Laboratory Monitoring of Mother, Fetus, and Newborn in Hemolytic Disease of Fetus and Newborn

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Abstract

Background: Laboratory monitoring of mother, fetus, and newborn in hemolytic disease of fetus and newborn (HDFN) aims to guide clinicians and the immunized women to focus on the most serious problems of alloimmunization and thus minimize the consequences of HDFN in general and of anti-D in particular. Here, we present the current approach of laboratory screening and testing for prevention and monitoring of HDFN at the Copenhagen University Hospital in Denmark.

Summary: All pregnant women are typed and screened in the 1st trimester. This serves to identify the RhD-negative pregnant women who at gestational age (GA) of 25 weeks are offered a second screen test and a non-invasive fetal RhD prediction. At GA 29 weeks, and again after delivery, non-immunized RhD-negative women carrying an RhD-positive fetus are offered Rh immunoglobulin. If the 1st trimester screen reveals an alloantibody, antenatal investigation is initiated. This also includes RhD-positive women with alloantibodies. Specificity and titer are determined, the fetal phenotype is predicted by non-invasive genotyping based on cell-free DNA (RhD, K, Rhc, RhC, RhE, ABO), and serial monitoring of titer commences. Based on titers and specificity, monitoring with serial peak systolic velocity measurements in the fetal middle cerebral artery to detect anemia will take place. Intrauterine transfusion is given when fetal anemia is suspected. Monitoring of the newborn by titer and survival of fetal red blood cells by flow cytometry will help predict the length of the recovery of the newborn.

Introduction

Alloimmunization is the process where an individual lacking a specific antigen of a blood group is exposed to the antigen and responds by producing specific antibodies. Exposure might occur by transfusion with donor blood, or by accidental transfer of fetal red blood cells (RBCs) to the pregnant woman, such as fetomaternal hemorrhage [1–3].

Active transplacental transfer of maternal antibodies via the neonatal Fc receptor [4] will take place when the antibody production has switched from the initial IgM response to IgG. Transfer will accelerate in the 2nd and
3rd trimester and can lead to hemolytic disease of fetus and newborn (HDFN) [5].

The essential clinical manifestation of HDFN is fetal and neonatal anemia. This is observed as erythroblastosis fetalis, hepatic dysfunction leading to hypoalbuminemia, ascites, hydrops fetalis, congestive heart failure, intravenous growth retardation, abdominal and pericardial edema, antenatal asphyxia, acute bilirubin encephalopathy, and kernicterus spectrum disorder [6]. The concentration of unconjugated bilirubin might surpass albumin bilirubin binding capacity and translocate across the brain-blood barrier with subsequent accumulation in basal ganglia resulting in neuronal-cell death. Bilirubin encephalopathy, or kernicterus, may lead to minor neurodevelopmental disabilities, nerve deafness, spastic cerebral palsy, or even death [7].

Laboratory monitoring is an important tool to predict a potential risk; but it cannot with certainty forecast clinical severity for the fetus. However, crucially, the laboratory setup removes the urgency of the diagnosis of HDFN and allows timely implementation of tested diagnostic and therapeutic measures. Cases of unpredicted HDFN still occur and they are consequently not included in the antenatal fetal screening program [2, 3, 8].

It is important at an early gestational age (GA) to determine if the woman is RhD negative and thus is at risk of producing the most frequent alloantibody, anti-D, and whether or not she is amenable to preventive treatment, Rh prophylaxis, later in pregnancy. At the same time all women are screened for the presence of any alloantibody.

In Denmark, if the woman is typed RhD negative and has no alloantibodies in the 1st trimester antibody screen, she will be offered a non-invasive prediction of the fetal RhD blood group at GA 25 weeks at routine consultation with her general practitioner. At the same time, an antibody screen test is performed. If the fetus is RhD positive and the women is non-immunized, anti-D immunoglobulin (RhIg) is offered at GA 29 weeks and again after delivery. This is described below (see Antenatal RHD Screening).

In contrast, if a potentially harmful maternal antibody is detected in the screen in the 1st trimester, it is essential to determine if the present fetus carries the allele for the antigen targeted by the maternal antibody. Only in this case is the fetus at risk of developing HDFN. In cases where the fetus is at risk, intensified pregnancy monitoring and treatment by transfusion can be instituted, and in cases where risk of HDFN due to known alloantibodies can be excluded, a less intensive and financially less burdensome approach can be taken. Also, much anxiety from the prospective parents can be avoided. If the woman is alloimmunized to the RhD antigen when tested in the 1st trimester in the antibody screen, we immediately use the same antenatal RHD screening assay as we use for non-immunized women in GA 25 weeks. Detection of the RHD gene is based on selective amplification of fetal DNA encoding the RHD gene.

