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ELSEVIER

Contents lists available at ScienceDirect

European Journal of Medical Genetics

journal homepage: www.elsevier.com/locate/ejmg





Genome-wide variant calling in reanalysis of exome sequencing data uncovered a pathogenic *TUBB3* variant

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ARTICLE INFO

Keywords: TUBB3 Exome sequencing Genome-wide variant calling Solve-RD ERN ITHACA

ABSTRACT

Almost half of all individuals affected by intellectual disability (ID) remain undiagnosed. In the Solve-RD project, exome sequencing (ES) datasets from unresolved individuals with (syndromic) ID (n = 1,472 probands) are systematically reanalyzed, starting from raw sequencing files, followed by genome-wide variant calling and new data interpretation. This strategy led to the identification of a disease-causing de novo missense variant in TUBB3 in a girl with severe developmental delay, secondary microcephaly, brain imaging abnormalities, high hypermetropia, strabismus and short stature. Interestingly, the TUBB3 variant could only be identified through reanalysis of ES data using a genome-wide variant calling approach, despite being located in protein coding sequence. More detailed analysis revealed that the position of the variant within exon 5 of TUBB3 was not targeted by the enrichment kit, although consistent high-quality coverage was obtained at this position, resulting from nearby targets that provide off-target coverage. In the initial analysis, variant calling was restricted to the exon targets \pm 200 bases, allowing the variant to escape detection by the variant calling algorithm. This phenomenon may potentially occur more often, as we determined that 36 established ID genes have robust off-target coverage in coding sequence. Moreover, within these regions, for 17 genes (likely) pathogenic variants have been identified before. Therefore, this clinical report highlights that, although compute-intensive, performing genomewide variant calling instead of target-based calling may lead to the detection of diagnostically relevant variants that would otherwise remain unnoticed.

1. Introduction

Approximately 1% of the population is affected by intellectual disability (ID) (Maulik et al., 2011), but as the disorder is genetically and clinically

highly heterogeneous, individual genetic causes are very rare, which complicates the diagnostic process and the identification of new disease-gene associations (Vissers et al., 2016). As a result, ~50–70% of individuals with ID remain undiagnosed (Srivastava et al., 2019). To overcome the

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https://doi.org/10.1016/j.ejmg.2021.104402

Received 1 September 2021; Accepted 28 November 2021 Available online 1 December 2021

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problem of rarity and aiming to provide individuals with a diagnosis, the Solve-RD project (Zurek et al., 2021) comprises long-term sharing and systematic reanalysis of next generation sequencing and phenotypic data of 19, 000 unsolved rare disease cases from four European Reference Networks (ERNs), including ~5,000 cases with (syndromic) ID from ERN-ITHACA (Intellectual Disability, TeleHealth and Congenital Anomalies; htt ps://ern-ithaca.eu/). In contrast to recent studies on unsolved cohorts that reanalyzed variant calling files (Satterstrom et al., 2020) or pooled (de novo) variants (Kaplanis et al., 2020), in Solve-RD, new mapping and genome-wide variant calling is the starting point for reanalysis. Reanalysis strategies using genome-wide calling potentially increase diagnostic yield, as enrichment kits are often incomplete, despite the efforts aimed at the enrichment of all protein coding sequences (Ballester et al., 2016). By not restricting variant calling to (extended) targets of enrichment, the intrinsic imperfection of enrichment strategies to generate off-target coverage (Ballester et al., 2016) can be exploited to rescue some of the missing regions.

TUBB3 encodes the neuron-specific β-tubulin-3. Pathogenic TUBB3 missense variants alter microtubule stability (Poirier et al., 2010; Tischfield et al., 2010) and impair axonal guidance in brain development (Huang et al., 2018), causing a spectrum of neuronal migration disorders (Poirier et al., 2010; Tischfield et al., 2010; Huang et al., 2018; Dentici et al., 2020; Fukumura et al., 2016; Shimojima et al., 2016; Whitman et al., 2016). Pathogenic variants can clinically and functionally be distinguished, and either increase microtubule stability, causing congenital fibrosis of extraocular muscles type 3A (MIM#600638) (Tischfield et al., 2010), or impair microtubule stability, leading to the highly variable phenotype of complex cortical dysplasia with other brain malformations type 1 (MIM#614039) (Poirier et al., 2010). A merge of both phenotypes has also been reported (Whitman et al., 2016).

