Technical report

Volume 3 – Additional literature

Prepared for National Blood Authority

Project

Update of the 2003 Guideline on the Prophylactic Use of Rh D Immunoglobulin (Anti-D) in Obstetrics

The Commonwealth of Australia as represented by the National Blood Authority

Technical report prepared by Health Technology Analysts Pty Ltd

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Note

This volume ('the 2021 udpate') presents additional literature published and identified after a systematic literature review on use of Rh D Immunoglobulin (Anti-D) in RhD negative pregnant women. Volume 1 presents the main body of evidence. Volume 2 present the appendixes (Appendix A to Appendix F) that document the evidence synthesis (published in 2018). Together the three volumes cover all research questions and evidence reviewed for this topic.

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Findings of the systematic review

Results of the literature search

The medical literature was searched on 27-28 September 2021 to identify relevant studies and systematic reviews published between 2018 to the literature search date. Searches were conducted of the databases and sources described previously (**Section 3.2, Technical report, Volume 1**). Manual searches of the reference lists of relevant articles were also performed.

Search terms are as described in **Appendix A**, with methodological filters applied to identify specific study types. Studies were excluded based on hierarchical, prespecified exclusion criteria as described previously (see **Technical report, volume 1**), with all citations returned by the literature searches reviewed based on information in the publication title and, where available, the abstract. Relevant publications were retrieved and reviewed in full text before a final decision was made on their inclusion or exclusion for the review. The expert group was consulted in cases where further judgement was required.

The results of the screening process and the application of the study selection criteria is provided in **Appendix B**. A PRISMA flow summarising the screening results is provided in **Figure 1** (all questions) and **Figure 2** (subquestion 3, diagnostic accuracy).

A total of 12 new studies were identified and included in the review (Alshehri, 2021, Jernman, 2021, Legler, 2021, Ontario Health, 2020, Parchure, 2021, Pazourkova, 2021, Runkel, 2020, Schmidt-Hansen, 2020, White, 2019, Wikman, 2021, Xie, 2020, Yang, 2019).

Studies that technically met the inclusion criteria (or potentially) but were later excluded (e.g., contained insufficient or inadequate data for inclusion, were considered incompatible with the Australian context) are listed in **Appendix C**.



Search conducted 27-28 Sept 2021, including Embase, MEDLINE, PubMed, Cochrane and CINAHL

Figure 2 Literature screening results. Questions 3.



Search conducted 27-28 Sept 2021, including Embase, MEDLINE, PubMed, Cochrane and CINAHL

Question 1 - Routine antenatal Rh D immunoprophylaxis

Question 1 – (Intervention)

In Rh D negative pregnant women with no preformed anti-D, does universal *routine* antenatal prophylaxis with Rh D immunoglobulin (1 or 2 doses) prevent Rh D alloimmunisation?

Subquestion 1 – (Intervention)

In Rh D negative pregnant women with no preformed anti-D, is universal *routine* antenatal prophylaxis with one dose of Rh D immunoglobulin as effective at preventing Rh D alloimmunisation as universal *routine* prophylaxis with two doses of Rh D immunoglobulin?

Background

The 2018 review identified four systematic reviews (Chilcott, 2003, McBain, 2015, Pilgrim, 2009, Turner, 2012) and one Level III study (Koelewijn, 2008) that evaluated the effectiveness of RAADP in Rh D negative women. The reviews identified two Level II studies (Huchet, 1987, Lee, 1995) and nine Level III studies (Bowman, 1978, Bowman, 1978, 1987, Hermann, 1984, MacKenzie, 1999, Mayne, 1997, Parsons, 1998, Tovey, 1983, Trolle, 1989) meeting their search criteria.

Summary of evidence

The 2021 update found one additional systematic review (Xie, 2020) that evaluated the effectiveness of RAADP in Rh D negative women. One Level II study (White, 2019) was also included that reported on serum anti-D antibody levels in Rh D negative women who had received one or two doses of RAADP.

Xie 2020 was a network meta-analysis that examined varying doses of Rh D immunoglobulin compared to no treatment in Rh D negative women. The authors searched multiple databases (including a Chinese database) up to 7 July 2019 and included studies that examined both antenatal and postnatal administration Rh D immunoglobulin that were published between 1958 and 2004. Doses of Rh D immunoglobulin administered varied between a single dose (250 µg) at 28 weeks through to two doses (300 µg) at 28 and 34 gestational weeks, with or without administration of 100 to 300 µg up to 72 hours postnatally. No new studies were found. Treatments were ranked using surface area under the curve analysis of cumulative probability of preventing Rh D alloimmunisation.

White 2019 is the published report of the Australian trial previously included in the 2018 review (see Pennell 2017 conference abstract). White 2019 compared two doses of Rh(D) immunoglobin-VF 625 (IU) administered at 28 and 32 weeks' gestation with a single dose of 1500 IU given at 28 weeks' gestation. Recruitment occurred through randomising Rh D pregnant women who intended to give birth at a tertiary obstetric referral hospital in Perth between May 2013 and November 2015. The main outcome assessed was the presence of Rh(D) immunoglobin antibodies in maternal blood at the time of delivery.

Results

Incidence of Rh D alloimmunisation

As reported by previous SRs, the network meta-analysis by Xie 2020 also showed an effect favouring RAADP compared to no treatment in preventing Rh D alloimmunisation. The analyses included different doses and timing of Rh D immunoglobin but favoured RAADP in all cases (odds ratio ranging from 0.00 to 0.15).

Based on analysis of the surface area under the cumulative ranking curve (SUCRA), Xie 2020 suggested that two dose of 1500 IU of Rh D immunoglobulin given at 28 and 34 gestational weeks' is better than other dosing regimens (SUCRA = 96.8%), with the second alternative being a single dose (1500 IU) given at 28

gestational weeks (SUCRA = 89.2%), followed by two doses (500 IU) given between 28 and 34 gestational weeks (SUCRA = 75.1%).

Serum anti-D antibody levels

White 2019 reported similar numbers to that reported in the 2018 review, noting that the number of women with anti-D antibody present at birth was higher in those women who received the two-dose regime compared to the one-dose regimen (86% v 56%; OR 4.91; CI 2.67, 9.02; p < 0.001). Concerns about the effect estimate exist, relating to missing antibody screening data (8%) and that twelve women in the single dose group (9%) received an incorrect dose (625 IU) at 28–30 weeks and were therefore given a second dose at 34–36 weeks to avoid potential late antenatal sensitisation.

As previously noted, the relationship between a lack of detectable circulating anti-D antibody following Rh D immunoprophylaxis and risk of alloimmunisation detected in a subsequent pregnancy is not known.

Discussion

If and how the 2021 search has impacted on evidence base?

The 2021 search provided two additional studies relevant to Question 1 (Xie 2020 and White 2019). Both studies provided solidified the existing evidence in favour of issuing universal routine antenatal Rh D immunoglobin to prevent Rh D alloimmunisation.

If and how the 2021 search has created changes in the evidence?

The studies found did not conflict or contradict any of the existing evidence, therefore no changes should be made to the 2018 recommendations. Questions regarding the effectiveness of a single dose of Rh D immunoglobulin compared to two doses remain unanswered.

Question 2 - Universal sensitising event prophylaxis in the first trimester

Question 2 – (Intervention)

In Rh D negative women with no preformed anti-D who have experienced one of the following first trimester sensitising events – abdominal trauma, molar pregnancy, ectopic pregnancy, spontaneous miscarriage, threatened miscarriage or medical termination of pregnancy (with/without a curette) – does universal first trimester sensitising event prophylaxis with Rh D immunoglobulin prevent Rh D alloimmunisation?

Background

The 2018 review identified two systematic reviews (Karanth, 2013, NCCWCH, 2012) that evaluated the effectiveness of prophylactic Rh D immunoglobulin in response to a first trimester sensitising event. The reviews included one Level II study (Visscher, 1972) and two Level III studies meeting the PICO criteria (Gavin, 1972, Simonovits, 1974).

No additional studies evaluating the use of prophylactic Rh D immunoglobulin in women with first trimester ectopic pregnancy, threatened miscarriage, or molar pregnancy were found.

Summary of evidence

The 2021 update found one additional systematic review (Schmidt-Hansen, 2020) that searched for evidence relating to sensitising events in women undergoing either medical abortion with mifepristone and misoprostol or surgical abortion using vacuum aspiration of a pregnancy up to 13⁺⁶ weeks' gestation. The review was used to inform the 2019 NICE guidelines on abortion care (NICE, 2019).

Results

In the absence of evidence, the following expert consensus guide was developed:

- Offer anti-D prophylaxis to women who are rhesus D negative and are having an abortion after 10⁺⁰ weeks' gestation.
- Do not offer anti-D prophylaxis to women who are having a medical abortion up to and including 10⁺⁰ weeks' gestation.
- Consider anti-D prophylaxis for women who are rhesus D negative and are having a surgical abortion up to and including 10⁺⁰ weeks' gestation.
- Providers should ensure that:
 - rhesus status testing and anti-D prophylaxis supply does not cause any delays to women having an abortion
 - o anti-D prophylaxis is available at the time of the abortion.

Discussion

If and how the 2021 search has impacted on evidence base?

The 2021 search provided one additional systematic review relevant to Question 2 (Schmidt-Hansen 2020), which found no new evidence relating to administration of antenatal Rh D immunoglobin to prevent Rh D alloimmunisation in women undergoing medical or surgical abortion.

If and how the 2021 search has created changes in the evidence?

No new studies were found therefore no changes should be made to the 2018 recommendations. In the absence of evidence, the precise benefits and risks of anti-D prophylaxis relating to medical termination of pregnancy before 10 weeks of gestation remain unclear.

Question 3 - Targeted routine antenatal or sensitising event prophylaxis

Question 3 – (Screening intervention)

In Rh D negative pregnant women with no preformed anti-D, does *targeted* routine antenatal or sensitising event prophylaxis to women with a Rh D positive fetus increase the incidence of Rh D alloimmunisation compared with *universal* routine antenatal or sensitising event prophylaxis?

Subquestion 3 – (diagnostic accuracy)

In Rh D negative pregnant women with no preformed anti-D, what is the diagnostic accuracy of noninvasive prenatal screening to identify fetal Rh D status?

Background

The 2018 review identified one systematic review (Saramago, 2018) that searched for evidence regarding the comparative effectiveness of targeted antenatal Rh D immunoprophylaxis against universal routine Rh D immunoprophylaxis. The report did not identify any head-to-head studies of targeted versus routine antenatal prophylaxis regimes that met the criteria for the review.

There were four systematic reviews that examined the diagnostic accuracy of NIPT to identify fetal Rh D status (Geifman-Holtzman, 2006, Mackie, 2017, Saramago, 2018, Zhu, 2014). The reviews included over 90 studies meeting their search criteria. Five additional Level II studies (Haimila, 2017, Macher, 2012, Manfroi, 2018, Moise, 2016, Picchiassi, 2015) and six additional Level III study (Hyland, 2017, Jakobsen, 2018, Orzińska, 2015, Papasavva, 2016, Ryan, 2017, Sorensen, 2018) were identified and subsequently included in the evidence review. Studies that were of small sample size (N<200), conference abstracts that did not provide sufficient data, and those in which the NIPT was not conducted in the context considered similar to Australia were excluded (see **Technical report, volume 1**).

Summary of evidence

The 2021 update found four systematic review that searched for evidence regarding the comparative effectiveness of targeted RAADP against universal RAADP and/or examined the diagnostic accuracy of NIPT to identify fetal Rh D status (Alshehri, 2021, Ontario Health, 2020, Runkel, 2020, Yang, 2019). Three of the reviews were published reports of health technology assessments used to inform the Canadian Agency for Drugs and Technologies in Health (Ontario Health, 2020) the German Institute for Quality and Efficiency in Health Care (Runkel, 2020) and the NHS (Yang, 2019). Alsheri 2021 was a systematic review focused on the diagnostic accuracy of NIPT to identify fetal Rh D status. The authors identified 16 studies, 11 of which were included in a meta-analysis.

One additional Level III study was identified that examined the effectiveness of targeted antenatal Rh D immunoprophylaxis against no routine prophylaxis (Jernman, 2021). Jernman 2021 reported the results of a nationwide cohort study conducted in all pregnant women with anti-D antibodies detected in the Finnish Red Cross (FRC) Blood Service between January 1, 2014 and December 31, 2017.

Two Level II studies (Parchure, 2021, Pazourkova, 2021) and one Level III study (Legler, 2021) that examined the diagnostic accuracy of NIPT to identify fetal Rh D status were also identified and included in the evidence review. Studies that were of small sample size (N<200), conference abstracts that did not provide sufficient data, and those in which the NIPT was not conducted in the context considered similar to Australia were excluded (see **Appendix C**).

Results

Incidence of Rh D alloimmunisation

Similar to the evidence found previously, the data reported in the SRs and that reported by Jernman 2021 suggests that the risk of Rh D alloimmunisation is lower in the cohort that received targeted RAADP compared with the historic reference cohort that received postnatal and antenatal Rh D immunoglobulin prophylaxis following any potentially sensitising events.

Utilisation of Rh D immunoglobulin

The Ontario health report noted that across studies, 25.3% to 39% of all Rh D negative pregnancies avoided unnecessary Rh D immunoglobulin after noninvasive fetal RhD blood group genotyping. Among the Rh D negative mothers carrying an Rh D negative fetus, over 90% avoided unnecessary Rh D immunoglobulin.

Diagnostic performance

Similar to the evidence reported in 2018, the data reported in the SRs and newly included studies suggests that the diagnostic performance of NIPT to identify fetal Rh D status is good, with the bivariate analysis reported by Runkel 2020 (12 studies, 60 011 participants) estimating high sensitivity 99.9% (95% CI 99.5, 100) and high specificity 99.2% (95% CI 98.5, 99.5).

Discussion

If and how the 2021 search has impacted on evidence base?

The 2021 search provided four additional systematic review (Alshehri 2021, Ontario Health 2020, Runkel 2020, Yang 2019), one cohort study (Jernman 2021) and three diagnostic accuracy studies (Parchure 2021, Pazourkova 2021, Legler 2021) relating to the effectiveness of non-invasive diagnostic testing of fetal Rh D status. The 2021 update has provided studies that impact on the evidence base through consolidating non-invasive techniques with high sensitivity, specificity and diagnostic accuracy.

If and how the 2021 search has created changes in the evidence?

The 2021 update does not change any of the findings from the evidence base. The additional studies have outcomes and findings similar to that of the previous search. Questions remain regarding the true effectiveness of NIPT on patient-relevant outcomes (i.e. the incidence of Rh D sensitisations or HDFN.

Question 4 - Risk of failure of Rh D immunoprophylaxis due to increased BMI

Question 4 – (Prognostic)

In Rh D negative pregnant or postpartum women with no preformed anti-D, does increasing BMI increase the risk of failure of Rh D immunoglobulin administration?

Background

The 2018 review identified two Level II studies (MacKenzie, 2004, Woelfer, 2004) and two Level III studies (Bichler J., 2003, Koelewijn, 2009) that provided some evidence relating maternal body weight to Rh D immunoglobulin administration.

Summary of evidence

The 2021 update found one additional Level III study (Wikman, 2021) that retrospectively examined the proportion of women with undetectable levels of prophylactic Rh D immunoglobulin at the time of delivery after RAADP (single dose of 1500 IU at28-29 gestational weeks'). It was noted that 16.5% had BMI > 30 and 4.4% had BMI > 35.

Results

During the retrospective study period (Oct 2010 to Oct 2012), Wikman 2021 found there were 876 (20.5%) cases among 4280 Rh D negative women carrying an RHD positive fetus in which the antibody screen result was negative (i.e., not detectable at delivery). In the prospective cohort, 7/39 (18%) women did not have detectable levels of anti-D at screening (38 gestational weeks), and in 10/39 (26%), the anti-D levels were below the lower limit of quantification.

After administration of the second dose at 38 gestational weeks', the mean increase in anti-D concentration (IU/mL) was 0.066 (SD 0.045) and showed a significant correlation with body mass index (p = 0.0118). The authors noted a large interindividual variation of anti-D concentration at delivery, which is suggested to depend on individual IgG clearance from plasma and consumption of anti-D, giving a variability in residual anti-D levels and in half-life. Uptake from muscular compartments and fat tissue may vary as well.

The incidence of FMH was analysed after delivery and the results were negative in all 25 of 39 (64%) patients tested (i.e., test result was below the limit of detection being 1 ml fetal blood in maternal circulation). Data were missing for 14/39 (36%) patients.

Discussion

If and how the 2021 search has impacted on evidence base?

The 2021 search provided one additional cohort study that show a correlation between anti-D levels and BMI. It enhances the evidence relating to the proportion of RhD negative pregnant women at risk of Rh D sensitisation with no detectable anti-D at delivery, despite RAADP

If and how the 2021 search has created changes in the evidence?

The 2021 update does not change any of the findings from the evidence base.

Appendix A Literature search results

This appendix documents the literature search strategy for a systematic review on the prophylactic use of Rh D Immunoglobulin (Anti-D) in pregnant women.

A search strategy to address all questions was developed via Ovid for both Embase and MEDLINE. An additional search for studies reporting diagnostic accuracy specific to subquestion 3 was also conducted. Both search strategies were then translated for PubMed (limited to in-process citations and citations not indexed in MEDLINE) and CINAHL.

A1 Questions 1 to 4

Embase

Table A.1	Search results Questions 1 to 4: Embase	(via Ovid) for Level I. Level II and Level III studies

#	Searches	Results ^a 19 July 2018	Results ^b 27 Sept 2021
1	exp "obstetrics"/ or exp "obstetric care"/ or exp "pregnancy"/ or exp "pregnancy disorder"/ or exp "prenatal disorder"/	1138534	1224786
2	(obstetric or obstetrics or pregnancy or maternal).ti,ab,kw.	688903	798197
3	(prepartum or pre partum or pre-partum or intrapartum or intra partum or intra-partum or perinatal or peri natal or peri-natal).ti,ab,kw.	98059	115994
4	(antenatal or ante natal or ante-natal or prenatal or pre natal or pre-natal).ti,ab,kw.	151232	178341
5	(postnatal or post natal or post-natal or postpartum or post partum or post-partum).ti,ab,kw.	194295	231147
6	1 or 2 or 3 or 4 or 5	1434130	1566408
7	exp "fetus"/	189819	199261
8	(fetu* or fetal* or f?etu* or f?etal*).ti,ab,kw.	375459	416482
9	7 or 8	428990	466603
10	exp alloimmunization/	4373	5319
11	exp Rh Isoimmunization/	1604	1500
12	(Rh* Isoimmuni?ation or Rh* D Isoimmuni?ation).ti,ab.	719	588
13	(Rh* alloimmuni?ation or Rh* D alloimmuni?ation).ti,ab.	381	456
14	(Rh* incompatibility or Rh* D incompatibility or blood group incompatibility).ti,ab.	1102	1022
15	(((Rh* adj3 incompatib*) or Rh* D) adj3 incompatibl*).ti,ab.	203	194
16	((Rh or RhD or rhesus) adj5 sensiti*).ti,ab.	1325	265
17	((fetomaternal or feto-maternal or foetomaternal or foeto-maternal) adj2 immuni?ation).ti,ab.	81	54
18	((rh or RhD or rhesus) adj2 (immuni?ation or autoimmuni?ation)).ti,ab.	862	541
19	10 or 11 or 12 or 13 or 14 or 15 or 16 or 17 or 18	9257	8692
20	exp rhesus D antigen/	1158	1532
21	rhesus D antigen.ti,ab.	55	57
22	rh* D antigen.ti,ab.	234	241
23	(RhD or rhesus D or Rh?D or Rh-?D or Rh D).ti,ab.	7552	9309
24	(Rh-negative or Rh-positive).ti,ab.	1312	1376
25	(Rhesus negative or Rhesus positive).ti,ab.	362	382
26	((rh or rhesus) adj2 (factor or factors or antigen\$ or system or group)).ti,ab.	4806	5015
27	20 or 21 or 22 or 23 or 24 or 25 or 26	12535	14306
28	(Macaca mulatta or Simian Immunodeficiency Virus or zika or macaque?).ti,ab.	38080	48547
29	27 not 28	12360	14114
30	(isoimmuni?ation or alloimmuni?ation).ti,ab,kw.	5921	6513
31	(isoimmuni* or iso-immuni* or isoimmune or iso-immune).ti,ab,kw.	2122	1570
32	(alloimmuni* or allo-immuni* or alloimmune or allo-immune).ti,ab,kw.	11072	13031
33	(unsensiti?ed or un-sensiti?ed or non-sensiti?ed).ti,ab,kw.	2409	2617

