Genome-wide sequencing and the clinical diagnosis of genetic disease: The CAUSES study

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Summary

Genome-wide sequencing (GWS) is a standard of care for diagnosis of suspected genetic disorders, but the proportion of patients found to have pathogenic or likely pathogenic variants ranges from less than 30% to more than 60% in reported studies. It has been suggested that the diagnostic rate can be improved by interpreting genomic variants in the context of each affected individual’s full clinical picture and by regular follow-up and reinterpretation of GWS laboratory results. Trio exome sequencing was performed in 415 families and trio genome sequencing in 85 families in the CAUSES study. The variants observed were interpreted by a multidisciplinary team including laboratory geneticists, bioinformaticians, clinical geneticists, genetic counselors, pediatric subspecialists, and the referring physician, and independently by a clinical laboratory using standard American College of Medical Genetics and Genomics (ACMG) criteria. Individuals were followed for an average of 5.1 years after testing, with clinical reassessment and reinterpretation of the GWS results as necessary. The multidisciplinary team established a diagnosis of genetic disease in 43.0% of the families at the time of initial GWS interpretation, and longitudinal follow-up and reinterpretation of GWS results produced new diagnoses in 17.2% of families whose initial GWS interpretation was uninformative or uncertain. Reinterpretation also resulted in rescinding a diagnosis in four families (1.9%). Of the families studied, 33.6% had ACMG pathogenic or likely pathogenic variants related to the clinical indication. Close collaboration among clinical geneticists, genetic counselors, laboratory geneticists, bioinformaticians, and individuals’ primary physicians, with ongoing follow-up, reanalysis, and reinterpretation over time, can improve the clinical value of GWS.

Introduction

Genome-wide sequencing (GWS; exome sequencing [ES] or genome sequencing [GS]) has transformed the ability to diagnose patients with genetic diseases. Many studies of the diagnostic rate or clinical utility of GWS have used the finding of variants classified as “pathogenic” or “likely pathogenic” according to the American College of Medical Genetics and Genomics (ACMG) standards and guidelines1 to provide a surrogate “molecular diagnosis” of genetic disease in patients,2–5 although this practice is inconsistent with the ACMG guidelines, which state:

In general, a variant classified as pathogenic using the proposed classification scheme has met criteria informed by empirical data such that a health-care provider can use the molecular testing information in clinical decision making. Efforts should be made to avoid using this as the sole evidence of Mendelian disease; it should be used in conjunction with other clinical information when possible.

A more recent ACMG clinical practice guideline6 puts this in a clinical, rather than laboratory-focused, context:

Clinical genetic testing by ES/GS can assist clinicians in confirming or establishing a clinical diagnosis that may lead to changes in management, obviate the need for further testing, and/or end the diagnostic odyssey.

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Establishing the diagnosis of a genetic disease in an affected individual requires knowledge of that individual’s medical history and disease course over time, family history, physical examination, specialist physician evaluations, and imaging studies, as well as the results of GWS and other laboratory tests. GWS is by far the best test for disease-causing genetic variants that has ever been available, but genotype-phenotype correlation is essential for establishing the diagnosis of, rather than just the risk for, a genetic disease in an affected individual. A recent multidisciplinary consensus statement on the use of ES as a first-tier clinical diagnostic test for neurodevelopmental disorders put it this way: “Ultimately, final variant interpretation of the ES test result requires review by expert clinicians who can reassess the patient’s phenotype in light of the suggested molecular diagnosis.”

To avoid confusion, we shall eschew use of the term “diagnosis” or “molecular diagnosis” to describe a laboratory interpretation based only on standard ACMG variant classification and the limited phenotypic information typically provided on a test requisition. We shall refer to such laboratory-based interpretations as “variant classifications” and reserve use of the term “diagnosis” to mean our multidisciplinary research team’s (or, in practice, the affected individual’s physician’s) interpretation of GWS findings as probably or definitely disease-causing for a particular genetic condition in the context of all of the available clinical information about the affected individual.

Previous studies have found that interpretation of GWS variants by a multidisciplinary team with clinical as well as laboratory expertise and deep knowledge of each affected individual’s entire phenotype can produce a higher rate of genetic disease diagnosis than can be obtained through a laboratory report based on variant interpretation with limited clinical information alone.8–17 Other factors that have been associated with higher rates of genetic disease diagnosis include trio, rather than singleton, GWS;17,18 the use of GS rather than ES;14,17 and reanalysis or reinterpretation of GWS data years after the initial analysis. Substantial rates of variant reclassification, both upward to pathogenic/likely pathogenic and downward from pathogenic/likely pathogenic, have been reported with reinterpretation, and these changes may have a substantial impact on clinical diagnosis and management.17,19,20

Here we report our experience using GWS in a longitudinal study of 500 families of children with suspected genetic diseases. GWS results were interpreted by a multidisciplinary clinical research team, and individuals were followed for an average of 5.1 years after testing, with clinical reassessment and reinterpretation of the GWS results as necessary.

Material and methods

Ethics and consent
The CAUSES study was approved by the University of British Columbia-BC Children’s and Women’s Hospital Research Ethics Board (protocol H15-00092). Consent or assent was obtained from probands (when possible) and from their parents.

Recruitment of participants
Individuals were enrolled through a genomic consultation service to identify individuals for whom there was high suspicion of an underlying monogenic disorder that had not been established through conventional genetic testing.21,22 As the CAUSES GWS Study was trio based, the availability of both parents was required for enrollment. Almost all individuals were <19 years of age at the time of enrollment. An algorithm describing patient and sample flow in the CAUSES Research Study has previously been published.23 All individuals received pre- and post-test genetic counseling by an experienced genetic counselor, who also obtained informed consent for GWS, opting in or out for incidental findings for parents, genetic counseling research, and health economic evaluation of GWS.22,23 The approach to incidental findings followed the Canadian College of Medical Geneticists guidelines.24 Consent for data sharing through DECIPHER was obtained from each family, and parents were offered the option to have whole blood in excess of that needed for DNA extraction banked for future research on their child at the BC Children’s Hospital Biobank. For individuals who had not been previously evaluated by a clinical geneticist, a brief clinical examination was performed by a geneticist and phenotypic information was captured. Telemedicine was offered for the pre-test genetic counseling session for families who lived outside of the Vancouver area and had previously been seen by a clinical geneticist.25

Individuals who received GWS were classified into one of four major phenotypic categories: isolated intellectual disability (ID), syndromic ID, unexplained disorders of organ dysfunction without ID, and multiple congenital anomalies without ID. Information collected on all individuals included age, sex, self-described ethnicity, relevant family history, consanguinity, phenotypic findings, and previous investigations.

Sequencing and bioinformatic analysis
Genomic DNA was isolated from peripheral blood using standard techniques on each member of the trio/quad at the Division of Genome Diagnostics at BC Children’s Hospital. In total, 415 families underwent ES and 85 families underwent GS (81 of whom were selected for ID as part of a collaboration with Genomics England). Hybridization-based capture was used to enrich for exomes on each member of the trio. ES was performed on Illumina platforms at Ambry Genetics, Centogene, or Canada’s Michael Smith Genome Sciences Centre. Read alignment and single-nucleotide variant (SNV)/insertion or deletion (indel) calling were performed as previously described.26,27

PCR-free genome libraries were sequenced on an Illumina platform at Canada’s Michael Smith Genome Sciences Centre or the McGill Genome Centre.28 Read alignment and SNV/indel calling utilized BWA-0.7.6,29 Samtools-0.5.5,30 GATK HaplotypeCaller 3.5,31 BCFTools-1.9,28 and HTSlib-1.9.28 VarSeq (Golden Helix, Inc., Bozeman, MT, www.goldenhelix.com) was used for SNV/indel annotation and filtering. CNVnator-0.3.3,32 ERDS-1.1,33 LUMPY-0.2.13,34 and Manta-1.1.35 were used for SV calling. EverythingSV (https://github.com/FriedmanLab/StructuralVariantAnalysis), Samtools-1.5,28 SURVIVOR-1.0.2,36 and ANNOVAR-2018-04-16 were used for SV merging, annotation, and filtering. Annotation databases used for SVs included HPO,38 OMIM,39 and RefSeq-105.40
Interpretation of GWS

GWS FASTQ files were analyzed by a bioinformatician (genomic analyst) who generated a short list of the most promising candidate variants for each family. Criteria used for variant filtration were as follows:

1. Variants were included if their reference allele frequency was consistent with either dominant or recessive disorders.
2. Variants were included if they were predicted to have an impact on the protein's level, structure, or function or were recorded in ClinVar as “pathogenic.”
3. Variant prioritization was largely based on the trio family structure. Specifically, those variants identified as de novo, compound heterozygous, homozygous, or hemizygous in the proband were the main focus of analysis.
4. Variant prioritization also included a list of all variants with reference frequencies consistent with an autosomal recessive disorder and in genes known to be associated with a disorder. The aim of this list was to identify variants seemingly outside of the trio family structure filter that might still be disease causing as a result of imprinting, incomplete penetrance, variable expressivity, mosaicism in a parent, a parent being affected (but unknown at the time of intake), or a hemizygous variant called as heterozygous.

