Need for transfusion (probability)	14.43% (SD: 3.14)	24.19% (SD: 4.57)	P<0.05		

			1
11.77 (SD: 0.1047) 19.44		0.1600)	P<0.05
5.51 (SD: 0.0863) 9.70 (SD: 0.		0.1712)	P<0.05
2.30% (SD: 0.88)	2.30% (SD: 0.88) 4.07% (SD:		P<0.05
2.70% (SD: 0.85)	6.01% (SD): 1.73)	P<0.05
Between \$2495-\$7073 p	er patient		NA
Clinical importance		Clinical relevance	
1		1	
1		1	
1		1	
1	1		
1	2		
1	1		
1	1		
1		1	
NA	NA		
	5.51 (SD: 0.0863) 2.30% (SD: 0.88) 2.70% (SD: 0.85) Between \$2495–\$7073 p Clinical importance 1 1 1 1 1 1	5.51 (SD: 0.0863) 9.70 (SD: 0.230% (SD: 0.88) 4.07% (SD: 0.88) 2.70% (SD: 0.85) 6.01% (SD: 0.85) Between \$2495-\$7073 per patient Clinical importance 1 1 1 1 1 1 1	5.51 (SD: 0.0863) 9.70 (SD: 0.1712) 2.30% (SD: 0.88) 4.07% (SD: 1.34) 2.70% (SD: 0.85) 6.01% (SD: 1.73) Between \$2495-\$7073 per patient Clinical importance Clinical relevance 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1

Generalisability

This review included studies of patients undergoing a variety of surgical procedures, in which several methods of hypothermia prevention were used. The maintenance of normothermia had a positive effect across a number of clinically important outcomes, as such, is likely to be beneficial in part to the general surgical population (excluding extreme hypothermia cases).

Applicability

The studies were conducted in developed countries (similar to Australia). The surgical procedures included are relevant in the Australian context. As such, this intervention and its findings are likely to be applicable in Australia.

Comments

This review suggests that the maintenance of normothermia results in fewer adverse outcomes, and lower overall hospital cost.

Abbreviations: ICU, intensive care unit; NA, not applicable; NR, not reported; RCT, randomised clinical trial; SD, standard deviation.

Rajagopalan S, Mascha E, Na J, Sessler DI. The effects of mild perioperative hypothermia on blood loss and transfusion requirement. Anesthesiology 2008;108(1):71-77.

Affiliation/Source of funds

Department of Outcomes Research, Cleveland Clinic, Cleveland Ohio.

Funding: National Institutes of Health Grant and the Joseph Drown Foundation.

Study design	Level of evidence	Location/setting
Systematic review of 18 RCTs that compared normothermic patients to those who had mild intraoperative hypothermia.	Level I	Hospitals

Intervention	Comparator
Maintenance of normothermia	Mild hypothermia (34–36°C)

Population characteristics

Studies which include patients undergoing any surgery. Patient population does not include studies where the core temperature is less than 34°C.

Length of follow-up	Outcomes measured
NR.	Blood loss and need for blood transfusion.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
All included studies were randomised. Method NR.	Analysis of baseline characteristics of each study not reported.	NR	Funnel plots used did not indicate substantial publication bias for either outcome. No significant study effect on treatment effect size observed.	ITT was specified in 10 studies.

Overall quality assessment (descriptive)

Good. This review clearly defined the research question and strategy. Although detailed characteristics of included studies were absent, quality scores were assigned for each study. The statistical analyses were well conducted and clearly presented. The presences of publication bias and study effect were also examined.

RESULTS						
Outcome	Intervention group Comparator		or group	Statistical significance		
Blood loss (Intervention vs control)	NR NR			Ratio: 0.84 (0.74, 0.96)		
Need for transfusion	NR NR			RR=0.78 (0.63, 0.97)		
Outcome	Clinical importance	Clinical importance		vance		
Blood loss	2		1			
Need for transfusion	2		1			

Generalisability

This review included studies of patients undergoing a variety of surgical procedures. However, this review excluded studies where the core temperature was less than 34°C, or if hypothermia had been induced. As such, the patient population may not be representative of patients undergoing surgery in which cooling methods are employed.

Applicability

The studies were conducted in developed countries (similar to Australia). The surgical procedures included are relevant in the Australian context. As such, this intervention and its findings are likely applicable in Australia.

Comments

This review suggests that the maintenance of normothermia results in less blood loss, and a reduced need for blood transfusions.

Abbreviations: NR, not reported; RCT, randomised clinical trial; RR, relative risk; SD, standard deviation.

Scott EM, Buckland R. A systematic review of intraoperative warming to prevent postoperative complications (Structured abstract). AORN Journal 2006;83:1090-1104.

Affiliation/Source of funds

University of Durham, Stockton-on-Tees, England. Easington Primary Care Trust, Country Durham, England. Funding: Actamed ltd, Wester Yorkshire, UK, and Pegasus ltd, Hampshire, UK.

Study design	Level of evidence	Location/setting
Systematic review of 26 RCTs that examined if preventing hypothermia during surgery prevents postoperative complications such as need for blood transfusion.	Level I	Hospitals

Intervention	Comparator
Prevention of hypothermia during surgery	No prevention of hypothermia.

Population characteristics

Patients having surgical procedures (other than cardiac procedures) under regional or general anaesthesia.

Length of follow-up		Outcomes measured
	NR, studies must include follow-up beyond	Need for blood transfusion, morbid cardiac events (eg myocardial
	intraoperative phase (ie during the post-anesthesia	infarction, angina, tachycardia) and pain.
	care unit, or hospital stay).	

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
All included studies were randomised. NR in 13 studies. 6 studies used computer generated numbers for allocation. 4 used sealed opaque envelopes. 1 used sealed envelopes. 1 used flip of a coin. 1 used random number generator blocking every 10 patients.	Baseline characteristics of intervention and control groups were comparable all except two studies, where the age and weight differed between treatment and control groups. All except three studies showed significant differences in the temperature between the treatment and control group.	8 studies were double-blinded. 8 studies were single-blinded. 8 studies NR. 2 studies stated blinding not possible.	Publication bias or bias of individual studies not reported.	NR

Overall quality assessment (descriptive)

Fair. This review provides clear description for the randomisation, inclusion and exclusion criteria, and quality assessment of the included studies. However, the pooled estimates for morbid cardiac events were derived from just two studies, while the need for blood transfusion was derived from four studies. No assessment of publication bias was performed.

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RESULTS						
Outcome	Intervention group Comparato		or group	Statistical significance		
Morbid cardiac events	NR	NR NR		RR=0.34 (0.20, 0.57)		
Need for blood transfusion (yes/no)	NR	NR NR		RR=0.39 (0.22, 0.68)		
Pain	No significant differences in pain between groups					
Outcome	Clinical importance	Clinical importance		vance		
Morbid cardiac events	1: reduced risk	1: reduced risk				
Need for blood transfusion	1: reduced need for transfusion		1			
Pain	4		1			

Generalisability

This review included studies of patients undergoing a variety of surgical procedures, in which several methods of hypothermia prevention were used. The review excluded studies involving cardiac procedures; as such it may not be representative of cardiac surgery patients.

Applicability

The studies in this review were not conducted in Australia; however, they were conducted in developed countries with similarly developed healthcare systems.

Comments

This review suggests that preventing hypothermia during surgery may reduce the likelihood of morbid cardiac events and the need for blood transfusion.

Abbreviations: NR, not reported; RCT, randomised clinical trial; RR, relative risk.

Jeong SM, Hahm KD, Jeong YB, Yang HS, Choi IC. Warming of intravenous fluids prevents hypothermia during off-pump coronary artery bypass graft surgery. Journal of Cardiothoracic and Vascular Anesthesia 2008;22:67-70.

Affiliation/Source of funds

University of Ulsan, Seoul Korea.

Funding: NR

Study design	Level of evidence		Location/setting
RCT	Level	II	Hospital
Intervention		Comparator	
Warmed (41°C) intravenous fluids		Conventional treatment	

Population characteristics

40 patients undergoing isolated off-pump coronary artery bypass (OPCAB) surgery.

Length of follow-up		Outcomes measured
Biochemical measurements, up to 24 hours after	Blood transfusion dose, surgery duration, temperature, ICU stay, hospital stay.	

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomised. Method NR.	No difference in preoperative characteristics between patient groups.	Investigators not blinded	Standardised treatment and management applied to all subjects irrespective of treatment group.	No lost to follow-up.

Overall quality assessment (descriptive)

Poor. The authors recognise that the small sample size was likely underpowered to detect changes in clinical data, as the study was designed to be powered to detect a change in patient temperature. Investigators were not blinded the treatment group of the subjects, this may have led to bias.

RESULTS				
Outcome	Intervention group	Comparat	or group	Statistical significance
Blood transfusion dose (mL)	400.5 ± 622.8	400.5 ± 622.8 365.0 ± 43°		P>0.05
Surgery duration (minutes)	247 ± 59	245 ± 49		P>0.05
Bladder temperature at 4 hours into operation (°C)	36.6 ± 0.32 35.8 ± 0.7			P<0.05
ICU stay (hours)	59.6 ± 19.6	70.5 ± 17.	8	P>0.05
Hospital stay (days)	10.6 ± 2.2	11.6 ± 2.7		P>0.05
Outcome	Clinical importance		Clinical rele	vance
Blood loss	4	4		
Surgery duration	4	4		
Bladder temperature	1: higher temperature		1	

ICU stay	4	2
Hospital stay	4	2

Generalisability

The small sample size of this study, and the specific patient population examined (patients undergoing OPCAB) makes generalising the findings from this study difficult.

Applicability

This study was conducted in Korea. Differences in the level of healthcare between Korea and Australia may limit the applicability of the findings of this study.

Comments

Warming of intravenous fluids does not reduce blood loss, ICU stay or hospital stay in patients undergoing OPCAB.

Abbreviations: NR, not reported; OPCAB, off-pump coronary artery bypass; RCT, randomised clinical trial.

Kim YS, Lee JY, Yang SC, Song JH, Koh HS, Park WK. Comparative Study of the Influence of Room-Temperature and Warmed Fluid Irrigation on Body Temperature in Arthroscopic Shoulder Surgery. Arthroscopy – Journal of Arthroscopic and Related Surgery 2009;25(1):24-29.

Affiliation/Source of funds

College of Medicine, Catholic University of Korea, Seoul, Korea. The Armed Forces Capital Hospital, Korea.

Funding: NR

Study design	Level of evidence		Location/setting
RCT	Level II		Hospital
Intervention		Comparator	
Warm irrigation fluid (35–37°C)		Irrigation fluid at room temperature	

Population characteristics

50 ASA I or II patients undergoing arthroscopic shoulder surgery.

Length of follow-up	Outcomes measured
1 days post-operation	Change in haemoglobin, surgery duration, pain, hypothermia, body temperature

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomised. Method NR.	No difference in preoperative characteristics between patient groups.	Postoperative measurements were recorded by an independent observer blinded to the study.	All operative procedures were performed by the same orthopaedic surgeon.	Four patients exclude because of incomplete data. Excluded from analysis

Overall quality assessment (descriptive)

Fair. This RCT had clearly defined research questions and methods. Based on the results of previous studies, their power calculation indicated that they had over 80% power.

RESULTS						
Outcome	Intervention group	Comparat	or group	Statistical significance		
Change in Haemoglobin (g/dL)	1.7 ± 0.7	1.7 ± 0.7 1.4 ± 0.6		1.4 ± 0.6		P=0.165
Surgery duration (minutes)	94.5 ± 21.9	94.5 ± 21.9 91.1 ± 32.4		P=0.68		
Postoperative pain (VAS)	5.0 ± 1.7 4.9 ± 1.6			P=0.927		
Final body temperature (°C)	36.2 ± 0.3	36.2 ± 0.3 35.5 ± 0.3		P<0.001		
Hypothermia (%)	17.4	91.3		P<0.001		
Outcome	Clinical importance		Clinical relevance			
Change in Haemoglobin	4	4				
Surgery duration	4		2			
Postoperative pain	4		1			

Hypothermia	1: reduced incidence of	1
	hypothermia	

Generalisability

This RCT was conducted in a specific patient population (undergoing arthroscopic shoulder surgery). As such, they may not share clinical characteristics with a general surgical patient population.

Applicability

This study was conducted in a military hospital in Korea. Differences in the demographics and exposure of the patients and the level of healthcare, as compared to Australia may limit the applicability of the findings of this study.

Comments

This review suggests that the warm fluid irrigation reduces perioperative hypothermia, however it does not significantly influence the change haemoglobin following surgery.

Abbreviations: NR, not reported; RCT, randomised clinical trial; VAS, visual analogue scale.

Melling AC, Ali B, Scott EM, Leaper DJ. Effects of preoperative warming on the incidence of wound infection after clean surgery: A randomised controlled trial. Lancet 2001;358(9285):876-880.

Affiliation/Source of funds

University Hospital of North Tees, Stockton-on-Tees, UK.

Funding: Action Research and Smith & Nephew Foundation.

Study design	Level of evidence		Location/setting
RCT	Level II		Hospital
Intervention		Comparator	
Preoperative warming		Standard care (no preoperative warming)	

Population characteristics

421 patients having clean surgery (breast, varicose vein, or hernia), that would result in a scar longer than 3 cm. Exclusion criteria: Pregnant, under 18 years, long term steroids, had received radiotherapy or chemotherapy in the last 4 weeks, or had an infection at the time of the surgery.

Length of follow-up	Outcomes measured
6 weeks	Wound infection, ASEPSIS score.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomised. Method NR.	No difference in preoperative characteristics between patient groups.	Treatment allocation concealed in opaque envelopes.	A single train observer, blinded to the treatment allocation assessed subjects at 2 and 6 weeks for wound infection.	6 patients lost to follow-up at 6 weeks. Outcomes evaluated on an ITT basis.

Overall quality assessment (descriptive)

Good. Prospective power calculations indicated that the sample size provided 90% power to detect a 5% change in infection rates. Statistical analyses performed were appropriate, with multivariate analysis used to identify possible risk factors.

RESULTS				
Outcome	Intervention group Comparato		or group	Statistical significance
Wound infection	13 (5%)	19 (14%)		P=0.001
ASEPSIS score				
0–10	259 (94%)	115 (83%)		
11–20	8 (3%)	7 (5%)		P=0.007
21–30	6 (2%)	9 (7%)		P=0.007
31–40	2 (0.7%)	6 (4%)		
>40	2 (0.7%)	2 (1%)		
Outcome	Clinical importance		Clinical relev	vance
Wound infection	1		1	
ASEPSIS	1		2	

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Generalisability

This RCT was conducted in patients undergoing a variety of clean surgery, as such findings of this study may be generalisable to other surgical patients undergoing similar clean surgeries. The study was adequately powered, adding credibility to the study findings.

Applicability

This study was conducted in the UK, which has a similar healthcare system to Australia. Also, the surgical procedures examined are performed in Australia, as such, findings from this study are likely applicable in Australia.

Comments

This study suggests that preoperative warming reduces the incidence of wound infection in patients undergoing clean surgical procedures.

Abbreviations: RCT, randomised clinical trial.

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Yau TM, Carson S, Weisel RD, Ivanov J, Sun Z, Yu R, Glynn MF, Teasdale SJ. The effect of warm heart surgery on postoperative bleeding. The Journal of Thoracic and Cardiovascular Surgery 1992;103:1155-1162.

Affiliation/Source of funds

The Toronto Hospital and the University of Toronto, Toronto, Ontario, Canada.

Funding: Medical Research Council of Canada

Study design	Level of evidence		Location/setting
Double-blinded RCT.	Level II		Hospital
Intervention		Comparator	
Warm systemic perfusion (35–37°C)		System perfusion at 25–29°C	

Population characteristics

146 consecutive patients undergoing isolated primary coronary artery bypass grafting (CABG).

Length of follow-up	Outcomes measured
Up to six days after operation	Blood loss, transfusion requirements, haemoglobin levels.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomised allocation reported using a randomisation table.	No difference in preoperative characteristics between patient groups.	Patient and surgeon blinded to the treatment group.	Transfusion of blood ordered by surgeon or anaesthetist who was not aware of the patient's treatment group.	Three patients who had reoperation were excluded from analysis.

Overall quality assessment (descriptive)

Fair. This double-blinded RCT has clearly defined research questions, with several post-operation follow-up carried out. Antifibrinolytic therapy was concurrently used by some patients in this study, and was not controlled for in this study; however, the use of such therapy had no bearing on the assignment to treatment group and as such would have had a non-differential effect, if any. The authors recognised that their sample size may have been underpowered to detect differences between treatment groups.

RESULTS				
Outcome	Intervention group	Comparat	or group	Statistical significance
Post operative blood loss :				
6 hours	409 ± 36 mL	418 ± 41 r	mL	
12 hours	591 ± 38 mL	596 ± 50 r	mL	P>0.05
24 hours	864 ± 42 mL	918 ± 68 r	mL	
Blood transfusion frequency	55%	64%		P=0.24
Haemoglobin levels	NR	NR		P>0.05
Outcome	Clinical importance		Clinical rele	vance
Blood loss at 6 hours	4	4		
Blood loss at 12 hours	4		1	
Blood loss at 24 hours	4		1	

Blood transfusion frequency	4	1
Haemoglobin levels	4	1

Generalisability

This RCT was conducted in a specific patient population (undergoing CABG surgery). As such, they may not share clinical characteristics with a general surgical patient population. In addition, the small sample size and the resulting lack of power, as recognised by the authors, make it difficult to draw conclusions with any certainty.

Applicability

This study was conducted in Canada, which has comparable healthcare system. CABG surgeries are regularly performed in Australia. As such, these findings may have some relevance to a similar patient population in Australia.

Comments

This review suggests that the use of warm system perfusion during heart surgery does not reduce postoperative bleeding.

Abbreviations: CABG, coronary artery bypass grafting; NR, not reported; RCT, randomised clinical trial.

Zhao J, Luo AL, Xu L, Huang YG. Forced-air warming and fluid warming minimize core hypothermia during abdominal surgery. Chinese medical sciences journal / Chinese Academy of Medical Sciences 2005;20:261-264.

Affiliation/Source of funds

Peking Union Medical College Hospital, Chinese Academy of Medical Sciences, Beijing.

Funding: NR

Study design	Level of evidence	Location/setting
RCT	Level II	Hospital

Intervention	Comparator
Warming with air-forced warming blanket and intravenous fluid warming system	Conventional treatment (Covered in cotton sheet)

Population characteristics

40 patients undergoing abdominal surgery lasting at least 2 hours (ASA class I and II).

Length of follow-up	Outcomes measured
NR	Blood loss, surgery duration, red blood cell transfusion, plasma transfusion and core temperature.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomised. Method NR.	No difference in preoperative characteristics between patient groups.	NR	Standardised treatment and management applied to all subjects irrespective of treatment group.	No lost to follow-up.

Overall quality assessment (descriptive)

Fair. This RCT did not provide a description of the blinding or randomisation methods employed.

RESULTS				
Outcome	Intervention group	Comparat	or group	Statistical significance
Blood loss (mL)	639 (SD: 441)	421 (SD: 2	249)	P>0.05
Surgery duration (minutes)	204 (SD: 76)	230 (SD: 8	38)	P>0.05
Red blood cell transfusion (units)	2.6 (SD: 2.5)	1.6 (SD: 2	.4)	P>0.05
Plasma transfusion (units)	220 (SD: 460)	240 (SD: 4	180)	P>0.05
Core temperature (°C)	36.4 (SD: 0.4)	35.3 (SD:	0.5)	P<0.001
Outcome	Clinical importance		Clinical rele	vance
Blood loss	4		1	
Surgery duration	4	4		
Red blood cell transfusion	4	4		
Plasma transfusion	4	4		
Core temperature	1: increased temperatur	1: increased temperature		

Generalisability

This study examined patients undergoing selective abdominal surgery (ASA Class I and II), as such it is likely most generalisable to patients undergoing such surgical procedures.

Applicability

This study was conducted in China, which has a different healthcare system and demographics to Australia. As such, it is difficult to assess the applicability and feasibility of such an intervention in Australia.

Comments

This study found that warming prevents hypothermia during abdominal surgery; however, no significant effect was observed on blood loss and transfusion requirements.

Abbreviations: ASA, American Society of Anaesthesiologists; NR, not reported; RCT, randomised clinical trial; SD, standard deviation.

Intervention 7 – Point-of-care testing using thromboelastography

Level II evidence

Citation

Ak, K., Isbir, SC., et al., Thromboblastography-based algorithm reduces blood product use after elective CABG: a prospective randomised study. Journal of Cardiac Surgery 2009;24:404-410.

Affiliation/Source of funds

Department of Cardiovascular Surgery, Marmara University, Istanbul, Turkey

Department of Biochemistry, Marmara University, Istanbul, Turkey

Cardiovascular Surgery Unit, Academic Hospital, Istanbul, Turkey

No conflict of interest

Study design	Level of evidence	Location/setting
Prospective RCT	II	Turkey, Academic hospital.
Randomisation process not described		

Intervention	Comparator
Thromoboelastography (TEG) based algorithm guided transfusion (N=114) (comprising kaolin-activated (k) TEG and h-kTEG analyses) which was a modified version of the one proposed by Royston and Kier ⁶ using tranexamic acid instead of aprotinin.	Clinician-directed transfusion (CDT, n=110) using criteria obtained from abnormal conventional laboratory tests (PT, APTT and platelet count) absence of visible clots and presence of generalized oozing-type bleeding in the surgical field to determine blood product administration.

Population characteristics

224 patients undergoing elective first-time coronary artery bypass grafting (CABG) with cardiopulmonary bypass. For CDT and TEG respectively: mean ages 65.9 and 63.2, males 79% and 75%, diabetes mellitus 35% and 29%.

Additional treatment: 65% of CDT group and 59% of TEG group on aspirin therapy prior to the operation Additional doses of protamine sulphate were given to some patients Transamine 10% initial dose was given at 10mg/kg over 20 mins followed by an infusion of 1mg/(kg.h) PRBCs transfused when hemocrit>25% (18% during CPB) –if patient was intolerant to anaemia or older age the threshold for blood transfusion was raised.

Length of follow-up	Outcomes measured
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⁶ Royston, D., von Kier S. Reduced hemostatic factor transfusion during heparinise-modified thromboelstaography during cardiopulmonary by-pass. Br J Anaesth 2001;86:575-578.

T4=On admission to the intensive care unit T5=6 hours after CPB T6=24 hours after CPB INTERNAL VALIDITY Allocation Results Blinding analysis Treatment/measurement bias Random No significant difference between groups in demographics and perioperative variables but hypertension was significantly higher in T4=On admission to the intensive care unit T5=6 hours after CPB Treatment/measurement bias as There was the possibility of measurement bias as the two groups did not have the same coagulation tests which may have been advantageous to the TEG group. Table 10 Table 11 Table 12 Table 13 Table 14 Table 14 Table 15 Table 15 Table 16 Table 16 Table 17 Table 17 Table 17 Table 17 Table 17 Table 18 Table 19 Table			
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significantly higher in need for transfusion). group. discharge 6.3 days.			
the TEG group - It is not clear if the clinician directing transfusion or Transfu			
P=0.000. Assessing other I here was also the Patient s	atus followed		
interventions was aware of the desiration up for all	least 30 days		
the study group the patient belonged to.	alion.		
Overall quality assessment (descriptive)			
Fair			
RESULTS			
Outcome Intervention group Comparator group Statistical sign	ificance		
Thromoboelastography Clinician directed transfusion (CDT)			
(kTEG & h-kTEG assays) (tests=PT, aPTT, platelets) N=114 N=110			
Mean (SD) mediastinal chest tube drainage, mL 480.5 (351) 591.4 (339.2) P=0.087			
Early mortality 3 patients (2.7%) 2 patients (1.7%) Not reported			
(defined as death within 30 days of operation)(low cardiac output=2, multiple organ failure=1)(mediastinitis=1, respiratory insufficiency=1)			
Clinical importance Clinical relevance	Clinical relevance		
Reported by Authors as 'no difference' 1			
Re-exploration for bleeding 6 patients 5 patients Not reported			
(causes all surgical) (Causes, 2= surgical, 3 inappropriate surgical intervention for bleeding)			

Clinical importance		Clinical relevance		
Reported by Authors as 'no difference'		1		
Patients transfused with PRBC	52/114 (45.6%)	60/110 (54.5%)	P=0.181	
Patients transfused with FFP	19/114 (16.6%)	31/110 (28.1%)	P=0.038	
Patients transfused with platelets	17/114 (14.9%)	29/110 (26.3%)	P=0.033	
Median (IQR) units of PRBCs transfused intraoperatively	1 (0, 1)	0 (0, 1)	P=0.581	
Median (IQR) units of PRBCs transfused postoperatively	1 (0, 1)	1 (0, 1)	P=0.741	
Median (IQR) units of PRBCs transfused both intra-and postoperatively	1 (0, 1)	1 (1, 2)	P=0.599	
Clinical importance		Clinical relevance		
Median (IQR) units of FFP transfused intraoperatively	0 (0, 1)	1 (0, 1)	P=0.008	
Median (IQR) units of FFP transfused postoperatively	1 (0, 1)	1 (0, 1)	P=0.034	
Median (IQR) units FFP transfused both intra- and postoperatively	1 (1, 1)	1 (1, 2)	P=0.001	
Clinical importance		Clinical relevance		
Median (IQR) units of platelets transfused intraoperatively	0(0, 1)	1 (0, 1)	P=0.004	
Median (IQR) units of platelets transfused postoperatively	1 (0, 1)	1 (0, 1)	P=0.028	
Median (IQR) units of platelets transfused both intra- and postoperatively	1 (1, 1)	1 (1, 2)	P=0.001	
1	l .	1		
Median (IQR) allogeneic units transfused	2 (1-3)	3 (2-4)	P=0.001	
(PRBC, FFP, and platelets)				
Clinical importance				
1		1		
EXTERNAL VALIDITY				
Generalisability				

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The study population were all undergoing surgery i.e. were the same as the target population. Two relevant groups were compared i.e. one undergoing POC testing and one undergoing clinician determined transfusion based on conventional lab tests. However, eligibility criteria prevented the inclusion of the full range of patients that may be expected in a normal practice.

Applicability

Reduced, study performed in Istanbul Turkey – study population characteristics and healthcare system likely to differ from that in Australia/New Zealand. This patient population was undergoing elective surgery for first time CABG with cardiopulmonary bypass and was without severe co-morbidities.

Comments

FU period was not clear and at 30 days there was no difference in early clinical outcome – late clinical outcomes not reported. The two groups were not completely matched for tests which may have favoured the TEG group.

Summary

Differentiation between surgical and non-surgical bleeding in patients with excessive MCTD was more successful in the TEG group and there were fewer transfusion units (FFP and platelets) required in the TEG group. There were no differences between the two groups in terms of consumption of red blood cells (packed cells), no of patients re-explored for bleeding, and early mortality.

Citation

Avidan M.S., Alcock E.L. et al. Comparison of structured use of routine laboratory tests or near-patient assessment with clinical judgement in the management of bleeding after cardiac surgery. British Journal of Anaesthesia. 2004; 2:176-86.

Affiliation/Source of funds

Department of Anesthesiology and Blood Bank, Washington University School of Medicine, St Louis, USA.

Departments of Anaesthesia and Cardiothoracic Surgery, King's College Hospital, London, UK.

Department of Haematology, Guy's, King's and St Thomas' Medical School (GKT), London, UK

The Royal College of Anaesthetists (British Journal of Anaesthesia Small Project Grant, Oxford, UK) and the National Blood Services (London, UK) provided financial support for this research. Medicell UK provided thromboelastography consumables at discounted prices and lent the investigators a thromboelastography machine. Medtronic provided the consumables for the Hepcon machine for the purposes of the study. Additional funds were raised by the Academic Department of Anaesthesia, King's College London.

Study design	Level of evidence	9	Location/setting		
Patients were randomised into two groups (point of care vs laboratory test). Blood loss and transfusion was compared between these two groups and with a retrospective case-control group, in which management of bleeding had been according to the clinician's discretion.	II and III (depending on comparator)		UK		
Intervention	Intervention		Comparator		
Algorithm based on near-patient haemostatic testing (v. clinical algorithm). POC devices used include ACT+/Junior, Hepcon HMS Hemostasis Management System, PFA-100 platelet function analyser; and two dual-channel TEG coagulation analysers used in parallel. N=51		Randomised comparator: algorithm using routine laboratory haemostatic tests. N=51 Historical comparator: Adults who had undergone routine CABG surgery with the same clinical team over a four-month period preceding the interventional study. They had received blood components on the basis of individual clinician's discretion. N=108			
Population characteristics		1			
Adults undergoing elective, first-time CA	ABG with CPB				
Length of follow-up		Outcomes measured			
24 hours		Transfusion (incidence), transfusion (volume), Blood loss, Haemoglobin concentration, platelet count			

INTERNAL VALI	DITY						
Allocation	Results		Blinding analys	is	Treatment/measuren bias	nent	Follow-up (ITT)
It is not clear whether allocation was concealed from those responsible for recruiting subjects.	The treatm had similar characteris	baseline	The study was n blinded	ot	None detected		No patients were lost to follow-up.
Overall quality a	ssessment	(descriptive))				
Fair							
RESULTS							
Outcome		Intervention	on group		nparator group ntrol)	Sta	tistical significance
Patients transfuse	ed with	POC: 34/5	1 (67%)	Lab	oratory: 35/51 (69%)	Chi-	-square test: P=0.01

RESULTS				
Outcome	Intervention group	Comparator group (Control)	Statistical significance	
Patients transfused with PRBCs	POC: 34/51 (67%)	Laboratory: 35/51 (69%) Clinician discretion: 92/108 (85%)	Chi-square test: P=0.01	
Patients transfused with FFP	POC: 2/51 (4%)	POC: 2/51 (4%) Laboratory: 0/51 (0%) Clinician discretion: 16/108 (15%)		
Patients transfused with platelets	nsfused with POC: 2/51 (4%) Laboration Clini (13%)		Chi-square test: P=0.02	
Median (IQR) 24 hour postoperative blood loss, mL			NR	
Median (IQR) postoperative haemoglobin concentration, g/dL	POC: 9.3 (8.4, 10.3)	Laboratory: 9.3 (8.5, 9.7) Clinician discretion: Not available	NR	
Median (IQR) postoperative 24 hour haemoglobin, g/dL			NR	
Median (IQR) postoperative platelet count, X109/L	POC: 131 (110, 165)	Laboratory: 140 (111, 168) Clinician discretion: Not available	NR	
Median (IQR) postoperative 24 hour platelet count, X10 ⁹ /L	POC: 149 (123, 187)	Laboratory: 159 (135, 200) Clinician discretion: 144 (121, 174)	NR	
Total units of PRBCs transfused	POC: 99	Laboratory: 93 Clinician discretion: 285	NR	
Median (IQR) volume of PRBCs transfused, mL		Laboratory: 495 (0, 612) Clinician discretion: 512 (286, 962)	Kruskal-Wallis ANOVA: P=0.03	

Total units of platelets transfused	POC: 3	Laboratory: 2 Clinician discretion: 14	NR				
Total units of FFP transfused	POC: 6	Laboratory: 0 Clinician discretion: 65	NR				
Reoperation for bleeding	POC: 1/51 (2%)	Laboratory: 1/51 (2%) Clinician discretion: 3/108 (3%)	POC vs. Laboratory RR (95% CI): 1.00 (0.06, 15.56); P=1.00 POC vs. clinician discretion RR (95% CI): 0.71 (0.08, 6.62); P=0.76				
EXTERNAL VALIDITY							
Generalisability							
The study was conducted in a surgeries.	adults undergoing CABG with CP	B but it may be somewhat gener	ralisable to other elective				
Applicability							
The study was conducted in t	he UK, however it is likely to also	be applicable to the Australian	context.				
Comments							

Citation

Royston D. and von Kier S. Reduced haemostatic factor transfusion using heparinise-modified thromboelastography during cardiopulmonary bypass. British Journal of Anaesthesia. 2001; 4:575-8.

Affiliation/Source of funds

Department of Anaesthesia and Critical Care, Royal Brompton and Harefield NHS Trust, Harefield Hospital, Harefield, Middlesex, UK

Study design	Level of evidence	Location/setting
RCT	II	UK
Intervention	Cou	mnarator

Intervention	Comparator
Heparinase-modified thromboelastogram guided intraoperative algorithm N=30	Clinical criteria and laboratory-based tests N=30

Population characteristics

Adults undergoing cardiac surgery. Ten per cent of the patients in each series had a heart transplantation and were taking aspirin and/or warfarin immediately before surgery. About 50% of the patients in each group had revascularisation and were also taking aspirin, and required multiple grafts with a bypass time estimated to be greater than 100 minutes. The remaining 40% of the patients were having the Ross procedure, multiple valve or valve and revascularisation surgery. No patients were having repeat operations and none received prophylactic aprotinin, epsilon aminocaproic or tranexamic acid.

Length of follow-up	Outcomes measured		
12 hours	Transfusion (incidence), transfusion (volume), blood loss		

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Allocation was concealed from those responsible for recruiting subjects	Patient details were not significantly different between groups but showed a wide distribution of values for age (21 to 83 years) and bypass time (48 to 167 minutes)	The study was not blinded	None detected	There was no loss to follow-up

Overall quality assessment (descriptive)

Poor

RESULTS

Outcome	Intervention group	Comparator group (Control)	Statistical significance
Patients transfused with blood components	5/30 (17%)	10/30 (33%)	P<0.05
Volume of blood components transfused	Five units of FFP and one pool of platelets	16 units of FFP and nine platelet pools	P<0.05
Median (IQR) 12 hour chest tube loss, mL	470 (295, 820)	390 (240, 820)	NR

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Generalisability

The study was conducted in adults undergoing cardiac surgery, but the results are likely to be generalisable to other elective surgeries

Applicability

The study was conducted in the UK, and is likely to be applicable to the Australian context.

Comments

Citation

Shore-Lesserson L., Manspeizer H.E. et al. Thromboelastography-guided transfusion algorithm reduces transfusions in complex cardiac surgery. Anesthesia and analgesia. 1999; 88:312-9.

Affiliation/Source of funds

Departments of Anesthesiology and Cardiothoracic surgery, Mount Sinai Medical Center, New York, New York.

Study design	Level of evidence	Location/setting
RCT	II	USA hospital

Intervention

TEG-guided transfusion algorithm. Data from the celiteactivated TEG was used to guide transfusion therapy 10 minutes after protamine completion. Transfusion therapy was prescribed in the presence of bleeding:

- Additional protamine (50 mg) was given if the heparinise modified TED R time was less than one half of the non-heparinise R time.
- If bleeding persisted, 6 U of platelets was transfused if platelet count <100,000/µL and TEG MA <45mm.
- 3. If bleeding persisted, 2 U of FFP was given if R time was > 20 mm.
- 4. If bleeding persisted, 10 U of cryoprecipitate was transfused if fibrinogen level <100 mg/dL.
- 5. If bleeding persisted and if the TEG showed evidence of fibrinolysis (LY30 > 7.5%), additional antifibrinolytic therapy (EACA 10g) was given at the discretion of the physicians caring for the patient. In both groups, if a patient received a transfusion, the abnormal tests were repeated and treated in accordance with the algorithm as long as the patient was still in the operating room.

Comparator

Standard laboratory-based transfusion therapy (TEQ). Data from laboratory-based tests were used to guide transfusion therapy 10 minutes after protamine completion:

- Additional protamine (50 mg) was given if ACT exceeded baseline by 15%.
- If bleeding persisted, 6 U of platelets was transfused if platelet count <100,000/µL.
- 3. If bleeding persisted, 2 U of FFP was given if PT > 150% of control.
- 4. If bleeding persisted, 10 U of cryoprecipitate was transfused if fibrinogen level <100 mg/dL.
- If bleeding persisted and if above therapy failed to reduce bleeding, an additional bolus of antifibrinolytic therapy (EACA 10g) was given at the discretion of the physicians caring for the patient.

(N=52)

(N=53)

Population characteristics

Adults undergoing a cardiac surgical procedure with a moderate to high risk for requiring a transfusion.

Length of follow-up	Outcomes measured
Two days postoperative	Mortality, reoperation for bleeding, Morbidity, clotting time, platelet count, prothrombin time, activated partial thromboplastin time, fibrinogen concentration, transfusion (incidence), transfusion (volume)

Allocation	Results		Blinding analysis	•	Treatment/measurem	nt	Follow-up (ITT)
Randomisation achieved via a table of random numbers. It is unclear whether allocation was blinded from those responsible for recruiting subjects.	The baseline characteristic similar betwee treatment and	cs were een ms.	The anaesthesiolo and surgeon caring for the patient were blinded to the patient's group assignment. All intraoperative result of the TEG and laboratory coagulatests were interpreted by an anaesthesiologist investigator not directly involved with patient's care. recommended therapy according the patient's group assignment was communicated to the anaesthesiologist as surgeon by this investigator as appropriate.	g e late and the l	None detected		Two patients in the TEG group were not included in follow-up analyses. One patient enrolled but not studied was undergoing cardiac reoperation and was placed emergently on CPB because of massive hemorrhage during sternotomy. The patient was excluded from the study at theis time. The other patient who did not complete the protocol was excluded due to a severe protamine reaction that required immediate reinstitution of CPB.
Overall quality a	ssessment (d	iescriptive)					
RESULTS							
Outcome		Interventio	tion group Cor (Co		parator group rol)	Stati	istical significance
Mortality (ITT)	(ITT) 0/53 (0%) 2/52 (4%)1	Revl	Man: RR (95% CI): 0.20		

KESUL13					
Outcome	Intervention group	Comparator group (Control)	Statistical significance		
Mortality (ITT)	0/53 (0%)	2/52 (4%)1	RevMan: RR (95% CI): 0.20 (0.01, 3.99); P=0.29		
Reoperation for bleeding (ITT)	0/53 (0%)	2/52 (4%) ²	RevMan: RR (95% CI): 0.20 (0.01, 3.99); P=0.29		
Cerebrovascular ischemic event (ITT)	1/53 (2%)	0/52 (0%)	RevMan: RR (95% CI): 2.94 (0.12, 70.67) P=0.51		
Mean (SD) activated clotting time ³ (baseline), seconds	165 (34)	170 (49)	Instat: Mean difference (95% CI): 5.0 (-11.5, 21.5); P=0.55		
Mean (SD) activated clotting time ³ (post-protamine), seconds	158 (93)	149 (20)	Instat: Mean difference (95% CI): -9.0 (-35.2, 17.2); P=0.50		
Mean (SD) platelet count ⁴ (baseline), X1000/μL	203 (66)	200 (78)	Instat: Mean difference (95% CI): -3.0 (-31.3, 25.3); P=0.83		

Mean (SD) platelet count ⁴ (warming on CPB), X1000/μL	92 (79)	96 (79)	Instat: Mean difference (95% CI): 4.0 (-26.9, 34.9); P=0.80
Mean (SD) platelet count ⁴ (ICU), X1000/μL	111 (48)	120 (48)	Instat: Mean difference (95% CI): 9 (-9.8, 27.8); P=0.34
Mean (SD) prothrombin time ⁵ (baseline), seconds	13.0 (1.1)	12.9 (1.3)	Instat: Mean difference (95% CI): -0.1 (-0.6, 0.4); P=0.67
Mean (SD) prothrombin time ⁵ (post-protamine), seconds	18.1 (2.3)	21.3 (26)	Instat: Mean difference (95% CI): 3.2 (-4.1, 10.5); P=0.38
Mean (SD) prothrombin time ⁵ (ICU), seconds	16.1 (1.7)	15.7 (1.6)	Instat: Mean difference (95% CI): -0.4 (-1.0, 0.2); P=0.22
Mean (SD) activated partial thromboplastin time ⁶ (baseline), seconds	31.6 (6.9)	34.1 (13.1)	Instat: Mean difference (95% CI): 2.5 (-1.6, 6.6); P=0.23
Mean (SD) activated partial thromboplastin time ⁶ (post-protamine), seconds	52.2 (48.0)	43.0 (14)	Instat: Mean difference (95% CI): -9.2 (-23.0, 4.6); P=0.19
Mean (SD) activated partial thromboplastin time ⁶ (ICU), seconds	35.9 (6.1)	36.8 (10.2)	Instat: Mean difference (95% CI): 0.9 (-2.4, 4.2); P=0.59
Mean (SD) fibrinogen concentration ⁷ (baseline), mg/dL	409 (82)	416 (118)	Instat: Mean difference (95% CI): 7 (-32.8, 46.8); P=0.73
Mean (SD) fibrinogen concentration ⁷ (post-protamine), mg/dL	239 (86)	246 (86)	Instat: Mean difference (95% CI): 7.0 (-26.6, 40.6); P=0.68
Mean (SD) fibrinogen concentration ⁷ (ICU), mg/dL	259 (95)	263 (118)	Instat: Mean difference (95% CI): 4.0 (-37.9, 45.9); P=0.85
Patients transfused with allogeneic blood components (total)	22/53 (42%)	34/52 (65%)	P=0.01
Patients transfused with packed red blood cells (intraoperative)	17/53 (32%)	23/52 (44%)	P=0.2
Patients transfused with packed red blood cells (postoperative)	10/53 (19%)	16/52 (31%)	P=0.16
Patients transfused with packed red blood cells (total)	22/53 (42%)	31/52 (60%)	P=0.06
Patients transfused with FFP (intraoperative)	3/53 (6%)	8/52 (15%)	P=0.1
Patients transfused with FFP (postoperative)	2/53 (4%)	11/52 (21%)	P<0.007
Patients transfused with FFP (total)	4/53 (8%)	16/52 (31%)	P=0.002

Patients transfused with platelet concentrates (intraoperative)	5/53 (9%)	8/52 (15%)	P=0.4
Patients transfused with platelet concentrates (postoperative)	3/53 (6%)	9/52 (17%)	P=0.06
Patients transfused with platelet concentrates (total)	7/53 (13%)	15/52 (29%)	P<0.05
Mean (SD) volume of PRBCs transfused (intraoperative), mL	267 (423)	346 (449)	P=0.4
Mean (SD) volume of PRBCs transfused (postoperative), mL	103 (252)	177 (318)	P=0.27
Mean (SD) volume of PRBCs transfused (total), mL	354 (487)	475 (593)	P=0.12
Mean (SD) volume of FFP transfused (intraoperative), mL	22 (101)	113 (407)	P=0.4
Mean (SD) volume of FFP transfused (postoperative), mL	33 (169)	146 (378)	P=0.13
Mean (SD) volume of FFP transfused (total), mL	36 (142)	217 (463)	P<0.04
Mean (SD) volume of platelet concentrates transfused (intraoperative), mL	22 (75)	41 (122)	P=0.6
Mean (SD) volume of platelet concentrates transfused (postoperative), mL	11 (46)	42 (107)	P=0.3
Mean (SD) volume of platelet concentrates transfused (total), mL	34 (94)	83 (160)	P=0.16
Mean (SD) six-hour mediastinal drainage, mL	362 (274)	469 (637)	P=0.63
Mean (SD) 24-hour mediastinal drainage, mL	702 (500)	901 (847)	P=0.27
		•	•

Generalisability

The study is in patients undergoing cardiac surgery, but the results are somewhat generalisable to other surgery types.

Applicability

The study was conducted in New York, and although there are differences in health care systems, the results are likely to be applicable to the Australian context.

Comments

- ¹ Both patients died from hemodynamic causes in the postoperative period.
 ² In one patient, a specific surgical source of bleeding was discovered.
 ³ Normal range: 90 to 120 seconds
 ⁴ Normal range: 120 to 500 X1000/µL
 ⁵ Normal range: 12 to 14 seconds
 ⁶ Normal range: 25 to 34 seconds
 ⁷ Normal range: 150 to 500 mg/dL

Citation

Westbrook AJ., Olsen J. et al. Protocol based on thrombolestaograph (TEG) out-performs physician preference using laboratory coagulation tests to guide blood replacement during and after cardiac surgery: a pilot study. Heart, Lung and Circulation.2009;18:277-288.

Affiliation/Source of funds

Department of Intensive Care and Hyperbaric Medicine, Alfred Hospital, Melbourne, Australia.

Department of Epidemiology and Preventive Medicine, Monash University, Melbourne, Australia.

Department of Surgery, Monash University, Melbourne Australia.

Funding for the salary of the research nurse was provided by Ventracor PTY LTD, and Medtel PTY LTD, Australia.				
Study design	Level of evidence		Location/setting	
RCT	II		Australia, Hospital	
Randomisation method not reported				
Intervention		Comparator		
TEG (N=32)		Physician directed pro	oduct administration (control) (N=37)	
Time taken before clot formation begins	;	With reference to labor	oratory coagulation tests	
Time taken for clot to form		APTT		
Maximum strength of the clot	f the clot •		• INR	
Clot strength maintenance and clot lysis	S	Fibrinogen and platelet count		
		and physician's previous experience.		
A strict protocol for administration of blo used based on a computerised thromb coagulation analyser (Haemoscope Cor	oelastograph			
Population characteristics				
All patients presenting for cardiac surge	ry with the exceptio	n of lung transplantation	n were included (N=69)	
Heparin for cardiopulmonary bypass wa	s administered acco	ording to standard activ	ated clotting time in both groups	
Length of follow-up		Outcomes measured		
Not reported but at least until discharge	(7-13 days)	Re-sternotomy,		
		Minimum haemoglobin,		

Ttransfusion of RBC,PLTS,FFP,CRYO

Intubation time

ICU stay

INTERNAL VALI	DITY						
Allocation	Results		Blinding analysis	6	Treatment/measureme bias	nt	Follow-up (ITT)
Randomisation (method not reported)	Both group satisfactoril (i.e. ns difference most meast characteristic including agages for content of the first surgery who mostly content of the TEG group and present the TEG group of the TEG group of the test of the TEG group of the test of th	y matched erences) by ured tics ge (mean ntrol and and 66 gender 81%), re aspirin rin (5% arfarin 6 %) and (3% and otions were edo" ich was fined to the (p = 0.04) onate bypass higher in	Surgeons were blinded to the met of haemostasis. It is not stated whomade the decision transfuse/assess other outcomes at they were blinded the patient's group allocation, however appended protocoflow diagrams for various protocols used suggest that decision to transful was not blinded to intervention group	on to nd if to o er ol the the	A potential confounder in TEG group was reported be the "Hawthorn effect" which may have exaggerated the trend towards a better outcome this group.	to 7	All patients included in the analysis.
Overall quality a	issessment (aescriptive)					
Fair							
RESULTS							
Outcome		TEG N=32		(Cor Phys adm to Al	nparator group Introl) Sician directed product Inistration with reference PTT, INR, fibrinogen and Elet count	Stat	istical significance
Units of blood products transfused intraoperatively		44		ns (p	o value not reported)		
Units of blood pro transfused in ICU		18		46		ns (p	o value not reported)
Total units of bloc transfused	products	37		90		ns (p	o value not reported)

 7 A term referring to the tendency of some people to work harder and perform better when they are participants in an experiment.

Clinical importance		Clinical relevance		
Units of PRBCs transfused intraoperatively	11	15	ns (p value not reported)	
Units of PRBCs transfused in ICU	3	18	ns (p value not reported)	
Total units of PRBCs transfused	14	33	ns (p value not reported)	
Clinical importance		Clinical relevance		
2		1		
Units of FFP transfused intraoperatively	8	14	ns (p value not reported)	
Units of FFP transfused postoperatively	10	8	ns (p value not reported)	
Total units of FFP transfused	18	22	ns (p value not reported)	
Clinical importance 2		Clinical relevance		
Units of platelets transfused intraoperatively	0	10	ns (p value not reported)	
Units of platelets transfused postoperatively	5	5	ns (p value not reported)	
Total units of platelets transfused	5	15	ns (p value not reported)	
Clinical importance		Clinical relevance		
2		1		
Units of cryoprecipitate transfused intraoperatively	0	5	ns(p value not reported)	
Units of cryoprecipitate transfused postoperatively	0	15	ns(p value not reported)	
Total units of cryoprecipitate transfused	0	20	ns(p value not reported)	
Clinical importance		Clinical relevance		
2		1		
Median Blood loss in mls (25th & 75th percentile)	875 (755-1130)	960 (820-1200)	ns (p = 0.437)	
Clinical importance		Clinical relevance		
Median intubation time in hours 25th & 75th percentile)	8 (5.3-19.8)	10.3 (5.8-19.5)	ns (p value not reported)	
Clinical importance		Clinical relevance		
Median minimum (IQR) haemoglobin concentration, g/I	87 (83-94)	86 (82-104)	ns (p value not reported)	

Clinical importance		Clinical relevance	
3		1	
Median (IQR) length of ICU stay, hours	29.4 (14.3, 56.4)	32.5 (22, 74.5)	ns (p value not reported)
Median (IQR) length of hospital stay, days	9 (7-13) *Extra day not due to bleeding	8 (7-12)	ns (p value not reported)
Clinical importance		Clinical relevance	

Generalisability

Good-surgical population similar to target guideline population, POC compared to a relevant/appropriate control.

Applicability

High- the study reports a cohort of patients undergoing cardiac surgery in a large Australian hospital.

Comments

The authors point to an 'impressive reduction (52% per patient) in blood products administered peri operatively' and a 'clinically significant reduction in the administration of blood products'. Small group size and non-parametric testing and a high proportion of redo study patients (18.7%) in the TEG group were reported as possible reasons for the failure of the differences between the two study groups to reach significance. However, all outcome parameters (with the exception of the hospital LOS) were consistently more favourable for the TEG group.

