Added Value of Reanalysis of Whole Exome- and Whole Genome Sequencing Data From Patients Suspected of Primary Immune Deficiency Using an Extended Gene Panel and Structural Variation Calling

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Background: Knowledge of the genetic variation underlying Primary Immune Deficiency (PID) is increasing. Reanalysis of genome-wide sequencing data from undiagnosed patients with suspected PID may improve the diagnostic rate.

Methods: We included patients monitored at the Department of Infectious Diseases or the Child and Adolescent Department, Rigshospitalet, Denmark, for a suspected PID, who had been analysed previously using a targeted PID gene panel (457 PID-related genes) on whole exome- (WES) or whole genome sequencing (WGS) data. A literature review was performed to extend the PID gene panel used for reanalysis of single nucleotide variation (SNV) and small indels. Structural variant (SV) calling was added on WGS data.

Results: Genetic data from 94 patients (86 adults) including 36 WES and 58 WGS was reanalysed a median of 23 months after the initial analysis. The extended gene panel included 208 additional PID-related genes. Genetic reanalysis led to a small increase in the proportion of patients with new suspicious PID related variants of uncertain significance (VUS). The proportion of patients with a causal genetic diagnosis was constant. In total, five patients (5%, including three WES and two WGS) had a new suspicious PID VUS identified due to reanalysis. Among these, two patients had a variant added due to the expansion of the PID gene panel, and three patients had a variant reclassified to a VUS in a gene included in the initial PID gene panel. The total proportion of patients with PID related VUS, likely pathogenic, and pathogenic variants increased from 43 (46%) to 47 (50%), as...
WES/WGS data have been shown to increase the rate of genetic data is important to consider in previously undiagnosed cases. In strati...and heterozygous variants in genes with autosomal recessive inheritance in 8 patients.

Conclusion: These data indicate a possible diagnostic gain of reassessing WES/WGS data from patients with suspected PID. Reasons for the possible gain included improved knowledge of genotype-phenotype correlation, expanding the gene panel, and adding SV analyses. Future studies of genotype-phenotype correlations may provide additional knowledge on the impact of the new suspicious VUSs.

Keywords: structural variation analysis, primary immune deficiencies (PID), single nucleotide variant analysis, whole genome sequencing (WGS), reanalysis approach, whole exome sequencing, small INDELs, gene panel analysis

INTRODUCTION

Primary immune deficiency (PID) is defined as immune dysfunction due to a genetic cause, which causes either inhibition and/or gain of function of components within the innate and/or adaptive immune system (1). The prevalence of patients with PID is estimated to be as high as one out of every thousand individuals (2). Patients with PID have impacted quality of life and increased mortality primarily due to infectious complications (3). Currently the total number of different monogenetic PID disorders extends to over 400 and is continuously increasing (4). Nevertheless, targeted analysis of PID genes on whole exome- or whole genome sequencing (WES/WGS) data detects monogenic pathogenic variants in less than 50% of patients suspected of PID (5, 6). Due to the continuously increasing knowledge of genes and variants related to PID, a reanalysis of prior WES/WGS data with an extended PID gene panel in undiagnosed patients may improve the diagnostic performance. The potential benefit of performing reanalysis needs to be established, so that institutes can properly weigh up cost benefit.

There are wide ranging benefits of providing a genetic diagnosis for the individual patients. The confirmation or re-classification of a clinical diagnosis may provide the potential for treatment adjustments. In addition, a clarification of the inheritance pattern is important to provide genetic counselling for relatives at risk, as well as the opportunity for prenatal or preimplantation genetic tests in relevant cases. Furthermore, the gain in knowledge of the underlying causes of PID diseases can pave the way for the development of novel biomarkers in risk stratification and treatments. Therefore, reanalysis of WES/WGS data is important to consider in previously undiagnosed cases. In other disease groups, conducting a semiautomated reanalysis of WES/WGS data have been shown to increase the rate of genetic diagnoses (7–10), primarily due to the analysis of newly added genes (11–13).

In this study, we aimed to assess the increase in the diagnostic rate after reanalysis using an extended gene panel on WES/WGS data in a cohort consisting of patients over 16 years of age and suspected of PID. The reanalyses included a repeated analysis for single nucleotide variants (SNVs) and small indels in 94 patients and adding analysis for structural variants (SVs) in 56 patients analysed with WGS. We investigate the proportion of new genetic variants identified after the reanalysis due to either 1) improved knowledge of phenotype-genotype correlation leading to a variant re-classification 2) expansion of the PID gene panel or 3) addition of SV analysis. To our knowledge, this is the first study conducting a WES/WGS data reanalysis in a PID population.

