Application of Blood Group Genotyping by Next-Generation Sequencing in Various Immunohaematology Cases

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Abstract

\textbf{Background:} Next-generation sequencing (NGS) technology has been recently introduced into blood group genotyping; however, there are few studies using NGS-based blood group genotyping in real-world clinical settings. In this study, we applied NGS-based blood group genotyping into various immunohaematology cases encountered in routine clinical practice.

\textbf{Methods:} This study included 4 immunohaematology cases: ABO subgroup, ABO chimerism, antibody to a high-frequency antigen (HFA), and anti-CD47 interference. We designed a hybridization capture-based NGS panel targeting 39 blood group-related genes and applied it to the 4 cases.

\textbf{Results:} NGS analysis revealed a novel intronic variant (NM_020469.3:c.29-10T>G) in a patient with an \textit{A} \textit{el} phenotype and detected a small fraction of ABO*\textit{A1.02} (approximately 3–6%) coexisting with the major genotype ABO*\textit{B.01/S.01.02} in dizygotic twins. In addition, NGS analysis found a homozygous stop-gain variant (NM_004827.3:c.376C>T, p.Gln126*; ABCG2*\textit{01N.01}) in a patient with an antibody to an HFA; consequently, this patient’s phenotype was predicted as Jr(a−). Lastly, blood group phenotypes predicted by NGS were concordant with those determined by serology in 2 patients treated with anti-CD47 drugs.

\textbf{Conclusion:} NGS-based blood group genotyping can be used for identifying ABO subgroup alleles, low levels of blood group chimerism, and antibodies to HFAs. Furthermore, it can be applied to extended blood group antigen matching for patients treated with anti-CD47 drugs.

Introduction

For over a century, serological testing based on haemagglutination has been the gold standard for determining blood group phenotypes. However, there are immunohaematology problems that serological testing alone may not be able to resolve, such as ABO discrepancies. With advances in molecular methods and growing knowledge on the molecular basis of blood group antigens, blood group genotyping has emerged as a powerful tool to address such immunohaematology problems. The majority of blood group antigens result from single nucleotide variants (SNVs), rendering genotyping assay design and data interpretation straightforward [1]. This has resulted in the widespread application of SNV-based mo-
lular methods, including PCR-restriction fragment length polymorphism, PCR sequence-specific primers, high-resolution melting analysis, matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF MS), and DNA microarray, in blood group genotyping [2–5]. In particular, MALDI-TOF MS and DNA microarray, with a high-throughput capacity, provide a robust platform for large-scale blood group genotyping [2–5]. However, an inherent limitation of SNV-based molecular methods is that they can only test a limited number of SNVs; thus, comprehensive analysis of all blood group genes is not feasible [1]. To overcome this limitation, next-generation sequencing (NGS) technology has been recently introduced into blood group genotyping.

NGS-based blood group genotyping can be applied to various immunohaematology problems encountered in clinical practice. First, NGS-based blood group genotyping can be an efficient tool for identifying ABO subgroup alleles, as the complete ABO gene can be covered in a single assay. NGS-based blood group genotyping can also be used to detect blood group chimerism. The sensitivity of NGS for detecting chimerism is equivalent to that of real-time PCR and digital PCR (0.01–1%) and is superior to that of fragment analysis (>1%) [6]. Furthermore, NGS-based blood group genotyping can be used to predict rare blood group phenotypes in patients with antibodies to HFAs. SNV-based assays typically target only a handful of null alleles [7–11] whereas NGS-based assays have the potential to identify all null alleles, allowing accurate prediction of rare blood group phenotypes [12]. Lastly, NGS-based blood group genotyping can be utilized to resolve anti-CD47 interference in pretransfusion compatibility testing. Compared to SNV-based assays, NGS-based assays enable the provision of more extensively matched RBC units for patients treated with anti-CD47 drugs, which could reduce the risk of alloimmunization and haemolytic transfusion reactions.

Over the past several years, many groups have designed NGS-based blood group genotyping assays and addressed the technical issues encountered while validating these assays [13–22]. However, there are few studies using NGS-based blood group genotyping in real-world clinical settings. In this study, we applied NGS-based blood group genotyping into various immunohaematology cases encountered in routine clinical practice.

