A comprehensive bioinformatic analysis of 126 patients with an inherited platelet disorder to identify both sequence and copy number genetic variants

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A comprehensive bioinformatic analysis of 126 patients with an inherited platelet disorder to identify both sequence and copy number genetic variants

Ibrahim Almazni | Rachel J. Stapley | Abdullah O. Khan | Neil V. Morgan ©

Abstract
Inherited bleeding disorders (IBDs) comprise an extremely heterogeneous group of diseases that reflect abnormalities of blood vessels, coagulation proteins, and platelets. Previously the UK-GAPP study has used whole-exome sequencing in combination with deep platelet phenotyping to identify pathogenic genetic variants in both known and novel genes in approximately 40% of the patients. To interrogate the remaining “unknown” cohort and improve this detection rate, we employed an IBD-specific gene panel of 119 genes using the Congenica Clinical Interpretation Platform to detect both single-nucleotide variants and copy number variants in 126 patients. In total, 135 different heterozygous variants in genes implicated in bleeding disorders were identified. Of which, 22 were classified pathogenic, 26 likely pathogenic, and the remaining were of uncertain significance. There were marked differences in the number of reported variants in individuals between the four patient groups: platelet count (35), platelet function (43), combined platelet count and function (59), and normal count (17). Additionally, we report three novel copy number variations (CNVs) not previously detected. We show that a combined single-nucleotide variation (SNV)/CNV analysis using the Congenica platform not only improves detection rates for IBDs, suggesting that such an approach can be applied to other genetic disorders where there is a high degree of heterogeneity.

KEYWORDS
CNV, inherited bleeding, platelet disorders, SNV, thrombocytopenia, variant interpretation, whole-exome sequencing

1 INTRODUCTION
Inherited bleeding disorders (IBDs) are a heterogeneous group of diseases that reflect abnormalities in blood vessels, coagulation proteins, and platelets. They often present after birth or during childhood, and clinically manifest with variable bleeding tendencies (Blanchette et al., 1991). Although the majority of IBDs are known to be primarily associated with coagulation factor abnormalities such as hemophilia A and B, rarer disorders of platelet count and function are still poorly understood (Sivapalaratnam et al., 2017). Therefore,
to investigate the molecular mechanisms of this group of disorders, it is often best to address the gene(s) already implicated in these bleeding disorders in the first instance, and then specifically to investigate how the genetic variants can disrupt the gene function (Nurden et al., 2012; Peyvandi et al., 2006). An increasing number of new genes and their variants have been discovered, which are implicated in megakaryocyte differentiation and/or platelet production and function (Johnson, Fletcher, et al., 2016).

The UK Genotyping and Phenotyping of Platelets study (UK-GAPP; https://www.birmingham.ac.uk/research/cardiovascular-sciences/research/platelet-group/platelet-gapp/index.aspx) has recruited over 1000 patients based on a history of suspected bleeding disorders of unknown cause from over 25 collaborating hemophilia care centers across the United Kingdom. Recruited patients underwent a combination of platelet phenotyping and genotyping to determine the likely causative genes attributable to their specific defects (Jones et al., 2012; Watson et al., 2013). Gross hematological analysis and light transmission aggregometry and/or flow cytometry were used to identify thrombocytopenia (low platelet counts), platelet function, and cell signaling defects. Following this, targeted genetic analysis was employed and revealed variants, both novel and known, to be causative of bleeding in patients.

High-throughput sequencing technologies including whole-exome sequencing (WES) and whole-genome sequencing are valuable tools used to uncover novel variants in platelet-specific genes. Over the past 10 years, such techniques have revealed many causative variants, therefore assisting in providing a clear diagnosis for some patients with severe bleeding disorders (Bastida et al., 2018; Daly et al., 2014; Downes et al., 2019; Leinæe et al., 2017). In addition, targeted next-generation sequencing (NGS) panels can be used to highlight platelet-specific genes that have been previously implicated in bleeding disorders. NGS panels can be employed in a clinical diagnostic setting or used for prescreening, filtering out patients with variants in known genes, and subsequently employing WES for those who may harbor variants in novel genes (Johnson et al., 2018; Simeoni et al., 2016). This approach was applied in the UK-GAPP study where patients with known mutations in hemophilia A and B or coagulation mediated genes, known to cause bleeding were eliminated. However, many of these panels do not search for copy number variations (CNVs), and indeed we, and others have not found definitively causative variants in approximately 40%–50% patients despite a strongly indicative inherited component for their bleeding (Bastida et al., 2018; Johnson et al., 2018; Johnson, Lowe, et al., 2016; Leinæe et al., 2017; Lentaigne et al., 2016). In this study, we address this by applying a newly established, comprehensive genetic analysis software that detects both single-nucleotide variations (SNVs) and CNVs. Congenica (https://www.congenica.com) is an automated clinical decision support platform that was used to analyze and rapidly interpret the WES data of 126 patients recruited to the UK-GAPP study. Users are able to prioritize and review genetic variants, as well as assign pathogenicity, after which the software calculates overall pathogenicity based on the American College of Medical Genetics and Genomics (ACMG) guidelines (Richards et al., 2015). It collates all essential information to make an informed and robust decision for the identification of causal genetic variants.

The Congenica platform is primarily applied for genetic diagnostics and is routinely used in clinical laboratories for variant validation and reporting. For the first time, we show its utility in interrogating a large cohort of patients recruited to the UK-GAPP research study. Using this approach, we perform a robust and comprehensive analysis to find both known and novel genetic variants with plausible association with disease, including rare CNVs not previously detected. Combined with extensive patient phenotypic studies, this provides a potent tool for the dissection of the genetic causes of bleeding in a cohort which, thus far, remains genetically unresolved despite an extensive clinical presentation of familial bleeding.

2 | METHODS

2.1 | Hematological evaluation and platelet phenotyping of patients

To initially classify patients as having a platelet defect and determine their platelet defect subtypes, they underwent an initial hematological workup and extensive platelet function testing workflow. These methods can be seen in detail in the Supporting Information Methods section.

2.2 | WES

WES was performed on the genomic DNA of 117 patients in this study as previously described (Johnson, Lowe, et al., 2016). Briefly following enrichment of coding regions and intron/exon boundaries with the SureSelect human AllExon 50 Mb Kit (Agilent Technologies), captured libraries were sequenced on the Illumina HiSeq 2500 (Illumina) with 100-bp paired-end reads.

2.3 | Processing WES data using Congenica software

First, an Interpretation Request (IR) was completed which included information about the proband and any other family members and related clinical data including HPO terms (abnormal bleeding HP:0001892 and/or thrombocytopenia HP:0001873) for affected individuals. Relevant gene panels (Inherited Bleeding Disorder; High Evidence_Green, Medium Evidence Amber and Low Evidence Red, gene lists) containing 119 genes (Table S1) from Panel app (https://panelapp.genomicsengland.co.uk/) were applied in the project and deemed suitable for research purposes. However, of this gene panel only 88 genes from the Genomics England website (R90) are considered as suitable for clinical use at this present time. The WES data (either BAM or FASTQ files) of patients were then transferred to the Congenica SFTP server for processing. The Congenica pipeline could...
then be used for sequence alignment and variant calling of SNVs, small insertion/deletion (indels), CNVs (Figure 1), and coverage (Table S3).