This was a pilot study – a larger RCT was planned but slow recruitment to the pilot study and lack of funds precluded a larger RCT (personal communication from Professor Bob Salamonsen 12th March 2010.8

Note: ICU results not included.

⁸ 'After the pilot trial for the TEG (westbrook et al.) we did not perform or plan a followup trial. We were troubled by a very low recruitment rate and obtained only the patients reported in the paper during a whole year - hence our decision to call it a pilot trial. After one year the money dried up'.

Level III evidence

Citation

Spalding, GJ., Hartrumpf, M. et al. Cost reduction of peri operative coagulation management in cardiac surgery: value of "bedside" thrombelastography (ROTEM). European Journal of Cardiothoracic Surgery 2007;31:1052-1057.

Affiliation/Source of funds

Department of Cardiovascular Surgery, Heart Center Brandenburg, Bernau/Berlin.

Department of Anaesthesiology, Heart Center Brandenburg, Bernau/Berlin.

Conflict of interest/ source of funds =None declared

Study design	Level of evidence	Location/setting
Before and after cohort design, single	III-3	Germany, Hospital,
institution		

Intervention	Comparator
ROTEM ⁹ (bedside instrument measuring clot formation and dissolution indicating changes in coagulation, platelet function, platelet-fibrinogen interaction and fibrinolysis). N=693	No ROTEM N=729

Population characteristics

All patients in a single institution undergoing cardiosurgical therapy over a 12 month period. Age 67 years \pm 8-9years, males 70.6%-74.5%.

Length of follow-up	Outcomes measured	
1 year	Blood consumption	
	Blood product consumption	
	Coagulation product consumption	
	Cumulative costs	
	Operative risk/early mortality	
	Early Resternotomy	

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Within a 1 year period (exact dates not given)	The cohort did not all receive the same surgical procedure.	Not reported	Cannot be ruled out - time periods were different.	All study patients were reported no losses to follow up.
Patients undergoing surgery in the first six months were assigned to the No ROTEM control group. Patients	71% of the comparator and 72% of the ROTEM group had isolated coronary artery revascularization (CABG) 3.3% of the comparator and 1.9% of the ROTEM group had CABG without cardiopulmonary bypass			

⁹ ROTEM® stands for rotation thromboelastometry and is an enhancement of classical thromboelastography, which is a technique for the assessment of blood coagulation disorders.

undergoing surgery in the second six months were assigned to the ROTEM intervention group.	15% of the comparate and 12% of the ROTE group had isolated vasurgery 9.5% of the comparate and 10.6% of the ROTEM group had CABG + valve surger	EM lve or			
	2.9% of the comparat and 3.5% of the ROT group had aortic surg 1.4% of the comparat and 1.6% of the ROT group had other measures	EM ery or			
	Demographic data between the two groudid not differ statistica in terms of age (mear age =67yrs), gender (74% male) surgery ty early mortality and ear resternotomy for bleeding. The ROTEN group had a significar higher EuroSCORE (p=0.006).	ally n (71- pe, rly / /			
	ssessment (descriptiv	ve)			
Fair					
RESULTS Outcome		Intervention group	Con	nparator group	Statistical significance
Outcome		ROTEM (second six month period)		ROTEM (first six month	Statistical significance
Red blood cell un (€ 70.00)	it	368	439		ns
Clinical importar	nce		Clin	ical relevance	
3			1		
Platelet concentrate unit 28 (€ 500.0)		28	59		p = 0.000
Clinical importance		Clinical relevance			
1		1			
Fresh frozen plas (€ 51.00)	Fresh frozen plasma unit 116 (€ 51.00)		118		ns
Clinical importar	Clinical importance			ical relevance	
3			1		

¹⁰ Euro SCORE is a method of calculating predicted operative mortality for patients undergoing cardiac surgery.

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Pooled coagulation concentrates - 500 units (€120)	27	130	p = 0.000	
Clinical importance		Clinical relevance		
1		1		
Fibrinogen (1 g, €287.50)	55	14	ns (0.060)	
Clinical importance		Clinical relevance		
3		1		
rFactor VIIa (120 IU, € 1512.00)	1	11	p= 0.000	
Clinical importance		Clinical relevance		
1		1		
Factor XIII (1250 IU,€ 405.00)	8	17	p = 0.001	
Cumulative monthly costs total	€55,925	€125,828	-44%	
Cumulative monthly costs blood	€45,000	€66.000	-32%	
products	€30,000	€60,000	-50%	
Cumulative monthly costs coagulation factor				
Clinical importance	•	Clinical relevance		
Not reported		Not relevant		
Mean EuroSCORE (SD)	5.5 ± 3.1	5.0 ±3.3	p=0.006	
(European System for Cardiac Operative Risk Evaluation)				
Clinical importance		Clinical relevance		
1		3		
Early resternotomy (%)	5.5	6.6	ns (0.384)	
Clinical importance		Clinical relevance		
3		1		
Early mortality (%)	6.0	5.9	ns	
Clinical importance	•	Clinical relevance		
3		1		
EXTERNAL VALIDITY				

Generalisability

The patient population were undergoing cardiac surgery i.e. they were part of the guideline target population, a POC system was the focus of the cost assessment and the comparator was normal clinical practice prior to the implementation of the ROTEM system.

Applicability

In general the results have reasonable applicability to any large specialist hospital carrying out this type of surgery. However, the cost analysis was representative only of the health system in which the study was carried out and may not pertain to other countries or health systems. Moreover, a full cost-effectiveness analysis was not carried out.

Comments

ROTEM was performed with a blood loss of over 200ml/h which did not cease after 2 hr. ROTEM was not performed in the majority of patients (65%) who presented with a regular postoperative drainage loss. The study is subject to the limitations of pre-post/before-after studies. Cost analysis was very rudimentary- only direct cost reductions were considered.

Intervention 8 - Administration of antifibrinolytics & DDAVP

Level I evidence

STUDY DETAILS	STUDY DETAILS							
Citation								
	nuna. J Wona ((2009) T	opical application o	of ant	tifibrinolytic druas for	r on	n-pump cardiac surgery:	a systematic
review and meta-a					zz a. a. a. a. ge re.		. pamp saraias sargery.	a ojotomano
Affiliation/Source	e of funds							
							nto, Ontario, Canada.	
	ent of Anesthe	esia, Toi			University Health No			
Study design			Level of evidence	е			cation/setting	
Systematic review			Level I			Ho	spital	
that investigated t								
application of antible blood loss and tra		ys on						
requirements in pa		oina						
on-pump cardiac l								
Intervention	bypass surgery	у.		Co	mparator			
Topical application	n of aprotinin o	or tranex	ramic acid		acebo or active			
Poured into perica								
tissues at the end								
median sternotom				L				
Population chara								
Patients undergoi	ng on-pump ca	ardiac sı	urgery (primary ope	ration	n only)			
Length of follow-	-up				itcomes measured			
Not specified							loss, units of allogenic R	
				trar	nsfusion, incidence (of a	allogenic RBC transfusion	า
INTERNAL VALIDITY								
					T			
Allocation	Results				Blinding analysis		Treatment/ measurement bias	Follow-up (ITT)
Allocation Randomised	Results Baseline cha		tics of intervention		Five studies were		measurement bias Egger's test for bias	(ITT) States all 8
Allocation Randomised allocation was	Results Baseline cha and control g	groups n	ot reported.		Five studies were double blinded.		measurement bias Egger's test for bias was non-significant	(ITT) States all 8 studies had
Allocation Randomised allocation was reported in all	Results Baseline cha and control g Methodologic	groups n cal quali	ot reported. Ity of studies		Five studies were double blinded. One study was	,	measurement bias Egger's test for bias was non-significant (p=0.6 and p=0.5) for	States all 8 studies had "adequate"
Randomised allocation was reported in all eight studies.	Results Baseline cha and control g Methodologic assessed ind	groups n cal quali depende	ot reported. Ity of studies ently by two authors.		Five studies were double blinded. One study was not blinded.	,	measurement bias Egger's test for bias was non-significant (p=0.6 and p=0.5) for reduction of blood loss	(ITT) States all 8 studies had
Randomised allocation was reported in all eight studies. Method of	Results Baseline cha and control g Methodologic assessed ind Disagreemer	groups n cal quali depende nts resol	ot reported. Ity of studies ently by two authors. Ived by a third autho	or.	Five studies were double blinded. One study was not blinded. The blinding	,	measurement bias Egger's test for bias was non-significant (p=0.6 and p=0.5) for reduction of blood loss or amount of blood	States all 8 studies had "adequate"
Randomised allocation was reported in all eight studies. Method of randomisation	Results Baseline cha and control g Methodologic assessed ind Disagreemer Data extractic	groups n cal quali depende nts resol ion for e	ot reported. Ity of studies ently by two authors. Ived by a third autho ach study not stated	or.	Five studies were double blinded. One study was not blinded. The blinding status of two	,	measurement bias Egger's test for bias was non-significant (p=0.6 and p=0.5) for reduction of blood loss or amount of blood transfusion,	States all 8 studies had "adequate"
Randomised allocation was reported in all eight studies. Method of randomisation reported in five	Results Baseline cha and control g Methodologic assessed ind Disagreemer Data extractic A random eff	groups n cal quali depende nts resol ion for e fects mo	ot reported. Ity of studies ently by two authors. Ived by a third authorach study not stated odel was use in the	or.	Five studies were double blinded. One study was not blinded. The blinding status of two studies was	,	measurement bias Egger's test for bias was non-significant (p=0.6 and p=0.5) for reduction of blood loss or amount of blood transfusion, suggesting that there	States all 8 studies had "adequate"
Randomised allocation was reported in all eight studies. Method of randomisation	Results Baseline cha and control g Methodologic assessed ind Disagreemer Data extractic A random eff meta-analyse	groups n cal quali depende nts resol ion for e fects mo es as all	ot reported. Ity of studies ently by two authors. Ived by a third autho ach study not stated	or.	Five studies were double blinded. One study was not blinded. The blinding status of two		measurement bias Egger's test for bias was non-significant (p=0.6 and p=0.5) for reduction of blood loss or amount of blood transfusion, suggesting that there was no publication	States all 8 studies had "adequate"
Randomised allocation was reported in all eight studies. Method of randomisation reported in five studies.	Results Baseline cha and control g Methodologic assessed ind Disagreemer Data extractic A random eff meta-analyse heterogeneit	groups n cal quali depende nts resol ion for e fects mo es as all y.	ot reported. Ity of studies Intly by two authors. Ived by a third authorach study not stated I showed statistical	or.	Five studies were double blinded. One study was not blinded. The blinding status of two studies was		measurement bias Egger's test for bias was non-significant (p=0.6 and p=0.5) for reduction of blood loss or amount of blood transfusion, suggesting that there	States all 8 studies had "adequate"
Randomised allocation was reported in all eight studies. Method of randomisation reported in five studies. Overall quality as	Results Baseline cha and control g Methodologic assessed ind Disagreemer Data extractic A random eff meta-analyse heterogeneityssessment (de	groups n cal quali depende nts resol ion for e fects mo es as all y. escripti	ot reported. ity of studies ently by two authors. lved by a third author ach study not stated del was use in the showed statistical ive)	or. d.	Five studies were double blinded. One study was not blinded. The blinding status of two studies was unclear.		measurement bias Egger's test for bias was non-significant (p=0.6 and p=0.5) for reduction of blood loss or amount of blood transfusion, suggesting that there was no publication bias.	States all 8 studies had "adequate" follow-up
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Allocation Randomised allocation was reported in all eight studies. Method of randomisation reported in five studies. Overall quality as Good. This review strategy employed	Results Baseline cha and control g Methodologic assessed ind Disagreemer Data extractic A random eff meta-analyse heterogeneity ssessment (do clearly defined appeared robolided study w	groups no cal qualidepende nts resolution for effects modes as all y. escription de the repost, and was provinced to the repost, and the repost of the re	ot reported. Ity of studies ently by two authors. Ived by a third author ach study not stated adel was use in the showed statistical Ive) search question, so d the methods for an ided. This review pr	or. d. cope, nalys	Five studies were double blinded. One study was not blinded. The blinding status of two studies was unclear. search terms and ir sis were appropriate	nclu	measurement bias Egger's test for bias was non-significant (p=0.6 and p=0.5) for reduction of blood loss or amount of blood transfusion, suggesting that there was no publication bias. usion/exclusion criteria. The summary, as well as a	States all 8 studies had "adequate" follow-up
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Randomised allocation was reported in all eight studies. Method of randomisation reported in five studies. Overall quality as Good. This review strategy employed rating, for each inc meta-analysis of t	Results Baseline cha and control g Methodologic assessed ind Disagreemer Data extractic A random eff meta-analyse heterogeneity ssessment (do clearly defined appeared roboluded study whe data from in	groups n cal quali depende nts resol ion for e fects mc es as all y. escripti ed the re bust, and was prov ncluded	ot reported. Ity of studies Ently by two authors. Ived by a third author ach study not stated adel was use in the showed statistical Ive) Is search question, so id the methods for an ided. This review pr	or. d. cope, nalys	Five studies were double blinded. One study was not blinded. The blinding status of two studies was unclear. search terms and ir sis were appropriate	nclu e. Tr	measurement bias Egger's test for bias was non-significant (p=0.6 and p=0.5) for reduction of blood loss or amount of blood transfusion, suggesting that there was no publication bias. usion/exclusion criteria. The summary, as well as a	States all 8 studies had "adequate" follow-up The search quality nes, through
Randomised allocation was reported in all eight studies. Method of randomisation reported in five studies. Overall quality as Good. This review strategy employed rating, for each in meta-analysis of to RESULTS Outcome Topical aprotining	Results Baseline cha and control g Methodologic assessed ind Disagreemer Data extractic A random eff meta-analyse heterogeneity ssessment (do appeared roboluded study whe data from in session of the s	groups n cal quali depende nts resol ion for e fects mc es as all y. lescripti ed the re bust, and vas prov ncluded Interver	ot reported. Ity of studies Ently by two authors. Ived by a third authorach study not stated odel was use in the showed statistical Ive) Search question, so the methods for a lided. This review prestudies.	or. d. cope, nalys	Five studies were double blinded. One study was not blinded. The blinding status of two studies was unclear. search terms and ir sis were appropriate ed pooled data, for example of the comparator group	nclu e. Tr	measurement bias Egger's test for bias was non-significant (p=0.6 and p=0.5) for reduction of blood loss or amount of blood transfusion, suggesting that there was no publication bias. usion/exclusion criteria. The summary, as well as a th of the specified outcon Statistical significan	States all 8 studies had "adequate" follow-up The search quality nes, through
Randomised allocation was reported in all eight studies. Method of randomisation reported in five studies. Overall quality as Good. This review strategy employed rating, for each induced meta-analysis of the RESULTS Outcome Topical aprotinin to 24-hr postoperation.	Results Baseline cha and control g Methodologic assessed ind Disagreemer Data extractic A random eff meta-analyse heterogeneity ssessment (do appeared roboluded study whe data from in session of the s	groups n cal quali depende nts resol ion for e fects mc es as all y. escripti ed the re bust, and was prov ncluded	ot reported. Ity of studies Ently by two authors. Ived by a third authorach study not stated odel was use in the showed statistical Ive) Search question, so the methods for a lided. This review prestudies.	or. d. cope, nalys	Five studies were double blinded. One study was not blinded. The blinding status of two studies was unclear. search terms and ir sis were appropriate ed pooled data, for each studies was unclead the studies was unclear.	nclu e. Tr	measurement bias Egger's test for bias was non-significant (p=0.6 and p=0.5) for reduction of blood loss or amount of blood transfusion, suggesting that there was no publication bias. usion/exclusion criteria. The summary, as well as a sh of the specified outcon Statistical significan	States all 8 studies had "adequate" follow-up The search quality nes, through
Randomised allocation was reported in all eight studies. Method of randomisation reported in five studies. Overall quality as Good. This review strategy employed rating, for each indicate meta-analysis of the RESULTS Outcome Topical aprotinin value blood loss	Results Baseline cha and control g Methodologic assessed ind Disagreemer Data extractic A random eff meta-analyse heterogeneity ssessment (do appeared roboluded study whe data from in session of the s	groups n cal quali depende nts resol ion for e fects mc es as all y. lescripti ed the re bust, and vas prov ncluded Interver	ot reported. Ity of studies Ently by two authors. Ived by a third authorach study not stated odel was use in the showed statistical Ive) Search question, so the methods for a lided. This review prestudies.	or. d. cope, nalys	Five studies were double blinded. One study was not blinded. The blinding status of two studies was unclear. search terms and ir sis were appropriate ed pooled data, for example of the comparator group	nclu e. Tr	measurement bias Egger's test for bias was non-significant (p=0.6 and p=0.5) for reduction of blood loss or amount of blood transfusion, suggesting that there was no publication bias. usion/exclusion criteria. The summary, as well as a th of the specified outcon Statistical significan	States all 8 studies had "adequate" follow-up The search quality nes, through
Randomised allocation was reported in all eight studies. Method of randomisation reported in five studies. Overall quality as Good. This review strategy employed rating, for each in meta-analysis of t RESULTS Outcome Topical aprotinin to 24-hr postoperative blood loss 5 trials (N=324)	Results Baseline cha and control g Methodologic assessed ind Disagreemer Data extractic A random eff meta-analyse heterogeneity ssessment (do clearly defined appeared roboluded study whe data from in separation of the control of t	groups n cal quali depende nts resol ion for e fects m ces as all y. escripti ed the re oust, and vas prov ncluded Intervel NR	ot reported. Ity of studies Ently by two authors. Ived by a third authorach study not stated odel was use in the showed statistical Ive) Search question, so the methods for a lided. This review prestudies.	cope, nalys	Five studies were double blinded. One study was not blinded. The blinding status of two studies was unclear. search terms and ir sis were appropriate ed pooled data, for example of the comparator group of the comparator g	nclu e. Tr	Egger's test for bias Egger's test for bias was non-significant (p=0.6 and p=0.5) for reduction of blood loss or amount of blood transfusion, suggesting that there was no publication bias. Usion/exclusion criteria. The summary, as well as a significant Statistical significant WMD -204 ml (95%Cl P<0.001 (Phet=0.04)	States all 8 studies had "adequate" follow-up The search quality nes, through
Randomised allocation was reported in all eight studies. Method of randomisation reported in five studies. Overall quality as Good. This review strategy employed rating, for each in meta-analysis of t RESULTS Outcome Topical aprotinin to 24-hr postoperation tube blood loss 5 trials (N=324) Allogenic RBC tra	Results Baseline cha and control g Methodologic assessed ind Disagreemer Data extractic A random eff meta-analyse heterogeneity ssessment (do clearly defined appeared roboluded study whe data from in separation of the control of t	groups n cal quali depende nts resol ion for e fects mc es as all y. lescripti ed the re bust, and vas prov ncluded Interver	ot reported. Ity of studies Ently by two authors. Ived by a third authorach study not stated odel was use in the showed statistical Ive) Search question, so the methods for a lided. This review prestudies.	cope, nalys	Five studies were double blinded. One study was not blinded. The blinding status of two studies was unclear. search terms and ir sis were appropriate ed pooled data, for example of the comparator group	nclu e. Tr	measurement bias Egger's test for bias was non-significant (p=0.6 and p=0.5) for reduction of blood loss or amount of blood transfusion, suggesting that there was no publication bias. usion/exclusion criteria. The summary, as well as a th of the specified outcon Statistical significan WMD -204 ml (95%Cl P<0.001 (Phet=0.04)	States all 8 studies had "adequate" follow-up The search quality nes, through The search and t
Randomised allocation was reported in all eight studies. Method of randomisation reported in five studies. Overall quality as Good. This review strategy employed rating, for each in meta-analysis of t RESULTS Outcome Topical aprotinin to 24-hr postoperative blood loss 5 trials (N=324)	Results Baseline cha and control g Methodologic assessed ind Disagreemer Data extractic A random eff meta-analyse heterogeneity ssessment (do clearly defined appeared roboluded study whe data from in separation of the control of t	groups n cal quali depende nts resol ion for e fects m ces as all y. escripti ed the re oust, and vas prov ncluded Intervel NR	ot reported. Ity of studies Ently by two authors. Ived by a third authorach study not stated odel was use in the showed statistical Ive) Search question, so the methods for a lided. This review prestudies.	cope, nalys	Five studies were double blinded. One study was not blinded. The blinding status of two studies was unclear. search terms and ir sis were appropriate ed pooled data, for example of the comparator group of the comparator g	nclu e. Tr	Egger's test for bias Egger's test for bias was non-significant (p=0.6 and p=0.5) for reduction of blood loss or amount of blood transfusion, suggesting that there was no publication bias. Usion/exclusion criteria. The summary, as well as a significant Statistical significant WMD -204 ml (95%Cl P<0.001 (Phet=0.04)	States all 8 studies had "adequate" follow-up The search quality nes, through The search and t

	T == (10 =)		>	I == 0 == (0=0) 0	
Allogenic RBC transfusion	97/179 (13.5) 108/162 (6		66.7)	RR 0.72 (95%CI : 0.47, 1.08)	
(incidence)				P=0.11 (P <i>het</i> =0.008)	
3 trials (N=341)	<u> </u>				
Topical tranexamic acid vs pla		г.,_		1	
24-hr postoperative chest	NR	NR		WMD -250 ml (95%CI : -465, -35)	
tube blood loss				P=0.02 (Phet<0.001)	
4 trials (N=269)					
Allogenic RBC transfusion	NR	NR		WMD -1.58 (95%CI : -2.26, -0.90)	
(units)				P=<0.001 (Phet=0.29)	
3 trials (N=229)					
Allogenic RBC transfusion	54/117 (46.2)	55/116 (47	7.4)	RR 0.98 (95%CI : 0.75, 1.27)	
(incidence)				P=0.88 (Phet=0.69)	
2 trials (N=233)					
Outcome	Clinical importance		Clinical rel	evance	
Topical aprotinin vs placebo					
24-hr postoperative chest	1: Clinically important benefit,		1: Patient-re	elevant clinical outcome.	
tube blood loss	confidence limit does not inclu	de null			
	value.				
Allogenic RBC transfusion	1: Clinically important benefit,		1: Patient-relevant clinical outcome.		
(units)	confidence limit does not inclu	de null			
	value.				
Allogenic RBC transfusion	4: Range of estimates includes		1: Patient-re	elevant clinical outcome.	
(incidence)	important point estimate but al				
	compatible with no effect, or a	harmful			
	effect				
Topical tranexamic acid vs pla					
24-hr postoperative chest	1: Clinically important benefit,		1: Patient-re	elevant clinical outcome.	
tube blood loss	confidence limit does not inclu	de null			
	value.				
Allogenic RBC transfusion	1: Clinically important benefit,		1: Patient-re	elevant clinical outcome.	
(units)	confidence limit does not inclu	de null			
	value.				
Allogenic RBC transfusion	4: Range of estimates includes	s clinically	1: Patient-re	elevant clinical outcome.	
(incidence)	important point estimate but al				
	compatible with no effect, or a	harmful			
	effect				
EXTERNAL VALIDITY					
Generalisability					
		np cardiac s	urgery, who n	nay not share clinical characteristics	
with the general surgical patie	nt population.				
Applicability	Applicability				

This review suggests that topical aprotinin and tranexamic acid may decrease blood loss and transfusion requirements.

Abbreviations: CI, confidence interval; het, heterogeneity; ITT, intention-to-treat; NR, not reported; RBC, red blood cell; RCT, randomised controlled trial; RR, relative risk; WMD, weighted mean difference.

The studies in this review were mostly conducted in developed countries (Croatia, Czech Republic, Egypt, England, Germany, Italy, Turkey), comparable to Australia. The surgeries performed (and the possible benefits) are likely to be relevant in the Australian context.

Comments

STUDY DETAILS

Citation

JR Brown, NJO Birkmeyer, GT O'Connor (2007) Meta-analysis comparing the effectiveness and adverse outcomes of antifibrinolytic agents in cardiac surgery. Circulation 115: 2801-2813.

Affiliation/Source of funds

Center for the Evaluative Clinical Sciences and Departments of Medicine and of Community and Family Medicine, Dartmouth Medical School, Hanover, New Hampshire, USA; Michigan Surgical Collaboration for Outcomes Research and Evaluation, University of Michigan, Ann Arbor, Michigan, USA.

Funding: Not stated.

Study design	Level of evidence	Location/setting
Systematic review including 138	Level I	Hospital
RCTs that investigated the		
effectiveness and adverse outcomes		
of antifibrinolytic agents in cardiac		
surgery.		

Intervention	Comparator
Aprotinin, tranexamic acid, ε-aminocaproic acid	Placebo or active
Demulation above stariation	

Population characteristics

Adults undergoing CABG, isolated valve or combined CABG/valve surgery.

Length of follow-up	Outcomes measured	
Not specified	Bleeding; incidence of transfusion; incidence or re-operation;	
	adverse events (including mortality, stroke, MI); renal	
	complications; renal dysfunction.	

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/	Follow-up
			measurement bias	(ITT)
Randomised allocation was reported in all 138 studies. Method of randomisation not reported in SR.	Baseline characteristics of intervention and control groups not reported. Data extraction conducted by two independent reviewers. Quality assessment conducted using Jadad score and determined not to influence results. A random effects model was use in the meta-analyses.	The majority of included studies were double-blinded.	Funnel plots generated to assess publication bias. Evidence of bias for aprotinin for total blood loss and incidence transfusion with pRBCs. Evidence of bias for aminocaproic acid and tranexamic acid for incidence of transfusion with pRBCs.	36/138 trials excluded patients from the analysis following randomis- ation

Overall quality assessment (descriptive)

Fair. The search strategy employed was inadequate as it included Medline only. The methods for analysis were appropriate. The summary, as well as a quality rating, for each included study was provided, but no individual study results were provided. This review provided pooled data only, for each of the specified outcomes, through meta-analysis of the data from included studies. A number of data extraction errors were identified when examining one of the subgroup analyses.

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance Risk estimate (95% CI)
Aprotinin high dose vs place	cebo		
	Mean ± SD	Mean ± SD	
Total blood loss (mL) 22 trials (N=1760)	NR	NR	High dose WMD -348 (-416, -281) P<0.001 (Phet=NR)
	n/N (%)	n/N (%)	

Incidence of transfusion with	NR	NR	High dose
pRBC			RR 0.60 (0.53, 0.67)
49 trials (N=4379)			P<0.001 (Phet=NR)
Return to operating room	NR	NR	High dose
40 trials (N=3912)			RR 0.47 (0.32, 0.69)
40 that3 (14–3712)			P<0.001 (Phet=NR)
Mortality	NR	NR	High dose
	INK	INK	
43 trials (N=6175)			RR 0.89 (0.65, 1.21)
			P=0.46 ((P <i>het</i> =NR)
Stroke	NR	NR	High dose
22 trials (N=1737)			RR 0.67 (0.30, 1.47)
			P=0.32 (P <i>het</i> =NR)
Myocardial infarction	NR	NR	High dose
31 trials (N=3315)			RR 1.10 (0.83, 1.45)
,			P=0.52 (P <i>het</i> =NR)
Renal failure	NR	NR	High dose
27 trials (N=4681)	IVIX	TVIC	RR 1.09 (0.68, 1.77)
27 tildis (N=4001)			P=0.71 (P <i>het</i> =NR)
Development's	ND	ND	
Renal dysfunction	NR	NR	RR 1.47 (1.12, 1.94)
19 trials (N=1778)			P=0.006 (P <i>het</i> =NR)
Aprotinin low dose vs placebo			
	Mean ± SD	Mean ± SD	
Total blood loss (mL)	NR	NR	WMD -226 (-277, -175)
6 trials (N=515)			P<0.001 (Phet=NR)
	n/N (%)	n/N (%)	,
Incidence of transfusion with	NR	NR	RR 0.76 (0.66, 0.86)
pRBC	IVIX	IVIX	P<0.001 (Phet=NR)
20 trials (N=1645)			F<0.001 (FIIel=NR)
	ND	ND	DD 0 (0 (0 41 1 10)
Return to operating room	NR	NR	RR 0.69 (0.41, 1.18)
20 trials (N=1623)			P=0.18 (P <i>het</i> =NR)
Mortality	NR	NR	RR 1.37 (0.72, 2.59)
14 trials (N=1453)			P=0.34 (P <i>het</i> =NR)
Stroke	NR	NR	RR 0.47 (0.09, 2.36)
10 trials (N=1049)			P=0.36 (P <i>het</i> =NR)
Myocardial infarction	NR	NR	RR 0.94 (0.58, 1.54)
16 trials (N=1585)			P=0.82 (P <i>het</i> =NR)
Renal failure	NR	NR	RR 1.86 (0.07, 49.3)
	INIX	IVIX	
7 trials (N=786)	ND	ND	P=0.71 (Phet=NR)
Renal dysfunction	NR	NR	RR 1.01 (0.69, 1.49)
9 trials (N=1041)			P=0.96 (P <i>het</i> =NR)
E-amino caproic acid vs place			
	Mean ± SD	Mean ± SD	
Total blood loss (mL)	NR	NR	WMD -240 (-341, -140)
3 trials (N=144)			P<0.001 (Phet=NR)
\	n/N (%)	n/N (%)	
Incidence of transfusion with	NR	NR	RR 0.63 (0.44, 0.90)
	IVIX	INIX	P=0.01 (P <i>het</i> =NR)
pRBC			r=0.01 (riiei=NK)
10 trials (N=628)			
Return to operating room	NR	NR	RR 0.51 (0.15, 1.82)
9 trials (N=851)			P=0.30 (P <i>het</i> =NR)
Mortality	NR	NR	RR 1.82 (0.55, 5.98)
6 trials (N=735)			P=0.32 ((Phet=NR)
Stroke	NR	NR	RR 0.60 (0.13, 2.81)
8 trials (N=833)		1 *** *	P=0.52 (P <i>het</i> =NR)
Myocardial infarction	NR	NR	RR 1.14 (0.50, 2.60)
	IVIX	IVIX	
8 trials (N=839)			P=0.76 (P <i>het</i> =NR)

Renal failure	-	-		-
0 trials				
Renal dysfunction 0 trials	-	-		-
Tranexamic acid vs placebo				
•	Mean ± SD	Mean ± S	D	
Total blood loss (mL) 11 trials (N=1100)	NR	NR		WMD -285 (-394, -175) P<0.001 (Phet=NR)
,	n/N (%)	n/N (%)		, ,
Incidence of transfusion with pRBC 22 trials (N=2429)	NR	NR		RR 0.75 (0.60, 0.92) P=0.007 (Phet=NR)
Return to operating room 21 trials (N=2255)	NR	NR		RR 0.70 (0.44, 1.11) P=0.13 (Phet=NR)
Mortality 18 trials (N=2229)	NR	NR		RR 0.67 (0.33, 1.37) P=0.28 ((Phet=NR)
Stroke 15 trials (N=2098)	NR	NR		RR 1.31 (0.59, 2.93) P=0.51 (Phet=NR)
Myocardial infarction 16 trials (N=2219)	NR	NR		RR 0.94 (0.51, 1.74) P=0.85 (Phet=NR)
Renal failure 3 trials (N=840)	NR	NR		RR 1.43 (0.30, 6.85) P=0.66 (P <i>het</i> =NR)
Renal dysfunction 4 trials (N=684)	NR	NR		RR 2.02 (0.73, 5.60) P=0.18 (P <i>het</i> =NR)
Outcome	Clinical importance		Clinical rele	evance
Aprotinin high dose vs placebo			T = -:	
Total blood loss	Clinically important benefit, confidence limit does not inclu value.	de null		elevant clinical outcome.
Incidence of transfusion with pRBC	1: Clinically important <i>benefit</i> , confidence limit does not inclu value.	de null	1: Patient-relevant clinical outcome.	
Return to operating room	1: Clinically important <i>benefit</i> , confidence limit does not incluvalue.	de null	1: Patient-re	elevant clinical outcome.
Mortality	4: Range of estimates includes important effects but is also cowith no effect, or a harmful effect.	mpatible	1: Patient-relevant clinical outcome.	
Stroke	4: Range of estimates includes important effects but is also cowith no effect, or a harmful effect.	s clinically mpatible	1: Patient-relevant clinical outcome.	
Myocardial infarction	4: Range of estimates includes important effects but is also cowith no effect, or a harmful effect.	mpatible ect	1: Patient-relevant clinical outcome.	
Renal failure	4: Range of estimates includes clinically important effects but is also compatible with no effect, or a harmful effect		1: Patient-relevant clinical outcome.	
Renal dysfunction	Clinically important <i>harm</i> , confidence limit does not include null value.		1: Patient-relevant clinical outcome.	
Aprotinin low dose vs placebo				
Total blood loss	1: Clinically important <i>benefit</i> , confidence limit does not inclu value.	de null		elevant clinical outcome.
Incidence of transfusion with pRBC	1: Clinically important <i>benefit</i> , confidence limit does not inclu value.	de null	1: Patient-relevant clinical outcome.	

Return to operating room	2: Point estimate shows clinically	1: Patient-relevant clinical outcome.
Return to operating room	important <i>benefit</i> , but confidence limit	1. I diletti-relevant clinical outcome.
	includes null value.	
Mortality	4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Wortanty	important effects but is also compatible	The distriction of the districti
	with no effect, or a harmful effect	
Stroke	4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
	important effects but is also compatible	
	with no effect, or a harmful effect	
Myocardial infarction	4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
,	important effects but is also compatible	
	with no effect, or a harmful effect	
Renal failure	4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
	important effects but is also compatible	
	with no effect, or a harmful effect	
Renal dysfunction	4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
	important effects but is also compatible	
	with no effect, or a harmful effect	
E-aminocaproic acid vs placel		
Total blood loss	1: Clinically important benefit,	1: Patient-relevant clinical outcome.
	confidence limit does not include null	
	value.	
Incidence of transfusion with	1: Clinically important benefit,	1: Patient-relevant clinical outcome.
pRBC	confidence limit does not include null	
	value.	
Return to operating room	2: Point estimate shows clinically	1: Patient-relevant clinical outcome.
	important benefit, but confidence limit	
Manufalls.	includes null value.	1 Dellant advantallated as terms
Mortality	2: Point estimate shows clinically	1: Patient-relevant clinical outcome.
	important <i>harm</i> but confidence limit	
Stroke	includes null value. 2: Point estimate shows clinically	1: Patient-relevant clinical outcome.
Siroke	important <i>benefit</i> , but confidence limit	1: Patient-relevant clinical outcome.
	includes null value.	
Myocardial infarction	2: Point estimate shows clinically	1: Patient-relevant clinical outcome.
Wyocardiai iriiaictiori	important <i>harm</i> , but confidence limit	1. Fatient-relevant clinical outcome.
	includes null value.	
Renal failure	NA	1: Patient-relevant clinical outcome.
Renal dysfunction	NA	1: Patient-relevant clinical outcome.
Tranexamic acid vs placebo/n		The distriction of the districti
Total blood loss	1: Clinically important <i>benefit</i> ,	1: Patient-relevant clinical outcome.
Total blood 1033	confidence limit does not include null	1. I dilott folevalt diffical dateome.
	value.	
Incidence of transfusion with	1: Clinically important <i>benefit</i> ,	1: Patient-relevant clinical outcome.
pRBC	confidence limit does not include null	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
•	value.	
Return to operating room	2: Point estimate shows clinically	1: Patient-relevant clinical outcome.
ı	important benefit, but confidence limit	
	includes null value.	
Mortality	2: Point estimate shows clinically	1: Patient-relevant clinical outcome.
ý	important benefit but confidence limit	
	includes null value.	
Stroke	2: Point estimate shows clinically	1: Patient-relevant clinical outcome.
	important <i>harm</i> , but confidence limit	
	includes null value.	

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Myocardial infarction	2: Point estimate shows clinically important <i>benefit</i> , but confidence limit includes null value.	1: Patient-relevant clinical outcome.
Renal failure	2: Point estimate shows clinically important <i>harm</i> , but confidence limit includes null value.	1: Patient-relevant clinical outcome.
Renal dysfunction	2: Point estimate shows clinically important <i>harm</i> , but confidence limit includes null value.	1: Patient-relevant clinical outcome.
EXTERNAL VALIDITY		

Generalisability

This systematic review focuses on adults undergoing cardiac surgery (CABG, valve or CABG+valve), who are unlikely to share clinical characteristics with the general surgical patient population.

Applicability

The location of the included studies in this review was not stated, however it includes a large number of studies so likely to be largely applicable. The surgeries performed (and the possible benefits) are likely to be relevant in the Australian context.

Comments

Abbreviations: CABG, coronary artery bypass graft; CI, confidence interval; *het*, heterogeneity; ITT, intention to treat; MI, myocardial infarction; NA, not applicable; NR, not reported; pRBC, packed red blood cells; RCT, randomised controlled trial; RR, risk ration; SR, systematic review; WMD, weighted mean difference.

a Includes one trial in which aprotinin + ε-aminocaproic acid was compared with ε-aminocaproic acid only.

^b Discrepancy between Figure 2 and text. Data taken from Figure 2.

STUDY DETAILS Citation PA Carless, BJ Stokes, AJ Moxey, DA Henry (2004) Desmopressin use for minimising perioperative allogenic blood transfusion. Cochrane Database of Systematic Reviews. Issue 1. Article No.: CD001884. DOI: 10.1002/14651858.CD001884.pub2. Affiliation/Source of funds Discipline of Clinical Pharmacology, Faculty of Health, University of Newcastle, Newcastle, Australia; Institute of Clinical Evaluative Sciences, Toronto, Canada. Funding: Special purpose grant, Hunter Area Pathology Services, NSW, Australia; Australian Health Ministers' Advisory Committee, NHMRC, Australia. Study design Level of evidence Location/setting Systematic review including 29 RCTs Level I Hospital that investigated the effectiveness and adverse outcomes of desmopressin use in reducing perioperative blood loss. Intervention Comparator Desmopressin acetate administered intravenously as Placebo prophylactic therapy during the perioperative period. Variable doing regimens were used in the included trials. Population characteristics Adults undergoing surgery who did not have congenital bleeding disorders. Length of follow-up Outcomes measured Not specified Incidence of transfusion; volume of transfusion; blood loss; reoperation due to bleeding; mortality; myocardial infarction; stroke; thrombosis INTERNAL VALIDITY Allocation Blinding analysis Treatment/ Follow-up Results measurement bias (ITT) Randomised Baseline characteristics of intervention 26/29 trials Funnel plots 11/29 trials allocation was and control groups not reported. double-blind; 2/29 generated to assess reported no reported in all Study inclusion and quality assessment unclear, and 1/29 publication bias. Little exclusions 29 studies. conducted by two independent reviewers. not double-blind evidence of or ITT Quality assessment conducted using analysis; Method of publication bias for criteria proposed by Schulz, Number of randomisation incidence of blood 17/29 inadequate or reviewers carrying out data extraction not transfusion and blood reported not reported in exclusions stated. loss. 22 trials. Meta-analysis performed using Review but these Appears to be no Manager using a random effects model. measurement or judged

Overall quality assessment (descriptive)

Good. The search strategy employed was adequate and the methods for analysis were appropriate. The summary, as well as a quality rating, for each included study was provided. This review provided pooled data, for each of the specified outcomes, through meta-analysis of the data from included studies. The authors note that the methodological quality of most of the included studies was poor

RESULTS

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unlikely to cause bias; 1/29 exclusions not reported.

treatment bias.

Outcome No. trials (N) No. trials included in analysis (N) ^a	Intervention group	Comparator group	Statistical significance Risk estimate (95% CI)
Desmopressin vs placebo			
	n/N (%)	n/N (%)	
Incidence of transfusion 19 trials (N=1387) 17 trials (N=1308)	383/703 (54.5)	377/684 (55.1)	All studies RR 0.96 (0.87, 1.06) P=0.42 (Phet=0.19)
Incidence of transfusion 15 trials (N=1196) 14 trials (1137)	341/610 (55.9)	330/586 (56.3)	Cardiac surgery RR 0.95 (0.84, 1.07) P=0.39 (Phet=0.11)
Incidence of transfusion 4 trials (N=191) 3 trials (N=171)	42/93 (45.2)	47/98 (48.0)	Miscellaneous surgery RR 1.01 (0.81, 1.26) P=0.91 (Phet=0.59)
Incidence of transfusion 9 trials (N=586) 8 trials (N=527)	150/299 (50.2)	158/287 (55.1)	Cardiac surgery/Primary CABG RR 0.85 (0.73, 0.99) P=0.038 (Phet=0.43)
Incidence of transfusion 6 trials (N=610)	191/311 (61.4)	172/299 (57.5)	Cardiac surgery/CABG + valve ± combination/redo surgery RR 1.03 (0.88, 1.19) P=0.75 (Phet=0.14)
Incidence of transfusion 6 trials (N=399) 5 trials (N=340)	91/192 (47.4)	103/207 (49.8)	Cardiac surgery/ASA use RR 0.89 (0.64, 1.23) P=0.49 (Phet=0.12)
Incidence of transfusion 4 trials (N=286)	69/153 (45.1)	73/133 (54.9)	Cardiac surgery/no ASA use RR 0.79 (0.62, 1.01) P=0.056 (Phet=0.36)
Incidence of transfusion 10 trials (N=736)	180/373 (48.3)	190/363 (52.3)	Transfusion protocol RR 0.90 (0.77, 1.04) P=0.16 (Phet=0.25)
Incidence of transfusion 9 trials (N=651) 7 trials (N=572)	203/330 (61.5)	187/321 (58.3)	No transfusion protocol RR 1.03 (0.93, 1.14) P=0.60 (Phet=0.40)
Incidence of transfusion 10 trials (N=732) 9 trials (N=673)	242/382 (63.4)	237/350 (67.7)	No autologous techniques used RR 0.91 (0.78, 1.07) P=0.25 (Phet=0.04)
Incidence of transfusion 9 trials (N=655) 8 trials (N=635)	141/321 (43.6)	140/334 (41.9)	Autologous techniques used RR 1.00 (0.84, 1.19) P=0.97 (Phet=0.31)
Incidence of transfusion 3 trials (N=249) 2 trials (N=190)	73/124 (58.9)	74/125 (59.2)	Cochrane quality rating A RR 0.97 (0.75, 1.24) P=0.80 (Phet=0.50)
Incidence of transfusion 11 trials (N=766) 10 trials (N=746)	219/400 (54.8)	215/366 (58.7)	Cochrane quality rating B RR 0.88 (0.75, 1.03) P=0.12 (Phet=0.04)
Incidence of transfusion 5 trials (N=372)	91/179 (50.8)	88/193 (45.6)	Cochrane quality rating C RR 1.11 (0.94, 1.33) P=0.22 (Phet=0.75)
	Mean ± SD	Mean ± SD	,
Units of blood transfused 14 trials (N=885)	NR	NR	All patients WMD -0.30 (-0.60, -0.01) P=0.042 (Phet=0.07)
Units of blood transfused 10 trials (N=621)	NR	NR	Cardiac surgery WMD -0.39 (-0.77, -0.01) P=0.047 (Phet=0.03)

Halla af laboration of the	LND	LND	Outhorse
Units of blood transfused	NR	NR	Orthopaedic surgery
2 trials (N=129)			WMD -0.15 (-0.64, 0.33)
Units of blood transfersed	ND	ND	P=0.54 (Phet=0.43)
Units of blood transfused	NR	NR	Vascular surgery
2 trials (N=135)			WMD 0.06 (-0.89, 1.02)
Units of blood transfused	NR	NR	P=0.90 (Phet=0.40)
	INK	INK	No autologous techniques used WMD -0.22 (-0.55, 0.10)
10 trials (N=734)			P=0.18 (P <i>het</i> =0.19)
Units of blood transfused	NR	NR	Autologous techniques used
4 trials (N=151)	INIX	IVIX	WMD -0.47 (-1.15, 0.20)
4 11013 (11–131)			P=0.17 (Phet=0.08)
Units of blood transfused	NR	NR	Transfused patients
5 trials (N=211)	IVIX	IVIX	WMD -0.49 (-0.94, -0.04)
5 tildis (IV-211)			P=0.033 (Phet=0.49)
Intraoperative blood loss	NR	NR	All surgery
7 trials (N=493)	IVIX	IVIX	WMD -90.07 (-199.56, 19.42)
/ titals (11–473)			P=0.11 (Phet=0.17)
Intraoperative blood loss	NR	NR	Cardiac surgery
3 trials (N=229)	IVIX	WIX	WMD -119.79 (-314.57, 75.00)
5 that5 (14–227)			P=0.23 (Phet=0.06)
Postoperative blood loss	NR	NR	All surgery
18 trials (N=1201)	TWI C	TVIX	WMD -92.98 (-149.86, -36.11)
10 mais (11 1201)			P=0.0014 (Phet=0.001)
Postoperative blood loss	NR	NR	Cardiac surgery
16 trials (N=1107)	1	1	WMD -96.58 (-163.04, -30.12)
			P=0.0044 (Phet<0.001)
Postoperative blood loss	NR	NR	Cardiac surgery / 0-6 hours post-
1 trial (N=59)		1	op
			-98.00 (-304.99, 108.99)
			P=0.35 (Phet=NA)
Postoperative blood loss	NR	NR	Cardiac surgery/ 0-12 hours post-
3 trials (N=233)			ор
, ,			WMD -114.05 (-269.46, 41.36)
			P=0.15 (Phet=0.004)
Postoperative blood loss	NR	NR	Cardiac surgery/ 0-16 hours post-
2 trials (N=122)			ор
			WMD -18.01 (-113.34, 77.32)
			P=0.71 (Phet=0.42)
Postoperative blood loss	NR	NR	All surgery/0-24 hours post-op
12 trials (N=787)			WMD -100.41 (-176.48, -24.34)
			P=0.0097 (Phet=0.004)
Postoperative blood loss	NR	NR	Cardiac surgery/ 0-24 hours post-
10 trials (N=693)			ор
			WMD -107.46 (-207.12, -7.80)
			P=0.035 (Phet=0.002)
Postoperative +	NR	NR	All surgery
intraoperative blood loss			WMD -241.78 (-387.55, -96.1)
10 trials (N=669)			P=0.0012 (P <i>het</i> =0.002)
Postoperative +	NR	NR	Cardiac surgery
intraoperative blood loss			WMD -237.92 (-413.43, -62.40)
7 trials (N=496)			P=0.0079 (Phet<0.001)
	n/N (%)	n/N (%)	
Reoperation for bleeding	7/383	14/395	All surgery
11 trials (N=778)			RR 0.69 (0.26, 1.83)
9 trials (N=693)			P=0.45 (P <i>het</i> =0.39)

742

Mortality	13/534	7/527	All surgery
12 trials (N=1061)			RR 1.72 (0.68, 4.33)
8 trials (N=774)			P=0.25 (Phet=0.80)
Myocardial infarction	28/441	18/435	All surgery
12 trials (N=876)	20/111	10/100	RR 1.38 (0.77, 2.50)
9 trials (N=731)	0/4.0.4	0/47/	P=0.28 (Phet=0.87)
Stroke (CVA)	8/184	2/176	All surgery
5 trials (N=360)			RR 2.40 (0.68, 8.43)
			P=0.17 (P <i>het</i> =0.17)
Any thrombosis	14/361	10/330	All surgery
9 trials (N=691)			RR 1.46 (0.64, 3.35)
7 trials (N=591)			P=0.37 (Phet=0.78)
Hypotension during infusion	34/92	9/91	All surgery
	34/92	9/91	
requiring treatment			RR 2.81 (1.50, 5.27)
5 trials (N=183)			P=0.0013 (Phet=0.50)
Outcome	Clinical importance		Clinical relevance
Desmopressin vs placebo	•		
Incidence of transfusion	Any surgery		1: Patient-relevant clinical outcome.
moderice of transitionin		e clinically	1. Fation(Fibiovant clinical vulcome.
	4: Range of estimates include:		
	important effects but is also co	ımpatıble	
	with no effect		
	Cardiac surgery		
	4: Range of estimates include:	s clinically	
	important effects but is also co		
	with no effect		
	Miscellaneous surgery		
	4: Range of estimates include:		
	important effects but is also co	mpatible	
	with no effect		
	Cardiac surgery/Primary CAB	G	
	1: Clinically important benefit,		
	confidence limit does not inclu	de null	
	value.	ue nun	
	Cardiac surgery/CABG + valve	e surgery	
	± combination/redo surgery		
	4: Range of estimates include:	s clinically	
	important effects but is also co	mpatible	
	with no effect	•	
	Cardiac surgery/ASA use		
	4: Range of estimates include:	s clinically	
	important effects but is also co	лпраные	
	with no effect		
	Cardiac surgery/no ASA use		
	4: Range of estimates includes clinically		
	important effects but is also co	,	
	with no effect		
	Transfusion protocol		
	•	e elipiaellu	
	4: Range of estimates include:		
	important effects but is also co	ımpatıble	
	with no effect		
	No transfusion protocol		
	4: Range of estimates include:	s clinically	
	important effects but is also co		
	with no effect		
		nd	
1	No autologous techniques use		
1	1. Dongo of actimates include	a aliminally	
	4: Range of estimates include: important effects but is also co		

	with no effect Autologous techniques used 4: Range of estimates includes clinically important effects but is also compatible with no effect Cochrane rating A 4: Range of estimates includes clinically important effects but is also compatible with no effect Cochrane rating B 4: Range of estimates includes clinically important effects but is also compatible with no effect Cochrane rating C 4: Range of estimates includes clinically important effects but is also compatible with no effect	
Units of blood transfused	All surgery/all patients 1: Clinically important benefit, confidence limit does not include null value Cardiac surgery/all patients 1: Clinically important benefit, confidence limit does not include null value Orthopaedic surgery/all patients 4: Range of estimates includes clinically important effects but is also compatible with no effect Vascular surgery/all patients 4: Range of estimates includes clinically important effects but is also compatible with no effect No autologous techniques used/all patients 4: Range of estimates includes clinically important effects but is also compatible with no effect No autologous techniques used/all patients 4: Range of estimates includes clinically important effects but is also compatible with no effect Autologous techniques used 4: Range of estimates includes clinically important effects but is also compatible with no effect Autologous techniques used 4: Range of estimates includes clinically important effects but is also compatible with no effect All surgery/transfused patients 1: Clinically important benefit, confidence limit does not include null value.	1: Patient-relevant clinical outcome.
Intraoperative blood loss	All surgery 4: Range of estimates includes clinically important effects but is also compatible with no effect Cardiac surgery 4: Range of estimates includes clinically important effects but is also compatible with no effect	

Postoperative blood loss	All surgery 1: Clinically important benefit, confidence limit does not include null value Cardiac surgery 1: Clinically important benefit, confidence limit does not include null value Cardiac surgery/0-6 hours post-op 4: Range of estimates includes clinically important effects but is also compatible	
	with no effect Cardiac surgery/ 0-12 hours post-op 4: Range of estimates includes clinically important effects but is also compatible with no effect Cardiac surgery/0-16 hours post-op 4: Range of estimates includes clinically important effects but is also compatible with no effect All surgery/0-24 hours/post-op 1: Clinically important benefit, confidence limit does not include null value Cardiac surgery/0-24 hours post-op 1: Clinically important benefit, confidence limit does not include null	
Postoperative + intraoperative blood loss	value All surgery 1: Clinically important benefit, confidence limit does not include null value Cardiac surgery 1: Clinically important benefit, confidence limit does not include null value	
Reoperation for bleeding	All surgery 4: Range of estimates includes clinically important effects but is also compatible with no effect, or a harmful effect	1: Patient-relevant clinical outcome.
Mortality	All surgery 4: Range of estimates includes clinically important harmful effects but is also compatible with no effect, or a beneficial effect	1: Patient-relevant clinical outcome.
Myocardial infarction	All surgery 4: Range of estimates includes clinically important harmful effects but is also compatible with no effect, or a beneficial effect	1: Patient-relevant clinical outcome.
Stroke	All surgery 4: Range of estimates includes clinically important harmful effects but is also compatible with no effect, or a beneficial effect	1: Patient-relevant clinical outcome.