MATERIALS AND METHODS

Patient Population

Patients monitored at the Department of Infectious Diseases, Rigshospitalet, Copenhagen, Denmark, for a suspected PID between January 2016 and September 2020 were offered WES/WGS analysed with a targeted PID gene panel (PID-1), after clinical evaluation and most often also following immunological analyses. Patients who accepted the offer and gave informed consent were included in the study (Figure 1). The study was approved by the national ethics committee. Patients fulfilling the diagnostic criteria for Common Variable Immunodeficiency (CVID) (14, 15) were subsequently excluded from this analysis, as a separate prospective study focused on CVID is ongoing. However, one child, who fulfilled the diagnostic criteria for CVID, was included as the child was part of a family including a parent and sibling with antibody deficiency. The excluded patients with CVID constituted approximately half of the patients suspected of PID and referred for genetic analysis at our institute.

Phenotype data were collected from patients records and assigned the Human Phenotype Ontology (HPO) terms (16). Patients enrolled between January 2016 and January 2018 were analysed with WES (n=37) and patients enrolled between January 2018 and September 2020 were analysed with WGS (n=58). One of the patients analysed with WES, who had a causal pathogenic variant and had died, was excluded from reanalysis. In total 94 patients had WES/WGS data reanalysed for SNVs and small indels. Two of the patients with WGS data were excluded.
from SV analysis, one of these had poor quality control of WGS data and one had a causal pathogenic SV reported in the initial analysis. In total, WGS data from 56 patients were analysed for SVs. The genetic analysis consisted of three different types of entities: 1) 18 Trio analysis (the analysis of a patient and the patient’s parents), 2) 60 Singletons (one patient analysis) and 3) Seven Families (one patient and one or more relatives also presenting symptoms compatible with a PID).

**Extended PID Gene Panel**
The initial gene panel, PID-1, was developed in 2015 and contained 457 genes. PID-1 genes were primarily from the 2016 European Society for Immunodeficiencies online registry of diseases genes (17) and Practice parameter for the diagnosis and management of primary immunodeficiency (18). References included in the development of PID-1 are listed in Supplementary. Based on a literature search in October 2020, 208 genes were added resulting in a total of 665 genes in PID-2. The majority of genes were added from Genomics England Panel App (19) (Supplementary). PID-1 and PID-2 included both known disease-associated PID genes and candidate genes for which the association with PID needs further study. PID-2 was implemented for clinical use in our laboratory after 1 November 2020 (Supplementary Table 1).

**Whole Exome Sequencing**
Genomic DNA was extracted from whole blood. 500 ng of DNA was fragmented to an average size of 300 base pairs using Covaris S2 (Woburn, MA, USA). Adaptor ligation was performed using the KAPA HTP Library Preparation Kit (Roche Diagnostics, Basel, Switzerland) on a Sciclone G3 robot (Perkin Elmer, Waltham, MA, USA). Sequence capture targeting the exome was done with the SureSelectXT Clinical Research Exome kit (Agilent) and followed sequencing on HiSeq2500 or NextSeq500 (Illumina, San Diego, CA, USA). For WES the sequencing protocol aimed to achieve at least 95% of positions within the capture covered by minimum 10x and average coverage of 50x.

**Whole Genome Sequencing**
Genomic DNA was isolated from whole blood samples using the liquid handling automated station (Tecan Freedom EVO HSM 2.0 Workstation) according to the manufacturer’s instructions.
MaxEntScan (28). In the analysis of SNVs and small indels we reanalysed WES/WGS data from all patients with variant, reported pathogenic in The Human Gene Mutation boundaries and +/- 10 kb for the 1st exon of the genes for variants. For SVs, we used the same +/- 20 bp for intron used the phenotypic information to evaluate the relevance of variants in coding regions (+/− minor allele frequency of less than 2% in GnomAD (25) data and of variants. In SNV and small indels analysis only variants with a family members was included in the applied in Ingenuity. In the trios and families, the information of genome data was analysed, or disease speci all patients. In a minor proportion of patients all exome or calling was genotype driven as the PID gene panels were used for sequencing unless they did not fulfi the required quality measures (coverage, variant allele frequency etc.). The variants were classified in accordance with the American College of Medical Genetics and Genomics and the Association for Molecular Pathology (ACMG/AMP) guidelines (29, 30). For each patient, data was analysed and variants were classified by two molecular biologists and/or clinical geneticists.

**Variant Calling**

Single nucleotide variants and small insertions and deletions were called with GATK Haplotype Caller version 4.1.3.0 in accordance with the GATK best practice guidelines developed by the Broad Institute (20). Structural variants were called by Manta version 1.5.0 (21), Lumpy version 0.3.0 (22), and CNVnator version 0.3.3 (23). Thereafter, sequences called with at least two of three tools were manually inspected.