Selective amplification is, however, not reliably achievable for other blood group polymorphisms [9]. We examine women alloimmunized to the other prevalent antigens (K, RhC, Rhc, RhE, and ABO) by non-invasive antenatal molecular diagnostics that amplify single nucleotide variants (SNVs) potentially present in the cell-free DNA (cfDNA), maternal as well as fetal. We describe our clinically implemented non-invasive methods based on cfDNA for 1st/2nd trimester determination of the genes encoding the clinically most important targets of alloantibodies (see Non-Invasive Prediction of Fetal K, RhC, Rhc, RhE, and ABO Blood Group).

Prediction of other phenotypes based on antenatal genotyping has not yet been implemented in our laboratory. Instead, we do a paternal phenotype if antibodies to the relevant antigens are available and make a statistical risk assessment based on that. Blood group antibodies anti-A and anti-B are responsible for neonatal hemolysis and hyperbilirubinemia, which in rare cases necessitate treatment with transfusion (see Maternal ABO Antibodies).

Flow cytometry (FC) is a useful method for small population detection and quantification, for example, after intrauterine transfusion (IUT) for detection of fetal RBCs and donor RBCs. Also, minute samples of fetal blood can be examined for multiple parameters improving laboratory guidance (see FC in HDFN, Fetus and Newborn). In this paper, we present the procedures related to laboratory screening and monitoring in HDFN as currently performed at the Copenhagen University Hospital, Rigshospitalet, in Denmark.

**Routine Blood Group Typing and Screening for Irregular Antibodies, Rh Prophylaxis**

Blood samples from the 1st trimester initial pregnancy consultation with the general practitioner are typed for ABO and RhD blood groups and an antibody screen is performed. We use automated equipment and Capture R® Ready-Screen (I and II) for detection of IgG antibodies to RBC antigens. Typing identifies the RhD-negative women who can develop anti-D antibodies. Antibody screening identifies those who have already developed alloantibodies in the 1st trimester, whether RhD positive or RhD negative.

At GA 25 weeks, RhD-negative women with a negative 1st trimester antibody screen are offered routine non-invasive fetal antenatal RHD screening, and the repeated routine antibody screening is offered only to RhD-negative women (see Antenatal RHD Screening). At GA 29 weeks, the nonimmunized pregnant woman is offered intramuscular injection of 250–300 µg RhIg by the midwife if RHD is detected in the cell-free fetal DNA (cfDNA).
from plasma. The 250–300 μg RhIg injection is repeated within 72 h after delivery. Investigation for fetomaternal hemorrhage and quantification by flowcytometry is only made if hemorrhage is suspected.

**Investigating and Monitoring Irregular Antibodies**

For alloimmunized women, identification of the target antigen of the alloantibody will be conducted to provide further information on the potential clinical impact of the alloimmunization. Maternal antibodies targeting a distinct blood group antigen often lead to a known distinct pattern of clinical manifestations. This knowledge guides the planning of the laboratory monitoring and the fetal specialist surveillance.

For the antibody identification we use 11 different reagent in-house single donation glycerol frozen-thawed RBCs and anti-IgG column agglutination technique (CAT). The woman's own RBCs are included in the panel to distinguish allo- and autoantibodies. If Rh antibodies are suspected the examination is extended with a panel of papain-treated RBCs. Per definition, for an alloantibody to be present, a phenotype of the woman's own RBCs should demonstrate absence of the target antigen of the alloantibodies.

Some antibodies have empirically been found to be of no clinical significance (anti-N, -Lea, -Leb, -A1, -IH, -I), whereas others are known to be of potential dire consequences (anti-K, -c) and referral to a fetal medicine center should be considered regardless of titer, and yet another group (anti-D, -C, -E, -e, -C\(^*\), -Kp\(^*\), -Kp\(^+\), -K, -Jk\(^*\), -Jk\(^+\), -Fy\(^a\), -Fy\(^b\), -S, -s, -Wr\(^a\), -M, -P1, -Lu\(^a\), and -Lu\(^b\)) is referred to a fetal medicine center if a titer above 16 is measured.

At GA 25 and 32 weeks, the alloimmunized woman is routinely examined with a measurement of titer and with a screening for additional antibodies. A titer above 16 for the latter group of antibodies is empirically determined as the threshold value indicating increased risk of HDFN and warrants closer surveillance by a fetal medicine specialist with serial Doppler ultrasound measurements of the peak systolic velocity (PSV) in the fetal middle cerebral artery (MCA) [10, 11].

Semi-quantification of alloantibody is done by a serial 2-step dilution of plasma in saline followed by examination with CAT. The titer is defined as the inverse of the antibody content of the pregnant woman at the time of sampling; the titer might increase rapidly because of continuing exposure to fetal or donor RBCs. Within days additional antibodies may also develop. Therefore, a serial monitoring is important.

**Antenatal RHD Screening**

As part of a targeted RhIg prophylaxis program for non-immunized RhD-negative women, knowledge of the fetal RhD type helps restrict prophylaxis to those women only who carry an RhD-positive fetus [12, 13]. This restriction avoids superfluous exposure to prophylaxis in women carrying an RhD-negative fetus and reduces the overall use of RhIg, which is a limited resource [14, 15].