Any thrombosis	All surgery 4: Range of estimates includes clinically important harmful effects but is also compatible with no effect, or a beneficial effect	1: Patient-relevant clinical outcome.
Hypotension during infusion requiring treatment	All surgery 1: Clinically important harm, confidence limit does not include null value.	1: Patient-relevant clinical outcome.

Generalisability

This systematic review focuses on adults who do not have congenital bleeding disorders undergoing any elective or non-urgent surgery so should be generalisable to other similar patients undergoing similar surgery types.

Applicability

The studies in this review were mostly conducted in developed countries (US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and the UK), comparable to Australia. The surgeries performed (and the possible benefits) are likely to be relevant in the Australian context.

Comments

This review suggests that desmopressin has no benefit in reducing the requirement for perioperative allogenic RBC transfusion in patients who do not have congenital bleeding disorders. There is some evidence of benefit reduction in perioperative blood loss, but these were small and not considered clinically important.

Abbreviations: ASA, acetylsalicylic acid; CABG, coronary artery bypass graft; CI, confidence interval; CVA, cerebrovascular accident; het, heterogeneity; ITT, intention to treat; NA, not applicable; NHMRC, National Health and Medical Research Council; NR, not reported; pRBC, packed red blood cells; RCT, randomised controlled trial; RR, relative risk; SR, systematic review; WMD, weighted mean difference.

a Where individual studies had either 100% events in both treatment arms, no events in both treatment arms or no SD was reported, a risk estimate for that individual study could not be calculated, and it could not be included in the pooled analysis. Where this has occurred, the actual number of studies and subjects included in the analysis will be stated.

STUDY DETAILS Citation G Crescenzi, G Landoni, G Biondi-Zoccai et al (2008) Desmopressin reduces transfusion needs after surgery. Anesthesiology 109: 1063-1076. Affiliation/Source of funds Department of Cardiothoracic Anesthesia and Intensive Care, Università Vita-Salute San Raffaele, Milan, Italy; Division of Cardiology, University of Turin, Turin, Italy; Department of Cardiothoracic Anestheia and Intensive Care, IRCCS Policinico S. Donato, San Donato Milanese, Italy. Funding: Solely from institutional and/or departmental sources. Level of evidence Location/setting Study design Systematic review including 42 RCTs Level I Hospital (4/38 trials reported data on different patient populations so counted as separate trials hence 42 trials in total) that investigated the effectiveness and safety of desmopressin in patients undergoing surgery. Intervention Comparator Desmopressin Placebo Dose varied slightly across included studies, being mostly a single 0.3 µg/kg dose administered 15-30 minutes during surgical haemostasis. In six studies the dose was repeated and in 8 studies it was administered immediately before surgery Population characteristics Adult surgical patients Length of follow-up Outcomes measured Not specified Transfusion (units); blood loss (mL); re-operation for bleeding; incidence of transfusion (blood products); incidence of transfusion (platelets only); hypotension; death; myocardial infarction; thromboses (other than MI). INTERNAL VALIDITY Allocation Results Blinding analysis Treatment/ Follow-up measurement bias (ITT) Randomised Baseline characteristics of intervention 33/38 double Publication bias not Not stated. allocation was and control groups not reported. blinded. assessed. Treatment reported in all Selection of studies carried out by four or measurement bias studies. reviewers. Data extracted independently not apparent. No further by five reviewers. No quality assessment details carried out. Analysis of units of blood transfused and provided. blood loss carried out using standardised mean difference. Fixed-effects model used if $I^2 < 50\%$ and random effects model used if $I^2 > 50\%$ Overall quality assessment (descriptive) Fair. The search strategy employed as well as study selection and extraction of data were adequate. No formal quality assessment of included studies was performed. No assessment of reasons for substantial heterogeneity carried out. RESULTS Statistical significance Outcome Intervention group Comparator group No. trials (N) Risk estimate (95% CI) No. trials included in analysis (N)a Desmopressin vs placebo

	Mean ± SD	Mean ± SD	
Units of blood products	NR	NR	All studies
transfused			SMD -0.29 (-0.52, -0.06)
34 trials (N=2065)			P=0.01 (Phet<0.001)
Units of blood products	NR	NR	Cardiac surgery
transfused			SMD -0.22 (-0.52, 0.08)
23 trials (N=1607)			P=0.14 (Phet<0.001)
Units of blood products	NR	NR	Non-cardiac surgery
transfused			SMD -0.45 (-0.77, -0.13)
11 trials (N=458)			P=0.006 (Phet=0.003)
Blood loss (mL)	NR	NR	All studies
40 trials (N=2445)			SMD -0.20 (-0.34, -0.06)
			P=0.004 (Phet<0.001)
Blood loss (mL)	NR	NR	Cardiac surgery
29 trials (N=1928)			SMD -0.23 (-0.40, -0.05)
			P=0.01 (Phet<0.001)
Blood loss (mL)	NR	NR	Non-cardiac surgery
11 trials (N=517)			SMD -0.10 (-0.28, 0.07)
	In . (0.1)	(n. (o.)	P=0.25 (P <i>het</i> =0.45)
D " () "	n/N (%)	n/N (%)	All I P
Re-operation for bleeding	21/763 (2.8)	34/779 (4.4)	All studies
25 trials (N=1542)			OR 0.65 (0.39, 1.09)
15 trials (N=1186)	10// 47 (2.0)	21// [7 / 4 7)	P=0.11 (Phet=0.50)
Re-operation for bleeding	18/647 (2.8)	31/657 (4.7)	Cardiac surgery
18 trials (N=1304)			OR 0.63 (0.36, 1.08)
14 trials (N=1136)	2/11/ (2 /)	2/122 /2 5\	P=0.09 (Phet=0.44)
Re-operation for bleeding	3/116 (2.6)	3/122 (2.5)	Non-cardiac surgery
7 trials (N=238) 1 trial (N=50)			OR 1.00 (0.18, 5.51) P=1.0 (P <i>het</i> =NA)
Incidence of transfusion with	411/746 (55.1)	430/742 (58.0)	All studies
blood products (pRBCs,	411/740 (33.1)	430//42 (30.0)	OR 0.88 (0.70, 1.10)
fresh frozen plasma,			P=0.26 (P <i>het</i> =0.19)
platelets)			1 -0.20 (1 1101-0.17)
22 trials (N=1488)			
21 trials (N=1429)			
Incidence of transfusion with	350/638 (54.9)	367/634 (57.9)	Cardiac surgery
blood products (pRBCs,	000,000 (0 117)	0077001 (0717)	OR 0.87 (0.68, 1.11)
fresh frozen plasma,			P=0.26 (Phet=0.07)
platelets)			,
17 trials (N=1272)			
16 trials (N=1213)			
Incidence of transfusion with	61/108 (56.5)	63/108 (58.3)	Non-cardiac surgery
blood products (pRBCs,			OR 0.93 (0.48, 1.79)
fresh frozen plasma,			P=0.83 (Phet=0.81)
platelets)			
5 trials (N=216)			
Incidence of transfusion with	37/386 (10.0)	53/383 (13.8)	Cardiac surgery
platelets			OR 0.64 (0.41, 1.01)
11 trials (N=769)	07/450 (0.0)	0/400 (0.1)	P=0.06 (Phet=0.22)
Risk of hypotension	37/450 (8.2)	9/432 (2.1)	All studies
18 trials (N=882)			OR 4.84 (2.31, 10.13)
7 trials (N=320)	10/240 (5.2)	1/240 (0.2)	P<0.001 (Phet=0.85)
Risk of hypotension	19/368 (5.2)	1/349 (0.3)	Cardiac surgery
13 trials (N=717)			OR 8.92 (2.54, 31.37) P<0.001 (Phet=0.94)
5 trials (N=221)			r<0.001 (<i>PHet</i> =0.94)

Risk of hypotension 5 trials (N=165) 2 trials (N=99)	18/82 (22.0)	8/83 (9.6)		Non-cardiac surgery OR 3.04 (1.18, 7.87) P=0.02 (Phet=0.64)
Risk of death 23 trials (N=1539) 8 trials (N=673)	9/771 (1.2)	7/768 (0.9		All studies 1.25 (0.51, 3.04) P=0.63 (Phet=0.76)
Risk of death 19 trials (N=1334) 7 trials (N=582)	7/674 (1.0)	7/660 (1.1)	Cardiac surgery 1.00 (0.38, 2.62) P=1.00 (Phet=0.81)
Risk of death 4 trials (N=205) 1 trial (N=50)	2/97 (2.1)	0/108 (0)		Non-cardiac surgery 5.84 (0.27, 125.19) P=0.26 (Phet=NA)
Risk of myocardial infarction 27 trials (N=1609) 13 trials (N=916)	31/816 (3.8)	23/793 (2.	9)	All studies 1.27 (0.73, 2.20) P=0.40 (Phet=0.88)
Risk of myocardial infarction 19 trials (N=1262) 11 trials (N=775)	28/648 (4.3)	19/614 (3.	,	Cardiac surgery 1.36 (0.75, 2.48) P=0.31 (Phet=0.86)
Risk of myocardial infarction 8 trials (N=347) 2 trials (N=141)	3/168 (1.2)	4/179 (2.2		Non-cardiac surgery 0.84 (0.20, 3.53) P=0.81 (Phet=0.35)
Risk of thromboses (other than myocardial infarction) 26 trials (N=1776) 14 trials (N=1151)	26/899 (2.9)	22/877 (2.	5)	All studies 1.20 (0.68, 2.09) P=0.53 (Phet=0.82)
Risk of thromboses (other than myocardial infarction) 18 trials (N=1400) 11 trials (N=931)	18/717 (2.5)	14/683 (2.	0)	Cardiac surgery 1.27 (0.64, 2.50) P=0.49 (Phet=0.86)
Risk of thromboses (other than myocardial infarction) 8 trials (N=376) 3 trials (N=220)	8/182 (4.4)	8/194 (4.1)	Non-cardiac surgery 1.06 (0.39, 2.84) P=0.92 (Phet=0.24)
Outcome	Clinical importance	1	Clinical rele	evance
Desmopressin vs placebo			,	
Units of blood transfused	All studies 1: Clinically important benefit, confidence limit does not include null value. Cardiac surgery 4: Range of estimates includes clinically important effects, but also compatible with no effect Non-cardiac surgery 1: Clinically important benefit, confidence limit does not include null value.		1: Patient-re	elevant clinical outcome.

Pland loss	All studios	1. Dationt relevant clinical outcome
Blood loss	All studies	1: Patient-relevant clinical outcome.
	1: Clinically important <i>benefit</i> ,	
	confidence limit does not include null	
	value.	
	Cardiac surgery	
	1: Clinically important benefit,	
	confidence limit does not include null	
	value.	
	Non-cardiac surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no effect	
Re-operation for bleeding	All studies	1: Patient-relevant clinical outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no effect	
	Cardiac surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no effect	
	Non-cardiac surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
In although a file of fire and the	with no effect	1. Dell'est selecced ell'ellectes acces
Incidence of transfusion with	All studies	1: Patient-relevant clinical outcome.
blood products	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no effect	
	Cardiac surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no effect	
	Non-cardiac surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no effect	
Incidence of transfusion with	Cardiac surgery	1: Patient-relevant clinical outcome.
platelets	4: Range of estimates includes clinically	
F	important effects, but also compatible	
	with no effect	
Risk of hypotension	All studies	1: Patient-relevant clinical outcome.
Task of Hypoterision	1: Clinically important <i>harm</i> , confidence	1. 1 duont folovant olimical outcome.
	limit does not include null value.	
	Cardiac surgery	
	1: Clinically important <i>harm</i> , confidence	
	limit does not include null value.	
	Non-cardiac surgery	
	1: Clinically important <i>harm</i> , confidence	
	limit does not include null value.	

Risk of death	All studies 4: Range of estimates includes clinically important effects, but also compatible with no effect Cardiac surgery 4: Range of estimates includes clinically important effects, but also compatible with no effect Non-cardiac surgery 4: Range of estimates includes clinically important effects, but also compatible with no effect	1: Patient-relevant clinical outcome.
Risk of myocardial infarction	All studies 4: Range of estimates includes clinically important effects, but also compatible with no effect Cardiac surgery 4: Range of estimates includes clinically important effects, but also compatible with no effect Non-cardiac surgery 4: Range of estimates includes clinically important effects, but also compatible with no effect	1: Patient-relevant clinical outcome.
Risk of thromboses (other than myocardial infarction)	All studies 4: Range of estimates includes clinically important effects, but also compatible with no effect Cardiac surgery 4: Range of estimates includes clinically important effects, but also compatible with no effect Non-cardiac surgery 4: Range of estimates includes clinically important effects, but also compatible with no effect	1: Patient-relevant clinical outcome.

Generalisability

This systematic review focuses on patients who have undergone various types of surgery as such it may be relevant to a general surgical population. Subgroup analyses are also provided for cardiac and non-cardiac surgery so these results are likely generalisable to these specific surgical populations.

Applicability

This review does not report the locations of the included studies and as such the applicability of the results to the Australian setting is unclear. However, this review include all studies included in a previous Cochrane review which stated the countries in which the included trials were carried out included the US, Canada, Spain, Sweden, Germany, Turkey, Israel, China, Norway, Finland and UK.

Comments

This review suggests that desmopressin may slightly reduce blood loss and transfusion volume without reducing the proportion of patients who require transfusion, while increasing the risk of hypotension. The authors note this is due to a mild vasodilating effect of desmopressin and does not result in changes in filling pressure, heart rate or right ventricular function.

Abbreviations: CI, confidence interval; het, heterogeneity; ITT, intention to treat; MI, myocardial infarction; OR, odds-ratio; pBRC, packed red blood cells; RCT, randomised controlled trial; SD, standard deviation; SMD, standardised mean difference.

^a Where individual studies had either 100% events in both treatment arms, no events in both treatment arms or no SD was reported, a risk estimate for that individual study could not be calculated, and it could not be included in the pooled analysis. Where this has occurred, the actual number of studies and subjects included in the analysis will be stated.

STUDY DETAILS Citation KS Gurusamy, D Sharma, BR Davidson (2009) Pharmacological interventions to decrease blood loss and blood transfusion requirements for liver resection. Cochrane Database of Systematic Reviews. 009,Issue 4.Art.No.: CD008085. DOI:10.1002/14651858. CD008085. Affiliation/Source of funds University Department of Surgery, Royal Free College and University College School of Medicine, London, UK Funding: None Level of evidence Location/setting Study design Systematic review including 6 trials (1 Level I Hospital aprotinin, 1 tranexamic acid, 1 aminocaproic acid; 3 non-relevant interventions) that investigated the effectiveness of pharmacological interventions to reduce blood loss and transfusion requirements in patients undergoing liver resection. Intervention Comparator Tranexamic acid, aprotinin, desmopressin, recombinant No treatment, placebo or active factor VIIa, antithrombin III Population characteristics Patients undergoing liver resection Length of follow-up Outcomes measured Perioperative mortality; survival; liver failure; perioperative Not specified morbidity; transfusion requirements; operating time; hospital stay; intensive therapy unit; blood loss; liver function; biochemical markers of liver injury INTERNAL VALIDITY Allocation Results Blinding analysis Treatment/ Follow-up measurement bias (ITT) All trials 1 trial free of baseline imbalance, 2 trials 3 studies Publication bias not Some adequately randomised. unclear assessed. Treatment patients blinded 2/3 trials had Studies included and data extracted or measurement bias excluded adequate independently by two reviewers. Risk of not apparent. from sequence bias assessed by two reviewers. analysis of generation and Meta-analysis methods as per the efficacy in 3/3 trials Cochrane Collaboration. Only one trials for included inadequately each comparison. studies reported (12/109)allocation aprotinin concealment study; 3/217 tranexamic acid; 0/60 desmopress Overall quality assessment (descriptive) Good. Comprehensive literature search carried out. Quality assessment undertaken. Single study only available for each comparison. Authors note all studies at high risk of bias. **RESULTS** Outcome Intervention group Comparator group Statistical significance Risk estimate (95% CI) Aprotinin vs placebo n/N (%) n/N (%) RR 1.18 (0.18, 7.48) Mortality 2/17 2/20 1 trial (N=37) P=0.86 (Phet=NA)

Incidence of blood transfusion 1 trial (N=97)	8/48	19/49		RR 0.43 (0.21, 0.89) P=0.02 (Phet=NA)
,	Mean ± SD	Mean ± S	D	
Operating time (mins) 1 trial (N=97)	232 ± SD 75	233 ± SD	71	MD -1.00 (-30.08, 28.08) P=0.95 (Phet=NA)
Operative blood loss (mL) 1 trial (N=97)	1217 ± SD 966	1653 ± 12	21	MD -436.00 (-873.67, 1.67) P=0.05 (Phet=NA)
Tranexamic acid vs placebo				
	n/N (%)	n/N (%)		
Mortality 1 trial (N=217)	0/109	0/108		RR 0.0 (0.0, 0.0)
Incidence of blood transfusion 1 trial (N=214)	0/108	17/106		RR 0.03 (0.00, 0.46) P=0.01 (Phet=NA)
,	Mean ± SD	Mean ± S	D	
Operating time (mins) 1 trial (N=214)	253.8 ± SD 126.7	306 ± 126	.7	MD -52.20 (-86.15, -18.25) P=0.003 (Phet=NA)
Hospital stay (days) 1 trial (N=NR)	8 ± 7.66	9 ± 7.66		MD NR P=0.34 (P <i>het</i> =NA)
Transection blood loss (mL) 1 trial (N=214)	190 ± 653	450 ± 653		MD -260.00 (-434.99, -85.01) P=0.0036 (Phet=NA)
Operative blood loss (mL) 1 trial (N=214)	300 ± SD 754	600 ± SD	754	MD -300.00 (-502.05, -97.95) P=0.0036 (Phet=NA)
Desmopressin vs placebo				
	n/N (%)	n/N (%)		
Incidence of blood transfusion 1 trial (N=59)	3/30 (10.0%)	5/29 (17.2	%)	RR 0.58 (0.15, 2.21) P=0.42 (Phet=NA)
Red cell transfusion (units) 1 trial (N=59)	0.23 ± SD 0.82	0.72 ± 2.0	9	SMD -0.31 (-0.82, 0.21) P=0.24 (Phet=NA)
	Mean ± SD	Mean ± S		
Operating time (minutes) 1 trial (N=59)	405 ± 162	435 ± 162		MD -30.00 (-112.69, 52.69) P=0.48 (P <i>het</i> =NA)
Transection blood loss (mL) 1 trial (N=59)	405 ± 1140	450 ± 114		MD -45.00 (-626.86, 536.86) P=0.88 (P <i>het</i> =NA)
Operative blood loss (mL) 1 trial (N=59)	832.5 ± 1426.7	800 ± 142		MD 32.50 (-695.69, 760.69) P=0.93 (Phet=NA)
Outcome	Clinical importance		Clinical rel	evance
Aprotinin vs placebo	1.5	, , ,	145"	
Mortality	4: Range of estimates include important effects, but also cor with no or harmful effect		1: Patient-re	elevant clinical outcome.
Incidence of blood transfusion	1: Clinically important <i>benefit</i> , confidence limit does not incluvalue (p=0.02)		1: Patient-re	elevant clinical outcome.
Operating time (mins)	4: Range of estimates include important effects, but also cor with no or harmful effect	npatible	1: Patient-re	elevant clinical outcome.
Operative blood loss (mL)	2: Point estimate indicates clir important effects, but range o also compatible with no clinica important effect (p=0.05)	f estimates	1: Patient-re	elevant clinical outcome.
Tranexamic acid vs placebo				

Mantality	4 Dange of estimates includes aliminally	1 Delicut relevant slinical systems
Mortality	4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
	important effects, but also compatible	
	with no or harmful effect	
Incidence of blood	1: Clinically important benefit,	1: Patient-relevant clinical outcome.
transfusion	confidence limit does not include null	
	value (p=0.01)	
Operating time (mins)	1: Clinically important benefit,	2: Predictive surrogate outcome.
	confidence limit does not include null	
	value (p=0.01)	
Hospital stay (days)	4: Range of estimates includes clinically	2: Predictive surrogate outcome.
(uaye)	important effects, but also compatible	2.1. Touroute ourrogate outdome.
	with no effect	
Transection blood loss (mL)	1: Clinically important <i>benefit</i> ,	1: Patient-relevant clinical outcome.
Transection blood loss (IIIE)	confidence limit does not include null	1. I dione relevant dimedi datedine.
	value (p=0.0036)	
Operative blood loss (mL)	1: Clinically important <i>benefit</i> ,	1: Patient-relevant clinical outcome.
Operative blood loss (IIIE)	confidence limit does not include null	1. I dicht-relevant cimical outcome.
	value (p=0.0036)	
Desmopressin vs placebo	ναίας (μ=0.0000)	
Incidence of blood	4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
transfusion	important effects, but also compatible	The distriction of the districti
tidisidsion	with no or harmful effect	
Red cell transfusion (units)	4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
red cell transidation (dritta)	important effects, but also compatible	1. I diletti relevant cimical outcome.
	with no or harmful effect	
Operating time (minutes)	4: Range of estimates includes clinically	2: Predictive surrogate outcome.
	important effects, but also compatible	2. Fredictive surrogate outcome.
Transation blanding (C.1)	with no or harmful effect	1 Delicutual containing a containing
Transection blood loss (mL)	4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
	important effects, but also compatible	
	with no or harmful effect	
Operative blood loss (mL)	4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
	important effects, but also compatible	
	with no or harmful effect	
EXTERNAL VALIDITY		

Generalisability

This systematic review focuses on patients who have undergone liver transection so may be generalisable only to this specific surgery group.

Applicability

This review does not report the locations of the included studies and as such the applicability of the results to the Australian setting is unclear.

Comments

The authors conclude that none of the included interventions seem to decrease perioperative morbidity or offer long-term survival benefit. Aprotinin and tranexamic seem to reduce blood transfusion requirements however as the data is based on few trials and small sample sizes the results should be interpreted with caution.

Abbreviations: CI, confidence interval; het, heterogeneity; ITT, intention to treat; MD, mean difference; NA, not applicable; RCT, randomised controlled trial;

RR, risk ratio; SD, standard deviation; SMD, standardised mean difference.

Citation

DA Henry, PA Carless, AJ Moxey et al (2007) Antifibrinolytic use for minimising perioperative allogenic blood transfusion. Cochrane Database of Systematic Reviews 2007, Issue 4. Art. No.: CD001886. DOI: 10.1002/14651858.CD001886.pub2.

Affiliation/Source of funds

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Funding: Special purpose grant, Hunter Area Pathology Service, Australia (Internal grant); Australian Health Minister's Advisory Committee. NHMRC, Australia.

Advisory Committee. NHMRC, Australia	a.			
Study design	Level of evidence)	Location/setting	
Systematic review including 211 placebo-controlled and head-to-head trials (116 aprotinin, 45 tranexamic acid and 11 aminocaproic acid) that investigated the effectiveness and safety of pharmacological interventions for minimising perioperative allogenic blood transfusion.	Level I		Hospital	
Intervention		Comparator		
Aprotinin, tranexamic acid, ε-aminocapr	Aprotinin, tranexamic acid, ε-aminocaproic acid		Placebo/no treatment or active	
Population characteristics				
Adult surgical patients (or patients < 18	years if undergoing	a procedure predominantly performed in adults)		
Length of follow-up		Outcomes measured		
Not specified		Primary outcomes:		
		Proportion of patients transfused with allogenic blood,		
		autologous blood, or both; amount of allogenic and autologous		
		blood transfused; per	rioperative blood loss.	
		Secondary outcomes		
			bleeding; mortality; postoperative	
			ardial infarction, stroke, deep vein	
		thrombosis, pulmonary embolism, any thrombosis, renal		
		failure); length of hos	pital stay.	

INTERNAL VALIDITY				
Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
All trials randomised. 56 trials had adequate allocation method and remaining trials did not. 32% judged to have adequate concealment, 14% did not have adequate concealment and 53% were not adequately described.	Comparison of baseline characteristics not assessed. Studies included and data extracted independently by at least two reviewers. Risk of bias assessed by two reviewers. (note: 16 trials not assessed for quality by both reviewers) Meta-analysis methods as per the Cochrane Collaboration.	70% trials double- blind, 7% double- blind but method unclear, 24% not double-blind	Possible evidence of publication bias particularly with aprotinin trials but further investigation revealed unlikely to overestimate results and not produce a false-positive result.	45% of trials reported no exclusions or used ITT, 37% exclusions were judged not likely to cause bias, 17% exclusions judged excessive and likely to cause bias, or not reported.

Overall quality assessment (descriptive)

Good. Comprehensive literature search carried out. Quality assessment undertaken. Subgroup analyses performed on *a priori* categories including surgery, transfusion protocol, dose and trial quality.

RESULTS			
Outcome	Intervention group	Comparator group	Statistical significance Risk estimate (95% CI)
Aprotinin vs placebo/no tre			
Exposed to allogenic bloo			
	n/N (%)	n/N (%)	
99 trials (N=10,144) 96 trials (N=9949) ^a	2521/5750 (43.8)	2827/4394 (64.3)	All studies RR 0.66 (0.62, 0.71) P<0.001 (Phet<0.001)
77 trials (N=8837) 76 trials (N=8793)	2279/5003 (45.6)	2535/3834 (66.1)	Cardiac surgery RR 0.66 (0.61, 0.72) P<0.001 (Phet<0.001)
14 trials (N=794) 13 trials (N=771)	111/480 (23.1)	138/314 (43.9)	Orthopaedic surgery RR 0.69 (0.56, 0.85) P<0.001 (Phet=0.23)
2 trials (N=62)	4/30 (13.3)	16/32 (50.0)	Thoracic surgery RR 0.28 (0.11, 0.74) P=0.011(Phet=0.54)
2 trials (N=188) 1 trial (N=60)	94/105 (89.5)	77/83 (92.8)	Vascular surgery RR 1.01 (0.72, 1.40) P=0.98 (Phet=1.00)
2 trials (N=177)	21/87 (24.1)	39/90 (43.3)	Liver surgery RR 0.58 (0.37, 0.90) P=0.015 (Phet=0.31)
1 trial (N=56)	11/30 (36.7)	13/26 (50.0)	Neuro surgery RR 0.73 (0.40, 1.35) P=0.32 (Phet=NA)
1 trial (N=30)	1/15 (6.7)	9/15 (60.0)	Orthognathic surgery 0.11 (0.02, 0.77) P=0.026 (Phet=NA)
16 trials (N=1251)	345/649 (53.2)	394/602 (65.4)	Prime dose RR 0.83 (0.71, 0.96) P=0.014 (Phet<0.001)

46 trials (N=3268)	648/1733 (37.4)	882/1535 (57.5)	Low dose
43 trials (N=3073)			RR 0.66 (0.59, 0.74) P<0.001 (Phet<0.001)
56 trials (N=6569)	1522/3320 (45.8)	2204/3249 (67.8)	High dose
			RR 0.65 (0.60, 0.71)
15 trials (N=1191)	317/610 (52.0)	379/581 (65.2)	P<0.001 (Phet<0.001) Cardiac surgery and prime dose
13 (11415 (11=1191)	317/010 (32.0)	379/301 (03.2)	RR 0.81 (0.69, 0.96)
			P=0.012 (Phet<0.001)
25 trials (N=2039)	438/1043 (42.0)	605/996 (60.7)	Cardiac surgery and low dose
24 trials (N=1995)			RR 0.67 (0.58, 0.77) P<0.001 (Phet<0.001)
55 trials (N=6533)	1518/3302 (46.0)	2193/3231 (67.9)	Cardiac surgery and high dose
55 thais (14-6555)	1010/0302 (40.0)	2173/3231 (07.7)	RR 0.66 (0.60, 0.72)
			P<0.001 (Phet<0.001)
Units of allogenic blood t		T	1
(2 Ed. M. (220)	Mean ± SD	Mean ± SD	All and broken
63 trials (N=6820)	NR	NR	All patients WMD -1.07 (-1.31, -0.83)
			P<0.001 (Phet<0.001)
38 trials (N=3388)	NR	NR	Transfused patients
35 trials (N=3363)			WMD -0.96 (-1.24, -0.68)
Diagram (tatal)			P<0.001 (Phet<0.001)
Blood loss (total)	Mean ± SD	Mean ± SD	
15 trials (N=1577)	NR	NR	All studies
			WMD -414.48 (-520.13, -308.82)
			P<0.001 (Phet=0.003)
5 trials (N=1147)	NR	NR	Cardiac surgery
			WMD -489.06 (-571.32, -406.80) P<0.001 (Phet=0.62)
10 trials (N=430)	NR	NR	Orthopaedic surgery
,			WMD -399.09 (-562.81, -235.37)
			P<0.001 (Phet=0.01)
Blood loss (intraoperative	,	Mean ± SD	
13 trials (N=722)	Mean ± SD NR	NR	All studies
10 thais (14 722)		TVIX	WMD -185.32 (-280.23, -90.41)
			P<0.001 (Phet<0.001)
5 trials (N=360)	NR	NR	Cardiac surgery
			WMD -140.00 (-244.42, -35.59) P=0.0086 (P=0.01)
5 trials (N=201)	NR	NR	Orthopaedic surgery
, ,			WMD -151.05 (-317.63, 15.52)
4 1 1 (1) 0 (1)	ND.		P=0.076 (Phet=0.16)
1 trial (N=24)	NR	NR	Thoracic surgery WMD -532.0 (-863.00, -199.00)
			P=0.0016 (Phet=NA)
2 trials (N=137)	NR	NR	Liver surgery
			WMD -1200.40 (-2943.39, -542.59)
Pland lass (nactor and)	0)		P=0.18 (P <i>het</i> =0.02)
Blood loss (postoperative	Mean ± SD	Mean ± SD	
79 trials (N=7414)	NR	NR	All studies
			WMD -358403.64, -312.62)
			P<0.001 (Phet<0.001)

68 trials (N=6948)	NR	NR	Cardiac surgery
, ,			WMD -385.43 (-432.36, -338.50)
			P<0.001 (Phet<0.001)
7 trials (N=318)	NR	NR	Orthopaedic surgery
			WMD -113.58 (-223.69, -3.46)
4 14 1 (N - 04)	ND	ND	P=0.043 (Phet=0.005)
1 trial (N=24)	NR	NR	Thoracic surgery
			WMD -441.0 (-786.40, -95.60)
1 trial (N=30)	NR	NR	P=0.012 (Phet=NA) Orthognathic surgery
i iiiai (iv=30)	INK	IVIX	WMD -513.0 (-717.21, -308.79)
			P<0.001 (Phet=NA)
1 trial (N=44)	NR	NR	Liver surgery
			WMD -105.0 (-194.36, -15.64)
			P=0.021 (Phet=NA)
1 trial (N=50)	NR	NR	Vascular surgery
			WMD -203.00 (-404.93, -1.07)
			P=0.049 (P <i>het</i> =NA)
15 trials (N=1158)	NR	NR	Cardiac surgery and prime dose
			WMD -343.08 (-458.13, -228.04)
	ND.	ND	P<0.001 (Phet<0.001)
21 trials (N=1781)	NR	NR	Cardiac surgery and low dose
			WMD -293.24 (-348.67, -237.81) P<0.001 (Phet<0.001)
48 trials (N=4819)	NR	NR	Cardiac surgery and high dose
40 tilais (IV-4017)	IVIX	IVIX	WMD -428.09 (-485.38, -370.80)
			P<0.001 (Phet<0.001)
Re-operation for bleeding			1 30.001 (1 110130.001)
	n/N (%)	n/N (%)	
51 trials (N=5384)	58/3030 (1.9)	110/2354 (4.7)	All trials
36 trials (N=4715)			RR 0.48 (0.35, 0.68)
			P<0.001 (Phet=0.51)
47 trials (N=5153)	55/2915 (1.9)	101/2238 (4.5)	Cardiac surgery
33 trials (N=4534)			RR 0.49 (0.34, 0.70)
Mantalita			P<0.001 (Phet=0.41)
Mortality	n/N (%)	n/N (%)	
52 trials (N=7721)	105/4319 (2.4)	87/3402 (2.6)	All trials
37 trials (N=6645)	103/4317 (2.4)	07/3402 (2.0)	RR 0.90 (0.67, 1.20)
37 that3 (14-00+3)			P=0.47 (Phet=0.95)
45 trials (N=7078)	99/3907 (2.5)	77/3171 (2.4)	` ,
45 trials (N=7078) 31 trials (N=6058)	99/3907 (2.5)	77/3171 (2.4)	Cardiac surgery
45 trials (N=7078) 31 trials (N=6058)	99/3907 (2.5)	77/3171 (2.4)	` ,
		77/3171 (2.4)	Cardiac surgery RR 0.95 (0.70, 1.28)
31 trials (N=6058) Myocardial infarction	n/N (%)	n/N (%)	Cardiac surgery RR 0.95 (0.70, 1.28) P=0.72 (Phet=0.93)
31 trials (N=6058) Myocardial infarction 40 trials (N=6107)			Cardiac surgery RR 0.95 (0.70, 1.28) P=0.72 (Phet=0.93) All trials
31 trials (N=6058) Myocardial infarction	n/N (%)	n/N (%)	Cardiac surgery RR 0.95 (0.70, 1.28) P=0.72 (Phet=0.93) All trials RR 0.92 (0.72, 1.18)
31 trials (N=6058) Myocardial infarction 40 trials (N=6107) 34 trials (N=5758)	n/N (%) 153/3523 (4.3)	n/N (%) 118/2584 (4.6)	Cardiac surgery RR 0.95 (0.70, 1.28) P=0.72 (Phet=0.93) All trials RR 0.92 (0.72, 1.18) P=0.50 (Phet=0.91)
31 trials (N=6058) Myocardial infarction 40 trials (N=6107) 34 trials (N=5758) 37 trials (N=5628)	n/N (%)	n/N (%)	Cardiac surgery RR 0.95 (0.70, 1.28) P=0.72 (Phet=0.93) All trials RR 0.92 (0.72, 1.18) P=0.50 (Phet=0.91) Cardiac surgery
31 trials (N=6058) Myocardial infarction 40 trials (N=6107) 34 trials (N=5758)	n/N (%) 153/3523 (4.3)	n/N (%) 118/2584 (4.6)	Cardiac surgery RR 0.95 (0.70, 1.28) P=0.72 (Phet=0.93) All trials RR 0.92 (0.72, 1.18) P=0.50 (Phet=0.91) Cardiac surgery RR 0.95 (0.74, 1.22)
31 trials (N=6058) Myocardial infarction 40 trials (N=6107) 34 trials (N=5758) 37 trials (N=5628) 31 trials (N=5279)	n/N (%) 153/3523 (4.3)	n/N (%) 118/2584 (4.6)	Cardiac surgery RR 0.95 (0.70, 1.28) P=0.72 (Phet=0.93) All trials RR 0.92 (0.72, 1.18) P=0.50 (Phet=0.91) Cardiac surgery
31 trials (N=6058) Myocardial infarction 40 trials (N=6107) 34 trials (N=5758) 37 trials (N=5628)	n/N (%) 153/3523 (4.3) 152/3204 (4.7)	n/N (%) 118/2584 (4.6) 113/2424 (4.7)	Cardiac surgery RR 0.95 (0.70, 1.28) P=0.72 (Phet=0.93) All trials RR 0.92 (0.72, 1.18) P=0.50 (Phet=0.91) Cardiac surgery RR 0.95 (0.74, 1.22)
31 trials (N=6058) Myocardial infarction 40 trials (N=6107) 34 trials (N=5758) 37 trials (N=5628) 31 trials (N=5279) Stroke	n/N (%) 153/3523 (4.3) 152/3204 (4.7)	n/N (%) 118/2584 (4.6) 113/2424 (4.7) n/N (%)	Cardiac surgery RR 0.95 (0.70, 1.28) P=0.72 (Phet=0.93) All trials RR 0.92 (0.72, 1.18) P=0.50 (Phet=0.91) Cardiac surgery RR 0.95 (0.74, 1.22) P=0.69 (Phet=0.92)
31 trials (N=6058) Myocardial infarction 40 trials (N=6107) 34 trials (N=5758) 37 trials (N=5628) 31 trials (N=5279)	n/N (%) 153/3523 (4.3) 152/3204 (4.7)	n/N (%) 118/2584 (4.6) 113/2424 (4.7)	Cardiac surgery RR 0.95 (0.70, 1.28) P=0.72 (Phet=0.93) All trials RR 0.92 (0.72, 1.18) P=0.50 (Phet=0.91) Cardiac surgery RR 0.95 (0.74, 1.22)

11 trials (N=1303) 9 trials (N=1163)	10/773 (1.3)	10/530 (1.9)	Cardiac surgery RR 0.76 (0.30, 1.93)
			P=0.57 (Phet=0.40)
Deep vein thrombosis			
	n/N (%)	n/N (%)	
15 trials (N=1104)	36/679 (5.3)	23/425 (5.4)	All trials
11 trials (N=986)			RR 0.79 (0.46, 1.34) P=0.38 (P <i>het</i> =0.80)
2 trials (N=272)	4/170 (2.4)	1/102 (1.0)	Cardiac surgery
			RR 2.52 (0.41, 15.45) P=0.32 (Phet=0.71)
Pulmonary embolism	1	,	,
-	n/N (%)	n/N (%)	
3 trials (N=233)	4/129 (3.1)	2/104 (1.9)	All trials
2 trials (N=175)			RR 1.98 (0.38, 10.46)
			P=0.42 (Phet=0.95)
Other thrombosis			
	n/N (%)	n/N (%)	
9 trials (N=736)	5/402 (1.2)	8/334 (2.4)	All trials
7 trials (N=583)			RR 0.73 (0.25, 2.15)
			P=0.57 (P <i>het</i> =0.64)
4 trials (N=426)	2/245 (0.8)	4/181 (2.2)	Cardiac surgery
3 trials (N=370)			RR 0.62 (0.11, 3.36)
			P=0.58 (P <i>het</i> =0.50)
Coronary artery graft occi		T	
	n/N (%)	n/N (%)	
2 trials (N=728)	54/369 (14.6)	39/359 (10.9)	Cardiac surgery
			RR 0.76 (0.10, 5.67)
D 16" /1 6 "			P=0.79 (Phet=0.13)
Renal failure/dysfunction	/N1 /O/)	/N1 /O/)	
21 totals (NL 4412)	n/N (%)	n/N (%)	All totals
21 trials (N=4412)	75/2525 (3.0)	42/1887 (2.2)	All trials
14 trials (N=3908)			RR 1.16 (0.79, 1.70)
18 trials (N=4174)	68/2395 (2.9)	39/1779 (2.2)	P=0.46 (Phet=0.88) Cardiac surgery
11 trials (N=3670)	00/2393 (2.9)	39/11/19 (2.2)	RR 1.12 (0.74, 1.67)
11 tilais (N=3070)			P=0.60 (P <i>het</i> =0.85)
Hospital length of stay (da	avs)		1 -0.00 (1 Het-0.03)
Trospital length of stay (ut	Mean ± SD	Mean ± SD	
17 trials (N=1570)	NR	NR	All trials
17 tilais (11–1570)	IVIC	IVIX	WMD -0.01 (-0.50, 0.48)
			P=0.96 (Phet=0.19)
13 trials (N=1412)	NR	NR	Cardiac surgery
			WMD -0.10 (-0.64, 0.44)
			P=0.73 (Phet=0.12)
Tranexamic acid vs place	bo	<u> </u>	, , ,
Exposed to allogenic bloc			
	n/N (%)	n/N (%)	
53 trials (N=3836)	546/2020 (27.0)	796/1816 (43.8)	All trials
51 trials (N=3751)	, ,	, ,	RR 0.61 (0.54, 0.70)
			P<0.001 (Phet<0.001)
29 trials (N=2488)	367/1322 (27.8)	476/1166 (40.8)	Cardiac surgery
28 trials (N=2443)		, , ,	
20 IIIais (IV=2443)			RR 0.69 (0.60, 0.79)

21 trials (N=993)	139/520 (26.7)	247/473 (52.2)	Orthopaedic surgery
20 trials (N=953)	137/320 (20.7)	2471473 (32.2)	RR 0.44 (0.33, 0.60)
20 11415 (14 700)			P<0.001 (Phet<0.001)
2 trials (N=296)	29/148 (19.6)	54/148 (36.5)	Liver surgery
			RR 0.16 (0.00, 32.47)
			P=0.50 (Phet<0.001)
1 trial (N=59)	11/30 (36.7)	19/29 (65.5)	Vascular surgery
			RR 0.56 (0.33, 0.96)
			P=0.035 (Phet=NA)
16 trials (N=926)	162/495 (32.7)	204/431 (47.3)	Cardiac surgery/total dose < 2.0 g
			RR 0.72 (0.59, 0.88)
			P=0.0013 (Phet=0.05)
14 trials (N=1616)	205/827 (24.8)	286/789 (36.2)	Cardiac surgery/total dose 2-10 g
13 trials (N=1571)			RR 0.67 (0.55, 0.83)
			P<0.001 (Phet=0.09)
Units of allogenic blood to		T	
4 () () () ()	Mean ± SD	Mean ± SD	
16 trials (N=1071)	NR	NR	All patients
14 trials (N=965)			WMD -1.12 (-1.59, -0.64)
11 totals (N. 400)	ND	ND	P<0.001 (Phet<0.001)
11 trials (N=429)	NR	NR	<i>Transfused patients</i> WMD -0.51 (-1.06, 0.04)
			P=0.071 (Phet<0.001)
Blood loss (total)			P=0.071 (PHet<0.001)
biood ioss (total)	Mean ± SD	Mean ± SD	
18 trials (N=955)	NR	NR	All studies
10 (11a13 (11-755)	IVIX	IVIX	WMD -443.53 (-572.08, -314.98)
			P<0.001 (Phet<0.001)
3 trials (N=245)	NR	NR	Cardiac surgery
0 thdis (14 2 10)	TWI C	TWI C	WMD -439.82 (-606.50, -273.15)
			P<0.001 (Phet=0.82)
14 trials (N=690)	NR	NR	Orthopaedic surgery
(WMD -439.51 (-590.93, -288.09)
			P<0.001 (Phet<0.001)
1 trial (N=20)	NR	NR	Liver surgery
, ,			WMD -6552.0 (-14329.54,
			1225.54)
			P=0.099 (P <i>het</i> =NA)
Blood loss (intraoperative			
12.11.7	Mean ± SD	Mean ± SD	
10 trials (N=553)	NR	NR	All studies
			WMD -54.89 (-105.31, -4.48)
0.12-1- (N. 444)	ND	MD	P=0.033 (Phet=0.26)
3 trials (N=144)	NR	NR	Cardiac surgery
			WMD -287.16 (-481.57, -92.75)
7 trials (NL 400)	ND	ND	P=0.0038 (Phet=0.66)
7 trials (N=409)	NR	NR	Orthopaedic surgery
			WMD -29.52 (-69.17, 10.14) P=0.14 (Phet=0.69)
Blood loss (postoperative	<u> </u>		F -0.14 (F1161=0.07)
טוטטע וטאא (אָטאַנע דייטטע ווער ווער	Mean ± SD	Mean ± SD	
23 trials (N=1423)	NR	NR	All studies
25 mais (N-1425)	IVIX	IVIX	WMD -247.90 (-313.07, -182.73)
			P<0.001 (Phet<0.001)
	<u> </u>		1 10.001 (1 110110.001)

17 trials (N=1130)	NR	NR	Cardiac surgery WMD -262.60 (-318.62, -206.59) P<0.001 (Phet=0.01)
6 trials (N=293)	NR	NR	Orthopaedic surgery WMD -209.72 (-384.28, -35.16) P=0.019 (Phet<0.001)
9 trials (N=302)	NR	NR	Cardiac surgery/total dose < 2.0 g WMD -251.77 (-352.27, -151.26) P<0.001 (Phet=0.07)
8 trials (N=828)	NR	NR	Cardiac surgery/total dose 2.0-10.0 g WMD -272.85 (-340.79, -204.90) P<0.001 (Phet=0.03)
Re-operation for bleeding	1	 	
<u> </u>	n/N (%)	n/N (%)	
20 trials (N=1676) 18 trials (N=1598)	25/872 (2.9)	40/804 (5.0)	All studies RR 0.67 (0.41, 1.09) P=0.11 (Phet=0.92)
19 trials (N=1618) 17 trials (N=1540)	23/843 (2.7)	38/775 (4.9)	Cardiac surgery RR 0.65 (0.39, 1.08) P=0.097 (Phet=0.90)
Mortality			·
-	n/N (%)	n/N (%)	
24 trials (N=2210) 16 trials (N=1684)	14/1129 (1.2)	26/1081 (2.4)	All studies RR 0.60 (0.32, 1.12) P=0.11 (Phet=0.84)
18 trials (N=1702) 11 trials (N=1390)	8/872 (0.9)	16/830 (1.9)	Cardiac surgery RR 0.55 (0.24, 1.25) P=0.15 (Phet=0.73)
Myocardial infarction		 	
,	n/N (%)	n/N (%)	
17 trials (N=1718) 12 trials (N=1344)	15/885 (1.7)	16/833 (1.9)	All studies RR 0.96 (0.48, 1.90) P=0.91 (Phet=0.96)
15 trials (N=1632) 9 trials (N=1048)	13/841 (1.5)	15/791 (1.9)	Cardiac surgery 0.91 (0.44, 1.88) P=0.79 (Phet=0.91)
Stroke			
	n/N (%)	n/N (%)	
14 trials (N=1403) 7 trials (N=937)	10/740 (1.4)	7/663 (1.1)	All studies RR 1.25 (0.47, 3.31) P=0.65 (Phet=0.79)
13 trials (N=1345) 5 trials (N=841)	9/711 (1.3)	5/634 (0.8)	Cardiac surgery RR 1.52 (0.52, 4.41) P=0.44 (Phet=0.78)
Deep vein thrombosis	·		<u> </u>
	n/N (%)	n/N (%)	
18 trials (N=1109) 10 trials (N=681)	11/565 (1.9)	16/544 (2.9)	All studies RR 0.77 (0.37, 1.61) P=0.49 (Phet=0.81)
4 trials (N=442) 2 trials (N=291)	0/221 (0.0)	2/201 (1.0)	Cardiac surgery RR 0.37 (0.04, 3.47) P=0.38 (Phet=0.95)
Pulmonary embolism			
	n/N (%)	n/N (%)	