**Variant Exploration and Classification**

For both WES and WGS datasets variant filtering was performed using the Ingenuity Variant Analysis or QCI software (Qiagen Bioinformatics, Redwood City, CA). Integrative Genomics Viewer (IGV) Genome Browser (24) was used to manually verify findings in the aligned raw sequencing data. The variant calling was genotype driven as the PID gene panels were used for all patients. In a minor proportion of patients all exome or genome data was analysed, or disease specific search words were applied in Ingenuity. In the trios and families, the information of family members was included in the filtering and interpretation of variants. In SNV and small indels analysis only variants with a minor allele frequency of less than 2% in GnomAD (25) data and variants in coding regions (+/− 20 base pairs) were selected for further analysis unless an established Pathogenic common variant, reported pathogenic in The Human Gene Mutation Database (26) or ClinVar (27) or predicted a splice effect by MaxEntScan (28). In the analysis of SNVs and small indels we reanalysed WES/WGS data from all patients with PID-2 and used the phenotypic information to evaluate the relevance of variants. For SVs, we used the same +/- 20 bp for intron boundaries and +/- 10 kb for the 1st exon of the genes for promoter/regulatory regions. By WGS-based SV-analysis we could pick up as small as 30bp SVs. SV analysis on WGS data was performed genome wide as an explorative analysis. Only SVs affecting genes included in PID-2 and/or genes related to key PID-2 characteristics are presented in Table 1 and detailed phenotype data is in Supplementary Table 2. The major clinical subgroups in our PID cohort were unclassified antibody deficiencies and unclassified autoinflammatory diseases. The median time between the initial- and the reanalysis of WES/WGS data was 23 months (range 1-57 months).

**Immunological Analysis**

The standard immunological analysis included flow cytometry-based leukocyte count and subtyping of maturation- and activation stages, somatic hypermutation (measuring in kappa light chain transcripts), T-lymphocyte proliferation (for patients between the ages of 0 and 10 or for adults suspected of a T-cell defect) and complement activation screen. Further specific immunological analyses were performed based on the patient’s clinical presentation and results of the basic immunological analysis after conference with clinical immunologists (31).

**RESULTS**

**Patient Population**

In total, 95 individuals suspected of having PID in 85 entities consisting of 18 trios, 60 singletons and 7 families were included in the study. Patients were categorized and sub-grouped according to ESID (2019) clinical criteria (17) based on the presentation at the time of the initial genetic analysis (PID-1). The patients’ clinical and immunological phenotypes were assessed again prior to the reanalysis in order to include any additional information obtained between the initial- and reanalysis. Baseline characteristics are presented in Table 1 and detailed phenotype data is in Supplementary Table 2. The major clinical subgroups in our PID cohort were unclassified antibody deficiencies and unclassified autoinflammatory diseases. The median time between the initial- and the reanalysis of WES/WGS data was 23 months (range 1-57 months).

**Results of Genetic Analysis**

The results of the initial genetic analyses (PID-1) and reanalyses (PID-2) are presented in Supplementary Table 3 and Table 2, respectively. The number of patients with genetic findings identified before and after reanalysis are summarized in Table 3 and described further below. In the initial analysis, a causal PID diagnosis was detected in eight patients, including

| TABLE 1 | Baseline Characteristics. |
|-----------------|--------------------------|
| Total number of patients, (n) | 95 |
| Adults, ≥18 years (n), Children (n) | 87, 8 |
| Age at inclusion (years), median (range) | 37 (16-75) |
| Symptom onset at age 0-10 years (n (%)) | 44 (46) |
| Symptom onset at age 10-20 years (n (%)) | 12 (13) |
| Symptom onset at age 20-40 years (n (%)) | 18 (19) |
| Symptom onset at age > 40 years (n (%)) | 20 (21) |
| Symptom onset at unknown age | 1 (1) |
| Female (n (%)) | 65 (68) |
| Time between the initial- and the reanalysis of WES/WGS data (months), median (range) | 23 (1-57) |
| PID related Clinical Characteristics* | | |
| Recurrent/opportunistic infections (n (%)) | 78 (81) |
| Autoimmunity (n (%)) | 15 (16) |
| Malignancy (n (%)) | 5 (5) |

*Patients can have more than one PID related clinical characteristic registered.