36 30 or 31 or 20 or 33 or 34 132314 148922 36 xep Exprimediations, Felavi 1103 265 37 ((erythrobiations), reflavi) 1103 265 38 ((Frandytic desaer' or 1/handytic desorder') is, ab, w. 1103 265 39 ((Frandytic desaer' or 1/handytic desorder') is, ab, w. 1169 1533 40 36 or 37 or 38 or 39 14943 12898 41 6 or 5 or 19 or 29 or 36 or 40 17072755 15568775 42 exp Rho D Immuno Globulin/ 3931 4222 43 op Rho D Immuno Globulin/ 3831 4222 44 exp Table Ormano Globulin/ 3831 422 45 rob arth Ormano Globulin/ 3831 425 46 Rho D Immuno Globulin/ 381 425 47 frimmuno/globulin or fri dimmuno/globulin b, ab. 574 425 48 christian and arthody distian 111 111 50 exp rhesus D anthody fi, ab. 110 111 50 rof 1 or 52 or 53	34	(sensiti?ation* or sensiti?ed).ti,ab,kw.	119508	134779
36 app Expthrobistoss, relat/ 11406 9270 37 ((erythrobistoss, relat/) 1103 265 38 (In/Rem/yic desses or ent/rem/yic disorder'), to, b, w. 1204 4940 38 G or 30 ar 30 ar 39 14193 1208 41 6 or 9 or 19 or 29 or 30 or 40 1702766 1208 42 app Rho Dimmunoglobulin/ 3931 4252 43 app Rho Dimmunoglobulin/ 3931 4252 44 app Rho Dimmunoglobulin/ 3931 4252 45 app Rho Dimmunoglobulin/ 3931 4252 46 RP in munoglobulin or n'd immunoglobulin/ b, b. 310 322 47 (h'mmunoglobulin or n'd immunoglobulin) b, b. 331 4252 50 or prinsus D antbody f, b. 331 4252 51 rhesus D antbody f, b. 111 11 52 (h'D antbody or n'D antbody b) a. 108 107 53 sap reflex D antbody f, b. 108 107 54 60 or 51 or 52 or 53 60 or 5	35	30 or 31 or 32 or 33 or 34	132314	148922
37 ((e)ryhodulations or eyrhodulations) add (left) (i.j.a), kw. 1103 268 38 (f)römnölic disses" or h7ömölic disorder") i.j.a), kw. 1109 1533 40 36 or 37 or 38 or 39 14943 12886 41 6 or 57 or 38 or 39 14943 12886 41 6 or 57 or 38 or 39 14943 12886 41 6 or 57 or 38 or 39 14943 12886 41 6 or 57 or 38 or 39 14943 12886 41 6 or 57 or 38 or 39 14922 1885 42 exp RhD Immuno Globulin7 3831 4252 44 exp RhD Immuno Globulin7 3831 4252 45 exp arb D Immuno Globulin 1, ab. 130 32 46 Rh O Immuno Globulin 1, ab. 131 11 50 or phress D antbody 1, ab. 111 11 51 11 11 11 11 52 11 11 11 11 52 11 128 11 11 11	36	exp Erythroblastosis, Fetal/	11405	9270
38 (Promolytic disease" or h?emolytic disorder") h, ab, w. 5204 4940 39 (HD1 N or HDN) h, ab, w. 1168 1533 41 6 or 3 or 3 or 3 or 3 or 3 or 4 11286 11286 42 exp Fb D Immuno Globulin / 3331 4252 43 exp Fb D Immuno Globulin / 3331 4252 44 exp Fb D Immuno Globulin / 3331 4252 45 exp Fb D Immuno Globulin / 3331 4252 46 RP D Immuno Globulin / 3331 4252 47 dr * dr or 4 do or 4 do or 4 do or 4 do r 4 do	37	((erythroblastoses or erythroblastosis) adj2 (fetal* or f?etal*)) ti,ab,kw.	1103	266
39 [HOPN or HON] is allow. 1169 1533 40 36 or 37 or 38 or 39 14943 1229 41 6 or 5 or 19 or 29 or 36 or 40 11702755 1866675 42 exp Rh Dimmune Globulin? 3331 4252 43 exp Rh Dimmune Globulin? 3331 4252 44 exp Rh Dimmune Globulin? 3331 4252 45 exp and Dimmune Globulin? 3331 4252 46 Rb Dimmune Globulin ja.b. 33 88 47 (h* immun2dobulin or h* dimmun2dobulin ja.b. 310 332 48 (h* immun2dobulin or h* dimmun2dobulin ja.b. 311 111 50 exp rhesus D antibody 3331 4252 51 frees and antibody ja.b. 1108 107 53 (h* Dantibody is.b. 1108 107 54 60 of 51 of 20 or 83 6309 6674 55 exp rhogan/ 3931 4252 56 exp rhogan/ 3931 4252 56 exp rhogan/ 3931 4252 56 exp rhogan/	38	(h?emolytic disease* or h?emolytic disorder*).ti,ab,kw.	5204	4940
40 36 or 37 or 38 or 39 14943 12988 11 6 or 9 ar 19 or 25 or 45 11956/5 41 6 or 97 or 15 or 47 11956/5 42 exp Rho D Immune Globulin* 3931 4252 43 exp Rho D Immune Globulin* 3931 4252 44 exp Rho D Immune Globulin* 3931 4252 45 exp ant-D Immune Globulin* hab 310 332 46 Rh D Immune Globulin hab 310 332 47 (fr 4 minune/ation ft 4 immune/ation [h, ab. 574 425 49 42 or 43 or 46 or 46 or 46 or 47 or 48 4584 4753 50 exp rhosus D antbody (h, ab. 11 11 11 51 dness D antbody (h, ab. 168 107 53 exp rhogan / ab. 4652 5158 54 50 or 51 or 52 or 53 6399 6974 55 exp rhogan / ab. 47 46 56 ehogan / ab. 47 46 57 exp winch 3931 4252 58 exp rhogan / ab. 54 <td< td=""><td>39</td><td>(HDFN or HDN).ti,ab,kw.</td><td>1169</td><td>1533</td></td<>	39	(HDFN or HDN).ti,ab,kw.	1169	1533
11 6 or 9 or 10 or 29 or 35 or 40 1702755 1858675 12 exp Rh: D Immuno Globulin/ 3931 4252 13 exp Rh: D Immuno Globulin/ 3931 4252 14 exp Rh: D Immuno Globulin/1 3931 4252 15 exp Rh: D Immuno Globulin/1 3931 4252 16 Rh: To Immuno Globulin/1 3931 4252 16 Rh: To Immuno Globulin/1, gab. 574 425 16 (h' immuno Globulin 1, gab. 574 425 17 (h' immuno Globulin) 1, gab. 574 425 18 (h' immuno Globulin) 1, gab. 11 11 11 11 11 11 11 12 (h' immuno Globuly) 1, gab. 100 107 16 exp rhogan/ 3931 4252 16 for 50 or 51 or 52 or 53 6309 6374 16 exp rhogan/ 3931 4252 16 hypositic 516 516 516 16 exp rhogan/ 3931 4252 17 exp hogan/ 3931	40	36 or 37 or 38 or 39	14943	12898
42 exp Rho D Immune Globulin/ 452 43 exp Rho D Immune Globulin/ 3931 4252 44 exp Rho D Immune Globulin/ 3931 4252 45 exp ant-D Immune Globulin/ 1, ab. 3931 4252 46 exp ant-D Immune Globulin 1, ab. 3931 4252 47 (n' mmune/alcoulin 1, ab. 310 332 48 (n' mmune/alcoulin 1, ab. 574 425 49 42 or 43 or 46 or 46 or 47 or 48 4584 4753 50 exp rhesus D antibody 1, ab. 11 11 11 51 dec nd 3 or 46 or 47 or 48 6454 4753 50 exp rhesus D antibody 1, jab. 108 107 51 dent-D or anti D or ant/D0, jab. 6309 6374 55 exp rhophysica 6331 4252 56 exp rhophysica 641 65 57 exp winch b, ab. 21 21 58 winch b, ab. 21 21 21 58	41	6 or 9 or 19 or 29 or 35 or 40	1702755	1858675
13 exp Rho D Immune Globulin/ 147 146 107 11 1	42	exp Rh D immunoglobulin/	3931	4252
44 exp TbhO[) Immune Globulin? 3331 4252 45 exp anb.D Immune Globulin is ab. 3331 4252 46 fr.D Immune Globulin is ab. 310 332 47 (rh' immune Globulin is ab. 310 332 48 (rh' immune Globulin or rh' d immune/abon) is ab. 574 425 49 40 ar 4 ar 46 ar 46 or 47 or 48 4584 4763 50 exp rhesus D anibody [ab. 111 11 51 rh are on anibody [ab. 108 107 53 (nh' D anibody or nh'D anibody] 54. 4252 5158 54 50 or 51 or 52 or 53 6309 6974 4252 56 exp rhogan/ 3931 4252 4252 58 wintho (nab. 64 65 59 59 59 59 59 531 4252 64 65 50 50 50 50 50 50 50 50 50 50 50 50 50 50 50<	43	exp Rho D Immune Globulin/	3931	4252
45 exp arti-D immunoglobulin/ 3831 4252 46 Rit D Immunoglobulin r, fab. 93 88 47 (fr. immunoglobulin or fr. d. immunoglobulin.) i, ab. 310 332 48 (fr. immunoglobulin or fr. d. immunoglobulin.) i, ab. 574 425 49 42 or 43 or 44 or 45 or 46 or 47 or 48 4584 4783 50 exp freesus D antibody 3831 4252 51 rhssus D antibody fs.ab. 108 107 52 (rh.*) D antibody or rh*D antibody.) i, ab. 108 107 53 (rh.*) D antibody or rh*D antibody.) i, ab. 4552 5158 54 50 or 51 or 52 or 53 6309 6974 55 exp rhogam 3931 4252 56 exp winrho.1 3931 4252 58 winrho.1, ab. 64 65 59 exp rhogam 3931 4252 58 winrho.1, ab. 1331 4252 59 exp rhogam 3931 4252 50 <td>44</td> <td>exp "Rho(D) Immune Globulin"/</td> <td>3931</td> <td>4252</td>	44	exp "Rho(D) Immune Globulin"/	3931	4252
46 Rh* D Immune Globulin Is ab. 30 332 47 (h* immunoglobulin or h* d immunoglobulin) is ab. 310 332 48 (h* immunoglobulin or h* d immunoglobulin) is ab. 574 425 49 42 or 43 or 44 or 45 or 46 or 47 or 48 4584 4753 50 exp rhesus D antibody [s.b. 111 11 51 rhsus D antibody [s.b. 108 107 53 (nh* D or ant D or ant/D) [s.b. 4652 5158 54 50 or 51 or 52 or 53 6309 6974 55 exp rhogan 1 3931 4252 56 hogan 1, ab 47 46 57 exp rindpland 3931 4252 58 winnto 1, ab 64 65 59 exp rhoghylac/ 3931 4252 58 winnto 1, ab 21 21 21 51 exp dimblands. 64 65 3331 4252 58 winnto 1, ab 3331 4252 26 45 3331 4252 53 exp thesphylac/ 3331 <td>45</td> <td>exp anti-D immunoglobulin/</td> <td>3931</td> <td>4252</td>	45	exp anti-D immunoglobulin/	3931	4252
47 (th* immun?aloin or th* d immun?aloin jt,ab. 310 332 48 (th* immun?aloin or th* d immun?aloin jt,ab. 574 425 50 exp rhesus D antibody/ 3831 4252 51 nhesus D antibody/ 3831 4252 51 nhesus D antibody/ 3831 4252 51 fibresus D antibody/ 108 107 53 (ant-D or anti D) rant/D) it,ab. 4652 6758 54 50 or 51 or 52 or 53 6309 6974 55 exp rhogam/ 3931 4252 56 hogam it,ab. 47 46 59 exp rhoghylac/ 3931 4252 60 hophylac/i.ab. 21 21 61 exp throkofam/ 3931 4252 62 exp banyRito-D/ 3931 4252 63 exp thresonaliv/ 3931 4252 64 exp thresonaliv/ 3931 4252 65 exp thresonaliv/ 3931 4252 64 exp wintho/ 39331 4252 <	46	Rh* D Immune Globulin.ti,ab.	93	88
48 (h* immun?ation or h* d immun?ation) it,ab. 574 425 49 42 or 43 or 44 or 45 or 46 or 47 or 48 4763 50 exp thesus D antibody (r.ab. 3931 4252 51 rhesus D antibody (r.ab. 11 11 52 (rh* D antibody (r.ab.) 108 107 53 (am: D or anti?D) (r,ab. 4652 5158 54 50 or 51 or 52 or 53 6309 6974 55 exp thogam // 3931 4252 56 hogam // 3931 4252 58 winthol, rab. 64 65 59 exp thoghylac/ 3931 4252 50 exp thoghylac/ 3931 4252 59 exp thoghylac/ 3931 4252 50 map MicRhoGam/ 3931 4252 51 exp MicRhoGam/ 3931 4252 50 for phylac/ fab 3931 4252 51 exp MicRhoGam/ 3931 4252 52 exp MicRhoGam/ 3931 4252 54 for 56 o	47	(rh* immunoglobulin or rh* d immunoglobulin).ti,ab.	310	332
49 42 or 43 or 44 or 45 or 46 or 47 or 48 4564 4753 50 exp thesus D antibody 3931 4252 51 hesus D antibody 11 11 52 (h* D antibody or h*D antibody) j,ab 108 107 53 (anti-D or ant) 07 ant) 70 thab. 4652 5158 54 60 or 51 or 52 or 53 6309 6974 55 exp hingam, ab. 47 46 56 hogam, ab. 47 46 57 exp winhol 3931 4252 58 winhol, j,ab. 64 65 59 exp thoghylac/ 3931 4252 61 exp MicRho-Cann 3931 4252 62 exp thoghylac/ 3931 4252 63 exp thosonaliv/ 3931 4252 64 RND immunglobulin vfi,ab. 0 0 65 56 or 57 or 88 or 99 or 60 or 61 or 62 or 63 or 64 3965 5152 68 exp thosonaliv/ 3931 4252 5152 68 exp thosonaliv/ or comparative study wing or e	48	(rh* immuni?ation or rh* d immuni?ation).ti,ab.	574	425
50 exp rhesus D anibody/ 3331 4252 51 mesus D anibody of the D anibody (b, ab. 11 11 11 52 (hT) D anibody of the D anibody (b, ab. 108 107 53 (ani-D or anti D or anti/D) (b, ab. 4652 5158 54 50 or 51 or 52 or 53 6309 6974 55 exp rhegam/ 3331 4252 56 fmogan b, ab. 47 46 58 winch b, ab. 64 65 59 exp hophylac/ 3931 4252 60 hopphylac b, ab. 21 21 61 exp Bic/Rho-D/ 3931 4252 62 exp Bic/Rho-D/ 3931 4252 63 exp Hos/RHo-D/ 3931 4252 64 RbD immunoglobuli nvf b, ab. 0 0 0 65 sor 56 or 56 or 50 or 50 or 60 or 61 or 62 or 63 or 64 3952 4257 66 49 or 54 or 65 6855 7163 67 11 and 66	49	42 or 43 or 44 or 45 or 46 or 47 or 48	4584	4753
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52 (h* D antibody or h*D antibody) li,ab. 108 107 53 (ant-D or ant D or ant/D) li,ab. 4652 5158 64 50 or 51 or 52 or 53 6309 6974 56 exp thogam/ 3331 4252 56 thogam li,ab. 47 46 57 exp winthol 3931 4252 58 winch ti,ab. 64 65 59 exp MICRhoGam/ 3931 4252 60 rhophylac/ 3931 4252 61 exp Bi/RN-b1/ 3931 4252 62 exp Bi/RN-b1/ 3931 4252 63 exp MiCRhoGam/ 3931 4252 64 RbD immunoglobulin vf. fi,ab. 0 0 65 to 75 or 58 or 59 or 60 or 61 or 62 or 63 or 64 3952 4267 66 49 or 54 or 65 6855 7133 67 41 and 66 4985 51152 exp comparative study or oreadowir/set on np, or exp single bind procedure m, or exp consoler work or exp consoler mp, or exp consoler sindy or exp consoler mp, o	51	rhesus D antibody.ti,ab.	11	11
53 (anti-D or anti PD) ti,ab. 4652 5158 54 50 or 51 or 52 or 53 6309 6974 55 exp rhogam/ 3931 4252 56 exp winthol 3931 4252 58 winrho ti,ab. 64 65 59 exp rhophylac/ 3931 4252 58 winrho ti,ab. 64 65 59 exp rhophylac/ 3931 4252 60 rhophylac. ti,ab. 21 21 61 exp MICRhoGam/ 3931 4252 62 exp BayRHo-D/ 3931 4252 63 exp rhesonativ/ 3931 4252 64 rbsD immunoglobulin vft i,ab. 0 0 65 or 58 or 59 or 60 or 61 or 62 or 63 or 64 6855 7363 67 41 and 66 6855 7363 5152 68 exp meta analysis/ or meta analysis mp. or exp cinical trial/ or clinical trial mp. or andomized controlled trial/ mp. or exp triple blind procedurer mp. or exp triple blind mp. or (exp prospective st	52	(rh* D antibody or rh*D antibody).ti,ab.	108	107
54 50 or 51 or 52 or 53 6309 6974 55 exp rhogam/ 3931 4252 56 rhogam li, ab. 47 46 57 exp winrho 3931 4252 58 winrho li, ab. 64 65 59 exp rhophylac/ 3931 4252 60 rhophylac li, ab. 21 21 61 exp MICRhoCam/ 3931 4252 62 exp BayRHo-D/ 3931 4252 63 exp rhesonativ/ 3931 4252 64 RhD immuoglobulin vf. it, ab. 0 0 65 r5 or 56 or 57 or 58 or 57 or 58 or 57 or 58 or 57 or 58 or 51 or 52 or 63 or 64 3952 4267 66 49 or 54 or 65 6855 7363 67 41 and 66 6855 7363 68 exp omparative study or omparative study mp. or exp randomized controlled trial. mp. or randomization or randomization mp. or randomization mp. or exp or parative study mp. or exp randomization or exp randomization or exp randomization or exp randomization or exp randomizatior or exp randomizatior or exp randomization or exp random	53	(anti-D or anti D or anti?D).ti,ab.	4652	5158
55 exp rhogan/ 3931 4252 56 rhogam i, ab. 47 46 57 exp winrho/ 3331 4252 58 winrho (ab. 64 65 59 exp rhophylac/ 3931 4252 60 rhophylac (i, ab. 21 21 61 exp MICRhoCam/ 3931 4252 62 exp BayRHo-D/ 3931 4252 63 exp rhesonaliv/ 3931 4252 64 (FkD) immunoglobulin vf fi, ab. 0 0 65 55 or 55 or 57 or 58 or 59 or 69 or 60 or 61 or 62 or 63 or 64 3952 4267 66 49 or 54 or 65 6855 7363 67 41 and 66 4895 5152 68 exp meta analysis mp. or exp systematic review/ or systematic review/ or exp and/miced or exp randomized controlled final, pp. or randomi2ed controlled final, pp. or exp and/miced or por or shop or 60 or 61 or 62 or 63 or 64 4955 69 procedure, or exp or 60 or 61 or 62 or 63 or 64 3952 4207 69 procedure, or exp catodinated or por trebied final, pp. or randomi2ed controled trial/ pr. or exp randomized controlled trial/ pp. or	54	50 or 51 or 52 or 53	6309	6974
56 rhogam ti,ab. 47 46 57 exp winrho/ 3331 4252 58 winrho ti,ab. 64 65 59 exp hophylac/ 3331 4252 60 rhophylac ti,ab. 21 21 61 exp MRCRhoCam/ 3331 4252 62 exp BayRHo-D/ 3931 4252 63 exp resonativ/ 3331 4252 64 'TRD immunoglobulin vf' ti,ab. 0 0 65 of 56 or 57 or 58 or 59 or 60 or 61 or 62 or 63 or 64 3952 4267 66 49 or 54 or 65 6855 7363 67 67 41 and 66 4895 5152 68 68 exp mela analysis/ or mela analysis.mp. or exp systematic review/ or systematic review.mp. or pooled analysis.mp. or (exp review/ or review.mp.) and (systemat' or pool").mp. 424029 643063 69 exp clinical trial, or or randomi?ation.mp. or randomi?ation.mp. or exp sigle blind procedure/ or onspecified blind procedure/ or onspecified blind procedure/ or onspecified blind procedure/ or or sover procedure/ or onspecified blind procedure/ or or sover procedure/ or onspecified blind procedure/ or onspecified blind procedure/ or onspecified blind procedure/ or or sover pro	55	exp rhogam/	3931	4252
57 exp winrho/ 3931 4252 58 winrho.it,ab. 64 65 59 exp rhophylac/ 3931 4252 60 rhophylac.it,ab. 21 21 21 61 exp BayRHo-D/ 3931 4252 62 exp BayRHo-D/ 3931 4252 63 exp rhesonativ/ 3931 4252 64 'RhD immunoglobulin vf.ti,ab. 0 0 65 55 or 56 or 57 or 58 or 59 or 60 or 61 or 62 or 63 or 64 3952 4267 66 49 or 54 or 65 6855 7363 67 41 and 66 4895 5152 68 exp review/ or review.mp.) and (systemat' or pool').mp.) 424029 643063 69 procedure mp. or exp duble blind procedure/ or orangle blind procedure/ or single blind 3989862 4922439 61 spp procedure/ or triple blind procedure/ or duble blind procedure/ or consover procedure/ or orangle blind 3989862 4922439 70 reforspective study/ or exp control study/ or exp family study/ or exp ingle blind mp. or aduble blind procedure/ or single blind 9057811 11236854 71	56	rhogam.ti,ab.	47	46
58 wintho.ti,ab. 64 65 59 exp rhophylad/ 3931 4252 60 rhophylac/ti,ab. 21 21 61 exp MiCRhoGam/ 3931 4252 62 exp BayRHo-D/ 3931 4252 63 exp rhesonativ/ 3931 4252 64 RFND immunoglobulin vf ti,ab. 0 0 0 65 55 or 56 or 59 or 59 or 60 or 61 or 62 or 63 or 64 3952 4267 66 49 or 54 or 55 7363 67 41 and 66 4895 5152 68 exp mela analysis/ or mela analysis.mp. or exp systematic review/ or systematir or pool").mp.) 424029 643063 69 procedure mp. or endom?ation mp. or exp andomized controlled trial mp. or random?ation mp. or exp isingle bind mp. or exp procedure/ or single bind procedure/ or single bind procedure/ or single bind procedure/ or single bind procedure/ or exp prospective study/ or exp case control study' or	57	exp winrho/	3931	4252
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60 rhophylac.li,ab. 21 21 21 61 exp MICRhoGam/ 3931 4252 62 exp hesonativ/ 3931 4252 63 exp thesonativ/ 3931 4252 64 RthD immunoglobulin vf.1,ab. 0 0 65 55 or 56 or 57 or 58 or 59 or 60 or 61 or 62 or 63 or 64 3952 4267 66 49 or 54 or 65 6855 7363 67 41 and 66 4895 5152 68 exp meta analysis/or meta analysis.mp. or exp systematic review/mp.or pool").mp.) 424029 643063 69 exp comparative study or comparative study mp. or exp candomized controlled trial.mp. or randomized controlled trial.mp. or randomized controlled trial.mp. or randomized controlled trial.mp. or randomized controlled trial.mp. or exp andomized controlled trial.mp. or exp andomized controlled trial.mp. or exp triple blind procedure/ or sup analysis.mp. or (exp review/ or prospective study/ or prospective study/ or exp contanatysis or or tot.mp. or single blind procedure/ or exp analysis.mp. or (exp review study/ or exp contanatysis or or tot.mp. or single blind mp. or exp triple blind mp. or exp analysis.mp. or (exp explexed study/ or exp contanatysis.mp.) or (follow up adj1 stud').mp. or (baservational adj1 stud').mp. or (follow up adj1 stud').mp. or (cobservational adj1 stud').mp. or (fobservational adj1 stud').mp. or (fo	59	exp rhophylac/	3931	4252
61 exp MICRhoGam/ 3931 4252 62 exp BayRHo-D/ 3931 4252 63 exp rhesonativ/ 3931 4252 64 'RhD immunoglobulin vf.ti,ab. 0 0 65 55 or 56 or 57 or 58 or 59 or 60 or 61 or 62 or 63 or 64 3952 4267 66 49 or 54 or 65 6855 7363 67 41 and 66 4895 5152 68 exp meta analysis.mp. or review.mp. or exp systematic review or systematic review.mp. or pooled analysis.mp. or ((exp review/ or review.mp.) and (systemat' or pool").mp.) 424029 643063 69 exp comparative study or comparative study mp. or exp chinical trial or clinical trial mp. or randomization.mp. or randomi?ed controlled trial.mp. or exp single blind procedure.mp. or exp triple blind procedure.mp. or exp double blind procedure.mp. or exp triple blind procedure.mp. or exp double blind procedure.mp. or exp triple blind procedure.mp. or exp double blind procedure.mp. or exp triple blind mp. or double blind mp. or double blind procedure.mp. or exp triple blind mp. or exp prospective study/ or exp contra analysis.mp. or (follow up adj1 stud*).mp. or (exp prospective study/ or exp contra analysis.mp.) or (follow up adj1 stud*).mp. or (exp prospective study/ or exp contra analysis.mp.) or (follow up adj1 stud*).mp. or (observational adj1 stud*).mp. or (epidemiologic* adj1 stud*).mp. or (exp sereport*/ 2319441 2688800 <td>60</td> <td>rhophylac.ti,ab.</td> <td>21</td> <td>21</td>	60	rhophylac.ti,ab.	21	21
62 exp BayRHo-D/ 3931 4252 63 exp rhesonativ/ 3931 4252 64 TkD immunoglobulin vf. ti, ab. 0 0 65 55 or 56 or 57 or 58 or 59 or 60 or 61 or 62 or 63 or 64 3952 4257 66 49 or 54 or 65 6855 7363 67 41 and 66 4895 5152 68 exp meta analysis/ or meta analysis.mp. or exp systematic review/ or systematic review.mp. or pooled analysis.mp. or (lexp review/ or review.mp.) and (systemat' or pool").mp.) 424029 643063 69 exp comparative study/ or comparative study.mp. or exp clinical trial / or clinical trial mp. or randomized controlled trial mp. or exp andomized controlled trial mp. or randomized controlled trial mp. or exp andomized controlled trial mp. or randomized controlled trial mp. or randomized controlled trial mp. or randomized controlled trial mp. or exp ingle blind procedure.mp. or exp placebo' or placebo' mp. or randomized mp. or exp cassover procedure.mp. or exp prospective study/ or exp case control study/ or exp family study' or exp longitudinal study' mp. or (exp sectional adj1 stud*) mp. or (cose sectional adj1 stud*) mp. or (observational adj1 stud*) mp. or (cores sectional adj1 stud*) mp. or (observational adj1 stud*) mp. or (cores sectional adj1 stud*) mp. or (observational adj1 stud*) mp. or (cose sectional adj1 stud*) mp. or (baservational adj1 stud*) mp. or (cose secontrol adj1 stud*) mp. or (observatio	61	exp MICRhoGam/	3931	4252
63 exp thesonativ/ 3931 4252 64 'RhD immunoglobulin vf'.ti,ab. 0 0 65 55 or 56 or 57 or 58 or 59 or 60 or 61 or 62 or 63 or 64 3952 4267 66 49 or 54 or 65 6855 7363 67 41 and 66 4895 5152 68 exp meta analysis/ or meta analysis.mp. or exp systematic review/ or systematic review.mp. or pooled analysis.mp. or ((exp review/ or review.mp.) and (systemat* or pool*).mp.) 424029 643063 69 exp comparative study/ or comparative study.mp. or exp clinical trial.mp. or randomized controlled trial.mp. or randomi?ed controlled trial.mp. or exp single blind procedure or single blind procedure or triple blind procedure mp. or exp consover procedure.mp. or exp placebo' or placebo' mp. or randomi *mp. or rtc.mp. or single blind mp. or double blind mp. or double blind procedure mp. or exp family study/ or exp prospective study/ or exp case control study/ or exp family study/ or exp prospective study/ or exp case control study/ or exp family study/ or exp longitudinal stud*).mp. or (observational adj1 stud*).mp. or (epidemiologic* adj1 stud*).mp. or (aditorial or letter or comment or historical article).pt. 11236854 71 "case report*/	62	exp BayRHo-D/	3931	4252
64 'RhD immunoglobulin vf.ti,ab. 0 0 65 55 or 56 or 57 or 58 or 59 or 60 or 61 or 62 or 63 or 64 3952 4267 66 49 or 54 or 65 6855 7363 67 41 and 66 4895 5152 68 exp meta analysis/or meta analysis.mp. or exp systematic review/ or systematic review.mp. or pooled analysis.mp. or ((exp review/ or review.mp.) and (systemat' or pool').mp.) 424029 643063 69 exp comparative study/ or comparative study.mp. or exp clinical trial/ or clinical trial/ or exp randomization/ or randomi?ed controlled trial mp. or exp single blind procedure/ or single blind procedure mp. or exp double blind procedure/ or double blind procedure/ or osingle blind procedure or triple blind procedure/ or double blind procedure/ or cossover procedure/ or exp placebo' or placebo' mp. or random*mp. or rct.mp. or single blind.mp. or exp triple blind procedure study/ or exp case control study/ or exp family study/ or exp longitudinal study/ or exp retrospective study/ or exp cose control study/ or exp family study/ or exp longitudinal study/ or exp retrospective study/ or exp contort analysis or (cohort analysis or (follow up adj1 stud*).mp. or (observational adj1 stud*).mp. or (epidemiologic* adj1 stud*).mp. or (observational adj1 stud*).mp. or (epidemiologic* adj1 stud*).mp. or (observational adj1 stud*).mp. or (epidemiologic* adj1 stud*).mp. or (observational adj1 stud*).mp. or (follow up adj1 stud*).mp. 9057811 11236854 71 *case report*/ 2319441 2658800	63	exp rhesonativ/	3931	4252
65 55 or 56 or 57 or 58 or 59 or 60 or 61 or 62 or 63 or 64 3952 4267 66 49 or 54 or 65 6855 7363 67 41 and 66 4895 5152 68 exp meta analysis/ or meta analysis.mp. or exp systematic review/ or systematic review.mp. or pooled analysis.mp. or ((exp review/ or review.mp.) and (systemat* or pool").mp.) 424029 643063 68 exp comparative study/ or comparative study.mp. or exp clinical trial or clinical trial.mp. or randomized controlled trial.mp. or randomi2ation.mp. or randomi2ation.mp. or exp single blind procedure mp. or exp double blind procedure.mp. or exp single blind procedure.mp. or exp double blind procedure.mp. or exp single blind.mp. or triple blind.mp. or triple blind.mp. or triple blind.mp. or triple blind.mp. or double blind.mp. or double blind.mp. or trethe blind.mp. or triple blind.mp. or triple blind.mp. or exp placebo/ or placebo*.mp. or randomi?ation.inp. or (cloave prospective study/ or exp conspective study/ or exp conspective study/ or exp contor adj1 stud*).mp. or (exp prospective study/ or exp case control study/ or exp family study/ or exp longitudinal stud*).mp. or (observational adj1 stud*).mp. or (epidemiologic* adj1 stud*).mp. or (observational adj1 stud*).mp. or (epidemiologic* adj1 stud*).mp. or (cleatorial or letter or comment or historical article) pt. 11236854 71 *case report*/ 2319441 2658800 72 (editorial or letter or comment or historical article) pt. 1600355 1895537 73	64	'RhD immunoglobulin vf.ti,ab.	0	0
66 49 or 54 or 65 6855 7363 67 41 and 66 4895 5152 68 exp mela analysis/ or meta analysis. or exp systematic review/ or systematic review.mp. or pooled analysis.mp. or (exp review/ or review.mp.) and (systemat* or pool*).mp.) 424029 643063 68 exp comparative study/ or comparative study.mp. or exp clinical trial/ or clinical trial.mp. or randomized controlled trial.mp. or randomi2ed controlled trial.mp. or randomi2ed controlled trial.mp. or randomi2ed controlled trial.mp. or exp single blind procedure/ or single blind procedure/ or triple blind procedure mp. or exp crossover procedure.mp. or exp triple blind procedure/ or triple blind procedure/ or or single blind.mp. or single blind.mp. or single blind.mp. or exp triple blind procedure mp. or exp crossover procedure.mp. or exp triple blind procedure/ or double blind procedure/ or or single blind.mp. or single blind.mp. or exp triple blind 3989862 4922439 70 exp clinical study/ or exp case control study/ or exp family study/ or exp longitudinal study/ or exp cosecutor analysis/ or (cohort adj1 stud*).mp. or (case control adj1 stud*).mp. or (observational adj1 stud*).mp. or (cross sectional adj1 stud*).mp. or (observational adj1 stud*).mp. or (comment or historical article).pt. 9057811 11236854 71 "case report"/ 2319441 2658800 2658800 72 (editorial or letter or comment or historical article).pt. 1600355 1895537 3716272 4329515 74	65	55 or 56 or 57 or 58 or 59 or 60 or 61 or 62 or 63 or 64	3952	4267
67 41 and 66 4895 5152 68 exp meta analysis/ or meta analysis.mp. or exp systematic review/ or systematic review.mp. or pooled analysis.mp. or ((exp review/ or review.mp.) and (systemat* or pool*).mp.) 424029 643063 68 exp comparative study/ or comparative study.mp. or exp clinical trial.mp. or randomized controlled trial.mp. or randomized controlled trial.mp. or randomized controlled trial.mp. or randomization.mp. or exp candomized controlled trial.mp. or exp triple blind procedure/ or single blind procedure/ or single blind procedure/ or crossover procedure mp. or exp placebo' or placebo* mp. or random*.mp. or rt.mp. or single blind.mp. or single blind.mp. or exp prospective study/ or exp case control study/ or exp family study/ or exp retrospective study/ or exp case control study/ or exp family study/ or exp longitudinal study' or exp retrospective study/ or exp cohort analysis/ or (cohort adj1 stud*).mp. or (case control adj1 stud*).mp. or (observational adj1 stud*).mp. or (epidemiologic* adj1 stud*).mp. or (cross sectional adj1 stud*).mp. 9057811 11236854 70 (editorial or letter or comment or historical article).pt. 1600355 1895537 73 71 or 72 3716272 4329515 74 (animals/ or nohuman/) not humans/ 6126128 6443842 75 (67 and 68) not (72 or 74) 69 85 76 (67 and 68) not (73 or 74 or 75) 470 542 76 (67 and 69) not (73 or 74 or 75) <td< td=""><td>66</td><td>49 or 54 or 65</td><td>6855</td><td>7363</td></td<>	66	49 or 54 or 65	6855	7363
68exp meta analysis/ or meta analysis mp. or exp systematic review/ or systematic review.mp. or pooled analysis.mp. or ((exp review/ or review.mp.) and (systemati' or pool").mp.)42402964306369exp comparative study/ or comparative study.mp. or exp clinical trial/ or clinical trial/ or exp randomization/ or randomi?ed controlled trial mp. or exp single blind procedure/ or single blind procedure.mp. or exp double blind procedure/ or double blind procedure/ or crossover procedure/ or crossover procedure/ or crossover procedure/ or double blind mp. or double blind encedure.mp. or exp triple blind.mp. or single blind.mp. or single blind.mp. or double blind.mp. or tandomi?mp. or rext finile blind.mp. or single blind.mp. or single blinded.mp. or double blind.mp. or tandomi?ed controlled trial.s.mp.) or (case control adj1 stud*).mp. or (exp prospective study/ not randomi?ed controlled trials.mp.) or (follow up adj1 stud*).mp. or (cobservational adj1 stud*).mp. or (epidemiologic* adj1 stud*).mp. or (cross sectional adj1 stud*).mp.90578111123685470reace report*/23194412658800265880071*case report*/371627243295157371 or 7237162723716272432951574(animals/ or nonhuman/) not humans/6126128644384275(67 and 68) not (72 or 74)698576(67 and 70) not (73 or 74 or 75)47054277Itimit 75 to yr="2018-current"NA27	67	41 and 66	4895	5152
exp comparative study/ or comparative study mp. or exp clinical trial/ or clinical trial.mp. or randomized controlled trial.mp. or randomi?ed controlled trial.mp. or exp randomized controlled trial/ or exp randomization/ or randomi?ation.mp. or exp randomized controlled trial/ or exp procedure/ or triple blind procedure.mp. or exp consover procedure/ or single blind procedure/ or triple blind procedure.mp. or exp consover procedure/ or crossover procedure/ or exp placebo/ or placebo*.mp. or random*.mp. or triple blind.mp. or triple blinded.mp. or double blind.mp. or double blinded.mp. or treble blind.mp. or triple blind.mp. or single blinded.mp. or double blind.mp. or double blinded.mp. or treble blind.mp. or triple blinded.mp. or exp prospective study/ or exp cohort analysis/ or (cohort adj1 stud*).mp. or (exp prospective study/ not random?ed controlled trials.mp.) or (follow up adj1 stud*).mp. or (clobservational adj1 stud*).mp. or (cerss sectional adj1 stud*).mp.90578111123685470(editorial or letter or comment or historical article).pt.160035518955377171 or 723716272432951574(animals/ or nonhuman/) not humans/6126128644384275(67 and 69) not (73 or 74 or 75)47054277(67 and 70) not (73 or 74 or 75)990122278limit 75 to yr=*2018 - Current*NA27	68	exp meta analysis/ or meta analysis.mp. or exp systematic review/ or systematic review.mp. or pooled analysis.mp. or ((exp review/ or review.mp.) and (systemat* or pool*).mp.)	424029	643063
respective study/ or exp case control study/ or exp family study/ or exp longitudinal study/ or exp retrospective study/ or exp cohort analysis/ or (cohort adj1 stud*).mp. or (case control adj1 stud*).mp. or (exp prospective study/ not randomi?ed controlled trials.mp.) or (follow up adj1 stud*).mp. or (observational adj1 stud*).mp. or (epidemiologic* adj1 stud*).mp. or (cross sectional adj1 stud*).mp.90578111123685471"case report"/2319441265880072(editorial or letter or comment or historical article).pt.160035518955377371 or 723716272432951574(animals/ or nonhuman/) not humans/6126128644384275(67 and 68) not (72 or 74)698576(67 and 69) not (73 or 74 or 75)47054277(67 and 70) not (73 or 74 or 75 or 76)990122278limit 75 to yr="2018 -Current"NA27	69	exp comparative study/ or comparative study.mp. or exp clinical trial/ or clinical trial.mp. or randomized controlled trial.mp. or randomi?ed controlled trial.mp. or exp randomized controlled trial.mp. or randomization.mp. or randomi?ation.mp. or exp single blind procedure/ or single blind procedure.mp. or exp double blind procedure/ or double blind procedure.mp. or exp triple blind procedure.mp. or exp crossover procedure/ or crossover procedure.mp. or exp placebo ⁺ or placebo ⁺ .mp. or random ⁺ .mp. or rct.mp. or single blind.mp. or single blinded.mp. or exp prospective study/ or prospective study.mp.	3989862	4922439
71 "case report"/ 2319441 2658800 72 (editorial or letter or comment or historical article).pt. 1600355 1895537 73 71 or 72 3716272 4329515 74 (animals/ or nonhuman/) not humans/ 6126128 6443842 75 (67 and 68) not (72 or 74) 69 85 76 (67 and 69) not (73 or 74 or 75) 470 542 77 (67 and 70) not (73 or 74 or 75 or 76) 990 1222 78 limit 75 to yr="2018 -Current" NA 27	70	exp clinical study/ or exp case control study/ or exp family study/ or exp longitudinal study/ or exp retrospective study/ or exp cohort analysis/ or (cohort adj1 stud*).mp. or (case control adj1 stud*).mp. or (exp prospective study/ not randomi?ed controlled trials.mp.) or (follow up adj1 stud*).mp. or (observational adj1 stud*).mp. or (epidemiologic* adj1 stud*).mp. or (cross sectional adj1 stud*).mp.	9057811	11236854
72 (editorial or letter or comment or historical article).pt. 1600355 1895537 73 71 or 72 3716272 4329515 74 (animals/ or nonhuman/) not humans/ 6126128 6443842 75 (67 and 68) not (72 or 74) 69 85 76 (67 and 69) not (73 or 74 or 75) 470 542 77 (67 and 70) not (73 or 74 or 75 or 76) 990 1222 78 limit 75 to yr="2018 -Current" NA 27	71	"case report"/	2319441	2658800
73 71 or 72 3/162/2 4329515 74 (animals/ or nonhuman/) not humans/ 6126128 6443842 75 (67 and 68) not (72 or 74) 69 85 76 (67 and 69) not (73 or 74 or 75) 470 542 77 (67 and 70) not (73 or 74 or 75 or 76) 990 1222 78 limit 75 to yr="2018 -Current" NA 27	72	(editorial or letter or comment or historical article).pt.	1600355	1895537
74 (animais/ or nonhuman/) not humans/ 6126128 6443842 75 (67 and 68) not (72 or 74) 69 85 76 (67 and 69) not (73 or 74 or 75) 470 542 77 (67 and 70) not (73 or 74 or 75 or 76) 990 1222 78 limit 75 to yr="2018 -Current" NA 27	73		3/162/2	4329515
75 (o7 and o8) not (72 or 74) 69 85 76 (67 and 69) not (73 or 74 or 75) 470 542 77 (67 and 70) not (73 or 74 or 75 or 76) 990 1222 78 limit 75 to yr="2018 -Current" NA 27	74	(animais/ or nonhuman/) not humans/	6126128	6443842
70 (or and os) not (73 or 74 or 75) 470 542 77 (67 and 70) not (73 or 74 or 75 or 76) 990 1222 78 limit 75 to yr="2018 -Current" NA 27	70	(0/ and 08) not (/2 or /4) (67 and 60) not (72 or 74 or 75)	09	50 540
78 limit 75 to yr="2018 -Current" 990 1222	70	(07 and 03) not (73 or 74 or 75) (67 and 70) not (72 or 74 or 75 or 76)	470	04Z
	78	limit 75 to vr="2018 -Current"	NA	27

79	limit 76 to yr="2018 -Current"	NA	106
80	limit 77 to yr="2018 -Current"	NA	302

a. Embase <1974 to 18 July 2018>

b. Embase <1974 to 24 September 2021 >

MEDLINE

Table A.2 Search results Questions 1 to 4: Medline (via Ovid) for Level I, Level II and Level III studies

#	Searches	Results ^a 19 July 2018	Results ^b 27 Sept 2021
1	exp "obstetrics"/ or exp "obstetric care"/ or exp "pregnancy"/ or exp "pregnancy disorder"/ or exp "prenatal disorder"/	846826	945998
2	(obstetric or obstetrics or pregnancy or maternal).ti,ab,kw.	542671	643633
3	(prepartum or pre partum or pre-partum or intrapartum or intra partum or intra-partum or perinatal or perin	72542	87299
4	(antenatal or ante natal or ante-natal or prenatal or pre natal or pre-natal).ti,ab,kw.	113427	137293
5	(postnatal or post natal or post-natal or postpartum or post partum or post-partum).ti,ab,kw.	152703	181566
6	1 or 2 or 3 or 4 or 5	1127944	1280398
7	exp "fetus"/	151495	161305
8	(fetu* or fetal* or f?etal*).ti,ab,kw.	296734	333936
9	7 or 8	368759	409865
10	exp alloimmunization/	0	0
11	exp Rh Isoimmunization/	1672	1753
12	(Rh* Isoimmuni?ation or Rh* D Isoimmuni?ation).ti,ab.	602	614
13	(Rh* alloimmuni?ation or Rh* D alloimmuni?ation).ti,ab.	215	256
14	(Rh* incompatibility or Rh* D incompatibility or blood group incompatibility).ti,ab.	910	973
15	(((Rh* adj3 incompatib*) or Rh* D) adj3 incompatibl*).ti,ab.	154	166
16	((Rh or RhD or rhesus) adj5 sensiti*).ti,ab.	1197	1225
17	((fetomaternal or feto-maternal or foetomaternal or foeto-maternal) adj2 immuni?ation).ti,ab.	78	78
18	((rh or RhD or rhesus) adj2 (immuni?ation or autoimmuni?ation)).ti,ab.	755	780
19	10 or 11 or 12 or 13 or 14 or 15 or 16 or 17 or 18	4742	5020
20	exp rhesus D antigen/	0	0
21	rhesus D antigen.ti,ab.	37	38
22	rh* D antigen.ti,ab.	183	189
23	(RhD or rhesus D or Rh?D or Rh-?D or Rh D).ti,ab.	4497	5412
24	(Rh-negative or Rh-positive).ti,ab.	951	1026
25	(Rhesus negative or Rhesus positive).ti,ab.	241	258
26	((rh or rhesus) adj2 (factor or factors or antigen\$ or system or group)).ti,ab.	3883	4358
27	20 or 21 or 22 or 23 or 24 or 25 or 26	8684	10069
28	(Macaca mulatta or Simian Immunodeficiency Virus or zika or macaque?).ti,ab.	32787	41471
29	27 not 28	8527	9889
30	(isoimmuni?ation or alloimmuni?ation).ti,ab,kw.	3791	4313
31	(isoimmuni* or iso-immuni* or isoimmune or iso-immune).ti,ab,kw.	2001	2064
32	(alloimmuni* or allo-immuni* or alloimmune or allo-immune).ti,ab,kw.	6618	7647
33	(unsensiti?ed or un-sensiti?ed or non-sensiti?ed).ti,ab,kw.	1631	1750
34	(sensiti?ation* or sensiti?ed).ti,ab,kw.	90235	103333
35	30 or 31 or 32 or 33 or 34	98707	112853
36	exp Erythroblastosis, Fetal/	11582	12015
37	((erythroblastoses or erythroblastosis) adj2 (fetal* or f?etal*)) ti ab kw.	858	908
38	(h?emolytic disease* or h?emolytic disorder*).ti,ab,kw.	4553	4970
39	(HDFN or HDN).ti,ab,kw.	552	722
40	36 or 37 or 38 or 39	13563	14346
41	6 or 9 or 19 or 29 or 35 or 40	1361580	1539131