The analytical pipeline for the detection of CNVs in genes involved in the IBDs panel was employed using the ExomeDepth coverage approach. The exome read depth of the target patient’s sample was compared against the read depth of a reference panel (up to 10 WES samples of each gender) to detect regions with different coverage which could represent a CNV event.

Using the Congenica software, the lower limit that the ExomeDepth calling software uses for CNV calling is ≥20 sequence reads. This ensures that ExomeDepth does not consider low quality reads when comparing the reference samples to the target patient.

3 | RESULTS

3.1 | Platelet phenotyping

Recruited patients were subjected to an initial hematological analysis and extended deep platelet phenotyping using the previously published workflow (Johnson, Lowe, et al., 2016). Phenotyping outcomes can be seen in detail in the Supporting Information Results section (Figures S1 and S2; Table S2).

3.2 | Validation of WES analysis with known variants

Validation of the WES analysis in the GAPP study was performed using Congenica software. Five different known genetic variants were identified previously by WES in nine patients with a suspected IBD (Fletcher et al., 2015; Johnson, Lowe, et al., 2016; Table 1). We used two trios (one parent and two affected children) and three unrelated individuals, all with known or likely pathogenic variants in platelet- or megakaryocyte-related genes. This analysis was conducted in a blind manner to assess the reliability and robustness of the software in correctly highlighting all known genetic variants in these patients. Using panels of genes implicated in IBDs, the first trio (Family A) including exomes of patients 1–3 were found to share the same splicing sequence variant in RUNX1: c.98-1G>A. The second trio (Family B) including exomes of patients 4–6 all shared a variant c.503G>T p.(Cys168Phe) in GFI1B. Patient 7 displayed a homozygous missense mutation c.1246G>A p.(Gly416Arg) in the GNE gene.

**FIGURE 1** Congenica pipeline overview for processing of whole-exome sequencing (WES) data. Adapted from https://www.congenica.com/. The informatic strategy shown is used to incorporate both single-nucleotide variation (SNV) and copy number variation (CNV) analysis. The raw WES data are inputted as either FASTQ, or BAM files followed by alignment to the reference genome. SNV calling is then performed to generate VCF files and subsequent in silico tools to determine the pathogenicity of variants. Simultaneously, CNV analysis is performed using a predefined reference and sex-matched WES panel and fed into ExomeDepth for CNV calling in the WES samples.
patient 8, a missense variant c.659T>A p.(Val220Asp) in SLFN14 and finally in patient 9, a stop gain mutation c.1611C>A p.(Cys537Ter) located within THBD was identified. All known variants found in the patients were successfully verified by Congenica software against our previously analyzed WES data (Table 1).

### 3.3 WES analysis to identify new SNVs and CNVs using the Congenica platform

WES data of all 117 patients were analyzed by the Congenica platform based on the ExomeDepth coverage following the phenotyping and platelet function studies workflow. The Congenica pipeline was used for exome sequence alignment and variant calling of SNVs, indels, and CNVs to determine plausible candidate variants from the WES data. An average read depth sequencing coverage of 205 was observed at the site of each variation across all DNA samples analyzed by WES for SNVs (Table S3). The ExomeDepth integrated tool was used to determine CNV based on read coverage (Table 3). First, WES data of the 117 patients were analyzed by filtering using an IBDs gene panel (Table S1). Variants were then filtered within the software based on the exclusion criteria initially stated in the GAPP study. A rare variant cut-off or minor allele frequency (MAF) of <0.01 in each data set was used and synonymous and intron variants ±5 base pairs away from the exon-intron boundaries were excluded. Non-shared variants between the same affected family members were also eliminated.

Following exclusion of variants based on these criteria, a range of between two and six variants (SNVs, small indels, and splice site) were noted per patient (Table 2). In silico pathogenicity prediction tools that have been integrated into the Congenica software were employed for further analysis (Table 2). A total of 135 variants in genes implicated in bleeding disorders were identified across all the 117 patients and all variants were observed in a heterozygous state (Table 2). In total, 22 variants were classified as pathogenic and 26 were likely pathogenic when considering the ACMG consensus guidelines. The remaining 87 variants were classified as of uncertain significance. The graphical illustration of this summary is shown in Figure 2a. There was a marked difference in the number of reported variants between the four classes of variants in patient groups: platelet count (35); platelet function (43); combined platelet count and function (59); and normal count (17) (Figure 2b).

### 3.4 Candidate variants identified in patient’s cohort

A total of 48/135 (35.5%) variants with MAF of 0, unless otherwise stated, were identified across the 117 patients (Table 2). In total, 14/48 (29.1%) variants have been published previously. The number of variants found to be shared in the same affected family members were 21. Plausible candidate variants were present within the following genes (RUNXI, SLFN14, FLJ1, ETV6, HPS3, F10, P2RY12, SMAD4, TUBB1, G1B1A, GBA, CYCS, VWF, THBD, LYST, ADAMTS13, GFI1B, ITGA2B, NBEAL2, MECOM, and MYH9; Table 2). Two rare variants were noted between five related affected family members including RUNXI: c.611G>A p.(Arg204Gln) in patients 38.1 and 38.2; WAS: c.1456G>A p.(Glu486Lys) in patients 38.3, 38.4, and 38.5; and a stop gain variant within ADAMTS13: c.1315G>T p.(Glu439Ter) was shared between two related affected individuals 53.1 and 53.2. Two related affected individuals with macrothrombocytopenia shared a variant within the newly discovered gene (involved in platelet disorders) MECOM: c.951G>T p.(Lys317Asn). A novel stop gained variant within ETV6: c.1288C>T p.(Arg430Ter) was noted in patient 7, which was subsequently classified as pathogenic.

### 3.5 CNVs found in the patient cohort

Overall, the CNV analysis using the integrated ExomeDepth tool revealed an average of four CNVs per exome (n = 15; Table 3). There were three rare structural variants covering large regions on chromosomes 11 and 17 and encompassing numerous genes, including

| Patient | Gene | Variation | Type |
|---------|------|-----------|------|
| (Family A) 1 | RUNXI (Zhang et al., 2018) | c.98-1G>A | Splice acceptor |
| (Family A) 2 | RUNXI | c.98-1G>A | Splice acceptor |
| (Family B) 4 | GFI1B (Rabbolini et al., 2017) | c.503G>T p.(Cys168Phe) | Missense |
| (Family B) 5 | GFI1B | c.503G>T p.(Cys168Phe) | Missense |
| (Family B) 6 | GFI1B | c.503G>T p.(Cys168Phe) | Missense |
| 7 | GNE | c.1246G>A p.(Gly416Arg) | Missense |
| 8 | SLFN14 (Johnson, Lowe, et al., 2016) | c.659T>A p.(Val220Asp) | Missense |
| 9 | THBD (Rabbolini et al., 2017) | c.1611C>A p.(Cys537Ter) | Stop gain |