Materials and Methods

Study Subjects and DNA Extraction

This study included 4 immunohaematology cases at the blood bank of Samsung Medical Center, South Korea, between December 2018 and April 2020. These 4 cases were as follows: (1) ABO subgroup, (2) ABO chimerism, (3) antibody to a high-frequency antigen (HFA), and (4) anti-CD47 interference. In total, 8 Korean subjects participated in this study (ABO subgroup, 1 patient; ABO chimerism, dizygotic twins and their parents; antibody to an HFA, 1 pregnant woman; anti-CD47 interference, 2 patients). Genomic DNA was isolated from whole blood using a solution-based DNA extraction kit (Wizard Genomic DNA Purification Kit; Promega, Madison, WI, USA).

Serological Testing

ABO forward typing was performed using the tube method with anti-A, anti-B (Shin yang Diagnostics, Siheung, South Korea), anti-A, B (Ortho Clinical Diagnostics, Raritan, NJ, USA), and anti-H (Lorne Laboratories, Reading, UK) reagents. ABO reverse typing was conducted using the tube method with A₂ and B cells (Ortho Clinical Diagnostics). ABO typing was also done using the gel method (DiaClon ABO/D + Reverse Grouping; BIO-RAD, Cressier, Switzerland). Adsorption and elution tests were performed to confirm the presence of A or B antigens on RBCs. Antibody screening was carried out using the tube method with a two-cell panel (Selectogen I-II; Ortho-Clinical Diagnostics) at immediate spin, 37°C, and albumin-indirect antiglobulin test (IAT) phases. Antibody screening was also conducted using gel IAT (ID-Card Liss/Coombs; Bio-Rad) with a two-cell panel (ID-DiaCell I-II; Bio-Rad). In case of a positive screening result, antibody identification was performed using gel IAT with untreated and papain-treated 11-cell panels (Bio-Rad), and direct antiglobulin test (DAT) was performed using the gel method with monospecific anti-IgG and anti-C3d (DC-Screening II; Bio-Rad). Prior to initiating anti-CD47 therapy, extended blood group antigen typing was performed using the tube method with anti-C, -E, -c, -e, -K, -k, -Jk⁺, -Jk⁻, -Fy⁺, -Fy⁻, -M, -N, -S, and -s reagents (Diagast, Loos, France). Patients with anti-CD47 interference in pretransfusion compatibility testing received extended antigen-matched RBC units guided by pretreatment serology (matched for C, E, c, e, K, Jk⁺, Jk⁻, Fy⁺, Fy⁻, S, and s antigens). The strength of agglutination was graded as 0, 1+, 2+, 3+, or 4+.

Flow Cytometry Analysis

Samples exhibiting mixed-field agglutination in ABO forward typing were analyzed using flow cytometry (FC). After three washes with phosphate-buffered saline (PBS), approximately 500,000 RBCs were added to each well of a 96-well plate containing 50 μL of PBS. The RBCs were then fixed with 0.1% glutaraldehyde for 10 min at room temperature. After fixation, the plate was centrifuged at 300 g for 1 min, and the supernatant was discarded. To each well, 5 μL of anti-A or anti-B reagent (murine monoclonal IgM antibodies; Shin yang Diagnostics) was added, incubated for 15 min at room temperature, and washed twice with PBS. Then, 2 μL of phycoerythrin-conjugated anti-mouse IgM secondary antibodies (BD Biosciences, San Jose, CA, USA) were added, incubated for 15 min at room temperature, and washed twice with PBS. Finally, the RBCs were resuspended in 200 μL of PBS and analyzed on a flow cytometer (BD FACSVerse; BD Biosciences).