#	Searches	Results ^a 19 July 2018	Results ^b 27 Sept 2021
42	exp Rh D immunoglobulin/	0	0
43	exp Rho D Immune Globulin/	1271	1388
44	exp "Rho(D) Immune Globulin"/	1271	1388
45	exp anti-D immunoglobulin/	1271	1388
46	Rh* D Immune Globulin.ti,ab.	68	73
47	(rh* immunoglobulin or rh* d immunoglobulin).ti,ab.	215	236
48	(rh* immuni?ation or rh* d immuni?ation).ti,ab.	486	504
49	42 or 43 or 44 or 45 or 46 or 47 or 48	1885	2028
50	exp rhesus D antibody/	0	0
51	rhesus D antibody.ti,ab.	10	10
52	(rh* D antibody or rh*D antibody).ti,ab.	86	89
53	(anti-D or anti D or anti?D).ti,ab.	2820	3050
54	50 or 51 or 52 or 53	2890	3123
55	exp rhogam/	1271	1408
56	rhogam.ti,ab.	32	32
57	exp winrho/	0	0
58	winrho.ti,ab.	41	42
59	exp rhophylac/	1271	1390
60	rhophylac.ti,ab.	8	8
61	exp MICRhoGam/	1271	1388
62	exp BayRHo-D/	0	0
63	exp rhesonativ/	0	0
64	'RhD immunoglobulin vf.ti,ab.	0	0
65	55 or 56 or 57 or 58 or 59 or 60 or 61 or 62 or 63 or 64	1302	1421
66	49 or 54 or 65	3847	4144
67	41 and 66	2741	2972
68	exp meta analysis/ or meta analysis.mp. or exp systematic review/ or systematic review.mp. or pooled analysis.mp. or ((exp review/ or review.mp.) and (systemat* or pool*).mp.)	278527	432159
69	exp comparative study/ or comparative study.mp. or exp clinical trial/ or clinical trial.mp. or randomized controlled trial.mp. or randomi?ed controlled trial.mp. or exp randomized controlled trial/ or exp randomization/ or randomization.mp. or randomi?ation.mp. or exp single blind procedure/ or single blind procedure.mp. or exp double blind procedure/ or double blind procedure.mp. or exp triple blind procedure/ or triple blind procedure.mp. or exp consover procedure/ or crossover procedure.mp. or exp placebo/ or placebo*.mp. or random*.mp. or triple blind.mp. or triple blind.mp. or single blind.mp. or exp prospective study/ or prospective study.mp.	3476781	3994060
70	exp clinical study/ or exp case control study/ or exp family study/ or exp longitudinal study/ or exp retrospective study/ or exp cohort analysis/ or (cohort adj1 stud*).mp. or (case control adj1 stud*).mp. or (exp prospective study/ not randomi?ed controlled trials.mp.) or (follow up adj1 stud*).mp. or (observational adj1 stud*).mp. or (epidemiologic* adj1 stud*).mp. or (cross sectional adj1 stud*).mp.	2982544	3749193
71	"case report"/	1887103	2212725
12	(editorial or letter or comment or historical article).pt.	1969551	2342764
73	/1 or /2	3656632	4331387
74	(animals/ or nonhuman/) not humans/	4443785	4856723
75	(67 and 68) not (72 or 74)	29	40
76	(67 and 69) not (73 or 74 or 75)	316	345
77	(67 and 70) not (73 or 74 or 75 or 76)	190	239
69	limit 66 to yr="2018 -Current"	NA	12
70	limit 67 to yr="2018 -Current"	NA	37
71	limit 68 to yr="2018 -Current"	NA	53

a. Ovid MEDLINE(R) Epub Ahead of Print, In-Process & Other Non-Indexed Citations, Ovid MEDLINE(R) Daily, Ovid MEDLINE and Versions(R) 1946 to July 18, 2018

b. Ovid MEDLINE(R) ALL <1946 to September 24, 2021>

Evidence-Based Medicine Reviews

#	Searches	Results 19 July 2018	Results 27 Sept 2021
1	exp "obstetrics"/ or exp "obstetric care"/ or exp "pregnancy"/ or exp "pregnancy disorder"/ or exp "prenatal disorder"/	19993	23474
2	(obstetric or obstetrics or pregnancy or maternal).ti,ab,kw.	37479	71083
3	(prepartum or pre partum or pre-partum or intrapartum or intra partum or intra-partum or perinatal or peri natal or peri-natal).ti,ab,kw.	4 694	7790
4	(antenatal or ante natal or ante-natal or prenatal or pre natal or pre-natal).ti,ab,kw.	5834	10176
5	(postnatal or post natal or post-natal or postpartum or post partum or post-partum).ti,ab,kw.	8587	15414
6	1 or 2 or 3 or 4 or 5	50733	88340
7	exp "fetus"/	1614	1812
8	(fetu* or fetal* or f?etu* or f?etal*).ti,ab,kw.	8812	15176
9	7 or 8	9664	16152
10	exp alloimmunization/	0	0
11	exp Rh Isoimmunization/	30	32
12	(Rh* Isoimmuni?ation or Rh* D Isoimmuni?ation).ti,ab.	13	18
13	(Rh* alloimmuni?ation or Rh* D alloimmuni?ation).ti,ab.	6	14
14	(Rh* incompatibility or Rh* D incompatibility or blood group incompatibility).ti,ab.	25	52
15	(((Rh* adj3 incompatib*) or Rh* D) adj3 incompatibl*).ti,ab.	2	4
16	((Rh or RhD or rhesus) adj5 sensiti*).ti,ab.	23	28
17	((fetomaternal or feto-maternal or foeto-maternal) adj2 immuni?ation).ti.ab.	0	0
18	((rh or RhD or rhesus) adj2 (immuni?ation or autoimmuni?ation)) ti ab.	30	33
19	10 or 11 or 12 or 13 or 14 or 15 or 16 or 17 or 18	116	164
20	exp rhesus D antigen/	0	0
21	rhesus D antigen ti ab.	0	0
22	rh* D antioen ti ab.	0	0
23	(RhD or rhesus D or Rh?D or Rh-?D or Rh D).ti.ab.	139	217
24	(Rh-negative or Rh-positive).ti.ab.	23	46
25	(Rhesus negative or Rhesus positive).ti.ab.	17	22
26	((rh or rhesus) adj2 (factor or factors or antigen\$ or system or group)).ti.ab.	117	162
27	20 or 21 or 22 or 23 or 24 or 25 or 26	283	418
28	(Macaca mulatta or Simian Immunodeficiency Virus or zika or macague?).ti.ab.	151	281
29	27 not 28	283	418
30	(isoimmuni?ation or alloimmuni?ation) ti ab kw.	175	262
31	(isoimmuni* or iso-immuni* or isoimmune or iso-immune).ti.ab.kw.	46	65
32	(alloimmuni* or allo-immuni* or alloimmune or allo-immune).ti.ab.kw.	266	406
33	(unsensiti?ed or un-sensiti?ed or non-sensiti?ed).ti,ab,kw.	49	71
34	(sensiti?ation* or sensiti?ed).ti.ab.kw.	2899	4409
35	30 or 31 or 32 or 33 or 34	3200	4861
36	exp Ervthroblastosis Fetal/	70	76
37	((ervthroblastoses or ervthroblastosis) adi2 (fetal* or f?etal*)) ti ab kw.	14	10
38	(h?emolytic disease* or h?emolytic disorder*) ti ab kw	106	157
39	(HDEN or HDN) ti ab kw	21	32
40	36 or 37 or 38 or 39	150	210
41	6 or 9 or 19 or 29 or 35 or 40	55540	95795
42	exp Rh D immunoalobulin/	0	0
43	exp Rho D Immune Globulin/	169	240
44	exp "Rho(D) Immune Globulin"/	169	240
45	exp anti-D immunoalobulin/	169	240
46	Rh* D Immune Globulin fi ab	9	11
47	(rh* immunoalobulin or rh* d immunoalobulin) ti ab.	13	17

#	Searches	Results 19 July 2018	Results 27 Sept 2021
48	(rh* immuni?ation or rh* d immuni?ation).ti,ab.	28	32
49	42 or 43 or 44 or 45 or 46 or 47 or 48	208	288
50	exp rhesus D antibody/	0	0
51	rhesus D antibody.ti,ab.	0	0
52	(rh* D antibody or rh*D antibody).ti,ab.	5	9
53	(anti-D or anti D or anti?D).ti,ab.	145	216
54	50 or 51 or 52 or 53	150	223
55	exp rhogam/	169	240
56	rhogam.ti,ab.	2	4
57	exp winrho/	0	0
58	winrho.ti,ab.	5	7
59	exp rhophylac/	169	240
60	rhophylac.ti,ab.	5	7
61	exp MICRhoGam/	169	240
62	exp BayRHo-D/	0	0
63	exp rhesonativ/	0	0
64	'RhD immunoglobulin vf'.ti,ab.	0	0
65	55 or 56 or 57 or 58 or 59 or 60 or 61 or 62 or 63 or 64	176	252
66	49 or 54 or 65	310	457
67	41 and 66	102	149
68	limit 67 to yr="2018 -Current"	NA	19

a. EBM Reviews combines several resources into a single database and includes the following: ACP Journal Club <1991 to June 2018>; Cochrane Database of Systematic Reviews <2005 to July 18, 2018>; Database of Abstracts of Reviews of Effects <1st Quarter 2016>; Cochrane Clinical Answers <June 2018>; Cochrane Central Register of Controlled Trials <June 2018>; Cochrane Methodology Register <3rd Quarter 2012>; Health Technology Assessment <4th Quarter 2016>; NHS Economic Evaluation Database <1st Quarter 2016>.

b. EBM Reviews - Cochrane Database of Systematic Reviews <2005 to September 23, 2021>; EBM Reviews - ACP Journal Club <1991 to August 2021>; EBM Reviews - Database of Abstracts of Reviews of Effects <1st Quarter 2016>; EBM Reviews - Cochrane Clinical Answers <September 2021>; EBM Reviews - Cochrane Central Register of Controlled Trials <August 2021>; EBM Reviews - Cochrane Methodology Register <3rd Quarter 2012>; EBM Reviews - Health Technology Assessment <4th Quarter 2016>; EBM Reviews - NHS Economic Evaluation Database <1st Quarter 2016>

PubMed

The PubMed search is restricted to records that are not indexed for MEDLINE (i.e. in-process citations and citations from journals (or parts of journals) that are not currently MEDLINE-indexed) and to records added to PubMed since January 2006.

The search comprises free-text terms only and replicates the free-text sets in the Embase search (converted from the Ovid syntax).

#	Searches	Results 20 July 2018	Results 27 Sept 2021
#49	(#47 AND pubmednotmedline[sb]) from 2018 to 2021	NA	108
#48	(#47 AND pubmednotmedline[sb])	200	310
#47	(#32 AND #46)	4737	5,281
#46	(#36 OR #45)	8156	9,191
#45	(#38 OR #39 OR #40 OR #41 OR #42 OR #43 OR #44)	53	87
#44	RhD immunoglobulin vf[tiab]	0	0
#43	rhesonativ[tiab]	2	2
#42	BayRHo-D[tiab]	0	0
#41	MICRhoGam[tiab]	2	1
#40	RhD immunoglobulin-vf[tiab]	0	0

Table A.4 Search results Questions 1 to 4: Pubmed (not MEDLINE)

#39 rhophylac[lab] B	#	Searches	Results	Results 27 Sept 2021
Top-projection Participation Parino Participation Participation<	#39	rbophylac[tiab]	8	8
International Internat	#38	wintho[tiab]	42	44
Inspanyon Operating Operating <t< td=""><td>#37</td><td>rhogam[fiab]</td><td>33</td><td>36</td></t<>	#37	rhogam[fiab]	33	36
Description Description Description Description 856 (and [dab] OR ani (diab]) ADD antibody(iab) ADD antibody ADD antibody(iab) ADD antibody(iab)	#36	(#33 OR #34 OR #35)	8146	9 164
Interspace Data	#35	(anti-d[tiah] OR anti d[tiah])	2811	3,104
Intersection of minute (Minute) (Minute) (Minute) (Minute) (Minute) (Minute) 50.02 #33 ((fresential) OR minute) (OR RhD[lab]) (OR RhD[lab)) (AND (minute) (Minute) 1576 1.781 #34 (#8 OR #10 OR #20 OR #27 OR #31) 1018102 1.179.283 #34 (#28 OR #20 OR #20 OR #27 OR #31) 1018102 1.179.283 #34 (#28 OR #20 OR #20 OR #27 OR #31) 1018102 1.179.283 #35 (Indiniab) OR Indiuiab) 563 760 (Minute) Cor Rhaemolytic AND (disorder[liab] OR disorders[liab] OR disease[liab] OR 18492 21.044 #28 (individual Cor Rh2 OR #26 OR #26) 9817 113.839 #27 (#23 OR #24 OR #26 OR #26) 9817 113.839 #28 (individual OR rhotab) OR ensuitation[liab] OR sensitized[liab] OR sensitized[liab]) 99526 104.466 #26 (#50 Nift) OR Rhotab) OR Resultabiliab (OR assensitized[liab] OR sensitized[liab]) 90526 104.466 #27 (#20 OR #20 OR #25 OR #26) 9817 113.839 #27 (#20 OK #25 OR #26) 98871 113.839 #28 (monturn) Tilab) OR sensitization[liab] OR sensitization[liab] OR	#30	(anu-duab) ON anu duab) ((rhogueftiah) OR rh(fiah) AND antibodu(fiah))	1090	5,000
#33 (Interactional of Research and the second and the se	#04	((hesus[iiab] OR th[iiab] OR tho[iiab] OR ph[iiab] OR PhD[iiab]) AND (immunod[ohulin[iiab] OR immuno	4303	1 791
#22 (#50 R# 6) CR #14 OR #22 OR #27 OR #31) 1019102 1,179,283 #31 (#26 RE #20 R #30) 20920 23,548 #30 (hdm[itab] OR hapmolytic) AND (disorder[itab] OR disorders[itab] OR disoaso[itab] OR 14492 21,044 #29 (disorders[itab]) R4492 21,044 3,172 #28 (erythroblastossel[itab]) OR entrisized[itab] OR fetals[itab] OR 3161 3,172 #27 (#23 OR #24 OR #25 OR #26) 98817 113,839 #26 (sensitisation[itab] OR sensitized[itab] OR sensitized[itab] OR sensitized[itab]) 99526 104,466 #275 (#field DR freisultab] OR locanimum("lab) OR sensitized[itab] OR sensitized[itab]) 99527 17,435 #284 (aloimmum"lab) OR sinoimmum[lab) CR aloimmum-litab) CR aloimmum-litab) 6637 7,693 #294 (#20 NOT #21) 23409 27,073 33079 42,076 #294 (#10 R #10 R #19 OR #19) 2449 27,870 3449 30079 42,076 #204 (#15 OR #10 CR #10 CR #19 OR #19) 244 27,870 3887 4,762 31,897 4,762	#33		1576	1,701
#31 (#26 OR #20 OR #30) 20920 23,548 #30 (htdn[tab] OR hdn[tab]) 553 750 #29 (htemolytic) OR haemolytic) AND (disorder[tiab] OR disorders[tiab] OR disease[tiab] OR 18492 21,044 #28 (intervibrolastoses[tab] OR enythroblastosis[tiab] AND (fetal[tiab] OR fotal[tiab] OR 3161 3,172 #28 (intervibrolastose]tab] OR enythroblastosis[tiab] AND (fetal[tiab] OR fotal[tiab] OR 98617 113,839 #27 (#23 OR #24 OR #25 OR #26) #286 (ensistation" titab) OR sensitized[tiab] OR sensitized[tiab]) 90526 104,466 #28 (ensistation" titab) OR sensitized[tiab] OR sensitized[tiab]) 90526 114,466 #27 (#26 IOR #100 RH no[tab] OR neumann"[tab] OR sensitized[tiab] OR sensitized[tiab]) 1599 1,735 #28 (scimmum"[tab] OR sensitized[tiab] OR sensitized[tiab] OR macaque[tiab] 23409 27,073 #27 (#20 NOT #21) Caster mutal tibb) OR factors[tiab] OR antigens[tiab] OR thesus gealue[tiab] OR thesus gealue[tiab] OR thesus gealue[tiab] OR thesus[tiab] AND	#32	(#5 OR #6 OR #14 OR #22 OR #27 OR #31)	1018102	1,179,283
#30 (hdm[lab] OR hadm[lab]) 553 750 #29 (flemolytic: OR naemolytic) AND (disorder[liab] OR disorders[liab] OR disaese[liab] OR diseases[liab] OR 18492 21,044 #28 (feeldisplab)) 3161 3,172 #27 (#23 OR #24 OR #25 OR #26) 98817 113,839 #26 (senstisation"[liab] OR hemsul[tab] AND (senstisargl[tab] OR senstized[liab]) 90526 104,466 #27 (#23 OR #24 OR #25 OR #26) 98817 113,839 #26 (senstisation"[liab] OR hemsul[tab] AND (senstissed[liab] OR sensitized[liab]) 90526 104,466 #27 (#104 D) CR hothesul[tab] OR Not (senstissed[liab] OR sensitized[liab]) 2006 2.084 #28 (senstissation"[liab] OR senstissed[liab] OR liso-immune[liab] 2006 2.084 #23 (senstimmune[liab] OR missing Immunodeficiency Virus[liab] OR macaque[liab] 30079 42.075 #24 (#00 NOT #21) (#16 OR #17 OR #18 OR #19) 2449 27.870 #24 (#16 OR #17 OR #18 OR #19) 240 261 #17 (thesus negative[liab] OR hossus[liab] AND (actors[liab] OR missus[liab] OR missus[liab] OR hossus[liab] OR missus[l	#31	(#28 OR #29 OR #30)	20920	23,548
#29 (fhemolytic) CR haemolytic) AND (disorder[tiab] OR disorders[tiab] OR disease[tiab] OR 18492 21,044 #28 ((erythroblastoses[tiab]) CR erythroblastosis[tiab] AND (fetal[tiab] OR fotals[tiab] OR 3161 3,172 #27 (#23 OR #24 OR #25 OR #25) 98817 113,839 #265 (sensitisation*[tiab] OR sensitization*[tiab] OR sensitisated[tiab]) 90526 104,466 #254 (sensitisation*[tiab] OR sensitisated[tiab] OR sensitisated[tiab]) 99817 113,839 #244 (aliommun*[tiab] OR allo-immun*[tiab] OR sensitisated[tiab] OR sensitisated[tiab]) 6657 7,693 #224 (#20 NOT #21) 2409 27,073 #224 (#20 NOT #21) 2409 27,073 #221 (Macaca mulatal[tiab] OR factor[tiab] OR factors[tiab] OR antigens[tiab] OR 33079 42,075 #221 (#160 R #10 R #13 OR #19) 27458 31,807 #232 (#20 NOT #21) C#16 OR #10 R #13 24249 27,870 #241 (Macaca mulatal[tiab] OR factor[tiab] OR factors[tiab] OR antigens[tiab] OR 24249 27,870 #141 (thitab) OR #10 R #11 OR #12 OR #13) 1105	#30	(hdfn[tiab] OR hdn[tiab])	553	750
#28 ((erythroblastosselitab) CR erythroblastossiliab) AND (fetal[tiab] OR fetalis[tiab] OR 3161 3.172 #27 (#32 OR #/24 OR #/25 OR #/26) 98817 113,839 #26 (seratisation"tiab) OR sensitization"[tiab] OR sensitized[tiab] OR sensitized[tiab] OR 99526 104,466 #275 sensitization(tiab) OR sensitization[tiab] OR sensitized[tiab] OR sensitized[tiab] OR 1599 1.735 #24 (aloimmun"[tiab] OR aloi-immun"[tiab] OR aloimmune[tiab] OR aloi-immune[tiab]) 6637 7,693 #22 (#20 NR #40 OR #10 R #10 R #10) R isoimmune[tiab] OR isoi-immune[tiab]) 2006 2,084 #24 (aloimmun"[tiab] OR simia Immunodeficiency Virus[tiab] OR zita[tiab] OR macaque[tiab] OR 33079 27,073 #24 (#20 OF #10 C #14 OR #14 OR #19) 24249 27,870 #15 (rheque negative[tiab] OR couplicab] 4249 27,870 #16 (rhegue negative[tiab] OR rhouspositive[tiab] OR rhousposi	#29	((hemolytic OR haemolytic) AND (disorder[tiab] OR disorders[tiab] OR disease[tiab] OR diseases[tiab]))	18492	21,044
#27 (#23 OR #24 OR #26 OR #26) 98817 113,839 #26 (sensitisaton"[tab] OR sensitizaton"[tab] OR sensitizad[tab] OR antigen?[tab] OR provide tab] OR the sensitivad] OR the negative[tab] OR the sensitivad] OR the negative[tab] OR (neter-maternal[tab]) AND (incomp	#28	((erythroblastoses[tiab] OR erythroblastosis[tiab]) AND (fetal[tiab] OR foetal[tiab] OR fetalis[tiab] OR foetalis[tiab]))	3161	3,172
#26(sensitisation*[tab] OR sensitization*[tab] OR sensitized[tab])90526104,466#275sensitisation[tab] OR hot[tab] OR hot[tab] OR sensitized[tab])15991,735#244(alloimmun*[tab] OR alloimmune[tab] OR alloimmune[tab] OR alloimmune[tab])66377,693#232(isoimmun*[tibb] OR liso-immun*[tibb] OR liso-immune[tab])20062,084#243(idoimmun*[tab] OR liso-immun*[tibb] OR liso-immune[tab])20062,084#244(idoimmun*[tab] OR Simian Immunodeficiency Virus[tab] OR zika[tab] OR macaque[tab] OR macaque[tab])3307942,075#210(Macaca mulata[tab] OR fiso-immun/[tiba] OR factors[tab] OR antigens[tab] OR macaque[tab])244927,870#211(Macaca mulata[tab] OR fiso-grinu) (fator[tab] OR factors[tab] OR antigens[tab] OR system[tab] OR group[tab])240261#113(Infibal OR rhesus[tab]) AND (fator[tab] OR factors[tab] OR hostive[tab])240261#114(from use negative[tab] OR rhesus positive[tab] OR rhogative[tab] OR hostive[tab]3493,815#115(infibia) OR rhesus[tab] OR rhogative[tab] OR rhogative[tab] OR adus(tab] OR Rho [tab] OR rhogative[tab] OR rhogative[tab] OR rhogative[tab] OR adus(tab] OR Rho [tab] OR rhogative[tab] OR rhogative[tab] OR adus(tab] OR Rho [tab] OR rhogative[tab] OR rhogative[tab] OR 	#27	(#23 OR #24 OR #25 OR #26)	98817	113,839
#25((fn[tab] OR mol(tab] OR sensitis/and[tab] OR sensitiz/ad[tab])15991,735#24(alionmun"[tab] OR alio-immun"[tab] OR alio-immun"[tab]66377,693#20(#20 NOT #21)20402,0842,073#21(macaques[tab])(Mod Car a mulata[tab] OR finitian Immunodeficiency Virus[tab] OR antigens[tab] OR3,07942,075#20(#15 OR #16 OR #17 OR #16 OR #19)27,6703,8072,780#19(system)[tab] OR hesus[tab] OR factors[tab] OR antigens[tab] OR2,424927,870#18(thesus negative[tab] OR thesus positive[tab] OR the negative[tab] OR the positive[tab]9491,033#17(th-negative[tab] OR thesus positive[tab] OR the negative[tab] OR the positive[tab]9491,033#16(RhD[tab] OR thesus [tab] OR Integative[tab] OR the positive[tab] OR thesus [tab] OR thesus [tab] OR for the positive[tab] OR thesus [tab] OR for the positive[tab] OR thesus [tab] OR thesus [tab] OR for the positive[tab] OR	#26	(sensitisation*[tiab] OR sensitization*[tiab] OR sensitised[tiab] OR sensitized[tiab])	90526	104,466
#24 (alloimmuni*[tiab] OR allo-immune[tiab] OR allo-immune[tiab]) 6637 7,633 #23 (soimmuni*[tiab] OR iso-immuni*[tiab] OR isoimmune[tiab] OR iso-immune[tiab]) 2006 2,084 #22 (#20 NOT #21) 23409 27,073 #21 (Maccaa mulatifiab) OR simian Immunodeficiency Virus[tiab] OR zika[tiab] OR macaque[tiab] OR 33079 42,075 #20 (#15 OR #16 OR #17 OR #18 OR #19) 27458 31,807 #19 system[tiab] OR rough[tab]) AND (factor[tiab] OR factors[tiab] OR antigen*[tiab] OR zough[tab] 24249 27,870 #18 (rhesus negative[tiab] OR rhesus positive[tiab] 240 261 261 #17 (rh-negative[tiab] OR rhesus positive[tiab] OR rh negative[tiab] OR rhositive[tiab] OR rhositive[#25	((rh[tiab] OR rho[tiab] OR rhesus[tiab]) AND (sensitising[tiab] OR sensitizing[tiab] OR sensitization[tiab] OR sensitization[tiab] OR sensitized[tiab] OR sensitized[tiab]))	1599	1,735
#23 (isoimmuni*[tiab] OR iso-immune[tiab] OR iso-immune[tiab] OR iso-immune[tiab] 2006 2,084 #22 (#20 NOT #21) 23409 27,073 #21 (Macaca mulatla[tiab] OR Simian Immunodeficiency Virus[tiab] OR zika[tiab] OR macaque[tiab] OR macaques[tiab]) 30079 42,075 #20 (#15 OR #16 OR #17 OR #18 OR #19) 27458 31,807 #19 (/h[tiab] OR rhesus[tiab]) AND (factor[tiab] OR factors[tiab] OR antigen*[tiab] OR antigens[tiab] OR system[tiab] OR rhesus positive[tiab] 240 261 #17 (rhesus negative[tiab] OR rhesus positive[tiab]) 240 261 #18 (rhesus negative[tiab] OR rhesus[tiab]) AND (natigen[tiab]) 349 1,038 #16 (rhesus negative[tiab] OR rhesus[tiab]) AND antigen[tiab]) 3429 3,815 #15 ((/h[tiab] OR rhesus[tiab]) AND (minumisation[tiab] OR immunization[tiab] OR autoimmunisation[tiab] OR rhesus[tiab]) AND (minumisation[tiab] OR immunization[tiab] OR autoimmunisation[tiab] OR rhesus[tiab]) AND (encompatibility[tiab]) 1066 174 #11 ((Rh[tiab] OR rhesus[tiab]) AND (incompatibility[tiab]) 1307 1,428 #14 (#10 RR #10 OR #10	#24	(alloimmuni*[tiab] OR allo-immuni*[tiab] OR alloimmune[tiab] OR allo-immune[tiab])	6637	7,693
#22 (#20 NOT #21) 23409 27,073 #21 (Macaca mulatla[tiab] OR Simian Immunodeficiency Virus[tiab] OR zika[tiab] OR macaque[tiab] OR macaques[tiab] 33079 42,075 #20 (#15 OR #16 OR #17 OR #18 OR #19) 27458 31,807 #19 (h[tiab] OR rhesus[tiab]) AND (factor[tiab] OR factors[tiab] OR antigen*[tiab] OR antigens[tiab] OR system[tiab] OR proup[tiab] 240 261 #11 (rhesus negative[tiab] OR rh-positive[tiab] 240 261 #17 (rh-negative[tiab] OR rhesus positive[tiab] OR rh positive[tiab] 3487 4,752 #15 ((rh[tiab] OR rhesus d[tiab] OR rhesus[tiab]) AND antigen[tiab]) 3429 3,815 #14 (#7 OR #6 OR #0 OR #10 OR #11 OR #12 OR #13) 11065 12,721 #13 attribution munization[tiab] OR intesus[tiab] NAD (munisation[tiab] OR international (tiab] OR 2055 2,251 #14 (#7 OR #6 OR #0 CR #10 OR #11 OR #12 OR #13) 11065 12,721 #13 attribution Minimumization[tiab] OR internation[tiab] OR 2055 2,251 #11 ((Refurator anternational) AND (incompatib*[tiab])) 1307 1,428 #11 ((Refutab] OR rholes	#23	(isoimmuni*[tiab] OR iso-immuni*[tiab] OR isoimmune[tiab] OR iso-immune[tiab])	2006	2,084
#21(Macaca mulatla[tiab] OR Simian Immunodeficiency Virus[tiab] OR zika[tiab] OR macaque[tiab] OR macaques[tiab])3307942,075#20(#15 OR #16 OR #17 OR #18 OR #19)2745831,807#19(rht[tiab] OR nesus[tiab] ND (factor[tiab] OR factors[tiab] OR antigen*[tiab] OR antigens[tiab] OR system[tiab] OR group[tiab])240261#18(rhesus negative[tiab] OR rhesus positive[tiab])240261#17(rh-negative[tiab] OR rhesus positive[tiab] OR n positive[tiab])9491,038#16(RhD[tiab] OR rhesus d[tiab] OR n negative[tiab] OR rh positive[tiab])34293,815#116(RhD[tiab] OR rhesus d[tiab] OR Rh D[tiab]34293,815#116(rht[tiab] OR rhesus[tiab]) AND antigen[tiab]1106512,721#13(trh[tiab] OR RhD[tiab] OR rhesus[tiab] AND (immunisation[tiab] OR immunization[tiab] OR autoimmunisation[tiab] OR rhesus[tiab] AND (immunisation[tiab] OR factor-maternal[tiab] OR focto-maternal[tiab] OR focto-maternal[tiab] OR autoimmunisation[tiab] OR rhesus[tiab] AND (incompatib*[tiab]))166174#11((Rh[tiab] OR RhD[tiab] OR rhesus[tiab]) AND (incompatib*[tiab]))13071,428#19((rht[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatib*[tiab]))13071,428#19((rht[tiab] OR rhd[tiab] OR nesus[tiab]) AND (incompatib*[tiab]) OR (blood group ncompatbility[tiab]) OR aloimmunisation[tiab])6382#11((rht[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatib*[tiab] OR foetal*[tiab])25903,240#8((rht[tiab] OR rhd[tiab] OR neture]tiab] OR pretore]tiab] OR foetal*[tiab] OR pretore]tiab] <td>#22</td> <td>(#20 NOT #21)</td> <td>23409</td> <td>27,073</td>	#22	(#20 NOT #21)	23409	27,073
#20 (#15 OR #16 OR #17 OR #18 OR #19) 27458 31,807 #19 (rh[tiab] OR rhesus[tiab]) AND (factor[tiab] OR factors[tiab] OR antigen*[tiab] OR antigens[tiab] OR yespen[tiab] OR group[tiab]) 24249 27,870 #18 (rhesus negative[tiab] OR rhesus positive[tiab]) 240 261 #17 (rh-negative[tiab] OR rhesus ditab] OR Rh D[tiab]) 949 1,038 #16 (RhD[tiab] OR rhesus ditab] OR Rh D[tiab]) 3887 4,752 #15 ((rh[tiab] OR rhesus ditab] OR Rh D[tiab]) 3429 3,815 #14 (#7 OR #8 OR #9 OR #10 OR #11 OR #12 OR #13) 11065 12,721 #13 ((rh[tiab] OR RhD[tiab] OR hesus[tiab]) AND (monunisation[tiab] OR intransitation[tiab] OR autoimmunization[tiab]) 2055 2,251 #11 ((Rh[tiab] OR RhD[tiab] OR hesus[tiab]) AND (monpatb "[tiab])) 166 174 #11 ((Rh[tiab] OR RhD[tiab] OR rhesus[tiab]) AND (incompatb "[tiab])) 1307 1,428 #11 ((Rh[tiab] OR RhD[tiab] OR rhesus[tiab)) AND (incompatbility[tiab]) OR (blood group informatibility[tiab]) 1636 1,747 #11 ((Rh[tiab] OR rhesus[tiab]) AND (incompatbility[tiab]) OR (blood group informabitily[tiab]) 1636 1,74	#21	(Macaca mulatta[tiab] OR Simian Immunodeficiency Virus[tiab] OR zika[tiab] OR macaque[tiab] OR macaques[tiab])	33079	42,075
#19(rh[tiab] OR rhesus[tiab]) AND (factor[tiab] OR antigen*[tiab] OR antigens[tiab] OR system[tiab] OR group[tiab])2424927,870#18(rhesus negative[tiab] OR rhesus positive[tiab])240261#17(rh-negative[tiab] OR rhesus d[tiab] OR rhegative[tiab] OR rh positive[tiab])9491,038#17(rh-negative[tiab] OR rhesus d[tiab] OR Rh D[tiab])38874,752#15((rh[tiab] OR rhesus d[tiab] OR Rh D[tiab])34293,815#14(#7 OR #8 OR #9 OR #10 OR #11 OR #12 OR #13)1106512,721#13((rh[tiab] OR rhesus[tiab]) AND (immunisation[tiab] OR immunization[tiab] OR autoimmunisation[tiab] OR autoimmunization[tiab])20552,251#14(#7 OR #8 OR #9 OR #10 OR #11 OR #12 OR #13)1106512,721#13((rh[tiab] OR RhD[tiab] OR hesus[tiab]) AND (immunisation[tiab] OR internativation[tiab] OR autoimmunisation[tiab] OR autoimmunization[tiab])20552,251#11((Rh[tiab] OR RhD[tiab] OR rhesus[tiab]) AND (incompatibil/[tiab])13071,428#14((rh[tiab] OR rhol[tiab] OR rhesus[tiab]) AND (incompatibil/[tiab])13071,428#19((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatibil/[tiab]) OR (blood group 	#20	(#15 OR #16 OR #17 OR #18 OR #19)	27458	31,807
#18 (rhesus negative[tiab] OR rhesus positive[tiab]) 240 261 #17 (rh-negative[tiab] OR rh-positive[tiab] OR rh negative[tiab] OR rh positive[tiab]) 949 1,038 #16 (RhD[tiab] OR rhesus d[tiab] OR rh D[tiab]) 3887 4,752 #15 ((rh[tiab] OR rhesus d[tiab] OR rhesus[tiab]) AND antigen[tiab]) 3429 3,815 #14 (#7 OR #8 OR #9 OR #10 OR #11 OR #12 OR #13) 11065 12,721 #13 ((rh[tiab] OR RhD[tiab] OR rhesus[tiab]) AND (mmunisation[tiab] OR immunization[tiab] OR autoimmunisation[tiab] OR foto-maternal[tiab] OR	#19	(rh[tiab] OR rhesus[tiab]) AND (factor[tiab] OR factors[tiab] OR antigen*[tiab] OR antigens[tiab] OR system[tiab] OR group[tiab])	24249	27,870
#17 (rh-negative[tiab] OR rh-positive[tiab] OR rh negative[tiab]) 949 1,038 #16 (RhD[tiab] OR rhesus d[tiab] OR Rh D[tiab]) 3887 4,752 #15 ((rh[tiab] OR rhesus d[tiab] OR rhesus[tiab]) AND antigen[tiab]) 3429 3,815 #14 (#7 OR #8 OR #9 OR #10 OR #11 OR #12 OR #13) 11065 12,721 #13 ((rh[tiab] OR RhD[tiab] OR rhesus[tiab]) AND (mmunisation[tiab] OR immunization[tiab] OR autoimmunisation[tiab] OR foto-maternal[tiab] OR foto-maternaltiab] OR foto-maternal[tiab] O	#18	(rhesus negative[tiab] OR rhesus positive[tiab])	240	261
#16(RhD[tiab] OR rhesus d[tiab] OR Rh D[tiab])38874,752#15((rh[tiab] OR rhesus d[tiab] OR rhesus[tiab]) AND antigen[tiab])34293,815#14(#7 OR #8 OR #9 OR #10 OR #11 OR #12 OR #13)1106512,721#13((rh[tiab] OR RhD[tiab] OR rhesus[tiab]) AND (immunisation[tiab] OR immunization[tiab] OR autoimmunization[tiab])20552,251#12((rhetiabl OR RhD[tiab] OR rhesus[tiab]) AND (incompation[tiab]))166174#11((Rh[tiab] OR RhD[tiab] OR rhesus[tiab]) AND (incompatib^T[tiab]))49335,734#10((rh[tiab] OR rhesus[tiab]) AND (incompatib^T[tiab]))13071,428#9((rh(tiab) OR rhesus[tiab]) AND (incompatib^T[tiab]))13071,428#9((rh[tiab] OR rhesus[tiab]) AND (incompatib^T[tiab]) OR (blood group incompatibility[tiab])6382#7(aloimmunization[tiab] OR aloimmunisation[tiab])25903,240#6(fetus[tiab] OR fetu*[tiab] OR fetu*[tiab] OR fetal*[tiab] OR foetal*[tiab])29655337,336#5(#1 OR #2 OR #3 OR #4)738834867,966#4(postnatal[tiab] OR post natal[tiab] OR post-natal[tiab] OR port natal[tiab] OR port-natal[tiab] OR port-natal[tiab] OR port-natal[tiab] OR pre-natal[tiab] OR pre-n	#17	(rh-negative[tiab] OR rh-positive[tiab] OR rh negative[tiab] OR rh positive[tiab])	949	1,038
#15((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND antigen[tiab])34293,815#14(#7 OR #8 OR #9 OR #10 OR #11 OR #12 OR #13)1106512,721#13((rh[tiab] OR RhD[tiab] OR rhesus[tiab]) AND (immunisation[tiab] OR inmunisation[tiab] OR autoimmunisation[tiab] OR autoimmunization[tiab])20552,251#12((fetomaternal[tiab] OR feto-maternal[tiab] OR foetomaternal[tiab] OR foeto-maternal[tiab] AND (immunisation[tiab] OR rhesus[tiab]) AND (sensiti*[tiab]))166174#11((Rh[tiab] OR RhD[tiab] OR rhesus[tiab]) AND (sensiti*[tiab]))49335,734#10((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatib*[tiab]))13071,428#9((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatib*[tiab]))16361,747#9((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatib*[tiab]) OR (blood group incompatibility[tiab]))6382#7(aloimmunization[tiab] OR alloimmunisation[tiab])25903,240#6(fetus[tiab] OR fetu*[tiab] OR foetu*[tiab] OR fetal*[tiab] OR foetal*[tiab])29655337,336#5(#1 OR #2 OR #3 OR #4)738834867,966#4(postnatal[tiab] OR ante natal[tiab] OR post-natal[tiab] OR pre-natal[tiab] OR pre-natal[tiab] OR pre-natal[tiab] OR pre-natal[tiab] OR pre-natal[tiab] OR pre-natal[tiab] OR pre- natal[tiab])115113139,910#3(antenatal[tiab] OR pre-partum[tiab] OR pre-natal[tiab] OR pre-natal[tiab] OR pre- natal[tiab] OR pre-natal[tiab] OR pre- natal[tiab] OR pre-natal[tiab] OR pre- natal[tiab] OR pre-natal[tiab] OR pre- natal[tiab] OR pre- natal[tiab] OR pre-natal[tiab] OR pre- natal[t	#16	(RhD[tiab] OR rhesus d[tiab] OR Rh D[tiab])	3887	4,752
#14(#7 OR #8 OR #9 OR #10 OR #11 OR #12 OR #13)1106512,721#13((rh[liab] OR RhD[liab] OR rhesus[liab]) AND (immunisation[liab] OR immunization[liab] OR autoimmunisation[liab] OR autoimmunization[liab])20552,251#12((fetomaternal[liab] OR feto-maternal[liab] OR foeto-maternal[liab] OR foeto-maternal[liab])166174#11((Rh[liab] OR RhD[liab] OR rhesus[liab]) AND (sensit*[liab]))49335,734#10((rh[liab] OR rhd[liab] OR rhesus[liab]) AND (sensit*[liab]))13071,428#9((rh[liab] OR rhd[liab] OR rhesus[liab]) AND (incompatib*[liab]) OR (blood group incompatibility[liab]))16382#9((rh[liab] OR rhd[liab] OR rhesus[liab]) AND (soimmunization[liab] OR isoimmunisation[liab]))6382#7(alloimmunization[liab] OR foetu*[liab] OR foetu*[liab] OR foeta*[liab] OR foeta*[liab])25903,240#6(fetus[liab] OR foetus[liab] OR foetu*[liab] OR foeta*[liab] OR foeta*[liab])2590337,336#5(#1 OR #2 OR #3 OR #4)738834867,966#44(postnatal[liab] OR post natal[liab] OR post-natal[liab] OR prenatal[liab] OR pre natal[liab] OR pre- natal[liab] OR pre partum[liab] OR pre- natal[liab] OR pre- partum[liab] OR pre- partum[liab])115113139,910#2(prepartum[liab] OR pre partum[liab] OR pre- partum[liab] OR peri-natal[liab] OR peri-natal[liab] OR peri-natal[liab]7316988,660#1(Obstetric[liab] OR obstetrics[liab] OR pre- partum[liab] OR peri-natal[liab] OR peri-natal[liab]554296657,008	#15	((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND antigen[tiab])	3429	3,815
#13(Irh[tiab] OR RhD][tiab] OR hesus[tiab]) AND (immunisation[tiab] OR immunization[tiab] OR autoimmunisation[tiab] OR autoimmunization[tiab]))20552,251#12((fetomaternal[tiab] OR feto-maternal[tiab] OR foetomaternal[tiab]) AND (immunisation[tiab] OR rhosus[tiab]) AND (sensiti*[tiab]))166174#11((Rh[tiab] OR RhD[tiab] OR rhesus[tiab]) AND (sensiti*[tiab]))49335,734#10((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatib*[tiab]))13071,428#9((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatib*[tiab]))13071,428#9((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatib*[tiab]))16382#8((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (isoimmunization[tiab] OR isoimmunisation[tiab]))6382#7(alloimmunization[tiab] OR alloimmunisation[tiab] OR fetal*[tiab] OR foetal*[tiab])25903,240#6(fetus[tiab] OR foetus[tiab] OR foetu*[tiab] OR fetal*[tiab] OR foetal*[tiab])299655337,336#5(#1 OR #2 OR #3 OR #4)738834867,966#44(postnatal[tiab] OR post natal[tiab] OR post-natal[tiab] OR pre-natal[tiab]	#14	(#7 OR #8 OR #9 OR #10 OR #11 OR #12 OR #13)	11065	12.721
#13autoimmunisation[tiab] OR autoimmunization[tiab])1120551#12((fetomaternal[tiab] OR feto-maternal[tiab] OR foeto-maternal[tiab]) AND (immunisation[tiab] OR immunization[tiab]))166174#11((Rh[tiab] OR RhD[tiab] OR rhesus[tiab]) AND (sensiti*[tiab]))49335,734#10((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatib*[tiab]))13071,428#9((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatib*[tiab]))13071,428#9((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatibility[tiab]) OR (blood group incompatibility[tiab]))6382#8((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (isoimmunization[tiab] OR isoimmunisation[tiab])6382#7(alloimmunization[tiab] OR alloimmunisation[tiab] OR foeta*[tiab] OR foeta*[tiab])25903,240#6(fetus[tiab] OR foetus[tiab] OR foeta*[tiab] OR foeta*[tiab] OR foeta*[tiab])299655337,336#5(#1 OR #2 OR #3 OR #4)738834867,966#44(postnatal[tiab] OR post natal[tiab] OR post-natal[tiab] OR post partum[tiab] OR pre- natal[tiab])115113139,910#3(antenatal[tiab] OR ante natal[tiab] OR pre-partum[tiab] OR pre- natal[tiab])7316988,660#4(Dostetric[tiab] OR perinatal[tiab] OR pre- natal[tiab] OR perinatal[tiab] OR peri-natal[tiab] OR peri-natal[tiab]7316988,660#1(Obstetric[tiab] OR obstetrics[tiab] OR pregnancy[tiab] OR maternal[tiab])554296657,008		((rh[tiab] OR RhD[tiab] OR rhesus[tiab]) AND (immunisation[tiab] OR immunization[tiab] OR	0055	2.251
#12((fetomaternal[tiab] OR feto-maternal[tiab] OR foetomaternal[tiab] OR foeto-maternal[tiab]) AND (immunisation[tiab] OR inmunization[tiab]))166174#11((Rh[tiab] OR RhD[tiab] OR hnesus[tiab]) AND (sensiti*[tiab]))49335,734#10((rh[tiab] OR RhD[tiab] OR nhesus[tiab]) AND (incompatib*[tiab]))13071,428#9((rh[tiab] OR rhd[tiab] OR nhesus[tiab]) AND (incompatibility[tiab]) OR (blood group incompatibility[tiab]))6382#8((rh[tiab] OR nhd[tiab] nhesus[tiab]) AND (isoimmunization[tiab] OR isoimmunisation[tiab]))6382#8((rh[tiab] OR nhd[tiab] nhesus[tiab]) AND (isoimmunization[tiab] OR isoimmunisation[tiab]))6382#7(aloimmunization[tiab] OR aloimmunisation[tiab])6382#6(fetus[tiab] OR foetus[tiab] OR foetu*[tiab] OR foeta*[tiab] OR foeta*[tiab] OR foeta*[tiab])299655337,336#5(#1 OR #2 OR #3 OR #4)738834867,966#44(postnatal[tiab] OR post natal[tiab] OR post-natal[tiab] OR post partum[tiab] OR pre- natal[tiab])139,910139,910#33(antenatal[tiab] OR ante-natal[tiab] OR pre-partum[tiab] OR pre- natal[tiab])7316988,660#41(Dostetric[tiab] OR obstetrics[tiab] OR pregnancy[tiab] OR maternal[tiab])554296657,008	#13	autoimmunisation[tiab] OR autoimmunization[tiab]))	2055	
#11((Rh[tiab] OR RhD[tiab] OR rhesus[tiab]) AND (sensiti*[tiab]))49335,734#10((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatib*[tiab]))13071,428#9((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatibility[tiab]) OR (blood group incompatibility[tiab]))16361,747#8((rh[tiab] OR rhd[tiab] rhesus[tiab]) AND (isoimmunization[tiab] OR isoimmunisation[tiab]))6382#7(aloimmunization[tiab] OR alloimmunisation[tiab])25903,240#6(fetus[tiab] OR foetus[tiab] OR foetu*[tiab] OR foeta*[tiab] OR foetal*[tiab])299655337,336#5(#1 OR #2 OR #3 OR #4)738834867,966#4(postnatal[tiab] OR post natal[tiab] OR post-natal[tiab] OR post-natal[tiab] OR post partum[tiab] OR pre- post-partum[tiab])115113139,910#3(antenatal[tiab] OR pre partum[tiab] OR pre-partum[tiab] OR pre- natal[tiab])115113139,910#4(postnatal[tiab] OR pre partum[tiab] OR pre-partum[tiab] OR pre- natal[tiab] OR pre- natal[tiab] OR pre- partum[tiab] OR pre-partum[tiab] OR pre- partum[tiab] OR pre-partum[tiab] OR pre- partum[tiab] OR pre-natal[tiab] OR pre- natal[tiab] OR pre-natal[tiab] OR pre- partum[tiab] OR pre-partum[tiab] OR pre- partum[tiab] OR pre- partum[tiab] OR pre-partum[tiab] OR pre- partum[tiab] OR pre- partum[ti	#12	((fetomaternal[tiab] OR feto-maternal[tiab] OR foetomaternal[tiab] OR foeto-maternal[tiab]) AND (immunisation[tiab] OR immunization[tiab]))	166	174
#10((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatib*[tiab]))13071,428#9((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatibility[tiab]) OR (blood group incompatibility[tiab]))16361.747#8((rh[tiab] OR rhd[tiab] rhesus[tiab]) AND (isoimmunization[tiab] OR isoimmunisation[tiab]))6382#7(alloimmunization[tiab] OR alloimmunisation[tiab])25903,240#6(fetus[tiab] OR foetus[tiab] OR foetu*[tiab] OR foetu*[tiab] OR foetal*[tiab] OR foetal*[tiab])299655337,336#5(#1 OR #2 OR #3 OR #4)738834867,966#4(postnatal[tiab] OR post natal[tiab] OR post-natal[tiab] OR postpartum[tiab] OR post partum[tiab] OR post partum[tiab]1307139,910#3(antenatal[tiab] OR ante natal[tiab] OR pre-partum[tiab] OR intrapartum[tiab] OR intra partum[tiab]7316988,660#2(posteric[tiab] OR obstetrics[tiab] OR pregnancy[tiab] OR maternal[tiab] OR maternal[tiab]54296657,008	#11	((Rh[tiab] OR RhD[tiab] OR rhesus[tiab]) AND (sensiti*[tiab]))	4933	5,734
#9((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatibility[tiab]) OR (blood group incompatibility[tiab]))16361,747#8((rh[tiab] OR rhd[tiab] rhesus[tiab]) AND (isoimmunization[tiab] OR isoimmunisation[tiab]))6382#7(alloimmunization[tiab] OR alloimmunisation[tiab])25903,240#6(fetus[tiab] OR foetus[tiab] OR fetu*[tiab] OR fetu*[tiab] OR fetat*[tiab] OR foetal*[tiab])299655337,336#5(#1 OR #2 OR #3 OR #4)738834867,966#4(postnatal[tiab] OR post natal[tiab] OR post-natal[tiab] OR postpartum[tiab] OR post partum[tiab] OR post-partum[tiab])153623183,012#3(antenatal[tiab] OR ante natal[tiab] OR pre-partum[tiab] OR prenatal[tiab] OR pre-natal[tiab] OR pre-natal[tiab] OR pre-natal[tiab] OR pre- natal[tiab])115113139,910#2(prepartum[tiab] OR pre partum[tiab] OR pre- partum[tiab] OR pre- natal[tiab] OR pre- natal[tiab] OR pre- natal[tiab] OR pre- natal[tiab] OR peri natal[tiab] OR pre- natal[tiab] OR peri-natal[tiab]554296657,008	#10	((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatib*[tiab]))	1307	1,428
#8((rh[tiab] OR rhd[tiab] rhesus[tiab]) AND (isoimmunization[tiab] OR isoimmunisation[tiab]))6382#7(alloimmunization[tiab] OR alloimmunisation[tiab])25903,240#6(fetus[tiab] OR foetus[tiab] OR fetu*[tiab] OR foetu*[tiab] OR fetal*[tiab] OR foetal*[tiab])299655337,336#5(#1 OR #2 OR #3 OR #4)738834867,966#4(postnatal[tiab] OR post natal[tiab] OR post-natal[tiab] OR postpartum[tiab] OR post partum[tiab] OR post partum[tiab] OR post-partum[tiab] OR pre-natal[tiab] OR pre-natal[tiab]88,660#2(prepartum[tiab] OR obstetrics[tiab] OR pregnancy[tiab] OR maternal[tiab])554296657,008	#9	((rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatibility[tiab]) OR (blood group incompatibility[tiab]))	1636	1,747
#7(alloimmunization[tiab] OR alloimmunisation[tiab])25903,240#6(fetus[tiab] OR foetus[tiab] OR fetu*[tiab] OR foetu*[tiab] OR fetal*[tiab] OR foetal*[tiab])299655337,336#5(#1 OR #2 OR #3 OR #4)738834867,966#4(postnatal[tiab] OR post natal[tiab] OR post-natal[tiab] OR postpartum[tiab] OR post partum[tiab] OR post-partum[tiab])153623183,012#3(antenatal[tiab] OR ante natal[tiab] OR pre-partum[tiab] OR pre-partum[tiab]7316988,660#1(Obstetric[tiab] OR obstetrics[tiab] OR pregnancy[tiab] OR maternal[tiab])554296657,008	#8	((rh[tiab] OR rhd[tiab] rhesus[tiab]) AND (isoimmunization[tiab] OR isoimmunisation[tiab]))	63	82
#6(fetus[tiab] OR foetus[tiab] OR fetu*[tiab] OR foetu*[tiab] OR fetal*[tiab] OR foetal*[tiab])299655337,336#5(#1 OR #2 OR #3 OR #4)738834867,966#4(postnatal[tiab] OR post natal[tiab] OR post-natal[tiab] OR postpartum[tiab] OR post partum[tiab] OR post-partum[tiab])153623183,012#3(antenatal[tiab] OR ante natal[tiab] OR pre-natal[tiab] OR prenatal[tiab] OR pre natal[tiab] OR pre natal[tiab] OR pre- natal[tiab])115113139,910#2(prepartum[tiab] OR pre partum[tiab] OR pre-partum[tiab] OR pre- OR intra-partum[tiab] OR prenatal[tiab] OR prenatal[tiab] OR peri-natal[tiab])7316988,660#1(Obstetric[tiab] OR obstetrics[tiab] OR pregnancy[tiab] OR maternal[tiab])554296657,008	#7	(alloimmunization[tiab] OR alloimmunisation[tiab])	2590	3,240
#5(#1 OR #2 OR #3 OR #4)738834867,966#4(postnatal[tiab] OR post natal[tiab] OR post-natal[tiab] OR postpartum[tiab] OR post partum[tiab] OR post-partum[tiab])153623183,012#3(antenatal[tiab] OR ante natal[tiab] OR ante-natal[tiab] OR prenatal[tiab] OR pre natal[tiab] OR pre- natal[tiab])115113139,910#2(prepartum[tiab] OR pre partum[tiab] OR pre-natal[tiab] OR pre- OR intra-partum[tiab] OR perinatal[tiab] OR prenatal[tiab] OR perinatal[tiab] OR perinatal[tiab] OR perinatal[tiab] OR perinatal[tiab]7316988,660#1(Obstetric[tiab] OR obstetrics[tiab] OR pregnancy[tiab] OR maternal[tiab])554296657,008	#6	(fetus[tiab] OR foetus[tiab] OR fetu*[tiab] OR foetu*[tiab] OR fetal*[tiab] OR foetal*[tiab])	299655	337,336
#4(postnatal[tiab] OR post natal[tiab] OR post-natal[tiab] OR postpartum[tiab] OR post partum[tiab] OR post-partum[tiab])153623183,012#3(antenatal[tiab] OR ante natal[tiab] OR ante-natal[tiab] OR prenatal[tiab] OR pre natal[tiab] OR pre natal[tiab] OR pre- natal[tiab])115113139,910#2(prepartum[tiab] OR pre partum[tiab] OR pre-partum[tiab] OR preinatal[tiab] OR perinatal[tiab] OR perinatal[tiab] OR perinatal[tiab] OR perinatal[tiab] OR perinatal[tiab]7316988,660#1(Obstetric[tiab] OR obstetrics[tiab] OR pregnancy[tiab] OR maternal[tiab])554296657,008	#5	(#1 OR #2 OR #3 OR #4)	738834	867,966
#3(antenatal[tiab] OR ante natal[tiab] OR ante-natal[tiab] OR prenatal[tiab] OR pre natal[tiab] OR pre- natal[tiab])115113139,910#2(prepartum[tiab] OR pre partum[tiab] OR pre-partum[tiab] OR prinatal[tiab] OR perinatal[tiab] OR perinatal[tiab] OR perinatal[tiab]7316988,660#1(Obstetric[tiab] OR obstetrics[tiab] OR pregnancy[tiab] OR maternal[tiab])554296657,008	#4	(postnatal[tiab] OR post natal[tiab] OR post-natal[tiab] OR postpartum[tiab] OR post partum[tiab] OR post-partum[tiab])	153623	183,012
#2(prepartum[tiab] OR pre partum[tiab] OR pre-partum[tiab] OR intrapartum[tiab] OR intrapartum[tiab] OR intrapartum[tiab] OR intrapartum[tiab] OR intrapartum[tiab]7316988,660#1(Obstetric[tiab] OR obstetrics[tiab] OR pregnancy[tiab] OR maternal[tiab])554296657,008	#3	(antenatal[tiab] OR ante natal[tiab] OR ante-natal[tiab] OR prenatal[tiab] OR pre natal[tiab] OR pre- natal[tiab])	115113	139,910
#1 (Obstetric[tiab] OR obstetrics[tiab] OR pregnancy[tiab] OR maternal[tiab]) 554296 657,008	#2	(prepartum[tiab] OR pre partum[tiab] OR pre-partum[tiab] OR intrapartum[tiab] OR intra partum[tiab] OR intra partum[tiab] OR perinatal[tiab] OR pe	73169	88,660
	#1	(Obstetric[tiab] OR obstetrics[tiab] OR pregnancy[tiab] OR maternal[tiab])	554296	657,008