PID, Primary immune deficiency; WES, Whole Exome Sequencing; WGS, Whole Genome Sequencing.
likely pathogenic- (LP) and pathogenic (P) variants (Supplementary Table 3). One of these eight patients was excluded from reanalysis. No new PID diagnoses were detected upon reanalysis. Seven of the patients with a genetic diagnosis had a SNV or small indel (11S, 25S, 44T, 45S, 46S, 48T, 74T) and one patient had a SV (63S). One patient (52S) had a variant of uncertain significance (VUS) in NLRP3, which was compatible with the phenotype and segregated in the affected family members. Due to reduced penetrance of the phenotype this variant could not be classified as LP or P. Therefore, this patient was not counted as having a genetic diagnosis. Additionally, in the initial analysis eight patients (3S, 37S, 53S, 54S, 62FP, 62FC, 77S, 81S) had suspicious PID variants detected (LP/P), which were not explanatory for the PID phenotype (including heterozygous variants in PID genes with AR inheritance). Further, 28 patients had suspicious PID VUS(s) detected. In the reanalysis, five (5%) patients had new suspicious PID related VUS(s) identified. Four out of the five patients did not have any variant identified in the initial analysis. Due to the addition of VUSs, the reanalysis increased the total number of patients in the cohort with PID related VUS(s), LP, and P variant (s) from 43 (46%) to 47 (50%).

Single Nucleotide Variation and Small Indels Analysis

When the genetic data was reanalysed, a new suspicious SNV VUS was identified in five patients (5%). The genetic variant details are shown in Table 2. We divided the patients into four groups based on the category of the reanalysis responsible for the detection of the new finding, as shown in Table 3. Category 1) Three of the patients (19S, 49S, 59S) had a new VUS added due to improved knowledge of phenotype-genotype correlation in a gene also included in PID-1. Category 2) Two patients (29S, 31S) had a VUS added due to the expansion of the PID gene panel. Additionally, one patient had an incidental finding (7S). Of these five patients with new PID related VUS(s), two patients (19S and 59S) had a single VUS in an AR/autosomal dominant (AD) gene: FLG and SERPING1, respectively, and three patients (29S, 31S, 49S) had a single VUS in an AD gene: OAS1, TNFAIP3 and NFKB1, respectively. These five patients’ phenotypes were compatible with the AR/AD or AD heritable disorder and functional immunological analysis in addition to genetic analysis of parents was recommended (where possible).

Structural Variation Analysis

One patient had a pathogenic SV detected in the initial analysis (63S). Category 3) In the reanalysis we only detected SVs in either PID candidate genes and/or heterozygous in PID genes with autosomal recessive inheritance. These variants were included in Category 4 (below).

SNV and SV VUS(s) in PID Candidate Genes and/or Heterozygous in PID Genes With Autosomal Recessive Inheritance