We describe a disease-causing *TUBB3* variant identified in systematic reanalysis of exome sequencing (ES) data under the collaborative efforts of the Solve-RD project and ERN-ITHACA. Our clinical report illustrates the importance of starting reanalysis with raw sequencing data and stresses the diagnostic potential of genome-wide variant calling, especially in reanalysis.

2. Materials (subjects) and methods

2.1. Patient inclusion

In compliance with the local ethical guidelines and the Declaration of Helsinki, all individuals (or legal representatives) in the Solve-RD project provided informed consent. The Radboudumc institutional review board approved the study (2018–4986). For publication of photos, additional consent was obtained.

2.2. Initial diagnostic ES analysis

In 2014, diagnostic ES was performed locally as described previously (de Ligt et al., 2012). Of essence for this clinical report, exome enrichment was performed using the Agilent V4 Sure Select All Exons (GRCh37/Hg19) enrichment kit followed by sequencing on a HiSeq2000 Sequencing System (Illumina) to an anticipated median 75-fold sequence coverage on target regions. Variant calling was performed using the GATK HaplotypeCaller (v3.4) (Poplin et al., 2017), restricted to genomic coordinates of the exome targets±200bp.

2.3. Solve-RD data-sharing for systematic reanalysis

Human Phenotype Ontology (HPO)-standardized phenotypic data and BAM files of ES were uploaded to the RD-Connect Genome-Phenome Analysis Platform (GPAP; https://platform.rd-connect.eu/), and deposited at European Genome-Phenome Archive (EGA; accession numbers EGAZ00001527579, EGAZ00001534566 and EGAZ00001549096 for index, father and mother respectively), as part of the Solve-RD infrastructure (Zurek et al., 2021; Matalonga et al., 2021).

2.4. Variant identification

Systematic reanalysis of ES data in Solve-RD is described by Matalonga et al. (2021). The pipeline includes genome-wide variant calling (i.e. not restricted to the targets of the ES enrichment kit) with GATK (v3.6) (Poplin et al., 2017) to the GRCh37/Hg19 genome build, i.e. hs37d5, as used by Phase2 of the 1000 GP. Variants in genes associated with ID were annotated and prioritized by applying the following filtering parameters: 1) coverage depth ≥8 and genotype quality ≥20, 2) gnomAD (Karczewski et al., 2019) allele frequency <1%, 3) internal RD-Connect allele frequency <2%, 4) "Likely pathogenic" (ACMG, class 4) or "Pathogenic" (ACMG, class 5) in ClinVar (Landrum, 2018). Prioritized variants were interpreted in the context of the phenotype and visualized in the Integrative Genomics Viewer (IGV) (Robinson, 2011) alongside the exome data of the parents.

2.5. Sanger and amplicon-based deep sequencing

Variant validation by Sanger sequencing was performed using a routine diagnostic workflow (de Ligt et al., 2012). To exclude low-level mosaicism in the father, amplicon-based deep-sequencing was performed on the Ion Torrent platform from Life Technologies (Gilissen et al., 2014). PCR amplicons were generated in accordance with standard protocols, followed by read processing and alignment using JSI SeqNext Version 5.1.0 Build 503 (FW: GTGCGGAAGGAGTGTGAAAA; RV GGCATGAAGAAGTGCAGG).

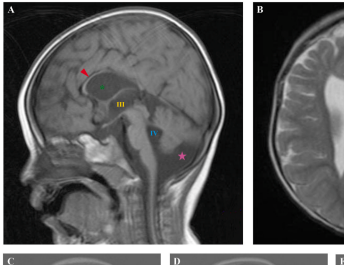
2.6. Detecting off-target coverage on coding sequence and intersection with HGMD variants

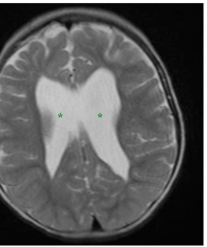
Occurrence of off-target coverage on coding sequence across individuals was analyzed for enrichment kits Agilent V4 and V5 (Sure Select All Exons; GRCh37/Hg19), as ~70% of ES datasets shared for ERN ITHACA in the Solve-RD consortium was generated after enrichment with Agilent V4 or V5. We used 20 representative ES datasets for both kits, and defined robust off-target coverage occurring consistently across individuals at depth sufficiently for variant calling if 95% of tested individuals showed coverage exceeding 20-fold at coding sequence (Gencode v31 (Frankish et al., 2019)) located outside extended targets of the respective enrichment kit (genomic coordinates of the exome targets±200bp of Agilent V4 and V5). For the resulting regions, an intersection was made with disease-causing (DM) or likely disease-causing (DM?) SNVs/indels reported in HGMD Professional (Stenson et al., 2003) with specific attention for genes associated to ID (Radboudumc diagnostic ES ID gene panel DG3.1.0, comprising 1,392 genes) (Radboudumc. Intellectual, 2021).