The complete, fully annotated list of selected variants was provided to the multidisciplinary research team for review; however, the genomic analyst flagged ClinVar “pathogenic” and predicted loss-of-function variants for particular attention. Each of the selected variants was discussed by a multidisciplinary research team that included genome analysts, MD clinical geneticists, genetic counselors, a PhD laboratory geneticist, and the referring physician. The team assigned a diagnostic category (“definitely causal,” “probably causal,” “uncertain,” or “uninformative”) for each individual. The categories were assigned by a consensus of clinical judgment based on the full clinical picture, including all of the information available on the variant(s) as well as the complete medical history, disease course over time, family history, physical examination findings, specialist consultations, imaging studies, and other laboratory test results. Individuals who had variants that were judged to be definitely or probably causal of the phenotype were considered to have been diagnosed with a genetic disease in our analysis.

The criteria that our multidisciplinary team used to determine if variants were probably or definitely disease-causing were independent of the ACMG variant classification. In fact, the clinical diagnostic categories were assigned before we obtained Sanger sequencing and standard ACMG classification of the variants through our clinical laboratory. Our bioinformatics analysis included annotation of all of the factors included in the ACMG classification, and this information was discussed by our multidisciplinary research team in the context of each individual’s complete clinical picture.

We adopted a convention of ≥99% certainty for “definitely causal” and ≥90% (but <99%) certainty for “probably causal” diagnoses. Individuals who had variants with bioinformatics evidence for pathogenicity but whose clinical features were not clearly compatible with those reported for the genotype in the published literature were categorized as “uncertain.” Individuals in whom our bioinformatics analysis found no variant that appeared to be causal for the clinical features were considered to have “uninformative” GWS. A research summary report was sent to the referring physician following the multidisciplinary team discussion.

Orthogonal validation of variants that were interpreted by our multidisciplinary research team as definitely or probably causal or that were uncertain but possibly causal was performed by Sanger sequencing through a clinical laboratory (Division of Genome Diagnostics at BC Children’s Hospital). The clinical laboratory independently assigned an ACMG classification to each variant and issued a standard report to the affected individual’s chart. Variants classified as “pathogenic” or “likely pathogenic” by the clinical laboratory were uploaded into the ClinVar database.

Definitely or probably causal variants were considered to be responsible for a “partial” diagnosis if the variant(s) appeared to explain only a portion of the individual’s phenotype. A dual diagnosis was assigned when probably or definitely causal variants of two different genetic loci were judged to have contributed to the individual’s phenotype. Variants that could not be classified as causal but were thought by the multidisciplinary team to be interesting candidates for further research were entered into GeneMatcher and considered for further study.

Post-test genetic counseling and reanalysis

The referring physician and CAUSES genetic counselor met with families of individuals in whom a diagnosis of genetic disease was made (i.e., whose GWS findings were judged to be probably or definitely causal of the clinical phenotype) to discuss these results when the clinical Sanger sequencing report was available. Families who did not receive a diagnosis of genetic disease through CAUSES were contacted by the genetic counselor, so informed, and told that reanalysis would occur periodically until the study end.

Variant call format files for GWS datasets from each family in whom no genetic diagnosis or only a partial diagnosis of genetic disease was made were reanalyzed through a Golden Helix VarSeq annotation and filtered workflow with the most up-to-date ClinVar and OMIM annotations every 1–2 years. Reanalysis was a planned part of the CAUSES project; it was not dependent on physician request. The approach used was similar to the primary analysis, with the focus on variants identified as de novo, compound heterozygous, homozygous, or hemizygous in the proband. Other variants were considered in the reanalysis only when important new clinical information had become available on the proband or the phenotype associated with the genetic locus had been expanded or characterized more fully in the published literature. The main focus of the routine reanalysis was to find variants for which the ACMG classification had changed to pathogenic or likely pathogenic in ClinVar or that occurred in genes that had been newly associated with a genetic disorder in OMIM or the literature. If the genomic analyst determined that results of the reanalysis might alter the diagnostic category (“definitely causal,” “probably causal,” “uncertain,” or “uninformative”) of an individual, it was reconsidered by the multidisciplinary research team and changed by consensus, if necessary. The reason for changing the diagnostic category was determined and recorded as a change in the bioinformatic pipeline, the emergence of a new disorder, a new publication, or the referring physician’s reinterpretation of the phenotype-genotype relationship. The genomic variants associated with such changes were Sanger sequenced in our clinical laboratory, reported through standard clinical protocols (including ACMG classification of variants), and returned to the affected individual’s medical record, referring physician, and family, as described above.
Referring physicians, who continued to follow their patients clinically; members of the CAUSES research team; or patients’ families (through recontact with CAUSES genetic counselors) could also request reanalysis of genomic variants or reinterpretation of CAUSES results by the multidisciplinary research team.

Statistical analysis
The Mantel-Haenszel test was used to compare the multidisciplinary team’s interpretation of the GWS results in each proband with the clinical laboratory’s ACMG classification of Sanger-sequenced variants. The cumulative probability of reinterpretation was calculated and plotted in IBM SPSS Statistics v.24 using the Kaplan-Meier module.

Results

CAUSES cohort demographics
We enrolled 500 families, including 531 children (probands and affected sibs). Trios, defined as two parents and one affected child, were usually studied. In 31 families, two affected sibs and both parents were studied; different combinations of affected and unaffected relatives were tested in a few other families. The mean age (± standard deviation) at referral of the children who received GWS was 8.0 (± 4.9) years. There were 246 females and 285 males. The individuals were ethnically diverse, with 48.5% of European, 16.2% of South Asian, 15.8% of East Asian, 4.3% of Middle Eastern, and 4.1% of First Nations ancestry. The most frequent indication for GWS was syndromic ID (85.1%), followed by multiple congenital anomalies without ID (5.3%), disorders of organ function (5.0%), and isolated ID (4.6%).

Diagnostic rate
ES was performed in 415 families and GS in 85. The CAUSES multidisciplinary research team diagnosed at least one genetic disorder in 261 (52.2%) of the families studied; 105 families were found to have variants that were probably causal and 156 families had variants that were considered to be definitely causal of a genetic disease in the child (Table S1). Of the 261 families diagnosed with at least one genetic disorder, 36 had variants that could not be classified according to the ACMG classification and 65 had variants that were classified as VUS. The rationale our multidisciplinary team used to diagnose a genetic disease in each of the individuals in whom an ACMG variant of uncertain significance or an unclassified variant was found is shown in Table 1.

Considering all families in which the CAUSES multidisciplinary research team diagnosed at least one genetic disorder, the diagnostic rate was 52.3% with ES and 51.8% with GS. In nine families, the probands had dual diagnoses (Table S2). Of the 261 probands who were diagnosed with a genetic disease, 219 had autosomal dominant (184 de novo), 27 autosomal recessive (one with isodisomy), and 13 X-linked recessive or X-linked dominant disorders. One proband had a Y-linked disorder, and one inherited an Xp25 genomic duplication from the mother. Partial diagnoses were established in 19 families (Table S3).

Affected sibs were tested in 31 of the families studied, and a diagnosis of genetic disease was established in 17 (55%) of these families by our multidisciplinary research team (Table S4). There were 11 sib pairs who were concordant for a definitely or probably causal variant associated with the individuals’ phenotypes. Six of these families exhibited autosomal recessive inheritance and three autosomal dominant inheritance, with parental mosaicism in two families and maternal inheritance in one. In addition, one sib pair was concordant for orofacial clefts and a paternally inherited PLEKHA7 variant and discordant for congenital NAD deficiency disorder 2/vertebral, cardiac, renal, and limb defects syndrome 2 (VCRL2; MIM: 617661) caused by compound heterozygosity for KYNU variants. In three families, the sibs were discordant for phenotype, and the diagnosis of a different genetic disease was established in each sib. In three other phenotypically discordant sib pairs, definitely or probably causal genetic variants were found in only one sib.