Note: NCBI reference sequences: RUNXI (NM_001001890.3); GFI1B (NM_001371908.1); GNE (NM_000157.4); SLFN14 (NM_001129820.1); THBD (NM_000361.2).
| Patient | Gene(s) | VEP | MAX AF | Genomic variation | Protein effect | PolyPhen | SIFT | Pathogenicity |
|---------|---------|-----|--------|------------------|---------------|----------|------|---------------|
| 1.1     | NVD     |     |        |                  |               |          |      |               |
| 1.2     | NVD     |     |        |                  |               |          |      |               |
| 1.3     | NVD     |     |        |                  |               |          |      |               |
| 1.4     | NVD     |     |        |                  |               |          |      |               |
| 2.1     | RUNX1 [Stockley et al, 2013] | Splice donor | 0 | c.508+1G>T | p.? | NA | NA | Pathogenic |
| 2.2     | RUNX1 [Stockley et al, 2013] | Splice donor | 0 | c.508+1G>T | p.? | NA | NA | Pathogenic |
| 3.1     | SLFN14 [Johnson, Lowe, et al, 2016] | Missense | 0 | c.659T>A | p.(Val20Asp) | Possibly_damaging | Deleterious | Pathogenic |
| 3.2     | SLFN14 [Johnson, Lowe, et al, 2016] | Missense | 0 | c.659T>A | p.(Val20Asp) | Possibly_damaging | Deleterious | Pathogenic |
| 4.1     | FLI1 [Johnson, Lowe, et al, 2016] | Frameshift | 0 | c.992_995del | p.(Asn331ThrfsTer4) | NA | NA | Pathogenic |
| 4.2     | FLI1 [Johnson, Lowe, et al, 2016] | Frameshift | 0 | c.992_995del | p.(Asn331ThrfsTer4) | NA | NA | Pathogenic |
| 5       | FGA     | Missense | 0.00951 | c.1366A>G | p.(Thr456Ala) | Possibly_damaging | Deleterious | Uncertain significance |
| 6.1     | NVD     |     |        |                  |               |          |      |               |
| 6.2     | MPK6B   | Missense | 0 | c.132G>C | p.(Trp44Cys) | Probably_damaging | Deleterious | Uncertain significance |
| VPS33B  | Missense | <0.0001 | c.434T>C | p.(Leu145Ser) | Probably_damaging | Deleterious | Uncertain significance |
| 7       | ETV6    | Stop gained | 0 | c.1288C>T | p.(Arg430Ter) | NA | NA | Pathogenic |
| 8       | VWF     | Frame shift | <0.0001 | c.2516del | p.(Gly839GlufsTer4) | NA | NA | Pathogenic |
| ANKRD26 | Missense | 0.00324 | c.3004G>A | p.(Glu1002Lys) | Possibly_damaging | NA | NA | Uncertain significance |
| 9       | SLC45A2 | Missense | <0.0001 | c.1471G>A | p.(Gly491Arg) | Probably_damaging | Deleterious | Uncertain significance |
| 10.1    | HPS3    | Missense | 0 | c.479G>A | p.(Ser160Asn) | Benign | Tolerated | Uncertain significance |
| 10.2    | HPS3    | Missense | 0 | c.479G>A | p.(Ser160Asn) | Benign | Tolerated | Uncertain significance |
| 11      | LYST    | Missense | 0.0005 | c.8960C>G | p.(Asp2987Arg) | Possibly_damaging | Deleterious | Uncertain significance |
| AP3D1   | Missense | 0.00116 | c.1246G>A | p.(Glu416Lys) | Possibly_damaging | Deleterious | Uncertain significance |
| 12      | COLS2A2 | Missense | 0.00264 | c.4067A>G | p.(Asp1356Gly) | Benign | Deleterious | Uncertain significance |
| 13      | F7      | Missense | <0.0001 | c.857C>T | p.(Ala286Val) | Benign | Deleterious | Uncertain significance |
| 14.1    | FI0     | Missense | 0 | c.1325G>A | p.(Gly442Asp) | Probably_damaging | Deleterious | Likely pathogenic |
| 14.2    | NBEAL2  | Missense | 0.0066 | c.6886G>A | p.(Arg2289Gln) | Possibly_damaging | Deleterious | Uncertain significance |
| G3A      | Missense | 0.00363 | c.1226A>G | p.(Asn409Ser) | Benign | Deleterious | Uncertain significance |
| F10     | Missense | 0 | c.1325G>A | p.(Gly442Asp) | Probably_damaging | Deleterious | Likely pathogenic |
| NBEAL2  | Missense | 0.0066 | c.6886G>A | p.(Arg2289Gln) | Possibly_damaging | Deleterious | Uncertain significance |
| Patient | Gene(s) | VEP | Genomic variation | Protein effect | PolyPhen | SIFT | Pathogenicity |
|---------|---------|-----|------------------|----------------|----------|------|--------------|
| GBA     | Missense| 0.00363 | c.1226A>G | p.(Asn409Ser) | Benign | Deleterious | Uncertain significance |
| 15.1 VWF | Missense | 0.00558 | c.1261G>A | p.(Arg544Gln) | Possibly_damaging | Deleterious | Likely pathogenic |
| 15.2 VWF | 5’-UTR | 0 | c.161C>T | p.? | NA | NA | Uncertain significance |
| NBEA    | Missense | 0.00558 | c.161G>A | p.? | NA | NA | Uncertain significance |
| 16 ACRVL1 | Missense | <0.0001 | c.653G>A | p.(Arg218Gln) | Benign | Deleterious | Likely pathogenic |
| RUNX1   | Stop gained | 0 | c.317G>A | p.(Trp106Ter) | NA | NA | Pathogenic |
| ITGB3   | Missense | <0.0001 | c.349C>T | p.(Arg117Trp) | Possibly_damaging | Deleterious | Likely pathogenic |
| 17 RUNX1 | Splice donor | 0 | c.351+1G>T | p.(Glu135Ter) | NA | NA | Pathogenic |
| F11     | Stop gained | 0 | c.1246G>T | p.(Arg383Ser) | Probably_damaging | Deleterious | Uncertain significance |
| SERPINC1 | Missense | 0.00276 | c.1469G>A | p.(Pro258Thr) | Probably_damaging | Deleterious | Uncertain significance |
| F13A1   | Missense | 0.00192 | c.1149G>T | p.(Ala139Val) | Probably_damaging | Deleterious | Uncertain significance |
| 18 PTPN11 | Missense | <0.0001 | c.922A>G | p.(Asn308Asp) | Benign | Deleterious | Likely pathogenic |
| 19 FGB   | Missense | 0.00674 | c.794C>T | p.(Pro265Leu) | Probably_damaging | Deleterious | Uncertain significance |
| 20 GP6   | Missense | 0.00209 | c.1274G>A | p.(Arg425Asn) | Possibly_damaging | Deleterious | Uncertain significance |
| THBD    | Missense | 0.00528 | c.197T>G | p.(Leu66Arg) | Possibly_damaging | Deleterious | Uncertain significance |
| 21 PLG   | Missense | 0.00297 | c.9017A>G | p.(Arg2663Pro) | Possibly_damaging | Deleterious | Uncertain significance |
| ARPC1B  | Missense | 0.00407 | c.904+3A>G | p.(Arg904Gln) | Probably_damaging | Deleterious | Uncertain significance |
| 22.1 P2RY12 | Missense | 0.00015 | c.365G>A | p.(Arg122His) | Probably_damaging | Deleterious | Likely pathogenic |
| 22.2 P2RY12 | Missense | 0.00015 | c.365G>A | p.(Arg122His) | Probably_damaging | Deleterious | Likely pathogenic |
| 23 P2RY12 | Missense | 0.00027 | c.416C>T | p.(Leu139Val) | Probably_damaging | Deleterious | Pathogenic |
| MCFD2   | Missense | 0.00002 | c.172G>A | p.(Ser117Asn) | Probably_damaging | Deleterious | Uncertain significance |
| 24 VPS33B | Missense | 0.00267 | c.9017A>G | p.(Arg301His) | Probably_damaging | Deleterious | Uncertain significance |
| ITGB3   | Missense | 0.00528 | c.197T>G | p.(Leu66Arg) | Possibly_damaging | Deleterious | Uncertain significance |
| LYST    | Missense | 0.00264 | c.9017A>G | p.(Leu3006Arg) | Probably_damaging | Deleterious | Uncertain significance |
| 25 RUNX1 | Missense | 0.00276 | c.403G>A | p.(Gly135Ser) | Probably_damaging | Deleterious | Likely pathogenic |
| VWF     | Missense | 0.00276 | c.7988G>C | p.(Arg2663Pro) | Possibly_damaging | Deleterious | Likely pathogenic |
| 26 RUNX1 | Missense | 0 | c.593A>T | p.(Asp198Val) | Possibly_damaging | Deleterious | Likely pathogenic |
| SMAD4   | Splice donor | 0 | c.904+1_904+2ins- sGCGTGTCCAAA | p.? | NA | NA | Uncertain significance |
| SMAD4   | Splice donor | 0 | c.904+3A>G | p.? | NA | NA | Uncertain significance |
| GCCX    | Missense | 0 | c.2012G>A | p.(Arg671His) | Benign | Deleterious | Uncertain significance |