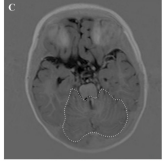
3. Results

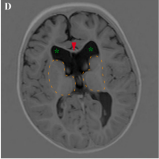
3.1. Clinical characteristics

We report on a 16-year-old female proband with severe ID, for whom extensive diagnostic testing in a university medical center in 2014 could not identify the (genetic) diagnosis. The proband was the first child from healthy, non-consanguineous parents and had a healthy brother and sister. Besides hyperemesis gravidarum, pregnancy was uneventful. Spontaneous vaginal delivery occurred at 42 weeks of gestation, and although delivery was complicated by meconium-stained amniotic fluid, there were no problems in the neonatal period (Apgar scores 9/9/10 after 1/5/10 min respectively; birth weight 3,480 g, -0.7 SD). At three months of age, developmental delay was evident, characterized by delayed motor and communicative milestones with notably quiet and undemanding behavior. Over the course of the first year, in which she was hospitalized for pneumonia and hypovolemic shock resulting from rotavirus gastro-enteritis, she gradually developed secondary microcephaly (head circumference at birth 0 SD, at 1 year -2 SD). Brain MRI (age 2 years and 2 months; Fig. 1) showed hypoplasia of the corpus callosum, a reduced volume of supratentorial white matter with delayed myelination, dysplasia of basal ganglia, thalami and cerebellum, and an enlarged ventricular system, mainly of the lateral









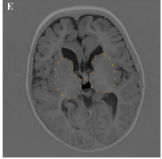


Fig. 1. Brain MRI displays structural brain abnormalities. Sagittal (A) and axial (C-E) T1-weighted and axial T2-weighted (B) MR images of the brain at age 2 years and 2 months show several structural brain abnormalities. There is general enlargement of the ventricular system, mostly affecting the lateral ventricles (indicated by green asterisks; panel A, B, D), but also the third ventricle (indicated by a yellow III; panel A) and the fourth ventricle (indicated by a blue IV; panel A), communicating with a megacisterna magna (indicated by a magenta star; panel A). The corpus callosum is hypoplastic (red arrowhead; panel A, D) and the cerebellum and cerebellar vermis show dysplasia, with abnormal morphology and orientation of folia (dotted white line; panel C). The basal ganglia and thalami are dysplastic, with an abnormal round appearance, and individual structures cannot be distinguished clearly (dashed orange line; panel D, E). Additionally, there is a reduced volume of supratentorial white matter (globally seen in panel B, D, E). . (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

ventricles (left more prominently enlarged than right), but also of the third and fourth ventricle, with the fourth ventricle communicating with an enlarged cisterna magna. There were no signs of cortical malformation. Electromyography and nerve conduction studies (age 2 years) did not show signs of an associated peripheral neuropathy or myopathy. There was no epilepsy. At ages 11 and 15 years, she was tested by SON-R 2.5–7, a nonverbal intelligence test, which showed a severe developmental delay and intellectual disability (chronological age 11 years and 11 months: developmental age of 2 years and 10 months; chronological age 15 years and 9 months: developmental age 2 years and 11 months, developmental age in fluid reasoning 3 years, developmental age at performance scale 2 years and

9 months). At 15 years of age, she had a vocabulary of 10 words, with a more developed receptive vocabulary than expressive vocabulary (WPPSI-III-NL 4–7;11 subscale General Language Composite (GLC) at chronological age 15 years and 9 months: developmental age 5 years and below 2 years and 7 months respectively). Other medical problems included high hypermetropia (+7D/+7.5D), convergent strabismus, short stature (–2.5SD), scoliosis, susceptibility to urinary and upper respiratory tract infections, constipation, factor V Leiden thrombophilia and behavioral problems (automutilation, sleep disturbances, phonophobia). She had minor facial dysmorphisms, consisting of a round face, with deep-set eyes and large palpebral fissures, a low nasal bridge and a prominent chin (Fig. 2), a mild pectus carinatum and