We reported incidental findings in one parent in 21 (4.4%) of the 478 families who opted for return of these results. Eight were pharmacogenomic variants (DPYD), and seven were cancer predisposition genes (BRCA2, BRCA1, BAP1, or CDK4). Single individuals had incidental findings in G6PD, LDLR, or APOB.

Diagnostic reinterpretation by the multidisciplinary team
In 4 (1.9%) of the 215 families initially diagnosed as having a genetic condition associated with a definitely or probably disease-causing genomic variant, our multidisciplinary research team reinterpreted the GWS findings as uncertain or uninformative as a result of additional information on the individual, gene, or variant that became available during the period of follow-up (Table 2). For individual G483, an ACMG pathogenic ALPL variant initially interpreted as definitely causal was reinterpreted as uncertain on the basis of a normal serum alkaline phosphatase level. In G369, a TRX1 variant initially interpreted as probably causal in a child with neurodevelopmental abnormalities consistent with velocardiofacial syndrome (VCFS; MIM: 192430) was reinterpreted as uncertain after an evaluation by a cardiologist was normal. In G550, a USP7 variant originally considered to be probably causal was reinterpreted as uncertain on the basis of a detailed physical examination by the referring physician.

An ATP2A2 variant in a child with global developmental delay, mild dysmorphic features, generalized hypotonia, and intention tremor (G103) was initially interpreted as definitely disease-causing but subsequently reinterpreted by our multidisciplinary research team as uncertain. This reinterpretation occurred when a de novo missense variant in DDX23, a gene that was not known to be disease associated at the time of initial analysis, was reinterpreted as probably disease-causing because of G103’s phenotypic...
| CAUSES ID no. | Sex | Gene | Variant | Mechanism | Disease | ACMG variant classification | Diagnostic interpretation by multidisciplinary research team | Rationale for diagnostic interpretation |
|--------------|-----|------|---------|-----------|---------|-----------------------------|-------------------------------------------------|---------------------------------------|
| G001-1       | F   | ASXL3| NP_085135.1:1.p. Asn1224Ter | de novo heterozygous | Bainbridge-Ropers syndrome | VUS | definitely causal | phenotype consistent with Bainbridge-Ropers syndrome; bioinformatics predict that variant is damaging |
| G004-1       | F   | TBCK | NP_001156907.2:p. Arg261Ser and NM_001163435.3:c.2060-2A>G | compound heterozygous | infantile hypotonia with psychomotor retardation and characteristic facies | VUS | probably causal | phenotype characteristic of reported cases; VUS predicted as damaging and allelic to likely pathogenic variant |
| G005-1       | F   | CAMK2G| NP_001354463.1:p. Arg297Trp | de novo heterozygous | mental retardation, autosomal dominant | unclassified | probably causal | phenotype consistent with reported cases; bioinformatics predict that variant is damaging |
| G010-1       | M   | SUZ12| NP_056170.2:p. Arg654Ter | de novo heterozygous | Imagawa-Matsumoto syndrome | unclassified | probably causal | phenotype consistent with reported cases; bioinformatics predict that variant is damaging |
| G022-1       | M   | NRXN1| NP_001317007.1:p.Ile382Met | de novo heterozygous | affective psychosis with severe obsessive compulsive disorder, onset at about 9 years of age, refractive to treatment; cognitive deterioration | VUS | probably causal | de novo variant predicted to cause haploinsufficiency; very unusual phenotype consistent with that observed in some individuals with NRXN1 haploinsufficiency reported with deletions |
| G025-1       | M   | CLTC | NP_004830.1:p. Pro890Leu | de novo heterozygous | mental retardation, autosomal dominant | unclassified | probably causal | phenotype consistent with reported cases; bioinformatics predict that variant is damaging |
| G027-1       | M   | SMC1A| NP_006297.2:p. Asp982Val | hemizygous (inherited from mosaic mother) | Cornelia de Lange syndrome 2 | VUS | probably causal | phenotype consistent with Cornelia de Lange syndrome; bioinformatics predict that variant is damaging |

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| CAUSES ID no. | Sex | Gene     | Variant               | Mechanism                          | Disease                    | ACMG variant classification | Diagnostic interpretation by multidisciplinary research team | Rationale for diagnostic interpretation |
|--------------|-----|----------|-----------------------|------------------------------------|----------------------------|-----------------------------|-------------------------------------------------------------|----------------------------------------|
| G035-1       | M   | MED12    | NP_005111.2:p.Arg911Leu | hemizygous (maternally inherited) | Ohdo syndrome             | VUS                         | probably causal                                               | phenotype consistent with Ohdo syndrome; similarly affected maternal uncle also carries variant; bioinformatics predict that variant is damaging |
| G036-1       | M   | SATB2    | NM_001172509.2:c.474-3C>G | de novo heterozygous             | Glass syndrome            | VUS                         | probably causal                                               | phenotype consistent with reported cases; bioinformatics predict that variant alters splicing |
| G044-1       | F   | SCN8A    | NP_001317189.1:p.Cys324Tyr | de novo heterozygous             | early infantile epileptic encephalopathy | VUS                         | probably causal                                               | phenotype consistent with reported cases; bioinformatics predict that variant is damaging |
| G050-1       | M   | EP300    | NP_001420.2:p.Gly2350HisfsTer52 | heterozygous (maternally inherited) | Rubinstein-Taybi syndrome 2 | VUS                         | definitely causal                                               | phenotype in both proband and mother consistent with Rubinstein-Taybi syndrome; bioinformatics predict that variant is damaging |
| G053-1       | M   | KCNQ5    | NP_0062816.2:p.Val145Gly  | de novo heterozygous             | autosomal dominant mental retardation 46 | unclassified                | definitely causal                                               | individual included in first published report of “new” genotype-phenotype association |
| G059-1       | M   | SLC16A2  | NP_006508.2:p.Arg371Leu  | hemizygous (maternally inherited) | Allan-Herndon-Dudley syndrome | VUS                         | definitely causal                                               | phenotype characteristic of Allan-Herndon-Dudley syndrome |
| G066-1       | M   | UBE2A    | NP_003327.2:p.Arg95Cys    | hemizygous (maternally inherited) | mental retardation, X-linked syndromic, Nascimento type | VUS                         | probably causal                                               | phenotype in both affected brothers consistent with reported cases; bioinformatics predict that variant is damaging |
| G066-4       | M   | UBE2A    | NP_003327.2:p.Arg95Cys    | hemizygous (maternally inherited) | mental retardation, X-linked syndromic, Nascimento type | VUS                         | probably causal                                               | phenotype in both affected brothers consistent with reported cases; bioinformatics predict that variant is damaging |

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| CAUSES ID no. | Sex | Gene | Variant | Mechanism | Disease | ACMG variant classification | Diagnostic interpretation by multidisciplinary research team | Rationale for diagnostic interpretation |
|------------|-----|------|---------|-----------|---------|-----------------------------|-------------------------------------------------|-------------------------------------|
| G070-1 M   | C2orf69 | NP_710156.3:p.Lys282GlnfsTer55 | homozygous | combined oxidative phosphorylation deficiency 53 | VUS | probably causal | phenotype consistent with reported cases; consanguineous family with two early infant deaths in cousins and two additional early childhood deaths in other relatives; bioinformatics predict that variant is damaging |
| G073-1 M   | SOD1 | NP_000445.1:p.Ala124del | homozygous | progressive spastic tetraplegia and axial hypotonia | unclassified | probably causal | very characteristic phenotype consistent with reported cases; similarly affected sib died in childhood; bioinformatics predict that variant is damaging |
| G075-1 F   | MAST1 | NP_055790.1:p.Gly98Val | de novo heterozygous | mega-corpus-callosum syndrome with cerebellar hypoplasia and cortical malformations | unclassified | probably causal | individual included in first published report of "new" genotype-phenotype association |
| G077-1 F   | BPTF | NP_872579.2:p.Arg655Ter | heterozygous (paternally inherited) | neurodevelopmental disorder with dysmorphic faces and distal limb anomalies | unclassified | probably causal | phenotype consistent with reported cases; bioinformatics predict that variant is damaging |
| G089-1 M   | NF1 | NP_001035957.1:p.Gly1190Val | heterozygous (maternally inherited) | neurofibromatosis 1 | VUS | probably causal | phenotype consistent with neurofibromatosis 1; bioinformatics predict that variant is damaging |
| G091-1 F   | TKT | NP_001055.1:p.Arg401His | compound heterozygous | short stature, developmental delay, and congenital heart defects | VUS | definitely causal | phenotype in both sibs consistent with reported cases; bioinformatics predict that both variants are damaging; biochemical assay demonstrates transketolase deficiency |
| G091-1 F   | TKT | NP_001055.1:p.Tyr564del | compound heterozygous | short stature, developmental delay, and congenital heart defects | VUS | definitely causal | phenotype in both sibs consistent with reported cases; bioinformatics predict that both variants are damaging; biochemical assay demonstrates transketolase deficiency |

