A novel splice-site mutation in RHD gene associated with RhD negative phenotype

Maria Antonietta Villa, Nicoletta Revelli, Cinzia Paccapelo, Francesca Truglio, Araitz Molano, Mónica López

ABSTRACT

Introduction: The Rh antigens are encoded by the highly polymorphic RHD and the RHCE genes. Individuals with D variants may make anti-D alloantibodies, which in pregnant women cause severe or fatal hemolytic disease of the fetus and newborn. Accurate D typing using reliable routine methods must be performed. A number of unusual RHD alleles are being currently reported due in large part to the growing implementation of molecular analyses. Case Report: A sample from a Caucasian male blood donor at Northern Italy was serologically classified as RhD negative (Ccee). Instead, the sample was genotyped with all SSP-PCR kits as RhD positive, Dd and the result obtained with Partial D-TYPE (Dva or Va like or Va associated or DBS) was consistent with a suggestion of a new variant. Through a microarray platform analysis (BLOODchip Reference), the sample was genotyped as DV/DBS [RHD-CE(S)-D hybrid], associated with a predicted partial D phenotype for the RhD group, due to the lack of a hybridization signal for RHD exon 5. The PCR analysis of the exon 5 of the RHD gene gave a positive amplification result and the sequencing revealed a polymorphism in hemizygous or homozygous state in position RHD*IVS5+1t. This novel splice-site mutation alters or completely abolishes the specific sequence where the splicing of RHD gene intron 5 takes place. Conclusion: The novel RHD mutation hereby described highlights the significance of using RHD genotyping to confirm and/or clarify D antigen uncertainties in order to prevent the unnecessary immunization of patients.

Keywords: D variants, Novel RHD mutation, Sequencing

INTRODUCTION

The Rh blood group is one of the most complex and polymorphic systems of red blood cells (RBCs) known in humans. Due to its high immunogenicity the Rh antigens play a crucial role in transfusion medicine, particularly in obstetrics, as they are the main cause of hemolytic...
disease of the fetus and newborn (HDFN) [1]. The Rh antigens expressed on the red blood cell (RBC) surface are defined by a trimeric complex of membrane polypeptides encompassing the Rh proteins and the Rh-associated glycoprotein (RhAG). The Rh antigens are encoded by two highly polymorphic genes located on the short arm of chromosome 1, the RHD and the RHCE genes. Each of these genes, originating from a duplication of a common gene ancestor, are comprised of 417 amino acids and 10 exons, form the Rh locus. The RHD gene encodes the RhD protein, whereas the RHCE gene encodes the RhCcEe polypeptides carrying the C (RH2)/c (RH4) and E (RH3)/e (RH5) antigens [2].

Due to their similarity, close proximity and opposite orientation, the two genes promote numerous genetic rearrangements, giving rise to variant alleles. The D variants encoded by these alleles can arise from different mechanisms such as nucleotide insertions or deletions, single or multiple missense mutations, splice site mutations, and gene conversions [3, 4]. The RHD alterations, affecting a partially or exclusively extracellular RhD section, result in the partial D variants, leading to type-specific D epitope lacking or diminishing the D antigen density. The partial D variants include partial D, weak D and DEL types, and their identification is of crucial importance in transfusion medicine [5]. Individuals with partial D variants may make alloantibodies anti-D against those D epitopes, sometimes in pregnant women causing severe or fatal HDFN [6–8]. Most RBCs known as weak D are thought to express D antigens altered in a quantitatively, but not qualitatively, form. Therefore, such individuals appear not to develop alloantibodies anti-D upon contact with normal D positive RBCs.

Reduced expression of D antigen, quantitatively or qualitatively, occurs in 0.3–1% of Caucasian people. Although the most common cause of the D negative in people of European origin is homozygosity for a complete deletion of RHD gene, the association of more than 115 mutations and RHD-RHCE gene conversion with this phenotype has also been described [9].

In past years, a number of unusual RHD alleles have been reported due to the growing use of several different known D epitope specificity monoclonal antibodies [10] and the evaluation of RHD alleles by the implementation of molecular analyses [11]. For that reason, D typing using reliable routine D typing methods must be performed in mother-to-be transfusion recipients, with special care in the case of blood donors with weak expression variants.

**CASE REPORT**

We describe a case of weak reactivity of D antigen found in a Caucasian male blood donor resulting from nucleotide change. The sample was sent, after obtaining a signed informed consent, to our Immunohematology Reference Laboratory (IRL) due to incongruent results detected in RhD typing from the Northern Italy Transfusion Blood Centre. Routine serological D typing was performed in hemagglutination with a fully automated system (Galileo, Immucor, Norcross, GA, USA) at room temperature by immediate spin employing a liquid-phase in microplate method (Select R, Immucor) using two different blended monoclonal reagents, the DUO IgM TH28 + IgG MS26 and NOVACLONE D175+D415 clones (Immucor). The sample was further tested by Innova Instrument (Ortho Biouv System, Ortho Clinical Diagnostics Raritan, NJ, USA) employing the ABOD1D2 card (Ortho Clinical Diagnostics) in column agglutination technology (CAT) using two anti-D monoclonal antibodies D7B8 and RUM1 (Ortho). The Weak D indirect antiglobulin test (IAT) was performed in solid-phase red cells adherence (SPRCA) with the same reagents of the microplate method and in CAT with the anti-D clones D7B8+H1121G6 +LORIFA (Ortho Clinical Diagnostics).