Sanger Sequencing of ABO Exons 6 and 7

Samples showing an ABO discrepancy in routine serology were subjected to Sanger sequencing of ABO exons 6 and 7 as described in our previous study [23]. Briefly, PCR amplification was carried out using the forward primer ABOe6F (5'-GCTGAGTGCG- GTTTCCAGGT-3') and the reverse primer ABOe7R (5'-AACAG-GACGGGACAAAAAGAAA-3'). The amplified products (2,080 bp) were sequenced using the internal primers ABOe6R (5'-CCA-CCCCAACCTGTCTTGGAA-3'), ABOe7F (5'-CTCTGTGCTCTT-AAGGCTTCC-3'), ABOe7SFI (5'-CTCTGAGGAGGAGATGAT- TAC-3'), and ABOe7SFI (5'-ACGGAGGAGGACCACCTGAG-3').
Confirmation of a Novel ABO Allele

When a novel intronic variant (NM_020469.3:c.29-10T>G) was found in a patient with a weak A phenotype, allele-specific PCR based on the c.99–288C>T polymorphism (rs687621; C: A allele, T: 0 allele) was performed using the primer pair described by Matzhold et al. [24]. Briefly, PCR amplification was conducted using the forward primer ABO_{In1}12.797_{Amp}_F(5'-GATCTGG-GACTGGTGGTTGGAG-3') and the reverse primer ABO_{In2}437C/T_{R}(5'-GCCCAACAGTGCTTGGTGGTGA-3'). To determine whether the c.29-10T > G variant was located on the A allele, the PCR products (655 bp) were sequenced using the same primer pair. Furthermore, three major regulatory regions of the ABO gene, the proximal promoter, the CCAAT-binding factor NF-Y (CBF/NF-Y) enhancer region, and the +5.8-kb site, were sequenced to exclude the possibility that a variant other than c.29-10T>G is responsible for the patient’s weak A phenotype. The proximal promoter of the ABO gene was amplified and sequenced using the primer pair primerno12F(5’-GGGCGGCTCCCTTCCCTACC-3’) and mo-12R(5’-CTCGGGTGAGCGGCCTTCCCTC-3’) as described previously [25]. The CBF/NF-Y enhancer region of the ABO gene was amplified and sequenced using the forward primer ABO_{In1}3043F (5’-ATGGCCACTCTAGGGTGGAC-3’) and the reverse primer ABO_{In1}3154R (5’-GAACTCCTCGTGTGGGTGAAGA-3’) as described previously [26]. To identify two types of deletions involving the +5.8-kb site of ABO intron 1, a 5.8-kb deletion (c.28 + 5110_10889del) and a 3.0-kb deletion (c.28 + 4077_7107del), PCR amplification was performed using the following primer pairs described by Sano et al. [27]: ABO+44195 (5’-TGGATGTTCTCTTCTTCTTCTTAGT-CC-3’) and ABO+11078AS (5’-GGTCTCTATCGCACCTGA-CAA-3’) for the 5.8-kb deletion; ABO+3043S (5’-GGAACTCCTCGTGTGGGTGAAGA-3’) and ABO+11078AS for the 3.0-kb deletion. To identify sequence variants in the +5.8-kb site, the PCR products were sequenced using the primer ABO+743S (5’-GAGCATTATCGTGAAGACATG-3’) described by Takahashi et al. [28].

NGS-Based Blood Group Genotyping

A blood group panel was designed to include 39 blood group-related genes (37 genes belonging to 34 blood group systems and 2 genes encoding an erythroid-specific transcription factor; online suppl. Table S1; for all online suppl. material, see www.karger.com/doi/10.1159/000517565). Target enrichment was performed using custom-designed capture probes (Celomics, Seoul, South Korea). The captured libraries were sequenced on an Illumina platform (Illumina, San Diego, CA, USA) with a 2 × 150-bp paired-end module. Sequence reads were aligned to the hg19 reference genome using BWA-MEM (version 0.7.10), and duplicated reads were removed using Picard (version 1.138). The Genome Analysis Toolkit (GATK, version 3.5) was used for local realignment and base quality score recalibration. Variants were called using the GATK HaplotypeCaller, followed by hard filtering using the following criteria: SNVs: QualByDepth (QD) < 20, FisherStrand (FS) > 60.0, RMSMappingQuality (MQ) < 40.0, MappingQualityRankSumTest (MQRankSum) < -12.5, or ReadPosRankSumTest (ReadPosRankSum) < -8.0; insertions and deletions: QD < 2.0, FS > 150.0, or ReadPosRankSum < -20.0. Variants that passed the hard filtering were further filtered using the following criteria: depth of coverage < 10x or variant allele fraction (VAF) < 20% (exceptionally, in case of chimerism, the VAF threshold was set at 1%). Variants passing through the filtering step were annotated using Variant Effect Predictor (version 94) and dbNSFP (version 4.0). After the annotated variant call format files were examined carefully, the blood group alleles of each individual were determined based on the International Society of Blood Transfusion allele database. Mapped reads were visualized using Integrative Genomics Viewer (IGV).