(Continues)
| Patient | Gene(s) | VEP | MAX AF | Genomic variation | Protein effect | PolyPhen | SIFT | Pathogenicity |
|---------|---------|-----|--------|-------------------|---------------|----------|------|--------------|
| 27      | TUBB1   | Missense | 0.0053 | c.13G>A, c.440C>T | p.(Val5Ile), p.(Ala147Val) | Probably damaging, Benign | Tolerated, Deleterious | Uncertain significance, Uncertain significance |
|         | TPM4    | Missense | 0.0053 | c.440C>T | p.(Ala147Val) | Benign | Tolerated, Deleterious | Uncertain significance, Uncertain significance |
| 28      | PLAT    | Missense | 0.00163 | c.928C>T | p.(Arg310Cys) | Possibly damaging | Deleterious | Uncertain significance |
| 29.1    | TUBB1    | Missense | 0.0008 | c.721C>T | p.(Arg241Trp) | Probably damaging | Deleterious | Uncertain significance |
| 29.2    | TUBB1    | Missense | 0.0008 | c.721C>T | p.(Arg241Trp) | Probably damaging | Deleterious | Uncertain significance |
| 30      | GP1BA   | Frameshift | 0 | c.1274_1275del, c.1277_1313del, c.653G>A | p.(Glu425AlafsTer72), p.(Pro426ArgfsTer34), p.(Trp218Ter) | NA | NA | Likely pathogenic |
|         | GP1BA   | Frameshift | 0 | c.1274_1275del, c.1277_1313del, c.653G>A | p.(Glu425AlafsTer72), p.(Pro426ArgfsTer34), p.(Trp218Ter) | NA | NA | Uncertain significance |
|         | GBA     | Stop gained | 0 | c.1274_1275del, c.1277_1313del, c.653G>A | p.(Glu425AlafsTer72), p.(Pro426ArgfsTer34), p.(Trp218Ter) | NA | NA | Pathogenic |
| 31      | MPI6G6B | Splice region | 0.00022 | c.621G>T | p.? | NA | NA | Uncertain significance |
|         | HPS6    | Inframe insertion | 0 | c.621G>T | p.? | NA | NA | Uncertain significance |
| 32.1    | CYCS    | Missense | 0.0014 | c.155C>T | p.(Ala52Val) | Benign | Tolerated | Uncertain significance |
|         | GGCX    | Missense | 0.0014 | c.155C>T | p.(Ala52Val) | Benign | Tolerated | Uncertain significance |
| 32.2    | CYCS    | Missense | 0.0014 | c.155C>T | p.(Ala52Val) | Benign | Tolerated | Uncertain significance |
|         | GGCX    | Missense | 0.0014 | c.155C>T | p.(Ala52Val) | Benign | Tolerated | Uncertain significance |
| 33      | COL5A1  | Missense | 0 | c.1715C>A | p.(Pro572His) | NA | NA | Uncertain significance |
|         | TUBB1   | Missense | 0 | c.1715C>A | p.(Pro572His) | NA | NA | Uncertain significance |
| 34      | KLKB1   | Frameshift | 0.00132 | c.772C>T | p.(Leu528Phe) | Benign | NA | Likely pathogenic |
|         | NBEAL2  | Missense | 0.00407 | c.772C>T | p.(Leu528Phe) | Benign | NA | Uncertain significance |
| 35      | COL5A1  | Missense | 0.00027 | c.145C>T | p.(His49Tyr) | Possibly damaging | NA | Uncertain significance |
|         | TBXAS1  | Missense | 0.00162 | c.1523A>T | p.(Glu508Val) | Possibly damaging | NA | Uncertain significance |
|         | THBD    | Missense | 0.00162 | c.405T>G | p.(Leu136Trp) | NA | NA | Uncertain significance |
| 36      | RASGRP2 | Missense | 0.00008 | c.281C>T | p.(Pro94Leu) | Benign | Tolerated | Uncertain significance |
|         | VWF     | Missense | 0.00024 | c.6424C>T | p.(Leu2142Phe) | Possibly damaging | Deleterious | Uncertain significance |
|         | VWF     | Missense | 0.00024 | c.6424C>T | p.(Leu2142Phe) | Possibly damaging | Deleterious | Uncertain significance |
|         | GP1BA   | Missense | 0.00024 | c.6424C>T | p.(Leu2142Phe) | Possibly damaging | Deleterious | Uncertain significance |
| 37.1    | GP1BA   | Missense | 0 | c.413G>T | p.(Gly138Val) | Probably damaging | Deleterious | Likely pathogenic |
| 37.2    | GP1BA   | Missense | 0 | c.413G>T | p.(Gly138Val) | Probably damaging | Deleterious | Likely pathogenic |
| 37.3    | GP1BA   | Missense | 0 | c.3493C>T | p.(Arg1165Cys) | Probably damaging | Deleterious | Pathogenic |
TABLE 2 (Continued)

| Patient | Gene(s) | VEP | MAX AF | Genomic variation | Protein effect | PolyPhen | SIFT | Pathogenicity |
|---------|---------|-----|--------|-------------------|----------------|----------|------|--------------|
| 38.1    | RUNX1   | Missense | 0 | c.611G>A | p.(Arg204Gln) | Possibly damaging | Deleterious | Likely pathogenic |
| 38.2    | RUNX1   | Missense | 0 | c.611G>A | p.(Arg204Gln) | Possibly damaging | Deleterious | Likely pathogenic |
| 38.3    | WAS      | Missense | 0 | c.1456G>A | p.(Glu486Lys) | Possibly damaging | Deleterious | Uncertain significance |
| 38.4    | WAS      | Missense | 0 | c.1456G>A | p.(Glu486Lys) | Possibly damaging | Deleterious | Uncertain significance |
| 38.5    | WAS      | Missense | 0 | c.1456G>A | p.(Glu486Lys) | Possibly damaging | Deleterious | Uncertain significance |