Units of allogenic blood	transfused		P=0.33 (P <i>net</i> =NA)
			RR 0.93 (0.80, 1.08) P=0.33 (P <i>het</i> =NA)
1 trial (N=82)	36/42 (85.7)	37/40 (92.5)	Liver surgery
			RR 0.96 (0.61, 1.50) P=0.85 (P <i>het</i> =0.64)
3 trials (N=122)	20/59 (33.9)	23/63 (36.5)	Orthopaedic surgery
			RR 0.65 (0.47, 0.91) P=0.011 (Phet=0.11)
10 trials (N=597)	82/313 (26.2)	113/284 (39.8)	Cardiac surgery
			RR 0.75 (0.58, 0.96) P=0.023 (Phet=0.03)
14 trials (N=801)	138/414 (33.3)	173/387 (44.7)	All studies
Exposed to dilogerile bit	n/N (%)	n/N (%)	
Exposed to allogenic blo			
E-aminocaproic acid vs	nlacoho		P=0.31 (P <i>he</i> t=0.64)
2 trials (N=116)	NR	NR	Cardiac surgery RR -0.23 (-0.67, 0.21)
2 trials (N. 117)	MD	ND	P=0.14 (Phet=0.66)
4 trials (N=176)	NR	NR	All studies RR -0.30 (-0.71, 0.10)
, ,	Mean ± SD	Mean ± SD	
Hospital length of stay		I	1. 0.00 (1.101 0.07)
4 trial (N=400)			RR 0.73 (0.16, 3.32) P=0.68 (Phet=0.69)
5 trials (N=444)	2/222 (0.9)	3/222 (1.4)	Cardiac surgery
Koriai Taliai 6/Uysiui 10101	n/N (%)	n/N (%)	
Renal failure/dysfunction	n		P=0.32 (P <i>het</i> =0.80)
2 trials (N=114)			RR 2.10 (0.49, 8.99)
7 trials (N=289)	5/148 (3.4)	2/141 (1.4)	All studies
Carol unionibosis	n/N (%)	n/N (%)	
Other thrombosis			P=0.34 (P <i>het</i> =0.98)
2 trials (N=289)			RR 0.33 (0.04, 3.15)
6 trials (N=569)	0/298 (0.0)	2/271 (0.7)	Cardiac surgery
7 triais (11–300)			P=0.31 (Phet=0.93)
13 trials (N=946) 7 <i>trials (N=568)</i>	2/487 (0.4)	6/459 (1.3)	All studies RR 0.55 (0.17, 1.76)

4 trials (N=171)	NR	NR	All studies WMD -142.02 (-284.95, 0.92)
			P=0.051 (Phet=0.19)
2 trials (N=79)	NR	NR	Cardiac surgery
			WMD -213.58 (-310.03, -117.13) P<0.001 (Phet=0.73)
2 trials (N=92)	NR	NR	Orthopaedic surgery
, ,			WMD 10.94 (-259.66, 281.54) P=0.94 (Phet=0.26)
Blood loss (postoperative)		· · · · · · · · · · · · · · · · · · ·
	Mean ± SD	Mean ± SD	
12 trials (N=940)	NR	NR	All studies WMD -202.08 (-273.64, -130.53) P<0.001 (Phet<0.001)
11 trials (N=894)	NR	NR	Cardiac surgery WMD -196.27 (-271.75, -120.79) P<0.001 (Phet<0.001)
1 trial (N=46)	NR	NR	Orthopaedic surgery WMD -276.00 (-448.83, -103.17) P=0.0017 (Phet=NA)
Re-operation for bleeding	1		, , ,
	n/N (%)	n/N (%)	
7 trials (N=740)	3/379 (0.8)	12/361 (3.3)	Cardiac surgery
5 trials (N=662)			RR 0.35 (0.11, 1.17) P=0.087 (Phet=0.78)
Mortality			,
	n/N (%)	n/N (%)	
6 trials (N=754)	10/388 (2.6)	7/366 (1.9)	All studies
5 trials (N=714)			RR 1.17 (0.47, 2.93) P=0.73 (Phet=0.78)
5 trials (N=672)	7/346 (2.0)	3/326 (0.9)	Cardiac surgery
4 trials (N=632)			RR 1.65 (0.50, 5.43) P=0.41 (P <i>het</i> =0.81)
Myocardial infarction	1		
	n/N (%)	n/N (%)	
5 trials (N=662) 4 trials (N=632)	12/340 (3.5)	4/322 (1.2)	Cardiac surgery RR 0.89 (0.37, 2.18) P=0.80 (Phet=0.33)
Stroke			P=0.00 (PHet=0.55)
Siroke	n/N (%)	n/N (%)	
6 trials (N=702)	2/361 (0.6)	3/341 (0.9)	Cardiac surgery
3 trials (N=541)	2.00. (0.0)	0,011 (017)	RR 0.59 (0.10, 3.44) P=0.55 (Phet=0.47)
Deep vein thrombosis	I	<u>L</u>	1
1	n/N (%)	n/N (%)	
3 trials (N=122)	3/59 (5.1)	3/63 (4.8)	All studies
1 trial (N=46)			RR 1.09 (0.25, 4.85) P=0.91 (Phet=1.00)
Pulmonary embolism	<u> </u>		
	n/N (%)	n/N (%)	
2 trials (N=92)	0/44 (0.0)	1/48 (2.1)	All studies
1 trial (N=46)			RR 0.36 (0.02, 8.46) P=0.53 (P <i>het</i> =1.00)
Other thrombosis	IN 1 (0/2)	N1 (0/)	
	n/N (%)	n/N (%)	

1 trial (N=82)	2/42 (4.8)	2/40 (5.0)		All studies RR 0.95 (0.14, 6.44) P=0.96 (Phet=NA)
Hospital length of stay	1			1 –0.70 (1 11ct–11/h)
Trespitationg at easy	Mean ± SD	Mean ± SD		
1 trial (N=46)	11.9 ± 7.3	9 ± 5.9		All studies MD 2.90 (-0.96, 6.76) P=0.14 (Phet=NA)
Outcome	Clinical importance		Clinical rele	evance
Aprotinin vs placebo				
Exposed to allogenic blood	All studies 1: Clinically important benefit, confidence limit does not incluvalue (p<0.001) Cardiac surgery 1: Clinically important benefit, confidence limit does not incluvalue (p<0.001) Cardiac surgery and prime does not incluvalue (p=0.014) Cardiac surgery and low dose not incluvalue (p=0.014) Cardiac surgery and low dose not incluvalue (p<0.001) Cardiac surgery and high dose not incluvalue (p<0.001) Cardiac surgery and high dose not incluvalue (p<0.001) Cardiac surgery and high dose not incluvalue (p<0.001) Cardiac surgery not incluvalue (p<0.001) Cardiac surgery not high dose not incluvalue (p<0.001) Cardiac surgery 1: Clinically important benefit, confidence limit does not incluvalue (p<0.001) Thoracic surgery 1: Clinically important benefit, confidence limit does not incluvalue (p=0.011) Vascular surgery 4: Range of estimates includes important effects, but also conwith no or harmful effect Liver surgery 1: Clinically important benefit, confidence limit does not incluvalue (p=0.015) Neuro surgery 4: Range of estimates includes important effects, but also conwith no or harmful effect Corthognathic surgery 1: Clinically important benefit, confidence limit does not incluvalue (p=0.015) Neuro surgery 1: Clinically important benefit, confidence limit does not incluvalue (p=0.026)	de null sclinically apatible de null de null		levant clinical outcome.
Units of allogenic blood	All patients		1: Patient-re	levant clinical outcome.
transfused	1: Clinically important benefit,			

	confidence limit does not include null value (p<0.001) Transfused patients 1: Clinically important benefit, confidence limit does not include null	
	value (p<0.001)	
Blood loss (total)	All studies 1: Clinically important <i>benefit</i> , confidence limit does not include null value (p<0.001)	1: Patient-relevant clinical outcome.
	Cardiac surgery 1: Clinically important <i>benefit</i> , confidence limit does not include null	
	value (p<0.001) Orthopaedic surgery 1: Clinically important <i>benefit</i> , confidence limit does not include null	
Dlood loss /introor!:\	value (p<0.001)	1. Detient relevant elinical cuterior
Blood loss (intraoperative)	All studies 1: Clinically important benefit, confidence limit does not include null value (p<0.001) Cardiac surgery 1: Clinically important benefit, confidence limit does not include null value (p=0.0086) Orthopaedic surgery 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Thoracic surgery 1: Clinically important benefit, confidence limit does not include null value (p=0.0016)	1: Patient-relevant clinical outcome.
	Liver surgery	
	1: Clinically important <i>benefit</i> , confidence limit does not include null	
	value (p<0.001)	
Blood loss (postoperative)	All studies 1: Clinically important benefit, confidence limit does not include null value (p<0.001) Cardiac surgery 1: Clinically important benefit, confidence limit does not include null	1: Patient-relevant clinical outcome.
	value (p<0.001) Cardiac surgery and prime dose 1: Clinically important benefit,	
	confidence limit does not include null value (p<0.001) Cardiac surgery and low dose	
	1: Clinically important <i>benefit</i> , confidence limit does not include null value (p<0.001)	
	Cardiac surgery and high dose 1: Clinically important benefit, confidence limit does not include null	

	value (p<0.001) Orthopaedic surgery 2: Clinically important benefit, but confidence limit may include non-clinically important benefit (p=0.043) Thoracic surgery 1: Clinically important benefit, confidence limit does not include null value (p=0.012) Orthognathic surgery 1: Clinically important benefit, confidence limit does not include null value (p<0.001) Liver surgery 1: Clinically important benefit, confidence limit does not include null value (p=0.021) Vascular surgery 2: Clinically important benefit, but confidence limit may include non-clinically important benefit, on 0.040)	
Re-operation for bleeding	clinically important benefit (p=0.049) All studies 1: Clinically important benefit, confidence limit does not include null value (p<0.001) Cardiac surgery 1: Clinically important benefit, confidence limit does not include null value (p<0.001)	1: Patient-relevant clinical outcome.
Mortality	All studies 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Cardiac surgery 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Myocardial infarction	All studies 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Cardiac surgery 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Stroke	All studies 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Cardiac surgery 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Deep vein thrombosis	All studies 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.

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	Cardiac surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
Pulmonary embolism	All studies	1: Patient-relevant clinical outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
Other thrombosis	All studies	1: Patient-relevant clinical outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
	Cardiac surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
Coronary artery graft	Cardiac surgery	1: Patient-relevant clinical outcome.
Coronary artery graft occlusion	A. Dange of estimates includes alinically	1: Patient-relevant clinical outcome.
OCCIUSION	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
Renal failure/dysfunction	All studies	1: Patient-relevant clinical outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
	Cardiac surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
Hospital length of stay	All studies	2: Predictive surrogate outcome.
3	4: Range of estimates includes clinically]
	important effects, but also compatible	
	with no or harmful effect	
	Cardiac surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
Tranexamic acid vs placebo	With no or naminal effect	
Exposed to allogenic blood	All studies	1: Patient-relevant clinical outcome.
Exposed to allogeriic blood		1. Patient-relevant clinical outcome.
	1: Clinically important <i>benefit</i> ,	
	confidence limit does not include null	
	value (p<0.001)	
	Cardiac surgery	
	1: Clinically important benefit,	
	confidence limit does not include null	
	value (p<0.001)	
	Cardiac surgery and dose <2.0 g	
	1: Clinically important benefit,	
	confidence limit does not include null	
	value (p=0.0013)	
	Cardiac surgery and dose 2-10 g	
	1: Clinically important benefit,	
	confidence limit does not include null	
	value (p<0.001)	
	Orthopaedic surgery	
	1: Clinically important <i>benefit</i> ,	
	confidence limit does not include null	
	value (p<0.001)	
	ναια ς (μ<υ.υυ1)	

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	Vascular surgery	
	1: Clinically important benefit,	
	confidence limit does not include null	
	value (p=0.035)	
	Liver surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
Units of allogenic blood	All patients	1: Patient-relevant clinical outcome.
transfused	1: Clinically important <i>benefit</i> ,	
i ansidsed	confidence limit does not include null	
	value (p<0.001)	
	Transfused patients	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no effect	
Blood loss (total)	All studies	1: Patient-relevant clinical outcome.
	1: Clinically important benefit,	
	confidence limit does not include null	
	value (p<0.001)	
	Cardiac surgery	
	1: Clinically important benefit,	
	confidence limit does not include null	
	value (p<0.001)	
	Orthopaedic surgery	
	1: Clinically important <i>benefit</i> ,	
	confidence limit does not include null	
	value (p<0.001)	
	Liver surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
Disadisa (intro or sortice)	with no effect	1 Delicut velevent elipical automas
Blood loss (intraoperative)	All studies	1: Patient-relevant clinical outcome.
	1: Clinically important benefit,	
	confidence limit does not include null	
	value (p=0.033)	
	Cardiac surgery	
	1: Clinically important benefit,	
	confidence limit does not include null	
	value (p=0.0038)	
	Orthopaedic surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
Blood loss (postoperative)	All studies	1: Patient-relevant clinical outcome.
(2.20.2b.c.a)	1: Clinically important <i>benefit</i> ,	3
	confidence limit does not include null	
	value (p<0.001)	
	Cardiac surgery	
	1: Clinically important <i>benefit</i> ,	
	confidence limit does not include null	
	value (p<0.001)	
	Cardiac surgery and dose < 2 g	
	1: Clinically important benefit,	
	confidence limit does not include null	
l .		•
	value (p<0.001) Cardiac surgery and dose 2-10 g	

	1	
	1: Clinically important <i>benefit</i> , confidence limit does not include null	
	value (p<0.001) Orthopaedic surgery	
	1: Clinically important benefit,	
	confidence limit does not include null	
Re-operation for bleeding	value (p=0.019) All studies	1: Patient-relevant clinical outcome.
re-operation for bleeding	4: Range of estimates includes clinically	1. I attent-relevant clinical outcome.
	important effects, but also compatible	
	with no or harmful effect	
	Cardiac surgery 4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
Mortality	All studies	1: Patient-relevant clinical outcome.
	4: Range of estimates includes clinically important effects, but also compatible	
	with no or harmful effect	
	Cardiac surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible with no or harmful effect	
Myocardial infarction	All studies	1: Patient-relevant clinical outcome.
,	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
	Cardiac surgery 4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
Stroke	All studies4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
	important effects, but also compatible	
	with no or harmful effect	
	Cardiac surgery	
	4: Range of estimates includes clinically important effects, but also compatible	
	with no or harmful effect	
Deep vein thrombosis	All studies	1: Patient-relevant clinical outcome.
	4: Range of estimates includes clinically important effects, but also compatible	
	with no or harmful effect	
	Cardiac surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible with no or harmful effect	
Pulmonary embolism	All studies	1: Patient-relevant clinical outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect Cardiac surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
Other though !	with no or harmful effect	4 Delicated and all the last
Other thrombosis	All studies4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
	1 4. Nange of estimates includes clinically	

	important effects, but also compatible with no or harmful effect	
Renal failure/dysfunction	Cardiac surgery	1: Patient-relevant clinical outcome.
Tronar failur or a yoranotton	4: Range of estimates includes clinically	The distriction of the control of the control
	important effects, but also compatible	
	with no or harmful effect	
Hospital length of stay	All studies	2: Predictive surrogate outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
	Cardiac surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
E-aminocaproic acid vs place		1
Exposed to allogenic blood	All studies	1: Patient-relevant clinical outcome.
	1: Clinically important <i>benefit</i> ,	
	confidence limit does not include null	
	value (p=0.023)	
	Cardiac surgery 1: Clinically important benefit,	
	confidence limit does not include null	
	value (p=0.011)	
	Orthopaedic surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
	Liver surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
Units of allogenic blood	All patients	1: Patient-relevant clinical outcome.
transfused	1: Clinically important benefit,	
	confidence limit does not include null	
	value (p<0.001)	
	Transfused patients	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
Dland lane (total)	With no effect.	1. Detient relevant elinical euteeme
Blood loss (total)	Orthopaedic surgery 1: Clinically important benefit,	1: Patient-relevant clinical outcome.
	confidence limit does not include null	
	value (p=0.0084)	
Blood loss (intraoperative)	All studies	1: Patient-relevant clinical outcome.
blood loss (intraoperative)	4: Range of estimates includes clinically	1. I diletti felevarit elimedi odteome.
	important effects, but also compatible	
	with no effect.	
	Cardiac surgery	
	1: Clinically important benefit,	
	confidence limit does not include null	
	value (p<0.001)	
	Orthopaedic surgery	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
Blood loss (postoperative)	All studies	1: Patient-relevant clinical outcome.
	1: Clinically important benefit,	

	confidence limit does not include null value (p<0.001) Cardiac surgery 1: Clinically important benefit, confidence limit does not include null value (p<0.001) Orthopaedic surgery 1: Clinically important benefit, confidence limit does not include null value (p=0.0017)	
Re-operation for bleeding	Cardiac surgery 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Mortality	All studies 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Cardiac surgery 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Myocardial infarction	Cardiac surgery 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Stroke	Cardiac surgery 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Deep vein thrombosis	All studies 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Pulmonary embolism	All studies 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Other thrombosis	All studies 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Hospital length of stay	All studies 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	2: Predictive surrogate outcome.
EXTERNAL VALIDITY		

Generalisability

This systematic review focuses on patients who have undergone many different types on surgery so is generalisable to the broad population undergoing surgery. It also contains numerous subgroup analyses which would allow it to be generalisable to specific patient groups.

Applicability

Included studies were performed in a wide range of countries. Eight of the 211 included studies were conducted in Australia. Given the wide range of included studies, the results are likely to be applicable to the Australian setting.

Comments

The authors conclude that antifibrinolytics provide worthwhile reductions in blood loss and transfusion requirements while not appearing to be offset by any serious safety issues. They also state that the lysine analogues appear to be generally as effective as aprotinin, but they are cheaper.

Abbreviations: CI, confidence interval; het, heterogeneity; ITT, intention to treat; MD, mean difference; NA, not applicable; NR, not reported; RCT, randomised controlled trial; RR, risk ratio; SD, standard deviation; WMD, weighted mean difference.

^a Trials (N) in italics denotes the number of trials (N) included in the analysis. Trials in which the events occurred in 100% in both treatment arms, 0% in both

treatment arms or in which a SD was not available were not estimable and were not included in the analysis.

Citation

DA Henry, PA Carless, D Fergusson, et al (2009) The safety of aprotinin and lysine-derived antifibrinolytic drugs in cardiac surgery: a meta-analysis. CMAJ 180(2): 183-193.

Affiliation/Source of funds

School of Medicine and Public Health, University of New castle, Newcastle, Australia; Ottawa Health Research Institute, The Ottawa Hospital, Ottawa, Canada; Keenan Research Centre, Li Ka Shing Knowledge Institute, St Michael's Hospital; the Institute for Clinical Evaluative Sciences; Faculty of Medicine, University of Toronto, Toronto, Canada.

Funding: No specific funding.

Study design	Level of evidence	Location/setting
Update of cardiac surgery subgroup from Henry 2007 Cochrane review (10 additional RCTs for aprotinin, 2 additional trials for tranexamic acid and 1 additional trial from ε-aminocaproic acid).	Level I	Hospital

Intervention	Comparator
Aprotinin, tranexamic acid, ε-aminocaproic acid	Placebo/no treatment or active

Population characteristics

Adults undergoing non-urgent cardiac surgery

Length of follow-up	Outcomes measured
Not specified	Myocardial infarction, mortality

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
All trials randomised.	Comparison of baseline characteristics not assessed.	Not reported for additional studies.	No evidence of publication bias for	Not reported for
No further details for additional studies.	Studies included and data extracted independently by at least two reviewers. Risk of bias assessed by two reviewers. (note: 16 trials not assessed for quality by both reviewers)		these outcomes. No evidence of treatment or measurement bias.	additional studies.
	Meta-analysis methods as per the Cochrane Collaboration.			

Overall quality assessment (descriptive)

Good. Comprehensive literature search carried out. Quality assessment undertaken. Some information missing for additional studies in publications, however the majority of the included studies were described in detail in the original Cochrane review (Henry 2007).

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance Risk estimate (95% CI)			
Aprotinin vs placebo/no treatment						

	n/N (%)	n/N (%)		
Incidence of blood transfusion 81 trials (N=9139) NRa	NR	NR		RR 0.66 (0.61, 0.72) P=NR (Phet=NR)
Re-operation due to bleeding NR	NR	NR		RR 0.48 (0.34, 0.67) P=NR (P <i>het</i> =NR)
Myocardial infarction 42 trials (N=5884) 34 trials (N=5441)	153/3329 (4.6)	115/2555	(4.5)	RR 0.94 (0.73, 1.21) P=NR (Phet=NR)
Mortality 49 trials (N=7439) 32 trials (N=6279)	101/4086 (2.5)	81/3353 (2	2.4)	RR 0.93 (0.69, 1.25) P=NR (P <i>het</i> =NR)
Tranexamic acid vs placeb	po/no treatment			
	n/N (%)	n/N (%)		
Incidence of blood transfusion NR	NR	NR		RR 0.70 (0.61, 0.80) P=NR (P <i>het</i> =NR)
Re-operation due to bleeding NR	NR	NR		RR 0.67 (0.41, 1.12) P=NR (P <i>het</i> =NR)
Myocardial infarction 16 trials (N=1732) 10 trials (N=1148)	13/891 (1.5)	16/841 (1.	9)	RR 0.86 (0.43, 1.75) P=NR (Phet=NR)
Mortality 19 trials (N=1802) 11 trials (N=1390)	8/922 (0.9)	16/880 (1.	8)	RR 0.55 (0.24, 1.25) P=NR (Phet=NR)
E-aminocaproic acid vs pla	acebo/no treatment			•
	n/N (%)	n/N (%)		
Incidence of blood transfusion NR	NR	NR		RR 0.75 (0.58, 0.96) P=NR (P <i>het</i> =NR)
Re-operation due to bleeding NR	NR	NR		RR 0.35 (0.11, 1.17) P=NR (P <i>het</i> =NR)
Myocardial infarction 5 trials (N=622)	12/340 (3.5)	14/322 (4.	.3)	RR 0.89 (0.37, 2.18) P=NR (Phet=NR)
Mortality 5 trials (N=672)	7/346 (2.0)	3/326 (0.9)		RR 1.65 (0.50, 5.43) P=NR (Phet=NR)
Outcome	Clinical importance		Clinical relevance	
Aprotinin vs placebo				
Incidence of blood transfusion	1: Clinically important bene confidence limit does not in value		1: Patient-relevant clinical outcome.	
Re-operation due to	1: Clinically important benefit,		1: Patient-relevant clinical outcome.	

bleeding	confidence limit does not include null value		
Myocardial infarction	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.	
Mortality	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.	
Tranexamic acid vs placebo	0		
Incidence of blood transfusion	Clinically important benefit, confidence limit does not include null value	1: Patient-relevant clinical outcome.	
Re-operation due to bleeding	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.	
Myocardial infarction	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.	
Mortality	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.	
ε-aminocaproic acid vs plac	cebo		
Incidence of blood transfusion	Clinically important benefit, confidence limit does not include null value	1: Patient-relevant clinical outcome.	
Re-operation due to bleeding	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	also compatible	
Myocardial infarction	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.	
Mortality	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.	
EVTEDNAL VALIDITY		·	

Generalisability

This systematic review focuses on patients who have undergone cardiac surgery so may only be generalisable to this group.

Applicability

Included studies were performed in a wide range of countries. Eight of the original 211 included studies were conducted in Australia. None of the 11 updated studies were conducted in Australia. Given the wide range of included studies, the results are likely to be applicable to the Australian setting.

Comments

The authors conclude that aprotinin has a higher risk of death than tranexamic acid and ε -aminocaproic acid, with no clear advantages. The risk of death for aprotinin compared with tranexamic acid in head-to-head trials was RR 1.43 (0.98, 2.08) while the risk of death for aprotinin compared with ε -aminocaproic acid in head-to-head trials was RR 1.49 (0.98, 2.28). These results are largely driven by the results of the large BART trial (Fergusson 2008).

Abbreviations: CI, confidence interval; het, heterogeneity: ITT, intention to treat; NR, not reported; RCT, randomised controlled trial; RR, risk ratio; SD, standard deviation.

^a Trials (N) in italics denotes the number of trials (N) included in the analysis. Trials in which the events occurred in 100% in both treatment arms, 0% in both treatment arms or in which a SD was not available were not estimable and were not included in the analysis.

STUDY DETAILS Citation Kagoma YK, Crowther MA, Douketis J et al (2009) Use of antifibrinolytic therapy to reduce transfusion in patients undergoing orthopaedic surgery: a systematic review of randomized trials. Thrombosis Research 123: 687-696. Affiliation/Source of funds Applied Science and Engineering, University of Toronto, Canada; Departments of Medicine and Hematology, McMaster University (St Josephs Hospital and Hamilton General Hospital), Hamilton, Canada; Department of Clinical Epidemiology and Biostatistics and Surgery (Division of Orthopedics), McMaster University, Hamilton, Canada. Funding not stated. Study design Level of evidence Location/setting Systematic review including 29 RCTs Hospital Level I that investigated the effectiveness and safety of aprotinin (6 RCTs), tranexamic acid (19 RCTs) and εaminocaproic acid (1 RCT); also combination studies (3 RCTs) in patients undergoing total hip replacement or total knee arthroplasty. Intervention Comparator Aprotinin: all aprotinin dosages were a variation on the Placebo/no treatment regimens recommended in the product monograph for CABG surgery. Tranexamic acid: Most doses were weight adjusted, ranging from 10-15 mg/kg. E-aminocaproic acid: doses ranged from 12.5 to 100 mg/kg, as well fixed doses of 5-10 g. Population characteristics Adult surgical patients undergoing total hip replacement or total knee arthroplasty. Length of follow-up Outcomes measured 3 days to 3 months Blood loss; transfusion incidence; VTE INTERNAL VALIDITY Follow-up Allocation Results Blinding analysis Treatment/ measurement bias (ITT) Randomised Baseline characteristics of intervention 20 studies double-Publication bias not No included allocation was and control groups reported. blind. assessed. Treatment studies reported in all Analysis of blood loss and VTE carried out or measurement bias reported ITT studies. using standardised mean difference analysis. not apparent. Randomisation 4.1% of (converted to RR for VTE). sequence subjects described in 18 excluded studies. from analysis. Overall quality assessment (descriptive) Good. The search strategy employed as well as study selection and extraction of data were adequate. Formal assessment of quality using Jadad Scale with 21/29 receiving a high rating RESULTS Outcome Intervention group Comparator group Statistical significance Risk estimate (95% CI) No. trials (N) No. trials included in analysis (N)a Aprotinin vs no aprotinin Mean ± SD Mean ± SD

Total blood loss (mL)	NR	NR	WMD -639 (-725, -536)	
6 trials (N=271)			P<0.05 (P <i>het</i> =NR)	
	n/N (%)	n/N (%)		
Transfusion incidence 5 trials (N=401)	NR	NR	RR 0.63 (0.50, 0.80) P<0.05 (P <i>het</i> =NR)	
VTE	NR	NR	RD -0.04 (-0.09, 0.02)	
7 trials (N=481)			P>0.05 (Phet=NR)	
Tranexamic acid vs no tran				
	Mean ± SD	Mean ± SD		
Total blood loss (mL) 20 trials (N=1157)	NR	NR	WMD -393 (-442, -345) P<0.05 (P <i>het</i> =NR)	
	n/N (%)	n/N (%)		
Transfusion incidence 21 trials (N=1237)	NR	NR	RR 0.47 (0.40, 0.55) P<0.05 (Phet=NR)	
VTE 21 trials (N=1237)	NR	NR	RD -0.01 (-0.04, 0.02) P>0.05 (Phet=NR)	
E-aminocaproic acid vs ε-a	minocaproic acid	L	· · · · · · · · · · · · · · · · · · ·	
•	Mean ± SD	Mean ± SD		
Total blood loss (mL)	NR	NR	WMD -331 (-544, -118)	
2 trials (N=150)			P<0.05 (P <i>het</i> =NR)	
	n/N (%)	n/N (%)		
Transfusion incidence 3 trials (N=180)	NR	NR	RR 0.64 (0.21, 1.93) P≥0.05 (Phet=NR)	
VTE 3 trials (N=180)	NR	NR	RD 0.00 (-0.07, 0.07) P>0.05 (Phet=NR)	
Outcome	Clinical importance	•	Clinical relevance	
Aprotinin vs no aprotinin	•			
Blood loss	1: Clinically important <i>benefit</i> , confidence limit does not include null value (p<0.05).		1: Patient-relevant clinical outcome.	
Transfusion incidence	1: Clinically important ben- confidence limit does not in value (p<0.05).		1: Patient-relevant clinical outcome.	
VTE	4: Range of estimates inclimportant effects, but also with no effect		1: Patient-relevant clinical outcome.	
Tranexamic acid vs no tran				
Blood loss	1: Clinically important <i>bene</i> confidence limit does not invalue (p<0.05).		1: Patient-relevant clinical outcome.	
Transfusion incidence	1: Clinically important <i>bend</i> confidence limit does not invalue (p<0.05).		1: Patient-relevant clinical outcome.	
VTE	4: Range of estimates includes clinically important effects, but also compatible with no effect		1: Patient-relevant clinical outcome.	
E-aminocaproic acid vs ε-a	minocaproic acid	a.		
Blood loss	1: Clinically important <i>benefit</i> , confidence limit does not include null value (p<0.05).		1: Patient-relevant clinical outcome.	
Transfusion incidence	4: Range of estimates incli important effects, but also with no effect		1: Patient-relevant clinical outcome.	
VTE	4: Range of estimates inclimportant effects, but also with no effect		1: Patient-relevant clinical outcome.	

Generalisability

This systematic review focuses on adult patients who have undergone total hip replacement or total knee arthroplasty so may only be generalisable to these specific surgical populations. An analysis including all studies which separated out the two surgery types found the blood loss results were similar for each (WMD -1.12; 95% CI: -1.31, -0.93for THR and WMD - 0.89; 95% CI: -1.05, -0.72 for TKA).

Applicability

This review does not report the locations of the included studies and as such the applicability of the results to the Australian/New Zealand setting is unclear.

Comments

Abbreviations: CABG, coronary artery bypass graft; CI, confidence interval; *het*, heterogeneity: ITT, intention to treat; NR, not reported; RCT, randomised controlled trial; RD, risk difference; RR, relative risk; SD, standard deviation; THR, total hip replacement; TKR, total knee replacement; VTE, venous thromboembolism; WMD, weighted mean difference.

^a Where individual studies had either 100% events in both treatment arms, no events in both treatment arms or no SD was reported, a risk estimate for that individual study could not be calculated, and it could not be included in the pooled analysis. Where this has occurred, the actual number of studies and subjects included in the analysis will be stated.

Citation

Systematic review: Kongnyuy EJ, Wiysonge CS (2009) Interventions to reduce haemorrhage during myomectomy for fibroids. Cochrane Database of Systematic Reviews 2009, Issue 3. Art. No.: CD005335. DOI: 10.1002/14651858.CD005335.pub3.

Single included RCT: Caglar GS, Tasci Y, Kayikcioglu F et al (2008) Intravenous tranexamic acid use in myomectomy: a prospective randomised double-blind placebo controlled study. European Journal of Obstetrics, Gynecology and Reproductive Biology 137(): 227-231.

Affiliation/Source of funds

Systematic review: Child and Reproductive Health Group, Liverpool School of Tropical Medicine, Liverpool, UK; Institute of Infectious Disease and Molecular Medicine, University of Cape Town, Cape Town, South Africa.

Funding: Saltonstall Fund for Pain Research, USA.

Single included RCT: Ankara Etlick Maternity and Women's Health Teaching Research Hospital, Ankara, Turkey.

Funding: not stated.

Study design	Level of evidence	9	Location/setting
Systematic review including 10 RCTs	Level II		Hospital
that assessed the effectiveness and			
safety of intervention to reduce			
haemorrhage during myomectomy for			
fibroids. Only 1 of the 10 included			
RCTs related to TXA. ^a			
Intervention		Comparator	
TXA: bolus injection of 10 mg/kg (max	1 g) 15 minutes	Placebo: saline (same	e regimen as TXA)

before incision followed by a continuous infusion of mg/kg/hr dissolved in 1 L saline for 10 h (max 1 g/10 hr).

Population characteristics

Women scheduled for myomectomy due to myoma uteri. Mean age 35 years, volume of myomas 457 cm³ ± SD 669 in the intervention group and 286 cm³ ± SD 259 in the control group. Authors state no difference in any baseline characteristics or number and volume of myomas between groups.

The state of the s					
Length of follow-up	Outcomes measured				
In hospital	Blood loss; duration of surgery, haemoglobin, haematocrit,				
	blood transfusion requirements on ward.				

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Patients randomised according to a computer- generate sequence. Allocation by sequentially numbered drug containers of identical appearance.	Authors state no difference in any baseline characteristics or number and volume of myomas between groups. Analysis not described.	Double-blind. Patients, surgeons and anaesthetists unaware of treatment assignment.	No suggestions of selective reporting. Unclear whether there are other sources of bias.	All patients included in analysis.

Overall quality assessment (descriptive)

RCT: Good.

Systematic review: Good. Extensive literature search, data assessed for quality.

RESULTS

Outcome No. trials (N) No. trials included in analysis (N) ^a	Intervention group	Comparator group		Statistical significance Risk estimate (95% CI)
Tranexamic acid vs placebo				
	N=50	N=50		Risk estimate (95% CI) P value
	n/N (%)	n/N (%)		
Transfusion incidence	15/50 (30)	10/50 (20)		OR 1.71 (0.68, 4.30) P=0.25
	Mean ± SD	Mean ± SD		
Blood loss (mL)	804 ± 482	1047 ± 617		MD -243.00 (-460.02, -25.98) P=0.028
Postoperative haemoglobin (g/dL)	9.97 ± 1.5	9.76 ± 1.4		MD 0.21 (-0.36, 0.78) P=0.47
			T	
Outcome	Clinical importance		Clinical rel	evance
Tranexamic acid vs placebo			1450	
Transfusion incidence	4: Range of estimates includes clinically important effects, but also compatible with no effect		1: Patient-re	elevant clinical outcome.
Blood loss	Clinically important benefit, confidence limit does not include null value (p=0.028).		1: Patient-re	elevant clinical outcome.
Postoperative haemoglobin	4: Range of estimates inc important effects, but also with no effect	ludes clinically compatible	1: Patient-relevant clinical outcome.	
EXTERNAL VALIDITY	•		1	
Generalisability				
This systematic review focuse	es on adult female patients u	ndergoing myon	nectomy for ut	erine fibroids and only includes one
relevant RCT for this interven				
Applicability				
	cted at a single centre in Tu	rkey so may not	be directly ap	plicable to the Australian/New
Zealand setting.				
Comments				

Abbreviations: CI, confidence interval; het, heterogeneity; ITT, intention to treat; MD, mean difference; RCT, randomised controlled trial; SD, standard deviation; TXA, tranexamic acid.

^a Where individual studies had either 100% events in both treatment arms, no events in both treatment arms or no SD was reported, a risk estimate for that individual study could not be calculated, and it could not be included in the pooled analysis. Where this has occurred, the actual number of studies and subjects included in the analysis will be stated.

^a All information relating to the single included RCT was taken from information provided in the Kongnyuy 2009 systematic review.

STUDY DETAILS								
Citation								
							ive outcomes in orthotopic	c liver
transplantation: a meta-analysis on aprotinin. World J Gatroenterol 14(9): 1425-1429.								
Affiliation/Source of funds								
							Medical University, Nanji	ng, China.
Funding: Supported	d by Grant (Committee			
Study design Level of evidence Location/setting								
Systematic review i			el I			H	ospital	
and 1 non-RCT tha	t assessed	the						
effectiveness and s	afety of apr	otinin						
in liver transplantati								
included tranexami								
and has been exclu		ne						
analyses presented	l here.							
Intervention					mparator			
Aprotinin: various d		ot stated.		Pla	icebo/no tre	eatment		
Population charac								
Adult patients unde		otopic liver trar	nsplantation; no	o fur	ther details	of patients	provided.	
Length of follow-u	ıp				tcomes m			
Not stated							s the only outcome includ	ed here as it
				inc	luded data	from RCTs	s only)	
INTERNAL VALIDI	TY							
Allocation	Results				Blinding	analysis	Treatment/	Follow-up
Daniel and a diameter	Nie detelle				Diania		measurement bias	(ITT)
Randomisation in		on compariso	n or baseline	3			Unclear	Unclear
RCTs not	characteri		an Davillan	studies not		Ol		
described. Analyses conducted using RevMan.		reported.						
1 included study								
non-randomised								
Overall quality ass			No. a. !				al assubusilis al buist. This store	d la a al a a
							d controlled trial. This stu	
							greater mortality and less	
							c acid. For this reason, th	is study was
included only for the	e unrombosi	is outcome wn	ich was based	OHC	iata irom z	piacebo-co	ontrolled RCTS.	
RESULTS		1					Challania da da directione	
Outcome		Intervention	group	C	mparator	group	Statistical significar	
No. trials (N)	l :						Risk estimate (95%	CI)
No. trials included	ı in							
analysis (N)a	- 2							
Aprotinin vs placebo	O ^a	Maan . CD		N 4.	CD			
		Mean ± SD			ean ± SD		Dials actimate (000)	CI)
Thursday and allege	n/N (%)						Risk estimate (95% CI)	
Thromboembolic ev	3/122 (2.5)		5/	78 (6.4)		OR 0.38 (0.09, 1.64)		
2 RCTs (N=200) P>0.05 (Phet=0.88)								
Outcome Clinical importance Clinical relevance								
Aprotinin vs placebo		1.5				4 5 11		
I hromboembolic ev	Thromboembolic events 4: Range of estimates includes clinically 1: Patient-relevant clinical outcome.							
			ects, but also c	omp	atible			
EVTERMAL ****	171/	with no effect	Į					
EXTERNAL VALIDITY								
Generalisability								
This systematic review focuses on adult patients undergoing orthotopic liver transplantation. It is likely to only be								
generalisable to this select patient group.								

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Applicability

There is no information provided at to the locations of the included studies so it is not possible to determine whether the results would be directly applicable to the Australian/New Zealand setting.

Comments

Abbreviations: CI, confidence interval; FFP, fresh frozen plasma; het, heterogeneity; ITT, intention to treat; NR, not reported; RBC, red blood cell; RCT, randomised controlled trial; SD, standard deviation; WMD, weighted mean difference.

a Where individual studies had either 100% events in both treatment arms, no events in both treatment arms or no SD was reported, a risk estimate for that individual study could not be calculated, and it could not be included in the pooled analysis. Where this has occurred, the actual number of studies and subjects included in the analysis will be stated.

^a Only the analysis excluding the RCT comparing aprotinin with tranexamic acid is presented here.

Citation

DR McIlroy, PS Myles, LE Phillips, JA Smith

Antifibrinolytics in cardiac surgical patients receiving aspirin: a systematic review and meta-analysis

Affiliation/Source of funds

Department of Anaesthesia and Perioperative Medicine, Alfred Hospital, Melbourne, Australia; Department of Epidemiology and Preventive Medicine, Monash University, Melbourne, Australia; Department of Surgery, Monash Medical Centre, Melbourne, Australia.

Funding: Alfred Hospital anaesthesia research fund.

Study design	Level of evidence	Location/setting
Systematic review including 17 RCTs	Level I	Hospital
that investigated the effects of		
aprotinin, tranexamic acid and ε-		
aminocaproic acid on blood loss and		
use of blood products in cardiac		
surgery patients receiving aspirin.		

Intervention	Comparator
Aprotinin, tranexamic acid or ε-aminocaproic acid	Placebo, no treatment or active

Population characteristics

Adult patients undergoing CABG \pm valve surgery where aspirin had been maintained or initiated through the prospective period.

Length of follow-up	Outcomes measured	
Not specified	Blood loss (chest tube drainage); incidence of transfusion; re-	
	operation; thrombotic complication;	

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomised allocation was reported in all 17 studies except one which was pseudorandomised. Method adequate for 5 trials, inadequate in 1 trial and unclear in the remaining trials.	Baseline characteristics of intervention and control groups not reported. Methodological quality of studies assessed independently by two authors. Disagreements resolved by consensus. Random effects model used for all analyses. Heterogeneity assessed.	12 trials double- blind, 4 open- label.	No evidence of publication bias found.	No studies judged to be at high risk of attrition bias, 4 judged to be at moderate risk, 7 judged to be at low risk. Remaining studies unable to judge.

Overall quality assessment (descriptive)

Good. Comprehensive literature search carried out. Quality assessment undertaken. Appropriate analysis methods used. Subgroup and sensitivity analyses undertaken.

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance		
Aprotinin vs placebo					
Blood loss (postoperative chest tube loss) mL 12 trials (N=992)	NR	NR	WMD -432.51 (-543.68, -321.35) P<0.001 (Phet<0.001)		
Incidence of transfusion 10 trials (N=856)	205/510 (40.2)	229/346 (66.2)	OR 0.34 (0.25, 0.46) P<0.001 (Phet=0.75)		

Re-operation 7 trials (N=352)	5/186 (2.7)	10/166 (6.0)		OR 0.42 (0.13, 1.36) P=0.15 (Phet=0.61)	
4 trials (N=198) ^a				P=0.15 (PHet=0.01)	
Thrombotic complication 8 trials (N=527)	10/269 (3.7)	17/258 (6.6)		OR 0.51 (0.21, 1.20) P=0.12 (Phet=0.76)	
3 trials (N=174)	acid or ε-aminocaproic acid) vs	nlaceho			
Blood loss (postoperative	NR	NR		WMD -189.35 (-287.24, -91.46)	
chest tube loss) mL 3 trials (N=259)	TVIC	INR		P<0.001 (Phet=0.05)	
Incidence of transfusion 1 trial (N=79)	8/40 (20.0)	8/39 (20.5)		OR 0.97 (0.32, 2.90) P=0.95 (Phet=NA)	
Re-operation 2 trials (N=109)	0/55 (0.0)	2/54 (3.7)		OR 0.31 (0.03, 3.14) P=0.32 (Phet=0.99)	
Thrombotic complication 3 trials (N=259) 1 trial (N=79)	0/155 (0.0)	1/104 (1.0)		OR 0.32 (0.01, 8.02) P=0.49 (Phet=NA)	
Outcome	Clinical importance		Clinical rel	evance	
Aprotinin vs placebo					
Blood loss	confidence limit does not inclivalue (P<0.001)	Clinically important benefit, confidence limit does not include null value (P<0.001)		1: Patient-relevant clinical outcome.	
Incidence of transfusion	Clinically important benefit, confidence limit does not include null value (P<0.001)		1: Patient-relevant clinical outcome.		
Re-operation	4: Range of estimates includes clinically important point estimate but also compatible with no effect, or a harmful effect		1: Patient-relevant clinical outcome.		
Thrombotic complication	4: Range of estimates includes clinically important point estimate but also compatible with no effect, or a harmful effect		1: Patient-relevant clinical outcome.		
Lysine analogue vs placebo					
Blood loss	1: Clinically important benefit, confidence limit does not include null value (P<0.001)		1: Patient-relevant clinical outcome.		
Incidence of transfusion	4: Range of estimates includes clinically important point estimate but also compatible with no effect, or a harmful effect		1: Patient-relevant clinical outcome.		
Re-operation	4: Range of estimates includes clinically important point estimate but also compatible with no effect, or a harmful effect		1: Patient-relevant clinical outcome.		
Thrombotic complication	4: Range of estimates includes clinically important point estimate but also compatible with no effect, or a harmful effect		1: Patient-relevant clinical outcome.		
EXTERNAL VALIDITY	•		•		
Generalisability					
review are likely to be genera	es on cardiac surgical patients w alisable only to this specific popu		eceiving aspir	rin. Therefore, the results of this	
Applicability The locations of the included setting.	studies are not reported so it is	not possible t	o comment o	n the applicability to the Australian	

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Comments

The authors concluded that antifibrinolytics were effective at reducing blood loss and transfusion requirements in cardiac surgery patients using aspirin.

Abbreviations: CABG, coronary artery bypass graft; *het*, heterogeneity; ITT, intention to treat; NR, not reported; OR, odds ratio; RCT, randomised controlled trial; SD, standard deviation; WMD, weighted mean difference.

^a Trials (N) in italics denotes the number of trials (N) included in the analysis. Trials in which the events occurred in 100% in both treatment arms, 0% in both

treatment arms or in which a SD was not available were not estimable and were not included in the analysis.

STUDY DETAILS Citation ES Schouten, AC van de Pol, ANJ Schouten et al. The effect of aprotinin, tranexamic acid and aminocaproic acid on blood loss and use of blood products in major pediatric surgery: a meta-analysis. Affiliation/Source of funds Division of Pediatric Intensive care and Division of Perioperative Care4 and Emergency Medicine, Wilhelmina Children's Hospital, University Medical Canter, Utrecht, The Netherlands Study design Level of evidence Location/setting Systematic review including 28 RCTs Level I Hospital that investigated the effects of aprotinin, tranexamic acid and εaminocaproic acid on blood loss and use of blood products Intervention Comparator Aprotinin, tranexamic acid or ε-aminocaproic acid Placebo or no treatment Population characteristics Children undergoing cardiac or scoliosis surgery Length of follow-up Outcomes measured Not specified Blood loss; transfusion (PRC, plasma or thrombo) INTERNAL VALIDITY Allocation Results Blinding analysis Treatment/ Follow-up measurement bias (ITT) Randomised Baseline characteristics of intervention 11/28 studies Publication bias 16/28 and control groups not reported. double-blind. studies had allocation was assessed. Methodological quality of studies 13/28 studies No evidence of reported in all good (> assessed independently by two authors. 80%) follow-28 studies. single-blind, and treatment/measureme Method of Disagreements resolved by discussion. remaining studies up, 5/28 nt bias. Model used not stated. Studies considered not blinded. studies had randomisation too heterogeneous for pooling if I2 statistic not described. moderate Only 3/28 ≥ 50%. follow-up studies scored (50-80%)2/2 for and the allocation, the remaining studies had remaining poor followstudies scored up. Overall quality assessment (descriptive) Good. Comprehensive literature search carried out. Quality assessment undertaken. Meta-regression analysis carried out to identify potential confounders for the cardiac studies due to heterogeneity (age, weight and time on cardiopulmonary bypass). **RESULTS** Intervention group Comparator group Statistical significance Outcome Aprotinin vs placebo Blood loss Mean ± SD Mean ± SD 14 trials (N=594) NR NR Cardiac surgery NR due to heterogeneity 1 trial (N=44) NR NR Scoliosis surgery WMD -385 mL (-727, -42) P=NR (I2=NA) Transfusion (packed red cells) Mean ± SD Mean ± SD

0.1.1. (\$1.050)	LND	LND	
3 trials (N=250)	NR	NR	Cardiac surgery WMD -4 mL/kg (-7, -2)
			P=NR (I ² =0%)
Transfusion (plasma)			1 -1414 (1 -070)
Translation (piasina)	Mean ± SD	Mean ± SD	
2 trials (N=228)	NR	NR	Cardiac surgery
			WMD -5 mL/kg (-8, -2)
			P=NR (I ² =0%)
Transfusion (thrombo)			•
	Mean ± SD	Mean ± SD	
NR (N=180)	NR	NR	Cardiac surgery
			NR due to heterogeneity
Tranexamic acid vs plac	cebo		
Blood loss			
	Mean ± SD	Mean ± SD	
6 trials (N=542)	NR	NR	Cardiac surgery
			WMD -11 mL/kg (-13, -8)
2 trials (NL 04)	ND	ND	P=NR (I ² =31%)
2 trials (N=84)	NR	NR	Scoliosis surgery
			WMD -682 mL (-1149, -214) P=NR (I ² =24%)
Transfusion (packed red	d colle)		P=NR (12=24%)
rransiusium (packeu let	Mean ± SD	Mean ± SD	
5 trials (N=460)	NR	NR	Cardiac surgery
5 thats (N=400)	IVIX	IVIX	WMD -7 mL/kg (-10, -5)
			P=NR (I ² =6%)
2 trials (N=84)	NR	NR	Scoliosis surgery
2 (14 0 1)			WMD -349 mL (-620, -77)
			P=NR (I ² =0%)
Transfusion (plasma)	'	II.	
,	Mean ± SD	Mean ± SD	
4 trials (N=419)	NR	NR	Cardiac surgery
			WMD -7 mL/kg (-9, -4)
			P=NR (I ² =0%)
2 trials (N=84)	NR	NR	Scoliosis surgery
			WMD -15 mL (-127, 98)
			P=NR (I ² =24%)
Transfusion (thrombo)			
ND (N. 270)	Mean ± SD	Mean ± SD	Conding support
NR (N=370)	NR	NR	Cardiac surgery
			WMD -5 mL/kg (-7, -3)
E-aminocaproic acid vs	nlacoho		P=NR (I ² =0%)
Blood loss	piaceno		
טטטע וטטט	Mean ± SD	Mean ± SD	
3 trials (N=410)	NR	NR	Cardiac surgery
3 111a13 (IN=41U)	INIX	INIX	Cardiac surgery NR due to heterogeneity
1 trial (N=36)	NR	NR	Scoliosis surgery
i iiiai (N-30)	INIX	INIX	WMD -59 mL (-262, 144)
			P=NR (I ² =NA)
Transfusion (packed red	d cells)	1	1
	Mean ± SD	Mean ± SD	
3 trials (N=410)	NR	NR	Cardiac surgery
		1	NR due to heterogeneity
Transfusion (plasma)	<u> </u>	<u> </u>	1
	Mean ± SD	Mean ± SD	
		1	

3 trials (N=410)	NR	NR		Cardiac surgery WMD -3 mL/kg (-5, -1) P=NR (I ² =20%)
Transfusion (thrombo)				1 111 (1 2370)
()	Mean ± SD	Mean ± SI	D	
3 trials (N=410)	NR	NR		Cardiac surgery NR due to heterogeneity
Outcome	Clinical importance		Clinical rel	evance
Aprotinin vs placebo				
Blood loss	Scoliosis surgery 1: Clinically important benefit, confidence limit does not incluvalue.	de null		elevant clinical outcome.
Transfusion (packed red cell)	Cardiac surgery 1: Clinically important benefit, confidence limit does not incluvalue.	de null	1: Patient-re	elevant clinical outcome.
Transfusion (plasma)	Cardiac surgery 1: Clinically important benefit, confidence limit does not incluvalue.	de null	1: Patient-r	elevant clinical outcome.
Tranexamic acid vs placebo				
Blood loss	Cardiac surgery 1: Clinically important benefit, confidence limit does not incluvalue. Scoliosis surgery 1: Clinically important benefit, confidence limit does not incluvalue.			elevant clinical outcome.
Transfusion (packed red cell)	Cardiac surgery 1: Clinically important benefit, confidence limit does not incluvalue. Scoliosis surgery 1: Clinically important benefit, confidence limit does not incluvalue.		1: Patient-ro	elevant clinical outcome.
Transfusion (plasma)	Cardiac surgery 1: Clinically important benefit, confidence limit does not incluvalue. Scoliosis surgery 4: Range of estimates includes important point estimate but al compatible with no effect, or a effect	s clinically so	1: Patient-ro	elevant clinical outcome.
Transfusion (thrombo) Tranexamic acid vs placebo	Cardiac surgery 1: Clinically important benefit, confidence limit does not incluvalue.	de null		
Tranonamie dela 73 piacebo				

Blood loss	Scoliosis surgery 4: Range of estimates includes clinically important point estimate but also compatible with no effect, or a harmful effect	1: Patient-relevant clinical outcome.					
Transfusion (plasma)	Cardiac surgery 1: Clinically important benefit, confidence limit does not include null value.	1: Patient-relevant clinical outcome.					
EXTERNAL VALIDITY							
Generalisability							
	This systematic review focuses on paediatric patients (<18 years) undergoing cardiac or scoliosis surgery. Therefore, the						
results of this review are likely to be generalisable only to this specific population.							
Applicability							

Applicability

The locations of the included studies are not reported so it is not possible to comment on the applicability to the Australian setting.