Category 4) We identified eight patients with SNV or SV VUS(s) in PID candidate genes (n=6) and/or heterozygous VUSs in PID genes with AR inheritance (n=2). Of the 56 individuals with WGS data, five patients (9%) had such a SV of uncertain significance identified (10FP, 27S, 32S, 33S, 39FA1). Four patients (9S, 16S, 27S, 75S) had a heterozygous VUS (SNV or SV) in a PID candidate gene. In one of these patients (27S) with Combined Immune Deficiency (CID), we detected a heterozygous, frameshift deletion in LAG3 (lymphocyte-activation gene 3), which is involved in T-cell receptor signalling (32). LAG3 was not included in the PID-2 gene panel. Further, one patient (75S) had a pathogenic incidental finding in addition to the new suspicious VUS in a PID candidate gene. In this patient, we detected a hemizygous pathogenic deletion in EDA, which was causal for the patient’s hypohidrotic ectodermal dysplasia phenotype. EDA is not included in the PID-2 gene panel. The EDA variant was categorized as an incidental finding. Notably, during the initial WGS analysis of this patient, who also presented a phenotype of recurrent infections, genes related to ectodermal dysplasia were included in addition to the PID-1 gene panel. SV analysis was not implemented at the time of the initial analysis. Furthermore, Array CGH had been performed which could not detect the deletion. Two patients (33S, 39FA1) had two VUSs in a PID candidate gene. These patients both had a SV and a SNV of uncertain significance in the same gene and potentially in trans (these SNVs were not included in the results of SNV analysis above). Two patients (10FP, 32S) had a single VUS in an autosomal recessive (AR) gene. These two patients both had a heterozygous tandem duplication in DOCK8. One of the patients presented a phenotype compatible with Hyper IgE syndrome. We added analysis for DOCK8 deep intrinsic SNVs and small indels in these two cases due to prior reports of intron variants in patients with DOCK8 deficiency. We did not identify LP or P variants in this analysis. Finally, one patient (7S) had an incidental finding (VUS) detected at reanalysis.
| ID* | WES/ WGS | Genetic result of SNV or SV analysis (gene, cDNA, Protein, RefSeq) | ACMG classification‡ | SNV/SV predicted coding effect | Zygosity | Inheritance | Allele frequency for NFE in GnomAD (%), and for SV DGV (%), (All) | CADD | Previous report of variant in HGMD (PMID) | Gene function or pathway involvement (PMID) | Comment | Category of reanalysis responsible for the detection of the new finding§ |
|-----|----------|---------------------------------------------------------------|---------------------|-------------------------------|---------|-------------|---------------------------------------------------------------|------|------------------------------------------|-----------------------------------------------|---------|-----------------------------------------------|
| 78  | WGS      | GSNV:c.487G>A, p.(Asp163Asn), NM_001353054.1†               | 3                   | Missense                     | Het     | Unknown     | 0.001                                                        | 26.5 | _                                        | Actin-binding protein (PMID: 30698126) | GSN gene function | _                                              |
| 98  | WGS      | BAKAV1:c.723del, p.(Met241Ilefs), NM_001275707.2             | 3                   | Frameshift                   | Het     | Unknown     | 0                                                           | N/A  | _                                        | B-cell receptor-, CD40-related-, TL7- and TL99 signaling (PMID: 34066164) | _             | 4                                             |
| 10 FP| WGS      | chr9:75297,381074dup Tandem duplication on 9p24.3 (~300kb) including exons 1-21 of DOCK9 (NM_203447.4) | 3                   | _                             | Het     | AR          | DGV: 0.14                                                    | _    | _                                        | TCR (PMID: 27599296), TL99 (PMID: 28882618) | The duplication is not present in the son (10FC). | 4                                             |
| 16S | WES      | NFXB6D (c.359C>T, p.(Ser120Leu), NM_139239.3)               | 3                   | Missense                     | Het     | Unknown     | <0.001                                                       | 24.5 | _                                        | NF-κB (PMID: 32578005) | _             | 4                                             |
| 19S | WGS      | FLG (c.10255C>T, p.(Arg3419*), NM_001127507.2)               | 3                   | Frameshift                   | Het     | Unknown     | 0                                                           | 39   | PMID: 31365035                           | Protein in the skin barrier (PMID: 31509236) | _             | 1                                             |
| 27S | WGS      | LAG3 (c.287_407 del, chr12:682943_6883063del A 135bp deletion in exon 3 of LAG3) | 3                   | _                             | Frameshift | Unknown     | GnomAD-SV: 0, DGV 0                                         | _    | _                                        | T-cell receptor signaling pathway (PMID: 31848144) | _             | 4                                             |
| 29S | WES      | GASP (c.163G>A, p.(Val55Met), NM_002534.3)                   | 3                   | Missense                     | Het     | AD          | 0.006                                                       | 23   | _                                        | Negative regulator of the expression of chemokines and interferon responsive genes in human macrophages (PMID: 30078389), RNA cleavage pathway (PMID: 3082544) | _             | 2                                             |
| 31S | WGS      | TNFAP3 (c.2140C>A, p.(Pro714Thr), NM_006290.4)               | 3                   | Missense                     | Het     | AD          | 0                                                           | 17.19| _                                        | NF-κB (PMID: 32334814) | _             | 2                                             |
| 32S | WGS      | chr9:75299,452698dup Tandem duplication on 9p24.3 (~400kb) including exons 1-44 of DOCK9 (NM_203447.4) | 3                   | _                             | Missense | Het         | AR                                                          | DGV: 0.11 | _                                        | Adaptor protein in RIG-I pathway (PMID: 25636800) | Unknown whether SV and SNV are in cis or trans. | 4                                             |
| 33S | WGS      | MAVS (c.1186C>T, p.(Asp396Glut), NM_001206491.2)             | 3                   | Missense                     | Het     | Unknown     | 0.007                                                       | 23.9 | _                                        | Adaptor protein in RIG-I pathway (PMID: 25636800) | The deletion is mediated by AU7 elements; therefore, it is not possible to identify the exact genomic coordinates. First degree relative with resembling phenotype does not carry the missense variant. Unknown whether SV and SNV are in cis or trans. | 4                                             |
| 33S | WGS      | chr20:g.3836764_384071del, ~4kb deletion including exon 3 of MAVS (c.118,292del, p.(Asp396Glu)*0) | 3                   | Missense                     | Het     | Unknown     | GnomAD-SV: 0, DGV 0                                         | _    | _                                        | Adaptor protein in RIG-I pathway (PMID: 25636800) | _             | _                                             |
| 33S | WGS      | MAVS (c.118,292del, p.(Asp396Glu)*0)                         | 3                   | Missense                     | Het     | Unknown     | GnomAD-SV: 0, DGV 0                                         | _    | _                                        | Adaptor protein in RIG-I pathway (PMID: 25636800) | _             | _                                             |
| 39FA1 | WGS     | FPR2 (c.207C>T, p.(Phe74Leu), NM_001462.3)                   | 3                   | Missense                     | Het     | Unknown     | 0.0046                                                      | 23   | _                                        | Annexin A1/FPR2 (PMID: 31908042) | Unknown whether SV and SNV are in cis or trans. | 4                                             |
| 39FA1 | WGS    | chr19:52271806_52617770dup Tandem duplication on 19q13.41 (~300kb) including exon 2 of FPR2 (NM_001206491.2) and the entire FPR2 gene (NM_001206491.2) | 3                   | Missense                     | Het     | Unknown     | GnomAD-SV: 0, DGV 0                                         | _    | _                                        | Annexin A1/FPR2 (PMID: 31908042) | Unknown whether SV and SNV are in cis or trans. | _                                             |
| 49S | WES      | NFXB7 (c.689G>T, p.(Asp230Leu), NM_0039986.4)                | 3                   | Missense                     | Het     | AD          | ≤ 0.001                                                    | 22.7 | _                                        | NF-κB (PMID: 26663363) | _             | 1                                             |
| 59S | WES      | SERPIN1 (c.507G>T, p.(Ala163Val), NM_0000062.3)              | 3                   | Missense                     | Het     | AD/AR       | 0.11                                                       | 23.6 | PMID: 32869191                           | C1-inhibitor, Major control of kallikrein-kinin system (PMID: 31517426) | _             | 1                                             |