39 NVD

40 NVD

41 ACTN1  | Missense | 0.00139 | c.2662G>C | p.(Gly888Arg) | Probably damaging | Deleterious | Tolerated | Uncertain significance |

42 ABG8   | Missense | 0.00157 | c.1924G>A | p.(Ala642Thr) | Benign | Tolerated | Uncertain significance |

43 RUNX1  | Splice acceptor | 0.001 | c.98-1G>A | NA | Pathogenic | Deleterious | NA | Pathogenic |
| VWF (Lester et al., 2007) | Missense | c.7390C>T | p.(Arg2464Cys) | NA | Pathogenic | NA | NA | Pathogenic |

44.1 RUNX1 (Stockley et al., 2013) | Splice donor | 0 | c.351+1G>T | p.? | NA | Pathogenic |

44.2 RUNX1 (Stockley et al., 2013) | Splice donor | 0 | c.351+1G>T | p.? | NA | Pathogenic |

45 HRG    | Missense | 0.00162 | c.1379G>A | p.(Arg460Gln) | Benign | Tolerated | Likely benign |

46 FLNA   | Missense | 0.0073 | c.5948C>T | p.(Ser1983Leu) | Probably damaging | Deleterious | Likely benign |

47 GGCG   | Missense | 0.0014 | c.1217G>A | p.(Arg406His) | Possibly damaging | Tolerated | Uncertain significance |

48 RUNX1  | Missense | 0 | c.586A>G | p.(Thr196Ala) | Possibly damaging | Deleterious | Likely pathogenic |
| SLFN14  | Missense | 0 | c.2686T>C | p.(Ser896Pro) | Possibly damaging | Deleterious | Uncertain significance |
| SLFN14  | Missense | 0 | c.1481A>G | p.(Gln494Arg) | Benign | Tolerated | Uncertain significance |
| SLFN14  | Frameshift | 0 | c.3_4insCTAGTC-GACTATA | p.(Glu2LeufsTer10) | NA | Pathogenic |

49 ABCG5  | Missense | <0.0001 | c.692T>C | p.(Ile231Thr) | Probably damaging | Deleterious | Uncertain significance |

(Continues)
| Patient | Gene(s) | VEP | MAX AF | Genomic variation | Protein effect | PolyPhen | SIFT | Pathogenicity |
|---------|---------|-----|--------|-------------------|---------------|----------|------|--------------|
| 50.1    | STXB2   |     |        |                   |               |          |      |              |
| 50.2    | NVD     |     |        |                   |               |          |      |              |
| 51.1    | ADAMTS13|     |        |                   |               |          |      |              |
| 51.2    | ADAMTS13|     |        |                   |               |          |      |              |
| 51.3    | ADAMTS13|     |        |                   |               |          |      |              |
| 52.1    | LYST    |     |        |                   |               |          |      |              |
| 52.2    | THBD    | (Dargaud et al., 2015) | |                   |               |          |      |              |
| 53.1    | ADAMTS13|     |        |                   |               |          |      |              |
| 53.2    | NVD     |     |        |                   |               |          |      |              |
| 54.1    | F10     |     |        |                   |               |          |      |              |
| 54.2    | MPL     |     |        |                   |               |          |      |              |
| 55      | ACVR1L  |     |        |                   |               |          |      |              |
|         | ITG4B2  | (Johnson et al., 2018) | |                   |               |          |      |              |
|         | THPD    | (Johnson et al., 2018) | |                   |               |          |      |              |
|         | HOXA11  |     |        |                   |               |          |      |              |
| 56      | GFI1B   | (Johnson, Lowe, et al., 2016) | |                   |               |          |      |              |
| 57      | NBEAL2  |     |        |                   |               |          |      |              |
| 58.1    | NBEAL2  |     |        |                   |               |          |      |              |
| 59      | MECOM   |     |        |                   |               |          |      |              |
| 60      | MPL     |     |        |                   |               |          |      |              |
| 61      | COL5A1  |     |        |                   |               |          |      |              |
| 62      | NVD     |     |        |                   |               |          |      |              |
| 63.1    | GFI1B   | (Rabbolini et al., 2017) | |                   |               |          |      |              |
| 63.2    | NVD     |     |        |                   |               |          |      |              |
| Patient | Gene(s) | VEP  | MAX AF | Genomic variation | Protein effect | PolyPhen        | SIFT       | Pathogenicity       |
|---------|---------|------|--------|-------------------|---------------|----------------|------------|---------------------|
| 63.3    | GFI1B   | Missense | 0.00438 | c.503G>T         | p.(Cys168Phe) | Probably damaging | Deleterious | Likely pathogenic   |
|         | (Rabbolini et al., 2017) |         |        |                   |               |                |            |                     |
| 64      | NVD     |        |        |                   |               |                |            |                     |
| 65.1    | MECOM   | Missense | 0.0001  | c.951G>T         | p.(Lys317Asn) | NA             | NA         | Uncertain significance |
| 65.2    | SLFN14  | Missense | 0.0014  | c.916G>C         | p.(Asp306His) | Benign        | Tolerated  | Uncertain significance |
|         | MECOM   | Missense | 0.0001  | c.951G>T         | p.(Lys317Asn) | NA             | NA         | Uncertain significance |
|         | SLFN14  | Missense | 0.0014  | c.916G>C         | p.(Asp306His) | Benign        | Tolerated  | Uncertain significance |
| 66.1    | MYH9    | Stop gained | 0.0001  | c.5797C>T       | p.(Arg1933Ter) | NA             | NA         | Pathogenic          |
| 66.2    | MYH9    | Stop gained | 0.0001  | c.5797C>T       | p.(Arg1933Ter) | NA             | NA         | Pathogenic          |
|         | (Savoia & Pecci, 2015) |         |        |                   |               |                |            |                     |
| 67      | NVD     |        |        |                   |               |                |            |                     |
| 68      | RASGRP2 | Missense | 0.000102 | c.1159C>T     | p.(Arg387Cys) | Probably damaging | Deleterious | Uncertain significance |
| 69      | F10     | Missense | 0.000547 | c.1406G>A      | p.(Arg469Lys) | Benign        | Tolerated  | Uncertain significance |
| 70.1    | ETV6    | Stop gained | 0      | c.313C>T        | p.(Arg105Ter) | NA             | NA         | Likely pathogenic   |
| 70.2    | NVD     |        |        |                   |               |                |            |                     |
| 71      | RUNX1   | Missense | 0      | c.1256T>G       | p.(Val419Gly) | Benign        | Possibly damaging | Deleterious | Likely pathogenic   |
|         | RUNX1   | Missense | 0      | c.1270T>C       | p.(Ser424Pro) | Benign        | Possibly damaging | Deleterious | Likely pathogenic   |
| 72      | FGG     | Missense | 0.000792 | c.323C>G       | p.(Ala108Gly) | Benign        | Tolerated  | Likely pathogenic   |
|         | STXBP2  | Missense | 0.000539 | c.499C>T       | p.(Arg167Trp) | Benign        | Tolerated  | Uncertain significance |
|         | TUBB1   | Missense | 0.0008  | c.721C>T        | p.(Arg241Trp) | Benign        | Tolerated  | Uncertain significance |
|         | (Johnson, Lowe, et al., 2016) |         |        |                   |               |                |            |                     |
| 73      | NVD     |        |        |                   |               |                |            |                     |
| 74      | RUNX1   | Missense | 0      | c.1265A>C       | p.(Glu422Ala) | Benign        | Deleterious | Uncertain significance |
|         | COL5A1  | Missense | 0.000121 | c.5411C>A      | p.(Thr1804Asn) | Benign        | Deleterious | Uncertain significance |
| 75      | GFI1B   | Missense | 0.00438 | c.503G>T       | p.(Cys168Phe) | Probably damaging | Deleterious | Likely pathogenic   |
|         | (Rabbolini et al., 2017) |         |        |                   |               |                |            |                     |
| 76      | THBD    | Missense | 0      | c.716C>T        | p.(Ala239Val) | Benign        | Tolerated  | Uncertain significance |
| 77      | THBD    | Missense | 0      | c.752G>A        | p.(Gly251Asp) | Probably damaging | Deleterious | Uncertain significance |
|         | COL5A2  | Missense | 0.0001  | c.2786C>T       | p.(Ala929Val) | Probably damaging | Tolerated  | Uncertain significance |