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| CAUSES ID no. | Sex | Gene | Variant | Mechanism | Disease | ACMG variant classification | Diagnostic interpretation by multidisciplinary research team | Rationale for diagnostic interpretation |
|-------------|-----|------|---------|-----------|---------|-----------------------------|-------------------------------------------------------------|----------------------------------------------------------------------------------|
| G091-4 M    | TKT | NP_001055.1:p.Arg401His/np_001055.1:p.Tyr564del | compound heterozygous | short stature, developmental delay, and congenital heart defects | VUS | definitely causal | phenotype in both sibs consistent with reported cases; bioinformatics predict that both variants are damaging; biochemical assay demonstrates transketolase deficiency |
| G092-1 M    | PAK3 | NP_002569.1:p.Ser105Cys | hemizygous (maternally inherited) | mental retardation, X-linked 30/47 | VUS | probably causal | phenotype consistent with reported cases; bioinformatics predict that variant is damaging |
| G103-1 F    | DDX23 | NP_004809.2:p.Ang754Cys | de novo heterozygous | DDX23-related disorder | unclassified | probably causal | phenotype consistent with reported cases; bioinformatics predict that variant is damaging |
| G105-1 M    | SMS | NP_004586.2:p.Met233Ile | hemizygous (maternally inherited) | X-linked mental retardation; Snyder-Robinson type | VUS | definitely causal | phenotype characteristic of Snyder-Robinson syndrome; bioinformatics predict that variant is damaging |
| G107-1 M    | SMARCA2 | NP_003061.3:p.Asp1571GlufsTer46 | de novo heterozygous | Nicolaides-Baraister syndrome | VUS | probably causal | phenotype characteristic of Nicolaides-Baraister syndrome; bioinformatics predict that variant is damaging |
| G114-1 F    | REST | NP_005603.3:p.Gln827Ter | de novo heterozygous | gingival fibromatosis | VUS | probably causal | phenotype characteristic of gingival fibromatosis; variant in last exon, where other variants that cause gingival fibromatosis lie; bioinformatics predict that variant is damaging |
| G117-1 F    | PIGG | NP_001120650.1:p.Asn138Ser | homozygous | autosomal recessive mental retardation 53 | VUS | probably causal | both sibs included in first published report of “new” genotype-phenotype association |
| G117-4 M    | PIGG | NP_001120650.1:p.Asn138Ser | homozygous | autosomal recessive mental retardation 53 | VUS | probably causal | both sibs included in first published report of “new” genotype-phenotype association |

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| CAUSES ID no. | Sex | Gene | Variant | Mechanism | Disease | ACMG variant classification | Diagnostic interpretation by multidisciplinary research team | Rationale for diagnostic interpretation |
|--------------|-----|------|---------|-----------|---------|-----------------------------|--------------------------------------------------------------|----------------------------------------------------------------------------------|
| G122-1       | F   | SRCAP| NP_006653.2:p.Val1835ProfsTer13 | de novo heterozygous | non-Floating-Harbor syndrome SRCAP-related neurodevelopmental disability | unclassified | probably causal | phenotype consistent with non-Floating-Harbor syndrome SRCAP-related neurodevelopmental disability; variant lies in exon 2S, outside of exons 33–34 involved in Floating-Harbor syndrome; bioinformatics predict that variant is damaging |
| G125-1       | M   | ZFYVE26| NP_056161.2:p.Lys741ArgfsTer3 | compound heterozygous | autosomal recessive spastic paraplegia 15 | likely pathogenic | probably causal | phenotype characteristic of reported cases; compound heterozygote with one likely pathogenic variant and allelic very rare |
|              |     |      | NP_056161.2:p.Arg2140Gln | VUS | | | | |
| G134-1       | F   | KCNQ2| NP_742105.1:p.Arg144Trp | de novo heterozygous | early infantile epileptic encephalopathy 7 | unclassified | definitely causal | phenotype consistent with reported cases; bioinformatics predict that variant is damaging |
| G141-1       | F   | KMT2C| NP_733751.2:p.Pro4843AlafsTer12 | de novo heterozygous | Kleefstra syndrome 2 | unclassified | probably causal | phenotype consistent with Kleefstra syndrome; bioinformatics predict that variant is damaging |
| G160-1       | M   | CIC | NP_001373227.1:p.Ala20156ProfsTer3 | de novo heterozygous | autosomal dominant mental retardation 45 | unclassified | probably causal | phenotype consistent with reported cases; bioinformatics predict that variant is damaging |
| G169-1       | M   | HSD17B3| NP_000188.1:p.Ala273Val | compound heterozygous | pseudohermaphroditism, male, with gynecomastia | pathogenic | probably causal | phenotype consistent with reported cases; compound heterozygote with one likely pathogenic variant and allelic rare | VUS predicted to be damaging |
|              |     |      | CTLA4 | heterozygous (maternally inherited) | autoimmune lymphoproliferative syndrome, type V | VUS | probably causal | phenotype characteristic of autoimmune lymphoproliferative syndrome; bioinformatics predict that variant is damaging |

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| CAUSES ID no. | Sex | Gene | Variant | Mechanism | Disease | ACMG variant classification | Diagnostic interpretation by multidisciplinary research team | Rationale for diagnostic interpretation |
|---------------|-----|------|---------|-----------|---------|-----------------------------|----------------------------------------------------------|----------------------------------------|
| G174-1        | M   | SET  | NP_003002.2:p.Arg44LeufsTer10 | heterozygous (maternally inherited) | autosomal dominant mental retardation 58 | unclassified | probably causal | individual included in first published report of “new” genotype-phenotype association |
| G175-1        | M   | ACTB | NP_001092.1:p.Leu110ArgfsTer10 | de novo heterozygous | Baraister-Winter syndrome 1 | VUS | probably causal | phenotype characteristic of Baraister-Winter syndrome; bioinformatics predict that variant is damaging |
| G194-1        | M   | MN1  | NP_002421.3:p.Trp1248Ter | de novo heterozygous | CEBALID syndrome | VUS | definitely causal | phenotype characteristic of CEBALID syndrome; bioinformatics predict that variant is damaging |
| G198-1        | M   | ARID2| NC_000012.11:g.46298857_46302229del | de novo heterozygous | Coffin-Siris syndrome 6 | VUS | definitely causal | phenotype consistent with Coffin-Siris syndrome; bioinformatics predict that deletion is damaging |
| G202-1        | M   | DLG4 | NP_001308004.1:p.Asn187ThrfsTer3 | de novo heterozygous | intellectual developmental disorder 62 | unclassified | probably causal | phenotype consistent with reported cases; bioinformatics predict that variant is damaging |
| G205-1        | F   | NBEA | NP_001371941.1:p.Glu2433ArgfsTer3 | de novo heterozygous | neurodevelopmental disorder with or without early-onset generalized epilepsy | unclassified | probably causal | individual included in first published report of “new” genotype-phenotype association |
| G216-1        | F   | COL12A1 | NM_004370.6:c.8319+1G>T | heterozygous (maternally inherited) | Bethlem myopathy | VUS | probably causal | phenotype consistent with Bethlem myopathy; bioinformatics predict that deletion is damaging; variant segregates with disease in family |
| G217-1        | M   | KAT6B | NP_036462.2:p.Arg153Gln | de novo heterozygous | SBYSS syndrome | VUS | probably causal | phenotype consistent with SBYSS syndrome; bioinformatics predict that variant is damaging |
| G218-1        | M   | ASH1L| NP_060995.2:p.Arg2691Ter | de novo heterozygous | autosomal dominant mental retardation 52 | unclassified | probably causal | phenotype consistent with reported cases; bioinformatics predict that variant is damaging |