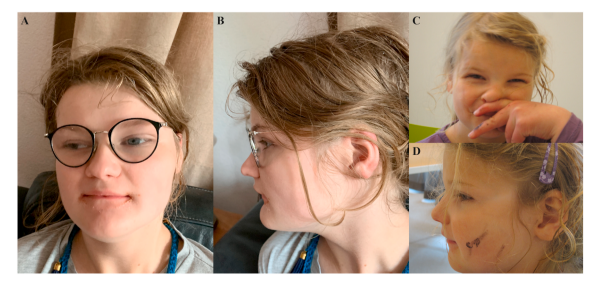


Fig. 2. Facial photographs of the proband. Frontal (panel A, C) and profile (panel B, D) facial photographs of the proband at age 17 years and 4 months (panel A, B) and at age 8 years and 2 months (panel C, D), showing mild facial dysmorphisms in the proband: she has a round face with deep-set eyes and large palpebral fissures, a low nasal bridge and a prominent chin.

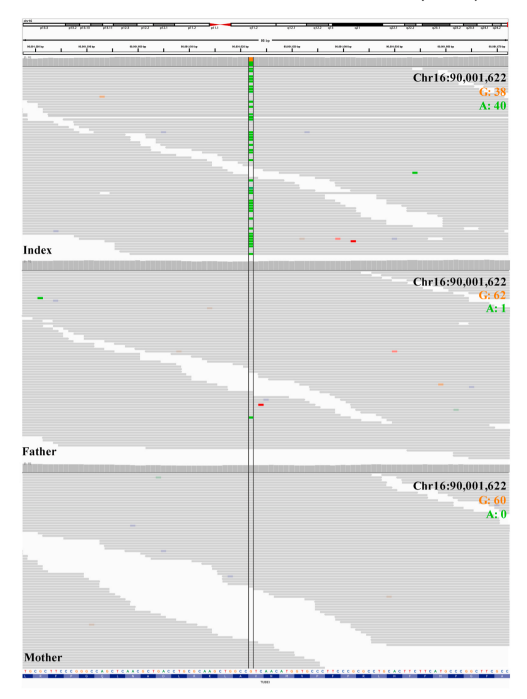


Fig. 3. IGV visualization of alignment files of index, father and mother. IGV visualization of alignment files of index (top), father (middle) and mother (bottom), showing the chr16:g.90001622G>A variant in the heterozygous state in the index (ref 38; alt 40), in one sequencing read of the father (ref 62; alt 1) and in no reads of the mother (ref 60; alt 0).

nipple retraction. Diagnostic evaluation consisted of karyotyping (at the age 1 of year), metabolic screening (age 1 year and 2 years), biochemical analysis of cerebrospinal fluid and blood (age 2 years), subtelomeric MLPA (age 2 years), 250K SNP array (age 4 years), targeted *GRIN2B* Sanger analysis (age 10 years), and trio-based ES (age 10 years), but did not result in an explanation for the phenotype.

3.2. Systematic reanalysis revealed a likely pathogenic TUBB3 variant

Data of the proband and parents were included in the Solve-RD project. Variant calling and annotation with the Solve-RD pipeline revealed a heterozygous *de novo* variant in *TUBB3*:Chr16(GRCh37/Hg19):g.90001622G>A;NM_006086.3:c.763G>A;p.(Val255Ile) in the

proband. The *TUBB3* missense variant was absent from gnomAD (v2.1.1) (Karczewski et al., 2019), had a CADD-PHRED score of 24.8 (Rentzsch et al., 2019) and *TUBB3* is highly intolerant to variation (Z-score missense 4.85; pLI 0.97) (Karczewski et al., 2019), all supporting variant pathogenicity. In addition, the variant was previously reported as "Likely pathogenic" in ClinVar (VCV000372654.1)(Landrum, 2018), but without any clinical description. Reanalysis did not identify other clinically relevant (*de novo*) variants.