CINAHL

Table A.5 Search results Questions 1 to 4: CINAHL

Searched conducted

#	Query	Results 20 July 2018	Results 27 Sept 2021
S1	(MH "Obstetrics") or (MH "Obstetric Care+") or (MH "Pregnancy+") or "pregnancy disorder" or "prenatal disorder"	128,235	240,984
S2	TI (obstetric or obstetrics or pregnancy or maternal) OR AB (obstetric or obstetrics or pregnancy or maternal) or ("obstetric" or "obstetrics" or "pregnancy" or "maternal")	157,109	308,803
S 3	TI (obstetric or obstetrics or pregnancy or maternal) OR AB (obstetric or obstetrics or pregnancy or maternal) or ("obstetric" or "obstetrics" or "pregnancy" or "maternal")	157,109	187,886
S4	TI (antenatal or ante natal or ante-natal or prenatal or pre natal or pre-natal) OR AB (antenatal or ante natal or ante-natal or prenatal or pre-natal) OR ("antenatal" or "ante natal" or "ante-natal" or "prenatal" or "pre-natal")	29,500	66,056
S5	TI (postnatal or post natal or post-natal or postpartum or post partum or post-partum) OR AB (postnatal or post-natal or postpartum or post partum or post-partum) OR ("postnatal" or "post natal" or "post natal" or "post partum" or "post-partum")	23,198	50,224
S6	S1 OR S2 OR S3 OR S4 OR S5	172,767	345,785
S 7	(MH "Fetus+")	17,301	26,623
S 8	TI (fetu* or fetal* or f#etu* or f#etal*) OR AB (fetu* or fetal* or f#etu* or f#etal*) OR ("fetu*" or "fetal*" or "f#etu*")	2,598,213	82,570
S9	S7 OR S8	2,598,237	82,893
S10	"alloimmuni?ation"	343	692
S11	(MH "RH Isoimmunization")	275	458
S12	TI (Rh* Isoimmuni?ation or Rh* D Isoimmuni?ation) OR AB (Rh* Isoimmuni?ation or Rh* D Isoimmuni?ation)	29	61
S13	TI (Rh* alloimmuni?ation or Rh* D alloimmuni?ation) OR AB (Rh* alloimmuni?ation or Rh* D alloimmuni?ation)	25	111
S14	TI (Rh* incompatibility or Rh* D incompatibility or blood group incompatibility) OR AB (Rh* incompatibility or Rh* D incompatibility or blood group incompatibility)	49	115
S15	TI ((Rh* N3 incompatib*) OR (Rh* D N3 incompatibl*)) OR AB ((Rh* N3 incompatib*) OR (Rh* D N3 incompatibl*))	37	82
S16	TI ((Rh or RhD or rhesus) N5 sensiti*) OR AB ((Rh or RhD or rhesus) N5 sensiti*)	60	124
S17	TI (fetomaternal or feto-maternal or foetomaternal or foeto-maternal) N2 immuni?ation) OR AB (fetomaternal or feto-maternal or foetomaternal or foeto-maternal) N2 immuni?ation)	2	2
S18	TI (((rh or RhD or rhesus) N2 (immuni?ation or autoimmuni?ation))) OR AB (((rh or rhesus) N2 (immuni?ation or autoimmuni?ation)))	10	17
S19	S10 OR S11 OR S12 OR S13 OR S14 OR S15 OR S16 OR S17 OR S18	656	1,247
S20	"rhesus D antigen"	2	4
S21	TI rhesus D antigen OR AB rhesus D antigen	3	9
S22	TI rh* D antigen OR AB rh* D antigen	36	64
S23	TI (RhD or rhesus D or Rh D or Rh-D) OR AB (RhD or rhesus D or Rh D or Rh-D)	528	1,117
S24	TI (Rh negative OR Rh positive) OR AB (Rh negative OR Rh positive))	88	194
S25	TI (Rhesus negative or Rhesus positive) OR AB (Rhesus negative or Rhesus positive)	32	78
S26	TI (rh or rhesus) N2 (factor or factors or antigen* or system or group)) OR AB (rh or rhesus) N2 (factor or factors or antigen* or system or group))	156	439
S27	S20 OR S21 OR S22 OR S23 OR S24 OR S25 OR S26	688	1,551
S28	TI (Macaca mulatta or Simian Immunodeficiency Virus or zika or macaque#) OR AB (Macaca mulatta or Simian Immunodeficiency Virus or zika or macaque#)	1,516	4,005
S29	S27 NOT S28	683	1,526
S30	TI (isoimmuni?ation or alloimmuni?ation) OR AB (isoimmuni?ation or alloimmuni?ation) OR ("isoimmuni?ation" or "alloimmuni?ation")	579	1,099
S31	TI (isoimmuni* or iso-immuni* or isoimmune or iso-immune) OR AB (isoimmuni* or iso-immuni* or isoimmune) OR ("isoimmuni*" or "iso-immuni*" or "isoimmune")	310	552
S32	TI (alloimmuni [*] or allo-immuni [*] or alloimmune or allo-immune) OR AB (alloimmuni [*] or allo-immuni [*] or alloimmune or allo-immune) OR ("alloimmuni ^{**} or "allo-immuni ^{**} or "alloimmune" or "allo-immune")	607	1,251

S33 TI (unsensiti?ed or un-sensiti?ed or non-sensiti?ed) OR AB (unsensiti?ed or un-sensiti?ed or non-sensiti?ed) OR ("unsensiti?ed" or "un-sensiti?ed") 20 61 S34 TI (sensiti?ation* or sensiti?ed) OR AB (sensiti?ation* or sensiti?ed) OR ("sensiti?ation*" or "sensiti?ed") 3,426 8,596 S35 S30 OR S31 OR S32 OR S33 OR S34 4,227 10,186 S36 (MH "Erythroblastosis, Fetal+") 616 1,202 TI ((erythroblastoses or erythroblastosis) N2 (fetal* or f#etal*))) OR AB (((erythroblastoses or erythroblastosis) N2 (fetal* or f#etal*))) OR AB ((t*tomolytic discoses* or b#tomolytic discoses* or b#tomo	-
S34 TI (sensiti?ation* or sensiti?ed) OR AB (sensiti?ation* or sensiti?ed) OR ("sensiti?ation*" or "sensiti?ed") 3,426 8,596 S35 S30 OR S31 OR S32 OR S33 OR S34 4,227 10,186 S36 (MH "Erythroblastosis, Fetal+") 616 1,202 TI (((erythroblastoses or erythroblastosis) N2 (fetal* or f#etal*))) OR AB (((erythroblastoses or erythroblastosis) N2 (fetal* or f#etal*))) OR AB (((erythroblastosis") N2 ("fetal*" or 240 437 TI ((https://tipe.com/site	
S35 S30 OR S31 OR S32 OR S33 OR S34 4,227 10,186 S36 (MH "Erythroblastosis, Fetal+") 616 1,202 T1 (((erythroblastosis) N2 (fetal* or f#etal*))) OR AB (((erythroblastoses or erythroblastosis) N2 (fetal* or f#etal*))) OR AB (((erythroblastosis") N2 ("fetal*" or 240 437 T1 (((http://trobulastosis) N2 (fetal* or f#etal*))) OR AB ((http://trobulastosis") N2 ("fetal*" or 240 437 T1 ((http://trobulastosis) N2 (fetal* or f#etal*))) OR AB ((http://tipelastosis") N2 ("fetal*" or 240 437	
S36 (MH "Erythroblastosis, Fetal+") 616 1,202 S37 TI (((erythroblastoses or erythroblastosis) N2 (fetal* or f#etal*))) OR AB (((erythroblastoses or erythroblastosis) N2 (fetal* or f#etal*))) OR ((("erythroblastoses" or "erythroblastosis") N2 ("fetal*" or 240 437 S17 TI ((herythroblastoses or erythroblastosis) N2 (fetal* or f#etal*))) OR ((("erythroblastoses" or "erythroblastosis") N2 ("fetal*" or 240 437 S17 TI ((herythroblastoses or erythroblastoses) or the erythroblastoses or or "erythroblastoses" or herythroblastoses or or erythroblastoses or ery	
S37 TI (((erythroblastoses or erythroblastosis) N2 (fetal* or f#etal*))) OR AB (((erythroblastoses or erythroblastosis) N2 (fetal* or f#etal*))) OR ((("erythroblastoses" or "erythroblastosis") N2 ("fetal*" or "f#etal*"))) 240 437 TI ((https://www.setalation.org/linear/setalatinatinter/setalatinatinter/setalation.org/linea	
TI ((httemolytic disease* or httemolytic disease*)) OR AB ((httemolytic disease* or httemolytic	
S38 If ((interinolytic disease of interinolytic disease in the interinolytic disease interi	
S39 TI (HDFN or HDN) OR AB (HDFN or HDN) OR ("HDFN" or "HDN") 57 141	
S40 S36 OR S37 OR S38 OR S39 873 1,824	
S41 S6 OR S9 OR S19 OR S35 OR S40 2,610,641 371,868	
S42 "Rh D immunoglobulin" 1 6	
S43 "Rho D Immune Globulin" 222 341	
S44 (MH "Rho(D) Immune Globulin") 219 338	
S45"anti-D immunoglobulin"3470	
S46 TI Rh* D Immune Globulin OR AB Rh* D Immune Globulin 27 36	
S47 TI (rh* immunoglobulin or rh* d immunoglobulin) OR AB (rh* immunoglobulin or rh* d immunoglobulin 56 155	
S48 TI (rh* immuni?ation or rh* d immuni?ation) OR AB (rh* immuni?ation or rh* d immuni?ation) 24 49	
S49 S42 OR S43 OR S44 OR S45 OR S46 OR S47 OR S48 298 536	
S50 "rhesus D antibody" 0 0	
S51 TI rhesus D antibody OR AB rhesus D antibody 4 12	
S52 TI (rh* D antibody or rh*D antibody) OR AB (rh* D antibody or rh*D antibody) 409 1,108	
S53TI (anti-D or anti D or anti?D) OR AB (anti-D or anti D or anti?D)4661,332	
S54 S50 OR S51 OR S52 OR S53 849 2,373	
S55 "rhogam" 7 17	
S56 TI rhogam OR AB rhogam 7 17	
S57 "winrho" 6 7	
S58 TI winrho OR AB winrho 6 7	
S59 TI rhophylac OR AB rhophylac OR "rhophylac" 2 2	
S60 TI RhD immunoglobulin vf OR AB RhD immunoglobulin vf OR "RhD immunoglobulin vf" 0 0	
S61 TI MICRhoGam OR AB MICRhoGam OR "MICRhoGam" 0 0	
S62 TI BayRHo-D OR AB BayRHo-D OR "BayRHo-D" 0 0	
S63 TI rhesonativ OR AB rhesonativ OR "rhesonativ" 0 0	
S64 TI RhD immunoglobulin vf OR AB RhD immunoglobulin vf 0 0	
S65 S55 OR S56 OR S57 OR S58 OR S59 OR S60 OR S61 OR S62 OR S63 OR S64 15 26	
S66 S49 OR S54 OR S65 1,019 2,660	
S67 S41 AND S66 973 657	
S68PT (Editorial or letter or comment or historical article)364,194689,076	
S69 S67 NOT S68 920 610	
S70 S67 NOT S68 Limiters - Date Published: 20180101-20211231 NA 147	