The fetal RhD status is predicted by analysis of cell-free DNA in the maternal plasma that also contains maternally derived cell-free DNA. Presence of the fetal RHD gene indicates that the fetus is RhD positive. Since the first reports of cell-free fetal RHD in maternal plasma [16, 17], non-invasive fetal RHD genotyping has become highly integrated into clinical medicine, and its accurate performance has been covered comprehensively in the literature [12, 13, 18–24].

As an antenatal screening to guide RhIg prophylaxis, non-invasive prenatal testing of fetal RHD has been introduced as a nationwide clinical service in several European countries [16–25]. Evaluations of national programs have demonstrated high test accuracy, with sensitivities of >99.9% around 25 weeks of gestation and >99% from GA 10 weeks [13]. Recent recommendations for validation and quality assurance of fetal RHD genotyping have been prepared [25].

The Copenhagen setup for antenatal RHD screening has been described in detail [26–28]. Briefly, blood samples are taken by the general practitioner at GA 25 weeks. Blood samples arrive at the laboratory after an average of 4 days in transport (up to 7 days are accepted). Plasma is separated and DNA is extracted from 1 mL of plasma. Eluted DNA is tested by real-time PCR targeting RHD exons 7 and 10 in a duplex manner with the same dye, which increases the analytical sensitivity [29]. The RHD PCR is sensitive enough to detect one genome equivalent (geq) per PCR [30], and the overall detection limit of the setup...
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Non-Invasive Prediction of Fetal K, RhC, Rhc, RhE and ABO Blood Group

We have recently reported a procedure based on next-generation sequencing (NGS) analysis of PCR-amplified cfDNA from maternal plasma for prediction of the fetal blood group [34–37]. As some fetuses may die from HDFN as early as GA 18 weeks, it is necessary to be able to predict the fetal blood group early in pregnancy. We use this general approach to predict fetal K, RhC, Rhc, RhE, and ABO blood groups in cases with a risk of HDFN due to maternal production of the corresponding antibodies [34–37].

The NGS based analysis can detect the presence or absence of alleles encoding incompatible antigens on the fetal RBCs. NGS is a powerful technology that enables the parallel sequencing of many million DNA sequences. We use this technology in a very simple approach: cfDNA is purified from 4 mL of maternal plasma and after PCR amplification of the genetic basis of the blood group, the PCR product is sequenced to great depth. The number of times that the blood group SNV in question occurs is counted and the relative frequency of the SNV, exceeding the background threshold, will be the basis of the prediction. As there are some background reads due to errors of PCR amplification and sequencing, this threshold is important to determine empirically. The background is remarkably low with an empirical threshold for a positive sample of approximately 0.05% positive reads.

Preanalytical conditions are important to address as these NGS-based assays rely on maintaining the in vivo ratio of fetal versus maternal SNVs. Thus, it is important to ensure that maternal cells do not contribute DNA after blood sampling. Taking blood samples in Streck tubes is highly recommended. Some factors are important, such as keeping the amplicons short and keeping spurious amplification to a minimum. The volume of plasma interrogated is approximately 1 mL. Finally, the data analysis we currently employ is two-pronged: one analysis of the fastq sequences is done using FastQC software and another analysis is performed using simple string searches with grep in a Linux formatted PC. We do not perform alignment-based analysis.

Even though the prediction of the different blood groups: K, RhC, Rhc, RhE, and ABO is based on the same generic method, there are important differences in respect to primer design and data analysis. For instance, in the case of RhC prediction, background reads from highly homologous sequences of RHD may complicate the prediction. The ABO prediction requires the combined results of two primer sets for an antigen prediction.

After implementation of the two-pronged data analysis, we have not yet had any discordant results from a small cohort of samples. As a postnatal blood group was not determined in many cases, a significant number of samples have not been used for formally validating the results of the prenatal fetal blood group prediction. Development of laboratory methods, validation as well as continuous quality control, is dependent on meticulous and continuous contribution from laboratories and clinicians. O’Brien et al. [38] have used digital PCR for fetal blood group prediction, and Orzińska et al. [39] have also used NGS for fetal blood group prediction.

Maternal ABO Antibodies

ABO incompatibility is now the most prevalent cause of HDFN with hyperbilirubinemia in developed countries due to the success of Rh prophylaxis [2, 6, 40]. A recent Danish study found ABO incompatibility in 15 of 21 cases with total serum bilirubin ≥600 μmol/L, comprising a significant risk of kernicterus spectrum disorder [41]. Furthermore, rare cases of fetal hemolysis, anemia, and hydrops fetalis caused by ABO antibodies have been described [42, 43].

Currently, we do not have a systematic screening procedure for maternal ABO antibodies harmful to the fetus and newborn [44–46]. Maternal anti-A and anti-B IgG titers are predictive of neonatal requirement for treatment of hyperbilirubinemia [47, 48]. However, we found the positive predictive values both in the 1st trimester (65%) and perinatally (73–76%) to be too low to be used clinically for routine screening and we aim for enhancement of predictive values [49] by on-going research.