Manual inspection of sequence alignment files in IGV clearly showed the presence of the variant in the proband (Fig. 3), which was subsequently confirmed by Sanger sequencing. Sanger analysis of both parental samples indicated the variant to be of *de novo* origin. However, the BAM file of the father showed the same variant in 1 of 63 reads (Fig. 3),

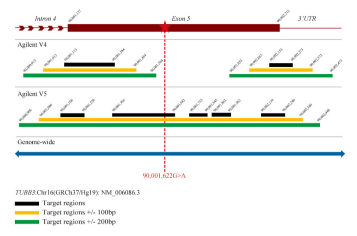


Fig. 4. Exon 5 of *TUBB3* and (extended) target regions used in variant calling. A schematic representation of the last part of *TUBB3*, including the variant identified in the proband. Shown in black are target regions of Agilent V4 and V5 enrichment kits, in orange target regions extended by 100bp on each side (the regions where sequence can be expected) and in green target regions extended by 200bp on each side (the regions used for variant calling in initial diagnostic analysis). The blue bar at the bottom visualizes genome-wide variant calling as done in the Solve-RD project. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

potentially indicative of low-level mosaicism (Wright et al., 2019). Due to the technical limitations of Sanger sequencing to detect such low-level mosaicism (Acuna-Hidalgo et al., 2015), we therefore continued with amplicon-based deep sequencing of the *TUBB3* variant on DNA derived from blood of the father to a sequence depth of 11,000-fold. This confirmed the absence of the variant in father (G: 99.89%; A: 0.05%; T: <0.01%; C: 0%, with percentages of A, T and C equal to background noise resulting from PCR artefacts), and its *de novo* origin in the proband.

3.3. The variant calling algorithm in initial ES analysis did not detect the TUBB3 variant

Next, we retrospectively analyzed why the TUBB3 variant was not identified before, as TUBB3 was included in the targets of the enrichment kit used for ES in this trio, and the gene was already described in literature in relation to intellectual disability and cortical dysplasia for 4 years at the time of initial diagnostic ES (Poirier et al., 2010). Additionally, in the diagnostic workflow used for individuals with intellectual disability, the interpretation of detected de novo variants in trio-based ES was performed invariably and with priority. Therefore, we first re-examined the output of the variant calling pipeline (annotated VCF) used in initial analysis and found that the variant was not called by the variant calling algorithm. Assessment of the regions considered in variant calling, showed that the variant is located in a region of the alignment not included in the extended targets for variant calling (target \pm 200bp), and therefore ignored by the variant calling algorithm (Fig. 4). Hence, we concluded that the combination of target-based calling and the exome enrichment kit used explains why the initial diagnostic ES did not lead to the detection of the TUBB3 variant.

3.4. Off-target covered regions disregarded in targeted variant calling coincide with known ID genes and previously reported pathogenic variants

To assess the overall impact of our conclusions, we checked the targets for *TUBB3* in Agilent V5, showing improved capture of *TUBB3*, with five additional targets compared to V4, of which three target exon 5 including the region of the variant (Fig. 4).

Next, we analyzed whether other regions of the coding sequence show off-target coverage but are disregarded in variant calling for both Agilent V4 and V5 enrichment kits (Table 1). First, we determined which protein coding sequences showed robust off-target coverage (≥20-fold), consistently present across multiple individuals (95% of 20 individuals), and then created the overlap with regions not included in targeted variant calling based on the extended targets (± 200 bp). This analysis detected a total of 840,347 and 761,594 nucleotides (1.95% and 1.76% of Gencode v31) in 1,537 and 1,298 of such regions, affecting 1,031 and 659 genes for Agilent V4 and V5, respectively. This included 36 established ID genes, including TUBB3 for V4, and 21 ID genes for V5. By intersecting the results of V4 with HGMD Professional (Stenson et al., 2003), we found that 262 unique (likely) pathogenic SNVs/indels (DM? 33; DM: 229) affect 17 of these 36 ID genes. To a lesser extent, this was also observed for V5, where we showed 71 (likely) pathogenic variants (DM? 6; DM: 65) to be located in these regions, affecting 8 of 21 ID genes (Table 1, Supplementary Table 1).

4. Discussion

Through systematic reanalysis in the Solve-RD project, we found a likely pathogenic missense variant in *TUBB3* (c.763G>A;p.(Val255Ile)) in a 16-year-old female proband with severe ID. The variant is causative to the proband's phenotype, which consists of severe developmental delay, strabismus, microcephaly and structural brain abnormalities and fits the

Table 1Genes and reported (likely) pathogenic variants in off-target covered regions disregarded in targeted variant calling.