A2 Subquestion 3

Embase

Table A.6	Search results subquestion 3: Embase	(via Ovid) for Level I, Level II and Level III studies
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#	Searches	Results ^a 19 July 2018	Results ^b 28 Sept 2021
1	exp Prenatal Diagnosis/	100703	114220
2	Maternal Serum Screening Tests/	232	301
3	Hematologic Tests/	12148	14896
4	((prenatal or pre-natal or antenatal or ante-natal) adj3 (test* or screen* or diagnos* or determin* or detect*)).ti,ab.	47618	55601
5	(f?etal adj3 (test* or screen* or diagnos* or determin* or detect*)).ti,ab.	23936	27181
6	((non-invasive adj7 screening) or (non?invasive adj7 screening)).ti,ab.	4412	6066
7	(NIPD or NIPT or NIPS or NIPA).ti,ab.	2110	3497
8	or/1-7	144660	167126
9	Cell-Free Nucleic Acids/	0	1254
10	(cffCDNA or cell-free f?etal DNA).ti,ab.	1056	1354
11	((cell free dna or cfDNA) adj3 (obstetric or obstetrics or pregnancy or maternal)).ti,ab.	443	672
12	((cell free dna or cfDNA) adj3 (fetu* or fetal* or f?etu* or f?etal*)).ab,ti.	267	425
13	Genotyping Techniques/	5856	8971
14	((genotype* or genotyping) adj3 (obstetric or obstetrics or pregnancy or maternal)).ti,ab.	1424	1685
15	((genotype* or genotyping) adj3 (fetu* or fetal* or f?etu* or f?etal*)).ti,ab.	1113	1352
16	(RHD adj3 gene).ti,ab.	667	794
17	or/9-16	9881	15162
18	8 or 17	152700	179708
19	exp "obstetrics"/ or exp "obstetric care"/ or exp "pregnancy"/ or exp "pregnancy disorder"/ or exp "prenatal disorder"/	1138381	1224786
20	(obstetric or obstetrics or pregnancy or maternal).kw,ab,ti.	688764	798197
21	(prepartum or pre partum or pre-partum or intrapartum or intra partum or intra-partum or perinatal or peri natal or peri-natal).kw,ab,ti.	98043	115994
22	(antenatal or ante natal or ante-natal or prenatal or pre natal or pre-natal).kw,ab,ti.	151208	178341
23	(postnatal or post natal or post-natal or postpartum or post partum or post-partum).kw,ab,ti.	194268	231147
24	or/19-23	1433924	1566408
25	exp "fetus"/	189793	199261
26	(fetu* or fetal* or f?etu* or f?etal*).kw,ab,ti.	375396	416482
27	or/25-26	428926	466603
28	exp alloimmunization/	4372	5319
29	exp Rh Isoimmunization/	1604	1500
30	(Rh* Isoimmuni?ation or Rh* D Isoimmuni?ation).ti,ab.	719	588
31	(Rh* alloimmuni?ation or Rh* D alloimmuni?ation).ti,ab.	381	456
32	(Rh* incompatibility or Rh* D incompatibility or blood group incompatibility).ti,ab.	1102	1022
33	(((Rh* adj3 incompatib*) or Rh* D) adj3 incompatibl*).ti,ab.	203	194
34	((Rh or RhD or rhesus) adj5 sensiti*).ti,ab.	1325	265
35	((fetomaternal or feto-maternal or foetomaternal or foeto-maternal) adj2 immuni?ation).ti,ab.	81	54
36	((rh or rhesus) adj2 (immuni?ation or autoimmuni?ation)).ti,ab.	770	541
37	or/28-36	9210	8692
38	exp rhesus D antigen/	1158	1532
39	rhesus D antigen.ti,ab.	55	57
40	rh* D antigen.ti,ab.	234	241
41	(RhD or rhesus D or Rh?D or Rh-?D or Rh D).ti,ab.	7551	9309
42	(Rh-negative or Rh-positive).ti,ab.	1311	1376
43	(Rhesus negative or Rhesus positive).ti,ab.	362	382
44	((rh or rhesus) adj2 (factor or factors or antigen* or system or group)).ti,ab.	4806	5015

#	Searches	Results ^a 19 July 2018	Results ^b 28 Sept 2021
45	or/38-44	12533	14306
46	(Macaca mulatta or Simian Immunodeficiency Virus or zika or macaque?).ti,ab.	38059	48547
47	45 not 46	12358	14114
48	(isoimmuni?ation or alloimmuni?ation).ti,ab,kw.	5920	6513
49	(isoimmuni* or iso-immuni* or isoimmune or iso-immune).ti,ab,kw.	2122	1570
50	(alloimmuni* or allo-immuni* or alloimmune or allo-immune).ti,ab,kw.	11071	13031
51	(unsensiti?ed or un-sensiti?ed or non-sensiti?ed).ti,ab,kw.	2409	2617
52	(sensiti?ation* or sensiti?ed).ti,ab,kw.	119495	134779
53	or/48-52	132300	148922
54	exp Erythroblastosis, Fetal/	11404	9270
55	((erythroblastoses or erythroblastosis) adj2 (fetal* or f?etal*)).kw,ab,ti.	1103	266
56	(h?emolytic disease* or h?emolytic disorder*).ti,ab,kw.	5204	4940
57	(HDFN or HDN).ti,ab,kw.	1169	1533
58	or/54-57	14942	12898
59	24 or 27	1568681	1707662
60	37 or 47 or 53 or 58	156987	172822
61	59 and 60	23151	21809
62	18 and 61	5024	5588
63	(diagnos*.mp. and (exp performance/ or yield.mp.)) or accura*.mp. or exp accuracy/ or exp diagnostic accuracy/ or sensitivity.mp. or specificity.mp. or exp "sensitivity and specificity"/ or exp "specificity and sensitivity"/ or exp precision/ or exp positive predictive value/ or exp negative predictive value/ or positive likelihood ratio.mp. or exp negative predictive value/ or positive likelihood ratio.mp. or negative predictive value/ or positive likelihood ratio.mp. or negative predictive value/ or positive likelihood ratio.mp. or exp negative predictive value/ or positive likelihood ratio.mp. or negative predictive value/ or positive likelihood ratio.mp. or negative ikelihood ratio.mp. or receiver operating.mp. or diagnostic odds.mp. or ppv.mp. or npv.mp. or plr.mp. or nlr.mp. or roc.mp. or exp sroc/ or dor.mp. or exp reliability/ or repeatability.mp. or exp reproducibility/ or reference standard.mp. or index test.mp. or reference test.mp. or exp gold standard/ or exp false positive result/ or exp false negative result/ or true positive.mp. or true negative.mp. or false positive.mp. or false negative.mp. or concord*.mp. or agreement.mp. or correlat*.mp. or accord*.mp. or (predictive adj4 value).mp.	5845182	7304460
64	62 and 63	1442	1702
65	(editorial or letter or comment or historical article).pt.	1600105	1895537
66	64 not 65	1415	1675
67	(animals/ or nonhuman/) not humans/	6124874	6443842
68	66 not 67	1402	1659
69	limit 68 to yr="2018 -Current"	NA	312

a. Embase <1974 to 2018 July 17>

b. Embase <1974 to 2021 September 24>

MEDLINE

Table A.7 Search results subquestion 3: Medline (via Ovid) for Level I, Level II and Level III studies

#	Searches	Results ^a 19 July 2018	Results ^b 28 Sept 2021
1	exp Prenatal Diagnosis/	68829	76931
2	Maternal Serum Screening Tests/	330	531
3	Hematologic Tests/	8696	9685
4	((prenatal or pre-natal or antenatal or ante-natal) adj3 (test* or screen* or diagnos* or determin* or detect*)).ti,ab.	36975	42346
5	(f?etal adj3 (test* or screen* or diagnos* or determin* or detect*)).ti,ab.	17863	20221
6	((non-invasive adj7 screening) or (non?invasive adj7 screening)).ti,ab.	2893	3832
7	(NIPD or NIPT or NIPS or NIPA).ti,ab.	1344	2172
8	or/1-7	104364	118015
9	Cell-Free Nucleic Acids/	198	1982
10	(cffCDNA or cell-free f?etal DNA).ti,ab.	666	847

#	Searches	Results ^a 19 July 2018	Results ^b 28 Sept 2021
11	((cell free dna or cfDNA) adj3 (obstetric or obstetrics or pregnancy or maternal)).ti,ab.	273	263
12	((cell free dna or cfDNA) adj3 (fetu* or fetal* or f?etu* or f?etal*)).ab,ti.	157	410
13	Genotyping Techniques/	5403	7844
14	((genotype* or genotyping) adj3 (obstetric or obstetrics or pregnancy or maternal)).ti,ab.	1115	1305
15	((genotype* or genotyping) adj3 (fetu* or fetal* or f?etu* or f?etal*)).ti,ab.	779	922
16	(RHD adj3 gene).ti,ab.	323	363
17	or/9-16	8282	12920
18	8 or 17	111426	129150
19	exp "obstetrics"/ or exp "obstetric care"/ or exp "pregnancy"/ or exp "pregnancy disorder"/ or exp "prenatal disorder"/	846400	945998
20	(obstetric or obstetrics or pregnancy or maternal).kw,ab,ti.	542881	643633
21	(prepartum or pre partum or pre-partum or intrapartum or intra partum or intra-partum or perinatal or peri natal or peri-natal).kw,ab,ti.	72582	87299
22	(antenatal or ante natal or ante-natal or prenatal or pre natal or pre-natal).kw,ab,ti.	113470	137293
23	(postnatal or post natal or post-natal or postpartum or post partum or post-partum).kw,ab,ti.	152751	181566
24	or/19-23	1127977	1280398
25	exp "fetus"/	151416	161305
26	(fetu* or fetal* or f?etu* or f?etal*).kw,ab,ti.	296787	333936
27	or/25-26	368773	409865
28	exp alloimmunization/	0	0
29	exp Rh Isoimmunization/	1672	1753
30	(Rh* Isoimmuni?ation or Rh* D Isoimmuni?ation).ti,ab.	602	614
31	(Rh* alloimmuni?ation or Rh* D alloimmuni?ation).ti,ab.	215	256
32	(Rh* incompatibility or Rh* D incompatibility or blood group incompatibility).ti,ab.	909	973
33	(((Rh* adj3 incompatib*) or Rh* D) adj3 incompatibl*).ti,ab.	155	166
34	((Rh or RhD or rhesus) adj5 sensiti*).ti,ab.	1195	1225
35	((fetomaternal or feto-maternal or foetomaternal or foeto-maternal) adj2 immuni?ation).ti,ab.	78	78
36	((rh or rhesus) adj2 (immuni?ation or autoimmuni?ation)).ti.ab.	713	780
37	or/28-36	4718	5020
38	exp rhesus D antigen/	0	38
39	rhesus D antigen ti ab.	37	0
40	rh* D antigen.ti.ab.	183	189
41	(RhD or rhesus D or Rh?D or Rh-?D or Rh D).ti.ab.	4499	5412
42	(Rh-negative or Rh-positive).ti.ab.	951	1026
43	(Rhesus negative or Rhesus positive) ti ab	238	258
44	((rh or rhesus) adi2 (factor or factors or antigen* or system or group)) ti ab.	3881	4358
45	or/38-44	8683	10069
46	(Macaca mulatta or Simian Immunodeficiency Virus or zika or macague?) ti ab	32790	41471
47	45 not 46	8526	9889
48	(isoimmuni?ation or alloimmuni?ation) ti ab.kw.	3791	4313
49	(isoimmuni* or iso-immuni* or isoimmune or iso-immune) ti ab kw	2000	2064
50	(alloimmuni* or allo-immuni* or alloimmune or allo-immune) ti ab kw	6616	7647
51	(unsensiti?ed or un-sensiti?ed or non-sensiti?ed) ti ab kw	1629	1750
52	(sensiti?ation* or sensiti?ed) ti ab kw	90214	103333
53	or/48-52	98682	112853
54	exp Ervthroblastosis. Fetal/	11580	12015
55	((ervthroblastoses or ervthroblastosis) adi2 (fetal* or f?etal*)) kw ab ti	858	908
56	(h?emolvtic disease* or h?emolvtic disorder*) ti ab kw	4554	4970
57	(HDEN or HDN) ti ab kw.	552	722
58	or/54-57	13562	14346
59	24 or 27	1260402	1423630
60	37 or 47 or 53 or 58	118578	134150

#	Searches	Results ^a 19 July 2018	Results ^b 28 Sept 2021
61	59 and 60	17363	18649
62	18 and 61	2898	3176
63	(diagnos*.mp. and (exp performance/ or yield.mp.)) or accura*.mp. or exp accuracy/ or exp diagnostic accuracy/ or sensitivity.mp. or specificity.mp. or exp "sensitivity and specificity"/ or exp "specificity and sensitivity"/ or exp precision/ or exp positive predictive value/ or exp negative predictive value/ or positive likelihood ratio.mp. or exp negative predictive value/ or positive likelihood ratio.mp. or negative likelihood ratio.mp. or exp sensitivity or exp receiver operating.mp. or diagnostic odds.mp. or ppv.mp. or npv.mp. or plr.mp. or nlr.mp. or roc.mp. or exp sroc/ or dor.mp. or exp reliability/ or repeatability.mp. or exp false positive result/ or exp false negative result/ or true positive.mp. or correlat*.mp. or false positive.mp. or false negative.mp. or concord*.mp. or agreement.mp. or correlat*.mp. or accord*.mp. or (predictive adj4 value).mp.	4531794	5539793
64	62 and 63	716	816
65	(editorial or letter or comment or historical article).pt.	1970016	2342764
66	64 not 65	702	802
67	(animals/ or nonhuman/) not humans/	4441716	4856723
68	66 not 67	699	799
69	limit 68 to yr="2018 -Current"	NA	106

a. Ovid MEDLINE(R) Epub Ahead of Print, In-Process & Other Non-Indexed Citations, Ovid MEDLINE(R) Daily, Ovid MEDLINE and Versions(R) <1946 to May 30, 2018>

b. Ovid MEDLINE(R) ALL <1946 to September 24, 2021>

Evidence-Based Medicine Reviews

Table A.8 Search results subquestion 3: EBM Reviews

#	Searches	Results ^a 19 July 2018	Results ^b 28 Sept 2021
1	exp Prenatal Diagnosis/	939	1055
2	Maternal Serum Screening Tests/	8	9
3	Hematologic Tests/	204	228
4	((prenatal or pre-natal or antenatal or ante-natal) adj3 (test* or screen* or diagnos* or determin* or detect*)).ti,ab.	852	1287
5	(f?etal adj3 (test* or screen* or diagnos* or determin* or detect*)).ti,ab.	692	1025
6	((non-invasive adj7 screening) or (non?invasive adj7 screening)).ti,ab.	159	260
7	(NIPD or NIPT or NIPS or NIPA).ti,ab.	137	281
8	or/1-7	2540	3613
9	Cell-Free Nucleic Acids/	3	18
10	(cffCDNA or cell-free f?etal DNA).ti,ab.	14	18
11	((cell free dna or cfDNA) adj3 (obstetric or obstetrics or pregnancy or maternal)).ti,ab.	11	16
12	((cell free dna or cfDNA) adj3 (fetu* or fetal* or f?etu* or f?etal*)).ab,ti.	11	12
13	Genotyping Techniques/	60	85
14	((genotype* or genotyping) adj3 (obstetric or obstetrics or pregnancy or maternal)).ti,ab.	31	57
15	((genotype* or genotyping) adj3 (fetu* or fetal* or f?etu* or f?etal*)).ti,ab.	13	24
16	(RHD adj3 gene).ti,ab.	3	4
17	or/9-16	122	2-Jun
18	8 or 17	2636	3779
19	exp "obstetrics"/ or exp "obstetric care"/ or exp "pregnancy"/ or exp "pregnancy disorder"/ or exp "prenatal disorder"/	19993	23474
20	(obstetric or obstetrics or pregnancy or maternal).kw,ab,ti.	37479	71083
21	(prepartum or pre partum or pre-partum or intrapartum or intra partum or intra-partum or perinatal or perinatal or perinatal or perinatal).kw,ab,ti.	4694	7790
22	(antenatal or ante natal or ante-natal or prenatal or pre natal or pre-natal).kw,ab,ti.	5834	10176
23	(postnatal or post natal or post-natal or postpartum or post partum or post-partum).kw,ab,ti.	8586	15414

#	Searches	Results ^a 19 July 2018	Results ^b 28 Sept 2021
24	or/19-23	50733	88340
25	exp "fetus"/	1614	1812
26	(fetu* or fetal* or f?etu* or f?etal*).kw,ab,ti.	8812	15176
27	or/25-26	9664	16152
28	exp alloimmunization/	0	0
29	exp Rh Isoimmunization/	30	32
30	(Rh* Isoimmuni?ation or Rh* D Isoimmuni?ation).ti,ab.	13	18
31	(Rh* alloimmuni?ation or Rh* D alloimmuni?ation).ti,ab.	6	14
32	(Rh* incompatibility or Rh* D incompatibility or blood group incompatibility).ti,ab.	25	52
33	(((Rh* adj3 incompatib*) or Rh* D) adj3 incompatibl*).ti,ab.	2	4
34	((Rh or RhD or rhesus) adj5 sensiti*).ti,ab.	23	28
35	((fetomaternal or feto-maternal or foetomaternal or foeto-maternal) adj2 immuni?ation).ti,ab.	0	0
36	((rh or rhesus) adj2 (immuni?ation or autoimmuni?ation)).ti,ab.	27	33
37	or/28-36	113	164
38	exp rhesus D antigen/	0	0
39	rhesus D antigen.ti,ab.	0	0
40	rh* D antigen.ti,ab.	0	0
41	(RhD or rhesus D or Rh?D or Rh-?D or Rh D).ti,ab.	139	217
42	(Rh-negative or Rh-positive).ti,ab.	23	46
43	(Rhesus negative or Rhesus positive) ti,ab.	17	22
44	((rh or rhesus) adj2 (factor or factors or antigen* or system or group)).ti,ab.	117	162
45	or/38-44	283	418
46	(Macaca mulatta or Simian Immunodeficiency Virus or zika or macague?).ti.ab.	151	281
47	45 not 46	283	418
48	(isoimmuni?ation or alloimmuni?ation).ti,ab,kw.	175	262
49	(isoimmuni* or iso-immuni* or isoimmune or iso-immune).ti,ab,kw.	46	65
50	(alloimmuni* or allo-immuni* or alloimmune or allo-immune) ti ab kw.	266	406
51	(unsensiti?ed or un-sensiti?ed or non-sensiti?ed).ti.ab.kw.	49	71
52	(sensiti?ation* or sensiti?ed).ti.ab.kw.	2899	4409
53	or/48-52	3200	4861
54	exp Erythroblastosis, Fetal/	70	76
55	((erythroblastoses or erythroblastosis) adj2 (fetal* or f?etal*)).kw,ab,ti.	14	10
56	(h?emolytic disease* or h?emolytic disorder*).ti,ab.kw.	106	157
57	(HDFN or HDN).ti,ab,kw.	21	32
58	or/54-57	150	210
59	24 or 27	52330	90952
60	37 or 47 or 53 or 58	3577	5408
61	59 and 60	367	565
62	18 and 61	38	48
63	(diagnos*.mp. and (exp performance/ or yield.mp.)) or accura*.mp. or exp accuracy/ or exp diagnostic accuracy/ or sensitivity.mp. or specificity.mp. or exp "sensitivity and specificity"/ or exp "specificity and sensitivity"/ or exp precision/ or exp positive predictive value/ or exp negative predictive value/ or positive likelihood ratio.mp. or exp negative predictive value/ or positive likelihood ratio.mp. or negative predictive value/ or positive likelihood ratio.mp. or por negative predictive value/ or positive likelihood ratio.mp. or preceiver operating.mp. or diagnostic odds.mp. or ppv.mp. or npv.mp. or plr.mp. or nlr.mp. or roc.mp. or exp sroc/ or dor.mp. or exp reliability/ or repeatability.mp. or exp reproducibility/ or reference standard.mp. or index test.mp. or reference test.mp. or exp gold standard/ or exp false positive result/ or true positive.mp. or false positive.mp. or false negative.mp. or concord*.mp. or gareement.mp. or correlat*.mp. or accord*.mp. or (predictive adi4	233823	340240
	value).mp.		
64	62 and 63	25	24
65	(editorial or letter or comment or historical article).pt.	7477	8703
66	64 not 65	25	24
67	(animals/ or nonhuman/) not humans/	25	27
68	66 not 67	25	24

#	Searches	Results ^a 19 July 2018	Results ^b 28 Sept 2021
69	limit 68 to yr="2018 -Current"	NA	2

a. EBM Reviews combines several resources into a single database and includes the following: ACP Journal Club <1991 to June 2018>; Cochrane Database of Systematic Reviews <2005 to July 18, 2018>; Database of Abstracts of Reviews of Effects <1st Quarter 2016>; Cochrane Clinical Answers <June 2018>; Cochrane Central Register of Controlled Trials <June 2018>; Cochrane Methodology Register <3rd Quarter 2012>; Health Technology Assessment <4th Quarter 2016>; NHS Economic Evaluation Database <1st Quarter 2016>.

b. EBM Reviews - Cochrane Database of Systematic Reviews <2005 to September 23, 2021>; EBM Reviews - ACP Journal Club <1991 to August 2021>; EBM Reviews - Database of Abstracts of Reviews of Effects <1st Quarter 2016>; EBM Reviews - Cochrane Clinical Answers <September 2021>; EBM Reviews - Cochrane Central Register of Controlled Trials <August 2021>; EBM Reviews - Cochrane Methodology Register <3rd Quarter 2012>; EBM Reviews - Health Technology Assessment <4th Quarter 2016>; EBM Reviews - NHS Economic Evaluation Database <1st Quarter 2016>

PubMed

The PubMed search is restricted to records that are not indexed for MEDLINE (i.e. in-process citations and citations from journals (or parts of journals) that are not currently MEDLINE-indexed) and to records added to PubMed since January 2006.

The search comprises free-text terms only and replicates the free-text sets in the Embase search (converted from the Ovid syntax).

#	Search terms	Results 20 July 2018	Results 28 Sept 2021
#1	(Maternal[tiab] OR obstetric[tiab] OR obstetrics[tiab] OR pregnant[tiab] OR pregnancy[tiab] OR prenatal[tiab] OR pre-natal[tiab]) AND (serum[tiab] OR sera[tiab]) AND (test[tiab] OR tests[tiab] OR testing[tiab] OR screen*[tiab] OR diagnos*[tiab] OR determin*[tiab] OR detect*[tiab])	21171	31,755
#2	(Blood[tiab] OR serum[tiab] OR sera[tiab] OR haematologic*[tiab] OR hematologic*[tiab]) AND (test[tiab] OR tests[tiab] OR testing[tiab])	344458	413,112
#3	(prenatal[tiab] OR pre-natal[tiab] OR antenatal[tiab] OR ante-natal[tiab]) AND (test[tiab] OR tests[tiab] OR tests[tiab] OR tests[tiab] OR determin*[tiab] OR detect*[tiab])	70051	85,001
#4	(foetal[tiab] OR fetal[tiab]) AND (test[tiab] OR tests[tiab] OR testing[tiab] OR screen*[tiab] OR diagnos*[tiab] OR determin*[tiab] OR detect*[tiab])	103239	119,205
#5	(noninvasive[tiab] OR non-invasive[tiab]) AND (screening[tiab])	8460	11,434
#6	NIPD[tiab] OR NIPT[tiab] OR NIPS[tiab] OR NIPA[tiab]	1368	2,356
#7	#1 OR #2 OR #3 OR #4 OR #5 OR #6	502555	603,704
#8	cffCDNA[tiab] OR cell free fetal DNA[tiab] OR cell free foetal DNA[tiab]	706	919
#9	(cell free dna[tiab] OR cfDNA[tiab]) AND (obstetric[tiab] OR obstetrics[tiab] OR pregnancy[tiab] OR maternal[tiab])	576	950
#10	(cell free dna[tiab] OR cfDNA[tiab]) AND (fetu*[tiab] OR fetal*[tiab] OR foetu*[tiab] OR foetal*[tiab])	593	976
#11	(genotype[tiab] OR genotyping[tiab] OR allele[tiab] OR alleles[tiab]) AND (test[tiab] OR tests[tiab] OR tests[tiab] OR tests[tiab] OR diagnos*[tiab] OR determin*[tiab] OR detect*[tiab])	180704	216,412
#12	(genotype*[tiab] OR genotyping[tiab]) AND (obstetric[tiab] OR obstetrics[tiab] OR pregnancy[tiab] OR maternal[tiab])	7324	8,971
#13	(genotype*[tiab] OR genotyping[tiab]) AND (fetu*[tiab] OR fetal*[tiab] OR foetu*[tiab] OR foetal*[tiab])	3628	4,398
#14	(RHD[tiab] AND gene[tiab])	607	730
#15	#8 OR #9 OR #10 OR #11 OR #12 OR #13 OR #14	186624	223,843
#16	#7 OR #15	675038	810,096
#17	Obstetric[tiab] OR obstetrics[tiab] OR pregnancy[tiab] OR maternal[tiab]	554296	657,138
#18	prepartum[tiab] OR pre partum[tiab] OR pre-partum[tiab] OR intrapartum[tiab] OR intra partum[tiab] OR intra-partum[tiab] OR perinatal[tiab] OR peri natal[tiab] OR peri-natal[tiab]	73169	88,677
#19	antenatal[tiab] OR ante natal[tiab] OR ante-natal[tiab] OR prenatal[tiab] OR pre natal[tiab] OR pre- natal[tiab]	115113	139,946
20#	postnatal[tiab] OR post natal[tiab] OR post-natal[tiab] OR postpartum[tiab] OR post partum[tiab] OR post-partum[tiab]	153623	183,050
#21	#17 OR #18 OR #19 OR #20	738834	868,132

Table A.9 Search results subquestion 3: Pubmed (not MEDLINE)

#	Search terms	Results 20 July 2018	Results 28 Sept 2021
#22	fetus[tiab] OR foetus[tiab] OR fetu*[tiab] OR foetu*[tiab] OR fetal*[tiab] OR foetal*[tiab]	299655	337,391
#23	alloimmunization[tiab] OR alloimmunisation[tiab]	2590	3,241
#24	(rh[tiab] OR rhd[tiab] rhesus[tiab]) AND (isoimmunization[tiab] OR isoimmunisation[tiab])	63	82
#25	(rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatibility[tiab]) OR (blood group incompatibility[tiab])	1636	1,747
#26	(rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND (incompatib*[tiab])	1307	1,428
#27	(Rh[tiab] OR RhD[tiab] OR rhesus[tiab]) AND (sensiti*[tiab])	4933	5,735
#28	(fetomaternal[tiab] OR feto-maternal[tiab] OR foetomaternal[tiab] OR foeto-maternal[tiab]) AND (immunisation[tiab] OR immunization[tiab])	166	174
#29	(rh[tiab] OR RhD[tiab] OR rhesus[tiab]) AND (immunisation[tiab] OR immunization[tiab] OR autoimmunization[tiab])	2055	2,251
#30	#23 OR #24 OR #25 OR #26 OR #27 OR #28 OR #29	11065	12,723
#31	(rh[tiab] OR rhd[tiab] OR rhesus[tiab]) AND antigen[tiab]	3429	3,816
#32	RhD[tiab] OR rhesus d[tiab] OR rh-d[tiab] OR Rh D[tiab]	3887	4,753
#33	rh-negative[tiab] OR rh-positive[tiab] OR rh negative[tiab] OR rh positive[tiab]	949	1,038
#34	rhesus negative[tiab] OR rhesus positive[tiab]	240	261
#35	(rh[tiab] OR rhesus[tiab]) AND (factor[tiab] OR factors[tiab] OR antigen*[tiab] OR antigens[tiab] OR system[tiab] OR group[tiab])	24249	27,871
#36	#31 OR #32 OR #33 OR #34 OR #35	27458	31,809
#37	Macaca mulatta[tiab] OR Simian Immunodeficiency Virus[tiab] OR zika[tiab] OR macaque[tiab] OR macaque[tiab]		42,084
#38	#36 NOT #37	23409	27,075
#39	isoimmuni*[tiab] OR iso-immuni*[tiab] OR isoimmune[tiab] OR iso-immune[tiab]	2006	2,084
#40	alloimmuni*[tiab] OR allo-immuni*[tiab] OR alloimmune[tiab] OR allo-immune[tiab]		7,695
#41	(rh[tiab] OR rho[tiab] OR rhesus[tiab]) AND (sensitising[tiab] OR sensitizing[tiab] OR sensitisation[tiab] OR sensitised[tiab] OR sensitized[tiab])		1,735
#42	sensitisation*[tiab] OR sensitization*[tiab] OR sensitised[tiab] OR sensitized[tiab]		104,472
#43	#39 OR #40 OR #41 OR #42		113,847
#44	(erythroblastoses[tiab] OR erythroblastosis[tiab]) AND (fetal[tiab] OR foetal[tiab] OR fetalis[tiab] OR foetalis[tiab])		3,172
#45	(hemolytic OR haemolytic) AND (disorder[tiab] OR disorders[tiab] OR disease[tiab] OR diseases[tiab])	18492	19,246
#46	hdfn[tiab] OR hdn[tiab]	553	750
#47	#44 OR #45 OR #46	20920	21,750
#48	#21 OR #22	887315	1,028,420
#49	#30 OR #38 OR #43 OR #47	143560	163,294
#50	#48 AND #49	12773	13,944
#51	#16 AND #50	3926	4,511
#52	Diagnos*[tiab] AND (performance[tiab] or yield[tiab]) OR accura*[tiab] OR diagnostic accuracy[tiab] OR sensitivity[tiab] OR specificity [tiab] OR precision[tiab] OR positive predictive value [tiab] OR negative predictive value[tiab] OR positive likelihood ratio[tiab] OR negative likelihood ratio[tiab] OR receiver operating[tiab] OR diagnostic odds[tiab] OR ppv[tiab] OR npv[tiab] OR plr[tiab] OR nlr[tiab] OR ROC[tiab] OR sroc[tiab] OR dor[tiab] OR reliability[tiab] OR repeatability[tiab] OR reproducibility[tiab] OR reference standard[tiab] OR index test[tiab] OR reference test[tiab] OR gold standard[tiab] OR false positive[tiab] OR false negative[tiab] OR true positive[tiab] OR true negative[tiab] OR concord*[tiab] OR agreement[tiab] OR correlate*[tiab] OR accord*[tiab] OR (predictive[tiab] AND value[tiab])	3585762	4,486,125
#53	#51 AND #52	981	1,150
#54	#53 AND pubmednotmedline[sb]	40	72
#55	#53 AND pubmednotmedline[sb] from 2018 - 2021	NA	28

CINAHL

Table A.10Search results subquestion 3: CINAHL

#	Query	Results 19 July 2018	Results 28 Sept 2021
S1	(MH "Prenatal Diagnosis+")	8232	20,759
S2	"Maternal Serum Screening Tests"	0	5
S 3	(MH "Hematologic Tests+")	22714	49,125
S 4	TI (((prenatal or pre-natal or antenatal or ante-natal) N3 (test* or screen* or diagnos* or determin* or detect*))) OR AB (((prenatal or pre-natal or antenatal or ante-natal) N3 (test* or screen* or diagnos* or determin* or detect*)))	4126	11,292
S5	TI ((f#etal N3 (test* or screen* or diagnos* or determin* or detect*))) OR AB ((f#etal N3 (test* or screen* or diagnos* or determin* or detect*)))	2104	5,707
S 6	TI (((non-invasive N7 screening) or (non#invasive N7 screening))) OR AB (((non-invasive N7 screening) or (non#invasive N7 screening)))	333	954
S7	TI (NIPD or NIPT or NIPA or NIPS) OR AB (NIPD or NIPT or NIPA or NIPS)	222	896
S8	S1 OR S2 OR S3 OR S4 OR S5 OR S6 OR S7	34153	78,684
S9	"Cell Free Nucleic Acids" OR "Cell Free dna"	247	1,519
S10	TI ((cffCDNA or cell free f#etal DNA)) OR AB ((cffCDNA or cell free f#etal DNA))	134	495
S11	TI (((cell free dna or cfDNA) N3 (obstetric or obstetrics or pregnancy or maternal))) OR AB (((cell free dna or cfDNA) N3 (obstetric or obstetrics or pregnancy or maternal)))	96	408
S12	TI (((cell free dna or cfDNA) N3 (fetu* or fetal* or f#etu* or f#etal*))) OR AB (((cell free dna or cfDNA) N3 (fetu* or fetal* or f#etal*)))	378	491
S13	(MH "Molecular Diagnostic Techniques")	729	2,491
S14	TI (((genotype* or genotyping) N3 (obstetric or obstetrics or pregnancy or maternal))) OR AB (((genotype* or genotyping) N3 (obstetric or obstetrics or pregnancy or maternal)))	128	284
S15	TI (((genotype* or genotyping) N3 (fetu* or fetal* or f#etu* or f#etal*))) OR AB (((genotype* or genotyping) N3 (fetu* or fetal* or f#etal*)))	3291	226
S16	TI RHD N3 gene OR AB RHD N3 gene	49	84
S17	S9 OR S10 OR S11 OR S12 OR S13 OR S14 OR S15 OR S16	4459	4,728
S18	S8 OR S17	38260	82,409
S19	(MH "Obstetrics") or (MH "Obstetric Care+") or (MH "Pregnancy+") or "pregnancy disorder" or "prenatal disorder"	128201	240,985
S20	TI (obstetric or obstetrics or pregnancy or maternal) OR AB (obstetric or obstetrics or pregnancy or maternal) or ("obstetric" or "obstetrics" or "pregnancy" or "maternal")		311,683
S21	TI (prepartum or pre partum or pre-partum or intrapartum or intra partum or intra-partum or perinatal or perinatal or peri-natal) OR AB (prepartum or pre partum or pre-partum or intrapartum or intra partum or intra-partum or perinatal or perinatal or perinatal or perinatal) OR ("prepartum" or "pre-partum" or "pre-partum" or "intra-partum" or "intra-partum" or "intra-partum" or "intra-partum" or "intra-partum" or "perinatal" or "peri		39,685
S22	TI (antenatal or ante natal or ante-natal or prenatal or pre natal or pre-natal) OR AB (antenatal or ante natal or ante-natal or ante-natal or pre natal or pre-natal) OR ("antenatal" or "ante natal" or "ante-natal" or "prenatal" or "pre-natal")	29496	66,056
S23	TI (postnatal or post natal or post-natal or postpartum or post partum or post-partum) OR AB (postnatal or post natal or post-natal or postpartum or post partum or post-partum) OR ("postnatal" or "post natal" or "post natal" or "post-natal" or "post-natal" or "post partum" or "post-partum")	23191	50,224
S24	S19 OR S20 OR S21 OR S22 OR S23	176163	352,395
S25	(MH "Fetus+")	17287	26,623
S26	TI (fetu* or fetal* or f#etu* or f#etal*) OR AB (fetu* or fetal* or f#etu* or f#etal*) OR ("fetu*" or "fetal*" or "f#etu*" or "f#etal*")	2597789	82,572
S27	S25 OR 26	40390	82,928
S28	"alloimmuni?ation" TI alloimmuni?ation OR AB alloimmuni?ation OR "alloimmuni?ation"	343	692
S29	(MH "RH Isoimmunization")	275	458
S30	TI (Rh* Isoimmuni?ation or Rh* D Isoimmuni?ation) OR AB (Rh* Isoimmuni?ation or Rh* D Isoimmuni?ation)		61
S31	TI (Rh* alloimmuni?ation or Rh* D alloimmuni?ation) OR AB (Rh* alloimmuni?ation or Rh* D alloimmuni?ation)	55	111
S32	TI (Rh* incompatibility or Rh* D incompatibility or blood group incompatibility) OR AB (Rh* incompatibility or Rh* D incompatibility or blood group incompatibility)	49	115