Results

Statistics of NGS Data

An average of 1.2 million reads were generated per sample, 42.0% of which were uniquely mapped to the target region (70 kb), resulting in a mean depth of coverage of 671×. On average, 96.1, 95.2, and 94.1% of the target region were covered at ≥5×, 30×, and 100×, respectively. The coverage information for each target gene is shown in online supplementary Table S1. In all but three genes (RHD, RHCE, and CR1), more than 97% of the target bases were covered at ≥30×. The poorly covered regions of RHD, RHCE, and CR1 (<30×) were as follows: RHD, exon 8; RHCE, exon 2 (in all samples with a CC phenotype); CR1, exons 7–13 and exons 18 and 19.

Case 1: ABO Subgroup

A blood sample from a 59-year-old male patient (patient 1) was tested for the ABO type. The patient had no recent history of blood transfusion and no history of malignancy (including lymphoma and leukaemia). Table 1 summarizes the results of serological and molecular testing, including NGS. The patient’s RBCs showed no agglutination with anti-A, anti-B, and anti-A,B reagents; however, the eluate displayed 2+ agglutination with A_1 cells in the adsorption and elution tests. Based on these serological results, the patient was considered to have an A_1 phenotype. Sanger sequencing of ABO exons 6 and 7 identified two heterozygous variants (NM_020469.3:c.261delG and c.467C>T); according to the sequencing results, the patient was presumed to carry the ABO^A1.02/O.01.01 genotype. Using our NGS assay covering all exons and flanking intronic regions of the ABO gene, we additionally identified a novel heterozygous variant in ABO intron 1.
1 (c.29-10T>G). Allele-specific PCR and Sanger sequencing of the PCR products confirmed that this intronic variant was located on the ABO*A1.02 allele. No variant was detected in the proximal promoter, CBF/NF-Y enhancer region, and +5.8-kb site of the ABO gene. The c.29-10T>G variant is absent in public databases, including dbSNP, 1000 Genomes, and gnomAD, and has not been reported in the literature previously.

**Case 2: ABO Chimerism**

Blood samples from 10-month-old dizygotic twins (patients 2 and 3) and their parents were tested for the ABO type. The patients were born from a monochorionic diamniotic twin pregnancy and had no recent history of blood transfusion. The results of serological testing, FC analysis, and NGS analysis are summarized in the pedigree (Fig. 1). In the gel and tube methods, the patients’ RBCs exhibited mixed-field agglutination with anti-A and anti-B reagents. In the gel method, agglutination with anti-B was stronger than that with anti-A (online suppl. Fig. S1). In the gel and tube methods, the patient plasma showed no agglutination with A₁ and B cells. FC analysis revealed two distinct RBC populations, group B RBCs (major population) and group A RBCs (minor population; online suppl. Fig. S1). The patients’ father and mother were group A and group AB, respectively. Sanger sequencing of ABO exons 6 and 7 was performed for the patients and their parents, and the resulting genotypes were as follows: patients 2 and 3: ABO*A1.02/O.01.02; father: ABO*A1.02/O.01.02; mother: ABO*A1.02/B.01. Despite a thorough review of the chromatograms obtained from Sanger sequencing, we failed to detect any peaks corresponding to c.297A and c.467C>T (present on ABO*A1.02 but absent on ABO*B.01 and ABO*O.01.02) in the twins. However, the two variants were successfully detected using our NGS assay (VAF: c.297A, 5.8% for patient 2 and 3.8% for patient 3; c.467C>T, 4.8% for patient 2, and 3.2% for patient 3) and were confirmed by visual inspection using IGV (Fig. 2). In addition, manual phasing in IGV revealed that c.297A was in cis with c.261 G belonging to ABO*A1.02 (Fig. 2). Taken together, NGS analysis identified a small fraction of ABO*A1.02 coexisting with ABO*B.01 and O.01.02.