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| CAUSES ID no. | Sex | Gene | Variant | Mechanism | Disease | ACMG variant classification | Diagnostic interpretation by multidisciplinary research team | Rationale for diagnostic interpretation |
|--------------|-----|------|---------|-----------|---------|-----------------------------|----------------------------------------------------------|------------------------------------------|
| G231-1       | F   | WDR26| NP_001366332.1:p.Ala150Val | de novo heterozygous | Skraban-Deardorff syndrome | VUS | probably causal | phenotype consistent with Skraban-Deardorff syndrome; bioinformatics predict that variant is damaging |
| G235-1       | M   | TRAF7| NP_115647.2:p.Arg524Trp | de novo heterozygous | cardiac, facial, and digital anomalies with developmental delay | unclassified | probably causal | phenotype consistent with reported cases; bioinformatics predict that variant is damaging |
| G239-1       | M   | HNRNPU| NP_114032.2:p.His451Pro | de novo heterozygous | early infantile epileptic encephalopathy 54 | unclassified | probably causal | phenotype consistent with reported cases; bioinformatics predict that variant is damaging |
| G241-1       | M   | SZT2 | NP_001352928.1:p.Glu2560SerfsTer92 | compound heterozygous | early infantile epileptic encephalopathy 18 | likely pathogenic | probably causal | phenotype consistent with reported cases; likely pathogenic variant allelic to VUS; bioinformatics predict that both variants are damaging |
| G248-1       | F   | CRYBA2| NP_476434.1:p.Gly65Arg | heterozygous (paternally inherited) | autosomal dominant cataract 42 | unclassified | probably causal | phenotype and family history characteristic of autosomal dominant congenital cataracts; variant segregates with cataracts in family; bioinformatics predict that variant is damaging |
| G256-1       | F   | KAT6A| NP_006757.2:p.Phe933Ser | de novo heterozygous | autosomal dominant mental retardation 32 | VUS | probably causal | phenotype consistent with reported cases; bioinformatics predict that variant may be damaging |
| G259-1       | F   | WDR45| NM_001029896.2:c.436+5G>C | de novo heterozygous | neurodegeneration with brain iron accumulation 5 | VUS | definitely causal | phenotype characteristic of neurodegeneration with brain iron accumulation; bioinformatics predict that variant is damaging |
| G260-1       | M   | MECP2| NP_001104262.1:p.Gln256Glu | de novo hemizygous | X-linked intellectual disability disorder, Lubs type | VUS | probably causal | phenotype consistent with reported cases; bioinformatics predict that variant is damaging |

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| CAUSES ID no. | Sex | Gene | Variant | Mechanism | Disease | ACMG variant classification | Diagnostic interpretation by multidisciplinary research team | Rationale for diagnostic interpretation |
|--------------|-----|------|---------|-----------|---------|-----------------------------|----------------------------------------------------------|----------------------------------------|
| G272-1       | M   | SETD1B | NM_001353345.2:c.5589+1G>A | de novo heterozygous | intellectual developmental disorder with seizures and language delay | unclassified | probably causal | phenotype consistent with reported cases; bioinformatics predict that variant is damaging |
| G280-1       | F   | CWF19L1 | NP_060764.3:p. Glu384Ter | compound heterozygous | autosomal recessive spinocerebellar ataxia 17 | likely pathogenic | probably causal | phenotype consistent with autosomal recessive spinocerebellar ataxia; likely pathogenic variant allelic to VUS; bioinformatics predict that both variants are damaging |
| G284-1       | F   | ABL1  | NP_005148.2:p. Thr117Met | de novo heterozygous | congenital heart defects and skeletal malformation syndrome | VUS | probably causal | phenotype consistent with reported cases; bioinformatics predict that variant is damaging; functional studies support effect |
| G285-1       | F   | GLRX5 | NP_057501.2:p. Met128Thr | homozygous | childhood-onset spasticity with hyperglycinemia | VUS | definitely causal | phenotype consistent with reported cases; bioinformatics predict that variant is damaging; biochemical studies consistent |
| G286-1       | M   | IQCE  | NP_689771.3:p. Asp112ValfsTer2 | compound heterozygous | post-axial polydactyly type A7 | unclassified | probably causal | phenotype characteristic of post-axial polydactyly; bioinformatics predict that both variants are damaging |
| G289-1       | F   | CLCN4 | NP_001821.2:p. Gly182Ser | de novo heterozygous | Raynaud-Claes syndrome | VUS | probably causal | phenotype characteristic of Raynaud-Claes syndrome; bioinformatics predict that variant is damaging |
| G291-1       | F   | KDM5A | NC_000012.11:g.460661_470642del | homozygous | autosomal recessive mental retardation 65 | VUS | probably causal | phenotype consistent with reported cases; bioinformatics predict that homozygous variant is damaging; homozygous variant also found in similarly affected sib |

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| CAUSES ID no. | Sex | Gene | Variant | Mechanism | Disease | ACMG variant classification | Diagnostic interpretation by multidisciplinary research team | Rationale for diagnostic interpretation |
|--------------|-----|------|---------|-----------|---------|-----------------------------|-------------------------------------------------------------|--------------------------------------------------------------------------------------|
| G291-4       | M   | KDM5A| NC_000012.11:g.460661_470642del | homozygous | autosomal recessive mental retardation 65 | VUS | probably causal | phenotype consistent with reported cases; bioinformatics predict that homozygous variant is damaging; homozygous variant also found in similarly affected sib |
| G292-1       | M   | SYT1 | NP_005630.1:p.Pro180Leu | de novo heterozygous | Baker-Gordon syndrome | unclassified | probably causal | phenotype consistent with Baker-Gordon syndrome; bioinformatics predict that variant is damaging |
| G297-1       | M   | JARID2| NP_004964.2:p.Arg1127Ter | de novo hemizygous | JARID2-neurodevelopmental syndrome | unclassified | probably causal | individual included in first published report of "new" genotype-phenotype association |
| G312-1       | F   | NAA10| NP_003482.1:p.Asn3101Lys | de novo heterozygous | Ogden syndrome | VUS | probably causal | phenotype consistent with Ogden syndrome; bioinformatics predict that variant is damaging |
| G323-1       | M   | GABBR2| NP_005449.5:p.Pro282Leu | de novo heterozygous | neurodevelopmental disorder with poor language and loss of hand skills | VUS | probably causal | phenotype consistent with reported cases; bioinformatics predict that variant is damaging |
| G336-1       | M   | KCNB1| NP_004966.1:p.Glu71Ter | heterozygous (maternally inherited) | early infantile epileptic encephalopathy 26 | VUS | probably causal | phenotype and family history consistent with reported cases; bioinformatics predict that variant is damaging |
| G338-1       | F   | KYNU | NP_003928.1:p.Lys121del; NP_003928.1:p.Ser345Arg | compound heterozygous | vertebral, cardiac, renal, and limb defects syndrome 2 | VUS | probably causal | phenotype characteristic of vertebral, cardiac, renal, and limb defects syndrome; variants are allelic and bioinformatics predict that both are damaging; functional studies demonstrated significant reduction in NAD levels |
|             |     |      | PLEKHA7 | heterozygous (paternally inherited) | Mendelian nonsyndromic cleft lip with or without cleft palate | unclassified | probably causal | phenotype consistent with reported cases; both sibs have orofacial clefting and variant; bioinformatics predict that variant is damaging |

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| CAUSES ID no. | Sex | Gene       | Variant                  | Mechanism                                      | Disease                                      | ACMG variant classification | Diagnostic interpretation by multidisciplinary research team | Rationale for diagnostic interpretation |
|---------------|-----|------------|--------------------------|-----------------------------------------------|----------------------------------------------|------------------------------|-------------------------------------------------------------|------------------------------------------|
| G338-4        | F   | PLEKHA7    | NP_001316559.1:p.Asp191Asn| heterozygous (paternally inherited)           | Mendelian non-syndromic cleft lip with or without cleft palate | unclassified                | probably causal                                              | phenotype consistent with reported cases; both sibs have orofacial clefting and variant; bioinformatics predict that variant is damaging |
| G345-1        | M   | HISG1H1E   | NP_005312.1:p.Gly124ArgfsTer71 | de novo heterozygous                          | Rahman syndrome                            | VUS                          | definitely causal                                             | phenotype consistent with Rahman syndrome; bioinformatics predict that variant is damaging |
| G350-1        | F   | TSC2       | NP_000539.2:p.His1543Arg  | heterozygous (paternally inherited)           | tuberous sclerosis 2                       | VUS                          | definitely causal                                             | phenotype characteristic of tuberous sclerosis; bioinformatics predict that variant is damaging |
| G356-1        | F   | COL4A3BP   | NP_001365958.1:p.Thr251Ala| de novo heterozygous                          | autosomal dominant mental retardation 34    | unclassified                | probably causal                                              | phenotype consistent with reported cases; bioinformatics predict that variant is damaging |
| G363-1        | M   | FLNA       | NM_00110556.2:c.4475+1G>T | hemizygous (maternally inherited)             | periventricular nodular heterotopia 1       | VUS                          | probably causal                                              | phenotype consistent with periventricular nodular heterotopia; bioinformatics predict that variant is damaging |
| G368-1        | M   | KDM5C      | NP_004178.2:p.Ser285Leu   | hemizygous (maternally inherited)             | X-linked syndromic mental retardation, Claes-Jensen type | VUS                          | probably causal                                              | phenotype consistent with X-linked syndromic mental retardation, Claes-Jensen type; bioinformatics predict that variant may be damaging |
| G370-1        | M   | RYR1       | NP_000531.2:p.Glu2987Gly  | compound heterozygous                         | King-Denborough syndrome                    | VUS                          | probably causal                                              | phenotype characteristic of King-Denborough syndrome; variants are allelic and bioinformatics predict that both are damaging |
| G385-1        | M   | CAMK2      | NP_057065.2:p.Ser341Thr   | de novo heterozygous                          | autosomal dominant mental retardation 53    | VUS                          | probably causal                                              | phenotype consistent with reported cases; bioinformatics predict that variant is damaging |