Kit	Regions with ≥20x coverage in 95% of individuals	All genes			Genes associated with ID			
		# unique genes	# genes with previously reported (likely) pathogenic variants in these regions	# of reported unique (likely) pathogenic variants in these regions	# ID genes (% of all known ID genes)	# ID genes with previously reported (likely) pathogenic variants in these regions	# of reported unique (likely) pathogenic variants in these regions	
Agilent V4	1,537	1,031	133	884	36 (2.6%)	17	262	
Agilent V5	1,298	659	58	426	21 (1.5%)	8	71	
Overlap V	4 and V5	420	34	94	16 (1.1%)	5	13	

phenotypic spectrum of complex cortical dysplasia with other brain malformations-1 (CDCBM1; MIM#614039) (Poirier et al., 2010; Dentici et al., 2020; Fukumura et al., 2016; Shimojima et al., 2016). Although the proband does not exhibit the typical cortical dysplasia phenotype, the peculiar combination of several of the other structural abnormalities seen in brain imaging of the proband, specifically dysplasia of the cerebellum, thalami and basal ganglia, hypoplasia of the corpus callosum, and enlargement of the ventricular system, is common in patients affected by pathogenic *TUBB3* missense variants (Dentici et al., 2020).

This variant, which explains the phenotype in this individual, could only be identified by subjecting sequencing data to genome-wide variant calling. Diagnostic variant calling pipelines were limited to detect this variant in exon 5 of TUBB3, as exon 5 was not completely targeted by the Agilent V4 enrichment kit, and the prior pipeline was dependent on targeted-based variant calling ± 200 bp. Of note, the decision to use ± 200 bp extended target calling was already double the recommendation of ± 100 bp done by Agilent.

Despite not being fully targeted by the capture kit, exon 5 of TUBB3 in this proband-parents trio showed sufficient off-target coverage, with the variant clearly detectable in the proband, enabling the genome-wide variant caller in the Solve-RD pipeline to identify the variant. To determine whether our observation was an isolated case, or whether this phenomenon may occur for multiple ID genes, we assessed the impact at exome-wide level. Exome-wide analysis for two enrichment kits across 20 individuals showed that interpretable off-target coverage in proteincoding sequence is generated for >1,000 genes in V4, and \sim 650 genes in V5. The use of target-based variant calling algorithms (± 200 bp) would disregard at least part of the coding sequence of these genes. The clinical relevance of these genes is highlighted by the fact that for multiple ID genes disease-causing variants similar to this variant in TUBB3 have been reported in regions escaping variant calling. The impact may even be underestimated as we have used targets ± 200 bp in diagnostic variant calling, already calling more off-target bases than recommended.

As reanalysis of existing ES data is a process used more often for individuals who remained undiagnosed at first diagnostic ES trajectory, the use of genome-wide variant calling should be considered. In particular if data were generated using 'older' exome enrichment kits, as it is to be expected that more recent enrichment kits keep improving and aim to resolve the issue reported here. Whether genome-wide variant calling should be used as a generalized approach for exome data during the first diagnostic analysis, is more difficult to assess as this depends on multiple factors, including the capture kit used and the (computational) infrastructure available.

In summary, we describe a female proband diagnosed with a *de novo* missense variant in *TUBB3*, through systematic, large-scale reanalysis on a cohort of unexplained (syndromic) ID cases within Solve-RD. We show that performing new variant calling enables detection of variants missed by variant calling in initial analysis, stressing the relevance of sharing raw data, as alternative approaches of data-sharing and reanalysis would have failed to identify these (potentially causative) variants. As the same issue also occurs for other (ID) genes, especially in older enrichment kits, this implies that genome-wide variant calling, although computationally demanding, expands the potential of ES and may effectuate new diagnoses in other unexplained cases of ERN ITHACA.

Funding

This work was financially supported by Aspasia grants of the Dutch Research Council (015.014.036 to TK and 015.014.066 to LELMV) and Netherlands Organization for Health Research and Development (917.183.10 to TK). The Solve-RD project has received funding from the European Union's Horizon 2020 research and innovation programme under grant agreement No 779257.

Declaration of competing interest

The authors declare no conflict of interest.

Acknowledgements

We are very grateful to all families participating in the Solve-RD project and in particular to the proband and parents described in this clinical report. The collaborations in this study were facilitated by ERN ITHACA [EU Framework Partnership Agreement ID: 3HP-HP-FPA ERN-01-2016/739516, one of the 24 European Reference Networks (ERNs) approved by the ERN Board of Member States, co-funded by the European Commission. For more information about the ERNs and the EU health strategy please visit https://ec.europa.eu/health/ern.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.ejmg.2021.104402.

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