#	Query	Results 19 July 2018	Results 28 Sept 2021
S33	TI ((Rh* N3 incompatib*) OR (Rh* D N3 incompatibl*)) OR AB ((Rh* N3 incompatib*) OR (Rh* D N3 incompatibl*))	37	82
S34	TI ((Rh or RhD or rhesus) N5 sensiti*) OR AB ((Rh or RhD or rhesus) N5 sensiti*)	60	124
S35	TI (fetomaternal or feto-maternal or foetomaternal or foeto-maternal) N2 immuni?ation) OR AB (fetomaternal or feto-maternal or foetomaternal or foeto-maternal) N2 immuni?ation)	2	2
S36	TI (((rh or RhD or rhesus) N2 (immuni?ation or autoimmuni?ation))) OR AB (((rh or rhesus) N2 (immuni?ation or autoimmuni?ation)))	10	17
S37	S28 OR S29 OR S30 OR S31 OR S32 OR S33 OR S34 OR S35 OR S36	656	1,247
S38	"rhesus D antigen"	2	4
S39	TI rhesus D antigen OR AB rhesus D antigen	3	9
S40	TI rh* D antigen OR AB rh* D antigen	36	64
S41	TI (RhD or rhesus D or Rh D or Rh-D) ORAB (RhD or rhesus D or Rh D or Rh-D)	528	1,117
S42	TI (Rh negative OR Rh positive) OR AB (Rh negative OR Rh positive))	88	194
S43	TI (Rhesus negative or Rhesus positive) OR AB (Rhesus negative or Rhesus positive)	32	78
S44	TI (rh or rhesus) N2 (factor or factors or antigen* or system or group)) OR AB (rh or rhesus) N2 (factor or factors or antigen* or system or group))	156	439
S45	S38 OR S39 OR S40 OR S41 OR S42 OR S43 OR S44	688	1,551
S46	TI (Macaca mulatta or Simian Immunodeficiency Virus or zika or macaque#) OR AB (Macaca mulatta or Simian Immunodeficiency Virus or zika or macaque#)	1514	4,005
S47	S45 NOT S46	683	1,526
S48	TI (isoimmuni?ation or alloimmuni?ation) OR AB (isoimmuni?ation or alloimmuni?ation) OR ("isoimmuni?ation" or "alloimmuni?ation")	579	1,099
S49	TI (isoimmuni* or iso-immuni* or isoimmune or iso-immune) OR AB (isoimmuni* or iso-immuni* or iso-immune) OR ("isoimmuni*" or "iso-immune" or "iso-immune")	310	552
S50	TI (alloimmuni* or allo-immuni* or alloimmune or allo-immune) OR AB (alloimmuni* or allo-immuni* or allo-immuni* or allo-immune) OR ("alloimmuni*" or "allo-immuni*" or "allo-immune")		1,251
S51	TI (unsensiti?ed or un-sensiti?ed or non-sensiti?ed) OR AB (unsensiti?ed or un-sensiti?ed or non- sensiti?ed) OR ("unsensiti?ed" or "un-sensiti?ed" or "non-sensiti?ed")		61
S52	TI (sensiti?ation* or sensiti?ed) OR AB (sensiti?ation* or sensiti?ed) OR ("sensiti?ation*" or "sensiti?ed")		8,596
S53	548 OR S49 OR S50 OR S51 OR S52		10,186
S54	(MH "Erythroblastosis, Fetal+")	616	1,202
S55	TI (((erythroblastoses or erythroblastosis) N2 (fetal* or f#etal*))) OR AB (((erythroblastoses or erythroblastosis) N2 (fetal* or f#etal*))) OR ((("erythroblastoses" or "erythroblastosis") N2 ("fetal*" or "f#etal*")))		437
S56	TI ((h#emolytic disease* or h#emolytic disorder*)) OR AB ((h#emolytic disease* or h#emolytic disorder*)) OR (("h#emolytic disease*" or "h#emolytic disorder*"))	376	825
S57	TI (HDFN or HDN) OR AB (HDFN or HDN) OR ("HDFN" or "HDN")	57	141
S58	S54 OR S55 OR S56 OR S57	873	1,824
S59	S24 OR S27	199864	408,390
S60	S37 OR S47 OR S53 OR S58	5450	12,886
S61	S59 AND S60	1142	2,427
S62	S18 AND S61	361	806
S63	(diagnos* and (performance or yield)) or (accura* or "diagnostic accuracy") or "sensitivity" or "specificity" or (MH "Sensitivity and Specificity") or (MH "Precision") or (MH "Predictive Value of Tests") or "positive predictive value" or "negative predictive value" or "positive likelihood ratio" or "negative likelihood ratio" or (MH "ROC Curve") or "receiver operating" or "diagnostic odds" or ppv or npv or plr or nlr or roc or sroc or dor or reliability or repeatability or reproducibility or "reference standard" or "index test" or "reference test" or "gold standard" or "false positive result" or (MH "False Positive Results") or "false negative result" or (MH "False Negative Results") or "true positive" or "true negative" or "false positive" or "false negative" or concord* or agreement or correlate* or accord* or (predictive N4 value) or (MH "Predictive Validity")		891,666
S64	S62 and S63	96	235
S65	PT (Editorial or letter or comment or historical article)	364150	689,083
S66	S64 NOT S65		230
S67	S64 NOT S65 Limiters - Date Published: 20180101-20211231	NA	49

Ovid syntax

Exp explodes controlled vocabulary term (i.e. includes all narrower terms in the hierarchy)
* denotes a term that has been searched as a major subject heading
/ denotes controlled vocabulary terms (EMTREE)
\$ truncation character (unlimited truncation)
\$n truncation limited to specified number (n) of characters (e.g. time\$1 identifies time, timed, timer, times but not timetable)
* truncation character (unlimited truncation)
? substitutes any letter (e.g. oxidi?ed identifies oxidised and oxidized)
adjn search terms within a specified number (n) of words from each other in any order
.ti. limit to title field
.ti,ab. limit to title and abstract fields
.kw,ti,ab. limit to keyword, title and abstract field
.pt limit to publication type

PubMed syntax

* truncation character (unlimited truncation)
[TI] limit to title field
[TIAB] limit to title and abstract fields
[EDAT] date citation added to PubMed
[SB] PubMed subset

CINHAL syntax

* truncation character (unlimited truncation)

wildcard character will replace 1 or 0 characters (e.g. f#etus will retrieve fetus and foetus)

? wildcard character will replace one character (e.g. wom?n will retrieve women and woman)

MH - Search the exact CINAHL® subject heading; searches both major and minor headings

MH"heading"+ Search an exploded subheading

TI search title fields

AB search abstract fields

Nn – Proximity "near" operator will find a result if the terms are within a certain number (n) words of each other, regardless of the order in which they appear. (e.g. eating N5 disorders for results that contain eating disorders, as well as mental disorders and eating pathology.)

PT limit to publication type

Appendix B Literature screening results (2021 update)

This appendix documents the updated literature search screening results for a systematic review on the prophylactic use of Rh D Immunoglobulin (Anti-D) in pregnant women.

A PRIMSA flow illustrating the screening results is provided in Figure 1 (all questions) and Figure 2 (subquestion 3, diagnostic accuracy).

	Questions 1-4		Q3	
Number of citations identified	Level I a	Level II (not Level I)	Level III (not Level II)	Diagnostic accuracy
Database				
Embase 1974 to 18 July 2018	27	106	302	312
MEDLINE 1946 to 18 July 2018	12	37	53	106
Cochrane 18 July 2018	0	19	0	2
PubMed	0	0	108	28
CINAHL	0	0	147	49
TOTAL	39	162	<mark>610</mark>	497
Date limit ^b	2018 to current	2018 to current	2018 to current	2018 to current
Duplicates removed in Endnote (across databases)	10	44	133	143
Duplicates removed by Covidence c	6	22	79	31
TITLE/ABSTRACT SCREENING				
Number of citations screened in Covidence	23	96	398	323
Additional duplicates identified	1	3	2	1
Nonhuman	0	0	14	0
Population out of scope	5	36	112	32
Intervention out of scope	8	27	202	214
Comparator out of scope	0	0	0	0
Outcome out of scope	0	0	0	0
Publication type out of scope. Not a systematic review.	0	0	0	2
Publication type out of scope. Opinion piece.	0	2	6	7
Publication type out of scope. Editorial.	0	0	0	0
Publication type out of scope. Other.	1	0	0	0
Study type out of scope. Level IV or below.	0	8	19	23
TOTAL irrelevant	15	76	355	279

Table B.1 Literature search and title/abstract screening results

a. NHMRC evidence level filters were applied in the Ovid interface. Studies identified in the Cochrane Collection and those retrieved via PubMed and CINAHL did not have filters applied but were screened in the first pass. (see Technical report, volume one)

b. A date limit was applied to studies based on the previous literature search date (July 2018), with the prior six months included to account for potential database changes (see Technical report, volume one).

c. https://www.covidence.org/home

Table B.2 Full text screening results

	Questions 1-4			
Number of citations identified	Level I a	Level II (not Level I)	Level III (not Level II)	Diagnostic accuracy
FULL TEXT REVIEW				
Number of citations screened in Covidence ^b	8	20	43	44
Duplicate citation	0	1	2	2
Not available in English	0	0	0	0
Population out of scope	0	1	3	3
Intervention out of scope	0	5	12	8
Comparator out of scope	0	2	0	0
Outcome out of scope	1	2	1	5
Publication type out of scope. Simple review.	0	0	1	0
Publication type out of scope. Opinion piece.	0	0	3	1
Publication type out of scope. Editorial.	0	0	2	0
Level II or III study already included in Level I	0	0	0	0
Study design out of scope (Level IV or below)	0	3	17	9
No usable data (conference abstract etc.)	0	0	2	9
Superseded	0	0	0	0
Duplicate data (published elsewhere)	0	0	0	0
Small sample size	0	3	0	4
Not comparable to the Australian context	0	1	0	0
TOTAL EXCLUDED	1	18	43	41
		-	-	
TOTAL INCLUDED	7	2	0	3

a. NHMRC evidence level filters were applied in the Ovid interface. Studies identified in the Cochrane Collection and those retrieved via PubMed and CINAHL did not have filters applied but were screened in the first pass. (see Technical report, volume one)

b. <u>https://www.covidence.org/home</u>

Appendix C Excluded studies

This appendix documents studies that are awaiting cliassification or those that met the prespecified inclusion criteria for a systematic review on the prophylactic use of Rh D Immunoglobulin (Anti-D) in pregnant women but were later excluded. These studies, and their reasons for exclusion, are listed below.

C1 Studies relevant to all Questions

No usable data (conference abstracts etc.)

Donohoe, O (2021). Cost-effectiveness of targeted antenatal anti-d in ireland. *BJOG: An International Journal of Obstetrics and Gynaecology* **128**(SUPPL 2): 125.

Donohoe, O, L Mulvany, E O'Connor, *et al.* (2019). One-year audit of targeted routine antenatal anti-d prophylaxis in portiuncula university hospital. *BJOG: An International Journal of Obstetrics and Gynaecology* **126**(Supplement 1): 99-100.

Gordon, L, R Flower and C Hyland (2018). Non-invasive fetal rhd genotyping of rhd negative pregnant women for targeted anti-d therapy in australia: A cost-effectiveness analysis. *Value in Health* **21**(Supplement 2): S93.

Matteocci, A, G Nespoli, K Castagna, *et al.* (2020). Cost and saving analysis of rhd genotyping and anti-d immuno-prophylaxis in d-variant women of childbearing age in central italy. *Vox Sanguinis* **115**(SUPPL 1): 279.

C2 Studies relevant to Question 3 (or subquestion 3)

No usable data (conference abstracts etc.)

Balsalobre, EL, RR Sanchez, MdMV Penas, *et al.* (2019). Implementation of the rhd fetal protocol in rhd negative gestants. *Clinica Chimica Acta* **493**(Supplement 1): S585-S586.

Bingulac-Popovic, J, V Dogic, I Babic, *et al.* (2018). Prenatal rhd genotyping: Automated extraction of cellfree fetal DNA using the qiasymphony sp platform. *Clinical Chemistry and Laboratory Medicine* **56**(6): eA111.

Choo, BL, M Williamson, EA Martindale, *et al.* (2019). Provision of a fetal rhd genotyping service: The east lancashire experience. *BJOG: An International Journal of Obstetrics and Gynaecology* **126**(Supplement 1): 83.

Doescher, A and C Vogt (2018). Pitfalls in prenatal diagnosis of fetal rhd: Frequency of maternal rhd variants as cause for a false positive genotype of the fetus. *Transfusion Medicine and Hemotherapy* **45**(Supplement 1): 37.

Joshi, N, S Bassiony, A Mathyalakan, *et al.* (2021). Re-audit of cell free foetal DNA (cffdna) screen to avoid administration of anti-d immunoglobulin in rhd-negative pregnant women with rhd-negative foetus. *British Journal of Haematology* **193**(SUPPL 1): 14.

Londero, D, D Bolzicco, M Candolini, *et al.* (2018). First trimester noninvasive fetal rhd genotyping using frozen DNA samples: Validation and optimization of the test to implement a screening program. *Vox Sanguinis* **113**(Supplement 1): 276-277.

Maric, I, K Zeleznik, I Bricl, *et al.* (2018). Targeted prophylaxis program for d-negative pregnant women based on genotyping fetal rhd from maternal blood. *Vox Sanguinis* **113**(Supplement 1): 277.
Small sample size (N<200)

Addai-Mensah, O, EY Afriyie, ME Annani-Akollor, *et al.* (2020). Fetal rhesus d genotyping and sex determination from maternal plasma of rhesus d-negative antenatal population: The usefulness of conventional polymerase chain reaction in resource-limited settings. *Obstetrics and Gynecology International* **2020**: 4913793.

Ahmadi, MH and N Amirizadeh (2018). Evaluation the sry to confirm the presence of fetal DNA in the fetal rhd genotyping using cffdna. *Vox Sanguinis* **113 (Supplement 1)**: 277.

Bingulac-Popovic, J, I Babic, V Dogic, *et al.* (2021). Prenatal rhd genotyping in croatia: Preliminary results. *Transfusion Clinique et Biologique* **28**(1): 38-43.

Blanco, S, MC Frutos, SV Gallego, *et al.* (2018). Usefulness of non-invasive fetal rhd genotyping towards immunoprophylaxis optimization. *Transfusion Medicine and Hemotherapy* **45**(6): 423-428.

Londero, D, D Bolzicco, M Candolini, *et al.* (2019). Fetal rhd detection from circulating cell-free fetal DNA in maternal plasma: Validation of a diagnostic kit using automatic extraction and frozen DNA. *Transfusion Medicine* **29**(6): 408-414.

Plesinac, S, D Plecas and I Babovic (2018). The determination of fetal rhd status from maternal blood in serbia. *Indian Journal of Hematology and Blood Transfusion* **34**(3): 486-490.

Rather, R, S Saha and V Dhawan (2019). Non-invasive prenatal rhesus d genotyping using cell-free foetal DNA. *Indian Journal of Medical Research* **150**(1): 62-66.

Not comparable to the Australian context

Bohmova, J, R Kratochvilova, E Krejcirikova, *et al.* (2020). Two reliable methodical approaches for non-invasive rhd genotyping of a fetus from maternal plasma. *Diagnostics* **10**(8): 564.

Appendix D Critical appraisal

D1 Question 1

Level I – Systematic review (of RCTs and cohort studies)

Question	Xie 2020	
1. Did the research questions and inclusion criteria for the review include the components of the PICO?	Yes	PICO and inclusion criteria provided.
2. Did the report of the review contain an explicit statement that the review methods were established prior to the conduct of the review and did the report justify any significant deviations from the protocol?	Partial yes	The authors refer to predetermined research objectives which note that they are searching for RCTs. They include cohort studies but do not provide a reason for this change.
3. Did the review authors explain their selection of the study designs for inclusion in the review?	No	No explanation provided
4. Did the review authors use a comprehensive literature search strategy?	Yes	A comprehensive search strategy was employed to search the PubMed, EMBASE, Web of Science, China National Knowledge Infrastructure (CNKI) and Wanfang databases. T
5. Did the review authors perform study selection in duplicate?	Yes	Two authors independently assessed all studies for inclusion and data extraction (p2)
6. Did the review authors perform data extraction in duplicate?	Yes	Two investigators collected data independently in accordance with predesigned tables (p3)
7. Did the review authors provide a list of excluded studies and justify the exclusions?	No	Reasons for exlcusion provided, but study details not provided.
8. Did the review authors describe the included studies in adequate detail?	Partial Yes	Table of study characteristics but not further described.
9. Did the review authors use a satisfactory technique for assessing the risk of bias (RoB) in individual studies that were included in the review?	No	RoB of included studies not conducted
10. Did the review authors report on the sources of funding for the studies included in the review?	No	Not reported
11. If meta-analysis was performed, did the review authors use appropriate methods for statistical combination of results?	Yes	random-effects model with parameters estimated using the Markov chain Monte Carlo method of Gibbs sampling
12. If meta-analysis was performed, did the review authors assess the potential impact of RoB in individual studies on the results of the meta-analysis or other evidence synthesis?	No	No RoB reported.
13. Did the review authors account for RoB in individual studies when interpreting/ discussing the results of the review?	Yes	No RoB reported, but authors note the inadequacies and age of the studies in their conclusions.
14. Did the review authors provide a satisfactory explanation for, and discussion of, any heterogeneity observed in the results of the review?	Yes	Fig 3 shows the inconsistency plot used to identify heterogeneity among studies in the closed loop of this network meta-analysis
15. If they performed quantitative synthesis did the review authors carry out an adequate investigation of publication bias (small study bias) and discuss its likely impact on the results of the review?	Yes	Funnel plots provided and discussed.
16. Did the review authors report any potential sources of conflict of interest, including any funding they received for conducting the review?	Yes	The authors have declared that no competing interests exist.
Overall risk of bias of the review	Moderate	More than one non-critical weakness – the systematic review has more than one weakness but no critical flaws. It may provide an accurate summary of the results of the available studies that were included in the review.

Source: Shea et al. 2017. BMJ 358: j4008 doi:10.1136 (https://doi.org/10.1136/bmj.j4008)

Level II- RCT

Study ID	White 2019		
Domain	Judgement	Description	Source
Random sequence generation (selection bias)	Low risk	Consenting participants were randomised 1:1 in blocks of ten. Presumed to be computer generated, but not reported.	p.262
Allocation concealment (selection bias)	Low risk	Sealed envelopes were used.	p.262
Blinding of participants and personnel (performance bias)	Unclear risk	No blinding used in study. Participants, clinicians, and researchers were aware of treatment allocation. Altough unlikely, this may have affected how participants were treated in followup routine care, including compliance. Twelve women in the single dose group (9%) received only 625 IU anti-D at 28–30 weeks; they were therefore given a second dose at 34–36 weeks, consistent with standard practice, to avoid potential late antenatal sensitisation.	p.262
Blinding of outcome assessment (detection bias)	Low risk	Not blinded but nature of the outcomes (objective measures relating to anti-D levels) makes it unlikely to have created bias in the results	p.262
Incomplete outcome data addressed (attrition bias)	Unclear risk	3/280 (1%) women lost to follow-up (low risk). Antibody screens were available for 254/277 (92%). No imputations/adjustments for missing data were made.	p.263
Selective reporting (reporting bias)	Low risk	Some outcomes missing as per trial registry (see ACTRN12613000661774) (total amount of Rh D lgG used per participant)	
Other sources of bias*	Low risk	Bias could exist in the results due to the high level of participants who did not receive the intervention in the required timeframe (high levels of non-compliance). A sensitivity analysis suggested this did not influence the primary outcome.	p.263
Overall risk of bias of the review	Unclear risk	One domain has some concerns raised, but none are found to b of bias	e at high risk

Source: Chapter 8 of the Cochrane Handbook for Systematic Reviews of Interventions, version 5.1.0

Note: a more recent version of the Cochrane Risk of bias tool is available (see <u>www.riskofbias.info</u>) however, we chose to use the tool specified and used in the 2018 review.

D2 Question 2

Level	I-Systematic	review	of	observational	studies

Study ID	Schmidt-Hansen 2020	
Question		
1 . Did the research questions and inclusion criteria for the review include the components of the PICO?	Yes	PICO elements are outlined (p2)
2. Did the report of the review contain an explicit statement that the review methods were established prior to the conduct of the review and did the report justify any significant deviations from the protocol?	Partial yes	No explicit statement was made about the establishment of prior review methods – except reference to the NICE guidelines. – which links to the full evidence review.
3. Did the review authors explain their selection of the study designs for inclusion in the review?	Partial yes	The study lists that it was able to select RCTs and observational studies but there wasn't an explanation of this. Full details provided in the NICE report.
4. Did the review authors use a comprehensive literature search strategy?	Yes	Yes details of the search strategy (p2)
5. Did the review authors perform study selection in duplicate?	Parital yes	Initial screening was only done by one author will full text screening performed by two
6. Did the review authors perform data extraction in duplicate?	No	Data extraction was to be performed by one author (p2)
7. Did the review authors provide a list of excluded studies and justify the exclusions?	Partial yes	Included in supplementary appendix 2 which was not to be seen in journal links. Found via NICE.
8. Did the review authors describe the included studies in adequate detail?	Partal yes	No studies were included
9. Did the review authors use a satisfactory technique for assessing the risk of bias (RoB) in individual studies that were included in the review?	Yes	Used the GRADE approach to assessing bias
10. Did the review authors report on the sources of funding for the studies included in the review?	Yes	Sources of funding are disclosed (p.5)
11. If meta-analysis was performed, did the review authors use appropriate methods for statistical combination of results?	Partial yes	No meta-analysis was performed but was planned. the The techniques planned were the Mantel-Haenszel statistical method for RRs and the inverse variance statistical method for MDs and SMDs
12. If meta-analysis was performed, did the review authors assess the potential impact of RoB in individual studies on the results of the meta-analysis or other evidence synthesis?	No	No plan for this and no meta-analysis actually performed
13. Did the review authors account for RoB in individual studies when interpreting/ discussing the results of the review?	Partial yes	No individual studies included but potentially would have used the ROB assessment from GRADE to discuss individual study bias in the results
14. Did the review authors provide a satisfactory explanation for, and discussion of, any heterogeneity observed in the results of the review?	No	No heterogeneity discussed
15. If they performed quantitative synthesis did the review authors carry out an adequate investigation of publication bias (small study bias) and discuss its likely impact on the results of the review?	No	No investigation of publication bias (no studies found)
16. Did the review authors report any potential sources of conflict of interest, including any funding they received for conducting the review?	Yes	Authors declared no conflicts of interest and funding details discussed (p5)
Overall risk of bias of the review	Low	No or one non-critical weakness – the systematic review provides an accurate and comprehensive summary of the results of the available studies that address the question of interest.

Source: Shea et al. 2017. BMJ 358: j4008 doi:10.1136 (<u>https://doi.org/10.1136/bmj.j4008</u>)

D3 Question 3

Level I - Systematic review of RCT, cohrot studies and/or diagnostic accuracy studies

Study ID	Alshehri 2021	
Question		
1. Did the research questions and inclusion criteria for the review include the components of the PICO?	Yes	PICO elements are outlined (p19)
2. Did the report of the review contain an explicit statement that the review methods were established prior to the conduct of the review and did the report justify any significant deviations from the protocol?	Yes	This health technology assessment was registered in PROSPERO, the international prospective register of systematic reviews (CRD42019128547), available at https://www.crd.york.ac.uk/PROSPERO.
3. Did the review authors explain their selection of the study designs for inclusion in the review?	Yes	Justification for the included study types is geven (p19)
4. Did the review authors use a comprehensive literature search strategy?	Yes	The comprehensie strategy used is outlined on p18. Multiple data bases were used with appropriate inclusion criteria
5. Did the review authors perform study selection in duplicate?	No	Only one reviewer screened the studies (p19)
6. Did the review authors perform data extraction in duplicate?	Partial yes	It is suggested that data extraction was performed by more than one reviewer but it isn't explicitly said (p20)
7. Did the review authors provide a list of excluded studies and justify the exclusions?	Partial yes	There is a list of excluded studies but the list is not complete. Justification for exclusion of studies is gien
8. Did the review authors describe the included studies in adequate detail?	Yes	The included studies are described in a good level of depth (p26/p33)
9. Did the review authors use a satisfactory technique for assessing the risk of bias (RoB) in individual studies that were included in the review?	Yes	Yes QUADAS-2 was used to assess the bias of included studies
10. Did the review authors report on the sources of funding for the studies included in the review?	No	No explicit statement around sources of funding
11. If meta-analysis was performed, did the review authors use appropriate methods for statistical combination of results?	No	No meta analysis performed
12. If meta-analysis was performed, did the review authors assess the potential impact of RoB in individual studies on the results of the meta-analysis or other evidence synthesis?	No	No meta analysis performed
13. Did the review authors account for RoB in individual studies when interpreting/ discussing the results of the review?	Yes	Appendix 2 demonstrates the authors sufficintly discussing RoB and how this could've affected results
14. Did the review authors provide a satisfactory explanation for, and discussion of, any heterogeneity observed in the results of the review?	Partial yes	Heterogeneity was investigated through the SROC plots for individual SRs. No discussion about heterogeneity in their own results was discussed
15. If they performed quantitative synthesis did the review authors carry out an adequate investigation of publication bias (small study bias) and discuss its likely impact on the results of the review?	Yes	Publication bias was adequatley investigated and the impact of such was considered (p23/p135)
16. Did the review authors report any potential sources of conflict of interest, including any funding they received for conducting the review?	Partial yes	The impact of conflicts of interest in the individual studies was investigated. However the conflicts of interest that could've occurred in the overarching study was not considered
Overall risk of bias of the review	Moderate	

Source: Shea et al. 2017. BMJ 358: j4008 doi:10.1136 (https://doi.org/10.1136/bmi.j4008)

Study ID	Ontario Health 2020	
Question		
1 . Did the research questions and inclusion criteria for the review include the components of the PICO?	Yes	PICO elements are outlined (p19)
2. Did the report of the review contain an explicit statement that the review methods were established prior to the conduct of the review and did the report justify any significant deviations from the protocol?	No	There is no explicit statement about the review methods
3. Did the review authors explain their selection of the study designs for inclusion in the review?	Yes	Justification for the included study types is geven (p19)
4. Did the review authors use a comprehensive literature search strategy?	Yes	The comprehensie strategy used is outlined on p18. Multiple data bases were used with appropriate inclusion criteria
5. Did the review authors perform study selection in duplicate?	No	Only one reviewer screened the studies (p19)
6. Did the review authors perform data extraction in duplicate?	Partial yes	It is suggested that data extraction was performed by more than one reviewer but it isn't explicitly said (p20)
7. Did the review authors provide a list of excluded studies and justify the exclusions?	Partial yes	There is a list of excluded studies but the list is not complete. Justification for exclusion of studies is gien
8. Did the review authors describe the included studies in adequate detail?	Yes	The included studies are described in a good level of depth (p26/p33)
9. Did the review authors use a satisfactory technique for assessing the risk of bias (RoB) in individual studies that were included in the review?	Yes	Yes QUADAS-2 was used to assess the bias of included studies
10. Did the review authors report on the sources of funding for the studies included in the review?	No	No explicit statement around sources of funding
11. If meta-analysis was performed, did the review authors use appropriate methods for statistical combination of results?	No	No meta analysis performed
12. If meta-analysis was performed, did the review authors assess the potential impact of RoB in individual studies on the results of the meta-analysis or other evidence synthesis?	No	No meta analysis performed
13. Did the review authors account for RoB in individual studies when interpreting/ discussing the results of the review?	Yes	Appendix 2 demonstrates the authors sufficiently discussing RoB and how this could've affected results
14. Did the review authors provide a satisfactory explanation for, and discussion of, any heterogeneity observed in the results of the review?	Partial yes	Heterogeneity was investigated through the SROC plots for individual SRs. No discussion about heterogeneity in their own results was discussed
15. If they performed quantitative synthesis did the review authors carry out an adequate investigation of publication bias (small study bias) and discuss its likely impact on the results of the review?	Yes	Publication bias was adequatley investigated and the impact of such was considered (p23/p135)
16. Did the review authors report any potential sources of conflict of interest, including any funding they received for conducting the review?	Partial yes	The impact of conflicts of interest in the individual studies was investigated. However the conflicts of interest that could've occurred in the overarching study was not considered
Overall risk of bias of the review	Low	

Source: Shea et al. 2017. BMJ 358: j4008 doi:10.1136 (https://doi.org/10.1136/bmi.j4008)

Study ID	Runkel 2020	
Question		
1. Did the research questions and inclusion criteria for the review include the components of the PICO?	Yes	PICO compoenents are outlined (p86)
2. Did the report of the review contain an explicit statement that the review methods were established prior to the conduct of the review and did the report justify any significant deviations from the protocol?	No	No statement that depicts when the review methods were established
3. Did the review authors explain their selection of the study designs for inclusion in the review?	Partial yes	The authors mention study designs that were excluded but have not described the reasoning behind exclusion
4. Did the review authors use a comprehensive literature search strategy?	Yes	Multiple databases were searched with search strategy shown.
5. Did the review authors perform study selection in duplicate?	No	There is no mention as to whether study selection was done in duplication or not
6. Did the review authors perform data extraction in duplicate?	Νο	The authors did not mention if data extraction was done in duplication or not
7. Did the review authors provide a list of excluded studies and justify the exclusions?	Νο	No list of excluded studies was given
8. Did the review authors describe the included studies in adequate detail?	Yes	Table one provides all of the relevant study characteristics (p87)
9. Did the review authors use a satisfactory technique for assessing the risk of bias (RoB) in individual studies that were included in the review?	Yes	Used the STROBE checklist to assess bias in individual studies
10. Did the review authors report on the sources of funding for the studies included in the review?	Yes	Authors declared no external sources of funding (p93)
11. If meta-analysis was performed, did the review authors use appropriate methods for statistical combination of results?	Partial yes	The method of meta-analysis is depicted but there isn't sufficient detail given for compatiblility of studies or why they wanted a single pooled effect
12. If meta-analysis was performed, did the review authors assess the potential impact of RoB in individual studies on the results of the meta-analysis or other evidence synthesis?	Partial yes	The authors described the potential bias in the individual studies to be quite low. This led to no discussion around the impact of individual studies on the meta-analysis
13. Did the review authors account for RoB in individual studies when interpreting/ discussing the results of the review?	Partial	Risk of bias in all included studies was deemed to be low. This created a lack of discussion about potential bias in indvividual studies in the results of the review
14. Did the review authors provide a satisfactory explanation for, and discussion of, any heterogeneity observed in the results of the review?	No	No discussion around potential heterogeneity in results
15. If they performed quantitative synthesis did the review authors carry out an adequate investigation of publication bias (small study bias) and discuss its likely impact on the results of the review?	No	No mention of publication bias
16. Did the review authors report any potential sources of conflict of interest, including any funding they received for conducting the review?	Yes	Authors declared no conflicts of interest (p93)
Overall risk of bias of the review	Low	

Source: Shea et al. 2017. BMJ 358: j4008 doi:10.1136 (<u>https://doi.org/10.1136/bmj.j4008</u>)

Study ID	Yang 2019	
Question		
1. Did the research questions and inclusion criteria for the review include the components of the PICO?	Yes	PICO components listed (p3)
2. Did the report of the review contain an explicit statement that the review methods were established prior to the conduct of the review and did the report justify any significant deviations from the protocol?	No	There was no explicit statement saying that the review methods were established prior to the review
3. Did the review authors explain their selection of the study designs for inclusion in the review?	No	The review authors did not mention the selection of study designs in the inclusion criteria
4. Did the review authors use a comprehensive literature search strategy?	Yes	The search strategy is comprehensive (p2-3, additional file 1)
5. Did the review authors perform study selection in duplicate?	Yes	Two authors involved with a third author used to settle disputes
6. Did the review authors perform data extraction in duplicate?	Partial yes	Data extraction was done by one author, with another author checked by another reviewer. Any disputes between these two authors was resolved by a third author
7. Did the review authors provide a list of excluded studies and justify the exclusions?	No	Reasons for exclusion are listed, but an actual list of excluded studies is not provided
8. Did the review authors describe the included studies in adequate detail?	Yes	PICO elements of included studies provided in table 1 (p5)
9. Did the review authors use a satisfactory technique for assessing the risk of bias (RoB) in individual studies that were included in the review?	Yes	Risk of bias of included studies was checked through the QUADAS-2 checklist
10. Did the review authors report on the sources of funding for the studies included in the review?	Yes	Yes details of funding are listed (p9)
11. If meta-analysis was performed, did the review authors use appropriate methods for statistical combination of results?	Yes	I values listed for combining results to assess heterogeneity. Pooled results from included studies listed (p6)
12. If meta-analysis was performed, did the review authors assess the potential impact of RoB in individual studies on the results of the meta-analysis or other evidence synthesis?	Partial yes	The effect of bias is assessed individually for each study but the effect on the meta analysis is not assessed
13. Did the review authors account for RoB in individual studies when interpreting/ discussing the results of the review?	Yes	The risk of bias is mentioned in the discussion, with certain studies valued more highly due to less bias (p8)
14. Did the review authors provide a satisfactory explanation for, and discussion of, any heterogeneity observed in the results of the review?	Yes	Reasons for the heterogeneity of results is listed (p6)
15. If they performed quantitative synthesis did the review authors carry out an adequate investigation of publication bias (small study bias) and discuss its likely impact on the results of the review?	No	No analysis for small study effects or publication bias was performed because there were too few studies identified to justify (p4)
16. Did the review authors report any potential sources of conflict of interest, including any funding they received for conducting the review?	Yes	Got external funding but declared no conflict of interest (p9)
Overall risk of bias of the review	Low	

Source: Shea et al. 2017. BMJ 358: j4008 doi:10.1136 (https://doi.org/10.1136/bmj.j4008)

Level III- Comparative Observational Studies

Study ID	Jernman 2021		
Domain	Judgement	Description	
Bias due to failure to develop and apply appropriate eligibility criteria	Low risk	Controls are from the same population as the exposed group – being all pregnant women with anti-D antibodies detected in the Finnish Red Cross (FRC) Blood Service between January 1, 2014 and December 31, 2017. There is a low risk for selection bias, with the intervention and outcome clearly defined.	
Bias due to flawed measurement of both exposure and outcome	Low risk	Outcomes are objective (alloimmunisation, HDFN) and comparable across groups, therefore unlikely to be significantly affected by bias. Any potential for flawed measurement of intervention status is carefully accounted for. Any deviations rom intended intervention likley refect usual practise.	
Bias due to failure to adequately control confounding	Serious risk	There are numerous potential confouding variables, some of which are not matched between groups (e.g. age, gravidity, parity). Details on BMI and potential sensitising events are not fully captured. There is also risk of potential time varying confouding (including change in obstetric practise) that are not accounted for. The authors conduct both univariate and multivariate analysis to identify potential risk factors associated with sensitisation, but numbers are low and residual confounding is expected.	
Bias due to incomplete or inadequately short follow-up	Moderate risk	Follow up was long enough to accurately determine the relevant outcomes (e.g. immunization incidence) but it is possible there is missing data that does not truly reflect the incidence of sensitisation prior to the introduction of targeted RAADP (ie the proportion of missing data between groups slightly differs)	
Overall risk of bias	Serious risk	The study appears to provide sound evidence for a non-randomised study but cannot be considered comparable to a well-performed randomised trial. There is potential for some serious residual confounding.	