We have described the use of two antibody screening methods: (i) solid phase red cell adherence assay (SPR-CA) only detecting IgG anti-A and anti-B and (ii) manual anti-IgG CAT detecting both IgG and IgM reacting at 37°C. The two methods yielded comparable results. SPR-CA is most suitable for batch analysis, whereas CAT is amenable for single sample analysis.
Standard infant transfusion practice in our health care region is ABO-identical RBCs. Therefore, in addition to an antibody screening test for irregular antibodies, we also perform a determination of regular anti-ABO antibodies of the IgG class of the incompatible newborn to be transfused. Detection of IgG anti-A and anti-B is followed by determination of the maternal IgG anti-A and anti-B titer. This is likely to lead to identification of more women with high-titer IgG anti-A and anti-B.

**Laboratory Monitoring, anti-A and anti-B**

In pregnancies with identified maternal high-titer IgG anti-A and anti-B or a history of a previous pregnancy where maternal anti-A/B was responsible for HDFN, the anti-A and anti-B IgG titer is determined in the 1st trimester as well as at GA 32 weeks. For the methods described a common cut-off value of 512 was initially found for anti-A/B [49]. However, additional studies (in preparation) showed that distinct cut-off values for anti-A and anti-B increased accuracy. Therefore, we now apply a cut-off value of 512 for maternal anti-A and 256 for anti-B. The cut-off value is used for recommendation of antenatal non-invasive fetal ABO blood group prediction and for fetal monitoring by ultrasound of MCA-PSV in case of incompatible antigens on fetal RBCs. The flow chart presented in Figure 1 presents the complete laboratory monitoring of HDFN.

**FC in HDFN, Fetus and Newborn**

Agglutination techniques are informative in most cases, but by supplementing with FC more detailed and semiquantitative information [50] can be produced also in unexpected urgent cases of suspected HDFN where diagnosis is initially uncertain. Determination of fetal and newborn antigens and direct antiglobulin test (DAT)-positive RBCs can be made impossible or inconclusive by access to a limited volume of sample, small surviving populations of fetal cells after multiple IUTs, and due to weak fetal expression of antigens [51]. FC enables quantification of subpopulations, for example several populations of distinct RBC phenotype in cases of mixed populations of donor and patient cells, enabling measurement of the survival of the infant’s own RBCs, as well as donor RBCs.

In Figure 2, we present an example of serial monitoring of various parameters of a severely anemic newborn, with hemoglobin (Hb) at birth of 6.3 g/dL (3.9 mmol/L). The RhD-positive woman unexpectedly delivered an anemic infant in GA 38 weeks. Upon investigation after delivery, the mother had an allo-anti-E, titer of 2,048. The anti-E developed between the 1st trimester antibody screening and delivery. The newborn was DAT positive.

Immediately after birth the newborn was given a transfusion with compatible donor RBCs, and again on day 10 and day 26 in accordance with guidelines for treatment of anemia in the newborn. Initially, a hepatic cause, Alagille syndrome, was suspected. However, only HDFN was found. To substantiate the diagnosis of HDFN, we used FC to determine a series of percentages of newborn E-positive RBC.

Newborn E-positive RBCs were identified in FC by reacting RBCs with reagent anti-E followed by anti-human IgG conjugated to a fluorophore, as previously described [52]. Total hemoglobin and reticulocytes were measured with a hematology analyzer. To corroborate E-positive results, we supplemented with measurement by FC of fetal hemoglobin (HbF) and obtained similar results (data not shown) [50].

We observed a close correlation between the waning of the allo-anti-E and an increasing survival of fetal RBCs. Figure 2 demonstrates that the effects of the anti-E is reflected in the newborn RBCs for more than 72 days and that survival of the newborn is dependent on transfusion therapy during the first 47 days.

Generally, if a fetus has received intrauterine transfusion it is possible to monitor the percentage of fetal versus donor RBCs. This can be done in several ways depending on the specific situation, but typically we use as a marker the antigen targeted by the maternal antibodies, positive RBCs are fetal, antigen-negative RBCs are from the donor due to the use of compatible blood. The percentage of fetal RBCs is also measurable with the marker HbF [50].

**Determining the Actual Clinical Course, Doppler Ultrasonography MCA-PSV for Non-Invasive Prediction of Fetal Anemia**

Maternal alloantibodies and fetal expression of the corresponding RBC antigen is the prerequisite for HDFN. However, a large variation in clinical impact is observed with identical laboratory findings. Even in the same woman clinical variation occurs from one antigen-positive fetus to another despite an unchanged alloantibody titer [53]. Supplementary modalities of monitoring are needed to determine the actual clinical consequence of the alloimmunization.

Measurement of MCA-PSV is the golden standard for non-invasive prediction of fetal anemia. Mari et al. [11] showed that a cut-off of 1.5 multiples of median on Doppler ultrasound measurement of MCA-PSV has 100% sensitivity with a false-positive rate of 12% in the prediction of moderate to severe anemia in the non-hydropic fetus. Timely identification of significant fetal anemia is the basis for therapeutic intervention with intrauterine blood transfusion or delivery, depending on GA and thereby preventing fetal demise.