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| CAUSES ID no. | Sex | Gene | Variant | Mechanism | Disease | ACMG variant classification | Diagnostic interpretation by multidisciplinary research team | Rationale for diagnostic interpretation |
|--------------|-----|------|---------|-----------|---------|----------------------------|---------------------------------------------------------------|------------------------------------------|
| G392-1       | M   | KCNK9| NP_001269463.1:p.p.Tyr205Cys | heterygous (maternally inherited) | Birk-Barel mental retardation dysmorphism syndrome | VUS | probably causal | phenotype and family history consistent with Birk-Barel mental retardation dysmorphism syndrome; bioinformatics predict that variant is damaging |
| G393-1       | M   | ERCC8| NP_000073.1:p.p.Val622PhefsTer20 | homozygous | Cockayne syndrome, type A | VUS | definitely causal | phenotype characteristic of Cockayne syndrome; parental consanguinity; bioinformatics predict that variant is damaging |
| G396-1       | M   | CHD3 | NP_001005273.1:p.p.Arg1172Gln | de novo heterozygous | Snijders Blok-Campeau syndrome | unclassified | definitely causal | phenotype characteristic of Snijders Blok-Campeau syndrome; bioinformatics predict that variant is damaging |
| G401-1       | F   | CTCF | NP_006556.1:p.p.Asp357Asn | de novo heterozygous | autosomal dominant mental retardation 21 | VUS | probably causal | phenotype consistent with reported cases; bioinformatics predict variant is damaging |
| G402-4       | F   | SMARCC2| NP_001317217.1:p.p.Tyr679Ter | de novo heterozygous | Coffin-Siris syndrome 8 | VUS | probably causal | phenotype consistent with reported cases; bioinformatics predict variant is damaging |
| G404-1       | M   | HNRNPU| NP_114032.2:p.p.Pnu506Leu | de novo heterozygous | early infantile epileptic encephalopathy 54 | VUS | probably causal | phenotype consistent with reported cases; bioinformatics predict variant is damaging |
| G406-1       | F   | H3F3A| NP_002098.1:p.p.Thr23Ile | de novo heterozygous | Bryant-Li-Bhoj neurodevelopmental syndrome 1 | VUS | probably causal | individual included in first published report of "new" genotype-phenotype association |
| G407-1       | M   | JARID2| NP_004964.2:p.p.Ile557ArgfsTer34 | de novo heterozygous | JARID2-neurodevelopmental disorder | unclassified | probably causal | individual included in first published report of "new" genotype-phenotype association |
| G421-1       | F   | NEUROD2| NP_006151.3:p.p.Glu130Iys | de novo heterozygous | early infantile epileptic encephalopathy 72 | unclassified | probably causal | phenotype consistent with reported cases; bioinformatics predict variant is damaging |

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| ID no. | Sex | Gene | Variant | Mechanism | Disease | ACMG variant classification | Diagnostic interpretation by multidisciplinary research team | Rationale for diagnostic interpretation |
|-------|-----|------|---------|-----------|---------|-----------------------------|-------------------------------------------------------------|----------------------------------------------------------------------------------|
| G422-1 | M   | DLL1 | NP_005609.3:p.Arg509Ter | heterozygous (paternally inherited) | neurodevelopmental disorder with non-specific brain abnormalities and with or without seizures | unclassified | definitely causal | individual included in first published report of “new” genotype-phenotype association |
| G447-1 | M   | SLC6A8 | NP_005620.1:p.Phe248del | de novo hemizygous (mosaic) | cerebral creatine deficiency syndrome 1 | VUS | definitely causal | phenotype consistent with reported cases; bioinformatics predict variant is damaging |
| G462-1 | F   | ASH1L | NP_060959.2:p.Glu1956Lys | de novo heterozygous | autosomal dominant mental retardation 52 | VUS | probably causal | phenotype consistent with reported cases; bioinformatics predict variant is damaging |
| G465-1 | F   | ANKR17 | NP_115593.3:p.Gln1787ArgfsTer5 | de novo heterozygous | Chopra-Amiel-Gordon syndrome | unclassified | definitely causal | individual included in first published report of “new” genotype-phenotype association |
| G468-1 | F   | GNAO1 | NP_066268.1:p.Asp151Asn | heterozygous (inherited from mosaic mother) | early infantile epileptic encephalopathy 17 | VUS | probably causal | phenotype consistent with reported cases; bioinformatics predict variant is damaging |
|       |     | PTCH1 | NM_000264.5:c.654+3A>G | de novo heterozygous | basal cell nevus syndrome | VUS | probably causal | phenotype consistent with basal cell nevus syndrome; bioinformatics predict variant is damaging |
| G472-1 | F   | GNB1 | NP_002065.1:p.Gly282Arg | de novo heterozygous | autosomal dominant mental retardation 42 | VUS | probably causal | phenotype consistent with reported cases; bioinformatics predict variant is damaging |
| G480-1 | M   | MSX1 | NP_002439.2:p.Trp219Arg | heterozygous (paternally inherited) | orofacial cleft 5 | VUS | probably causal | phenotype and family history typical of hereditary orofacial clefting; bioinformatics predict variant is damaging; variant segregates with phenotype in family |
| G482-1 | F   | SETD1B | NP_001340274.1:p.Gln1322Ter | de novo heterozygous | intellectual disability, epilepsy, and autism | unclassified | probably causal | phenotype consistent with reported cases; bioinformatics predict variant is damaging |

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| CAUSES ID no. | Sex | Gene | Variant | Mechanism | Disease | ACMG variant classification | Diagnostic interpretation | Rationale for diagnostic interpretation |
|--------------|-----|------|---------|-----------|---------|-----------------------------|--------------------------|----------------------------------------|
| G487-1       | F   | DDX3X| NP_001347.3:p. Ile190Ser | de novo heterozygous | X-linked mental retardation 102 | VUS | definitely causal | phenotype consistent with reported cases; bioinformatics predict variant is damaging |
| G494-1       | M   | EDEM3| NP_079467.3:p. Arg314Ser | homozygous (maternally inherited isodisomy) | EDEM3-related disorder | unclassified | probably causal | individual included in first published report of "new" genotype-phenotype association |
| G498-1       | F   | TANC2| NP_079461.2:p. Arg1770Gly | de novo heterozygous | global developmental delay, cerebellar atrophy, and dysmorphic features (non-clinome) | unclassified | probably causal | phenotype consistent with reported cases; bioinformatics predict variant is damaging |
| G504-1       | F   | TRRAP| NP_001362453.1:p.Thr10Met | de novo heterozygous | developmental delay with or without dysmorphic facies and autism | unclassified | probably causal | individual included in first published report of "new" genotype-phenotype association |
| G508-1       | M   | ERCC2| NP_000391.1:p. Leu581Pro | Compound heterozygous | trichothiodystrophy | pathogenic | definitely causal | phenotype consistent with trichothiodystrophy; VUS allelic to pathogenic variant and predicted to be damaging |
|             |     |      | NP_000391.1:p. Arg658Cys |           |         |                             |                          |                                        |
| G536-1       | M   | NF1  | NM_001042492.3:c.5609+1G>T | de novo heterozygous | neurofibromatosis 1 | unclassified | probably causal | phenotype consistent with neurofibromatosis 1; variant predicted as damaging; RNA studies demonstrated disruption of canonical splice site |
| G553-1       | F   | TRIP12| NP_001335252.1:p.Tyr1744Asp | heterozygous (inherited from mosaic father) | autosomal dominant mental retardation 49 | VUS | probably causal | phenotype consistent with reported cases; bioinformatics predict variant is damaging |
| G553-4       | M   | TRIP12| NP_001335252.1:p.Tyr1744Asp | heterozygous (inherited from mosaic father) | autosomal dominant mental retardation 49 | VUS | probably causal | phenotype consistent with reported cases; bioinformatics predict variant is damaging |
| G558-1       | F   | CDH2 | NP_001783.2:p. Asp627Tyr | de novo heterozygous | syndromic neurodevelopmental disorder | unclassified | definitely causal | phenotype consistent with newly described disorder, bioinformatics predict variant is damaging |