(Continues)
| Patient | Gene(s) | VEP | MAX AF | Genomic variation | Protein effect | PolyPhen | SIFT | Pathogenicity |
|---------|---------|-----|--------|-------------------|---------------|----------|------|--------------|
| 78      | STXBP2  | Missense | 0.000201 | c.911C>T | p.(Thr304Met) | Probably damaging | Deleterious | Uncertain significance |
|         | MCFD2   | Missense | 0       | c.364G>A | p.(Asp122Asn) | Probably damaging | Deleterious | Likely pathogenic |
| 79      | NBEAL2  | Missense | 0.000128 | c.3184G>A | p.(Val1062Ile) | Possibly damaging | Deleterious | Uncertain significance |
|         | AP3D1   | Missense | 0.0014  | c.1363G>A | p.(Ala455Thr) | Possibly damaging | Deleterious | Uncertain significance |
| 80      | RUNX1   | Missense | 0       | c.1270T>G | p.(Ser424Ala) | Possibly damaging | Deleterious | Uncertain significance |
|         | MPL     | Missense | 0       | c.305G>A | p.(Arg102His) | Possibly damaging | Deleterious | Uncertain significance |
| 81      | AP3B1   | Missense | 0.000809 | c.2188C>T | p.(Arg730Trp) | Benign | Deleterious | Uncertain significance |
| 82      | TUBB1   | Missense | 0.0001  | c.4C>T | p.(Arg2Cys) | Probably damaging | Deleterious | Uncertain significance |
|         | TUBB1   | Missense | 0.0002  | c.68T>C | p.(Met23Thr) | Benign | Tolerated | Uncertain significance |
| 83      | LPA     | Stop gained | 0.001  | c.5081C>G | p.(Ser1694Ter) | NA | NA | Uncertain significance |
| 84.1    | F5      | Missense | 0.00806 | c.5245C>G | p.(Leu1749Val) | Possibly damaging | Tolerated | Uncertain significance |
| 84.2    | F5      | Missense | 0.00806 | c.5054C>G | p.(Thr1685Ser) | Benign | Tolerated | Uncertain significance |
|         | NBEAL2  | Missense | 0.00276 | c.5021G>A | p.(Arg1674His) | Benign | Deleterious | Uncertain significance |
|         | HP55    | Missense | 0.00593 | c.345G>A | p.(Met115Ile) | Benign | Tolerated | Uncertain significance |

Note: Variants previously reported in the literature are indicated. NCBI reference sequences: ABCG5 (NM_022436.2); ABCG8 (NM_022437.2); ACTN1 (NM_001102.3); ACVRL1 (NM_001077401.1); ADAMTS13 (NM_139025.4); ANKR2D6 (NM_01256053.1); AP3B1 (NM_003664.4); AP3D1 (NM_003938.6); ARPC1B (NM_001987.4); F10 (NM_000504.3); F11 (NM_000128.3); F13A1 (NM_000130.4); F7 (NM_000132.3); FGA (NM_005050.3); FGB (NM_005141.4); FGG (NM_01870.2); FLN (NM_000157.3); GBA (NM_000157.3); GFCX (NM_000212.2); ITGA2B (NM_000419.3); ITGB3 (NM_000419.3); ITGAV (NM_000212.2); KIF1B (NM_000892.3); LPA (NM_000157.3); MP1G6B (NM_138272.2); MCL2 (NM_005373.2); MSH2 (NM_000247.4); MYH9 (NM_000247.4); NBEA (NM_001754.3); PLAT (NM_000930.3); PLG (NM_000301.3); PRO1 (NM_000313.3); PRO2 (NM_000313.3); PTPN11 (NM_000329.3); RASGR2P2 (NM_001754.3); SERPINC1 (NM_000488.3); SLCA4A (NM_016180.3); SLCN14 (NM_001129820.1); SMAD4 (NM_005359.5); STXB1P2 (NM_006949.3); THBD (NM_000361.2); THPO (NM_000460.2); TPM4 (NM_003290.2); TUBB1 (NM_030773.3); VPS33B (NM_018668.3); WIF (NM_000552.3); WAS (NM_000377.2).

Abbreviations: NA, not available; NVD, no variant detected; SIFT, sorting intolerance from tolerance; VEP, variant effect predictor.
candidate genes within the IBD gene panel. First, a rare 604-kbp CNV loss was noted on chromosome 11q24.3 in patient 35 which covered nine genes including FLI1. A further rare deletion was found on chromosome 11q24.3 in patient 71 which covered 31 genes including FLI1 (Figure 3a, b).

Following ExomeDepth alignment with a panel of controls the reads ratio was around 0.5 which indicates heterozygosity, as observed in Table 3. A rare CNV gain was noted in patient 45 within TBXA2R on chromosome 19p13.3 and containing four genes in total (Figure 3c). The CNV reads ratio was 2.72 which is indicative of a heterozygous insertion.

3.6 | Oligogenic findings in patient cohort

Within the patient cohort there were several examples of potential oligogenic inheritance involving either two or more gene variations from the IBD gene panel. Of particular interest was patient 16 who demonstrated an apparent pathogenic missense variant in RUNX1 and a likely pathogenic variant in ITGB3, both of which are plausible candidate variants to explain the thrombocytopenia and bleeding history observed. Patient 20 harbored two heterozygous missense variants within GP6 and THBD in which the patient had a platelet function disorder and episodes of bleeding. In patient 30, likely pathogenic and pathogenic variations were found in GPP1BA and GBA respectively. Again this patient had a low platelet count and a history of bleeding.