**The Newborn in HDFN**

When the fetus becomes a newborn it might still be suffering from anemia and the other pathophysiologic conse-
Fig. 1. The flow chart illustrates the use of individual components of the laboratory HDFN monitoring of all pregnant women. RhD-positive and RhD-negative women are screened for irregular antibodies against RBC antigens in the 1st trimester. Antibodies to ABO blood group antigens are only included in the examination if supplementary information gives an indication to do so. RhD-positive women who test negative at this first antibody screening are not examined later. All RhD-negative women are re-tested for antibodies at GA 25 weeks, and the antenatal RHD screening based on cfDNA is performed. RHD screening showing an RHD-positive fetus leads to the administration of Rh prophylaxis at GA 29 weeks and subsequently also the postnatal prophylaxis. Immunized RhD-negative women are screened for antibodies again at GA 32 weeks. At GA 25 weeks and GA 32 weeks, RhD-positive women with irregular antibodies detected in the first trimester are screened. Alternative timing of examination for both RhD-negative and positive women is followed if the clinician decides so.

* Shows a potential trigger for conducting antenatal antigen prediction. The specific criteria for antenatal antigen prediction are: C, c, K, or D at titer ≥1; E at titer >1; A at titer ≥512; B at titer ≥256. For blood groups A and B, also HDFN due to anti-A or anti-B in a previous pregnancy gives an indication for antigen prediction.

** Shows a potential trigger for referral to the fetal medicine center. The specific criteria for referral are: anti-D, -C, -E, -e, -Cw, -Kpa, -Kpb, -k, -Jka, -Jkb, -Pya, -Pyb, -S, -s, -Wra, -M, -P1, -Lua, -Lub titer >16, and anti-K, -c titer ≥1.

*** Designates postnatal monitoring of the newborn with titer, serological antigen detection, and flow cytometric quantification of fetal and donor RBCs. A black box designates an analysis, a grey box designates a result, and a white box designates Rh prophylaxis.
quences of the persisting maternal antibody [54] present in the newborn. In most cases the fetal RBCs will carry maternal antibodies detectable by the DAT. We routinely determine the titer of free alloantibody in the plasma of the newborn and determine the fetal blood group antigen targeted by the maternal antibody. The latter is routinely done to assess the quality of laboratory work. FC-based measurement of fetal versus donor cells is decided in each case.

The laboratory should be aware of the importance of information being shared with the team of neonatologists providing postnatal care for the newborn. It should be remembered that laboratory investigation of the mother might still be relevant and can yield valuable information, for example examination for antibodies, phenotype, determination of titers, fetomaternal hemorrhage, especially in the RhD-positive women who have not been tested since the 1st trimester.

Further Preventive Measures to Avoid Alloimmunization

Prevention of alloimmunization due to transfusion in girls and women of premenopausal age, or under the age of 50 years, has been implemented in some countries by matching a limited number of RBC antigens. Basic matching of ABO and RhD blood groups is supplemented by supplying K-negative RBC components for premenopausal women in Denmark. Matching has been extended to routinely encompass Rhc and E in some countries [3].

A study on the effect of matching donor and recipient in IUT indicates that an efficient prevention of alloimmunization (64%) can be achieved by an extended phenotypic match: C, c, E, K, Fy, Jk, S [55]. Another study in ordinary transfusion recipients demonstrated that matching for C, c, E, K, Jk could prevent 78% of immunizations, and enhanced matching for C, c, E, K, Fy, Jk, Cw improved prevention to 83.4% of immunizations [56]. We have implemented matching for IUT for C, c, E, K, Fy, Jk with a pragmatic view for the available supply. However, our extensive genotyping of donors helps making matches possible by access to ample donor genotype information [57].

Platelet transfusion seems to be a source of alloimmunization that could be taken into consideration. Small amounts of RBCs in the platelet component are enough to immunize. We administer RhIg if, for logistical reasons, D-positive platelet or plasma components must be given to a female RhD-negative recipient of premenopausal age.
Perspectives for Optimization and Future Developments

Several studies have addressed the feasibility of screening all pregnant women for irregular antibodies in the 3rd trimester, not only RhD-negative women. First trimester screening of all pregnant women is already implemented in many health care systems. A 3rd trimester repeated screening of RhC-negative women has been proposed. Focusing on individuals with a high risk of immunization would enhance cost benefit in comparison with screening all women [3, 8].

Some individuals develop alloantibodies after alloantigen exposure whereas others can be transfused repeatedly without being alloimmunized [58, 59]. Elucidation of the genetic background for individual propensity to develop alloantibodies as well as the genetic background for regulation of quantities of antibodies produced has been attempted by several groups [60–62]. Access to this information for pregnant women would potentially add useful guidance to the clinical risk assessment of a specific woman.