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| CAUSES ID no. | Sex | Gene | Variant | Mechanism | Disease | ACMG variant classification | Diagnostic interpretation by multidisciplinary research team | Rationale for diagnostic interpretation |
|--------------|-----|------|---------|-----------|---------|-----------------------------|---------------------------------------------------------------|----------------------------------------------------------------------------------|
| G559-1       | F   | MPZL2| NP_005788.1:p.Lle24MetsTer22 | compound heterozygous | autosomal recessive deafness 111 | pathogenic | probably causal | phenotype consistent with reported cases; VUS allelic to pathogenic variant and predicted to be damaging |
| G561-1       | F   | ETV6 | NP_001978.1:p.Lys409Glu     | heterozygous (maternally inherited) | thrombocytopenia 5 | VUS | probably causal | phenotype and family history consistent with thrombocytopenia 5; bioinformatics predict that variant is damaging; variant segregates with disease in family |
| G563-1       | F   | MTHFS| NP_006432.1:p.Ala9GlyfsTer42 | homozygous | neurodevelopmental disorder with microcephaly, epilepsy, and hypomyelination | unclassified | probably causal | phenotype consistent with neurodevelopmental disorder with microcephaly, epilepsy, and hypomyelination; bioinformatics predict variant is damaging |
| G575-1       | M   | COL5A1| NP_000084.3:p.Pro1566Leu    | heterozygous (maternally inherited) | Ehlers-Danlos syndrome type 1 | VUS | probably causal | phenotype consistent with Ehlers-Danlos syndrome; bioinformatics predict variant is damaging |
| CAUSES ID no. | Exome or genome sequencing | Gene | Variant (HGVS cDNA nomenclature) | Variant (HGVS protein nomenclature) | Diagnostic interpretation by multidisciplinary research team | ACMG variant classification | Mechanism | Disease (phenotype OMIM no.) | Reason for reinterpretation |
|---------------|-----------------------------|------|---------------------------------|------------------------------------|------------------------------------------------------------|-----------------------------|-----------|------------------------------|-----------------------------|
| G004-1        | F exome                     | TRCK | NM_001163435.3:c.783G>T; NM_001163435.3:c.2060-2A>G | NP_001156907.2:p.Arg261Ser          | uninformative probably causal                              | compound heterozygous       | likely pathogenic; VUS | infantile hypotonia with psychomotor retardation and characteristic facies 3 (616900) | newly described disorder |
| G007-1        | M exome                     | TUBA1A | NM_006009.4:c.1177C>T           | NP_006000.2:p.His393Tyr           | uninformative probably causal                              | de novo heterozygous        | likely pathogenic     | lissencephaly, AD 3 (611603) | improvement in bioinformatics pipeline |
| G010-1        | M exome                     | SUZ12 | NM_001535.5:4c.1960C>T          | NP_056170.2:p.Arg654Ter           | uncertain probably causal                                  | de novo heterozygous        | unclassified           | Imagawa-Matsumoto syndrome (618786) | newly described disorder |
| G025-1        | M exome                     | CLTC | NM_004859.4:c.2669C>T           | NP_004850.1:p.Pro890Leu           | uncertain probably causal                                  | de novo heterozygous        | unclassified           | mental retardation, autosomal dominant 56 (617814) | newly described disorder |
| G036-1        | M exome                     | SATB2 | NM_001172509.2:c.474-3C>G       | –                                 | uninformative probably causal                              | de novo heterozygous        | VUS                   | Glass syndrome (612313) | improvement in bioinformatics pipeline |
| G040-1        | M exome                     | ABCB7 | NM_001271696.3:c.1235T>C        | NP_001258625.1:p.Met412Thr       | uninformative definitely causal                            | de novo hemizygous          | likely pathogenic      | sideroblastic anemia and spinocerebellar ataxia (301310) | expansion of phenotype |
| G063-1        | M exome                     | POLR2A | NM_000937.5:4c.3373_3375del     | NP_000928.1:p.Lys1125del          | uninformative probably causal                              | de novo heterozygous        | likely pathogenic      | neurodevelopmental disorder with hypotonia and variable intellectual and behavioral abnormalities (618603) | newly described disorder |
| G067-1        | M exome                     | EBF3 | NM_001375380.1:c.616C>T         | NP_001362309.1:p.Arg206Ter       | uninformative definitely causal                            | heterozygous (inherited from mosaic parent) | likely pathogenic      | hypotonia, ataxia, and delayed development syndrome (617330) | new publication |
| G067-4        | F exome                     | EBF3 | NM_001375380.1:c.616C>T         | NP_001362309.1:p.Arg206Ter       | uninformative definitely causal                            | heterozygous (inherited from mosaic parent) | likely pathogenic      | hypotonia, ataxia, and delayed development syndrome (617330) | new publication |
| G070-1        | M exome                     | C20orf69 | NM_153689.6:4c.843_847del     | NP_710156.3:p.Lys282GlnfsTer55   | uninformative probably causal                              | VUS                         | homozygous            | combined oxidative phosphorylation deficiency 53 (619425) | new publications |
| G073-1        | M exome                     | SOD1 | NM_000454.5:4c.371_373del       | NP_0004445.1:p.Ala124del         | uninformative probably causal                              | homozygous                  | homozygous            | progressive spastic tetraplegia and axial hypotonia (618598) | new publication |
| G075-1        | F exome                     | MAST1 | NM_014975.3:4c.293G>T          | NP_055790.1:p.Gly98Val           | uninformative probably causal                              | de novo heterozygous        | unclassified           | mega-corpus-callosum syndrome with cerebellar hypoplasia and cortical malformations (618273) | new publication (includes this individual) |
| G077-1        | F exome                     | BPTF | NM_182641.4:4c.1957A>T         | NP_872579.2:p.Arg653Ter          | uninformative probably causal                              | de novo heterozygous        | heterozygous           | neurodevelopmental disorder with dysmorphic faces and distal limb anomalies (617755) | new publication |
| G078-1        | F exome                     | ASH1L | NM_018489.3:4c.3664_3667del    | NP_060959.2:p.Lys1222GlyfsTer10   | uninformative definitely causal                            | de novo heterozygous        | pathogenic             | autosomal dominant mental retardation 52 (617796) | new publication |