(Continued)
Subsequent analysis proved the variant to segregate with the phenotype in several family members. Additionally, functional immunological analysis confirmed the effect. Therefore, this variant was reclassified to a P variant. ID 46S had a heterozygous, missense variant (STAT1, c.823C>A, p.Gln275Lys), which was absent from controls. Subsequent analysis of the parents proved the variant to be de novo. Therefore, the variant was reclassified to a LP variant. We did not re-classify any VUSs to benign or likely benign variants.

**DISCUSSION**

This study was conducted to investigate the value of performing a reanalysis of WES/WGS data in a PID population consisting of patients over 16 years of age, after an inclusion period of approximately 4 ½ years. The re-analysis (PID-2) was performed a median of 23 months after the initial analysis (PID-1). Upon re-analysis new suspicious PID VUSs were detected in 5% of the patients. Since all new PID related variants (SNVs and SVs) were VUSs, the number of causal genetic PID diagnoses did not increase after the reanalysis. The total rate of patients having PID VUS(s), LP or P variant(s) increased by 4% to a total rate of 50%. Three patients had a VUS identified due to improved knowledge of genotype-phenotype interaction and two patients had a VUS in new PID genes added in PID-2. In addition, we identified five patients with new SVs of uncertain significance and three patients with SNVs of uncertain significance. These eight patients were described separately, as they have VUS(s) in either PID candidate genes or heterozygous in established PID genes with AR inheritance. Reasons for the identification of new variants include the updated tools for calling of SNVs/small indels and SVs as well as improvements of *in silico* analysis. Our findings indicate a possible diagnostic importance of analysing SVs in an adult PID population. Therefore, we aim to transfer the genetic analysis from WES to WGS to be able to perform SV analysis in selected cases with high suspicion of a PID. Furthermore, in selected cases presenting symptoms beyond a PID phenotype, a re-evaluation of the indication for adding additional gene panels or extending the full exome/genome analysis should be considered. The detection of VUSs in new PID candidate genes together with functional immunological analysis of affected pathway(s) can potentially improve the knowledge of genotype-phenotype correlations underlying PIDs.

When comparing rates of genetic findings between PID cohorts, the selection of patients is crucial. Including patients with more severe phenotypes, patients with symptom onset in childhood and a positive family history will lead to a higher molecular diagnostic rate (33). Further, a proportion of PIDs with symptom onset in adulthood are suspected to be caused by polygenetic (34, 35) and/or multifactorial inheritance (36). Additional knowledge of mono- and polygenic variation underlying these PIDs remains to be gained. Methods beyond analysis of single cases and smaller cohorts are needed to detect new genetic variants contributing to PID phenotypes (37, 38).
with copy number variation (deletions and duplications), led to the detection of explanatory variants in all cases analysed (39).