It has also been attempted to interfere with an established immune response by administration of peptides derived from the antigen to end the active production of antibody [63, 64]. Another approach is administration of non-destructive antibodies competing for the antigen to the alloimmunized woman [65, 66].

Conflict of Interest Statement

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Author Contributions

All authors contributed to writing of the text. All authors have read and accepted the final version of the manuscript.

References

1 Bowman JM, Pollock JM, Penston LE. Fetal-maternal transplacental hemorrhage during pregnancy and after delivery. Vox Sang. 1986; 51(2):117–21.
2 de Haas M, Thurik FF, Koelwijn JM, van der Schoot CE. Haemolytic disease of the fetus and newborn. Vox Sang. 2015 Aug;109(2): 99–113.
3 Koelwijn JM, Vrijkotte TG, de Haas M, van der Schoot CE. Integration of noninvasive prenatal testing for fetal RhD status in RhD-negative women: a systematic review and meta-analysis. Transfus Med Rev. 2021;35(2):85–94.
4 Finning K, Martin P, Summers J, Daniels G. Fetal genotyping for the K (Kell) and Rh C, c, and E blood groups on cell-free fetal DNA in maternal plasma. Transfusion. 2007 Nov; 47(11):2126–33.
5 Moise KJ. Red blood cell alloimmunization in pregnancy. Semin Hematol. 2005 Jul;42(3): 169–78.
6 Materi M, Deter RL, Carpenter RL, Raham F, Zimmerman R, Moise KJ Jr., et al. Noninvasive diagnosis by Doppler ultrasonography of fetal anemia due to maternal red-cell alloimmunization. Collaborative Group for Doppler Assessment of the Blood Velocity in Anemic Fetuses. N Engl J Med. 2000 Jan 6;342(1): 9–14.
7 Clausen FB. Lessons learned from the implementation of non-invasive fetal RHD screening. Expert Rev Mol Diagn. 2018 May;18(5): 423–31.
8 van der Schoot CE, de Haas M, Clausen FB. Genotyping to prevent Rh disease: has the time come? Curr Opin Hematol. 2017 Nov; 24(6):544–50.
9 Bills VL, Soothill PJ. Fetal blood grouping using cell free DNA – an improved service for RhD negative pregnant women. Transfus Apher Sci. 2014 Apr;50(2):148–53.
10 Kent J, Farrell AM, Soothill P. Routine administration of Anti-D: the ethical case for offering pregnant women fetal RHD genotyping and a review of policy and practice. BMC Pregnancy Childbirth. 2014 Feb 25;14:87.
11 Faas BH, Beuling EA, Christiaens GC, von dem Borne AE, van der Schoot CE. Detection of fetal RHD-specific sequences in maternal plasma. Lancet. 1998 Oct 10;352(9135):1196.
12 Clausen FB, Damkjaer MB, Dziegel MH. Noninvasive prenatal prediction of fetal blood group into clinical prenatal care. Prenat Diagn. 2014 May; 34(5):409–15.
13 Clausen FB, Dammacker MB, Dziegel MH. Noninvasive fetal RhD genotyping. Transfus Apher Sci. 2014 Apr;50(2):154–62.
14 Daniels G, Finning K, Martin P, Massey E. Noninvasive prenatal diagnosis of fetal blood group phenotypes: current practice and future prospects. Prenat Diagn. 2009 Feb;29(2): 101–7.
15 Runkel B, Bein G, Sieben W, Sow D, Polas S, Fleer D. Targeted antenatal anti-D prophylaxis for RhD-negative pregnant women: a systematic review. BMC Pregnancy Childbirth. 2020 Feb 7;20(1):83.
16 Yang H, Llewellyn A, Walker R, Harden M, Saramago P, Griffin S, et al. High-throughput, non-invasive prenatal testing for fetal rhesus D status in RhD-negative women: a systematic review and meta-analysis. BMC Med. 2019 Feb 14;17(1):37.
17 Alshehri AA, Jackson DE. 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. Transfus Med Rev. 2021;35(2):85–94.
18 Alshehri AA, Jackson DE. 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. Transfus Med Rev. 2021;35(2):85–94.
19 Clausen FB. Integration of noninvasive prenatal prediction of fetal blood group into clinical prenatal care. Prenat Diagn. 2014 May; 34(5):409–15.
20 Clausen FB, Damkjaer MB, Dziegel MH. Noninvasive fetal RhD genotyping. Transfus Apher Sci. 2014 Apr;50(2):154–62.
21 Daniels G, Finning K, Martin P, Massey E. Noninvasive prenatal diagnosis of fetal blood group phenotypes: current practice and future prospects. Prenat Diagn. 2009 Feb;29(2): 101–7.
22 Runkel B, Bein G, Sieben W, Sow D, Polas S, Fleer D. Targeted antenatal anti-D prophylaxis for RhD-negative pregnant women: a systematic review. BMC Pregnancy Childbirth. 2020 Feb 7;20(1):83.
23 Yang H, Llewellyn A, Walker R, Harden M, Saramago P, Griffin S, et al. High-throughput, non-invasive prenatal testing for fetal rhesus D status in RhD-negative women: a systematic review and meta-analysis. BMC Med. 2019 Feb 14;17(1):37.