Comments

The authors concluded that there was no evidence than tranexamic acid and ϵ -aminocaproic acid are less effective than aprotinin in major paediatric surgery.

Abbreviations: ITT, intention to treat; NA, not applicable; NR, not reported; PRC, packed red cell; RCT, randomised controlled trial; SD, standard deviation; WMD, weighted mean difference.

STUDY DETAILS Citation Tzortzopoulou A, Cepeda MS, Schumann R et al (2008) Antifibrinolytic agents for reducing blood loss in scoliosis surgery in children. Cochrane Database of Systematic Reviews 2008, Issue 3. Art. No.: CD006883. DOI: 10.1002/14651858.pub2. Affiliation/Source of funds Department of Anesthesia, Tufts medical Center, Boston, US; Pharmacoepidemiology, Johnson & Johnson Pharmaceutical Research and development, Titussville, US. Funding: Saltonstall Fund for pain Research, US. Study design Level of evidence Location/setting Systematic review including 6 RCTs Level I Hospital that investigated the effects of aprotinin (2 RCTs), tranexamic acid (2 RCTs) and ε-aminocaproic acid (2 RCTs) on blood loss and transfusion in children undergoing surgery for primary or secondary scoliosis. Intervention Comparator Aprotinin: High-dose regimen Placebo Tranexamic acid: 1 high-dose regimen and 1 low-dose regimen E-aminocaproic acid: high-dose regimen Population characteristics Children (< 18 years) undergoing primary or secondary scoliosis surgery. Mean age 14.1 years (idiopathic scoliosis) and 13.2 years (idiopathic and secondary scoliosis); 72.5% female (idiopathic scoliosis) and 44.1% female (idiopathic and secondary scoliosis) Length of follow-up Outcomes measured In-hospital Blood loss, transfusion incidence, transfusion volume, mortality, morbidity. INTERNAL VALIDITY Blinding analysis Follow-up Allocation Results Treatment/ measurement bias (ITT) Randomised Methodological quality of studies 5/6 studies No evidence of All studies allocation was assessed independently by two authors. double-blind. 1 treatment/measureme described reported in all 6 Disagreements resolved by discussion. study not nt bias. as having in studies; 4/6 studies rated A (low risk o bias) and described. ITT described in 5 2/6 rated B (moderate risk of bias). analysis. studies (1 computergenerated, 2 random number tables, 2 drawing numbers from container). Overall quality assessment (descriptive) Good. Comprehensive literature search carried out. Quality assessment undertaken. Studies rated as having low-moderate risk of bias. **RESULTS** Statistical significance Outcome Intervention group Comparator group Risk estimate (95% CI) P value (heterogeneity) Aprotinin vs placebo Transfusion incidence n/N (%) n/N (%)

1 RCT (N=43)	8/15 (53.3)	20/28 (71.4)	RR 0.75 (0.44, 1.27) P=0.28 (Phet=NA)
Transfusion volume (mL)			
	Mean ± SD	Mean ± SD	
2 RCTs (N=87)	NR	NR	WMD -361.42 (-583.88, -138.96) P=0.0015 (Phet=0.80)
Blood loss (mL)			•
	Mean ± SD	Mean ± SD	
2 RCTs (N=87)	NR	NR	WMD -450.32 (-726.35, -174.29) P=0.0014 (Phet=0.53)
Mortality			
2.5.25 (1) 25	n/N (%)	n/N (%)	
2 RCTs (N=87)	0/15 (0)	0/28 (0)	NA
Renal insufficiency	.	-	
	n/N (%)	n/N (%)	
2 RCTs (N=87)	0/15 (0)	0/28 (0)	NA
Tranexamic acid vs place	bo		
Transfusion incidence	n/N (%)	n/N (%)	
2 RCTs (N=84)	20/45 (44.4)	21/39 (53.8)	RR 0.84 (0.56, 1.27)
	20/45 (44.4)	21/39 (33.6)	P=0.41 (Phet=0.94)
Transfusion volume (mL)	111 00		
0.007 (N. 0.4)	Mean ± SD	Mean ± SD	WIAD 005 11 ((07.55, 100.70)
2 RCTs (N=84)	NR	NR	WMD -395.14 (-687.55, -102.73) P=0.0081 (Phet=0.51)
Blood loss (mL)	1		
	Mean ± SD	Mean ± SD	
2 RCTs (N=84)	NR	NR	WMD -681.81 (-1149.12, -214.49) P=0.0042 (Phet=0.25)
Mortality			
	n/N (%)	n/N (%)	
2 RCTs (N=84)	0/45 (0)	0/39 (0)	NA
Renal insufficiency	L	1	'
j	n/N (%)	n/N (%)	
2 RCTs (N=84)	0/45 (0)	0/39 (0)	NA
E-aminocaproic acid vs p	lacebo		
Transfusion incidence			
	n/N (%)	n/N (%)	
1 RCT (N=36)	14/19 (73.7)	12/17 (70.6)	RR 1.04 (0.69, 1.57) P=0.84 (Phet=NA)
Transfusion volume (mL)	L	I .	
	Mean ± SD	Mean ± SD	
1 RCT (N=84)	NR	NR	WMD -245.00 (-481.03, -8.97) P=0.042 (Phet=NA)
Blood loss (mL)		I	, ,
. ,	Mean ± SD	Mean ± SD	
1 RCT (N=36)	NR	NR	WMD -325.00 (-586.83, -63.17) P=0.015 (Phet=NA)
Mortality	1	1	

2 RCTs (N=83)	0/46 (0)	0/37 (0)		NA	
Renal insufficiency					
	n/N (%)	n/N (%)			
1 RCTs (N=36)	0/19 (0)	0/17 (0)		NA	
DVT					
	n/N (%)	n/N (%)			
1 RCTs (N=47)	0/27 (0)	3/20 (15)		Not estimable P=0.07	
Outcome	Clinical importance		Clinica	al relevance	
Aprotinin vs placebo			•		
Transfusion incidence	4: Range of estimates ir important point estimate compatible with no effect	e but also	1: Pati	ent-relevant clinical outcome.	
Transfusion volume	1: Clinically important be confidence limit does no value (p=0.0015).	ot include null		ent-relevant clinical outcome.	
Blood loss	1: Clinically important be confidence limit does no value (p=0.0014).	ot include null		ent-relevant clinical outcome.	
Mortality	There were no deaths in arm.	n either treatment	1: Patient-relevant clinical outcome.		
Renal insufficiency	There were no cases of insufficiency in either tre		1: Pati	ent-relevant clinical outcome.	
Tranexamic acid vs placebo					
Transfusion incidence	4: Range of estimates ir important point estimate compatible with no effect	e but also	1: Pati	ent-relevant clinical outcome.	
Transfusion volume	1: Clinically important be confidence limit does no value (p=0.0081).		1: Patient-relevant clinical outcome.		
Blood loss	1: Clinically important be confidence limit does no value (p=0.0042).		1: Patient-relevant clinical outcome.		
Mortality	There were no deaths in arm.	n either treatment	1: Patient-relevant clinical outcome.		
Renal insufficiency	There were no cases of insufficiency in either tre		1: Patient-relevant clinical outcome.		
E-aminocaproic acid vs plac			,		
Transfusion incidence	4: Range of estimates ir important point estimate compatible with no effect	e but also ct, or a harmful	1: Patient-relevant clinical outcome.		
Transfusion volume	1: Clinically important be confidence limit does no value (p=0.042).	ot include null	1: Patient-relevant clinical outcome.		
Blood loss	1: Clinically important be confidence limit does no value (p=0.015).	ot include null	1: Patient-relevant clinical outcome.		
Mortality	There were no deaths in arm.				
Renal insufficiency	There were no cases of insufficiency in either tre		1: Pati	ent-relevant clinical outcome.	

DVT	4: Range of estimates includes clinically important point estimate but also compatible with no effect, or a harmful effect (p=0.07)	1: Patient-relevant clinical outcome.						
EXTERNAL VALIDITY								
Generalisability								
	This systematic review focuses on paediatric patients (<18 years) undergoing surgery for primary or secondary scoliosis. Therefore, the results of this review are likely to be generalisable only to this specific population.							
Applicability								
The locations of the included studies are not reported so it is not possible to comment on the applicability to the Australian/New Zealand setting.								
Comments	Comments							

Abbreviations: CI, confidence interval; DVT, deep vein thrombosis; *het*, heterogeneity; ITT, intention to treat; NA, not applicable; NR, not reported; RCT, randomised controlled trial; RR, risk ratio; SD, standard deviation; WMD, weighted mean difference.

Level II evidence

Level II evidel	100							
STUDY DETAILS								
Citation								
Alvarez JC, Santive	ri FX, Ramo	s I et al (200	08) Tranexamic a	acid redu	ces blood tran	sfusion in to	tal kn	ee arthroplasty even
when a blood conse								. ,
Affiliation/Source	of funds							
Department of Anes	sthesiology a	and the Depa	artment of Ortho	pedic Su	rgery, Univers	ity Hospital c	of Mar	, Barcelona, Spain
No details of funding	g reported.							·
Study design		L	evel of evidence	е		Location/s	ettino	9
Double-blind RCT		- 1				Hospital (si	ngle-d	centre)
Intervention				Compa				
Tranexamic acid: a				Placeb	o: regimen as	per tranexan	nic ac	id
research anaestheti								
tourniquet followed								
	the end of the operation and continuing during the first 6							
	postoperative hours							
Population charac								
ASA-I to IIIa patients			rthritis and unde	rgoing ur	nilateral bicond	lylar cementa	al tota	ıl knee arthroplasty.
Mean age 72; fema		31.						
Length of follow-u					nes measure			
Up to 3 months pos	toperative (f	or thrombos	IS)		, outcome: trai			
					lary outcome:		e bloc	od Ioss
INTERNAL VALIDI	TV			Salety	outcome: thror	ndosis		
INTERNAL VALIDI Allocation	Results		Dlinding analy	roio.	Trootmontin	naacuraman		Follow up /ITT\
Allocation	Results		Blinding analy	ysis	Treatment/n bias	neasuremer	IL	Follow-up (ITT)
Computer-	Results de	termined	Double-blind. N	leither Staff blinded to treatment			t	15 subjects excluded
generated	by anaesth	etist not	patient not the	allocation so not likely to b		be	from analysis following	
random	aware of tr	eatment	anaesthetist	treatment or measurement		nt	randomisation (9 in	
numbers/sealed	assignmen	t	assessing resu	ılts	bias.			treatment group and 6
envelopes.			aware of treatr	nent				in control group) due
Treatment			assignment.					to release of
prepared by an								tourniquet (7), no
anaesthetist not								epidural catheter (7)
otherwise								and error during blood
engaged in the								sampling (1)
study.								
Overall quality ass				/O. I		171		\
						and 6 in cont	rol gr	oup). These subjects
were not further des	scribea. The	autnors no	te this is a ilmitat	ion of the	e study.			
RESULTS		Intonionti	on group	Con	anaratar arau	n	Ctot	ictical cignificance
Outcome		Interventi N=46	on group	N=4	nparator grou	þ	Stat	istical significance
Tranexamic acid		11-40		111-4	7			
Transamic aciu		n/N (%)		n/N	(%)		P va	nlue
Transfusion inciden	CP	1/46 (2.2)			(12.2)			(post-hoc)
(allogeneic and aut		1170 (2.2)		0/47	(14.4)		0.00	(2006 1100)
blood)								
Transfusion inciden	ce	2/46 (4.3)		36/4	9 (73.5)		<0.0	0001
(recovered blood)								
		Mean ± S	D		n ± SD		P va	ılue
Total RBC transfusi		1		1.8			NR	
transfused patients	only	(1 unit in 1	patient)	(11	units in 6 patie	nts)		
(units)								

Allogeneic RBC transfusion in	11	NR		NR		
transfused patients only	(1 unit in 1 patient)	(8 unit; numbe	r of patients			
(units)		NR)	,			
Autologous RBC transfusion	0	NR		NR		
in transfused patients only		(3 unit; numbe	r of patients			
(units)		NR)				
Chest tube blood loss at 0-6	159 ± 110	534 ± 351		<0.0001		
hr postoperative (mL)						
Chest tube blood loss at 6 hr	132 ± 151	132 ± 150		0.98		
 4 day postoperative (mL) 						
Total chest tube blood loss	170 ± 109	551 ± 352		<0.001		
(mL)						
	n/N (%)	n/N (%)		P value		
Thrombosis	0/46 (0)	0/49 (0)		NA		
1. Haemoglobin						
	Mean ± SD	Mean ± SD		P value		
Haemoglobin (preoperative)	13.5	13.6		NS		
Haemoglobin (end of	11.9	11.9		NS		
surgery)						
Haemoglobin (6 hr	11.5	10.9		P<0.05		
postoperative)						
Haemoglobin (4 day	10.4	9.9		P<0.05		
postoperative)						
Clinical importance	•		Clinical relev	ance		
Tranexamic acid						
Transfusion incidence	Allogeneic and autologous bl	ood	1: Patient-rele	vant clinical outcome.		
	4: Range of estimates include	es clinically				
	important effects, but also co	mpatible with				
	no or harmful effect					
	Recovered blood					
	1: Clinically important benefit					
	limit does not include null vali	ue (p<0.001)				
Transfusion volume	Total RBC transfusion in tran		1: Patient-relevant clinical outcome.			
	Unclear. No formal statistical	comparison				
	made.					
	Allogeneic RBC transfusion					
	Unclear. No formal statistical	comparison				
	made.					
	Autologous RBC transfusion					
	Unclear. No formal statistical	comparison				
	made.					
Blood loss	Chest-tube blood loss (0-6 hr	-)	1: Patient-rele	vant clinical outcome.		
	1: Clinically important <i>benefit</i> , confidence					
	1: Clinically important benefit					
	1: Clinically important benefit limit does not include null value.	ue (p<0.001)				
	1: Clinically important benefit limit does not include null value. Chest-tube blood loss (6 hr –	ue (p<0.001) · <i>4 day)</i>				
	1: Clinically important benefit limit does not include null value. Chest-tube blood loss (6 hr – 4: Range of estimates include	ue (p<0.001) • <i>4 day)</i> es clinically				
	1: Clinically important benefit limit does not include null value. Chest-tube blood loss (6 hr – 4: Range of estimates include important effects, but also co	ue (p<0.001) • <i>4 day)</i> es clinically				
	Clinically important benefit limit does not include null value. Chest-tube blood loss (6 hr – 4: Range of estimates include important effects, but also cono or harmful effect.	ue (p<0.001) • <i>4 day)</i> es clinically				
	1: Clinically important benefit limit does not include null value Chest-tube blood loss (6 hr – 4: Range of estimates include important effects, but also cono or harmful effect Chest-tube blood loss (total)	ue (p<0.001) 4 day) es clinically mpatible with				
	1: Clinically important benefit limit does not include null value Chest-tube blood loss (6 hr – 4: Range of estimates include important effects, but also cono or harmful effect Chest-tube blood loss (total) 1: Clinically important benefit	ue (p<0.001) 4 day) es clinically mpatible with c, confidence				
	1: Clinically important benefit limit does not include null value Chest-tube blood loss (6 hr – 4: Range of estimates include important effects, but also cono or harmful effect Chest-tube blood loss (total) 1: Clinically important benefit limit does not include null value.	ue (p<0.001) 4 day) es clinically mpatible with c, confidence				
Morbidity	1: Clinically important benefit limit does not include null value Chest-tube blood loss (6 hr – 4: Range of estimates include important effects, but also cono or harmful effect Chest-tube blood loss (total) 1: Clinically important benefit	ue (p<0.001) 4 day) es clinically mpatible with c, confidence	1: Patient-rele	vant clinical outcome.		

Haemoglobin	End of surgery	2: Predictive surrogate outcome.
Tidemoglobin	0 3	2.1 Todictive Surrogate outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible with	
	no or harmful effect	
	6 hr postoperative	
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p<0.05)	
	4 day postoperative	
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p<0.05)	

EXTERNAL VALIDITY

Generalisability

Study conducted specifically in total knee arthroplasty in patients with ASA-I – III so likely to only be generalisable to this select population. The authors note the dosing in this study is lower that seen is other studies and so results may not be generalisable to higher doses of tranexamic acid.

Applicability

Study conducted in a single centre in Spain so may not be completely applicable to the Australian/New Zealand setting.

Comments

33 patients underwent preoperative blood conservation programme (includes autologous blood transfusion, treatment with rHuEPO, and administration of elemental iron). The authors state there was no difference between groups regarding the use of these treatments. The authors note that the study was not sufficiently powered to show a difference between tranexamic acid and placebo with regards to thrombosis.

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: ASA, American Society of Anaesthesiologist; BMI, body mass index; ITT, intention-to-treat; NA, not applicable, NR, not reported; NS, not significant; RBC, red blood cell; RCT, randomised controlled trial; rHuEPO, recombinant human erythropoietin; SD, standard deviation.

^a A system used by anaesthesiologists to stratify severity of patients' underlying disease and potential for suffering complications from general anaesthesia. American Society of Anesthesiology (ASA) patient classification status defined as follows: ASA I – normal healthy patient; ASA II – patient with mild systemic disease; no functional limitation; ASA III – patient with severe systemic disease; definite functional impairment; ASA IV – patient with severe systemic disease that is a constant threat to life; ASA V – unstable moribund patient who is not expected to survive 24 hours with or without the operation; ASA VI –brain-dead patient whose organs are removed for donation to another; E – emergency operation of any type, which is added to any of the 6 above categories.

STUDY DETAILS									
Citation									
Apostolakis E, Panago	noulos N. K	alateie FM	Crockott I Star	mou Ko	uki H. Souraia	ndaki E Eilos	K Do	ougonis D (2008)	
								ournal of Cardiothoracic	
Surgery 3:14.			J						
Affiliation/Source of f	unds								
Departments of Cardio		rgery and A	nesthesiology,	Patras l	Jniversity Sch	ool of Medicir	ne, Pa	atras, Greece.	
Study design			evel of evidence		,	Location/s			
Single-blind RCT		П				Hospital (sir			
Intervention				Comp	arator	1	9		
Ultra-low dose aprotini	n: test dose	of 1mL foll	owina		o: 0.9% salin	e (regimen as	s per	aprotinin)	
intubation; 500,000 IU						(19		,	
dose following closure.			•						
Population character									
Adult patients undergo		oracic surg	ery. Mean age 5	58 years	s; female 10%	, BMI ~ 25 ka	/m².		
Length of follow-up		<u>J</u>	<i>, ,</i>	_	mes measur				
Hospitalisation period					loss, transfus		ents, i	postoperative	
					ications.	,	. 1		
INTERNAL VALIDITY									
Allocation	Results		Blinding anal	ysis	Treatment/i	measuremen	nt	Follow-up (ITT)	
Randomised using	Standard :	statistical	Single-blind.			ring outcome	S	All patients included in	
randomisation	methods u		Anaesthetist a	ware			3	analysis.	
tables.	momodo		of treatment	allocation so not likely to			he	analysis.	
tubios:			allocation. Sur	aeons					
			unaware until	bias. Possible potential for					
			patients transf	erred			-		
			to ward.		knowing allocation.				
Overall quality asses									
Fair. Random treatme	nt allocation	ı, treating a	naesthetist awa	re of as	signment so p	ossibility of b	ias; s	mall trial.	
RESULTS									
Outcome		Intervention group		Pla	Placebo		Statistical significance		
		N=29		N=30					
Ultra-low dose aprotini	n (IV)								
		Mean ± SD		Mean ± SD			P value		
Intraoperative pRBCs t (units)	ransfused	0.17 ± 0.539		0.1	0.17 ± 0.531		0.967		
Postoperative pRBCs t (units)	ransfused	0.00 ± 0.00		0.03	0.03 ± 0.183		0.97		
Intraoperative FFP tran	nsfused	0.21 ± 0.0	620	0.20	0.20 ± 0.761		0.330		
(units) Postoperative FFP tran	nsfused	0.21 ± 0.0	620	0.8	0.87 ± 1.525		0.03	5	
(units)									
Day 1 postoperative thoracic 4		412.6 ± 1	99.2	764	.3 ± 213.9		<0.0	001	
drainage (mL) Day 2 postoperative the	noracia	2402 : 1	70 E	455	0 . 274 /		0.00	11	
	IOLACIC	248.3 ± 1	70.5	455	0.0 ± 274.6		0.00	VI	
drainage (mL)		0/29 (0)		0/2/) (0)		NΙΛ		
In-hospital mortality	ina	0/29 (0)			0 (0)		NA NA		
Re-operation for bleed	ırıy	0/29 (0)		0/3	0 (0)	linical releva			
Clinical importance Ultra-low-dose aprotini	n (I\/)				1 6	linical releva	ince		
onia-iow-dose apronni	II (IV)								

r=	1	
Transfusion volume	Intraoperative pRBCs	1: Patient-relevant clinical outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
	Postoperative pRBCs	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect <i>Intraoperative</i>	
	FFP	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
	Postoperative FFP	
	1: Clinically important <i>benefit</i> , confidence	
Disadisas	limit does not include null value (p=0.035)	1 Deticat relevant eliminal automas
Blood loss	Day 1 postoperative thoracic drainage	1: Patient-relevant clinical outcome.
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p<0.001)	
	Day 2 postoperative thoracic drainage	
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p=0.001)	
Mortality	In hospital mortality	1: Patient-relevant clinical outcome.
	There were no deaths in either treatment	
	arm	
Reoperation for bleeding	In hospital mortality	1: Patient-relevant clinical outcome.
	There was no reoperation due to bleeding	
	in either treatment arm	
EXTERNAL VALIDITY		
Generalisability		
Study conducted specifically in	adult patients undergoing thoracic surgery (or w	hich most received a lateral thoracotomy for
lung resection) so likely to only	be generalisable to this select surgical population	n.
Applicability	<u> </u>	
	ntre in Greece so may possibly be applicable to	he Australian/New Zealand setting.
Comments		

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: BMI, body mass index; FFP, fresh frozen plasma; ITT, intention-to-treat; IV, intravenous; pRBCs, packed red blood cells; RCT, randomised controlled trial; SD, standard deviation.

STUDY DETAILS								
Citation								
Athanasiadis T, B	eule AG Wo	rmald P I (20	007) Effects of to	nical antif	ihrinolytics in	endosconic	sinus	surgery: a nilot
randomized contro				picai aritii	ibililolytics ili	cridoscopic .	Jirius	surgery, a phot
	Affiliation/Source of funds							
Department of Oto	Department of Otorhinolaryngology, Head and Neck Surgery, The Queen Elizabeth Hospital, University of Adelaide,							
								eifswald, Germany.
Study design			evel of evidence		,	Location/s		
Double-blind RCT						Hospital		,
Intervention		<u> </u>		Compa	rator			
Topical ε-aminoca	proic acid 2.	5 g; topical tr	ranexamic acid		(saline) – use	ed in contral	ateral	side.
100 mg; topical tra					` ,			
Population chara	cteristics	<u> </u>						
Aged > 18 years; undergoing bilateral endoscopic sinus surgery (ESS) involving complete sphenoethmoidectomy and frontal								
recess clearance								,
Length of follow-	-up			Outcon	nes measured			
Efficacy measured								rmald grading scale and
for safety analysis	taken 6 hou	rs after appli	cation of				d at 0,	2, 4, 6, 8 and 10 mins
treatment. following application of								
					erative advers			
				Postope	erative coagula	ation parame	eters.	
INTERNAL VALID								
Allocation	Results		Blinding analy	ysis	Treatment/r bias	neasuremei	nt	Follow-up (ITT)
Randomised to	Grading per		Double-blind. S	Surgical Surgeon and observer				All patients included in
treatment and	surgeon an		team and inde					analysis.
placebo (ie, left	independer	nt	observer blinde	,				
or right side).	observer.		treatment alloc			nt		
Method of			Treatments pre		bias.			
randomisation			by anaesthetis					
not reported.			given to nurse					
			labelled left an for left and righ					
Overall quality as	ssessment (descrintive)		it sirius.				
				t nrenare	treatment so	notential for	· unhir	nding. Use of rating
scales that have n			นาน นานธรมเซมร	r hi chai ci	a a camient 30	Potential IOI	uribli	iding. Use of falling
RESULTS	ot boom valle	iaiou.						
Outcome		Intervention	n group	Com	parator grou	n	Stat	istical significance
E-aminocaproic ac	cid 2.5 a		g. op	0011	.parator grou	r	Ciul	.ccar organitourioo
Wormald grading		No differen	ce between ACA	and plac	cebo		NR	
Boezaart grading			ce between ACA				NR	
Epistaxis			ce between ACA				NR	
Tranexamic acid 1	100 ma	710 United	J DOWNOON NOP	. and plac				
Wormald grading		Significant	difference betwe	en TA an	d placebo at 1	mins 4	P<0.	.05
270111ala grading	23410	mins and 6		. 511 171 UII	- piacono al z	- /1/J ₁ T	. 、 、	
Boezaart grading	scale		difference betwe	en TA an	d placebo at 2	2 mins. 4	P<0.	.05
		mins and 6			. r	=/ '		
Epistaxis			ce between TA a	and place	bo		NR	
Tranexamic acid 1	l g							
Wormald grading		No significa	ant difference be	tween TA	and placebo		P>0.	.05
Boezaart grading			ant difference be				P>0.	
Epistaxis			ce between TA a				NR	
The difference between Triana placebo								

800

Clinical importance		Clinical relevance
E-aminocaproic acid 2.5 g		
Wormald grading scale	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	5: Evidence confined to unproven surrogate outcomes.
Boezaart grading scale	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	5: Evidence confined to unproven surrogate outcomes.
Epistaxis	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Tranexamic acid 100 mg		
Wormald grading scale	1: Clinically important <i>benefit</i> , confidence limit does not include null value (p<0.05)	5: Evidence confined to unproven surrogate outcomes.
Boezaart grading scale	1: Clinically important <i>benefit</i> , confidence limit does not include null value (p<0.05)	5: Evidence confined to unproven surrogate outcomes.
Epistaxis	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Tranexamic acid 1 g	<u> </u>	
Wormald grading scale	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	5: Evidence confined to unproven surrogate outcomes.
Boezaart grading scale	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	5: Evidence confined to unproven surrogate outcomes.
Epistaxis	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
EXTERNAL VALIDITY	•	1
Generalisability		
population.	y in endoscopic sinus surgery using topical agents :	so likely to only be generalisable to this select
Applicability		
	a so likely to be applicable to the Australian/New Ze	ealand setting.
Comments		

Abbreviations: ACA, aminocaproic acid; ESS, endoscopic sinus surgery; ITT, intention-to-treat; NR, not reported; RCT, randomised controlled trial; TA, Tranexamic acid.

STUDY DETAILS							
Citation							
Berenholtz SM, Pham JC, Garrett-Mayer E et al (2009)							
Affiliation/Source of funds							
		ol of Medicin	e· Blombera Sch	nol of Pu	hlic Health: Me	edical University	of South Carolina
John Hopkins University School of Medicine; Blomberg School of Public Health; Medical University of South Carolina. Federal funds received in support of this work.					or South Carolina.		
Study design		Le	evel of evidence	9		Location/setti	ing
Double-blind RCT						Hospital (single	e-centre)
Intervention				Compa	rator		,
E-aminocaproic ad	cid 100 mg/k	g administer	ed	Placebo			
immediately following anaesthesia followed by an infusion							
10 mg/kg/hr contir	nued for 8 ho	urs after sur	gery				
Population chara							
Aged > 18 years;							
		g surgical pro	ocedures: anterio	or spinal f	fusion, posterio	or spinal fusion,	anterior-posterior spinal
fusion or osteotom				T			
Length of follow-					nes measured		
Up to postoperativ	∕e day 8						BC transfusion and
					erative RBC tra		
						blood loss, oth	
					ments, laborato	ory results, com	plications, length of stay,
INTERNAL VALID	NTV			cost.			
Allocation	Results		Dlinding analy	ucic	Troatmont/n	neasurement	Follow-up (ITT)
Allocation	Nesulis		Blinding analysis Treatment/m bias		neasurement	i ollow-up (i i i)	
Computer	Intraoperat	ive blood	Double-blind. A	All study		d to treatment	
generated block	loss determ		personnel, pati				analysis. Similar
randomisation,	anaesthesi		and care provide			measurement	proportions did not
stratified by		J	blinded. Adequ		bias.		receive allocated
surgeon.			blinding tested				intervention (6 in each
			surveying staff	to test			arm)
			their ability to				
			determine alloc				
			(κ=0.06; P=0.7	(2)			
Overall quality as							
Good. Secure met	tnod of rando	omisation use	ed, double-blindi	ng secure	e, all patients i	ncluded in analy	/SIS.
RESULTS		In the second Co		10			
Outcome		Intervention	on group		nparator grou	p St	atistical significance
E-aminocaproic ad	rid 100 ma/k	N=91		N=9	1		
L-aminocaproic ac	sia 100 mg/k	Mean ± SI)	Mea	n ± SD	P	value
Total allogeneic R	BC.	5.9 ± 4.7			± 5.4		18
transfusion (units)		J.7 I 4.1		U. / ± U.4			
Postoperative RB0		2.0 ± 1.8		2.8 ± 2.8		0.	03
transfusion (units)				2.0 ± 2.0			
Total autologous F	` '		0.6 ± 1.4		0.	27	
transfusion (units)							
Total allogeneic ar	Total allogeneic and 6.4 ±4.9			7.6	± 5.5	0.	12
	autologous RBC transfusion						
(units)							
Total FFP transfus		2.8 ± 3.9			3.5 ± 6.0		37
Total platelets tran	nsfusion	1.2 ± 3.1		1.2	± 4.8	0.	23
(units)							

Total blood products transfused (units)	10.4 ± 10.8	13.0 ±14.9		0.17
transiusea (aritis)	Mean ± SD	Mean ± SD		P value
Intraoperative blood loss (mL)	2938 ± 2315	3273 ± 2195		0.32
Post-surgery to POD 1 blood loss (mL)	3265 ± 2416	3695 ± 2341		0.23
	n/N (%)	n/N (%)		P value
In-hospital mortality	0/91 (0)	1/91 (1.1)		0.32
Deep vein thrombosis	0/91 (0)	2/91 (2.2)		0.16
Cerebral infarction/transient ischaemic attack	0/91 (0)	1/91 (1.1)		0.32
Myocardial infarction	0/91 (0)	0/91 (0)		NA
Pulmonary embolism	1/91 (1.1)	3/91 (3.3)		0.31
Acute renal failure	1/91 (1.1)	1/91 (1.1)		1.00
Any thrombotic complication	2/91 (2.2)	6/91 (6.6)		0.15
Re-operation due to bleeding	0/91 (0)	2/91 (2.2)		0.16
	Mean ± SD	Mean ± SD		P value
ICU length of stay (days)	1.8 ± 1.6	2.8 ± 4.6		0.04
Hospital length of stay (days)	8.5 ± 3.9	9.5 ± 8.6		0.32
Total hospital charges (US\$)	62,344 ± 27,497	68,670 ± 32,1	41	0.16
Clinical importance		1	Clinical relev	ance
E-aminocaproic acid 100 mg/k	g			
	Total allogeneic RBC transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative RBC transfusion 1: Clinically important benefit, confidence limit does not include null value (p=0.03) Total autologous RBC transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total allogeneic and autologous RBC transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total FFP transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total plasma transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total blood products transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total blood products transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect			

		T
Blood loss	Intraoperative blood loss 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Post-surgery to POD 1 blood loss 4: Range of estimates includes clinically important effects, but also compatible with no	1: Patient-relevant clinical outcome.
	or harmful effect	
Mortality	In-hospital mortality 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Morbidity	Deep vein thrombosis 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Cerebral infarction/transient ischaemic attack 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Myocardial infarction 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Pulmonary embolism 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Acute renal failure 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Any thrombotic complication 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Re-operation due to bleeding 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Re-operation due to bleeding 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Other	ICU length of stay (days) 1: Clinically important benefit, confidence limit does not include null value (p=0.04) Hospital length of stay (days) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total hospital charges (US\$) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	2: Predictive surrogate outcome.
EXTERNAL VALIDITY	or narmar cirect	
Generalisability		
5	n certain types of spinal surgery which are known	to result in the most blooding (in anterior
	fusion, anterior-posterior spinal fusion or osteotor	

Applicability

Study conducted in the US so may not be completely applicable to the Australian/New Zealand setting.

Comments

The authors note that the study was not sufficiently powered to show a difference between ϵ -aminocaproic acid and placebo. They state that due to the variability between results (which may have related to differences in clinical practice between the individual surgeons involved) a sample size of approximately 1088 would have been required to show a 1-unit reduction in total RBC transfusion.

Abbreviations: FFP, fresh frozen plasma; ICU, intensive care unit; ITT, intention-to-treat; POD, postoperative day; RBCs, red blood cells; RCT, randomised controlled trial; SD, standard deviation.

STUDY DETAILS Citation Chen CC, Wang CC, Wang, CP et al (2008) Prospective, randomized, controlled trial of tranexamic acid in patients who undergo head and neck surgery. Otolaryngology – Head and Neck Surgery 138: 762-767. Affiliation/Source of funds Department of Otolaryngology and Hematology Division, Department of Internal Medicine, Taichung Veterans General Hospital, Taichung, Taiwan; Faculty of Medicine, School of Medicine, National Yang-Ming University, Taipei; Department of Information Technology, Overseas Chinese Institute of technology, Taichung, Taiwan. Supported by a grant from Taichung Veterans General Hospital. Level of evidence Location/setting Study design Double-blind RCT Hospital (single-centre) Intervention Comparator Tranexamic acid: preoperative dose of IV TXA 10 mg/kg Placebo: regimen as per tranexamic acid followed by continuous infusion of 1 mg/kg/hr during the Population characteristics Aged 20-80 years; scheduled to undergo head and neck surgery. Mean age ~ 48; BMI ~24.5; 44% female. Outcomes measured Length of follow-up Hospitalisation period. Primary outcome: drainage duration Secondary outcome: drainage volume (blood loss); perioperative bleeding Other outcomes: hospitalisation, coagulation profiles INTERNAL VALIDITY Allocation Blinding analysis Treatment/measurement Follow-up (ITT) Results bias Computer-Results determined Double-blind. Neither Staff blinded to treatment 5 patients excluded patient not the generated from analysis following by anaesthetist not allocation so not likely to be aware of treatment randomisation (3 randomisation anaesthetist treatment or measurement list. Treatment assignment. assessing results bias. treatment/2 control) Standard statistical aware of treatment prepared by staff not involved in methods used. assignment. the study. Overall quality assessment (descriptive) Fair. 5/60 (8%) subjects excluded from analysis post-randomisation (3 in treatment group and 2 in control group). These subjects were not further described. RESULTS Outcome Intervention group Comparator group Statistical significance N=26 N = 29Tranexamic acid Mean ± SD Mean ± SD P value 115.5 ± 120.3 86.5 ± 128.5 0.392 Perioperative bleeding (mL) Drainage amount (mL) 49.7 ± 32.6 88.8 ± 89.9 0.041 n/N (%) n/N (%) P value Deep vein thrombosis 0/26 (0) 0/29 (0) NA Mean ± SD Mean ± SD P value Length of hospital stay (days) 4.81 ± 0.80 5.31 ± 1.26 0.087

Clinical importance

Tranexamic acid

Clinical relevance

Blood loss	Perioperative bleeding	1: Patient-relevant clinical outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible with	
	no or harmful effect	
	Drainage amount	
	1: Clinically important benefit, confidence	
	limit does not include null value (p=0.041)	
Morbidity	Thrombosis	1: Patient-relevant clinical outcome.
_	No events in either group	
Length of hospital stay	Length of hospital stay	2: Predictive surrogate outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible with	
	no or harmful effect	
EXTERNAL VALIDITY		
Generalisability		
Study conducted specifically	in patients undergoing head and neck surgery so	likely to only be generalisable to this select

population. **Applicability**

Study conducted in a single centre in Taiwan so may not be completely applicable to the Australian/New Zealand setting.

Comments

The authors note that head and neck surgery generally results in less blood loss than cardiac and orthopaedic surgery so that tranexamic acid may not be as efficacious is head and neck surgery. Study was underpowered to detect a difference as powered based on blood loss seen in orthopaedic surgery.

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: BMI, body mass index; ITT, intention-to-treat; IV, intravenous; RCT, randomised controlled trial; SD, standard deviation; TXA, tranexamic acid.

STUDY DETAILS Citation Choi WS, Irwin MG, Samman N (2009) The effect of tranexamic acid on blood loss during orthognathic surgery: a randomized controlled trial. J Oral Maxillofac Sug 67: 125-133. Affiliation/Source of funds Faculty of Dentistry and Medicine, University of Hong Kong, Hong Kong, P. R. China. Funding not reported. Study design Level of evidence Location/setting Double-blind RCT Hospital (single-centre) Intervention Comparator Tranexamic acid: 20 mg/kg immediately prior to surgery Placebo: regimen as per tranexamic acid Population characteristics Aged 16-40 years; scheduled for bimaxillary osteotomy at Queen Mary Hospital; ASA class Ia. Mean age ~ 23 years, 66% female; mean weight ~57 kg. Length of follow-up Outcomes measured Blood loss; patients requiring transfusion; length of hospital Hospitalisation period. stay; haematology. INTERNAL VALIDITY Allocation Results Blinding analysis Treatment/measurement Follow-up (ITT) bias Computer-Results determined Double-blind. Neither Staff blinded to treatment 12 patients (16%) by anaesthetist not allocation so not likely to be excluded from generated patient nor the randomisation aware of treatment anaesthetist, surgeon treatment or measurement analysis following list/sealed assignment. or nurse aware of bias. randomisation (7 envelopes. Standard statistical treatment treatment/5 control) Treatment methods used. assignment. prepared by Blood analysis surgeon not adjusted for involved in the operation time. study and then transferred to anaesthetist. Overall quality assessment (descriptive) Fair. 12/73 (16%) subjects excluded from analysis post-randomisation (7in treatment group and 5 in control group). These subjects were not further described. RESULTS Outcome Intervention group Comparator group Statistical significance Tranexamic acid n/N (%) n/N (%) P value Transfusion incidence 4/32 (12.5) 7/29 (24.1) 0.32 (post-hoc) Mean ± SD Mean ± SD P value Intraoperative blood loss 277.0 ± 211.7 415.9 ± 314.2 NS during anterior mandibular N = 21N = 23surgery (mL) Intraoperative blood loss 428.0 ± 233.3 643.8 ± 430.0 < 0.05 during maxillary surgery (mL) N = 32N = 29Intraoperative blood loss 329.3 ± 233.4 287.0 ± 216.3 NS during ramus surgery (mL) N = 24N = 17

1257 ± 817.8

N=29

n/N (%)

0/29(0)

Mean ± SD

< 0.05

P value

P value

NA

878.6 ± 577.7

N=32

n/N (%)

0/32(0)

Mean ± SD

Total intraoperative blood loss

(mL)

Thrombosis

Length of hospital stay (days)	7.2 ± 2.1	7.5 ± 2.3	0.32
Clinical importance			Clinical relevance
Tranexamic acid			
Transfusion incidence	Transfusion incidence 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect		1: Patient-relevant clinical outcome.
Blood loss	Anterior mandibular surgery 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Maxillary surgery 1: Clinically important benefit, confidence limit does not include null value (p<0.05) Ramus surgery 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total 1: Clinically important benefit, confidence		1: Patient-relevant clinical outcome.
Morbidity	limit does not include null value (p<0.05) Thrombosis No events in either group		1: Patient-relevant clinical outcome.
Length of hospital stay	Length of hospital stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect		2: Predictive surrogate outcome.
EXTERNAL VALIDITY	1		•
Generalisability			
Study conducted specifically in	patients classified as ASA I un	dergoing orthog	nathic surgery (bimaxillary osteotomy) so lik

to only be generalisable to this select population.

Applicability

Study conducted in a single centre in Hong Kong so may not be completely applicable to the Australian/New Zealand setting.

Comments

The authors did not measure postoperative bleeding as no suction drains were used in intraoral wounds so there may have been significant concealed blood loss during the early postoperative period which has not been accounted for

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: ASA, American Society of Anaesthesiologist; ITT, intention-to-treat; NS, not significant; RCT, randomised controlled trial; SD, standard

a A system used by anaesthesiologists to stratify severity of patients' underlying disease and potential for suffering complications from general anaesthesia. American Society of Anesthesiology (ASA) patient classification status defined as follows: ASA I – normal healthy patient; ASA II – patient with mild systemic disease; no functional limitation; ASA III – patient with severe systemic disease; definite functional impairment; ASA IV – patient with severe systemic disease that is a constant threat to life; ASA V – unstable moribund patient who is not expected to survive 24 hours with or without the operation; ASA VI –brain-dead patient whose organs are removed for donation to another; E – emergency operation of any type, which is added to any of the 6 above categories.

STUDY DETAILS

Citation

Colwell Jr CW, Chelly JE, Murkin JM, Stevens D, O'Keefe TJ, Hall R, Parvizi J (2007) Randomized study of aprotinin effect on transfusions and blood loss in primary THA. Clinical Orthopaedics and Related Research 465: 189-195.

Affiliation/Source of funds

Shiley Centre for Orthopaedic Research & Education at Scripps Clinic, La Jolla, CA; the Department of Anesthesiology, University of Pittsburgh School of Medicine, Pittsburgh, PA; The department of Anesthesia, London Health Sciences Centre-UC, University of Western Ontario, Ontario Canada; Grand River Hospital, Kitchener, Ontario, Canada; Michigan Orthopaedic Centre, Ypsilanti, MI; the Department of Anesthesiology, Queen Elizabeth II Health Sciences Centre, Halifax Infirmary, Halifax, Nova Scotia, Canada; and the Roman Institute, Philadelphia, PA.

Hospital	
Comparator	
n Placebo: saline (regimen as per aprotinin)	
0	

Population characteristics

Adult patients undergoing elective, unilateral, primary THA; > 18 years. Mean age 64 years; female 51%.

Length of follow-up	Outcomes measured
Hospitalisation period	Blood loss, transfusion requirements, ICU stay.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement	Follow-up (ITT)
			bias	
Randomised using computer-generated codes and managed by IVRS.	Standard statistical methods used.	Described as double-blind. Patient and staff unaware of treatment	No evidence that there may be treatment/measurement bias.	All patients included in analysis.
		assignment.		

Overall quality assessment (descriptive)

Good. Random treatment allocation, double-blind, all patients included in analysis, large trial.

RESULTS

Outcome	Intervention group N=175	Placebo N=177	Statistical significance
Aprotinin (IV)			
	n/N (%)	n/N (%)	P value
Transfusion incidence (whole blood or RBCs)	30/175 (17)	57/177 (32)	0.0009
Transfusion incidence (allogeneic blood)	19/175 (11)	39/177 (22)	0.006
Transfusion incidence (whole blood or RBCs without donation)	18/140 (13)	33/138 (24)	0.02
Transfusion incidence (whole blood or RBCs with donation)	12/37 (32)	23/37 (62)	ND (small sample size)
	Mean ± SD	Mean ± SD	P value
Transfusion volume (whole blood or RBCs; units)	0.27a	0.63 a	0.0003
Transfusion volume (allogeneic blood; units)	0.17 a	0.42 a	0.004
Transfusion volume (whole blood or RBCs without donation; units)	0.21 a	0.46 a	0.0153

810

Transfusion volume (whole blood or RBCs with donation;	0.52 a	1.21 a		ND (small sample size)
units)				
	LSM (95% CI)	LSM (95% CI)		P value
Intraoperative blood loss (mL)	331 (297, 368)	385 (346, 429		0.0217
0-6 hr drainage (mL)	96 (72, 129)	177 (133, 235		0.0003
Total drainage (mL)	276 (216, 353)	390 (307, 494	,	0.0141
Total fluid loss (mL)	709 (618, 813)	957 (837, 109		0.0002
	n/N (%)	n/N (%)	,	P value
Mortality	0/175 (0)	1/177 (0.6)		NS
Deep vein thrombosis	2/175 (1.1)	3/177 (1.7)		NS
Pulmonary embolism	2/175 (1.1)	2/177 (1.1)		NS
Myocardial infarction	1/175 (0.6)	1/177 (0.6)		NS
Renal failure	2/175 (1.1)	2/177 (1.1)		NS
Clinical importance			Clinical relev	ance
Aprotinin (IV)				
Transfusion incidence	Whole blood or RBCs	a. a.	1: Patient-rele	evant clinical outcome.
	1: Clinically important bene			
	limit does not include null va	alue		
	(p=0.0009)			
	Allogeneic blood	fit confidence		
	1: Clinically important <i>benefit</i> , confidence limit does not include null value (p=0.006) <i>Whole blood or RBCs (– donation)</i>			
		Clinically important <i>benefit</i> , confidence limit does not include null value (p=0.02)		
	Whole blood or RBCs (+ do			
	1: Clinically important <i>bene</i>			
	limit does not include null va			
Transfusion volume	Whole blood or RBCs	а. с. (р. 112)	1: Patient-rele	evant clinical outcome.
Transferent Volume	1: Clinically important bene	fit. confidence		van em near e a teenner
	limit does not include null va			
	(p=0.0003)			
	Allogeneic blood			
	1: Clinically important <i>benefit</i> , confidence limit does not include null value (p=0.004) <i>Whole blood or RBCs (– donation)</i> 1: Clinically important <i>benefit</i> , confidence			
		limit does not include null value		
	(p=0.0153)			
	Whole blood or RBCs (+ do			
	1: Clinically important bene			
	limit does not include null va	aiue (p=ND)		

Blood loss	Intraoperative 1: Clinically important benefit, confidence limit does not include null value (p=0.0217) 0-6 hr drain 1: Clinically important benefit, confidence limit does not include null value (p=0.0003) Total drainage 1: Clinically important benefit, confidence limit does not include null value (p=0.0141) Total fluid loss 1: Clinically important benefit, confidence limit does not include null value (p=0.0002)	1: Patient-relevant clinical outcome.
Mortality	Mortality 4: Range of estimates includes clinically important point estimate but also compatible with no effect, or a harmful effect	1: Patient-relevant clinical outcome.
Deep vein thrombosis	Deep vein thrombosis 4: Range of estimates includes clinically important point estimate but also compatible with no effect, or a harmful effect	1: Patient-relevant clinical outcome.
Pulmonary embolism	Pulmonary embolism 4: Range of estimates includes clinically important point estimate but also compatible with no effect, or a harmful effect	1: Patient-relevant clinical outcome.
Myocardial infarction	Myocardial infarction 4: Range of estimates includes clinically important point estimate but also compatible with no effect, or a harmful effect	1: Patient-relevant clinical outcome.
Renal failure	Renal failure 4: Range of estimates includes clinically important point estimate but also compatible with no effect, or a harmful effect	1: Patient-relevant clinical outcome.
EXTERNAL VALIDITY		
Generalisability Study conducted specifically in population. Applicability	adult patients undergoing total hip arthroplasty	so likely to be generalisable to this surgical
Study conducted in the US and	Canada so likely to be applicable to the Austral	lian/New Zealand setting.
Comments	-	-
Note: All post-hoc calculations perform	ad using Craph Dad software	

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: CABG, coronary artery bypass graft; FFP, fresh frozen plasma; ITT, intention-to-treat; IV, intravenous; IVRS, interactive voice response system; LSM, least squares mean; ND, not determined; NS, not significant; pRBCs, packed red blood cells; RCT, randomised controlled trial; SD, standard deviation.