Source: Table 5.5 GRADE handbook http://gdt.guidelinedevelopment.org/app/handbook/handbook.html#h.m9385o5z3li7

D4 Question 3b

Study ID	Parchure 2021	
Domain	Risk of bias	Applicability
Patient selection	Yes (consecutive selection, case-control design was avoided and the study did not inappropriately exclude participants)	Yes (patients match those targetted by the review)
Index test	Yes (reference standard was interpreted without knowledge of the reference standard, prespecified threshold used)	Yes (No difference in the interpretation or variability of test technology)
Reference standard	Yes (Reference standard is likely to correctly classfiy target audience and the result of the reference standard was not known beforehand)	Yes (the target condition as defined by the reference standard matches the index test)
Patient flow	No (All patients received the same reference standard but not all patients were tested at the same point in time)	

Level II – Consecutive patients with valid reference standard

Source: QUADAS-2 (Whiting et al., 2011)

Level III-1 - Non-consecutive patients with valid reference standard

Study ID	Legler 2021	
Domain	Risk of bias	Applicability
Patient selection	No (case-control design, no innappropriate exclusions, consecutive selection)	Yes (patients are applicable to research question)
Index test	Yes (Knowledge of reference standard was known but not likely to have impacted results, prespecified threshold used)	Yes (no real variation in test technology or difference in interpretation)
Reference standard	Unclear (Reference standard is likely to correctly classify target audience but the result of the reference standard was known prior to the index test)	Yes (the target condition as defined by the reference standard matches the index test)
Patient flow	Unclear (All patients received the same reference standard but not all patients tested were the exact same)	

Source: QUADAS-2 (Whiting et al., 2011)

D5 Question 4

Level III- Retrospective Cohort studies

Study ID	Wikman 2021		
Domain	Judgement		
Bias due to failure to develop and apply appropriate eligibility criteria	Critical	Both cohorts are from different places and from different times. Large disparity between the numbers of each cohort group	
Bias due to flawed measurement of both exposure and outcome	Moderate risk	The measurement of the outcome could have potentially been different in both cohorts but it is unlikely that this introduced significant bias	
Bias due to failure to adequately control confounding	Serious risk	Confounders are accounted for in the intervention group (BMI) but not discussed in the comparative group	
Bias due to incomplete or inadequately short follow-up	Low risk	Both groups were followed up at the same time	
Overall risk of bias	Critical risk	The study is too problematic to provide any useful evidence on the outcome of interest.	

Source: Table 5.5 GRADE handbook http://gdt.guidelinedevelopment.org/app/handbook/handbook.html#h.m9385o5z3li7

Appendix E Data extraction forms

E1 Question 1

Level I – Systematic review (RCTs and cohort studies)

STUDY DETAILS: SR/MA

Citation

Xie 2020

Xie, X., Qiurong, F., Bao, Z., Zhang, Y. & Zhou, D. (2020). Clinical value of different anti-D immunoglobulin strategies for preventing Rh hemolytic disease of the fetus and newborn: A network meta-analysis. *PLoS ONE 15*(3). pp. 1-14.

https://doi.org/10.1371/journal.pone.0230073

Affiliation/Source of funds

Department of Obstetrics and Gynecology, the First People's Hospital of Neijiang, Neijiang, Sichuan

Province, P. R. China (X.X. & D.Z.)., Department of Nursing, The first Affiliated Hospital of Hainan Medical University, Haikou, Hainan Province, P. R. China (Q.F.)., Department of medicine, Southwest Medical University, Luzhou, Sichuan Province, P. R. China (Z.B.). & Department of General Surgery, the First People's Hospital of Neijiang, Neijiang, Sichuan Province, P. R. China (Y.Z.)

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Study design	Level of evidence	Location	Setting	
SR and MA of Level II and III	Level I-III	USA, Canada, Scotland, Holland,	Obstetrics and maternal care	
studies (RCTs and cohoet		England, France, Denmark,		
studies)		Sweden		
Intervention		Comparator		
Various dosage amounts of Rh D immunoglobulin administered		No treatment; or a placebo; or comparisons of different anti-D		
antenatal or postpartum		regimens.		

Population characteristics

Rh negative women with Rh positive fetuses, reported positive incidence of anti-D antibody in postpartum mothers

Length of follow-up	Outcomes measured		
Studies published between 1968-2004	Effectiveness of dose and timing of anti-D immunoglobin in		
	preventing maternal antibody sensitisation		

INTERNAL VALIDITY

Overall risk of bias (descriptive)

Rating: Moderate

Description: Network meta-analysis of low quality studies. The authors do not provide risk of bias assessments or consider the quality of the cohort studies within the analysis. The review may provide an accurate assessment of the available evidence, but the results should be interpreted with caution.

RESULTS:

Outcome	RAADP	No therapy	Risk estimate	Statistical significance
No. patients (No. trials)	n/N (%)	n/N (%)	OR (95% CI)	p-value
N = 64860 (24 studies)				Heterogeneity ^a
				I ² (<i>p</i> -value)
One dose (250 μg within 28 wee	ks' gestation) v placebo/	no treatment		
Incidence of Rh D	NR	NR	0.05 (0.01, 0.18)	Favours intervention
alloimmunisation				p < 0.05
N = 9295 (1 study)				
SUCRA (surface area under the	NR	NR	50.3%	Rank = 5
cumulative ranking curve)				
One dose (300 μg within 28 wee	ks' gestation) v placebo/	no treatment		
Incidence of Rh D	NR	NR	0.01 (0.00, 0.01)	Favours intervention
alloimmunisation				p < 0.05
N = 16 639 (4 studies)				

SUCRA (surface area under the	NR	NR	89.2%	Rank = 2			
cumulative ranking curve)							
Two dose (50 μg within 28 and .	34 weeks' gestation) v pl	acebo/no treatment		1			
Incidence of Rh D	NR	NR	0.15 (0.09, 0.24)	Favours intervention			
alloimmunisation				p < 0.05			
N = 1180 (1 study)							
SUCRA (surface area under the	NR	NR	17.5%	Rank = 8			
cumulative ranking curve)							
Two dose (100 µg between 28 a	nd 34 weeks' gestation)	v placebo/no treatment					
Incidence of Rh D	NR	NR	0.01 (0.01 0.03)	Favours intervention			
alloimmunisation			0.01 (0.01, 0.00)	n < 0.05			
N = 19 684 (4 studies)							
SUCRA (surface area under the	NR	NR	75.1%	Rank = 3			
cumulative ranking curve)							
Two dose (300 µg between 28 g	nd 34 weeks' aestation)	v placebo/no treatment					
Insidence of Ph D	ND			Equation			
alloimmunisation			0.00 (0.00, 0.04)				
N = 2361 (1 study)				p < 0.05			
N = 2501 (1 study)	ND	ND	00.00/	Deals 1			
SUCRA (surface area under the	NK	NK	96.8%	Rank = 1			
	200						
Administered 100 μg ≤ dosage <	200 μg within 72 h post	partum v placebo/no tred	atment				
Incidence of Rh D	NR	NR	NR	NR			
alloimmunisation							
NR (NR)							
SUCRA (surface area under the	NR	NR	40.1%	Rank = 6			
cumulative ranking curve)							
Administered 200 μg ≤ dosage <	< 300 μg within 72 h post	partum v placebo/no treo	atment				
Incidence of Rh D	NR	NR	0.11 (0.04, 0.31)	Favours intervention			
alloimmunisation				p < 0.05			
NR (NR)							
SUCRA (surface area under the	NR	NR	24.1%	Rank = 7			
cumulative ranking curve)							
Administered 300 μg ≤ dosage <	< 500 μg within 72 h post	partum v placebo/no treo	atment				
Incidence of Rh D	NR	NR	0.04 (0.02, 0.06)	Favours intervention			
alloimmunisation				p < 0.05			
NR (NR)							
SUCRA (surface area under the	NR	NR	57.0%	Rank = 4			
cumulative ranking curve)							
EXTERNAL VALIDITY							
Generalisability (relevance of th	e study population to th	e Guidelines target popu	lation)				
The evidence is generalisable to	the target population wit	h few caveats. All studies	were conducted in 'West	tern' countries.			
Applicability (relevance of the e	vidence to the Australian	health care system)					
The suideness is applicable the A	ustralian health care cont	avt with some coverts					
The evidence is applicable the A	ustralian nealth care cont	ext with some caveats.					
Additional comments							
Statistical analysis							
Each closed loop in the network was assessed for inconsistency. Inconsistency factor (IF) was 0.11 = 2.13. 95% CI crossed line of no effect</td							
(contained 0, <i>p</i> > 0.05). Node an	(contained 0, $p > 0.05$). Node analysis showed direct and indirect effect estimates also did not differ.						
Treatments were ranked using SUCRA analysis of cumulative probability of preventing Rh D alloimmunisation.							
Authors conclusions							
In conclusion, this study showed	that the current first-line	recommendation is two	300-μg prenatal immuniz	ations at 28 and 34 gestational			
weeks. If the anti-D immunoglob	oulin supply is inadequate	, the second alternative s	hould be a single 300-µg	prenatal immunization at 28			

gestational weeks.

Included studies

Ascari 1968, Ascari 1969, Bryant 1969, Jennings 1968, Pollack 1968, Robertson 1969, Stenchever 1971, White 1970, Dudok 1968, Clarke 1968, Buchanan 1969, Chown 1969, John 1969, Tovey 1983, Huchet 1987, Bowam 1987, Trolle 1989, Mayne 1997, Mackenzie 1999, Mackenzie 2004, Lee 1995, Bowam 1978, Bowam 1978, Hermann 1984

CI, confidence interval; IU international units; MA, meta-analysis; µg, microgram; RAADP, routine antenatal anti-D prophylaxis; RCT, randomised controlled trial; RR, relative risk; SD, standard deviation; SR, systematic review; UK, United Kingdom

a. Heterogeneity defined as follows: (i) no significant heterogeneity if P_{het} > 0.1 and I² < 25%; (ii) mild heterogeneity if I² < 25%; moderate heterogeneity if I² between 25–50%; substantial heterogeneity I² >50%

Level II- RCT

STUDY DETAILS: RCT

Citation

White 2019

White, SW., Cheng, JC., Penova-Vaselinovic, B., Wang, C., White, M., Ingleby, B., Arnold, C. & Pennell, CE. (2019). Single dose v two-dose antenatal anti-D prophylaxis: a randomised controlled trial. *Medical Journal of Australia*. 221(6). pp.261-265. Doi:10.5694/mja2.50266

Affiliation/Source of funds

Author Affiliations: University of Western Australia, Perth, WA (SWW. & BPV)., King Edward Memorial Hospital for Women, Perth, WA (SWW., BI. & CA)., Royal Perth Hospital, Perth, WA (JCC)., University of Newcastle, Newcastle, NSW (CW., MW. & CEP)., Hunter Medical Research Institute, Newcastle, NSW (CEP).

Sources of Funding: The study was funded in part by a grant to Scott White from the Women and Infants Research Foundation (Perth). Conflicts of Interest: Authors declared no conflicts of interest

Study design	Level of evidence	Location	Setting	
RCT	Level II	King Edward Memorial Hospital,	Obstetrics and maternity care	
		WA, Australia		
Intervention		Comparator		
1500 IU Rh(D) Immunoglobulin-VF at 28 weeks gestation		625 IU Rh(D) Immunoglobulin-VF at 28 and 34 weeks gestation		

Population characteristics

277 women who attended a tertiary obstetric referral hospital in Perth for antenatal care and were at least 18 years of age, less than 30 weeks pregnant and yet to receive RAADP, Rh(D)-negative (negative antibody screen), and who intended to deliver their baby at the hospital. Exclusion criteria were prior anti-D sensitisation, any contraindication of anti-D administration, and a history of isolated IgA deficiency.

Mean age of 30.9 and 31.2 years, 2% to 3% with multiple pregnancy, median BMI of 26.2 and 24.3 and 27% to 31 % had caesarean delivery.

Length of follow-up	Outcomes measured
Between May 2013 and November 2015.	 Detectability anti-D levels in maternal blood at the time of delivery Non-compliance with allocated Rh(D) immunoglobulin prophylaxis regimen

INTERNAL VALIDITY

Overall risk of bias (descriptive)

Rating: Unclear

Description: One domain, relating ot blinding of the participants and researchers has some concerns raised, but none are found to be at high risk of bias. 9% of women in the single dose group were given a second dose, which may bias the results in favour of the single dose.

RESULTS

RESOLIS					
Population analysed	Intervention (one dose)		Comparator (two dose)		
Randomised	140		140		
Efficacy analysis (ITT)	125		129		
Efficacy analysis (PP)	65		75		
Safety analysis	138		139		
Outcome	1500 IU Rh D IgG at 28	625 IU Rh D Ig G at 28 and	Risk estimate (95% CI)	Statistical significance	
	weeks	34 weeks		p-value	
	n/N (%)	n/N (%)			
One-dose (1500 IU at 28 v	weeks) versus two-dose (625	IU at 28 and 34 weeks)			
Proportion with	70/125 (56%)	111/129 (86%)	OR 4.91 (2.67, 9.02)	Favours two-dose	
detectable anti-D at				<i>p</i> < 0.001	
delivery (ITT)	Univariate analyses:	Univariate analyses:			
N = 254	increasing maternal weight [per kg]		OR 0.84 (0.76, 0.93)	<i>p</i> < 0.001	
	interval between final dose and birth [per day]		OR 0.96 (0.95, 0.98)	<i>p</i> < 0.001	
	gestaton at birth [per day]		OR 0.99 (0.99, 1.01)	<i>p</i> = 0.20	

	Multivariate analysis (adjust	ting for maternal weight and	OR 1.55 (0.62, 3.87)	No difference		
	Interval between final dose	and birth)		<i>p</i> = 0.35		
Proportion with	57/65 (88%)	42/75 (56%)	NNR	NR		
detectable anti-D at						
delivery (PP)						
N = 140						
Non compliant (total)	52/138 (38%)	69/139 (50%)	NR	No significant difference		
				<i>p</i> = 0.06		
Safety	No major adverse events ob	oserved.				
	The greater injection volum	e (> 5 mL) for the single dose	group initially made it			
	more painful than for the st	andard regimen; which was al	lleviated by using a more			
	concentrated product, delivering the same dose in a smaller volume (2 mL).					
	Twelve women in the single					
	weeks; they were therefore	5 weeks, consistent with				
	standard practice, to avoid					
EXTERNAL VALIDITY	·					
Generalisability (relevand	ce of the study population to	the Guidelines target popula	tion)			
The evidence is directly ge						
Applicability (relevance o						
The evidence is directly a	oplicable to the Australian hea	alth care system.				
Additional comments						

This is the final published report of the previously included conference abstract (Pennell 2017) that was considered in the 2018 review.

ANZCTR, Australian New Zealand Clinical Trials Registry; CI, confidence interval; IgG, immunoglobulin; ITT, intent to treat; IU, international units; NR, not reported; OR, odds ratio; PP, per-protocol; RCT, randomised controlled trial; WA, Western Australia

E2 Question 2

Level I – Systematic review of observational studies

STUDY DETAILS: SR/MA

Citation

Schmidt-Hansen, 2020

Schmidt-Hansen, M., Lord, J., Hawkins, J., Cameron, S., Pandey, A., Hasler, E. & Regan, F. (2020). Anti-D prophylaxis for rhesus D (RhD)-negative women having an abortion of a pregnancy up to 13⁺⁶ weeks' gestation: a systematic review and new NICE consensus

guidelines. BMJ Sexual Reporductive Health 0(0), 1-6, doi:10.1136/bmjsrh-2019-200536

Affiliation/Source of funds

National Guideline Alliance, Royal College of Obstetricians & Gynaecologists, London UK (MSH, JH, EH)

Department of Obstetrics & Gynaecology, Royal Cornwall Hospitals NHS Trust, Truro, UK (JL)

Sexual and Reproductive Health Services, NHS Lothian, Edinburgh, UK (SC)

Department of Haematology, Imperial College Healthcare NHS Trust and NHS Blood & Transplant, London, UK (FR)

Funding: The study was undertaken by the National Guideline Alliance (NGA) at the Royal College of Obstetricians & Gynaecologists (RCOG), which received funding from the National Institute for Health and Care Excellence (NICE).

The authors declared no conflict of interest

Study design	Level of evidence	Location	Setting	
SR and MA of Level II and Level III	Level I-III	NR	Obstetrics and maternal care	
studies (RCTs and non-				
randomised trials)				
Intervention		Comparator		
Intramuscular anti-D prophylaxis		No anti-D prophylaxis		
(minimum dose of 250 IU/50 μg within 72 hours of medical or				
surgical abortion)				

Population characteristics

Women who are RhD (or D) negative and undergoing either medical abortion with mifepristone and misoprostol or surgical abortion using vacuum aspiration of a pregnancy up to 13⁺⁶ weeks' gestation.

Length of follow-up	Outcomes measured		
Searched Embase, Medline and the Cochrane Library on 19 October	- anti-D isoimmunisation/sensitisation or subsequent affected		
2018.	pregnancy.		
Studies ranged from 1947-2018			

INTERNAL VALIDITY

Overall risk of bias (descriptive)

Rating: Low

Description: No critical weaknesses – the systematic review provides an accurate and comprehensive summary of the results of the available studies that address the question of interest.

RESULTS:						
Outcome	RAADP	No therapy	Risk estimate	Statistical significance		
No. patients (No. trials)	n/N (%)	n/N (%)	RR (95% CI)	p-value		
				Heterogeneity ^a		
				I² (p-value)		
No studies found.						
Recommendation 1						
Offer anti-D prophylaxis to wom	en who are rhesus D nega	itive who are having an al	portion after 10 ⁺⁰ weeks' g	gestation		
Recommendation 2						
Do not offer anti-D prophylaxis to women who are having a medical abortion up to and including 10 ⁺⁰ weeks' gestation						
Recommendation 3						
Consider anti-D prophylaxis for v gestation	vomen who are rhesus D	negative and are having a	surgical abortion up to a	nd including 10 ⁺⁰ weeks'		

Recommendation 4

Providers should ensure that: rhesus status testing and anti-D prophylaxis supply does not cause any delays to women having an abortion **Recommendation 5**

Providers should ensure that anti-D prophylaxis is availableat the time of the abortion

EXTERNAL VALIDITY

Generalisability (relevance of the study population to the Guidelines target population)

The evidence is generalisable to the target population with some caveats.

Applicability (relevance of the evidence to the Australian health care system)

The evidence is applicable the Australian health care context with some caveats.

Additional comments

The systematic review ended up producing 0 studies that were relevant to the inclusion material. Outcomes were to be analysed as risk ratios in Review Manager 5.3 using the Mantel-Haenszel statistical method and a fixed or random effect model. The overall quality of the evidence was planned to be assessed using GRADE.

The results were based off an expert committee that generate the 2019 NICE guidelines on abortion care

CI, confidence interval; IU international units; MA, meta-analysis; µg, microgram; RAADP, routine antenatal anti-D prophylaxis; RCT, randomised controlled trial; RR, relative risk; SD, standard deviation; SR, systematic review; UK, United Kingdom

a. Heterogeneity defined as follows: (i) no significant heterogeneity if P_{het} > 0.1 and I² < 25%; (ii) mild heterogeneity if I² < 25%; moderate heterogeneity if I² between 25–50%; substantial heterogeneity I² >50%

E3 Question 3

Level I – Systematic review (of RCTs, cohort studies and/or diagnostic studies)

STUDY DETAILS: Systematic review of diagnostic studies

Citation

Alshehri 2021

Alshehri, AA. & Jackson, DE. 2021. Non-Invasive Prenatal Fetal Blood Group Genotype and Its Application in the Management of Hemolytic Disease of Fetus and Newborn: Systematic Review and Meta-Analysis. *Transfusion Medicine Reviews* 35(1). 85-94. https://doi.org/10.1016/j.tmrv.2021.02.001

Study design	Lev	el of evidence		Location		Setting	Setting	
Systematic review and	meta- Lev	el I		India, France, N	ance, Netherlands, Great C		Obstetrics and maternity	
analysis of diagnostic s	studies			Britain, Denmark, Spain, Sweden,		,		
				Belgium				
Index test	Ехо	n(s) sequenced		Internal contro	bl(s)	Reference st comparator	tandard or	
High-throughput, NIPT	cell-free 4, 5	, 7, 10 (depends	on the study)	Not specified		Serologic co	rd blood testing	
fetal DNA tests of mate	ernal							
plasma								
Population characteris	stics							
Pregnant Rh negative	women who cou	ld be alloimmun	ised					
Number of studies				Outcomes mea	sured			
16 studies investigatin	g NIPT			Specificity, sens	sitivity			
11 sudies included in t	he meta-analysis	5						
Method of analysis								
Meta-analysis of sensit	tivity and specifi	city was done th	rough DerSimo	nian-Liard rando	m effect model			
INTERNAL VALIDITY								
Overall risk of bias (de	escriptive)							
Rating: Moderate								
Description: More tha	n one non-critic	al weakness – th	e systematic re	view has more t	han one weaknes	s but no critical i	flaws. It may provide	
an accurate summary	of the results of	the available stu	dies that were	included in the r	eview.			
Included studies: All st	udies assessed b	y the authors to	fulfill STROBE	quality standards	s, but details not _l	provided.		
RESULTS								
Outcome	Sensitivity	Specificity	PPV	NPV	LR+	LR-	Diagnostic	
	% (95% CI)	% (95% CI)	% (95% CI)	% (95% CI)	% (95% CI)	% (95% CI)	accuracy	
							% (95% CI)	
Diagnostic performan	ce NIPT against	birth blood sam	ple (inconclusiv	ve as positive)				
N= 31 441	99.3% (98.7,	98.4% (97.4,	NR	NR	NR	NR	NR	
(11 studies)	99.7)	99.0)						
Rather 2019 (India)	99.2%	92.3%	NR	NR	12.88 (NR)	0.0087 (NR)	NR	
	(99.4, 99.9)	(60.9, 98.9)						
Darlington 2018	99.7%	93.2%	NR	NR	14.66 (NR)	0.0032 (NR)	NR	
	(98.1, 100)	(87.7, 96.3)						
Soothill 2015	99.8%	99.2%	NR	NR	124.75 (NR)	0.0020 (NR)	NR	
	(97.5, 100)	(96.1, 99.8)						
Banch-Clausen 2014	99.5%	99.8%	NR	NR	497.5 (NR)	0.0050 (NR)	NR	
	(99.3, 99.6)	(99.6, 99.9)						
Chitty 2014	99.3%	99.1%	NR	NR	110.33 (NR)	0.0071 (NR)	NR	
	(99.0, 99.6)	(98.6, 99.4)						
Grande 2013	99.7%	98.4%	NR	NR	62.31 (NR)	0.0030 (NR)	NR	

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NRR

NR

88.73 (NR)

(92.3, 99.7)

98.9%

(96.0, 100)

97.6%

Wikman 2012

NR

0.0242 (NR)

	(96.9, 98.2)	(98.2, 99.3)					
Akolekar 2011	98.1%	99.7%	NR	NR	327 (NR)	0.0191 (NR)	NR
	(95.9, 99.1)	(95.3, 100)					
Minon 2008	99.9%	99.7%	NR	NR	333 (NR)	0.0010 (NR)	NR
	(97.8, 100)	(95.9, 100)					
Finning 2008	99.7%	98.0%	NR	NR	49.85 (NR)	0.0031 (NR)	NR
	(99.2, 99.9)	(96.7, 98.8)					
Finning 2007	98.1%	99.5%	NR	NR	196.2 (NR)	0.0191 (NR)	NR
	(91.0, 99.6)	(91.8, 100)					
EXTERNAL VALIDITY							
Generalisability (relev	ance of the study	y population to t	he Guidelines t	arget populatior	ı)		
The evidence is genera	lisable to the Au	stralian populatio	on with some ca	veats.			
Studies enrolled Rh D r	negative pregnan	t women but sor	ne may not be d	lirectly applicabl	e in terms of RH	D prevalence.	
Applicability (relevanc	e of the evidenc	e to the Australi	an health care s	ystem)			
The evidence is application	ble to the Austra	lian health care	context with fev	v caveats.			
Includes both high thro	oughput studies (automated) and	those with man	ual DNA extracti	on.		
Additional comments							

Included studies:

Rather 2019; Darlington 2018; Soothill 2015; Banch-Clausen 2014; Grande 2013; Wikman 2012; Akolekar 2011; Minon 2008; Finning 2008; Finning 2007

---. data not reported; cffDNA, cell free fetal DNA; CI, confidence interval; DNA, deoxyribonucleic acid; LR+, positive likelihood ratio; LR-, negative likelihood ratio; NIPT, non-invasive prenatal testing; NPV, negative predictive value; NR, not reported; PPV, positive predictive value; RNA, ribonucleic acid; RT-PCR, real-time polymerase chain reaction.

STUDY DETAILS: Systematic review of diagnostic studies

Citation

Ontario Health 2020

Ontario Health. 2020. Noninvasive fetal RhD blood group genotyping: a health technology assessment. *Ontario Health Technology Assessment Series* [Internet]. 20(15), 1–160. Available from: https://www.hqontario.ca/evidence-to-improve-care/health-technologyassessment/reviews-and-recommendations/noninvasive-fetal-rhd-blood-group-genotyping

Study design	Level of evidence	Location	Setting
Systematic review and meta-	Level I	UK, France, Finland, Cyprus,	Obstetrics and maternity
analysis of diagnostic studies		Netherlands, Denmark, Sweden,	
		Spain, US	
Index test	Exon(s) sequenced	Internal control(s)	Reference standard or
			comparator
cffDNA NIPT testing	4, 5, 7, 10 (depends on the study)	Not specified	Serologic cord blood testing
including laboratory-developed			
tests or commercial test kits			

Population characteristics

Pregnant Rh negative women (who could be alloimmunised) with singleton or multiple pregnancies.

Number of studies	Outcomes measured
6 systematic reviews	Diagnostic accuracy, sensitivity, specificity, PPV, NPV, diagnostic
11 cohort studies	accuracy,
	Unnecessary RhIG avoided; Risk of alloimmunization; Compliance
	with RhIG prophylaxis; Maternal quality of life; Adverse effects such
	as infections from or reactions to RhIG; Implementation outcomes
	such as uptake of testing, uptake of RhIG; Avoidance of cord blood
	RhD testing

Method of analysis

No meta-analysis, the type of analysis differs relevant to the individual study

INTERNAL VALIDITY

Overall risk of bias (descriptive)

Rating: Low

Description: No critical weaknesses – the systematic review provides an accurate and comprehensive summary of the results of the available studies that address the question of interest.

Risk of bias of included systematic reviews assessed using the Risk of Bias in Systematic Reviews (ROBIS) tool. For nonrandomized studies, the risk of bias of each included study using the Risk of Bias Assessment tool for Non-randomized Studies (ROBINS).

Assessments included in GRADE summary of findings

RESULTS

Outcome	Sensitivity	Specificity	PPV	NPV	LR+	LR-	Diagnostic
	% (95% CI)	% (95% CI)	% (95% CI)	% (95% CI)	% (95% CI)	% (95% CI)	accuracy
							% (95% CI)
Diagnostic performance	e of cffDNA NIP	T v blood sample	e at birth				
Mackie 2017	99.3%	98.4%	NR	NR	61 (22,167)	-0.007	NR
N = 10 290 tests	(98.2, 99.7)	(96.4, 99.3)				(0.003, 0.186)	
Zhu 2014	98.5%	97.7%	98.7	98.0	42.83	0.015	98.5% (98.2, 98.7)
N = 10 777 tests	(98.2, 98.7)	(0.87, 1.83)	(98.4, 98.9)	(97.5, 99.0)			
(excludes 352							
inconclusive tests)							
Geifman-Holtzman	95.4%	98.6%	99%	92.1%	17.42 (NR)	0.002 (NR)	94.8%
2006	(90.6, 97.8)	(96.4, 99.5)	(97.9 <i>,</i> 99.6)	(80.9, 97.0)			(NR)
N = 3 078 tests							
(excludes 183 duplicate							
samples, studies with							
N<10, and where							

excluded by primary											
Bivariate meta-	False-positive	rate			Ealso-ne	arativ	ve rate				
analysis	% (95% CI)	late			% (95%		verate		% (95% CI)	% (95% CI)	
	70 (5576 Ci)				/0 (55/0	CI)			70 (3570 Cl)		
Yang 2018											
Inconclusive (8 studies)											
treated as positive	3.86 (2.54–5.8	2)			0.34 (0.	15–0	.76)				
exlcuded	1.26 (0.87–1.8	3)			1.26 (0.	87–1	.83)				
Universal v targeted ar	nti-D										
Outcome	Targeted RAA	OP	No F	RAADP			Risk estim	ate			
	n/N (%)		n/N	(%)			OR (95% C	CI)			
	% (95% CI)		% (9	5% CI)							
Incidence of Rh D	24/9380		86/1	.8 546			RR 0.55 (0	.35, 0.87)	The risk of alloimn	nunization was 45%	
alloimmunisation	0.26% (0.15, 0	.36)	0.46	% (0.37	7 <i>,</i> 0.56)		Absolute F	RD: 0.20	lower in the genot	yping cohort	
(1 study, N=27 926)							NNT 500		compared with the	e historic reference	
Tiblad 2013									antenatal Rh D im	munoglobulin	
									prophylaxis follow	ing any potentially	
									sensitising events.		
Utilisation of Rh D	Pregnancies Carr	ying RhD	All Pr	regnanci	es, % (n/N)	Narrative	summary (re	esults not pooled)		
immunoglobulin	negative Fetus (9	% women									
(8 studies)	immunoglobulin)									
Darlington 2018 (N=850)	479/515 (93%)		90/3	35 (27%))		Across stud	ies, 25.3% to 3	9% of all RhD- pregna	ncies (with an RhD+ or	
Haimila 2017 (N=10 814)	3626/3641 (99.6	%)	3626	/10 814	(33.7%)		RhD– fetus)	avoided unne	cessary RhIG after non	invasive fetal RhD	
Papasavva 2016 (N=71)	18/18 (100%)		18/7	1 (25.3%	6)		blood group) genotyping. A (i.o. pot PhD ii	Among the RhD– pregn neompatible por at risk	ancies carrying an	
Soothill 2015 (N=529)	17/18 (94%)		NR (3	35%)			alloimmuni	zation). over 90	0% avoided unnecessa	rv RhlG.	
Clausen 2014 (N=12 668)	NR (97.3%)		NR (3	37.1%)			Darlington (et al reported 9	93% of not-at-risk RhD-	, – pregnancies avoided	
Tiblad 2013 (N= 27 926)	NR (100%)		3270	/8374 (3	39%)		unnecessar	, y RhIG in the g	enotyping arm, compa	red with only 27% in	
Grande 2013 (N=302)	90/95 (95%)		NR				the control	arm (P value o	r confidence intervals	not provided).	
Damkjaer 2012 (N=239)	68/69 (98.6%)		68/2	16 (31.5	%)		After nonin	vasive fetal Rh	D blood group genotyp	ing in the studies, a	
							small propo request45 o	ortion of people or when test re	e (range: 0.4%—10%) re sults were inconclusive	ceived RhIG upon e.	
EXTERNAL VALIDITY	1		1								
Generalisability (releva	nce of the study	populatio	on to t	he Guio	delines ta	rget	population)			
The evidence is general	isable to the Aus	tralian no	pulatio	on with	some cav	/eats					

Applicability (relevance of the evidence to the Australian health care system)

The evidence is applicable to the Australian health care context.