Author Contributions

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Transfus Med Hemotheor 2021;48:306–315
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24 Zhu YJ, Zheng YR, Li L, Zhou H, Liao X, Guo JX, et al. Diagnostic accuracy of non-invasive fetal RhD genotyping using cell-free fetal DNA: a meta analysis. *J Matern Fetal Neonatal Med*. 2014 Dec;27(18):1839–44.

25 Clausen FB, Hellberg A, Bein G, Bugert P, Schwartz D, Dierksev TD, et al. Recommendation for validation and quality assurance of non-invasive prenatal testing for foetal blood groups and implications for IVD risk classification according to EU regulations. *Vox Sang*. 2021. doi: 10.1111/vox.13172.

26 Clausen FB, Christiansen M, Steffensen R, Jørgensen S, Nielsen C, Jakobsen MA, et al. Report of the first nationally implemented clinical routine screening for fetal RHD in D–pregnant women to ascertain the requirement for antenatal RhD prophylaxis. *Transfusion*. 2012 Apr;52(4):752–8.

27 Clausen FB, Rieneck K, Krog GR, Bundgaard BS, Dziegiel MH. Noninvasive antenatal screening for fetal RHD in RhD negative women to guide targeted anti-D prophylaxis. *Methods Mol Biol*. 2019;1885:347–59.

28 Clausen FB, Steffensen R, Christiansen M, Rudby M, Jakobsen MA, Jakobsen TR, et al. Routine noninvasive prenatal screening for fetal RHD in plasma of RhD-negative pregnant women – 2 years of screening experience from Denmark. *Prenat Diagn*. 2014 Oct;34(10):1000–5.

29 Clausen FB, Krog GR, Rieneck K, Råsmark M, Steffensen R, Christiansen M, et al. Next generation sequencing-based fetal abo blood group prediction by analysis of cell-free DNA from maternal plasma. *Transfus Med Hemother*. 2020 Feb;47(1):45–53.

30 Bőr J, Halász F, Helmanty A. Non-invasive prenatal testing (NIPT) for fetal Kell, Duffy and Rh blood group antigen prediction in alloimmunised pregnant women: power of droplet digital PCR. *Br J Haematol*. 2020 May;189(3):e90–e.

31 van de Watering M, Verheggen L, Abelen W, et al. DNA from maternal plasma. *Transfus Med*. 2015 Sep;123(9):731–9.

32 Haimila K, Sulin K, Kuosmanen M, Sareneva T, et al. Maternal IgG anti-A and anti-B titres predict the requirement for antenatal RhD prophylaxis. *J Perinat Med*. 2015 Oct;43(5):e90–4.

33 Wilde J, Benes T, Srb DJ, et al. Maternal IgG anti-A titre and maternal age and parity as risk factors for ABO haemolytic disease of the newborn (HDN). *Arch Dis Child Fetal Neonatal Ed*. 1998 May;78(3):F220–1.

34 Foundations for Research on Inborn Errors of Metabolism. Fetal alloimmunization and phenotype from cell-free fetal DNA in maternal plasma. *Transfusion*. 2013 Nov;53(11 Suppl 2):2892–8.

35 Rieneck K, Clausen FB, Dziegiel MH. Next-generation sequencing for antenatal prediction of KEL1 blood group status. *Methods Mol Biol*. 2015;1310:115–21.

36 Rieneck K, Clausen FB, Dziegiel MH. Noninvasive antenatal determination of fetal blood group status using next-generation sequencing. *Cold Spring Harb Perspect Med*. 2015 Oct 28;6(1):a023093.

37 Rieneck K, Egeberg Hothier C, Clausen FB, Jakobsen MA, Bergholt T, Hellmuth E, et al. Next generation sequencing-based fetal abo blood group prediction by analysis of cell-free DNA from maternal plasma. *Transfus Med Hemother*. 2020 Feb;47(1):45–53.

38 O'Brien H, Hyland C, Schoeman E, Flower R, Daly J, Gardener G. Non-invasive prenatal testing (NIPT) for fetal Kell, Duffy and Rh blood group antigen prediction in alloimmunised pregnant women: power of droplet digital PCR. *Br J Haematol*. 2020 May;189(3):e90–e.

39 Van der Ploeg CP, Thurik FF, de Haas M, Janssen H, Huiskes E, Goldschmeding R, Overbeke MA, et al. Protection against immune haemolytic disease of newborn infants by maternal monocyte-reactive IgG alloantibodies (anti-HLA-DR). *Lancet*. 1995 Feb 25;345(8910):1067–70.

40 Chen YJ, Ling UP. Prediction of the development of neonatal hyperbilirubinemia in ABO incompatibility. *Zhonghua Yi Xue Za Zhi*. 1994 Jan;53(1):13–8.