4 | DISCUSSION

NGS approaches have increasingly been used over the last decade in the molecular diagnosis of IBD. Here, we present a large-scale application of WES analysis by using a robust molecular diagnostics platform for diagnosis of 117 patients recruited to the UK-GAPP study. The aim was to assess the ability of Congenica software to analyze WES data of the patients for both sequence and structural variants by targeting a known panel of IBD genes. Subsequently, patients with variants in known bleeding disorder genes can be eliminated by a series of filtration steps and WES data targeted for those with undetected variants who may harbor a variant in novel genes. We included and applied a total of 119 genes to our patient cohort for filtering; it is, however, important to note that currently only 88 of these genes are considered clinical-grade genes according to Genomics England. The remainder of the genes may become clinical grade once more variants are identified in patients and deemed pathogenic over time.

Phenotypic presentation and platelet counts varied considerably among our recruited patients, which is consistent with the variability of clinical presentation between patients with suspected IBDs. However, the majority of patients 33/117 (28.2%) were noted with a platelet function defect and 23/117 (19.6%) patients represented thrombocytopenia. Of the 117 recruited patients, 15 (12.8%) were deemed to have a macrothrombocytopenia. Platelet function studies revealed the presence of a combination of platelet defects in addition to thrombocytopenia in 36/117 (30%) of patients. The majority of the patients with platelet defects displayed both secretion and Gi defects (Supplementary figure S2). However, a previous study has shown that some patients with normal lumi-aggregometry results have platelet spreading defects, indicating the difficulties faced when diagnosing patients with IBDs and the multitude of assays required for platelet phenotype disorders to be diagnosed (Khan et al., 2020).

Overall, a total of 135 variants in genes implicated in bleeding disorders were identified across all 117 patients and all variants were observed in a heterozygous state, implicating dominant inheritance patterns. The study has shown that the majority of
### Table 3: Copy number variations detected in 15 patients by using ExomeDepth calling approach

| Patient | Gene | Band  | Location | Size | Type | Reads expected | Reads observed | Reads ratio | CNV | Bays factor | Patients overlap |
|---------|------|-------|----------|------|------|----------------|---------------|-------------|-----|-------------|-----------------|
| 21      | ANKR- | 10p12.1 | X:102780843-27389421 | 10.8 kbp | Loss | 254           | 157           | 0.62        | 1236 | 8.26        | 1               |
| 31      | ANKR- | 10p12.1 | X:102780843-27389421 | 10.8 kbp | Gain | 271           | 369           | 1.36        | 272  | 6.13        | 1               |
| 21      | PIGA  | Xp22.31-p21.3 | X:15337573-15353676 | 21.0 Mbp | Loss | 187,776       | 105,800       | 0.56        | 1126 | 30.50       | 16              |
|         | GATA1 | Xp21.1-q13.3 | X:48644962-48652716 | 41.5 Mbp | Loss | 458,560       | 256,748       | 0.56        | 112  | 6580        | 27              |
|         | WAS   | Xp21.1-q13.3 | X:48539485-48549818 | 41.5 Mbp | Loss | 458,560       | 256,748       | 0.56        | 112  | 6580        | 27              |
|         | F9    | Xq25-q27.2 | X:138612917-138645617 | 19.0 Mbp | Loss | 153,271       | 86,478        | 0.56        | 1128 | 23.60       | 24              |
|         | F8    | Xq27.3-q28  | X:545064063-154255215 | 8.3 Mbp | Loss | 165,645       | 90,968        | 0.55        | 1098 | 24.10       | 31              |
|         | FLNA  | Xq27.3-q28  | X:153576892-153603006 | 8.3 Mbp | Loss | 165,645       | 90,968        | 0.55        | 1098 | 24.10       | 31              |
| 26      | SMAD4 | 18q21.2 | X:488494410-48611415 | 2.0 kbp | Gain | 355           | 495           | 1.39        | 278  | 10.8        | 0               |
| 35      | FL1   | 11q24.3 | X:1128556430-128683162 | 604.6 kbp | Loss | 9862          | 5420          | 0.55        | 11   | 422         | 1               |
| 45      | SLFN14 | 17q12 | X:1733875144-33885117 | 5.2 kbp | Loss | 833           | 402           | 0.48        | 0.966 | 6.35        | 4               |
|         | TBX2R | 19p13.3 | X:193594504-3606838 | 43.5 kbp | Gain | 1467          | 1993          | 1.36        | 272  | 12.2         | 0               |
|         | GP6   | 19q13.42 | X:195525073-5549632 | 77.3 kbp | Gain | 1667          | 843           | 0.51        | 1012 | 39.2         | 0               |
| 46      | FYB1  | 5p13.1 | X:539105338-39274630 | 145.7 kbp | Gain | 2146          | 2630          | 1.23        | 246  | 12.6         | 2               |
|         | NBEA  | 13q13.3 | X:133551642-36247159 | 98.7 kbp | Gain | 527           | 742           | 1.41        | 282  | 6.86         | 0               |
|         | SLFN14 | 17q12 | X:1733875144-33885117 | 9.9 kbp | Loss | 7036          | 3693          | 0.52        | 105  | 15.1         | 4               |
| 47      | SLFN14 | 17q12 | X:1733875144-33885117 | 9.9 kbp | Gain | 4964          | 2886          | 0.58        | 1162 | 6.03         | 4               |
|         | GP1BB | 22q11.21 | X:2219730468-19712294 | 37.8 kbp | Gain | 421           | 224           | 0.53        | 1064 | 8.47         | 0               |
| 48      | SLFN14 | 17q12 | X:1733875144-33885117 | 9.9 kbp | Gain | 4090          | 11381         | 2.78        | 556  | 13.3         | 4               |
| 52.1    | PIGA  | Xp22.2-p22.13 | X:15337573-15353676 | 3.6 Mbp | Loss | 49,781        | 32,738        | 0.66        | 1316 | 27.5         | 3               |
|         | GATA1 | Xp11.3-p11.22 | X:48644962-48652716 | 5.3 Mbp | Loss | 154,142       | 94,557        | 0.61        | 1226 | 75.6         | 15              |
|         | WAS   | Xp11.3-p11.22 | X:48539485-48549818 | 5.3 Mbp | Loss | 154,142       | 94,557        | 0.61        | 1226 | 75.6         | 15              |
|         | F9    | Xq27.1 | X:136612917-136645617 | 47.6 kbp | Loss | 11,569        | 7853          | 0.68        | 1358 | 53.5         | 1               |
|         | F8    | Xq28   | X:154064063-154255215 | 4.4 Mbp | Loss | 148,620       | 91,530        | 0.62        | 1232 | 7.30         | 15              |
|         | FLNA  | Xq28   | X:154064063-154255215 | 4.4 Mbp | Loss | 148,620       | 91,530        | 0.62        | 1232 | 7.30         | 15              |
| 52.2    | PIGA  | Xp22.33-p213 | X:15337573-15353676 | 23.5 Mbp | Gain | 205,946       | 296,796       | 1.45        | 29   | 28.9         | 21              |
|         | GATA1 | Xp11.3-q13.3 | X:48644962-48652716 | 29.3 Mbp | Gain | 361,570       | 519,986       | 1.44        | 288  | 41.50        | 25              |
|         | WAS   | Xp11.3-q13.3 | X:48539485-48549818 | 29.3 Mbp | Gain | 361,570       | 519,986       | 1.44        | 288  | 41.50        | 25              |
|         | F9    | Xq25-q27.3 | X:136612917-136645617 | 14.4 Mbp | Gain | 134,092       | 194,039       | 1.45        | 29   | 16.9         | 21              |
|         | F8    | Xq27.3-q28 | X:154064063-154255215 | 9.9 Mbp | Gain | 153,704       | 221,043       | 1.44        | 288  | 18.30        | 38              |
| 57      | COL5A1 | 9q34.3 | X:137533620-137736686 | 190 bp | Loss | 158           | 108           | 0.68        | 1368 | 5.27         | 0               |
plausible candidate variants were associated with IT genes which explain the association of thrombocytopenia with platelet defects in the majority of patients. When considering pathogenicity prediction, 22 patients were classified as pathogenic and 26 patients as likely pathogenic, while 87 patients had uncertain pathogenicity and therefore classified as uncertain significance. A targeted WES analysis was previously carried out on some patients which identified genetic variants in inherited thrombocytopenia with or without secondary qualitative defects (Johnson et al., 2018; Johnson, Lowe, et al., 2016). This study has conclusively identified these genetic variants, which indicates the ability of the Congenica platform to analyze and provide suitable validation of WES data in these patients.