**Case 3: Antibody to an HFA**

A blood sample was collected from a pregnant woman at 29 weeks of gestation (patient 4) prior to an emergency caesarean section and submitted to our blood bank for type and screen. The patient had no previous history of blood transfusion. The patient’s ABO and RhD types were group A, D-positive, and her plasma reacted with all reagent RBCs in tube IAT (1+) and gel IAT (2+). Her plasma also reacted with all the reagent RBCs treated with papain (3+). DAT was negative for IgG and C3d. NGS analysis identified a homozygous stop-gain variant in ABCG2 exon 4 (NM_004827.3:c.376C>T, p.Gln126*; rs72552713; Fig. 3). As this variant was detected in the homozygous form, the patient’s genotype was determined to be ABCG2*01N.01/01N.01. Anti-Jr₄ human serum (from 1 group AB donor) and Jr(a−) RBCs (from 1 group O donor) from the SCARF exchange program were used for confirmatory testing. The patient’s RBCs showed no agglutination with human anti-Jr₄, and the patient’s plasma exhibited no agglutination with Jr(a−) RBCs. Consequently, the patient’s phenotype and antibody specificity were confirmed as Jr(a−) and anti-Jr₄, respectively.

**Case 4: Anti-CD47 Interference**

Our blood bank was informed that 2 patients with diffuse large B-cell lymphoma (patients 5 and 6) were scheduled to begin taking ALX148, a novel anti-CD47 drug. These 2 patients enrolled in the phase 1 clinical trial of ALX148 (AT148001) were assigned to receive ALX148 at a dose of 10 mg/kg weekly. None of the patients had a recent history of blood transfusion. Prior to commencing ALX148 therapy, the patients’ type and screen results were group A, D-positive with a negative antibody screen. The patients’ extended blood group phenotypes determined by serology were as follows: M+N+S−s+s; C+c−E−
In the initial analysis, the serologically determined blood group phenotypes were concordant with the blood group phenotypes predicted from NGS data, except for the MN phenotype (serology: M+N+; NGS: M−N+). In further analysis using IGV, a few reads containing all three variants on the GYPA*01 (c.59C, c.71G, and c.72T) were identified (Fig. 4). Accordingly, the patients’ MN phenotypes were correctly predicted as M+N+ using NGS (Table 2). After an initial ALX148 infusion, the patients’ plasma exhibited panreactivity in both gel IAT (3+) and tube IAT (2+). DAT was positive for IgG (4+), but negative for C3d. During the course of ALX148 therapy, each patient received 2 units of extended antigen-matched RBCs (negative for E, c, K, Fy\(^b\), and S). Haemolytic transfusion reaction was not reported in any of the patients.
Discussion

In all samples included in this study, most of the target region was covered at a high depth, allowing reliable detection of sequence variants in the region. On the contrary, a significant proportion of the target bases in three genes (RHD, RHCE, and CR1) were rarely covered; thus, we could not rule out the possibility that sequence variants may be present in the underrepresented region. The RHD and RHCE genes have high nucleotide sequence homology (approx. 92%) [29]. In particular, the nucleotide sequence of RHCE exon 2 in the RHCE*C allele is identical to that of RHD exon 2, and the nucleotide sequences of exon 8 in RHD and RHCE are identical [15]. Because of this sequence homology, RHCE exon 2 (in all samples with a CC phenotype) and RHD exon 8 (in all samples regardless of their RhCE phenotype) were rarely covered, which is similar to the findings of previous studies [15, 16, 21]. The CR1 gene encodes a protein carrying 30 short consensus repeats, 28 of which are arranged in four long homologous repeats [30]. Because of these repetitive sequences, a part of the CR1 gene was poorly covered by our NGS assay; however, the CR1 exons 26 and 29 where all Knops blood group polymorphisms are located were adequately covered (mean depth of coverage: 252× for exon 26 and 425× for exon 29), allowing for the reliable prediction of Knops blood group phenotypes. An inherent limitation of short-read sequencing is that false-positive or false-negative variant calls can arise from the misalignment of reads to homologous regions in the genome [31]. Such a limitation can be overcome by short-read sequencing based on PCR enrichment, discriminating between homologous regions or long-read sequencing such as nanopore sequencing. A short-read sequencing method based on long-range PCR enrichment, specifically amplifying the RHD gene, has already been described [18, 32].