^a Calculated post-hoc. Approximation based on the proportion of patients who received 1, 2, 3 or 4 units of transfusion.

STUDY DETAILS Citation Elwatidy S, Jamjoon Z, Elgamal E et al (2008) Efficacy and safety of prophylactic large dose of tranexamic acid in spine surgery: a prospective, randomized, double-blind, placebo-controlled study. Spine 33(24): 2577-2580. Affiliation/Source of funds Division of Neurosurgery and Department of Anaesthesia, College of Medicine King Aaud University, Riyadh, Saudi Arabia. Funded by CRMC, College of Medicine, King Saud University. Drug and placebo provided by pharmacy department King Khalid University Hospital. Study design Level of evidence Location/setting Double-blind RCT Hospital (single-centre) Intervention Comparator Placebo: 0.9% saline; regimen as per tranexamic acid Tranexamic acid: loading dose of 2 g (adults) or 30 mg/kg (children) followed by continuous infusion of 100 mg/hr (adults) or 1 mg/hr/kg (children) during surgery and 5 hours following surgery. Population characteristics Patients (adults or children) undergoing spine surgery with expectant significant blood loss. Mean age ~ 50 years, 39% female; mean weight ~71 kg. Length of follow-up Outcomes measured Up to 12 months – 3 years. Blood loss; patients requiring transfusion; transfusion volume; length of hospital stay; haematology. INTERNAL VALIDITY Allocation Results Blinding analysis Treatment/measurement Follow-up (ITT) bias Randomised Anaesthetist and Double-blind, Neither Staff blinded to treatment All 64 randomised using odd/even surgeon unaware of patient, the allocation so not likely to be patients included in numbers. treatment anaesthetist or treatment or measurement analysis. Treatment assignment. surgeon aware of bias. Standard statistical treatment prepared by pharmacy staff methods used. assignment. who did not know patients. Overall quality assessment (descriptive) Fair. Non-secure method of randomisation used (odd/even numbers). Study double-blind. All patients included in analysis. **RESULTS** Outcome Intervention group Statistical significance Comparator group Tranexamic acid n/N (%) n/N (%) P value 4/32 (12.5) 12/32 (37.5) 0.021 Transfusion incidence Mean ± SD Mean ± SD P value Amount of transfusion (mL) 93.75 ± 267.53 531.25 ± 1275.94 800.0 Units transfused/patient 1.5 2.8a NA (transfused patients only) Intraoperative blood loss (mL) 311.25 ± 412.49 584.69 ± 797.30 0.03 Wound drain blood loss (mL) 97.94 ± 136.28 215.31 ± 276.04 0.004 Total blood loss (mL) 406.13 ± 495.31 800.00 ± 1034.25 0.007 n/N (%) n/N (%) P value Thrombosis 0/32(0) 0/32 (0) Mean ± SD Mean ± SD P value Length of hospital stay (days) 8.45 ± 5.79 10.69 ± 8.27 0.21 Mean ± SD Mean ± SD P value Postoperative haemoglobin 11.35 ± 1.57 12.39 ± 1.28 0.006 (a/dL) Clinical importance Clinical relevance Tranexamic acid

Transfusion incidence	Transfusion incidence	1: Patient-relevant clinical outcome.
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p=0.021)	
Transfusion volume	Amount of transfusion	1: Patient-relevant clinical outcome.
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p=0.008)	
Blood loss	Intraoperative blood loss	1: Patient-relevant clinical outcome.
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p=0.03)	
	Wound drain blood loss	
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p=0.004	
	Total blood loss	
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p=0.007)	
Morbidity	Thrombosis	1: Patient-relevant clinical outcome.
	No events in either group	
Length of hospital stay	Length of hospital stay	2: Predictive surrogate outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible with	
	no or harmful effect	
Haemoglobin	Postoperative haemoglobin	2: Predictive surrogate outcome.
	1: Clinically important benefit, confidence	
	limit does not include null value (p<0.05)	
EXTEDNIAL MALIDITA		

EXTERNAL VALIDITY

Generalisability

Study conducted specifically in adult and paediatric patients classified undergoing spine surgery (so likely to only be generalisable to this select surgical population.

Applicability

Study conducted in a single centre in Saudi Arabia so may not be completely applicable to the Australian/New Zealand setting.

Comments

The authors used a larger than usual dose of tranexamic acid in this study in both adults and children. They state that there were no haemodynamic disturbances, apparent thromboembolic events, or other drug disturbances (including disturbed colour vision, numbness or weakness, confusion or allergic reaction) associated with its use. However, the small sample size should be noted when assessing the safety of this dose of tranexamic acid.

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: ITT, intention-to-treat; NA, not applicable; NS, not significant; RCT, randomised controlled trial; SD, standard deviation.

^a One patient received 14 units of blood. If this person is excluded the mean units of blood transfused per transfused patient is 1.8.

STUDY DETAILS Citation Fawzy H, Elmistekawy E, Bonneau D et al (2009) Can local application of tranexamic acid reduce post-coronary bypass surgery blood loss? A randomized controlled trial. Journal of Cardiothoracic Surgery 4:25. Affiliation/Source of funds Division of Cardiovascular and Thoracic Surgery, St Michael's Hospital, University of Toronto, Toronto, Ontario, Canada. Not funded. Study design Level of evidence Location/setting Double-blind RCT Hospital (single-centre) located in Saudi Ш Arabia (West Armed Forces Hospital, Tabuk) Intervention Comparator Tranexamic acid: 1 g TXA diluted in 100 ml normal saline Placebo: 100 mL normal saline; regimen as per tranexamic applied locally into the pericardial and mediastinal acid cavities. Population characteristics Patients scheduled for primary isolated elective CABG. Mean age ~ 57 years, 5% female; mean weight ~72 kg. Length of follow-up Outcomes measured Hospitalisation period Blood loss; transfusion volume; length of hospital stay; haematology. INTERNAL VALIDITY Allocation Results Blinding analysis Treatment/measurement Follow-up (ITT) All 38 randomised Randomised Staff measuring Double-blind. Neither Staff blinded to treatment using random outcomes unaware the patient, surgeon, allocation so not likely to be patients included in numbers table. anaesthetist, scrub treatment or measurement of treatment analysis. nurse nor perfusionist Treatment assignment. bias. Standard statistical knew treatment prepared by research methods used. assignment. pharmacist who delivered to operating room. Overall quality assessment (descriptive) Good. Random treatment allocation, double-blind, all patients included in analysis RESULTS Outcome Intervention group Statistical significance Comparator group Tranexamic acid (topical) Median Median P value Postoperative transfusion of 0.82 pRBCs (units) Postoperative transfusion of 0 2 0.42 FFP (units) Postoperative transfusion of 0 2 0.03 platelets (units) 24-hour chest-tube loss (mL) 626 1040 0.04 656 (range 248-2105) Total chest-tube loss (mL) 1056 (range 210-3010) NR n/N (%) n/N (%) P value In-hospital mortality 0/19(0) 0/19 (0) NA Re-operation for bleeding 1/19 (5.3) 0/19 (0) NR In-hospital myocardial 0/19(0) 0/19 (0) NA infarction Mean ± SD Mean ± SD P value Length of hospital stay (days) 7.5 ± 3 7.8 ± 2 0.68 29 ± 26 Length of ICU stay (hours) 49 ± 20 0.02

Postoperative haemoglobin	10 ± 1.3	10 ± 1.3		0.39
(g/dL)				
Clinical importance			Clinical relev	ance
Tranexamic acid (topical)				
Transfusion volume	Postoperative pRBCs 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative platelets 1: Clinically important benefit, confidence limit does not include null value (p=0.03)			evant clinical outcome.
Blood loss	24-hour chest-tube loss 1: Clinically important benefit, limit does not include null value Total chest-tube loss NR		1: Patient-rele	evant clinical outcome.
Mortality	In hospital mortality No events in either group		1: Patient-rele	vant clinical outcome.
Morbidity	Re-operation for bleeding NR Myocardial infarction No events in either group		1: Patient-rele	evant clinical outcome.
Length of hospital stay	Length of hospital stay 4: Range of estimates include important effects, but also con no or harmful effect		1: Patient-rele	vant clinical outcome.
Length of ICU stay	Length of ICU stay 1: Clinically important benefit, limit does not include null value.		2: Predictive s	surrogate outcome.
Haemoglobin	Postoperative haemoglobin 4: Range of estimates include important effects, but also con no or harmful effect		2: Predictive s	surrogate outcome.
EXTERNAL VALIDITY Generalisability				
Study conducted specifically ir population. The authors note the further limit the generalisability	n patients undergoing primary CA hat their strict inclusion/exclusion of the results within the populati	n criteria, small s	sample size and	surgical methods used may
Applicability Study conducted in a single ce	entre in Saudi Arabia so may not	be completely a	applicable to the	Australian/New Zealand
setting.			11	
Comments				

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: CABG, coronary artery bypass graft; FFP, fresh frozen plasma; ICU, intensive care unit; ITT, intention-to-treat; NA, not applicable; NR, not reported; pRBCs, packed red blood cells; RCT, randomised controlled trial; SD, standard deviation; TXA, tranexamic acid.

STUDY DETAILS Citation Gharabaghian M, Eghtesadi-Araghi P (2006) The efficacy of epsilon-aminocaproic acid and its timing in reducing blood loss in major cardiac coronary bypass surgery: a randomized double-blinded placebo-controlled study. International journal of Pharmacology 2(1): 131-135. Affiliation/Source of funds Imam Khomeini Hospital Complex, Tehran University of Medical Sciences, Tehran, Iran; Parsteb Pajouheshyar Medical Sciences Research Institute, Tehran, Iran Funding not stated. Study design Level of evidence Location/setting RCT Hospital (single-centre) Intervention Comparator ε-aminocaproic acid administered using two different Placebo regimens: (i) Post-heparin group – normal saline preincision and post-incision and ACA at 150 mg/kg over 10 minutes after heparin injection and 15 mg/kg/hr from three minutes following heparin injection to the end of CPB; (ii) Pre-incision group – 150 mg/kg over 10 mins as pre-incision bolus, 15 mg/kg/hr as post-incision infusion and saline as post-heparin and 15 mg/kg/hr ACA as 3 min following heparin to end of CPB. Population characteristics Aged > 18 years; undergoing primary CABG requiring at least 4 grafts with CPB. Length of follow-up Outcomes measured Not stated Blood loss (chest-tube drainage) INTERNAL VALIDITY Allocation Results Blinding analysis Treatment/measurement Follow-up (ITT) bias Randomised. Blood loss Double-blinding Unclear if there was Assumed that all No method determined by cheststated. No further potential for treatment or patients included in tube drainage at 6 reported. details provided. measurement bias. analysis. hrs, 12 hrs and on removal of drainage. Overall quality assessment (descriptive) Fair. Unclear if treatment allocation robust or full follow-up of patients RESULTS Outcome Intervention group Comparator group Statistical significance N=20 (each group) N = 20E-aminocaproic acid (post-heparin group) Mean ± SD Mean ± SD P value 6-hr chest tube loss (mL) \sim 300 ± NR \sim 600 ± NR < 0.05 12-hr chest tube loss (mL) ~500 ± NR ~650 ± NR >0.05 ~2000 ± NR Final chest tube loss (mL) ~800 ± NR < 0.05 E-aminocaproic acid (pre-incision group) Mean ± SD Mean ± SD P value 6-hr chest tube loss (mL) \sim 300 ± NR $\sim 600 \pm NR$ < 0.05 ~650 ± NR >0.05 12-hr chest tube loss (mL) \sim 500 ± NR Final chest tube loss (mL) ~1000 ± NR ~2000 ± NR < 0.05 Clinical importance Clinical relevance E-aminocaproic acid (post-heparin group)

· · ·		
Blood loss	6-hr chest tube loss	1: Patient-relevant clinical outcome.
	1: Clinically important benefit, confidence	
	limit does not include null value (p<0.05)	
	12-hr chest tube loss	
	4: Range of estimates includes clinically	
	important effects, but also compatible with no	
	or harmful effect	
	Total chest tube loss	
	1: Clinically important benefit, confidence	
	limit does not include null value (p<0.05)	
E aminocaproic acid (pre-incis		
Blood loss	6-hr chest tube loss	1: Patient-relevant clinical outcome.
	1: Clinically important benefit, confidence	
	limit does not include null value (p<0.05)	
	12-hr chest tube loss	
	4: Range of estimates includes clinically	
	important effects, but also compatible with no	
	or harmful effect	
	Total chest tube loss	
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p<0.05)	
EXTERNAL VALIDITY	mint does not molade han value (p 10.00)	
Generalisability		
	dergoing major CABG surgery so likely to be ge	neralisable to this population only.
Applicability		
Study conducted in Iran so ma	y not be applicable to the Australian/New Zealan	d setting.
Comments		
The authors note that there wa	s no difference in chest tube blood loss between	the two different ε-aminocaproic acid
regimens.		·

Abbreviations: ACA, aminocaproic acid; CABG, coronary artery bypass graft; CPB, cardiopulmonary bypass; FFP, fresh frozen plasma; ICU, intensive care unit; ITT, intention-to-treat; NR, not reported; RCT, randomised controlled trial; SD, standard deviation.

No evidence that there may

be treatment/measurement

bias.

130 randomised; 10

not analysed (5 each

decision to use CPB.

4 patients did not receive CT

angiographic follow-

arm) due to

intraoperative

STUDY DETAILS Citation Grant MC, Kon Z, Joshi A, Christenson E, Kallam S, Burris N, Gu J, Poston RS (2008) Is aprotinin safe to use in a cohort at increased risk for thrombotic events; results from a randomized, prospective trial in off-pump coronary artery bypass. Ann Thorac Surg 86: 815-822 Affiliation/Source of funds Division of Cardiac Surgery, Department of Surgery, University of Maryland Medical System, University of Maryland, Baltimore County, Baltimore, Maryland, US Level of evidence Location/setting Study design Double-blind RCT \parallel Hospital Intervention Comparator Aprotinin: loading dose 2 million KIU followed by 0.5 million Placebo: saline (regimen as per aprotinin) KIU per hours until the end of surgery. Population characteristics Adult patients undergoing off-pump coronary artery bypass surgery. Baseline demographic data not reported. Intraoperative data similar between groups. Outcomes measured Length of follow-up Renal function; blood loss; mortality; morbidity Up to 1 year INTERNAL VALIDITY Allocation Results Blinding analysis Treatment/measurement Follow-up (ITT) bias

Overall quality assessment (descriptive)

Powered to detect

major cardiac and

cerebrovascular

events (MACCE).

methods used.

Standard statistical

Computer-generated

randomisation based

on permuted blocks

of 4.

Fair. Randomised, described as double-blind, no baseline demographics reported; 5 patients from each arm not included in analysis.

Described as

double-blind.

delivered to

Placebo use; drug

operating room in

unlabelled bottle.

Outcome	Intervention group	Placebo	Statistical significance
	N=59	N=61	
Aprotinin (IV)			
	Mean ± SD	Mean ± SD	P value
Intraoperative blood loss (mL)	867 ± 413a	1252 ± 380	<0.02
	870 ± 383 ^b		
Postoperative blood loss (mL)	415 ± 330 ^a	716 ± 336	< 0.003
. , ,	427 ± 171 ^b		
	n/N (%)	n/N (%)	P value
MACCE	7/59 (11.8)	21/61 (34.4)	<0.005
1-year mortality	3/59 (5.1)	8/61 (13.1)	NS
6-month acute occlusion	3/80 SVG (3.8)	8/90 SVG (8.9)	NS
In-hospital stroke	0/59 (0)	1/61 (1.6)	NS
In-hospital myocardial infarction	1/59 (1.7)	4/61 (6.6)	NS
Postoperative acute kidney	27/59 (45.8)	15/61 (24.6)	<0.03
injury	, ,	, ,	
Acute renal failure within 6	2/59 (3.4)	2/61 (3.3)	NS
months			

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Clinical importance		Clinical relevance
Aprotinin (IV)		
Blood loss	Intraoperative blood loss 1: Clinically important benefit, confidence limit does not include null value (p<0.02) Postoperative blood loss 1: Clinically important benefit, confidence limit does not include null value (p<0.003)	1: Patient-relevant clinical outcome.
MACCE	MACCE 1: Clinically important <i>benefit</i> , confidence limit does not include null value (p<0.005)	1: Patient-relevant clinical outcome.
Mortality	Mortality 4: Range of estimates includes clinically important point estimate but also compatible with no effect, or a harmful effect	1: Patient-relevant clinical outcome.
Stroke	In-hospital stroke 4: Range of estimates includes clinically important point estimate but also compatible with no effect, or a harmful effect	1: Patient-relevant clinical outcome.
Myocardial infarction	In-hospital myocardial infarction 4: Range of estimates includes clinically important point estimate but also compatible with no effect, or a harmful effect	1: Patient-relevant clinical outcome.
Renal effects	Acute kidney injury 1: Clinically important harm, confidence limit does not include null value (p<0.03) Renal failure 4: Range of estimates includes clinically important point estimate but also compatible with no effect, or a harmful effect	1: Patient-relevant clinical outcome.
EXTERNAL VALIDITY		
Generalisability		
	in adult patients undergoing off-pump coronary art	ery bypass so likely to be generalisable to
this surgical population.		
Applicability		
3	o likely to be applicable to the Australian/New Zeal	and setting.
Comments	al failt an ann alt and with a stadio lead	

Authors note all cases of renal failure resolved without dialysis

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: ITT, intention-to-treat; IV, intravenous; MACCE, major adverse cardiac and cerebrovascular event; NS, not significant; RCT, randomised controlled trial; SD, standard deviation; SVG, saphenous vein graft.

a Patients with peak aprotinin levels > 271 KIU/ML.

b Deticate with peak aprotinin levels > 271 KIU/ML.

^b Patients with peak aprotinin levels < 271 KIU/mL.

STUDY DETAILS Citation Jabalami M, Zakeri K (2006) Evaluation of topical tranexamic acid on intraoperative bleeding in endoscopic sinus surgery. Iran J Med Sci 31(4): 221-223. Affiliation/Source of funds Department of Anesthesiology and Intensive Care, Alzahra General Hospital, Isafan University of Medical Sciences, Isfahan, Iran. Funding not stated. Level of evidence Location/setting Study design **RCT** Hospital (single centre) Ш Intervention Comparator Tranexamic acid: 1 g diluted in 20 mL saline applied Placebo – 20 mL saline applied topically topically Population characteristics Adult patients (18-55 years) undergoing elective endoscopic sinus surgery. Female 32%. Authors note no significant difference between groups in terms of age, MABP, preoperative pulse rate, duration of surgery and operation indications. Length of follow-up Outcomes measured Intraoperative. Intraoperative bleeding INTERNAL VALIDITY Allocation Results Blinding analysis Treatment/measurement Follow-up (ITT) bias Not stated but Unclear due to lack of Unclear. Assumed to Randomised, No. No details further details provided on assumed to be information provided on be all patients. provided. assessment of blinded due to use of allocation and blinding. placebo. outcomes or statistical analysis. Overall quality assessment (descriptive) Poor. Allocation concealment and blinding poorly reported. Assessment of outcomes poorly reported. **RESULTS** Outcome Intervention group Comparator group Statistical significance N=26 N=30 Tranexamic acid (topical) Mean ± SD Mean ± SD P value Intraoperative blood loss (mL) 174.0 ± 10.6 229.1 ± 23.8 < 0.05 Clinical importance Clinical relevance Tranexamic acid (topical) **Blood loss** Intraoperative blood loss 1: Patient-relevant clinical outcome. 1: Clinically important benefit, confidence limit does not include null value (p<0.05) **EXTERNAL VALIDITY** Generalisability Study conducted specifically in adult patients undergoing elective endoscopic sinus surgery so likely to be generalisable only to this select group of patients. **Applicability** Study conducted in a single centre in Iran so may not be applicable to the Australian/New Zealand setting. Comments

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: ITT, intention-to-treat; MABP, mean arterial blood pressure; RCT, randomised controlled trial; SD, standard deviation; TXA, tranexamic acid.

STUDY DETAILS Citation Jimenez JJ, Iribarren JL, Lorente L et al (2007) Tranexamic acid attenuates inflammatory response in cardiopulmonary bypass surgery through blockade of fibrinolysis: a case control study followed by a randomized controlled trial. Critical care: Affiliation/Source of funds Intensive Care Department, Hematology Department, Research Unit, Cardiac Surgery Department and Biochemistry and Central laboratories, Hospital Universitario de Canarias, Ofra s/n La Cuesta, La Laguna, Spain. Funded by the Fundación Canaria de Investigación y Salud (FUNCIS) Level of evidence Location/setting Study design Double-blind RCT Hospital (single-centre) Intervention Comparator Tranexamic acid: 2g before and after intervention. Placebo: 0.9% saline Population characteristics Adult patients undergoing elective CPB surgery; age 67 years, female 46%, BMI 28 mg/kg². Outcomes measured Length of follow-up Hospitalisation period Primary: inflammatory response and vasoplegic shock Secondary outcomes: inflammation, coagulation and fibrinolysis parameters INTERNAL VALIDITY Allocation Treatment/measurement Follow-up (ITT) Results Blinding analysis bias Randomly assigned Double-blind. Staff blinded to treatment All 50 randomised Staff measuring by independent outcomes unaware allocation so not likely to be patients included in pharmacists using a of treatment treatment or measurement analysis. list of assignment. bias. pseudorandomised Standard statistical numbers to receive methods used. coded infusions of either TXA or placebo. Overall quality assessment (descriptive) Good. Random treatment allocation, double-blind, all patients included in analysis **RESULTS** Outcome Intervention group Comparator group Statistical significance N=24N=26 Tranexamic acid (IV) n/N (%) n/N (%) P value Incidence of RBC and plasma 0.39 1/24 (4.2) 2/26 (7.6) transfusion in first 4 hours Incidence of RBC and plasma 9/24 (37.5) 19/26 (73.1) 0.01 transfusion until chest tube withdrawal Incidence of plasma transfusion 1/24 (4.2) 8/26 (30.8) 0.02 until chest tube withdrawal Mean (95% CI) Mean (95% CI) P value 24-hour blood loss (mL) 464 (308, 620) 1037 (771, 1303) < 0.01 Total blood loss (mL) 835 (407, 1263) 1466 (1116, 1818) < 0.01 n/N (%) n/N (%) P value In-hospital mortality 0/24 (0) 0/26 (0) NA Mean (95% CI) Mean (95% CI) P value Length of hospital stay (days) 4.5(3,6)4(2, 5)0.34 Mean (95% CI) Mean (95% CI) P value

Length of ICU stay (hours)	3 (2, 5.5)	3.5 (2, 5)	0.96				
Clinical importance		1 , , , ,	Clinical relevance				
Tranexamic acid (IV)							
Transfusion incidence	important effects, but with no or harmful end withdrawn 1: Clinically importate limit does not include Chest-tube withdrawn 1: Clinically importate limit does not include limit does not include limit does not include withdrawn 1: Clinically importate limit does not include with limit limit does not include with limit limit does not include with limit	es includes clinically ut also compatible ffect wal RBCs and plasma nt benefit, confidence e null value (p=0.01)	1: Patient-relevant clinical outcome.				
Blood loss	limit does not includ Total blood loss 1: Clinically importa	nt <i>benefit</i> , confidence e null value (p<0.01) nt <i>benefit</i> , confidence e null value (p<0.01)	1: Patient-relevant clinical outcome.				
Mortality	In hospital mortality No events in either	, , ,	1: Patient-relevant clinical outcome.				
Length of hospital stay	Length of hospital s	tay es includes clinically ut also compatible	1: Patient-relevant clinical outcome.				
Length of ICU stay	Length of ICU stay	es includes clinically ut also compatible	1: Patient-relevant clinical outcome.				
EXTERNAL VALIDITY							
Generalisability							
surgical population.	adult patients undergoin	g CPB surgery so likely	to only be generalisable to this select				
Applicability							
Study conducted in a single cer Zealand setting.	ntre in the Canary Islands	s (Spain) so may not be	completely applicable to the Australian/New				
Comments							
The state of a constant and a substitute	er de die elikkele en maar van de		and the there are a second of the control of the co				

The study was stopped early due to the higher proportion of severe bleeding seen in the placebo group during follow-up.

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: BMI, body mass index; CI, confidence interval; CPB, cardiopulmonary bypass; FFP, fresh frozen plasma; ICU, intensive care unit; ITT, intention-to-treat; IV, intravenous; pRBCs, packed red blood cells; RCT, randomised controlled trial; TXA, tranexamic acid.

STUDY DETAILS Citation Later AFL, Maas JJ, Engbers FHM et al (2009) Tranexamic acid and aprotinin in low- and intermediate risk cardiac surgery: a non-sponsored, double-blind, randomised placebo-controlled trial. European Journal of Cardiothoracic Surgery 36: 322-Affiliation/Source of funds Departments of Cardiothroacic Surgery, Anaesthesiology and Intensive Care Medicine, Lelds Universitair Medisch Centrum (LUMC), The Netherlands. Funded by intramural sources only. Level of evidence Location/setting Study design Double-blind RCT Hospital (single-centre) Intervention Comparator Tranexamic acid: 1g loading dose, 500 mg added to the Placebo: 0.9% saline (regimen as per TXA) CPB system p0rime, and a continuous infusion of 400 High dose aprotinin (Hammersmith protocol): 2 x 106 KIU aprotinin loading dose, 2 x 106 KIU added to the CPB system prime, and a continuous infusion of 5 x 105 KIU/h during CPB) Population characteristics Adult patients undergoing first-time, non-complex (one or two procedures) heart surgery (ie, low to moderate risk) with the use of CPB. Mean age 65 years; female 30%, BMI 26.5 kg/m² Length of follow-up Outcomes measured Hospitalisation period Primary: total postoperative blood loss and transfusion requirements. Secondary: in-hospital mortality, re-exploration, perioperative myocardial infarction; mediastinitis, renal failure, neurological complications, sepsis, length of ICU and hospital stay. INTERNAL VALIDITY Allocation Results Blinding analysis Treatment/measurement Follow-up (ITT) bias Opaque envelopes Double-blind. All Staff blinded to treatment 35/333 (10.5%) Staff measuring patients randomised prepared by outcomes unaware caretakers were allocation so not likely to be independent treatment or measurement excluded from of treatment blinded to analysis, mostly due statistician. assianment. medication bias. Standard statistical medication prepared allocation. to patients who did by independent methods used. not subsequently use anaesthesia CPB. assistant into syringes marked with patient number Overall quality assessment (descriptive) Good. Random treatment allocation, double-blind, not all patients included in analysis but reasonably large trial. **RESULTS** Intervention group Outcome Placebo Statistical significance N=99 TXA N=103 N=96 aprotinin Tranexamic acid (IV) P value n/N (%) n/N (%) 73/103 (70.9) PRBC transfusion 57/99 (57.6) 0.057 69/99 (69.7) 81/103 (78.6) Blood products transfusion 0.15 Median (IQR) Median (IQR) P value

Total units nDDCs transfused	1 0 (2 0)	2.0 (2.0)		0.038
Total units pRBCs transfused (units)	1.0 (2.0)	2.0 (3.0)		0.038
Total mediastinal chest tube loss	760 (540)	860 (740)		0.034
(mL)	700 (340)	000 (740)		0.034
(IIIL)	n/N (%)	n/N (%)		P value
In-hospital mortality	1/99 (1.0)	1/103 (1.0)		1.00
Re-operation for any reason	14/99 (14.1)	14/103 (13.6)		1.00 (post-hoc)
Re-operation due to surgical	3/99 (3.0)	3/103 (2.9)		1.00 (post-hoc)
bleeding	3/99 (3.0)	3/103 (2.9)		1.00 (<i>post-noc)</i>
Re-operation due to non-surgical	2/99 (2.0)	4/103 (3.9)		0.68 (post-hoc)
bleeding	2/99 (2.0)	4/103 (3.9)		0.06 (<i>pusi-nuc)</i>
Perioperative myocardial	0/99 (0)	8/103 (7.8)		0.004
infarction	0/99 (0)	0/103 (7.6)		0.004
Renal failure by Mangano ^a	3/99 (3.0)	3/103 (2.9)		1.00 (post-hoc)
Danal complication DIFLE				
Renal complication RIFLE	8/99 (8.1)	18/103 (17.5)		0.059 (post-hoc)
Stroke	1/99 (1.0)	1/103 (1.0)		1.00 (post-hoc)
	Mean ± SD	Mean ± SD		P value
Length of hospital stay (days)	9.4 ± 8.6	8.5 ± 7.4		0.43 (post-hoc)
Length of ICU stay (hours)	49.2 ± 89.6	60.1 ± 116.6		0.46 (post-hoc)
High-dose aprotinin (IV)	1	I		
	n/N (%)	n/N (%)		P value
PRBC transfusion	48/96 (50.0)	73/103 (70.9)		0.004
Blood products transfusion	59/96 (61.5)	81/103 (78.6)		0.009
	Median (IQR)	Median (IQR)		P value
Total units pRBCs transfused	0.5 (1.0)	2.0 (3.0)		<0.001
(units)				
Total mediastinal chest tube loss	546 (405)	860 (740)		<0.001
(mL)				
	n/N (%)	n/N (%)		P value
In-hospital mortality	2/96 (2.1)	1/103 (1.0)		0.61 (post-hoc)
Re-operation for any reason	5/96 (5.2)	14/103 (13.6)		0.054 (post-hoc)
Re-operation due to surgical	4/96 (4.2)	3/103 (2.9)		0.71 (post-hoc)
bleeding	, ,	, ,		,
Re-operation due to non-surgical	0/96 (0)	4/103 (3.9)		0.12 (post-hoc)
bleeding		, ,		7
Perioperative myocardial	1/96 (1.0)	8/103 (7.8)		0.023
infarction	,	,		
Renal failure by Mangano ^a	3/96 (3.1)	3/103 (2.9)		1.0 (post-hoc)
Renal complication RIFLE	10/96 (10.4)	18/103 (17.5)		0.011
Stroke	1/96 (1.0)	1/103 (1.0)		1.0 (post-hoc)
5510	Mean ± SD	Mean ± SD		P value
Length of hospital stay (days)	7.8 ± 6.7	8.5 ± 7.4		0.49 (post-hoc)
Length of ICU stay (hours)	55.4 ± 134.2	60.1 ± 116.6		0.79 (post-hoc)
Clinical importance	JU-1 1 101.Z	00.1 ± 110.0	Clinical releva	
Tranexamic acid (IV)			Cililical Televi	unoc
	nDDCc		1. Dationt roles	vant clinical outcome.
Transfusion incidence	pRBCs4: Range of estimates inclu	doc clinically	1. Pauent-reie	vani ciinicai oulconie.
	3	,		
	important effects, but also of with no or harmful effect	ompannie		
	Blood products	doc clinically		
	4: Range of estimates inclu			
	important effects, but also o	ompatible		
	with no or harmful effect			

Transfusion volume	pRBCs	1: Patient-relevant clinical outcome.
Transiusion volume	•	1. Patient-relevant clinical outcome.
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p=0.038)	
Blood loss	Total mediastinal chest-tube loss	1: Patient-relevant clinical outcome.
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p=0.034)	
Mortality	In hospital mortality	1: Patient-relevant clinical outcome.
· · · · · · · · · · · · · · · · · · ·	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
Museardial inferetion		1. Detient relevant clinical autoens
Myocardial infarction	Myocardial infarction	1: Patient-relevant clinical outcome.
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p=0.007)	
Re-operation	Any	1: Patient-relevant clinical outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
	Surgical bleeding	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect Non-surgical	
	bleeding	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
Renal failure	Renal failure	1: Patient-relevant clinical outcome.
Renarranure		1. Patient-reievant cimical outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
Renal complication	Renal complication	1: Patient-relevant clinical outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
Stroke	Stroke	1: Patient-relevant clinical outcome.
SHOKE		1. Fatient-relevant clinical outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
Length of hospital stay	with no or harmful effect Length of hospital stay	2: Predictive surrogate outcome.
Length of hospital stay		2: Predictive surrogate outcome.
Length of hospital stay	Length of hospital stay 4: Range of estimates includes clinically	2: Predictive surrogate outcome.
Length of hospital stay	Length of hospital stay 4: Range of estimates includes clinically important effects, but also compatible	2: Predictive surrogate outcome.
	Length of hospital stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	· ·
	Length of hospital stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Length of ICU stay	Predictive surrogate outcome. Predictive surrogate outcome.
Length of hospital stay Length of ICU stay	Length of hospital stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Length of ICU stay 4: Range of estimates includes clinically	· ·
	Length of hospital stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Length of ICU stay 4: Range of estimates includes clinically important effects, but also compatible	· ·
Length of ICU stay	Length of hospital stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Length of ICU stay 4: Range of estimates includes clinically	· ·
	Length of hospital stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Length of ICU stay 4: Range of estimates includes clinically important effects, but also compatible	· ·
Length of ICU stay	Length of hospital stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Length of ICU stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	· ·
Length of ICU stay High-dose aprotinin (IV)	Length of hospital stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Length of ICU stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect pRBCs	2: Predictive surrogate outcome.
Length of ICU stay High-dose aprotinin (IV)	Length of hospital stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Length of ICU stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect pRBCs 1: Clinically important benefit, confidence	2: Predictive surrogate outcome.
Length of ICU stay High-dose aprotinin (IV)	Length of hospital stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Length of ICU stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect pRBCs 1: Clinically important benefit, confidence limit does not include null value (p=0.004)	2: Predictive surrogate outcome.
Length of ICU stay High-dose aprotinin (IV)	Length of hospital stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Length of ICU stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect pRBCs 1: Clinically important benefit, confidence limit does not include null value (p=0.004) Blood products	2: Predictive surrogate outcome.
Length of ICU stay High-dose aprotinin (IV)	Length of hospital stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Length of ICU stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect pRBCs 1: Clinically important benefit, confidence limit does not include null value (p=0.004) Blood products 1: Clinically important benefit, confidence	2: Predictive surrogate outcome.
Length of ICU stay High-dose aprotinin (IV) Transfusion incidence	Length of hospital stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Length of ICU stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect pRBCs 1: Clinically important benefit, confidence limit does not include null value (p=0.004) Blood products 1: Clinically important benefit, confidence limit does not include null value (p=0.009)	2: Predictive surrogate outcome. 1: Patient-relevant clinical outcome.
Length of ICU stay High-dose aprotinin (IV)	Length of hospital stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Length of ICU stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect pRBCs 1: Clinically important benefit, confidence limit does not include null value (p=0.004) Blood products 1: Clinically important benefit, confidence	2: Predictive surrogate outcome.
Length of ICU stay High-dose aprotinin (IV) Transfusion incidence	Length of hospital stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Length of ICU stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect pRBCs 1: Clinically important benefit, confidence limit does not include null value (p=0.004) Blood products 1: Clinically important benefit, confidence limit does not include null value (p=0.009)	2: Predictive surrogate outcome. 1: Patient-relevant clinical outcome.

Blood loss	Total mediastinal chest-tube loss 1: Clinically important benefit, confidence limit does not include null value (p<0.001)	1: Patient-relevant clinical outcome.
Mortality	In hospital mortality 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Myocardial infarction	Myocardial infarction 1: Clinically important benefit, confidence limit does not include null value (p=0.023)	1: Patient-relevant clinical outcome.
Re-operation	Any 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Surgical bleeding 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Non-surgical bleeding 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Renal failure	Renal failure 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Renal complication	Renal complication 1: Clinically important benefit, confidence limit does not include null value (p=0.011)	1: Patient-relevant clinical outcome.
Stroke	Stroke 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Length of hospital stay	Length of hospital stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	2: Predictive surrogate outcome.
Length of ICU stay	Length of ICU stay 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	2: Predictive surrogate outcome.
EXTERNAL VALIDITY		
generalisable to this select sur	n adult patients undergoing first-time, non-comple gical population.	ex CPB surgery so likely to only be
Applicability Study conducted in a single co Comments	entre in the Netherlands so may be applicable to	the Australian/New Zealand setting.
Note: All post has calculations perform		

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: BMI, body mass index; CPB, cardiopulmonary bypass; FFP, fresh frozen plasma; ICU, intensive care unit; IQR, inter-quartile range; ITT, intention-to-treat; IV, intravenous; pRBCs, packed red blood cells; RCT, randomised controlled trial; SD, standard deviation; TXA, tranexamic acid.

a Mangano et al (2006) The risk associated with aprotinin in cardiac surgery. NEJM 354: 353-365.

STUDY DETAILS							
Citation							
Leijdekkers VJ, Vahl AC, Mackaay AJC, Huijgens PC, Rauwerda JA (2006) Aprotinin does not diminish blood loss in elective							
operations for infrarena	al abdomina	I aneurysm	is: a randomized	l, double	e-blind control	led trial. Ann Vas	sc Surg 20: 322-329.
Affiliation/Source of f	unds						
Department of Surgery, Onze Leve Vrouwe Gasthius, Amsterdam, The Netherlands; Department of Surgery, Meander							
Medical Center, Amers	foort, The N	letherlands	; Departments of	of Surge	ry and Haema	tology, Vrije Uni	versitiet Medical Center,
Amsterdam, The Netherlands.							
Study design		L	evel of evidence	е		Location/setti	ng
Double-blind RCT		II				Hospital	
Intervention					arator		
Aprotinin: Test dose of	500,000 KI	U, followed	by 2,000,000	Placeb	o: 0.9% salin	e (regimen as pe	er aprotinin)
KIU in 15 mins. During	surgery pat	tients receiv	ved a				·
continuous infusion of	500,000 KIL	J per hour,	to a maximum				
of 2,000,000 KIU.							
Population character							
							e or bifurcation prosthesis
for asymptomatic infrar	renal abdom	ninal aneury	ysm. Mean age				
Length of follow-up					mes measur		
Hospitalisation period						on requirements	s, postoperative
				compli	cations.		
INTERNAL VALIDITY							
Allocation	Results		Blinding anal	ysis		measurement	Follow-up (ITT)
					bias		
Randomised using	Standard :		Described as		No details p		All patients included in
standard	methods ເ	ısed.	double-blind. I		assess poss		analysis.
randomization list.			not to be open		treatment/m	easurement	
			until study incl		bias.		
			finalised. No fu				
			details provide	ed.			
Overall quality asses							
Fair. Random treatme	nt allocation	ı, double-bl	ind but not well	describe	ed, small trial.		
RESULTS		I					
Outcome			tion group		cebo	St	atistical significance
		N=16		N=1	19		
Aprotinin (IV)		Π				T =	
		Mean ± S			an ± SD		value
Mean total infusion (ml		7845 ± 4	888		5 ± 4776	0.9	
Mean pRBCs transfuse		4.1 ± 3.1			± 2.9	0.0	
Mean FFP transfused	(units)	0.5 ± 0.9			± 0.8	0.3	
Mean blood loss (mL)		2362 ± 1		_	6 ± 1370	0.8	
Mortality		1/16 (6.3)			9 (5.3)		00 (post-hoc)
Re-operation for bleed	ing	1/16 (6.3))	1/19	9 (5.3)		00 (post-hoc)
Clinical importance					C	linical relevanc	e
Aprotinin (IV)							

Transfusion volume	Mean total infusion (mL) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Mean pRBCs (units) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Mean FFP (units) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Blood loss	Mean blood loss (mL) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Mortality	In hospital mortality 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Reoperation for bleeding	In hospital mortality 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
EXTERNAL VALIDITY	·	
Generalisability		
	n adult patients undergoing elective surgery for in	nfra-renal abdominal aneurysm so likely to only
be generalisable to this select	surgical population.	
Applicability		
	lands so likely to be applicable to the Australian	/New Zealand setting.
Comments		

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: FFP, fresh frozen plasma; ITT, intention-to-treat; IV, intravenous; pRBCs, packed red blood cells; RCT, randomised controlled trial; SD, standard deviation.

STUDY DETAILS								
Citation								
	or MC (200	7\ Transvar	min anid and aris		conomi orton	, b		nroon oath o otudu
Maddali MM, Rajakuma Asian Cardiovascular a				nary co	onary artery	/ bypass surge	ery: a	prospective study.
Affiliation/Source of f								
Departments of Anesth		ardiothorac	ic Surgery, Roya	al Hospi	tal. Muscat.	Sultanate of C)man	
Funding not stated			.o o u. go. y,o y	л. г.оор.	tai, massat,	ounanate of o		
Study design			evel of evidence	:e		Location/s	settino	7
Double-blind RCT						Hospital (s		
Intervention		Comparator					301111 0)	
Tranexamic acid: 10 m	n/kn loading	n dose befo	re skin			aline (regimen	as ne	r TXΔ)
incision followed by a c				1 lacck	o. normai s	anne (regimen	us po	7 170 9
until commencement of								
from CPB.	. p. ota	. ov or our ar	to. ooparation					
Population characteri	istics							
Adult patients undergo		non-emera	ency CABG. Me	an age	58, female 3	32%, weiaht 65	ī ka.	
Length of follow-up	<u> </u>	9			mes measi		<u> </u>	
Hospitalisation period							ısion r	requirements, re-
								hemical data, operative
				data.	•	J 1		
INTERNAL VALIDITY								
Allocation	Results		Blinding anal	ysis	Treatmen	t/measureme	nt	Follow-up (ITT)
				,	bias			, , ,
Randomisation	Staff meas	suring	Double-blind.	With	Staff blind	ed to treatmen	t	All patients included in
based on computer-	outcomes	unaware	the exception	of the allocation so no		so not likely to	be	analysis.
generated code and	of treatme	nt	nurse who pre	pared treatment or measureme				
sequentially	assignmer	nt.	the treatment,					
numbered, sealed,	Standard s	statistical	in the operating	g				
opaque envelopes	methods u	sed.	room and pos					
opened by a nurse in			surgical unit w	ere				
the operating room			not aware of					
who prepared the			treatment					
infusions.			assignment.					
Overall quality assess								
Good. Random treatme	ent allocatio	n, double-b	olind, all patients	include	d in analysi	S		
RESULTS								
Outcome			tion group		nparator gr	oup	Stat	istical significance
		N=111		N=1	111			
Tranexamic acid (IV)								
		Mean ± S			an ± SD		Pva	
Total pRBCs transfuse		608.6 ± 2			.4 ± 292.1		0.00	
Total units FFP transfu		$0.72 \pm 1.$			± 2.4		<0.0)]
	platelets transfused 0.7 ± 1.9				± 2.3		NS	
Total drainage (mL)		633.0 ± 1	183.2		.9 ± 267.2		0.00	
		n/N (%)	- \		(%)		Pva	ilue
Re-operation due to ble	eeding	3/111 (2.			11 (2.7)		NS	
		Mean ± S			an ± SD		Pva	ilue
Postoperative haemog	Iobin	10.1 ± 1.	1	10.4	4 ± 1.3		NS	
(g/dL)					Т	<u> </u>		
Clinical importance						Clinical relev	ance	
Tranexamic acid (IV)								

Transfusion volume	 pRBCs 1: Clinically important benefit, confidence limit does not include null value (p=0.001) FFP 1: Clinically important benefit, confidence limit does not include null value (p<0.01) Plasma 4: Range of estimates includes clinically 	1: Patient-relevant clinical outcome.
	important effects, but also compatible with no or harmful effect	
Blood loss	Total drainage 1: Clinically important benefit, confidence limit does not include null value (p=0.001)	1: Patient-relevant clinical outcome.
Re-operation	Re-operation due to bleeding 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	1: Patient-relevant clinical outcome.
Haemoglobin	Postoperative haemoglobin 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect	2: Predictive surrogate outcome.
EXTERNAL VALIDITY		

Generalisability

Study conducted specifically in adult patients undergoing coronary artery bypass surgery so likely to only be generalisable to this select surgical population.

Applicability

Study conducted in a single centre in the Sultanate of Oman so may not be completely applicable to the Australian/New Zealand setting.

Comments

The authors note that as this was the only tertiary cardiac centre in the country, the surgeons involved in the study were at various stages of expertise and training and this might have led to higher than expected blood drainage and transfusion requirements.

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: BMI, body mass index; CABG, coronary arterial bypass graft; CPB, cardiopulmonary bypass; FFP, fresh frozen plasma; ICU, intensive care unit; ITT, intention-to-treat; IV, intravenous; NS, not significant; pRBCs, packed red blood cells; RCT, randomised controlled trial; SD, standard deviation; TXA, tranexamic acid.

^a Article states units rather than mL, but amounts reported correspond to mL.

STUDY DETAILS								
Citation								
								ring and after cesarean
section: a randomized		lled prospe	ective study. The	e Journa	I of Obstetrics	and Gyneco	ology (of India 57(3): 227-230.
Affiliation/Source of f								
Department of Obstetri			edical College a	and SSC	G Hospital, Gu	jarat, India.		
No pharmaceutical con	npany fundi							
Study design			evel of eviden	ce		Location/s		
Pseudo-RCT				1 0		Hospital (s	ingle-	centre)
Intervention	10 \ \ \	/ F!-			oarator			
Tranexamic acid: 1 g (i minutes prior to incision		over 5 mir	iules 20	No TX	λA			
Population character								
Full term primiparas, m		th sinalatar	n nrognancy: mg	an and	24 mean w	aight =50 kg		
Length of follow-up	iuitiparas Wi	ui siriyicili	i pregnancy, me		mes measur			
Hospitalisation period							sians	haemoglobin.
INTERNAL VALIDITY				1 03(-)	a turr riuorriu	mayo, vitai	oigi io,	naomogiobin.
Allocation	Results		Blinding ana	lvsis	Treatment/	measureme	nt	Follow-up (ITT)
				.,	bias			т спот ар (т т)
Pseudorandomised	Statistical	methods	Open-label		Objective m	easurements	of	All patients included in
based on odd/even	not descri	bed.			volume and	weight used	for	analysis.
numbers.						haemorrhag		
						ot be subject	to	
					bias.			
Overall quality assess						.1		
Poor. Pseudo-random	isation usin	g oaa/even	numbers (not v	ery seci	ure), open-iab	el.		
RESULTS		Intonion	tion avoirs	Co			Ctot	intinal ciamificanas
Outcome		N=50	tion group	Comparator group N=50		Statistical significance		
Tranexamic acid (IV)		14=30		11=	JU			
Post-partum hae	morrhage							
z. Tost partam nac	morriage	Mean ± 3	SD	Me	an ± SD		P va	nlue
Post-partum haemorrh	age	299.21 ±			l l		0.05	
(placental delivery to e							2.00	
surgery; mL)								
Post-partum haemorrh	age (end	75.71 ± 2	20.02	133.03 ± 14.68		0.001		1
of surgery to 2 hours p	ost							
partum; mL)								
Post-partum haemorrh		374.92 ±	51.46	472	2.79 ± 43.54		0.00	3
(placental delivery to 2	hours							
post-partum; mL)	La la Sa	ND		N.I.			NIC	
Postoperative haemog	Iobin	NR		NR			NS	
(g/dL)		n/N /0/\		/N	(0/)		D	dua
Thrombooic		n/N (%) 0/50		0/5	(%)		P va NA	iiue
Thrombosis Clinical importance		0/30		0/5		linical relev		
Tranexamic acid (IV)					10	in iicai reiev	ance	
Transparinc acid (IV)								

Blood loss	Placental delivery to end of surgery	1: Patient-relevant clinical outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
	End of surgery to 2 hours post-partum	
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p=0.001)	
	Placental delivery to 2 hours post partum	
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p=0.003)	
Thrombosis	Thrombosis	1: Patient-relevant clinical outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
Haemoglobin	Postoperative haemoglobin	2: Predictive surrogate outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
EXTERNAL VALIDITY		
Generalisability		
	ally in women delivering full-term, single pregnancies b	by lower segment caesarean section so
generalisable only to this	specific population.	
Applicability		
	gle centre in India so may not be completely applicable	e to the Australian/New Zealand setting.
Comments		
There was no significant of	difference in Apgar scores between the TXA and place	ebo arms (p=0.5).

There was no significant difference in Apgar scores between the TXA and placebo arms (p=0.5).

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: ICU, intensive care unit; ITT, intention-to-treat; IV, intravenous; NA, not applicable; NR, not reported; RCT, randomised controlled trial; SD, standard deviation; TXA, tranexamic acid.