Unclassified antibody deficiencies and unclassified autoimmune diseases constitute the major clinical subgroups in our PID cohort (Table 2). The proportion of antibody deficiencies corresponds to the proportion in a large study of primarily adult PID patients (6). Furthermore, the proportion consisting of CIDs was relatively low (7%), whereas the proportion of pathogen specific immunodeficiencies was relatively high (9%). These deviations could be due to a larger proportion of children in the study by Smith et al. However, these subgroups contain small numbers of patients. Notably, in the subgroup of patients, who did not fulfil ESID clinical criteria, a variant was identified in six of the 13 patients, of which one had a homozygous, pathogenic variant in \textit{ERAP1} (74T) and five had VUS(s) in PID genes. In addition, three patients had familial chronic mucocutaneous candidiasis (CMC) due to pathogenic gain of function variants in \textit{STAT1}.

Two studies from the UK that investigated a semiautomated reanalysis of WES data in more than 1000 families with developmental delay (13) and more than 2000 consecutive cases of WES (12), showed increases of molecular diagnostic rates of 13% (27 to 40%) and 11.5% (25 to 36%), respectively. However, different disease groups are clearly not directly

- **TABLE 3** | Number of variants of uncertain significance (VUS), likely pathogenic- (LP) and pathogenic variants (P) before and after reanalysis (n = 94).

| Total patients identified in the initial analysis (PID-1) | Patients with new variants identified in the reanalysis (PID-1 & PID-2) |
|--------------------------------------------------------|-------------------------------------------------------------|
| Genetic diagnosis causal for PID (LP/P)                |                                                                      |
| 7 (7%)                                                 | 7 (7%)                                                      | 0 |
| Suspicious PID variants (LP/P)                         | 8 (8%)                                                      | 8 (8%) |
| 32 (34%)                                               | 28 (30%)                                                   | 5 (5%)* |
| All PID related VUS, LP and P                          | 47 (50%)                                                   | 43 (46%) | 5 (5%)* |

| Categories of the reanalysis responsible for the detection of genetic findings1 | |
| Category 1 and 2: | |
| Vars identified in analysis for SNV or small indels | |
| 46 (49%) | 42 (45%) | 5 (5%)* |

| Variants detected in analysis for SVs | |
| 8 (8%) | 8 (8%) | 8 (8%) |

| SNV and SV VUS(s) in PID candidate genes and/or heterozygous in PID genes with AR inheritance | |
|---|---|
| 4 (4%) | 2 (2%) | 2 (2%) |

*1 of these 5 patients also had another SNV identified in the initial analysis. **Categories of the reanalysis responsible for detection of the new finding: 1 = improved knowledge of genotype-phenotype, 2 = expansion of PID gene panel, 3 = SV analysis, or 4 = VUS(s) in candidate genes or heterozygous VUS(s) in genes with AR inheritance. **The one patient with a pathogenic SV is listed relative to the number of patients analysed for SVs (n = 56) in double brackets. **Patients with VUS(s) in PID candidate genes and/or heterozygous VUS(s) in PID genes with AR inheritance detected in the reanalysis were described separately and thus not counted in Suspicious PID variants (VUS). AR, Autosomal recessive; LP, likely pathogenic; P, Pathogenic; PID, Primary immune deficiency; SNV, Single nucleotide variant or small indel; SV, Structural variant; VUS, Variant of uncertain significance.

One patient with a causal variant identified with PID-1 was excluded from reanalysis and therefore not included in this table.
comparable. The two studies both included children and the molecular diagnostic rates did not include variants of uncertain significance. In contrast, our findings of new variants in PID genes were solely VUSs. Notably, our cohort primarily investigated adult patients and few children included in families with a suspected PID. As well as developmental delay, PID phenotypes are very heterogenous, especially if milder phenotypes of PID are included. Also, PID patients can present with a symptom onset in adulthood, making it more difficult to determine the impact and pathogenicity of genetic variants, which may be inherited from an apparently unaffected parent. Finally, both PID and developmental delay can potentially arise due to variants with variable expression and/or reduced penetrance.

Data analyses and variant classifications were performed by six molecular biologists and clinical geneticists during the 5-year study period. To limit the effect of interobserver variability, all interpreters followed the ACMG international guidelines (29), and all findings were confirmed by two interpreters. However, inevitably there will be interobserver variability of variant interpretation, which can affect the rate of PID related variants identified. Additionally, when rates of genetic variants are compared between PID cohorts, there can be inter-laboratory variability. Further, over time some monogenic PID disorders known to be caused by autosomal recessive inheritance have been proven also to be caused by autosomal dominant inheritance (40). Therefore, during this study period, the number of heterozygous variants reported has increased. Finally, more patients were included as trio analysis in the beginning of the inclusion period, and more singletons were included towards the end. Trio analysis are often preferable for WES/WGS data interpretation since trio analyses can provide information on potential de novo variants or compound heterozygosity for two variants in the same gene. Therefore, the change to a predominant inclusion of singletons later in the inclusion period may also have led to an increase of heterozygous variants reported, where subsequent analysis of parents is needed to unravel de novo variants or compound heterozygosity for two variants.