Extended Rh(D) typing for epitope detection was done using Galileo (Immucor) with the ALBACloner Advanced Partial RhD typing kit (Albabioscience, Ellen’s Glen Road, Edinburgh, UK) to differentiate partial RhD types and weak RhD types 1 and 2. The phenotype was determined using the microplate method (Galileo) with monoclonal antibodies specific for Rh antigens (CcEe). All tests were performed according to the manufacturers’ recommendations. These devices and reagents have CE Marking license in Europe.

A bench top apparatus for fully automated nucleic acid purification (QIAcube—Qiagen, Hilden, Germany) was used, according to the manufacturer’s instructions, to extract DNA from whole blood cells. DNA analysis was performed using single specific primer-polymerase chain reaction (SSP-PCR) with Ready Gene CDE kit (Inno-Train Diagnostik, GmbH, Kronberg im Taunus, Germany) and RH-TYPE, Weak D-TYPE, Partial D-TYPE kits (BAGene, BAG Health Care GmbH, Lich, Germany). Subsequently, to perform extended typing, the sample was sent to the Grifols-Progenika Reference Laboratory and tested by a microarray platform (BLOODchip Reference v4.0, Progenika Biopharma, Derio, Spain) designed to detect 128 polymorphisms, including RHD and RHCE variants. The result of BLOODchip Reference v4.0 test was further confirmed with DNA by the Sanger method [12].

Serologically, the sample was classified as RhD negative (Ccee). The reaction pattern of monoclonal anti-D’s is reported in Table 1. Instead, the sample was genotyped with all SSP-PCR kits as RhD positive, Dd and the result obtained with Partial D-TYPE (Dva or Va like or DBS) was consistent with a suggestion of a new variant.

Through the BLOODchip Reference v4.0 analysis, the sample was genotyped as DV/DBS [RHD-CE(S)-D hybrid], associated with a predicted partial D phenotype for the RhD group, due to the lack of a hybridization signal for RhD exon 5. Thereafter, the sequencing of RhD exon 5 with an alternative primer pair was performed, to check its presence/absence and its sequence. In addition, the sequencing of RHD gene exons 1–4 and 6–10 was also
carried out to search for any additional mutation not included in the BLOODchip Reference v4.0, and which could explain the serology results. The PCR analysis of the exon 5 of the RHD gene gave a positive amplification result and the sequencing revealed a polymorphism in hemizygous or homozygous state in position RHD*IVS5+1t. This nucleotide change is located in the BLOODchip Reference v4.0 RHD exon 5 primer reverse sequence, and it explains the lack of amplification; the rest of exon 5 sequence was consensus RHD. Sequencing of exons 1–4 and 6–10 of RHD gene did not reveal additional mutations (Figure 1).

### DISCUSSION

Considering weak D and partial D with frequencies lower than 1%, even if in the daily practice of most blood transfusion centers the status of Rh(D) antigens are determined by serology with agglutination tests. For the immunogenicity of D antigen and the high rate of immunization of D-negative individuals after the transfusion of D-positive RBCs, the determination of RHD alleles play a significant role in D negative pregnant women carrying D positive fetuses and in blood donors. Pregnant women expressing weak D type alleles or carrying D negative fetuses, which can be detected by RHD genotyping, can avoid RhIg prophylaxis [13]. Instead, RHD genotyping in blood donors is beneficial to transfusion recipients because it can exclude weak D, partial D and DEL donors among apparent D negative blood donors and avoid red cell alloimmunization. It is obvious that determining a donor’s phenotype and genotype is more decisive to obtain correct results than two or more serological tests alone [14].

Herein, we report the molecular background of a novel D mutation and their associated immunohematological features. This novel splice-site mutation alters or completely abolishes the specific sequence where the splicing of RHD gene intron 5 takes place. Therefore, the protein product would be aberrant explaining the RhD negative serology result. This hypothesis is supported by the previous description of another nucleotide change in the same position (RHD*IVS5+1g>a) which is also associated with a D negative phenotype in the Portuguese population [15, 16].

### CONCLUSION

The novel RHD mutation, RHD*IVS5+1t hereby described should affect the splicing site and generate an RhD negative phenotype and explain the serology results. In fact, at first, serological analysis for D appeared to be trustworthy negative and only the use of the genetic protocol was decisive for obtaining correct results. Strategies in blood banks regarding the weak D types and the immunization should be established, to reliably type these individuals as RhD-positive, not prone to anti-D immunization. The application of methods based on molecular biology could lead to a rapid and cost-effective large-scale detection in the near future. This case report highlights the significance of using RHD genotyping to confirm and/or clarify D antigen uncertainties in order to prevent the unnecessary immunization of patients.

### REFERENCES


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