Additional comments

Included studies

Diagnostic Accuracy: Mackie 2017; Zhu 2014; Geifman-Holtzman 2006; Yang 2019

Clinical Utility: Darlington 2018; Haimila 2017; Papasavva 2016; Soothill 2015; Clausen 2014; Tiblad 2013; Grande 2013; Damkjaer 2012

--. data not reported; cffDNA, cell free fetal DNA; CI, confidence interval; DNA, deoxyribonucleic acid; LR+, positive likelihood ratio; LR-, negative likelihood ratio; NIPT, non-invasive prenatal testing; NPV, negative predictive value; NR, not reported; PPV, positive predictive value; RNA, ribonucleic acid; RT-PCR, realtime polymerase chain reaction.

STUDY DETAILS: Systematic review of RCTs and diagnostic studies

Citation

Runkel 2020

Runkel, B., Bein, G., Sieben, W., Sow, D., Polus, S. & Fleer, D. 2020. Targeted antenatal anti-D prophylaxis for RhD-negative pregnant women: a systematic review. *BMC Pregnancy and Childbirth 20*(83). 1-10. https://doi.org/10.1186/s12884-020-2742-4

Affiliation/Source of funds

Author affiliations: Institute for Quality and Efficiency in Health Care, Cologne Germany, (BR, WS, DS & DF)., Institute for Clinical Immunology and Transfusion Medicine, Justus-Liebig-University, Giessen, Germany, (GB)., Institute for Research in Operative Medicine, Witten/Herdece University, Cologne, Germany (SP).

Study design	Level of evidence	Location	Setting
Systematic review and meta-	Level I	France, UK, Netherlands,	Obstetrics and maternity
analysis of RCTs and diagnostic		Denmark, Finland, Sweden,	
studies		Germany, Spain, Australia,	
		Belgium	
Index test	Exon(s) sequenced	Internal control(s)	Reference standard or
			comparator
NIPT testing with subsequent	4, 5, 7, 10 (depends on the study)	Not specified	Universal anti-D prophylaxis for
administration of anti-D			all non-sensitzed rh D-negative
prophylaxis depending on the			women
result			
Population characteristics		·	
Non-sensitized Rh D negative pre	gnant women		

Number of studies	Outcomes measured
2 RCTs (Rh D prophylaxis)	Sensitivity, specificity
Identified 70 relevant diagnostic accuracy studies - 58 had small	
numbers (between 2 and 467), therefore only 12 included in meta-	
analysis.	

Method of analysis

Meta-analysis was conducted of all the included studies

INTERNAL VALIDITY

Overall risk of bias (descriptive)

Rating: Low

Description: No critical weaknesses – the systematic review provides an accurate and comprehensive summary of the results of the available studies that address the question of interest.

Included studies:

Both off-label studies on anti-D prophylaxis showed a high risk of bias on the study and outcome level.

In 11 of the 12 diagnostic accuracy studies, the risk of bias was high in the total score. However, the pooled estimate of all studies were similar to the results of the study with the low risk of bias

RESULTS

Outcome	RAADP		No therapy		Risk estimate	9	Statistical significance
No. patients (No. trials)	n/N (%)		n/N (%)		OR (95% CI)		p-value
							Heterogeneity ^a
							I ² (p-value)
Incidence of Rh D	NR		NR		Knapp-Hartu	ng method	<i>p</i> = not significant
alloimunisation					OR 0.33 (0, 1	23851)	l ² = 52% (NR)
N = 2297 (2 studies)					Mantel-Haen	szel method	
					OR 0.37 (0.13	8, 1.06)	l ² = 51% (NR)
					Beta-binomia	ıl model	
					OR 0.30 (0.07	7, 1.26)	
Outcome	Sensitivity	Specificity	PPV	NPV	LR+	LR-	Diagnostic accuracy
	% (95% CI)	% (95% CI)	% (95% CI)	% (95% CI)	% (95% CI)	% (95% CI)	% (95% CI)

Diagnostic performance o	of cffDNA NIPT v	v blood sample	at birth				
bivariate meta-analysis	99.9% (99.5;	99.2%	NR	NR	NR	NR	NR
N = 60 011 (12 studies)	100)	(98.5; 99.5)					
De Haas 2016	99.9	97.7	NR	NR	NR	NR	NR
N = 25789	(99.9, 100)	(97.4, 98.0)					
Clausen 2014	99.9	99.1	NR	NR	NR	NR	NR
N = 12668	(99.7, 99.9)	(98.8, 99.4)					
Haimila 2017	100	99.8	NR	NR	NR	NR	NR
N = 10814	(99.9, 100)	(99.6, 99.9)					
Wikman 2012	97.6	98.9	NR	NR	NR	NR	NR
N = 3652	(96.9, 98.2)	(98.2, 99.4)					
Chitty 2014	99.3	99.1	NR	NR	NR	NR	NR
N = 2288	(98.9, 99.6)	(98.5, 99.4)					
Finning 2008	99.7	98.0	NR	NR	NR	NR	NR
N = 1869	(99.2, 99.9)	(96.6, 98.9)					
Muller 2008	99.7	99.2	NR	NR	NR	NR	NR
N = 1022	(98.9, 100)	(97.6, 99.8)					
Macher 2012	100	98.2	NR	NR	NR	NR	NR
N = 2012	(99.4, 100)	(96.4, 99.3)					
Hyland 2017	100	99.6	NR	NR	NR	NR	NR
N = 599	(99, 100)	(97.6, 100)					
Akolekar 2011	98.2	100	NR	NR	NR	NR	NR
N = 586	(96.2, 99.3)	(97.8, 100)					
Minon 2008	100	100	NR	NR	NR	NR	NR
N = 545	(99, 100)	(98, 100)					
Soothill 2015	100	99.4	NR	NR	NR	NR	NR
N = 499	(98.6, 100)	(96.8, 100)					
EXTERNAL VALIDITY							

Generalisability (relevance of the study population to the Guidelines target population)

The evidence is generalisable to the Australian population with some caveats.

Applicability (relevance of the evidence to the Australian health care system)

The evidence is applicable to the Australian health care context.

Additional comments

Evidence is to inform the German Institute for Quality and Efficiency in Health Care (IQWiG).

The current policy of universal antenatal anti-D administration leads to approximately 50,000 RhD negative pregnant women per year in Germany receiving anti-D prophylaxis even though they are carrying an RhD negative fetus.

Included studies:

Effectiveness: Hutchet 1987; Lee 1995

Diagnostic accuracy: De Haas 2016; Clausen 2014; Haimila 2017; Wikman 2012; Chitty 2014; Finning 2008; Muller 2008; Macher 2012; Hyland 2017; Akolekar 2011; Minon 2008; Soothill 2015

--- data not reported; cffDNA, cell free fetal DNA; CI, confidence interval; DNA, deoxyribonucleic acid; LR+, positive likelihood ratio; LR-, negative likelihood ratio; NIPT, non-invasive prenatal testing; NPV, negative predictive value; NR, not reported; PPV, positive predictive value; RNA, ribonucleic acid; RT-PCR, realtime polymerase chain reaction.

STUDY DETAILS: Systematic review of diagnostic studies

Citation

Yang, 2019

Yang, H., Llewellyn, A., Walker, R., Harden, M., Saramago, P., Griffin, S. & Simmonds, M. (2019). High-throughput, non-invasive prenatal testing for fetal rhesus D status in RhD negative women: a systematic review and meta-analysis. *BMC Medicine* 17(37). pp. 1-10. https://doi.org/10.1186/s12916-019-1254-4

11(195.)/ 001.016/ 10.1100/ 5	12310 013 1	234 4					
Study design	Lev	el of evidence		Location		Setting	
Systematic review and me	ta- Lev	el I		London, Denma	rk, Bristol, Spain,	Obstetrics a	nd maternity
analysis of diagnostic stud	ies			Netherlands, Sv	veden		
Index test	Ехо	n(s) sequenced		Internal contro	l(s)	Reference s comparator	tandard or
High-throughput, NIPT cel	l-free 4, 5	, 7, 10 (depends or	n the study)	Not specified		Serologic co	rd blood testing
fetal DNA tests of materna	al						
plasma							
Population characteristics	5						
Pregnant Rh negative wor	nen who cou	ld be alloimmunise	ed				
Number of studies				Outcomes meas	sured		
Diagnostic accuracy: 8 stu	dies included	l in the review. Cor	nbined	Diagnostic accu	racy, sensitivity, s	pecificity, PPV,	NPV, accuracy at
sample of 42491.				gestational age			
Method of analysis							
Meta-analysis of all eight s	studies to de	termine overall fals	se positive an	d false negative	rates.		
INTERNAL VALIDITY							
Overall risk of bias (descri	iptive)						
Rating : Low							
Description: No critical fla	ws. The syste	ematic review prov	ides an accur	ate and compreh	ensive summary	of the results of	f the available
studies that address the q	uestion of in	terest.					
RESULTS							
Outcome S	Sensitivity	Specificity	PPV	NPV	LR+	LR-	Diagnostic
9	% (95% CI)	% (95% CI)	% (95% CI)	% (95% CI)	% (95% CI)	% (95% CI)	accuracy
	. ,		. ,			. ,	% (95% CI)

Diagnostic performance NIPT against birth blood sample (inconclusive as positive)								
N = NR	99.66%	96.14% (2.54-	NR	NR	25.82 (NR)	0.004 (NR)	NR	
	(0.15-0.76)	5.82)						
Diagnostic performance NIPT against birth blood sample (excluding inconclusive)								
N = NR	99.65%	98.74% (0.87-	NR	NR	79.09 (NR)	0.004 (NR)	NR	
	(0.15-0.82)	1.83)						
Diagnostic performance	e NIPT against b	irth blood samp	le (only including	g Bristol populat	tion)			
N = NR	99.79%	94.27% (4.58-	NR	NR	17.42 (NR)	0.002 (NR)	NR	
	(0.09-0.48)	7.16)						
EXTERNAL VALIDITY				•		•		
Generalisability (relevance of the study population to the Guidelines target population)								
Generalisability (releva	nce of the study	population to t	he Guidelines ta	rget population)			

Studies enrolled Rh D negative pregnant women but some may not be directly applicable in terms of *RHD* prevalence

Applicability (relevance of the evidence to the Australian health care system)

The evidence is probably applicable to the Australian health care system with some caveats.

Only high throughput studies were included. This may overestimate the sensitivity of the test.

Additional comments

Duplicate Data (this is published report of data included in our original 2018 search - see Saramago 2018

Included studies

Akolekar 2011; Banch-Clausen, 2014; Chitty 2014; Finning 2008; Grande 2013; Soothill 2015; Thurik 2015; Wikman 2012

--. data not reported; cffDNA, cell free fetal DNA; CI, confidence interval; DNA, deoxyribonucleic acid; LR+, positive likelihood ratio; LR-, negative likelihood ratio; NIPT, non-invasive prenatal testing; NPV, negative predictive value; NR, not reported; PPV, positive predictive value; RNA, ribonucleic acid; RT-PCR, realtime polymerase chain reaction.

Level III- Comparative Observational Studies

STUDY DETAILS: Cohort / Case-control

Citation

Jernman 2021

Jernman, R., Isaksson, C., Haimila, K., Kuosmanen, M., Makikallio-Anttila, K., Toivonen, S., Orden, MR., Sulin, K., Tihtonen, K., Vaarasmaki. & Sainio, S. (2021). Time points and risk factors for RhD immunizations after the implementation of targeted routine antenatal anti-D prophylaxis: A retrospective nationwide cohort study. *Acta Obstetricia et Gynecologica Scandinavica*. *100*(10). pp. 1868-1875. doi: 10.1111/aogs.14216

Affiliation/Source of funds

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The authors declared no conflict of interest

Author Affiliations: Department of Obstetrics and Gynecology, University of Helsinki and Helsinki University Hospital, Helsinki, Finland (RJ. & CI)., Finnish Red Cross Blood Service, Helsinki, Finland (KI., MK., ST., KS. & SS)., Department of Obstetrics and Gynecology, Turku University Hospital, Turku, Finland (KM)., Department of Obstetrics and Gynecology, Kuopio University Hospital, Kuopio, Finland (MR)., Department of Obstetrics and Gynecology, Colu University Hospital, Tampere, Finland (KT)., Department of Obstetrics and Gynecology, Oulu University Hospital, Oulu, Finland (MV).

Study design	Level of evidence	Location		Setting	
Retrospective cohort study	spective cohort study Level III-3 Finland Obstetrics and maternit				
Intervention		Comparator			
National screening program of Fin prophylaxis <i>Risk-based prophylaxis:</i> 250-300 to the event of a sensitising event (s weeks, all terminations of pregna chorionic villous sampling, amnio antenatal haemorrhage, external <i>Targeted:</i> 250-300 mcg anti-D im mothers with an RHD-positive fet unknown at 28-30 weeks of gesta <i>Postnatal:</i> 250-300 mcg given wit negative mothers with an RhD-po status of the newborn	nland routine antenatal anti-D ncg anti-D immunoglobin given in pontaneous abortions after 8 ncy, extrauterine pregnancies, centesis, abdominal trauma, version, intrauterine death munoglobin given to RhD-negative us or if the fetal RhD status is tion. hin 72 hours of delivery to RhD- ositive newborn or unclear RhD	Pre-introduction screening), get the same perion Welfare witho	on of routine anti-D i neral population of p od (obtained from the ut matching for parit	mmunoglobin (no routine anti-D regnant women in Finland during e Finnish Institute for Health and y).	
Population characteristics					
RhD negative pregnant women v	vith detected anti-D antibodies who g	ave birth in Finla	and (ave age of 27.3)	, median BMI of 24.5	
Length of follow-up		Outcomes me	asured		
Between 2014-2017		Incidence of a	nti-D immunization		
Between 2014-2017 Method of analysis		Incidence of a	nti-D immunization		
Between 2014-2017 Method of analysis A nationwide cohort study was co Service between January 1, 2014 The data were analysed using SPS and 95% CI was used. The number exact test and Student's t test, de 2014 and between time-points of	onducted of all pregnant women with and December 31, 2017. S Version 25.0 (IBM Corporation, Arm r of observations in the study group a pending on the variable. Logistic regro	Incidence of an anti-D antibodio onk, NY, USA). nd controls was ession was used	es detected in the Fir A p-value <0.05 was compared using Pea to sort out risk facto	nish Red Cross (FRC) Blood considered statistically significant, rson's chi-squared test, Fisher's or proportion before and after	
Between 2014-2017 Method of analysis A nationwide cohort study was co Service between January 1, 2014 The data were analysed using SPS and 95% CI was used. The number exact test and Student's t test, de 2014 and between time-points of INTERNAL VALIDITY	onducted of all pregnant women with and December 31, 2017. S Version 25.0 (IBM Corporation, Arm r of observations in the study group a pending on the variable. Logistic regre immunization	Incidence of an anti-D antibodio onk, NY, USA). nd controls was ession was used	nti-D immunization es detected in the Fir A p-value <0.05 was s compared using Pea I to sort out risk facto	unish Red Cross (FRC) Blood considered statistically significant, rson's chi-squared test, Fisher's or proportion before and after	
Between 2014-2017 Method of analysis A nationwide cohort study was co Service between January 1, 2014 The data were analysed using SPS and 95% CI was used. The number exact test and Student's t test, de 2014 and between time-points of INTERNAL VALIDITY Overall risk of bias (descriptive)	onducted of all pregnant women with and December 31, 2017. S Version 25.0 (IBM Corporation, Arm r of observations in the study group a pending on the variable. Logistic regro	Incidence of an anti-D antibodio onk, NY, USA). nd controls was ession was used	nti-D immunization es detected in the Fir A p-value <0.05 was s compared using Pea I to sort out risk facto	nish Red Cross (FRC) Blood considered statistically significant, rson's chi-squared test, Fisher's or proportion before and after	
Between 2014-2017 Method of analysis A nationwide cohort study was co Service between January 1, 2014 The data were analysed using SPS and 95% CI was used. The number exact test and Student's t test, de 2014 and between time-points of INTERNAL VALIDITY Overall risk of bias (descriptive) Rating: Serious Description: The study appears to performed randomised trial. The	onducted of all pregnant women with and December 31, 2017. S Version 25.0 (IBM Corporation, Arm r of observations in the study group a pending on the variable. Logistic regre immunization	Incidence of an anti-D antibodia onk, NY, USA). nd controls was ession was used ndomised study al confounding.	es detected in the Fir A p-value <0.05 was s compared using Pea I to sort out risk facto	unish Red Cross (FRC) Blood considered statistically significant, rson's chi-squared test, Fisher's or proportion before and after dered comparable to a well-	
Between 2014-2017 Method of analysis A nationwide cohort study was co Service between January 1, 2014 The data were analysed using SPS and 95% CI was used. The number exact test and Student's t test, de 2014 and between time-points of INTERNAL VALIDITY Overall risk of bias (descriptive) Rating: Serious Description: The study appears to performed randomised trial. The RESULTS	onducted of all pregnant women with and December 31, 2017. S Version 25.0 (IBM Corporation, Arm r of observations in the study group a pending on the variable. Logistic regre immunization	Incidence of an anti-D antibodio onk, NY, USA). nd controls was ession was used ndomised study al confounding.	es detected in the Fir A p-value <0.05 was compared using Pea to sort out risk facto	nish Red Cross (FRC) Blood considered statistically significant, rson's chi-squared test, Fisher's or proportion before and after dered comparable to a well-	
Between 2014-2017 Method of analysis A nationwide cohort study was co Service between January 1, 2014 The data were analysed using SPS and 95% CI was used. The number exact test and Student's t test, de 2014 and between time-points of INTERNAL VALIDITY Overall risk of bias (descriptive) Rating: Serious Description: The study appears to performed randomised trial. The RESULTS Population analysed	onducted of all pregnant women with and December 31, 2017. S Version 25.0 (IBM Corporation, Arm r of observations in the study group a pending on the variable. Logistic regre immunization o provide sound evidence for a non-ran re is potential for some serious residua Intervention	Incidence of an anti-D antibodia onk, NY, USA). nd controls was ession was used ndomised study al confounding.	es detected in the Fir A p-value <0.05 was a s compared using Pea to sort out risk facto y but cannot be consid	unish Red Cross (FRC) Blood considered statistically significant, rson's chi-squared test, Fisher's or proportion before and after dered comparable to a well-	

Outcome	Targeted RAADP	No RAADP	Risk estimate (95% CI)	Statistical significance		
	n/N	n/N		p-value		
	% (95% CI)	% (95% CI)				
Targeted RAADP vs no RAADP						
Prevalence of anti-D	54	174	NR	Favours intervention		
sensitisation among pregnant	0.88% (0.68%, 1.14%)	1.52% (1.26%, 1.84%)		<i>p</i> = 0.0009		
women (274 pregnancies of						
228 women)						
- Screening at 8-10 weeks	10/54 (18.5%)	NR (52%)				
 Screening at 24-26 weeks 	27/54 (50%)	NR (20%)				
- Screening at 36 weeks	17/54 (28%)	NR (28%)				
Incidence of anti-D	0.10% (0.05%, 0.22%)	0.33% (0.22%, 0.48%)	RR 0.29 (0.10, 0.71)	Favours intervention		
sensitisation among pregnant			[new sensitisations]	<i>p</i> = 0.0037		
women (NR pregnancies of 197			Absolute RD 0.20%			
women)	Univariate analysis sugge	sted the following risk fact	tors for sensitisation:			
	PPH ≥ 1000 mL, RBC tran	sfusion in previous pregna	ncy, twins in ongoing			
	pregnancy.					
	id not reach statistical					
	(low numbers may preve					
EXTERNAL VALIDITY						

Generalisability (relevance of the study population to the Guidelines target population)

The evidence is generalisable to the Australian population with some caveats.

Applicability (relevance of the evidence to the Australian health care system)

The evidence is applicable to the Australian health care context with some caveats.

Additional comments

*There were significant baseline differences between the intervention and comparator groups in relation to mean age (27.36 vs 30.7); gravidity (G1: 18.2% vs 29.6%); parity (P0: 25.5% vs 41.3%); and delivery complications (assisted delivery, transfusion, bleeding ≥1000mL, postmaturity ≥41 weeks).

*There is insufficient information on the incidence of potential sensitising events.

*It is noted that none of the sensitising events were attributed to false-negative fetal RHD typing.

CI, confidence interval; HDFN, haemolytic disease of the fetus and newborn; IgG, immunoglobulin; ITT, intention to treat; IU, international units; IUT, intrauterine transfusion; NNT, number needed to treat; PP, per-protocol; RAADP, routine antenatal anti-D prophylaxis; RCT, randomised controlled trial; RR, relative risk; SD, standard deviation

E4 Question 3b

Level II – Consecutive patients with valid reference standard

STUDY DETAILS: Diagnostic study

Citation

Parchure 2021

Parchure, D., Madkaikar. & Kulkarni, S. 2021. Algorithm development and diagnostic accuracy testing for non-invasive foetal RHD genotyping: an Indian experience. *Blood Transfusion*. 1-11. doi: 10.2450/2021.0022-21

Affiliation/Source of funds

Author Affliations: Department of Transfusion Medicine, ICMR-National Institute of Immunohaematology, Mumbai, India (DP, SK)., Department of Pediatric Immunology and Leukocyte Biology, ICMR-National Institute of Immunohaematology, Mumbai, India (MM). Funding was sort through an intramural grant received from the Indian Council of Medical Research.

The authors declared no sources of conflict

Study design		Level of evidence Location and study date			Setting		
Prospective obser	rvational study	Level II		Mumbai, India		Obstetrics and maternity	
Index test		Exon(s) sequen	ced	Internal control(s)		Reference standard or comparator	
PCR method in th cffDNA from mate various weeks of 34)	e extraction of ernal plasma in gestation (10-	RHD exons 4, 5 samples) RHD exons 5 an samples)	and 10 (initial 54 d 10 (163	CCR5, SRY and I were used as co	RASSF1A genes ontrols	Cord blood serol	ogy at delivery
Population chara	cteristics						
RhD negative prea	gnant Indian wor	nen aged betwee	n 19-42 with a me	an age of 32.5			
Number of studie	es or samples			Outcomes meas	ured		
217				Sensitivity, speci	ficity, diagnostic ac	curacy, alloimmur	nisation
Method of analys	sis						
Specificity, sensiti	ivity and diagnos	tic accuracy value	s of the diagnostic	methods were ca	alculated		
INTERNAL VALI	DITY						
Overall risk of bia	as (descriptive)						
RESULTS							
2x2 table with inc	conclusive result	s counted as test	positive ^a				
N = 217		Reference stand	lard positive	Reference stand	dard negative	Inconclusive res	ults
		n = 175 (86.21%)	n = 28 (13.79%)		n = 14	
Index text positiv n = 175 (86.21%)	/e	175		0		NR	
Index text negation n = 28 (13.79%)	ve	0		28		NR	
Index test inconc	lusive						
Outcome	Sensitivity % (95% Cl)	Specificity % (95% Cl)	PPV % (95% CI)	NPV % (95% CI)	LR+ (95% CI)	LR- (95% CI)	Diagnostic accuracy % (95% CI)
Diagnostic perfor	mance NIPT aga	inst birth blood s	ample ^d		•	•	
	100% (NR)	100%(NR)	NR	NR	NR	NR	100%
EXTERNAL VALI	DITY				•		
Generalisability (relevance of the study population to the Guidelines target population)							
The evidence is generalisable to the Australian population with some caveats							
Applicability (relevance of the evidence to the Australian health care system)							
The evidence is not applicable to the Australian health care system							

Additional comments

cffDNA, cell free fetal DNA; CI, confidence interval; DNA, deoxyribonucleic acid; GW, gestational week; LR+, positive likelihood ratio; LR-, negative likelihood ratio; NIPT, non-invasive prenatal testing; NPV, negative predictive value; NR, not reported; PPV, positive predictive value; RT-PCR, real-time polymerase chain reaction.

Level III-1 - Non-consecutive patients with a valid reference standard

STUDY DETAILS: Diagnostic study

Citation

Legler 2021

Legler, TJ., Luhrig, S., Korschineck, I. & Schwartz, D. (2021). Diagnostic performance of the noninvasive prenatal FetoGnost RhD assay for the prediction of the fetal RhD blood group status. *Archives of Gynecology and Obstetrics (304)*1. pp. 1191-1196. https://doi.org/10.1007/s00404-021-06055-1

Affiliation/Source of funds

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Author affiliations: Department of Transfusion Medicine, University Medical Center Göttingen, Robert Koch Str. 40, 37075 Göttingen, Germany (TJL & SL)., Ingenetix GmbH, Vienna, Austria (IK)., Department of Blood Group Serology and Transfusion Medicine, Medical University of Vienna, Vienna, Austria (DS).

Sources of conflict: T.L. receives consultation fees from LADR GmbH and participates in the revenue of his employer. I.K. is the owner and manager of the company Ingenetix GmbH. S.L. and D.S. do not have any conflicts of interest/competing interests to declare

Study design	Level of evidence	Location and study date	Setting
Retrospective observational	Level III-1	Level III-1 Vienna Medical University	
study		Obstetrics department. Between	
		2009-2020	
Index test	Exon(s) sequenced	Internal control(s)	Reference standard or
			comparator
FetoGnost RhD assay	RHD exon 5, exon 7	NR	Cord blood serology
Population characteristics			
Pregnant women aged between	16-50		
Number of studies or samples		Outcomes measured	
2968 pregnant women Sensitivity specificity PPV NPV diagnostic accuracy			

Method of analysis

Samples of EDTA blood of RhD negative women were received in the genetics laboratory within a maximum of 6 h of venipuncture. Plasma was separated by centrifugation at 3000 rpm/10 min and stored frozen at - 20C until the insulation. Free-floating DNA from the plasma was isolated from Macherey–Nagel commercial NucleoSpin Plasma kit according to the manufacturer's instructions. In parallel with the isolation of plasma sample in duplicate, and was isolated by the same amount of RNAse free water as a negative control monitored the entire procedure. CffDNA is eluted with 30 II of the elution buffer.

Statistically evaluated by reviewing NIPT-RhD results from the FetoGnost RhD assay with the reference standard of RhD blood group serology results from newborns from the Medical University of Vienna in a retrospective analysis

INTERNAL VALIDITY

Overall risk of bias (descriptive)

Rating: Some concerns

Description:

RESULTS

2x2 table with inconclusive results counted as test positive^a

N = 2968		Reference standard positive		Reference star	Reference standard negative		Inconclusive results	
		n = 1475 (63.71%)		n = 769 (33.59	n = 769 (33.59%)		n = 644	
Index text positiv	ve							
n = 1891 (65.48%)		1474		3	3		414	
Index text negati	ve	1		766	766		220	
n = 997 (34.52%)		1		700	700		230	
Index test inconc	lusive			NP		ND	NR	
n = 80 (2.70%)		NN						
Outcome	Sensitivity	Specificity	PPV	NPV	LR+	LR-	Diagnostic	
	% (95% CI)	% (95% CI)	% (95% CI)	% (95% CI)	(95% CI)	(95% CI)	accuracy	
							% (95% CI)	

Diagnostic performance NIPT against birth blood sample ^d							
	99.93%	99.61%	99.80 (NR)	99.87 (NR)	256.16 (NR)	0.0007 (NR)	99.82%
	(99.61, 99.99)	(98.86, 99.87)					(99.54, 99.93)
EXTERNAL VAL	DITY						
Generalisability (relevance of the study population to the Guidelines target population)							
The evidence is generalisable to the Australian population with some caveats							
Applicability (relevance of the evidence to the Australian health care system)							
The evidence is not applicable to the Australian health care system							
Additional comments							

cffDNA, cell free fetal DNA; CI, confidence interval; DNA, deoxyribonucleic acid; GW, gestational week; LR+, positive likelihood ratio; LR-, negative likelihood ratio; NIPT, non-invasive prenatal testing; NPV, negative predictive value; NR, not reported; PPV, positive predictive value; RT-PCR, real-time polymerase chain reaction.

E5 Question 4

Level III- Retrospective cohort studies

STUDY DETAILS: Case-control

Citation

Wikman, 2021

Wikman, A., Mortberg, A., Jalkesten, E., Jansson, Y., Karlsson, A., Tiblad, E. & Ajne, G. 2021. Altered strategy of prophylactic anti-D administration in pregnancy to cover term and post-term – a pilot study. *The international journal of transfusion medicine 116*(1) 1005-1011. https://doi.org/10.1111/vox.13092

Affiliation/Source of funds

Author Affiliations: Department of Clinical Immunology and Transfusion Medicine, Karolinska University Hospital, Stockholm, Sweden (AW, AM, EJ & AK)., Division of Immunology, Department of CLINTEC, Karolinska Institute, Stockholm, Sweden (AW & AM)., Pregnancy Care & Delivery, Karolinska University Hospital, Stockholm, Sweden (YJ & GA)., Center for Fetal Medicine, Department of Obstetrics and Gynecology, Karolinska University Hospital, Stockholm, Sweden (ET)., Clinical Epidemiology Division, Department of Medicine Solna, Karolinska Institute, Stockholm, Sweden (ET)., Division of Obstet & Gynecol, Department of CLINTEC, Karolinska Institute, Stockholm, Sweden (GA).

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Study design	Level of evide	nce	Location		Setting	
Case-control	Level III		Sweden, Gerr	nany	Maternity and obstetrics	
Intervention			Comparator			
RAADP of 1500 IU of anti-D given at GA 28 and another 1500 IU dose			RAADP with 1	.250 IU of anti-D give	n at GA 28-29	
given at GA 38						
Population characteristic	cs					
RhD negative women with a	a RhD positive fetus					
Length of follow-up			Outcomes n	neasured		
Retrospective cohort was co	ollected between Octob	er 2010 and	- Effect of	3MI on anti-D IgG det	tection in week 38	
October 2012 in Sweden			- Detection	of anti-D prophylaxi	s at delivery	
The prospective cohort was	collected between 2016	5 and 2018 in				
Germany						
Method of analysis						
Linear regression analysis w	vas conducted to show th	he effect of BMI or	n anti-D detecti	on		
INTERNAL VALIDITY						
Overall risk of bias (descrip	tive)					
Rating: Moderate						
Description: The study appe	ears to provide sound ev	idence for a non-ra	andomised stud	ly but cannot be con	sidered comparable to a well-	
performed randomised tria	I. Of key concern is an ov	ver-representation	of women from	n the primary setting	g (midwives, GPs) vs obstetric	
setting (3:1) in the controls	compared with cases. W	eighted data were	e used in the ar	alysis.		
RESULTS						
Population analysed	Cases			Controls		
Available	39			4280		
Analysed	39			4280		
Outcome	Cases	Controls		Risk estimate (95%	Statistical significance	
	n/N (%)	n/N (%)		CI)	p-value	
Linear Regression Analysis						
Detectability of anti-D at	7/39 (18%)	856/4280 (2	0.5%)	NR	NR	
delivery						
Incidence of FMH (>1mL)	None detected					
at delivery	0/25 (0%)					
BMI	23.9 (18.8, 34.8)	NR		NR	NR	

median (min, max)	Linear regression analysis				
	showed a significant				
	correlation to body mass				
	index (<i>p</i> = 0.0118)				
EXTERNAL VALIDITY					
Generalisability (relevance of the study population to the Guidelines target population)					
The results are somewhat generalisable to the Australian population					
Applicability (relevance of the evidence to the Australian health care system)					

The results are the study are applicable to the Australian context with some caveats

Additional comments

BMI, body mass index; CI, confidence interval; GP, general practitioner; im, intramuscular; IU, international units; OR, odds ratio; RAADP, routine antenatal anti-D prophylaxis; RBC, red blood cell; RCT, randomised controlled trial; SD, standard deviation

a. By design, the controls under primary care were overrepresented (with lower prevalence of potential risk factors for example previous medical intervention), which could overestimate the effect of potential risk factors. The authors therefore weighted the primary care controls (0.35) to restore the proportion of primary care pregnancies to the control group. All p-values are based on n=146.

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