One of the limitations of the study was, that analysis for low level of germline mosaicism (<20% of reads) was not performed, since WGS analysis is not ideal for identifying potential germline mosaicism. Another limitation of our study was in the analysis of genes known to have pseudogenes, e.g. IKBKG, NCF1, C4A, and C4B. For such regions specialized bioinformatics pipelines could be required to accurately call the variants in these genes, which was not applied in our study. This limitation is included as a disclaimer in the clinical reports. The targeted analysis performed in our study could also be considered as a limitation, since additional VUSs in new PID candidate genes may be detected by performing analysis of the entire exome or genome. Another limitation was, that we did not consider the deep intronic variants (located deeper then +/-20bp to the exons). However, as the performance of any kind of functional analysis for potential effect of splice- or other non-coding variants was not within our study scope, we decided to focus our analysis to the +/-20bp exon-flanking regions in order to simplify the clinical interpretation of the variants.

Performing a reanalysis of broad sequencing data is resource demanding. As such, a potential gain in diagnostic performance after data reanalysis needs to be balanced against the time and cost involved (41). In contrast to other studies including automated reanalysis, we did not detect new genetic diagnoses upon reanalysis. Yet our results showed that performing the reanalysis may improve genetic results in prior undiagnosed patients. Our findings included both SNVs and SVs in new PID genes (added in PID-2), which shows the importance of regularly extending gene panels for reanalysis in PID. The PID-2 gene panel contains many PID candidate genes (newly described in relation to PID) without prior reports in the literature of variants in patients. Therefore, until the knowledge of gene-disease interaction improves, variants identified in PID candidate genes within PID-2 will primarily be classified as VUSs. Within the Genomics England Panel App approximately half of the genes in the current PID gene panel (Primary immune deficiency Version 2.498) are categorized as red (not enough evidence) or amber (moderate evidence of gene-disease association), where additional evidence is needed to confirm the genotype-phenotype correlation. Thus, additional evidence of pathogenicity for the VUSs identified in our PID cohort may arise from individual, specific functional immunological analysis and segregation analysis.

Due to the high level of heterogeneity of PIDs, the interpretation of rare variants and planning of functional analysis, particularly of variants in PID candidate genes, can benefit from international collaborations, e.g. GeneMatcher (42). Due to costs of manual reanalysis an automated reanalysis of VUS and LP variants incorporating genomic, phenotypic, and pedigree data could ideally prioritize the variants in established PID genes for clinicians to review, in line with what has been implemented for other disease groups (43–46). Currently, new methods are being developed and implemented to improve automation of genetic reanalysis (8, 10, 47). In PID populations, where most genetic variants are VUSs, patients could benefit from regular semiautomated reanalysis of WES/WGS data with a reduced demand of time for manual variant interpretation. However, due to the large extent of PID candidate genes, VUSs detected by automated reanalysis methods in these genes will require additional evidence of pathogenicity from segregation studies, functional analysis and/or other reports of cases with similar phenotypes.

In conclusion, our reanalysis of WES/WGS data did not lead to novel genetic diagnoses causal for PID. We detected new suspicious PID VUS(s) in five (5%) patients suspected of PID. Additionally, eight patients had SNV/SV VUS(s) in PID candidate genes or heterozygous VUS(s) PID genes with AR inheritance. Our data indicate a possible diagnostic value of reassessing WES/WGS data with an extended gene panel and performing analysis of SV in a PID population of primarily adults. To our knowledge, this is the first study to investigate the impact of a WES/WGS data reanalysis and to include structural variation analysis in a PID population. Future improved
knowledge of genotype-phenotype correlation in PID candidate genes may provide additional certainty of the effect of the identified VUSs.

DATA AVAILABILITY STATEMENT

The datasets presented in this article are not readily available because of national legislation. Requests to access the datasets should be directed to sara.bohnstedt.moerup@regionh.dk.

ETHICS STATEMENT

The patients/participants legal guardian/next of kin provided written informed consent to participate in this study. The study was reviewed and approved by the Danish Regional Ethics Committee (H-15009663).

AUTHOR CONTRIBUTIONS

HM, RM, TK, and JL: Planned the study. SM, JR, HM, JL, DM, MH, LB, and TK: Contributed to the design. SM, TK, TM, and MH: Acquisition of the phenotype data. SM, MG, LN-P, RM, and LB: Interpretation, analyses, and quality control of genetic data. SM, HM, and HH: Interpretation of immunologic data. SM, JR, DM, MH, and LB: Wrote the manuscript. All authors critically reviewed and approved the final version of the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2022.906328/full#supplementary-material