The vast majority of ABO subgroup-causing variants reported to date are located within ABO exons 6 and 7. Therefore, when ABO subgroups are suspected in routine serology, the initial molecular target is limited to ABO exons 6 and 7 in many blood banks, including our own. However, as ABO subgroup-causing variants can occur in regions other than ABO exons 6 and 7 (including exons 1–5 and regulatory regions of the ABO gene), though rarely [25–28, 33–35], it is necessary to develop molecular assays to screen all these regions. Wu et al. [19] recently designed a hybridization capture-based NGS assay covering the entire ABO gene and identified various ABO subgroup alleles, including a 5.9-kb deletion in ABO intron 1. Similarly, our study attempted to identify an ABO subgroup allele by employing a hybridization capture-based NGS assay targeting 39 blood group-related genes including ABO; as a result, a novel intronic variant (c.29–10T>G in ABO intron 1) was detected in a patient with an A_2 phenotype. The present study demonstrated that a hybridization capture-based NGS assay is useful for identifying ABO subgroup alleles; however, unlike the NGS assay designed by Wu et al. [19], our NGS panel did not include important regulatory regions of the ABO gene, such as the +5.8-kb site, and thus required additional Sanger sequencing to cover these regions. The coverage of the regulatory regions of the ABO gene is particularly important in East Asians, in whom variants in these regions explain a significant proportion of ABO subgroups [34, 35]. Therefore, in future studies, we will add the regulatory regions to our NGS panel.

Table 2. Comparison of blood group phenotypes predicted from NGS data and those determined by serology in 2 patients (patients 5 and 6)

| Blood group (gene) | NGS genotype | phenotype | Serology phenotype |
|--------------------|--------------|-----------|--------------------|
| MNS (GYPA)         | GYPA*02/02 → GYPA*01/02* | M−N+ → M+N+* | M + N+ |
| MNS (GYPB)         | GYPB*04/04 | S−s+ | S−s+ |
| Rh (RHCE)          | RHCE*02/02* | C+c−e−e+ | C+c−E−e+ |
| Kell (KEL)         | KEL*02/02 | K−k+ | K−K+ |
| Duffy (ACKR1)      | FY*01/01 | Fy(a+b−) | Fy(a+b−) |
| Kidd (SLC14A1)     | JK*01/02 | Jk(a+b+) | Jk(a+b+) |

* In both patients, all three variants on the GYPA*01 allele (c.59G, c.71G, and c.72T in GYPA, exon 2) were not called by our pipeline. However, visual inspection using IGV identified a few reads containing all these variants; therefore, we corrected the patients’ genotypes from GYPA*02/02 to GYPA*01/02, which was correlated with their phenotypes determined by serology (M+N+). Both patients were presumed to carry the RHCE*02/02 (RHCE*Cc/Ce) genotype as two of seven variants on the RHCE*02 allele (NM_020485.6:c.48G>C in RHCE, exon 1 and c.676G in RHCE, exon 5) were identified in the homozygous form. However, detection of the remaining five variants on the RHCE*02 allele (NM_020485.6:c.150C>T, c.178C>A, c.201A>G, c.203A>G, and c.307C>T) in RHCE, exon 2) was not possible, as no reads were mapped to RHCE, exon 2.
Intronic variants outside the canonical splice sites may cause aberrant splicing, but the effect of these variants on the splicing process should be verified through functional studies such as minigene splicing assays [36]. Although the c.29-10T>G variant in patient 1 is absent in population databases including gnomAD, it is not certain whether this variant can cause splicing defects such as exon skipping as we did not perform functional studies. In addition to functional studies, detection of transferase activity in the patient’s plasma might be helpful. In this study, the presence of A antigens on the patient’s RBCs was demonstrated by adsorption and elution tests. However, it is important to understand that adsorption and elution tests are prone to false-positive results [37], and therefore further FC analysis may be necessary to confirm adsorption and elution test results.