STUDY DETAILS								
Citation								
	M Madani 4	shi M (200	7) Doos transva	mio onio	l raduas blass	l loce in off n	ımn	coronary artery bypass?
Asian cardiovascular a				IIIIC acid	reduce blood	1 1088 III 011-pt	лпр с	coronary artery bypass?
Affiliation/Source of f	unds							
		hariati Hos _l	oital, Research I	Departn	nent, Tehran H	Heart Centre,	Tehra	an University of Medical
Sciences, Tehran, Iran								
Study design			evel of eviden	ce		Location/se		
RCT							ngle-d	centre)
Intervention					arator			
Tranexamic acid: loadi				Placel	oo – saline sol	lution (same r	egim	en as TXA)
the beginning of surger								
infusion of heparin, at t	he end of si	urgery and	after					
protamine infusion.	. ,,							
Population characteristics Undergoing primary CABG, aged ≤ 70 years, LVEF ≥ 35%, BMI ≤ 25, no aspirin for 7 days prior, no preoperative heparin						0 1 2		
	ABG, aged ≤	≤ /0 years,	LVEF ≥ 35%, E	3MI ≤ 25	o, no aspirin fo	or / days prior	r, no l	preoperative neparin
infusion.				0.1.				
Length of follow-up					mes measur		1 -1-	
Hospitalisation period						0	od cr	nemistry parameters,
INTERNAL MALIRITM				morbio	dity outcomes	, mortality.		
INTERNAL VALIDITY	Dogulto		Diadiaa sast	i.o	Trootmoonth	** * * * * * * * * * * * * * * * * * * *		Follow up /ITT\
Allocation	Results		Blinding anal		bias	measuremen		Follow-up (ITT)
Randomly allocated.	Outcomes	1	Double-blind.	Staff Staff unaware of treat				
No further details	assessed	,	in the operatin				t be	analysis.
provided.	clinical sta		room and ICU			easurement		
Drug prepared by	aware of t		unaware of	bias.				
pharmacy staff not	assignmer		treatment					
involved in study into	Standard		assignment.					
coded infusion	methods ι	ısed.						
syringes.	. / 1	\						
Overall quality assess			Control of the con-	. 1 1 .				
Good. Randomised, d	ouble-blind,	all subjects	s included in ana	alysis.				
RESULTS		T		10		T	0	
Outcome			tion group		mparator gro	up	Stat	istical significance
T		N=33		N=3	33			
Tranexamic acid (IV)		m/N1 (0/)		/NI	(0/)		Dva	dua
Transfusion incidence	/whole	n/N (%) 5/33 (15	2)		(%)		P va 0.07	
Transfusion incidence		5/33 (15	2)	8/3.	3 (24.2)		0.07	
blood or pRBC transfus		0/22 (0)		6/33 (18.2)			0.05	
Transfusion incidence	(FFP	0/33 (0)		6/3.	3 (18.2)		0.05	
transfused)	/	0/22 (0)		0/2:	2 (0)		N.O.	
Transfusion incidence	(higieieiz	0/33 (0)		0/3.	3 (0)		NA	
transfused) Transfusion incidence (total 5/33 (15.2)		2)	10/	22 (26 1)		0.00	(nost hos)	
Transfusion incidence (total 5/3: patients transfused)		0/33 (13	4)	12/33 (36.4)			0.09 (post-hoc)	
patients transiuseu)		Moan		Mea	an		P va	luα
Whole blood or nDDC	transfusad							
	แสกรเนรษน	0.40		0.9	†		0.00	' I
per patient (units)		Mean + 9	SD	Mo	an + SD		D 1/2	nlue
Postonerative blood los	ss ∩-2 hr		<i>,</i> ,				P value <0.001	
•	55 U Z III	70 ± 23		100	· _ 01		\U.U	
Whole blood or pRBC per patient (units) Postoperative blood los (mL)		Mean 0.46 Mean ± SD 90 ± 25		0.94 Mean ± SD 180 ± 37			0.00 P va	1 Ilue

Postoperative blood loss 2-6 hr	190 ± 41	290 ± 78		0.001	
(mL) Total postoperative blood loss	320 ± 38	480 ± 75		0.001	
(mL)	320 ± 30	400 ± 73		0.001	
	n/N (%)	n/N (%)		P value	
Mortality	0/33	0/33		NA	
Surgical re-exploration for bleeding	0/33 (0)	1/33 (3.0)		>0.05	
Myocardial infarction	0/33	0/33		NA	
Renal dysfunction (creatinine > 2 mg/dL)	0/33 (0)	1/33 (3.0)		>0.05	
Hospital length of stay (days)	4.8 ± 0.4	4.8 ± 0.9		0.09	
ICU length of stay (hours)	10 ± 1.8	12 ± 3.2		<0.05	
First postoperative day	10.6 ± 1.9	10.4 ± 0.8		>0.05	
haemoglobin (g/dL)					
Clinical importance			Clinical relev	ance	
Tranexamic acid (IV)					
Transfusion incidence Transfusion volume Blood loss	Whole blood or pRBC 4: Range of estimates inclu important effects, but also of with no or harmful effect FFP 4: Range of estimates inclu important effects, but also of with no or harmful effect (psplatelets) 1: Clinically important benellimit does not include null via Any blood products 4: Range of estimates inclu important effects, but also of with no or harmful effect Whole blood or pRBC 1: Clinically important benellimit does not include null via Postoperative blood loss (O) 1: Clinically important benellimit does not include null via Postoperative blood loss (O) 1: Clinically important benellimit does not include null via Postoperative blood loss (O) 1: Clinically important benellimit does not include null via Postoperative blood loss (O) 1: Clinically important benellimit does not include null via Postoperative blood loss (O) 1: Clinically important benellimit does not include null via Postoperative blood loss (O) 1: Clinically important benellimit does not include null via Postoperative blood loss (O) 1: Clinically important benellimit does not include null via Postoperative blood loss (O) 1: Clinically important benellimit does not include null via Postoperative blood loss (O) 1: Clinically important benellimit does not include null via Postoperative blood loss (O) 1: Clinically important benellimit does not include null via Postoperative blood loss (O) 1: Clinically important benellimit does not include null via Postoperative blood loss (O)	des clinically compatible =0.05) fit, confidence alue (p<0.05) des clinically compatible fit, confidence alue (p=0.001) -2 hr) fit, confidence alue (p<0.01) -6 hr) fit, confidence alue (p<0.01)	1: Patient-relevant clinical outcome. 1: Patient-relevant clinical outcome. 1: Patient-relevant clinical outcome.		
Mortality	limit does not include null vi In-hospital mortality No deaths in either treatme		1: Patient-rele	vant clinical outcome.	
Re-operation	Re-operation for bleeding 4: Range of estimates inclu important effects, but also o with no or harmful effect	des clinically	1: Patient-rele	vant clinical outcome.	
Myocardial infarction	Thrombosis No MI in either treatment gr	oup	1: Patient-rele	vant clinical outcome.	
Renal dysfunction	Creatinine > 2 mg/dL 4: Range of estimates incluimportant effects, but also dwith no or harmful effect	des clinically	1: Patient-rele	vant clinical outcome.	

Hospital length of stay	Hospital length of stay	2: Predictive surrogate outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
ICU length of stay	ICU length of stay	2: Predictive surrogate outcome.
-	1: Clinically important <i>benefit</i> , confidence	-
	limit does not include null value (p<0.05)	
Haemoglobin	First day postoperative haemoglobin	2: Predictive surrogate outcome.
· ·	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
EXTERNAL VALIDITY		
Generalisability		
Study conducted specifically	in patients no older than 70 years who were under	going primary CABG so likely to be
generalisable only to this sele	ect group of patients.	
Applicability		
Study conducted in a single of	centre in Iran so may not be completely applicable	to the Australian/New Zealand setting.
Comments		

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: BMI, body mass index; CABG, coronary artery bypass graft; FFP, fresh frozen plasma; ICU, intensive care unit; ITT, intention-to-treat; IV, intravenous; LVEF, left ventricle ejection fraction; MI, myocardial infarction; NA, not applicable; pRBCs, packed red blood cells; RCT, randomised controlled trial; SD, standard deviation; TXA, tranexamic acid.

STUDY DETAILS									
Citation									
Mehraien A, Ghafari A	Mohamma	di SS (200	19) Effect of tonic:	al anrot	inin on ear	ly nostonerative	hleer	ling and ICLL stay after	
coronary artery bypass								and 100 stay after	
Affiliation/Source of t		1103. 1 4113	tan Journal of Bi	ological	001011003	12(10): 010 010			
Department of Anaesth		ehran I Iniv	versity of Medical	Scienc	es Dr Sha	riati Hospital Te	hran	Iran	
Study design	icsiology, i		Level of evidence		CS, DI SIIC	Location/s			
Double-blind RCT						Hospital	cun	9	
Intervention			<u> </u>	Comp	arator	Tiospilai			
Aprotinin: 500,000 KIU (50 mL) applied topica			ally to tho			regimen as per	anroti	nin)	
heart, pericardium and mediastinum prior to				riacei	o. saine (regimen as per	aproti	11111)	
Population character		iii piioi to .	sterria ciosure.						
Adult patients undergo		coronary	artory hypacs ar	aft cura	orv: ASA r	hycical status II	or III.	agod 50 70 years	
Mean age 58 years; fe		e coronary	artery bypass gr	ait sury	ciy, ASA p	iriysicai status ii	OI III,	aged 50-70 years.	
Length of follow-up	maic 32 70.			Outco	mes mea	sured			
Hospitalisation period						fusion requireme	nts	ICII stav	
INTERNAL VALIDITY				Dioou	1000, 110113	asion requirem	JIII.J _I	ioo siay.	
Allocation	Results		Blinding anal	vsis	Treatme	nt/measuremer	nt	Follow-up (ITT)	
	ounto		g undi	,	bias		- •	« P (///)	
Randomised using	Standard	statistical	Described as			nce that there m	nay	All patients included in	
computer-generated	methods used.		double-blind. (Coded	be treatn	be treatment/measurement bias.		analysis.	
codes			syringes prepa	ired	bias.				
			by independent						
		anaesthetist.							
			Outcomes determined by						
			blinded						
0 11 12		\	anaesthetist.						
Overall quality asses			111 1 11 11 11			•			
Good. Random treatm	ient allocation	on, aouble-	-biina, aii patients	s include	ed in anaiy	SIS.			
RESULTS		Intonion	tion group	Dla			Ctot	ictical cianificance	
Outcome		Intervention group N=64			Placebo N=64		Stat	istical significance	
Aprotinin (topical)		IN=04		IN=C)4				
Aprodillili (topical)		Mean ±	SD	Mos	an ± SD		P va	aluo	
Mean pRBCs transfuse	nd (units)	0.5 ± 0.7			± 1.0		0.00		
24-hour chest tube los		451 ± 21			± 269	0.00			
ICU length of stay (hou		48.8 ± 13			± 207 4 ± 16.6		0.003		
Clinical importance	113/	TU.U 1	5.0	07.4	1 ⊥ 10.0	Clinical releva		' I	
Aprotinin (topical)						Jiiiiicai reievi	ai ice		
Transfusion volume		Mean nF	RBCs (units)			1. Patient-rele	vant c	clinical outcome.	
Transiusion volume			ally important <i>ber</i>	nefit coi	nfidence	1. T diloni Tolo	varit	minear outcome.	
			s not include null						
Blood loss			chest tube loss (i		z 0.000j	1: Patient-relevant clinical outcome.		clinical outcome.	
			ally important <i>ber</i>		nfidence			,	
			s not include null						
ICU length of stay			th of stay (hours)		, ,	1: Patient-rele	vant c	clinical outcome.	
J ···J			ally important ber		nfidence				
			s not include null						
EXTERNAL VALIDITY	<i>'</i>								
EXTERNAL VALIDITI					_				
Generalisability									
	fically in ad	ult patients	undergoing first-	time C	ABG so like	ely to be genera	lisable	e to this surgical	

Applicability

Study conducted in Iran so may applicable to the Australian/New Zealand setting

Comments

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: CABG, coronary artery bypass graft; FFP, fresh frozen plasma; ITT, intention-to-treat; IV, intravenous; pRBCs, packed red blood cells; RCT, randomised controlled trial; SD, standard deviation.

STUDY DETAILS							
Citation							
	بمتاد ۵ (۲۵۸۸	O) Aprotinin	for notionts over	ocod to	alanidaaral h	oforo off nump oo	ranary hynass Asian
Nurözler F, Kutlu T, Kü Cardiovascular and Th				osea to	ciopidogrei b	erore on-pump cor	onary bypass. Asian
Affiliation/Source of f		113 10. 403-	407.				
Division of Cardiovasco		, Central H	Insnital Izmir Tı	ırkev			
Study design	ulai Julyery		evel of evidence			Location/setting	n
Double-blind RCT				,,,		Hospital	9
Intervention		"		Comp	arator	Поэрна	
Aprotinin: loading dose	1 million K	II I followed	hy 0.5 million			imen as per aproti	nin)
KIU per hour until the 6			by 0.5 million	1 lacck	o. same (reg	inien as per aprou	11111)
Population character		<i>.</i> ,					
Adult patients who had		lopidoarel v	vithin 5 days of s	uraerv	ındergoing of	f-pump coronary a	rtery bypass surgery.
Mean age 64 years; 27				a. go. j	and going on	. pap oo.oa.j a	
Length of follow-up		•	<i>y</i>	Outco	mes measur	ed	
In hospital						ce; transfusion vol	ume: blood loss:
I					ity; morbidity.	,	,
INTERNAL VALIDITY							
Allocation	Results		Blinding anal	ysis	Treatment/r	measurement	Follow-up (ITT)
			, and the second		bias		,
List of random	Standard	statistical	Described as		No evidence	that there may	All subjects included
treatment codes	methods u	used.	double-blind.		be treatmen	t/measurement	in analysis.
generated by a			Blinding of ICU		bias.		
biostatistician using			described as b				
a block design.			"conducted in				
		appropriate wa	ау".				
Overall quality assess							
Fair. Randomised, des	cribed as do	ouble-blind,	, similar at basel	ine; all p	atients includ	led; small study.	
RESULTS						1	
Outcome			tion group		cebo	Stat	istical significance
Λ Δ ! (I) ()	N=25			N=2	.6		
Aprotinin (IV)		/N1 (O/)		//\	(0/)	D.,,	N
Transfusion incidence	(DDCs)	n/N (%)))		<u>(%)</u>	P va	
Transfusion incidence	<u>, , , , , , , , , , , , , , , , , , , </u>	17/25 (68	•		26 (88) 26 (53)	0.01	
products)	(DIOOU	7/25 (28)		14/2	20 (33)	0.00	12
products)		Mean ± S	en en	Mod	n ± SD	P va	aluo.
Transfusion volume (pl	DDCc:	1.7 ± 1.4			± 1.8	0.01	
units)	NDC3,	1.7 ± 1.4		2.7	± 1.0	0.01	4
Transfusion volume (pl	atelets:	0.4 ± 0.6		2.3	± 1.2	0.00	12
units)	atcicis,	0.4 ± 0.0		2.5	± 1.2	0.00	72
Transfusion volume (F	FP: units)	0.6 ± 0.3		1 4	± 0.6	0.00	08
Drainage (mL/24 hr) 423 ± 178			748 ± 212		0.00		
=: aago (IIIE/E 1 III)		n/N (%)	-		(%)	P va	
Re-operation		0/25 (0)			(7.7)	0.15	
In-hospital myocardial	infarction	0/25 (0)			5 (0)	NA NA	
In-hospital stroke		1/25 (4.0))		5 (0)	0.31	7
		Mean ± S			n ± SD	P va	
ICU length of stay (hr)		28 ± 11			± 10	0.15	
Hospital length of stay	(days)	5.3 ± 1.6			± 1.4	0.66	
	<u> </u>					Partie at materials	
Clinical importance					C	linical relevance	

Transfusion incidence	Red blood cell	1: Patient-relevant clinical outcome.
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p=0.014)	
	Blood products	
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p=0.002)	
Transfusion volume (units)	Mean pRBCs	1: Patient-relevant clinical outcome.
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p=0.014)	
	Mean platelets	
	1: Clinically important <i>benefit</i> , confidence limit does not include null value (p=0.002)	
	Mean FFP	
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p=0.008)	
Blood loss	Drainage (mL/24 hr)	1: Patient-relevant clinical outcome.
2.004.000	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p=0.005)	
Re-operation	Re-operation	1: Patient-relevant clinical outcome.
·	4: Range of estimates includes clinically	
	important point estimate but also	
	compatible with no effect, or a harmful	
	effect	
Myocardial infarction	Myocardial infarction	1: Patient-relevant clinical outcome.
Charles	No events in either treatment arm.	4. Dalbad adams talbahada ataun
Stroke	Stroke	1: Patient-relevant clinical outcome.
	4: Range of estimates includes clinically important point estimate but also	
	compatible with no effect, or a harmful	
	effect	
Length of stay	ICU	1: Patient-relevant clinical outcome.
	4: Range of estimates includes clinically	
	important point estimate but also	
	compatible with no effect, or a harmful	
	effect	
	Hospital	
	4: Range of estimates includes clinically	
	important point estimate but also	
	compatible with no effect, or a harmful	
EVTEDNIAL VALIDITY	effect	
EXTERNAL VALIDITY Conoralisability		
Generalisability Study conducted specifically in a	dult patients undergoing off-pump coronary art	ory hypass who had boon on clonidogral
	to be generalisable to this surgical population.	ery bypass who had been on clopidogref
Applicability	to be generalisable to this surgical population.	
	ay be applicable to the Australian/New Zealand	l setting.
Comments	2) we approad to the restanding town Localine	

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: BMI, body mass index; FFP, fresh frozen plasma; ICU, intensive care unit; ITT, intention-to-treat; IV, intravenous; NS, not significant; RBC, red blood cell; RCT, randomised controlled trial; SD, standard deviation.

CTUDY DETAIL C								
STUDY DETAILS								
Citation	A (2007) D							
								ransfusion requirements I Iranica 45(6): 437-442.
Affiliation/Source of f		pective rai	idomised double	-billiu s	iddy iii o'r pat	ienis. Acia ivi	icuica	i ilaliica 45(0). 457-442.
Department of Anesthe		ariati Hosni	tal School of M	edicine	Medical Scien	res Univers	ity of	Tehran Iran
Funding not reported.	Slology, 511	anati mospi	tai, Scribbi bi ivi	culcille	Medical Scien	ices, Univers	ity Oi	Terilari, irari.
Study design			evel of evidence	20		Location/s	ottino	n
RCT				,,,		Hospital (si		
Intervention		!!		Comp	arator	1 103pitai (3i	rigic-c	senii e)
Tranexamic acid: single	holus dos	if 15 ma/k	a hoforo		oo – saline sol	ution (samo	rogim	on as TVA)
surgical incision.	z bolus uost	on 15 mg/N	y before	Tacei	00 – 3aii ie 30i	ution (same	regiiii	ciras inny
Population characteri	etice							
		a of the hin	nacassitation h	in surac	ary (Avtracansi	ılar fracturos	rogui	iring plating and nailing
and intracapsular fracti	ıracıul ırac ranılirir	a hamiarth	ronlastv) Moan	ane1	8 famala 10%	ulai Iraciules L RMI - 22	requ	ining plating and nationg
Length of follow-up	ires requirii	ig ricilliai ii i	ropiasty). Meari		mes measur			
Hospitalisation period							tran	sfusion volume,
Hospitalisation penou					ity, haematolo		i, liaii	Siusion volume,
INTERNAL VALIDITY				morta	ity, naematoic	99		
Allocation	Results		Blinding anal	vcic	Troatmont/r	neasuremer	nt.	Follow-up (ITT)
Allocation	Nesuits		Difficility arial	ysis	bias	ileasureillei	ıı	i ollow-up (i i i)
Randomly allocated	Outcomes		Double-blind.	Staff		re of treatme	nt	All patients included in
using a random	assessed	d by in the operatin					analysis.	
number technique.	clinical sta					l DC		
Drug prepared by	aware of t			unaware of		casarcincin		
pharmacy staff not	assignme		treatment		bias.			
involved in study into	Standard:		assignment.					
coded infusion	methods u		uosigiimoni.					
syringes.								
Overall quality assess	sment (des	criptive)						
Good. Randomised, de			s included in ana	alvsis.				
RESULTS	20010 200104	an oanjoon	,					
Outcome		Intervent	tion group	Cor	nparator gro	ın	Stat	istical significance
Outcome		N=32	ion group	N=3		- P	Otat	istical significance
Tranexamic acid (IV)		14 02		1				
Transfusion incid	dence							
o. Transiasion mon	401100	n/N (%)		n/N	(%)		P va	alue
Transfusion incidence	/whole	12/32 (37	' 5)		35 (57.1)		0.04	
blood or pRBC transfus		12/32 (37	.5)	207	33 (37.1)		0.01	
Transfusion incidence		1/32 (3.1)	1	0/3	5 (0)		NS	
transfused)	(1.1.	1/32 (3.1)	,	0/3	3 (0)		NO	
Transfusion incidence	nlatelets	0/33 (0)		0/3	3 (0)		NA	
transfused)	piatolots	0/33 (0)		0/3	3 (0)		14/1	
Transfusion incidence	'total	12/32 (37	' 5)	20/	35 (57.1)		0.04	
patients transfused)	(total	12/32 (37	.5)	207	33 (37.1)		0.01	
pationio tidribiusou)		Mean		Mea	an		P va	alue
Whole blood or pRBC t	ransfused	1.25		1.9			0.00	
per patient (units)	. anduocu	1.20		1.7	•		0.00	• •
por pationit (dinito)		Mean ± S	SD	Me	an ± SD		P va	alue
Perioperative blood los	s (ml)	652 ± 22			8 ± 372		0.00	
Postoperative blood los		111 ± 76			± 100		0.39	
(mL)		± ,0		''	_ 100		0.07	
···-/		J						

Destaurant a blandlag Oba	100 70	04/ 110		0.00
Postoperative blood loss 2 hr (mL)	192 ± 78	246 ± 113		0.28
Postoperative blood loss 5 hr (mL)	255 ± 59	323 ± 54		0.31
Postoperative blood loss 12 hr (mL)	296 ± 40	375 ± 30		0.20
Postoperative blood loss 24 hr (mL)	300 ± 54	390 ± 65		0.11
Total blood loss (mL)	960 ± 284	1484 ± 374		0.001
	n/N (%)	n/N (%)		P value
Mortality	0/32 (0)	1/35 (2.9)		NR
•	Mean ± SD	Mean ± SD		P value
Hospital length of stay (days)	4.3 ± 1.6	5.8 ± 1.5		<0.05
Postoperative day haemoglobin (g/dL)	10.1 ± 1.4	8.9 ± 2.1		<0.05
Clinical importance		Clinical relevance		
Tranexamic acid (IV)				
Transfusion incidence	1: Clinically important benefit, confidence limit does not include null value (p=0.04) FFP 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect (p<0.05) Platelets No patients in either group transfused with platelets Any blood products 1: Clinically important benefit, confidence		1: Patient-rele	evant clinical outcome.
Transfusion volume	limit does not include null value (p=0.04) Whole blood or pRBC 1: Clinically important benefit, confidence limit does not include null value (p=0.001)		1: Patient-rele	vant clinical outcome.

DI II	10 11 11	1 4 B 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
Blood loss	Perioperative blood loss	1: Patient-relevant clinical outcome.
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p=0.003)	
	Postoperative blood loss (1 hr)	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect Postoperative	
	blood loss (2 hr)	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect <i>Postoperative</i>	
	blood loss (5 hr)	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
	Postoperative blood loss (12 hr)	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
	Postoperative blood loss (24 hr)	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
	Total blood loss	
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p=0.001)	
Mortality	In-hospital mortality	1: Patient-relevant clinical outcome.
Wortality	4: Range of estimates includes clinically	1. I attent-relevant clinical outcome.
	important effects, but also compatible	
	with no or harmful effect	
Hospital length of stay	Hospital length of stay	1: Patient-relevant clinical outcome.
Hospital length of stay	1: Clinically important <i>benefit</i> , confidence	1. Falletit-felevarit cililical outcome.
	limit does not include null value (p<0.05)	
Haemoglobin		1: Patient-relevant clinical outcome.
паеттоуюыт	First day postoperative haemoglobin	1. Patient-relevant clinical outcome.
	1: Clinically important <i>benefit</i> , confidence limit does not include null value (p<0.05)	
EXTERNAL VALIDITY	IIIIIII does not include nuii value (p<0.05)	
Generalisability		
3	notionto undorgoina colocted hin curacrice for n	articular his fractures as likely to be
	patients undergoing selected hip surgeries for p	articular nip fractures so likely to be
generalisable only to this selec	a group or patients.	
Applicability		
	entre in Iran so may not be completely applicable	to the Australian/New Zealand setting.
Comments		
	difference in thrombotic complications, pulmona	ry dystunction and neurological deficits
however these are not quantific	ed.	

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: BMI, body mass index; FFP, fresh frozen plasma; ICU, intensive care unit; ITT, intention-to-treat; IV, intravenous; NA, not applicable; NR, not reported; NS, not significant; pRBCs, packed red blood cells; RCT, randomised controlled trial; SD, standard deviation; TXA, tranexamic acid.

STUDY DETAILS								
Citation								
	A Dalii M	ot al (2000) Efficacy of tran	ovomio	acid in rac	lucina blood loc	o ofto	r cesarean section. The
Journal of maternal-Fe					aciu iii iec	aucing blood los	ss arte	i cesarean section. The
Affiliation/Source of f		natai woak	51110 22(1): 72 70	,				
Department of Obstetri		ecology. St	nahid Sedughi H	lospital.	Shahid Se	eduahi Universit	v of M	ledical Sciences and
Health Services, Yazd,		ooologj, ol	iama oodagiii i	iospitai,	Oriania oc	adgiii diiivoisii	.j 01 111	iodiodi Golorioos diid
Not supported by a pha		I company.						
Study design			evel of evidence	ce.		Location/s	settin	n
RCT						Hospital (s		
Intervention			<u> </u>	Comp	arator	Trospital (c	migio .	001111 0)
Tranexamic acid: 1 g/1	0 mL IV slo	wlv infused	over 5			ucose dextrose	solutio	on (same regimen as
minutes, 10 minutes pr				TXA)	3			(** * * * * * * * * * * * * * * * * * *
Population character								
Women at full-term with		regnancy.	Mean age ~27 v	ears.				
Length of follow-up		<u> </u>	<u> </u>		mes meas	sured		
Hospitalisation period						natology, vital si	igns, tl	hrombosis.
INTERNAL VALIDITY						<i>JJ</i> ,	J - 1 - 1	
Allocation	Results		Blinding anal	ysis		nt/measureme	nt	Follow-up (ITT)
Pseudorandomised	Statistical	mothodo	Open-label		Objective	e measurement	c of	All patients included in
based on odd/even	not descri		Open-label					analysis.
numbers.	HOLUCSCII	ucu.		volume and weight used for post-partum haemorrhage			analysis.	
Humbers.						d not be subject		
					bias.		10	
Overall quality assess	sment (des	criptive)	I		Dias.			I
Poor. Pseudo-random			numbers (not ve	erv seci	ıre), open-	label.		
RESULTS		J 222/01/011		, 5000	3/1 3POIT			
Outcome		Interven	tion group	Coi	nparator	aroup	Stat	istical significance
	N=45				15 15	J. 9 4 P	0.0.	
Tranexamic acid (IV)								
,		Mean ± SD		Mean ± SD		P value		
Blood loss up to 2 hour	S	28.0 ± 5.		37.1 ± 9.0			<0.001	
postoperative (mL)								
		n/N (%)	V (%)		n/N (%)		P value	
Thrombosis		0/45 (0)			0/45 (0)		NA	
	Mean ± SD N		Mea	Mean ± SD		P va	alue	
Postoperative haemog	lobin	12.6 ± 1.	12.6 ± 1.3		11.7 ± 1.1		<0.01	
(g/dL)								
Change in haemoglobi		-0.1 ± 0.6	<u></u>	-2.5 ± 0.8			<0.0	001
preoperative to postop	erative							
(g/dL)								
Clinical importance						Clinical relev	ance	
Tranexamic acid (IV)								
Blood loss			r post-op		·	1: Patient-rele	evant c	clinical outcome.
			lly important be					
			not include nul	l value (p<0.001)			
Thrombosis		Thrombo				1: Patient-rele	evant c	clinical outcome.
			s in either treatn					
Haemoglobin			ostoperative ha			2: Predictive s	surrog	ate outcome.
			Illy important be					
EVEDNAL VALIETY	,	limit does	not include nul	ı value (p<0.001)			
EXTERNAL VALIDITY	•							

Generalisability

Study conducted specifically in women undergoing caesarean section so likely to be generalisable only to this select group of patients.

Applicability

Study conducted in a single centre in Iran so may not be completely applicable to the Australian/New Zealand setting.

Comments

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: BMI, body mass index; FFP, fresh frozen plasma; ICU, intensive care unit; ITT, intention-to-treat; IV, intravenous; NA, not applicable; pRBCs, packed red blood cells; RCT, randomised controlled trial; SD, standard deviation; TXA, tranexamic acid.

STUDY DETAILS Citation Taghaddomi RJ, Mirzaee A, Attar AS et al (2009) Tranexamic acid reduces blood loss in off-pump coronary artery bypass surgery. Journal of Cardiothoracic and Vascular Anaesthesia 23(3): 312-315. Affiliation/Source of funds Departments of Anesthesiology and Haematology, Ghaem Hospital, Mashhad University of Medical Sciences, Mashad, Iran; Department of Cardiosurgery, Chaem Hospital, Mashad University of Medial Sciences, Mashad, Iran. Funding not stated. Level of evidence Study design Location/setting **RCT** Hospital (single-centre) Ш Intervention Comparator Tranexamic acid: 1 g given 20 minutes prior to skin incision Placebo – saline solution (same regimen as TXA) and 400 mg/h during the entire surgical procedure. Population characteristics Patients scheduled for off-pump CABG. Mean age ~ 62, female 28%, weight 75 kg. No statistically difference in any demographic or perioperative variables between treatment arms. Outcomes measured Length of follow-up Hospitalisation period Blood loss, transfusion, haematology, morbidity. INTERNAL VALIDITY Allocation Results Blinding analysis Treatment/measurement Follow-up (ITT) bias Random allocation Assessed by staff Double-blind. Double-blind so 108 patients enrolled; 8 not analysed (7.4%; using envelopes. not aware of Operating room treatment/measurement bias staff and ICU staff 4 converted to on-Independent treatment should not be an issue. anaesthesiologist assignment. unaware of pump surgery and 4 prepared coded treatment required reinfusions; not directly exploration). assignment. involved in clinical treatment of randomised patients. Overall quality assessment (descriptive) Fair. Randomised, double-blind, not all patients included in analysis. RESULTS

Outcome	Intervention group N=50	Comparator group N=50	Statistical significance
Tranexamic acid (IV)			·
	n/N (%)	n/N (%)	P value
Patients transfused with pRBCs (intraoperative)	0/50 (0)	3/50 (6.0)	0.24 (post-hoc)
Patients transfused with pRBCs (0-4 hrs)	0/50 (0)	15/50 (30.0)	<0.001 (post-hoc)
Patients transfused with pRBCs (4-24 hrs)	8/50 (16.0)	9/50 (18.0)	1.00 (post-hoc)
Patients transfused with FFP (0-4 hrs)	2/50 (4.0)	2/50 (4.0)	1.00 (post-hoc)
Patients transfused with FFP (4-24 hrs)	0/50 (0)	0/50 (0)	NA (post-hoc)
Total number of transfused patients	8/50 (16.0)	27/50 (54.0)	<0.001 (post-hoc)
	Mean	Mean	P value
Intraoperative pRBC transfusion (units per transfused patients)	0	1	0.36

Postoperative pRBC transfusion 0-4 hrs (units per transfused patients)	0	1.3		<0.001
Postoperative pRBC transfusion 4-24 hrs (units per transfused patients)	1	1		0.5
Postoperative FFP transfusion 0- 4 hrs (units per transfused patients)	3	2.5		0.8
Postoperative FFP transfusion 4- 24 hrs (units per transfused patients)	0	0		1.00
Total transfusion (units per transfused patients)	1	1.1		NR
	Mean ± SD	Mean ± SD		P value
Intraoperative bleeding (mL)	467 ± 170	531 ± 164		0.62
Postoperative bleeding (0-4 hrs; mL)	87 ± 62	210 ± 195		0.005
Postoperative bleeding (4-24 hrs; mL)	462 ± 118	570 ± 184		0.07
Total bleeding within 24 hrs (mL)	471 ± 182	844 ± 363		<0.001
,	n/N (%)	n/N (%)		P value
Myocardial infarction	0/50 (0)	0/50 (0)		NA
Myocardial ischaemia	0/50 (0)	0/50 (0)		NA
Thrombosis	0/50 (0)	0/50 (0)		NA
Neurologic dysfunction	0/50 (0)	0/50 (0)		NA
	Mean ± SD	Mean ± SD		P value
24-hr postoperative haemoglobin	11.2 ± 0.96	11.1 ± 1.2		0.96
(g/dL)				
Clinical importance			Clinical relev	ance
Tranexamic acid (IV)				
Transfusion incidence	Intraoperative pRBC transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative 0-4 hrs pRBC transfusion 1: Clinically important benefit, confidence limit does not include null value (p<0.001) Postoperative 4-24 hrs pRBC transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative 0-4 hrs FFP transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative 4-24 hrs FFP transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total number transfused patients 1: Clinically important benefit, confidence limit does not include null value (p<0.001)		1: Patient-rele	evant clinical outcome.

Transfusion volume Blood loss	Intraoperative RBC 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative RBC (0-4 hr) 1: Clinically important benefit, confidence limit does not include null value (p<0.001) Postoperative RBC (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (0-4 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	Patient-relevant clinical outcome. 1: Patient-relevant clinical outcome.
Blood loss	important effects, but also compatible with no or harmful effect Postoperative RBC (0-4 hr) 1: Clinically important benefit, confidence limit does not include null value (p<0.001) Postoperative RBC (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (0-4 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	with no or harmful effect Postoperative RBC (0-4 hr) 1: Clinically important benefit, confidence limit does not include null value (p<0.001) Postoperative RBC (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (0-4 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	Postoperative RBC (0-4 hr) 1: Clinically important benefit, confidence limit does not include null value (p<0.001) Postoperative RBC (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (0-4 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	1: Clinically important benefit, confidence limit does not include null value (p<0.001) Postoperative RBC (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (0-4 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	limit does not include null value (p<0.001) Postoperative RBC (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (0-4 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	Postoperative RBC (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (0-4 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (0-4 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	important effects, but also compatible with no or harmful effect Postoperative FFP (0-4 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	with no or harmful effect Postoperative FFP (0-4 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	Postoperative FFP (0-4 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Postoperative FFP (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	important effects, but also compatible with no or harmful effect Postoperative FFP (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	important effects, but also compatible with no or harmful effect Postoperative FFP (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	with no or harmful effect Postoperative FFP (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	FFP (4-24 hr) 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	important effects, but also compatible with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	with no or harmful effect Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	Total transfusion 4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	4: Range of estimates includes clinically important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	important effects, but also compatible with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	with no or harmful effect Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Blood loss	Intraoperative blood loss 4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
Rioog loss	4: Range of estimates includes clinically	1: Patient-relevant clinical outcome.
		1
	important effects, but also compatible	
	with no or harmful effect	
	Blood loss up to 0-4 hr postoperative	
	(mL)	
	1: Clinically important benefit, confidence	
	limit does not include null value (p=0.005)	
	Blood loss up to 4-24 hr postoperative	
	(mL)	
	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
	Total bleeding within 24 hr	
	1: Clinically important <i>benefit</i> , confidence	
	limit does not include null value (p<0.001)	
Myocardial infarction	In-hospital myocardial infarction	1: Patient-relevant clinical outcome.
	No episode of MI reported in either	
	treatment arm	
Myocardial ischaemia	In-hospital myocardial ischaemia	1: Patient-relevant clinical outcome.
<u> </u>	No episode of myocardial ischaemia	
	reported in either treatment arm	
Thrombosis	In-hospital thrombosis	1: Patient-relevant clinical outcome.
1111011100313	No episode of thrombosis reported in	1. 1 ation: Tolovant olinical outcome.
Nouvologio di efi e elice	either treatment arm	1. Detient relevant allulast automo
Neurologic dysfunction	In-hospital neurologic dysfunction	1: Patient-relevant clinical outcome.
	No episode of neurologic dysfunction	
	reported in either treatment arm	
Haemoglobin	24-hr postoperative haemoglobin	1: Patient-relevant clinical outcome.
-	4: Range of estimates includes clinically	
	important effects, but also compatible	
	with no or harmful effect	
EXTERNAL VALIDITY		
Generalisability		

Study conducted specifically in patients undergoing off-pump CABG so likely to be generalisable only to this select group of patients.

Applicability

Study conducted in a single centre in Iran so may not be completely applicable to the Australian/New Zealand setting.

Comments

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: BMI, body mass index; CABG, coronary artery bypass graft; FFP, fresh frozen plasma; ICU, intensive care unit; ITT, intention-to-treat; IV, intravenous; MI, myocardial infarction; NA, not applicable; NR, not reported; pRBCs, packed red blood cells; RCT, randomised controlled trial; SD, standard deviation; TXA, tranexamic acid.

STUDY DETAILS Citation Wong J, El Beheiry H, Rampersaud YR et al (2008) Tranexamic acid reduces perioperative blood loss in adult patients having spinal fusion surgery. Anesth Analg 107: 1479-1486. Affiliation/Source of funds Departments of Anaesthesia and Orthopaedics, Toronto Western Hospital, University Health Network, Toronto, Canada; Department of Orthopaedics, St Michael's Hospital, Toronto, Canada; Departments of Anaesthesia and Orthopaedics, Trillium Hospital, Mississauga, Canada. Funded by the Physicians' Services Incorporated Foundation, Toronto, Canada. Study design Level of evidence Location/setting RCT Hospital (3 centres) Intervention Comparator Tranexamic acid: bolus 10 mg/kg after induction followed Placebo – saline solution (same regimen as TXA) by a maintenance infusion of 1 mg/kg/hr. Note: all patients at one of the three centres received DVT Note: all patients at one of the three centres received DVT prophylaxis. prophylaxis. Population characteristics Adult patients (≥ 18 years) undergoing elective posterior thoracic/lumbar instrumented spinal fusions. Age 57 years TXA and 50 years placebo (p=0.011)a, female 68%, weight ~73 kg. No differences in any other baseline demographics or patient or surgical factors. Length of follow-up Outcomes measured Up to 3 months for thrombosis assessment. Primary: total estimated and calculated perioperative blood loss (intraoperative and 24 h postoperative). Secondary: transfusion incidence and volume (RMCs, coagulation components), haemoglobin, hospital stay, thrombosis. INTERNAL VALIDITY Allocation Blinding analysis Treatment/measurement Follow-up (ITT) Results bias Double-blind. Double-blind so 151 patients enrolled; Computer-generated Assessed by staff 4 (2.6%) withdrawn random numbers; not aware of Research personnel, treatment/measurement bias stratified by surgeon treatment anaesthesiologists, should not be an issue. due to excessive and and number of assignment. surgeons and initially uncontrollable Standard statistical operating room staff surgical bleeding as vertebrae fused. per a priori exclusion Patient assignment tests used. Also blinded to treatment placed into sealed performed linear assignment. criteria. envelopes. and logistic Independent regression. pharmacist prepared infusions which were identical in appearance Overall quality assessment (descriptive) Good. Randomised, double-blind, majority of patients included in analysis (missing patients not likely to bias results. **RESULTS** Outcome Intervention group Comparator group Statistical significance N=74 N = 73Tranexamic acid (IV) Transfusion incidence n/N (%) n/N (%) P value Perioperative Perioperative Patients transfused with pRBCs 23/73 (31) 30/74 (40) 0.25 Patients transfused with AWB 27/74 (36) 24/73 (32) 0.65

Patients transfused with cell- saver blood	33/73 (45)	47/74 (63)	0.026
Patients transfused with FFP	5/73 (7)	9/74 (12)	0.27
Patients transfused with	2/73 (3)	2/74 (3)	0.99
platelets	2113 (3)	2117 (3)	0.77
Intraoperative			
Patients transfused with pRBCs	14/73 (19)	17/74 (23)	0.57
Patients transfused with AWB	18/73 (25)	21/74 (28)	0.61
Patients transfused with cell-	33/73 (45)	46/74 (62)	0.039
saver blood	00/70 (10)	10/7 1 (02)	0.007
Patients transfused with FFP	4/73 (5)	7/74 (9)	0.36
Patients transfused with	2/73 (3)	2/74 (3)	0.99
platelets			
Postoperative			
Patients transfused with pRBCs	11/73 (15)	21/74 (28)	0.051
Patients transfused with AWB	10/73 (13)	10/74 (13)	0.97
Patients transfused with cell-	2/73 (3)	3/74 (4)	0.66
saver blood			
Patients transfused with FFP	0/73 (0)	0/74 (0)	NA
Patients transfused with	0/73 (0)	0/74 (0)	NA
platelets			
5. Transfusion volume			
	Mean ± SD	Mean ± SD	P value
Perioperative			
Patients transfused with pRBCs (mL)	266 ± 541	406 ± 649	0.16
Patients transfused with AWB (mL)	222 ± 343	315 ± 672	0.30
Patients transfused with cell- saver blood (mL)	218 ± 347	334 ± 450	0.083
Intraoperative	<u> </u>		
Patients transfused with pRBCs	169 ± 486	208 ± 436	0.61
(mL)	107 ± 400	200 ± 430	0.01
Patients transfused with AWB	150 ± 278	249 ± 656	0.24
(mL)			
Patients transfused with cell- saver blood (mL)	210 ± 343	323 ± 443	0.086
Postoperative Postoperative			
Patients transfused with pRBCs	97 ± 239	198 ± 384	0.057
(mL)	1. = 20,		
Patients transfused with AWB	72 ± 200	66 ± 198.2	0.85
(mL)	0 40	11 /4	0.70
Patients transfused with cell- saver blood (mL)	8 ± 49	11 ± 64	0.73
6. Blood loss			
<u>Bioda ida</u>	Mean ± SD	Mean ± SD	P value
Perioperative	1 2 0 2	1 2 2 2	1
Estimated blood loss (mL)	1592 ± 1315	2138 ± 1607	0.026
Calculated blood loss (mL)	3079 ± 2558	4363 ± 3030	0.017
Calculated RBC loss (mL)	1078 ± 895	1527 ± 1060	0.017
` '	1070 ± 070		•
Intraoperative	•		0.044
Intraoperative Estimated blood loss (mL)	1203 ± 1060	1600 ± 1301	0.044
Intraoperative	•		0.044

	m/NL (0/)	n/N (0/)		Duralua
Myocardial infarction	n/N (%) 1/73 (0) ^b	n/N (%) 0/74 (0)		P value NS
Myocardial infarction 8. Thrombosis	11/3 (U) ²	0/74 (0)		CVI
o. HIIIUHIDUSIS	n/N (%)	n/N (%)		P value
Thrombosis	0/73 (0)	1/74 (0)		NS
9. Hospital length of stay	0173 (0)	1//4 (0)		1/13
7. Hospital length of stay	Mean ± SD	Mean ± SD		P value
Hospital longth of stay (days)	9.19 ± 5.48	8.47 ± 4.12		0.38
Hospital length of stay (days) 10. Haemoglobin	9.19 ± 3.40	0.47 ± 4.12		0.30
10. Haeiliogiobili	Mean ± SD	Mean ± SD		P value
Lowest postoperative	9.4 ± 1.4	8.9 ± 1.3		0.033
haemoglobin (g/dL)	7.4 ± 1.4	0.7 ± 1.3		0.033
Percentage decrease from	31.1 ± 14.2	3/15 + 13.7		0.154
preoperative to lowest	31.1 ± 14.2	34.3 ± 13.7		0.134
postoperative haemoglobin (%)				
Clinical importance		1	Clinical relev	ance
Tranexamic acid (IV)			1 234113101	
Transfusion incidence	Perioperative RBC transfus	sion	1: Patient-rele	evant clinical outcome.
	4: Range of estimates incluimportant effects, but also on or harmful effect Perioperative AWB transfus 4: Range of estimates incluimportant effects, but also on or harmful effect Perioperative cell-saver transfus 1: Clinically important benelimit does not include null v. Perioperative FFP transfus 4: Range of estimates incluimportant effects, but also on or harmful effect Perioperative platelet transfus 4: Range of estimates incluimportant effects, but also on or harmful effect Intraoperative RBC transfus 4: Range of estimates incluimportant effects, but also on or harmful effect Intraoperative AWB transfus 4: Range of estimates incluimportant effects, but also on or harmful effect Intraoperative cell-saver transfus 4: Range of estimates incluimportant effects, but also on or harmful effect Intraoperative FFP transfus 4: Range of estimates incluimportant effects, but also on or harmful effect Intraoperative platelet transfus 4: Range of estimates incluimportant effects, but also on or harmful effect Intraoperative platelet transfus 4: Range of estimates incluimportant effects, but also on or harmful effect Intraoperative platelet transfus 4: Range of estimates incluimportant effects, but also on or harmful effect Intraoperative Platelet transfus 4: Range of estimates incluimportant effects, but also on or harmful effect	includes clinically also compatible with ansfusion includes clinically also compatible with ar transfusion openefit, confidence all value (p=0.026) and includes clinically also compatible with ansfusion includes clinically also compatible with		

	4: Range of estimates includes clinically	
	important effects, but also compatible with	
	no or harmful effect (p=0.051)	
	Postoperative AWB transfusion	
	4: Range of estimates includes clinically	
	important effects, but also compatible with	
	no or harmful effect	
	Postoperative cell-saver transfusion	
	4: Range of estimates includes clinically	
	important effects, but also compatible with	
	no or harmful effect	
	Postoperative FFP transfusion	
	No postoperative FFP transfusion in	
	either treatment arm	
	Postoperative platelet transfusion	
	No postoperative platelet transfusion in	
	either treatment arm	
Transfusion volume	Perioperative RBC transfusion	1: Patient-relevant clinical outcome.
Transiusion voidine	•	1. Fatient-relevant cimical outcome.
	4: Range of estimates includes clinically	
	important effects, but also compatible with	
	no or harmful effect	
	Perioperative AWB transfusion	
	4: Range of estimates includes clinically	
	important effects, but also compatible with	
	no or harmful effect	
	Perioperative cell-saver transfusion	
	4: Range of estimates includes clinically	
	important effects, but also compatible with	
	no or harmful effect (p=0.083)	
	Perioperative FFP transfusion	
	4: Range of estimates includes clinically	
	important effects, but also compatible with	
	no or harmful effect	
	Perioperative platelet transfusion	
	4: Range of estimates includes clinically	
	important effects, but also compatible with	
	no or harmful effect	
	Intraoperative RBC transfusion	
	4: Range of estimates includes clinically	
	important effects, but also compatible with	
	no or harmful effect	
	Intraoperative AWB transfusion	
	4: Range of estimates includes clinically	
	important effects, but also compatible with	
	no or harmful effect	
	Intraoperative cell-saver transfusion	
	4: Range of estimates includes clinically	
	important effects, but also compatible with	
	no or harmful effect (p=0.086)	
	Intraoperative FFP transfusion	
	4: Range of estimates includes clinically	
	important effects, but also compatible with	
	no or harmful effect	
	Intraoperative platelet transfusion	
	4: Range of estimates includes clinically	
	important effects, but also compatible with	
	no or harmful effect	
	1	i .

Applicability

Study conducted in three Canadian centres so likely to be applicable to the Australian/New Zealand setting.

Comments

Multiple linear regression found TXA to be significantly related to reduced perioperative blood loss (Mean difference -580 mL (95% CI: -949, -211; p=0.002) while logistic regression found a trend for TXA to be related to reduced incidence of perioperative allogeneic pRBC transfusion (OR 0.43; 95% CI: 0.17, 1.07); p=0.068). Authors note the study not powered to find a reduction in incidence or volume of transfusion.

Note: All post-hoc calculations performed using GraphPad software.

Abbreviations: AWB, autologous white blood; DVT, deep vein thrombosis; FFP, fresh frozen plasma; ITT, intention-to-treat; IV, intravenous; NA, not applicable; OR, odds-ratio; pRBCs, packed red blood cells; RCT, randomised controlled trial; SD, standard deviation; TXA, tranexamic acid.

^a Regression analysis showed age not significantly associated with blood loss or transfusion requirements.

^b Asymptomatic MI only; diagnosed via elevation of cardiac enzymes.

Intervention 9 - Appropriate patient positioning

Level II evidence

Citation

De Sio M, Autorino R, Quarto G, Calabro F, Damiano R, Giugliano F, Mordente S, D'Armiento M. Modified Supine versus Prone Position in Percutaneous Nephrolithotomy for Renal Stones Treatable with a Single Percutaneous Access: A Prospective Randomized Trial. European Urology 2008;54(1):196-203.

Affiliation/Source of funds

Seconda Universita degli Studi, Naples, Italy. University of Cassino, Cassino, Italy. Universita Magna Graecia, Catanzaro, Italy.

Funding: NR

Study design	Level of evidence		Location/setting
RCT	Level II		Medical institutions
Intervention		Comparator	
Modified supine position		Prone position	

Population characteristics

75 patients undergoing nephrolithotomy.

Length of follow-up	Outcomes measured
Until discharge	Blood loss, hospital stay, surgery duration, postoperative complications.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomised. Computer generated numbers	No difference in preoperative characteristics between patient groups.	Not possible for surgeons to be blinded of intervention.	Analyst unaware of treatment allocation,	Analysed based on ITT

Overall quality assessment (descriptive)

Good. This study was well described and with appropriate statistical and subgroup analysis conducted.

RESULTS					
Outcome	Intervention group	Comparator group	Statistical significance		
Change in Haemoglobin (g/dL)	-2.3 (-3.5, -0.4)	-2.2 (-3.3, -0.5)	0.23		
Mean hospital stay (days)	4.3 (Range: 2.2–8.4)	4.1 (Range: 2.4–7.8)	0.18		
Operating room time (minutes)	43 (25–120)	68 (55–140)	<0.001		
Major complications (loss of nephrostomy tract, fever)	1/39 patients	0/36 patients	0.2		
Minor complications (transient fever, renal colic, significant bleeding)	7 patients	5 patients	0.16		

Outcome	Clinical importance	Clinical relevance
Change in Haemoglobin	4	1
Mean hospital stay	4	2
Operating room time	1	2
Major complications	4	1
Minor complications	4	1

EXTERNAL VALIDITY

Generalisability

This RCT was conducted in patients undergoing nephrolithotomy. As such, generalisability is likely limited to patients undergoing such a procedure, with similar clinical characteristics.