41 Gourdin MF, Lejonc JL, Cartron JP, Gourdin MF, Lejonc JL, Cartron JP. The heterogeneity of erythrocyte antigen distribution in human newborns. *Curr Opin Hematol*. 2006 Nov;13(6):490–5.

42 Reyes F, Gourdin MF, Lejonc JL, Cartron JP, Gourdin MF, Lejonc JL, Cartron JP. The heterogeneity of erythrocyte antigen distribution in human newborns. *Curr Opin Hematol*. 2006 Nov;13(6):490–5.

43 Krog GR, Donneborg ML, Hansen BM, Eeg O, Johannessen H, Clausen FB, Jensen KV, et al. Prediction of ABO hemolytic disease of the newborn according to EU regulations. *Vox Sang*. 2021. doi: 10.1111/vox.13172.

44 Kaplan M, Hammerman C, Vreman HJ, Mikkola M, Brand A, Zijlstra M, Schonewille H. Severe fetal anaemia. *Transfus Med*. 2005 Feb;15(1):57–60.

45 McDonnell M, Hannam S, Devane SP, O'Neill M, Marmont D, O'Neill M, Marmont D. Antenatal RhD screening for antenatal prediction of fetal phenotype from cell-free fetal DNA in maternal plasma. *Transfus Med*. 2015 Sep;123(9):731–9.

46 Sarici SU, Yurdakök M, Serdar MA, Oran O, Gürsoy G, TUNC N, TUNC N. Diagnostic accuracy of non-invasive prenatal testing (NIPT) for fetal Kell, Duffy and Rh blood group antigen prediction in alloimmunised pregnant women: power of droplet digital PCR. *Br J Haematol*. 2020 May;189(3):e90–e.

47 Böhringer M, Hondeghem JL, Batista MG, et al. Non-invasive prenatal testing (NIPT) for fetal Kell, Duffy and Rh blood group antigen prediction in alloimmunised pregnant women: power of droplet digital PCR. *Br J Haematol*. 2020 May;189(3):e90–e.

48 Orzińska A, Guz K, Mikula M, Kluska A, Balducci B, et al. Sensitivity of fetal RHD screening for safe guidance of targeted anti-D immunoglobulin prophylaxis: prospective cohort study of a nationwide programme in the Netherlands. *BMJ*. 2016 Nov 7;355:i5789.

49 Orzińska A, Guz K, Mikula M, Kluska A, Balducci B, et al. Sensitivity of fetal RHD screening for safe guidance of targeted anti-D immunoglobulin prophylaxis: prospective cohort study of a nationwide programme in the Netherlands. *BMJ*. 2016 Nov 7;355:i5789.

50 Dziegiel MH, Nielsen LK, Berkowicz A. Determination of IgG anti-D titre in ABO incompatible pregnancies. *Transfus Med*. 2003 Sep;13(3):201–4.

51 Reyes F, Gourdin MF, Lejonc JL, Cartron JP, Gourdin MF, Lejonc JL, Cartron JP. The heterogeneity of erythrocyte antigen distribution in human newborns. *Curr Opin Hematol*. 2006 Nov;13(6):490–5.

52 Gourdin MF, Lejonc JL, Cartron JP, Gourdin MF, Lejonc JL, Cartron JP. The heterogeneity of erythrocyte antigen distribution in human newborns. *Curr Opin Hematol*. 2006 Nov;13(6):490–5.
61 Tan JCG, Yuan FF, Daley J, Marks K, Flower RL, Dyer WB. D-immunized blood donors who are female and who possess at least one HLA-DRB1*15 allele show a propensity for high serum RhIG production. Transfusion. 2018 May;58(5):1182–8.

62 Verduin EP, Brand A, van de Watering LM, Roelen DL, Kanhai HH, Doxiadis II, et al. The HLA-DRB1*15 phenotype is associated with multiple red blood cell and HLA antibody responsiveness. Transfusion. 2016 Jul;56(7):1849–56.

63 Hall AM, Cairns LS, Altmann DM, Barker RN, Urbaniak SJ. Immune responses and tolerance to the RhD blood group protein in HLA-transgenic mice. Blood. 2005 Mar 1;105(5):2175–9.

64 Hall LS, Hall AM, Pickford W, Vickers MA, Urbaniak SJ, Barker RN. Combination peptide immunotherapy suppresses antibody and helper T-cell responses to the RhD protein in HLA-transgenic mice. Haematologica. 2014 Mar;99(3):588–96.

65 Mathiesen L, Nielsen LK, Andersen JT, Grevys A, Sandlie I, Michaelsen TE, et al. Materno-fetal transplacental transport of recombinant IgG antibodies lacking effector functions. Blood. 2013 Aug 15;122(7):1174–81.

66 Nielsen LK, Green TH, Sandlie I, Michaelsen TE, Dziegiel MH. In vitro assessment of recombinant, mutant immunoglobulin G anti-D devoid of hemolytic activity for treatment of ongoing hemolytic disease of the fetus and newborn. Transfusion. 2008 Jan;48(1):12–9.