This study proved the utility of the NGS approach in detecting low levels of blood group chimerism. Our NGS assay accurately detected the minor allele ABO* A1.02 co-existing with the major genotype ABO* B.01/O.01.02 in dizygotic twins (patients 2 and 3). Considering the VAFs of c.297A and c.467C>T unique to ABO* A1.02, the percentage of ABO* A1.02 was estimated to be 3–6%, which is below the detection limit of Sanger sequencing (15–20%). This explains why unlike NGS, Sanger sequencing failed to detect ABO* A1.02 in the twins. Allele-specific PCR selectively amplifying minor alleles in chimeric samples can be utilized to overcome the low sensitivity of Sanger sequencing [38, 39]; however, such procedures are time-consuming and laborious, thus limiting their use in clinical practice. In addition to molecular methods, FC analysis can be used for detecting and quantifying blood group chimerism [40]. In the present study, FC analysis identified a small fraction of group A RBCs in the twins, which was correlated well with the findings of our NGS assay.

One option to determine the antibody specificity in a patient with an antibody to an HFA is to predict the patient’s rare phenotype (HFA-negative phenotype or null phenotype) using molecular methods. In this NGS study, patient 4 was found to carry the ABCG2*01N.01 allele (encoding the M antigen); therefore, the patients’ phenotypes were erroneously predicted to be M–N+ (serology: M+N+). This genotyping error has been reported repeatedly and is presumed to occur due to the misalignment of GYPB exon 2 reads in GYPB*01 (GYPB, GYPB, and GYPE are highly homologous) [13, 16, 21, 22]. For accurate identification of GYPB*01, Fichou et al. [14] developed a short-read sequencing method based on PCR enrichment, specifically amplifying the exons 1 and 2 of GYPB and GYPB [13]. In addition, GYPB*01 could be identified by relaxing variant filtering criteria such as the VAF threshold and minimum depth requirement [16, 22] or by detecting misaligned GYPB exon 2 reads [21]. The present study showed that visual inspection using IGV could be useful for detecting GYPB*01. As reads carrying all three variants unique to GYPB*01 (c.59C, c.71G, and c.72T in GYPB exon 2) are highly likely to originate from GYPB*01, the false-positive rate of our method is presumed to be very low.

**Conclusion**

Our study demonstrated that NGS-based blood group genotyping is an effective tool for identifying ABO subgroup alleles, low levels of blood group chimerism, and antibodies to HFAs. Furthermore, our study proved that NGS-based blood group genotyping can be used to resolve anti-CD47 interference in pretransfusion compatibility testing. Our findings thus suggest that NGS-based blood group genotyping can be applied to various immunohematology cases encountered in routine clinical practice.
Blood Group Genotyping Using NGS

Author Contributions

J.-H.J. and D.C. contributed to the conception and design of the study. H.Y. and M.-T.T.P. performed serological and molecular studies. T.Y.K. and J.-H.J. contributed to data analysis. T.Y.K., H.Y., and D.C. drafted the first version of the manuscript. All authors reviewed and edited the manuscript.

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Statement of Ethics

The study protocol was approved by the Institutional Review Board of Samsung Medical Center, and written informed consent was obtained from all the participants.

Conflict of Interest Statement

All authors declare that there are no conflicts of interest.

Author Contributions

J.-H.J. and D.C. contributed to the conception and design of the study. H.Y. and M.-T.T.P. performed serological and molecular studies. T.Y.K. and J.-H.J. contributed to data analysis. T.Y.K., H.Y., and D.C. drafted the first version of the manuscript. All authors reviewed and edited the manuscript.
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