Applicability

This study was conducted in Italy, where the healthcare system is likely comparable to Australia.

Comments

This study showed that the supine position was similar to the prone position for percutaneous stone removal.

Abbreviations: NR, not reported; RCT, randomised clinical trial.

Citation

Ko MT, Chuang KC, Su CY. Multiple analyses of factors related to intraoperative blood loss and the role of reverse Trendelenburg position in endoscopic sinus surgery. Laryngoscope 2008;118(9):1687-1691.

Affiliation/Source of funds

Chang Gung Memorial Hospital, Kaohsiung Medical Centre, Chang Gung University College of Medicine, Kaohsiung, Taiwan.

Funding: NR

Study design	Level of evidence		Location/setting
RCT	Level II		Hospital
Intervention		Comparator	
Reverse Trendelenburg position		Supine position	

Population characteristics

60 patients undergoing Endoscopic sinus surgeries.

Length of follow-up	Outcomes measured
NR	Blood loss, surgery duration

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomised. Method NR	No difference in preoperative characteristics between patient groups.	Not possible for surgeons to be blinded of intervention.	All operations performed by same surgeon.	No loss to follow- up

Overall quality assessment (descriptive)

Fair. This study was well described and utilised appropriate statistical tests.

RESULTS				
Outcome	Intervention group	Comparator gr	oup	Statistical significance
Total blood loss (mL)	126 (SD: 85.8)	251.67 (SD: 139	9.1)	P<0.001
Blood loss per minute	0.87 (SD: 0.6)	1.74 (SD:1.0)		P<0.001
Surgery duration (minute)	138.5 (SD: 50.8)	165.5 (SD: 56.1))	P=0.056
Outcome	Clinical importance		Clinical relevance	
Total blood loss	2: reduced blood loss, however difference may not be clinically significant		1	
Blood loss per minute	2: reduced blood loss, however difference may not be clinically significant		1	
Surgery duration	4		2	

Generalisability

This RCT was conducted in patients undergoing Endoscopic sinus surgery. As such, generalisability is likely limited to patients undergoing such a procedure, with similar clinical characteristics.

Applicability

This study was conducted in Taiwan, while differences may exist, the level and quality of healthcare is likely comparable to Australia. As such the findings should be applicable.

Comments

This study showed that RTP may reduce intraoperative blood loss.

Abbreviations: NR, not reported; RCT, randomised clinical trial; SD, standard deviation.

Citation						
Ong SM, Taylor	GJSC. Can knee position	save blood fol	lowing total kr	nee replacement?	Knee 2003;10	(1):81-85.
Affiliation/Sour	ce of funds					
University Hospi	tal of Leicester NHS Trust,	, Glenfield Hos	spital, UK.			
Funding: NR						
Study design		Lev	el of evidenc	e	Location/set	tting
RCT		Lev	el II		Hospital	
Intervention				Comparator		
Intervention 1: Leg elevated 35° at the hip with knee flexed to 70° for 6 hours post-operation Intervention 2: Leg elevated 35° at the hip with knee extended Knee extended and level with bed after operation					bed after operation	
Population cha	racteristics					
60 (20 in each tr	eatment group) patients ur	ndergoing prim	nary unilateral	total knee replace	ement for osteo	oarthritis.
Length of follow	v-up		Outcomes	measured		
Until discharge			Haemoglobi	in loss, blood trans	sfusion require	ments, morbidity.
INTERNAL VAL	IDITY					
Allocation	Results	Blinding ar	nalysis	Treatment/ measuremer	nt bias	Follow-up (ITT)
Randomised. Using sealed envelopes Using sealed envelopes Benvelopes with treasure allocation only operation. Envelopes with treasure allocation only operation.			nly opened	NR		No loss to follow- up
Overall quality	assessment (descriptive)		•		•
Fair. This study	was well described and ut	ilised appropri	iate statistical	tests.		

RESULTS							
Outcome	Interv 1	ention group	Intervention group 2	Com	parator group	Statistical significance	
Haemoglobin loss (g/dL)	3.8 (95%Cl 3.3, 4.3)		3.6 (95%Cl 3.0, 4.2)	4.8 (95%	CI 4.0, 5.6)	P=0.018	
Blood transfusion (median number of units transfused)	0 (Range 0, 2)		0 (Range 0,2)	2 (Ra	inge 0, 3.5)	P=0.3	
Blood transfusion incidence	7/20 patients		7/20 patients	11/20) patients	P=0.3	
Knee swelling (cm) 3.4 (Rang		e 1.0, 7.0)	3.3 (Range 1.5, 8.0)	3.8 (Ran	ge 1.5, 8.0)	P=0.6	
Incidence of DVT 1/20 pa		atients	1/20 patients	0/20	patients	NR	
Outcome		Clinical importance			Clinical relevance		
Haemoglobin loss			2: reduced haemoglobin loss, difference may not be clinically significant		1		

Blood transfusion (median number of units transfused)	4	1
Blood transfusion incidence	4	1
Knee swelling (cm)	4	2
Incidence of DVT	NR	1

Generalisability

This RCT was conducted in patients undergoing total knee replacement. As such, generalisability is likely limited to patients undergoing such a procedure, with similar clinical characteristics.

Applicability

This study was conducted in the UK, where the health system is similar to that in Australia. As such, findings are likely applicable.

Comments

This study recommends the elevation of the leg at 35° from the hip with knee extended following total knee replacement for more favourable patient outcomes.

Abbreviations: NR, not reported; RCT, randomised clinical trial.

Pace A, Yousef A. The effect of patient position on blood loss in primary cemented total hip arthroplasty. Archives of Orthopaedic and Trauma Surgery 2008;128(10):1209-1212.

Affiliation/Source of funds

Queens Medical Centre, Derby Road, Nottingham, UK.

Funding: NR

Study design	Level of evidence		Location/setting
RCT	Level II		Hospital
Intervention		Comparator	
Lateral position		Supine position	

Population characteristics

101 patients undergoing hip arthroplasty

Length of follow-up	Outcomes measured
NR, at least up to five days post-operation	Blood loss, change in haemoglobin, transfusion requirements, wound infection, deep vein thrombosis.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomised. Using sealed envelopes opened before surgery	No difference in preoperative characteristics between patient groups.	Not possible to blind surgeon from patients' treatment group.	Standardised transfusion protocol use for both groups.	No loss to follow- up

Overall quality assessment (descriptive)

Fair. This study was well described and utilised appropriate statistical tests.

RESULTS				
Outcome	Intervention group	Comparator	group	Statistical significance
Total blood loss (mL)	1129 (95%CI 989, 1310)	1156 (95%CI	954, 1265)	P=0.41
Surgery duration (minute)	74 (95%CI 63, 89)	69 (95%CI 5	5, 79)	P=0.31
Transfusion incidence	5/51 patients	8/50 patients		P=0.65
Transfusion rate (units/patient)	0.39	0.32		P=0.56
Change in haemoglobin (g/dL) 1 day after operation 5 days after operation	3.6 (95%Cl 2.9, 5.0) 3.7 (95%Cl 2.6, 5.1)	3.9 (95%CI 2 3.75 (95%CI	•	P=0.24 P=0.92
Wound infection	0/51 patients	2/50 patients		NS
Deep vein thrombosis	1/51 patients	0/50 patients		NS
Outcome	Clinical importance		Clinical rele	vance
Total blood loss	4		1	
Surgery duration	4		2	
Transfusion requirement	4		1	

Transfusion rate	4	1
Change in haemoglobin	4	1
Wound infection	4	1
Deep vein thrombosis	4	1

Generalisability

This RCT was conducted in patients undergoing spinal surgery. As such, generalisability is likely limited to patients undergoing such a procedure, with similar clinical characteristics.

Applicability

This study was conducted in the UK where the level and quality of healthcare is likely comparable to Australia. As such the findings should be applicable.

Comments

This study shows that patient positioning in supine or lateral during hip arthroplasty surgery has no bearing on the amount of blood loss.

Abbreviations: CI, confidence interval; NR, not reported; NS, not statistically significant; cRCT, randomised clinical trial.

Park CK. The effect of patient positioning on intraabdominal pressure and blood loss in spinal surgery. Anesthesia and Analgesia 2000;91(3):552-557.

Affiliation/Source of funds

Eulji University School of Medicine, Taejon, Korea.

Funding: NR

Study design	Level	of evidence	Location/setting
RCT	Level	II	Hospital
Intervention		Comparator	
Narrow pad width of Wilson spinal supporting frame		Wide pad width of Wilson s	pinal supporting frame

Population characteristics

40 ASA I and II patients undergoing posterior lumbar spinal surgery.

Length of follow-up	Outcomes measured		
NR	Blood loss, blood transfusion dose and frequency, surgery duration, change in haemoglobin levels.		

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomised. Method NR	No difference in preoperative characteristics between patient groups.	Double-blinded. Surgical team blinded to treatment group	All operations performed by same surgeon, and same anaesthesiologist. Standardised transfusion protocol use for both groups.	No loss to follow- up

Overall quality assessment (descriptive)

Good. This study was well described and utilised appropriate statistical tests.

RESULTS				
Outcome	Intervention group	Comparator gr	oup	Statistical significance
Total blood loss (mL)	878 (SD: 521)	436 (SD: 159)		P<0.05
Blood loss per vertebra (mL)	381 (SD: 236)	190 (SD: 65)		P<0.05
Surgery duration (minute)	136.8 (SD:23.7)	134.0 (SD: 27.8))	NS
Transfusion frequency	5 patients	1 patient		NS
Transfusion dose per patient	2.2 units	2 units		NS
Haemoglobin (g/dL) Preoperative Postoperative	13.1 (SD: 1.0) 10.6 (SD: 1.1)	13.1 (SD: 1.4) 11.3 (SD: 1.1)		NS
Outcome	Clinical importance		Clinical relevance	
Total blood loss	2: increased blood loss, may not be clinically significant		1	

Blood loss per vertebra	2: increased blood loss, may not be clinically significant	1
Surgery duration	4	2
Transfusion frequency	4	1
Transfusion dose per patient	4	1
Change in haemoglobin	4	1

Generalisability

This RCT was conducted in ASA I and II patients undergoing spinal surgery. As such, generalisability is likely limited to patients undergoing such a procedure, with similar clinical characteristics.

Applicability

This study was conducted in Korea, while differences may exist, the level and quality of healthcare is likely comparable to Australia. As such the findings should be applicable.

Comments

This study shows that the use of wide pad on the Wilson frame decreases blood loss.

Abbreviations: NR, not reported; NS, not statistically significant; RCT, randomised clinical trial; SD, standard deviation.

Widman J, Isacson J. Lateral position reduces blood loss in hip replacement surgery: A prospective randomized study of 74 patients. International Orthopaedics 2001;25(4):226-227.

Affiliation/Source of funds

St Gorans Hospital, Stockholm, Sweden.

Funding: NR

Study design	Leve	l of evidence	Location/setting
RCT	Level	H	Hospital
Intervention		Comparator	
Lateral position		Supine position	

Population characteristics

74 patients undergoing hip replacement surgery

Length of follow-up	Outcomes measured
NR	Blood loss, surgery duration, blood transfusion frequency and dose.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/ measurement bias	Follow-up (ITT)
Randomised, using a table of random numbers	Uneven gender distribution between groups	Not possible for the surgeon to be blinded to the treatment group of the patient.	Dropouts from study were random. The surgeons and surgical technique were the same in the two groups.	Drop-outs not included in analysis.

Overall quality assessment (descriptive)

Fair.

RESULTS				
Outcome	Intervention group	Comparator gro	up	Statistical significance
Intraoperative blood loss (mL)	508 (SD:316)	723 (SD: 316)		P<0.001 (Gender adjusted)
Blood loss after 24 hours (mL)	1273 (SD:407)	1374 (SD: 458)		P=0.043 (Gender adjusted)
Blood transfusion frequency	17/30 patients	30/44 patients		P=0.336
Blood transfusion volume (mL)	321 (SD: 341)	407 (SD: 362)		P=0.307
Surgery duration (minutes)	70 (SD: 11)	77 (SD: 19)		NR
Outcome	Clinical importance		Clinica	al relevance
Intraoperative blood loss	2: difference may not be clir	2: difference may not be clinically significant		
Blood loss after 24 hours	2: difference may not be clir	nically significant	1	
Blood transfusion frequency	4	4		
Blood transfusion volume	4	4		
Surgery duration	NR		2	

Generalisability

This RCT was conducted in patients undergoing hip replacement surgery. As such, generalisability is likely limited to patients undergoing such a procedure, with similar clinical characteristics.

Applicability

This study was conducted in the Sweden, where the level of healthcare is likely comparable to that in Australia. As such, findings are likely applicable.

Comments

Patients operated in the lateral position had significantly lower total blood loss (~201mL less)

Abbreviations: NR, not reported; RCT, randomised clinical trial; SD, standard deviation.

Intervention 10 – Preoperative autologous donation

Level I evidence

Citation			
Carless P, Moxey A, O'Connell D, and F efficacy. Transfusion Medicine 14:123-1		tologous transfusion to	echniques: A systematic review of their
Affiliation/Source of funds			
Research supported by a grant obtained purpose grant from the Hunter Area Pat			esearch Council of Australia and a special
Study design	Level of evidence	е	Location/setting
Systematic review of RCTs and observational studies with meta-analysis	I		NA
Search conducted July 2002			
Intervention		Comparator	
Autologous transfusion techniques: preoperative Autologous blood deposit (PAD), ANH, and cell salvage (CS).		Comparator: No Autologous transfusion technique (active versus active comparisons were excluded) Sample size (control for PAD) N=553	
NOTE: This form only contains RCT info PAD.	relevant for	Sample Size (contro	11 (11 FAD) N=333
Sample size (PAD) N=566			
Population characteristics			
Patients older than 18 years undergoing involved orthopaedic surgery, and one t			curative surgery for colorectal cancer, three
Length of follow-up		Outcomes measured	
NA		thrombosis, non-fat	on, infection, wound complication, al MI, rate of allogeneic red blood cell ume of allogeneic blood transfused

INTERNAL VALI	INTERNAL VALIDITY			
Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Concealment of treatment allocation and the method of randomisation were judged to be inadequate in all trials (kappa=1.0 for each of these quality items)	The trials studied an equal ratio of males to females. The mean (or median) age of the participants studied ranged from 57.5 to 71 years for those randomised to PAD and 60.5 to 71 years for those patients randomised to control. SR did not discuss the variability of other baseline characteristics between intervention groups.	Blinding was not reported in any trial.	Non detected	NR

Overall quality assessment (descriptive)

Good

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance
Number of subjects transfused with allogeneic blood 8 trials (N=1119; 566 PAD, 553 control)			RR (95% CI): 0.37 (0.26, 0.54) P<0.05 (Phet=0.0018)
Number of subjects transfused with allogeneic blood (cancer surgery) 5 trials (N=NR)			RR (95% CI): 0.49 (0.38, 0.63) P<0.05 (Phet=NR)
Number of subjects transfused with allogeneic blood (orthopaedic surgery) 3 trials (N=NR)			RR (95% CI): 0.16 (0.07, 0.36) P<0.05 (Phet=NR)
Number of subjects transfused with allogeneic blood (transfusion protocol) 5 trials (N=NR)			RR (95% CI): 0.49 (0.37, 0.63) P<0.05 (Phet=NR)
Number of subjects transfused with allogeneic blood (no transfusion protocol) 3 trials (N=NR)			RR (95% CI):0.15 (0.06, 0.37) P<0.05 (Phet=NR)

Number of subjects receiving any allogeneic and/or autologous RBC transfusion 7 trials (N=1088; 550 PAD, 538 control)			RR (95% CI): 1.29 (1.12, 1.48) P<0.05 (Phet=0.0049)
Preoperative haemoglobin concentration			On average, the preoperative Haemoglobin level of patients who deposited their blood was 1.23 gdL ⁻¹ (95% CI: 0.71, 1.74 gdL ⁻¹)
Mortality			Insufficient evidence
Morbidity: infection 3 tirals (N=NR)			RR (95% CI): 0.70 (0.34, 1.43) P>0.05 (P <i>het</i> =NR)
Morbidity: thrombosis 2 trials (N=NR)			RR (95% CI): 0.82 (0.21, 3.13) P>0.05 (P <i>het</i> =NR)
Morbidity: other			Insufficient evidence for stroke, DVT, and pulmonary embolus.
Clinical importance		Clinical relevance	
EXTERNAL VALIDITY			
Generalisability			
Patients considered similar to g	uideline target population		
Applicability			
All the studies included in this reAus/NZ).	eview were conducted in count	ries with well developed healthca	are systems (not specifically

Comments

The search found eight RCTs for PAD with a total of 1119 subjects.

Davies L, Brown TJ, Haynes S, Payne K, Elliott RA, and McCollum C. (2006) Cost-effectiveness of cell salvage and alternative methods of minimising perioperative allogeneic blood transfusion: a systematic review and economic model (Provisional abstract). Health Technology Assessment 10:1-228.

Affiliation/Source of funds

One of the authors received sponsorships from Haemonetics and AstraTech to attend conferences. The author has also given invited lectures for AstraTech Ltd and Unomedical with honoraria and expenses paid.

Study design	Level of evidence	Location/setting
Systematic review of reviews and a systematic update of Henry 2001.	1	NA

Intervention	Comparator
Transfusion strategies to minimise perioperative allogeneic blood transfusion: cell salvage, PAD, PAD p EPO, EPO, ANH, cell salvage plus ANH, AFs, FSs, restrictive transfusion thresholds or protocols. NOTE: this form only contains information relevant for PAD.	Control group that did not receive PAD.
Search conducted January 2004	

Population characteristics

For inclusion, the SRs had to only include adults undergoing elective, non-urgent surgery. Of the three studies included in the update, one study was performed in patients undergoing maxillofacial surgery, and two were carried out on patients undergoing joint arthroplasty (one for total hip arthroplasty and the other for a mixuture of of unilateral, bilateral, primary and revision hip and knee arthroplasty.

Length of follow-up	Outcomes measured
NA	Number of patients transfused with allogeneic blood, volume of allogeneic blood transfused, number of patients transfused with autologous blood, volume of autologous blood transfused, preoperative Hb, volume of autologous blood wasted and length of hospital stay.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
All of the studies in the update were randomised but only one of the studies described an adequate method to secure allocation concealment.		Blinding was unclear in all three studies, but in one study only the operating surgeon was blinded only to whether the patient had received EPO and the participants received open-label treatment.	Two of the studies in the systematic update reported using a transfusion protocol.	One study did not perform an ITT analysis as participants were excluded from the analysis by the authors and in one study it was unclear whether an ITT analysis had been used. One study did analyse all participants who were recruited.

Good			
RESULTS			
Outcome	Intervention group	Comparator group	Statistical significance
Number of patients transfused with allogeneic blood (all studies) 11 trials (N=1423)	149/716 (21%)	375/707 (53%)	RR (95% CI): 0.36 (0.25, 0.51) P<0.05 (Phet=0.0005)
Number of patients transfused with allogeneic blood (transfusion protocol) 7 trials (N=1196)	138/ 585 (24%)	299/611 (49%)	RR (95% CI): 0.48 (0.38, 0.60) P<0.05 (Phet=0.18)
Number of patients transfused with allogeneic blood (no transfusion protocol) 4 trials (N=217)	11/121 (9%)	76/96 (79%)	RR (95% CI): 0.12 (0.04, 0.33) P<0.05 (Phet=0.08)
Number of patients transfused with allogeneic blood (orthopaedic surgery) 5 trials (N=425)	21/221 (10%)	75/204 (37%)	RR (95% CI): 0.21 (0.11, 0.43) P<0.05 (Phet=0.07)
Number of patients transfused with allogeneic blood (oncology) 5 trials (N=950)	128/467 (27%)	280/483 (58%)	RR (95% CI): 0.49 (0.38, 0.63) P<0.05 (Phet=0.15)
Number of patients transfused with allogeneic blood (maxillofacial surgery) 1 trial (N=48)	0/28 (0%)	20/20 (100%)	RR (95% CI): 0.02 (0.00, 0.28) P<0.05 (P <i>het</i> =NA)
Number of patients transfused with allogeneic/autologous blood (all studies) 9 trials (N=1232)	496/620 (80%)	343/612 (56%)	RR (95% CI): 1.33 (1.10, 1.61) P<0.05 (Phet<0.00001)
Number of patients transfused with allogeneic/autologous blood (transfusion protocol) 5 trials (N=1015)	384/499 (77%)	267/516 (52%)	RR (95% CI): 1.48 (1.16, 1.89) P<0.05 (Phet=0.001)
Number of patients transfused with allogeneic/autologous blood (no transfusion protocol) 4 trials (N=217)	112/121 (93%)	76/96 (79%)	RR (95% CI): 1.10 (0.95, 1.29) P>0.05 (P <i>het</i> =0.26)
Number of patients transfused with allogeneic/autologous blood (orthopaedic surgery) 3 trials (N=234)	105/125 (84%)	43/109 (39%)	RR (95% CI): 1.78 (0.61, 5.20) P>0.05 (Phet<0.00001)

Number of patients transfused with allogeneic/autologous blood (oncology) 5 trials (N=950)	363/467 (78%)	280/483 (58%)	RR (95% CI): 1.38 (1.20, 1.58) P<0.05 (Phet=0.13)
Any thrombosis 3 trials (N=250)	6/140 (4%)	3/110 (3%)	RR (95% CI): 0.82 (0.21, 3.13) P>0.05 (Phet=0.53)
Any infection 3 trials (N=621)	74/309 (24%)	81/312 (26%)	RR (95% CI): 0.70 (0.34, 1.43) P>0.05 (Phet=0.07)
Preoperative Hb levels (g/dl) 5 trials (N=534)	N=267	N=267	WMD (95% CI): -1.16 (- 1.60, -0.73) P<0.05 (Phet=0.004)
Clinical importance		Clinical relevance	

Generalisability

Patients considered similar to guideline target population.

Applicability

All the studies included in this review were conducted in countries with well developed healthcare systems (not specifically Aus/NZ).

Comments

Three new RCTs were included in the update of the Henry 2001 PAD systematic review

Abbreviations: AvC, active versus control

Citation							
Duffy G and Neal KR. (1996) Differend allogeneic blood transfusion: a meta-a Transfusion Medicine 6:325-328.							
Affiliation/Source of funds							
None declared							
Study design	Le	evel of evidence	е		Location/se	tting)
SR of RCTs and retrospective studies Date of search NR	s I			NA			
Intervention			Compa	nrator			
Autologous transfusion (including PAE	or c	ell salvage)		neic blood tran	sfusion only		
Population characteristics	0. 0		7 mogor				
Patients undergoing any surgical oper	ation.						
Length of follow-up			Outcor	nes measure	d		
			Infections				
INTERNAL VALIDITY			ı				
Allocation Results		Blinding analy	ysis	Treatment/i	• • •		Follow-up (ITT)
NR Baseline characteristics NR	stics NR NR			SR did not define whether a transfusion protocol was used		а	NR
Overall quality assessment (descrip	tive)					ı	
Fair							
RESULTS							
Outcome Interv	entio	n group	Comparator group Statistical signific		istical significance		
Infections 4/44 (*1) 1 trial (N=77)	9%)		13/33 (39%) OR (95% CI): 6.5 (2.1, 20.7) P<0.05 (Phet=NA)				
Clinical importance			Clinical relevance				
EXTERNAL VALIDITY							
Generalisability							
Limited generalisability (due to uncertabias)	ninty i	regarding the sn	nall samp	le size, and la	ck of info rega	ırdinç	g potential sources of
Applicability							
The results are applicable to the Austr	alian	context (the stu	dy was c	onducted in G	ermany)		
Comments							
NB: only one RCT of PAD was include	d (He	eiss 1993)					

Forgie MA, Wells PS, Laupacis A, and Fergusson D. (1998) Preoperative autologous donation decreases allogeneic transfusion but increases exposure to all red blood cell transfusion: Results of a meta-analysis. Archives of Internal Medicine 158:610-616.

Affiliation/Source of funds

Dr Laupacis is the recipient of the First Fellowship from the International Society of Technology Assessment in Health Care, funded by the PPP Medical Trust, United Kingdom. Investigators of the International Study of Perioperative Transfusion (ISPOT) received funding from the following sources. The Ottawa Coordinating Centre: Janssen Ortho Inc, Don Mills, Canada; investigators from Australia: the National Health and Medical Research Council and the Hunter Area Pathology Services, Newcastle; investigators from France: Haemonetics France, Ortho Diagnostics France, and University Victor Segalen, Bordeaux; investigators from Scotland: the Scottish National Blood Transfusion Service and the Clinical Resources and Audit Group of the Scottish Health Service; and investigators from the United States: Baxter Healthcare Corporation Biotech Group, Deefield, III, and the Emory Center for Clinical Evaluation Scieinces, Atlanta, Ga.

	ip of the Scottish Health Ser Deefield, III, and the Emory					althcare Corporation
Study design	L	Level of evidence			Location/setting	
SR of RCTs ar studies	nd observational I			NA		
Search conduc	ted April 1996					
Intervention	•		Compa	rator		
PAD (defined as the process by which patients donate blood prior to elective surgery and subsequently receive their own blood in the perioperative period if transfusion is required.			Not PAD			
Population ch	aracteristics		l			
	going elective surgery. In tw went colon resection, and in					nree of the trials the
Length of follo	ow-up		Outcomes measured			
NR			Number of patients exposed to allogeneic blood; total number of patients who underwent transfusion with RBCs (including both allogeneic and autologous units); postoperative complications			
INTERNAL VA	LIDITY					
Allocation	Results	Blinding analysis		Treatment/r bias	neasurement	Follow-up (ITT)
	One of the trials had a	All 6 randomise	ed	Two of the tr	ials did not	Two of the trials had

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
	One of the trials had a large difference in the male-female ratio between control and PAD groups (5:10/8:8). Another trial did not report the male-female ratio. With the exception of one trial that did not report the mean age for the control group, the mean ages in the trials were balanced between the intervention groups.	All 6 randomised studies scored 2 on the Jadad quality scale. Since it is ethically inappropriate to blind autologous donors to the treatment they received, the maximum possible score for quality in these studies was 3.	Two of the trials did not report a transfusion protocol.	Two of the trials had patient withdrawals as follows (control/PAD): 0/14 and 5/26.

Overall quality assessment (. ,		
RESULTS			
Outcome	Intervention group	Comparator group	Statistical significance
Patients transfused with allogeneic blood 6 trials (N=1099; 613 PAD, 486 control)	NR	NR NR	OR (95% CI): 0.17 (0.08, 0.32) P<0.05 (Phet<0.008)
Patients transfused with allogeneic blood (colorectal surgery) 3 trials (N=977; 542 PAD, 435 control)	NR	NR	OR (95% CI): 0.26 (0.19, 0.37) P<0.05 (Phet>0.26)
Patients transfused with allogeneic blood (THA) 2 trials (N=91; 61 PAD, 30 control)	NR	NR	OR (95% CI): 0.20 (0.00, 0.28) P<0.05 (Phet>0.03)
Patients transfused with allogeneic blood (transfusion protocol) 4 trials (N=1008; 558 PAD, 450 control)	NR	NR	OR (95% CI): 0.25 (0.17, 0.37) P<0.05 (Phet>0.20)
Patients transfused with allogeneic blood (no transfusion protocol) 2 trials (N=91; 55 PAD, 36 control)	NR	NR	OR (95% CI): 0.02 (0.00, 0.24) P<0.05 (Phet>0.10)
Patients transferred with allogeneic and/or autologous blood 5 trials (N=1068; 597 PAD, 471 control)	NR	NR	OR (95% CI): 3.03 (1.70, 5.39)
Postoperative infection (colorectal surgery) 2 trials (N=595; 297 PAD, 298 control)	NR	NR	OR (95% CI): 1.44 (0.49, 4.26) P>0.05 (Phet=NR)
Other morbidity	NR	NR	Insufficient evidence for MI, angina, venous thrombosis, and prolonged hospital admission
Clinical importance		Clinical relevance	
EXTERNAL VALIDITY			
Generalisability			
This review is generalisable to	elective surgery with mode	rate blood loss.	
Applicability			
All the studies included in this Aus/NZ).	review were conducted in c	ountries with well developed hea	althcare systems (not specifically

Comments

6 RCTs were included.

Abbreviations: THA, total hip arthroplasty

Gurusamy KS, Li J, Sharma D, and Davidson BR. (2009) Cardiopulmonary interventions to decrease blood loss and blood transfusion requirements for liver resection. Gurusamy. Kurinchi. Selvan., Li Jun, Sharma. Dinesh., Davidson. Brian. R. Cardiopulmonary interventions to decrease. blood loss. and blood transfusion requirements. for liver resection. Cochrane Database of Systematic Reviews: Reviews 2009. Issue. 4 John. Wiley. & So. Affiliation/Source of funds

Citation

Study design	Level of evidence	Location/setting
SR	1	NA
Search conducted November 2008		

Intervention	Comparator
Any cardiopulmonary intervention aimed at reducing	Control: no intervention, placebo, or another intervention aimed
operative blood loss or perioperative allogeneic blood transfusion requirements	at reducing surgical blood loss or at decreasing allogeneic blood transfusion requirements during surgery.

Population characteristics

Patients undergoing liver resection.

Length of follow-up	Outcomes measured
NA	Perioperative mortality, survival, liver failure, perioperative morbidity (other than mortality and liver failure), transfusion requirements, operating time, hospital stay, ICU stay, transaction blood loss, operative blood loss, blood loss within 24 hours.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
See Hashimoto	See Hashimoto 2007	See Hashimoto 2007	See Hashimoto 2007 below	See Hashimoto 2007
2007 below	below	below		below

Overall quality assessment (descriptive)

Good

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance
Mortality	0/40	0/39	0 (0, 0)
			P=NA (P <i>het</i> =NA)
Morbidity: bile leak	0/40 (0%)	1/39 (3%)	RR (95% CI): 0.33 (0.01, 7.75)
			P=0.49 (P <i>het</i> =NA)
Morbidity: intra-abdominal bleeding	0/40 (0%)	1/39 (3%)	RR (95% CI): 0.33 (0.01, 7.75)
			P=0.49 (P <i>het</i> =NA)
Number requiring allogeneic	0/40 (0%)	0/39 (0%)	RR (95% CI): 0 (0, 0)
blood transfusion			P=NA (P <i>het</i> =NA)

Transection blood loss (mL)	140 (185)	230 (185)	WMD (95% CI): -90.0 (-171.60, -8.40) P=0.031 (Phet=NA)			
Operative blood loss (mL)	403 (144)	440 (144)	WMD (9%% CI): -37.0 (-100.51, 26.51) P=0.25 (Phet=NA)			
Clinical importance		Clinical relevance				
EXTERNAL VALIDITY						
Generalisability	Generalisability					
See Hashimoto 2007 below						
Applicability						
See Hashimoto 2007 below						
Comments						
SR found one trial assessing I	PAD (Hashimoto 2007)					

Citation						
Henry DA, Carless PA donation for minimisir						
Affiliation/Source of	funds					
None declared						
Study design	L	evel of evidence	9		Location/setting	
SR of RCT January 2004	I		-		The included trials were conducted in Germany (N=3), Greece (N=1), Japan (N=1), Sweden (N=3), the Netherlands (N=2), and the USA (N=2).	
Intervention			Compa	rator		· ,
PAD			Any			
Population characte	ristics		J			
Adult patients underg	oing elective or non-					r involved curative
Length of follow-up			Outcon	nes measure	d	
INTERNAL VALIDITY			Proportion of patients who were transfused with allogeneic blood or any blood transfusion (allogeneic or autologous); amounts of allogeneic and autologous blood transfused; adverse transfusion reactions; preoperative morbidity; preoperative haemoglobin levels; reported postoperative complications; and mortality.			
Allocation	Results	Blinding analy	Blinding analysis Treatn		neasurement	Follow-up (ITT)
Using the Cochrane grading system for allocation concealment, two trials were assessed as providing inadequate allocation concealment, and, in eight trials the allocation concealment was not clearly described. One trial was assessed to have adequate allocation concealment.		Blinding was n reported in any trials assessed methodologica quality.	y of the I for	Eight of the the use of tr	trials reported ansfusion ith a transfusion	Of the trials assessed for methodological quality, seven trials reported either no exclusions or reported the use of ITT analysis. In one trial, exclusions were reported; however, these were deemed unlikely to cause bias. In one trial, exclusions were deemed excessive and likely to cause bias
Overall quality asses	ssment (descriptive	e)				
Good						

880

Outcome	Intervention group	Comparator group	Statistical significance	
Patients transfused with allogeneic blood ¹ 11 trials (N=1423)	149/716 (21%)	375/707 (53%)	RR (95% CI): 0.36 (0.25, 0.51) RD (95% CI): -0.44 (-0.61, -0.27) NNT (95% CI): 2.3 (1.6, 3.7) P<0.05 (Phet=0.00052)	
Patients transfused with allogeneic blood (oncology) 5 trials (N=950)	128/467 (27%)	280/483 (58%)	RR (95% CI): 0.49 (0.38, 0.63) P<0.05 (Phet=0.15)	
Patients transfused with allogeneic blood (orthopaedic surgery) 5 trials (N=425)	21/221 (10%)	75/204 (37%)	RR (95% CI): 0.21 (0.11, 0.43) P<0.05 (Phet=0.07)	
Patients transfused with allogeneic blood (maxillofacial surgery) 1 trial (N=48)	0/28 (0%)	20/20 (100%)	RR (95% CI): 0.02 (0.00, 0.28) P<0.05 (P <i>het</i> =NA)	
Patients transfused with allogeneic blood (transfusion protocol) 7 trials (N=1206)	138/595 (23%)	299/611 (49%)	RR (95% CI): 0.48 (0.38, 0.60) P<0.05 (Phet=0.18)	
Patients transfused with allogeneic blood (no transfusion protocol) 4 trials (N=217)	11/121 (9%)	76/96 (79%)	RR (95% CI): 0.12 (0.04, 0.33) P<0.05 (Phet=0.08)	
Patients transfused with allogeneic and/or autologous blood 9 trials (N=1232)	496/620 (80%)	343/612 (56%)	RR (95% CI): 1.33 (1.10, 1.61) P<0.05 (Phet<0.000001)	
Patients transfused with allogeneic and/or autologous blood (orthopaedic surgery) 3 trials (N=234)	105/125 (84%)	43/109 (39%)	RR (95% CI): 1.78 (0.61, 5.20) P>0.05 (Phet<0.00001)	
Patients transfused with allogeneic and/or autologous blood (oncology) 5 trials (N=950)	363/467 (78%)	260/483 (54%)	RR (95% CI): 1.38 (1.20, 1.58) P<0.05 (Phet=0.13)	
Patients transfused with allogeneic and/or autologous blood (maxillofacial surgery) 1 trial (N=48)	28/28 (100%)	20/20 (100%)	RR (95% CI): 0 (0, 0) (Phet=NA)	

Patients transfused with allogeneic and/or autologous blood (transfusion protocol) 5 trials (N=1015)	384/499 (77%)	267/516 (52%)	RR (95% CI): 1.48 (1.16, 1.89) P<0.05 (Phet=0.001)
Patients transfused with allogeneic and/or autologous blood (no transfusion protocol) 4 trials (N=217)	112/121 (93%)	76/96 (79%)	RR (95%CI): 1.10 (0.95, 1.29) P>0.05 (Phet<0.00001)
Preoperative haemoglobin, g/dL 5 trials (N=534; 267 PAD, 267 control))	Mean (SD):	-	WMD (95% CI): -1.16 (-1.60, -0.73) P< 0.05 (Phet=0.004)
Infection 3 trials (N=621)	74/309 (24%)	81/312 (26%)	RR (95% CI): 0.70 (0.34, 1.43) P>0.05 (Phet=0.07)
Any thrombosis 3 trials (N=250)	6/140 (4%)	3/110 (3%)	RR (95% CI): 0.82 (0.21, 3.13) P>0.05 (Phet=0.53)
Other morbidity/mortality			There were insufficient data to draw any conclusions about the effect of PAD on mortality, stroke, DVP, and pulmonary embolus.
Clinical importance		Clinical relevance	

Generalisability

Patients considered similar to guideline target population

Applicability

All the studies included in this review were conducted in countries with well developed healthcare systems (not specifically Aus/NZ).

Comments

Twelve studies were included in the analysis. Lorenz 1991 was not assessed for methodological quality, however, as only an English abstract was available.

¹Of the 761 patients randomised to PAD, the majority (N=467) donated their blood prior to cancer surgery.

Citation								
Laupacis A and F (Structured abstra				ologies to	minimise peri	operative alloge	neic transfusion	
Affiliation/Source	e of funds							
NR								
Study design		L	evel of evidence	9		Location/sett	ng	
SR January 1997		1				NA		
Intervention		l .		Compa	rator			
Technologies to r transfusion: aprol epsilon aminocap	inin, desmopi roic acid, ery	ressin, trane thropoietin, l	xamic acid,	Any				
Population char	•							
-		tive surgery.	Types of surger	y included	d cardiac, colo	prectal, liver rese	ection and orthopaedic	
Length of follow	-up			Outcomes measured				
NR			Proportion of patients receiving at least one unit of allogeneic packed red blood cells, perioperative MI, re-operations because of bleeding.					
INTERNAL VALI	DITY			•				
Allocation	Results		Blinding analy	ding analysis Tre		measurement	Follow-up (ITT)	
NR	Baseline characteris	tics NR	NR		Use of transfusion protocol included in subgroup analysis.		NR	
Overall quality a	ssessment (descriptive)					
Fair								
RESULTS								
Outcome		Intervention group		group Comp		ip St	Statistical significance	
Proportion of pati receiving allogene transfusion 6 trials (N=933)		N=NR		N=NR		0.	R (95% CI): 0.17 (0.08, 32) :0.05 (P <i>het</i> =NR)	
Proportion of pati receiving allogend transfusion (color surgery) Number of trials N	eic ectal	-		-		0.	R (95% CI): 0.26 (0.19, 37) :0.05 (P <i>het</i> =NR)	

Proportion of patients receiving allogeneic transfusion (THA) Number of trials NR (N=NR)	-	-	OR (95% CI): 0.20 (0.00, 0.28) P<0.05 (Phet=NR)			
Proportion of patients receiving allogeneic transfusion (transfusion protocol) Number of trials NR (N=NR)	_	_	OR (95% CI): 0.25 (0.17, 0.37) P<0.05 (Phet=NR)			
Proportion of patients receiving allogeneic transfusion (no transfusion protocol) Number of trials NR (N=NR)	-	-	OR (95% CI): 0.02 (0.00, 0.24) P<0.05 (Phet=NR)			
Proportion of patients receiving allogeneic and/or autologous transfusion 5 trials (N=NR)	-	-	OR (95% CI): 3.03 (1.70, 5.39) P<0.05 (Phet=NR)			
Clinical importance		Clinical relevance				
EXTERNAL VALIDITY		<u> </u>				
Generalisability						
The SR is generalisable for ele	The SR is generalisable for elective, non urgent surgery.					
Applicability						
The studies were mostly from	countries with similar health-care	e systems to Australia				
Comments						

884

Citation								
Vamvakas EC. (2 recipients of allog							erative infection between	
Affiliation/Source	e of funds							
None declared								
Study design		L	evel of evidence	е		Location/set	ting	
SR		1				NA		
January 2002				1				
Intervention				Compa				
Autologous transfi salvage.				Transfu blood.	sion of non-W	/BC reduced all	ogeneic RBCs or whole	
NB: this form only Vamvakas is not analyses conflate with trials testing	discussed in results from	l1-4, becaus trials testing	se the meta-					
Population char	acteristics							
Patients undergo	ing any surgio	cal operation	1					
Length of follow	-up			Outcomes measured				
NA				Postoperative infection				
INTERNAL VALI	DITY							
Allocation	Results		Blinding anal	ysis	rsis Treatment/measurement Follow bias		Follow-up (ITT)	
Not reported					Use of transfusion protocol not reported			
Overall quality a	ssessment (descriptive	·)					
Poor								
RESULTS								
Outcome		Interventi	on group	Con	parator grou	ip S	tatistical significance	
Rate of infection 2 trials (N=590)		NR		NR		4	OR (95% CI): 1.35 (0.45, 4.08) P>0.05 (Phet=NR)	
Clinical importance			Clinical relevance					
EXTERNAL VAL	IDITY			T.				
Generalisability								
The populations i	n the trials ar	e similar to t	the guideline pop	ulation.				
Applicability								
The systematic re	eview is applic	cable to the	Australian conte	xt.				

Comments

NB: five studies met the criteria for the meta-analysis.

Level II evidence

Citation	
Citation	

Bouchard D, Marcheix B, Al Shamary S, Vanden Eynden F, Demers P, Robitaille D, Pellerin M, Perrault LP, and Carrier M. (2008) Preoperative autologous blood donation reduces the need for allogeneic blood products: A prospective randomized study. Canadian Journal of Surgery 51:422-427.

Affiliation/Source of funds

None declared

Study design	Level of evidence	Location/setting
RCT	=	Canada / hospital

Intervention	Comparator
PAD: protocol consisted of harvesting 2 units of 39 each (or 6 mL/kg when the patient's weight was be kg). 1 unit was harvested weekly. N=25	

Population characteristics

Patients undergoing elective cardiac surgery.

Length of follow-up	Outcomes measured

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
It is unclear whether allocation was blinded from those responsible for recruiting subjects.	The treatments arms had similar demographic characteristics	Neither the patient nor the surgeon was blinded to the group assignment; however, the ICU intensivist, nurses and residents were blinded.	A transfusion protocol was used.	All analyses were conducted ITT. PAD was not completed in 2 patients (8%) because of worsened angina pectoris.

Overall quality assessment (descriptive)

Fair

RESULTS

Outcome	Intervention group	Comparator group	Statistical significance
Mean (SD) perioperative blood loss, mL	416 (190)	450 (281)	P=0.62
Mean (SD) postoperative blood loss, mL	936 (583)	909.5 (576)	P=0.88
Mean (SD) duration of surgery, min	174.7 (44.9)	177.6 (62.3)	P=0.85
Patients transfused with autologous blood	6/25 (24%)	-	-
Mean (SD) units of autologous blood transfused (for those transfused)	2 (1.2)	-	-

Patients transfused with allogeneic blood products	4/25 (16%)	9/23 (39%)	RR=2.25 P=0.036
Patients transfused with allogeneic blood	0	7/23 (30%)	-
Mean (SD) units of allogeneic whole blood transfused (for those transfused)	0	2 (1.2)	-
Patients transfused with FFP	1/25 (4%)	5/23 (22%)	P=0.06
Mean (SD) units of FFP transfused (for those transfused)	4	2.8 (1)	NR
Patients transfused with platelets	3/25 (12%)	4/23 (17%)	P=0.59
Mean (SD) units of platelets transfused (for those transfused)	4.3 (2.9)	6 (0)	NR
Patients transfused with cryoprecipitate	0	1/23 (4%)	P=0.29
Mean (SD) units of cryoprecipitate transfused (for those transfused)	-	10	NR
Patients transfused with allogeneic and/or autologous blood products	11/25 (44%)	9/23 (39%)	P=0.036
Mean (SD) preoperative haemoglobin concentration, g/L	128.7 (14.4)	135.0 (12.9)	P=0.12
Mean (SD) haemoglobin concentration 24 hours after surgery, g/L	81.6 (12.1)	86.2 (13)	P=0.22
Mean (SD) haemoglobin concentration 5 d after surgery, g/L	102.8 (11.8)	107.7 (11.9)	P=0.16
Mean (SD) preoperative prothrombin time, sec	9.7 (2.8)	9.4 (1.1)	P=0.69
Mean (SD) prothrombin time 30 min after surgery, sec	13.2 (3.9)	13.5 (2.2)	P=0.76
Mean (SD) prothrombin time 24 h after surgery, sec	10.3 (1.3)	10.9 (1.7)	P=0.16
Mean (SD) preoperative fibrinogen concentration, g/L	4.3 (1.5)	3.1 (0.9)	P=0.004
Mean (SD) fibrinogen concentration 30 min after surgery, g/L	3.0 (0.9)	2.6 (0.7)	P=0.10
Mean (SD) fibrinogen concentration 24 h after surgery, g/L	6.2 (1.3)	5.1 (1.2)	P=0.006
Length of stay in hospital, d	5.4 (0.9)	5.4 (0.9)	P=1.00

Length of stay in ICU, d	1.5 (0.6)	1.5 (0.6) P=1.00		
Clinical importance		Clinical relevance		
EXTERNAL VALIDITY				
Generalisability				
The patients investigated are similar to the guideline population.				
Applicability				
The study was conducted in Montreal; however, the results are applicable to the Australian context.				
Comments				

¹The 1 patient received 2 autologous blood units.
²1 patient received 3 allogeneic blood units, and another patient received 6 platelet units.
³1 patient received 4 FFP units and 3 patients received platelets (mean 4.3, SD 2.9 units).
⁴5 patients received FFP units (mean 2.8, SD 1 units), 4 patients received 6 platelet units and 1 patient received 10 cryoprecipitate units.

Hashimoto T, Kokudo N, Orii R, Seyama Y, Sano K, Imamura H, Sugawara Y, Hasegawa K, and Makuuchi M. (2007) Intraoperative blood salvage during liver resection: A randomized controlled trial. Annals of Surgery 245:686-691.

Affiliation/Source of funds

Study supported by a grant from the Kanae Foundation for Life and Socio-medical Science, a grant from the Public Trust Surgery Research Fund, a grant from the Japanese Clinical Oncology Fund, a grant from the Public Trust Haraguchi Memorial Cancer Research Fund in Japan, and a Grant-in-Aid for Scientific Research from the Ministry of Education, Culture, Sports, Science and Technology of Japan.

RCT	II		
IXO1			Japan / hospital
Intervention		Comparator	
PAD: blood volume equal to approximately 0.7% of the patient's body weight was collected before the liver transection. The collected blood was reinfused into the patient after the graft procurement. N=40		Control: "control group using a minimization method" N=39	

Population characteristics

Patients scheduled to undergo liver graft procurement.

L. and the second of the secon	
Length of follow-up	Outcomes measured
NR	The primary outcome measure was blood loss during hepatic parenchymal division. Secondary outcome measures consisted of total blood loss, blood loss during hepatic parenchymal division per unit transaction area, CVP at the start of the hepatic parenchyma division, serum aspartate aminotransferase and total bilirubin levels on the third postoperative day, length of hospital stay, and morbidity.

INTERNAL VALIDITY

Allocation	Results	Blinding analysis	Treatment/measurement bias	Follow-up (ITT)
Participants were randomised to treatment; however, it is unclear whether allocation was concealed to those in charge of recruiting participants.	The treatment arms were similar in baseline and operative values.	The patients and surgeons were blinded to randomisation results.	None detected	One patient in the control group was excluded from analysis after randomisation because the operation was stopped due to an asthmatic attack.

Overall quality assessment (descriptive)

Fair

RESULTS			
Outcome	Intervention group	Comparator group	Statistical significance
Morbidity ¹	Major: 1/40 Minor: 6/40	Major: 2/39 Minor: 9/39	P>0.999
Median (range) length of hospital stay	14 (10 to 36)	14 (11 to 46)	P=0.476
Median (range) preoperative haemoglobin concentration, g/dL	13.0 (11.0 to 15.7)	13.6 (11.6 to 15.9)	P=0.455
Median (range) haemoglobin concentration 24-h postoperative, g/dL	12.3 (9.6 to 15.9)	12.5 (10.5 to 15.0)	P=0.280
Median (range) preoperative PT-INR	1.11 (0.95 to 1.34)	1.10 (0.91 to 1.31)	P=0.350
Median (range) PT-INR 24 h postoperative	1.76 (1.30 to 2.37)	1.77 (1.29 to 2.32)	P=0.456
Median (range) intraoperative blood loss	Total, mL: 403 (120 to 1240) During liver parenchymal division, mL: 140 (40 to 430) During liver parenchymal division per unit transactional area (mL/cm²): 2.15 (0.86 to 7.37)	Total, mL: 440 (130 to 1230) During liver parenchymal division, mL: 230 (40 to 660) During liver parenchymal division per unit transactional area (mL/cm²): 3.75 (0.64 to 7.93)	P-value Total: 0.257 During liver parenchymal division: 0.034 During liver parenchymal division per unit transactional area (mL/cm²): 0.012
Mean (range) total operation time (min)	473 (385 to 640)	470 (380 to 730)	P=0.883
Clinical importance		Clinical relevance	

Generalisability

The study population is limited to people undergoing liver graft procurement, however it is still somewhat generalisable to other elective surgeries associated with moderate blood loss.

Applicability

The study was conducted in Japan, which may limit it's applicability to the Australian context.

Comments

¹Complications requiring surgical or other interventions were defined as major complications, while those that resolved with conservative treatment but prolonged the hospital stay by more than 2 weeks (eg, wound infection, ileus, and minor bile leakage) were defined as minor complications.

Abbreviations: PT-INR, prothrombin time international normalisation ratio