If we compare the performance of the Congenica tool employed here with other bioinformatic platforms we observe the following: 25 variants were identified by the Congenica software as well as other bioinformatic tools and the majority of them were either pathogenic or likely pathogenic. A further 24 variants were classified as pathogenic or likely pathogenic by the Congenica software only. Therefore, this showed that the Congenica software is a more robust tool to analyze WES as it provides a higher variant detection rate compared with other bioinformatic tools. It is also important to note that we did not include variants of uncertain significance here in this evaluation as it is difficult to assign causality but are still plausible pathogenic variants. Congenica software also has the added benefit of detecting CNVs, a process which is notoriously difficult yet valuable in identifying rare causative variants in heterogeneous diseases. Congenica utilizes the integrated ExomeDepth tool to compare a target with reference and here, identified rare CNVs in this population. Congenica software alongside targeted gene panel searching, allows for efficient and accessible detection of variants and with some clinical interpretation will be a valuable tool when analyzing large datasets (Nowakowska, 2017; Valsesia et al., 2012).

### Table 3 (Continued)

| Patient | Gene   | Band Location       | Type  | Size        | CNV       | Reads expected | Reads observed | Reads ratio | Bays factor |
|---------|--------|---------------------|-------|-------------|-----------|----------------|---------------|-------------|-------------|
| 67      | PIGA   | X:1533775-1535367   | Gain  | 9.8 Mbp     | Gain      | 56.698         | 284           | 1.42        | 1070        |
|         | F9     | X:138612917-138645617 | Gain  | 5.4 Mbp     | Gain      | 25.745         | 284           | 1.42        | 386         |
|         | FLNA   | X:15357692-153603006 | Gain  | 6.0 Mbp     | Gain      | 32.057         | 284           | 1.49        | 1550        |
| 71      | FLI1   | 11:12855640-128683162 | Loss  | 2.5 Mbp     | Loss      | 10.050         | 284           | 0.53        | 912         |
| 72      | F9     | X:138612917-138645617 | Gain  | 5.4 Mbp     | Gain      | 32.672         | 284           | 1.42        | 469         |
|         | FLNA   | X:15357692-153603006 | Gain  | 6.0 Mbp     | Gain      | 32.667         | 284           | 1.43        | 322         |

Abbreviation: CNV, copy number variation.

Paris-Trousseau syndrome, characterized by a bleeding defect with large α-granules and abnormal megakaryocyte morphology is well documented, which is caused by a dominant inheritance of q23 deletion on chromosome 11 (Stevenson et al., 2015). Patients with this disorder have variable size of chromosomal deletion associated with different components of the syndrome. A hemizygous deletion of FLI1 was attributed to the platelet defect in two individuals of our cohort. These CNVs noted in patients 35 and 71 cover the deletion region in FLI1 and are also surrounded by several flanking genes. Both patients presented with thrombocytopenia and a secretion defect which suggest the platelet phenotype and the CNVs in FLI1 to be associated with their disorders. Thromboxane receptor deficiency is an autosomal recessive or dominant disorder characterized by bleeding symptoms associated with quantitative or qualitative defects within the thromboxane receptor (Mundell & Mumford, 2018). Although we did not find any plausible candidate SNVs in the 119 candidate genes or the thromboxane receptor in patient 45, we did note a rare CNV duplication in the TBX2A2R gene and deduce that either alone or in combination with variants in GP6 and SLFN14...
which were also detected, could be causative of the patient’s thrombocytopenia and bleeding. In the future, it would be interesting to investigate these CNVs further to determine the extent of the contiguous deletions or insertions by long-range polymerase chain reaction and sequencing to determine the breakpoints and mechanisms of the variant, as well as confirming these regions using multiplex ligation-dependent probe amplification, should kits be available for these genomic regions.

**FIGURE 3** Copy number variants found in cohort of GAPP patients. Screenshots from the Congenica software CNV/structural variant tab in patients showing (a) copy number variant (loss) found in patient 35 which includes FLI1 and eight other genes on chromosome 11q24.3; (b) copy number variant (loss) found in patient 71 which includes FLI1 and 30 other genes on chromosome 11q24.3; and (c) copy number variant (gain) found including TBXA2R in patient 45 and three other genes within chromosome 19p13.3

**FIGURE 3** Continued
In summary, we show validation and a practical approach of a robust diagnostic platform that can be employed for WES analysis. In this study, we use data from a cohort of patients with suspected IBDs: a broad category of diseases, well acknowledged in the hematology field as difficult to classify and associate to single causative genetic abnormalities. This study has shown the ability of the software to detect CNVs with high efficiency with the use of targeted gene panels as a replacement of traditional methods for detecting CNVs.

To conclude, our data reveals use of a highly sensitive and valuable tool which can be used for detecting SNVs and CNVs based on WES data. To our knowledge, this is one of the first studies, although in a research setting, to implement this software for both SNV and CNV analysis. We see this as a leap forward in the ability to classify highly complex disorders with a high degree of heterogeneity within the wider scientific community providing concise and definitive diagnosis for patients.

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CONFLICT OF INTERESTS
The authors declare that there are no conflict of interests.

AUTHOR CONTRIBUTIONS
Neil V. Morgan designed the research. Ibrahim Almazni performed the experiments and analysis. Rachel J. Stapley performed analysis of the data. All authors contributed to the writing of the manuscript and revised versions.

DATA AVAILABILITY STATEMENT
The variants reported in this manuscript have now been submitted to a public database and can be found at ClinVar (https://www.ncbi.nlm.nih.gov/clinvar/).

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SUPPORTING INFORMATION

Additional Supporting Information may be found online in the supporting information tab for this article.

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