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| CAUSES ID no. | Sex | Exome or genome sequencing | Gene | Variant (HGVS cDNA nomenclature) | Variant (HGVS protein nomenclature) | Diagnostic interpretation by multidisciplinary research team | ACMG variant classification | Mechanism | Disease (phenotype OMIM no.) | Reason for reinterpretation |
|-------------|-----|----------------------------|------|---------------------------------|------------------------------------|----------------------------------------------------------|-----------------------------|-----------|----------------------------|-----------------------------|
| G082-1 M    |     | exome                      | ASH1 | NM_018489.3:c.6803_6804delinsTTCCTCA | NP_060959.2:p.Cys2268PhefsTer7     | uninformative definitely causal                        | pathogenic                  | de novo    | heterozygous               | autosomal dominant mental retardation 54 (617796) | new publication |
| G089-1 M    |     | exome                      | NF1  | NM_001042492.3:c.3569G>T        | NP_001035957.1:p.Gly1190Val        | uninformative probably causal                        | VUS                         | heterozygous (maternally inherited) | neurofibromatosis 1 (162200) | improvement in bioinformatics pipeline |
| G092-1 M    |     | exome                      | PAK3 | NM_002578.5:c.314C>G            | NP_002569.1:p.Ser105Cys            | uncertain probably causal                        | VUS                         | hemizygous (maternally inherited) | intellectual developmental disorder, X-linked 30 (300558) | referring physician’s interpretation based on patient phenotype |
| G103-1 F    |     | exome                      | DDX23| NM_004818.3:c.314C>G            | NP_004809.2:p.Arg754Cys            | uncertain probably causal                        | unclassified                | de novo heterozygous | DDX23-related disorder | new publication |
| G117-1 F    |     | exome                      | PIGG | NM_001127178.3:c.413A>G         | NP_001120650.1:p.Asn138Ser        | uncertain probably causal                        | VUS                         | homozygous | autosomal recessive mental retardation 53 (616917) | new publication (includes this individual) |
| G117-4 M    |     | exome                      | PIGG | NM_001127178.3:c.413A>G         | NP_001120650.1:p.Asn138Ser        | uncertain probably causal                        | VUS                         | homozygous | autosomal recessive mental retardation 53 (616917) | new publication (includes this individual) |
| G122-1 F    |     | exome                      | SRCAP| NM_006662.3:c.5503_5515del      | NP_006663.2:p.Val1835ProfsTer13    | uncertain probably causal                        | unclassified                | de novo heterozygous | developmental delay, hypotonia, musculoskeletal defects, and behavioral abnormalities (619595) | new publication |
| G164-1 F    |     | exome                      | MECP2| NM_001110792.2:c.1200_1243del   | NP_001110862.1:p.Pro401Ter         | uninformative probably causal                     | pathogenic                  | de novo heterozygous | Rett syndrome (312750) | improvement in bioinformatics pipeline |
| G169-1 M    |     | exome                      | HSD1B3| NM_000197.2:c.277+4A>T;NM_000197.2:c.824C>T | NP_000188.1:p.Ala275Val          | uninformative probably causal                     | pathogenic; VUS             | compound heterozygous | pseudohermaphroditism, male, with gynecomastia (264300) | improvement in bioinformatics pipeline |
| G174-1 M    |     | exome                      | SET  | NM_0030011.4:c.128_131del       | NP_003002.2:p.Arg44LeufsTer10     | uninformative probably causal                     | unclassified                | heterozygous (maternally inherited) | autosomal dominant mental retardation 58 (618106) | new publication (includes this individual) |
| G194-1 M    |     | exome                      | MNJ  | NM_002430.3:c.3743G>A           | NP_002421.3:p.Trp1248Ter           | uncertain definitely causal                       | VUS                         | de novo heterozygous | CIEBALID syndrome (618774) | new publication |
| G198-1 M    |     | genome                     | ARID2| NC_000012.11:546298857_4630229del | –                                | uncertain definitely causal                       | VUS                         | de novo heterozygous | Coffin-Siris syndrome 6 (617808) | referring physician’s interpretation based on patient phenotype |

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| CAUSES ID no. | Sex | Exome or genome sequencing | Gene | Variant (HGVS cDNA nomenclature) | Variant (HGVS protein nomenclature) | Diagnostic interpretation by multidisciplinary research team | ACMG variant classification | Mechanism | Disease (phenotype OMIM no.) | Reason for reinterpretation |
|--------------|-----|-----------------------------|------|----------------------------------|-------------------------------------|---------------------------------------------------------------|-----------------------------|-----------|----------------------------|--------------------------------|
| G205-1       | F   | exome                       | NBEA | NM_001385012.1:c.7294_7295dup   | NP_001371941.1: p.Glu2433ArgfsTer3 | uninformative probably causal                                  | unclassified                | de novo heterozygous | neurodevelopmental disorder with or without early-onset generalized epilepsy (619157) | new publication (includes this individual) |
| G231-1       | F   | exome                       | WDR26| NM_001379403.1:c.449C>T         | NP_001366332.1: p.Ala150Val        | uninformative probably causal                                  | VUS                         | de novo heterozygous | Skraban-Deardorff syndrome (617616) | new publication |
| G235-1       | M   | exome                       | TRAF7| NM_002271.3:c.1570C>T           | NP_115647.2: p.Arg524Trp           | uninformative probably causal                                  | unclassified                | de novo heterozygous | cardiac, facial, and digital anomalies with developmental delay (618164) | new publication |
| G292-1       | M   | genome                      | SYT1 | NM_005639.3:c.539C>T            | NP_005630.1: p.Pro180Leu           | uncertain probably causal                                       | unclassified                | de novo heterozygous | Baker-Gordon syndrome (618218) | new publication |
| G297-1       | F   | exome                       | JARID2| NM_004973.4:c.3379C>T           | NP_004964.2: p.Asp1277Ter          | uncertain probably causal                                       | unclassified                | de novo hemizygous | JARID2-neurodevelopmental syndrome (includes this individual) | new publication |
| G328-1       | F   | exome                       | RRAS2| NM_012250.6:c.68G>A             | NP_036382.2: p.Gly23Asp            | uninformative definitely causal                                | pathogenic                  | de novo heterozygous | Noonan syndrome 12 (618624) | new publication |
| G338-1       | F   | exome                       | PLEKHA7| NM_001329630.2:c.571G>A         | NP_001316659.1: p.Asp191Asn        | uninformative probably causal                                  | unclassified                | heterozygous (paternally inherited) | Mendelian non-syndromic cleft lip with or without cleft palate | referring physician's interpretation based on patient phenotype |
| G338-4       | F   | exome                       | PLEKHA7| NM_001329630.2:c.571G>A         | NP_001316659.1: p.Asp191Asn        | uninformative probably causal                                  | unclassified                | heterozygous (paternally inherited) | Mendelian non-syndromic cleft lip with or without cleft palate | referring physician's interpretation based on patient phenotype |
| G363-1       | M   | exome                       | FLNA | NM_001110556.2:c.4475-1G>T      | –                                  | uncertain probably causal                                       | VUS                         | hemizygous (maternally inherited) | periventricular nodular heterotopia I (300049) | referring physician's interpretation based on patient phenotype |
| G368-1       | M   | genome                      | KDMS5| NM_004187.5:c.854C>T            | NP_004178.2: p.Ser285Lys           | uncertain probably causal                                       | VUS                         | hemizygous (maternally inherited) | X-linked syndromic mental retardation, Claes-Jensen type (300534) | referring physician's interpretation based on patient phenotype |
| G369-1       | M   | exome                       | TRX1 | NM_001379200.1:c.901G>A         | NP_001366129.1: p.Ala301Thr        | probably causal                                                | uncertain                   | de novo heterozygous | tetralogy of Fallot (187500) | referring physician's interpretation based on patient phenotype |
| G402-4       | F   | genome                      | SMARC2| NM_001330288.2:c.2037C>A        | NP_001317217.1: p.Tyr679Ter        | uncertain probably causal                                       | VUS                         | de novo heterozygous | Coffin-Siris syndrome 8 (618362) | new publication |
| G406-1       | F   | exome                       | H3F3A| NM_002107.7:c.68C>T             | NP_002098.1: p.Thr231le            | uninformative probably causal                                  | VUS                         | de novo heterozygous | H3F3A-related disorder (includes this individual) | new publication |
| G407-1       | M   | exome                       | JARID2| NM_004973.4:c.1668_1669dup      | NP_004964.2: p.Ile557ArgfsTer34    | uncertain probably causal                                       | unclassified                | de novo heterozygous | JARID2-neurodevelopmental disorder (includes this individual) | new publication |
| G421-1       | F   | exome                       | NEUROD2| NM_006160.4:c.388G>A            | NP_006151.3: p.Glu130Lys           | uninformative probably causal                                  | unclassified                | de novo heterozygous | developmental and epileptic encephalopathy 72 (618374) | new publication |

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| CAUSES ID no. | Exome or genome sequencing | Gene | Variant (HGVS cDNA nomenclature) | Variant (HGVS protein nomenclature) | Diagnostic interpretation by multidisciplinary research team | ACMG variant classification | Mechanism | Disease (phenotype OMIM no.) | Reason for reinterpretation |
|---------------|-----------------------------|------|----------------------------------|------------------------------------|------------------------------------------------------------|----------------------------|-----------|----------------------------|-----------------------------|
| G422-1 M      | exome                       | DLL1 | NM_005618.4:c.1525C>T            | NP_005609.3: p.Arg509Ter           | uninformative                                             | definitely causal          | heterozygous (paternally inherited) | variable neurodevelopmental disorder with multisystem features (618709) | new publication (includes this individual) |
| G465-1 F      | exome                       | ANKRD17 | NM_032217.5:c.5360_5363del       | NP_115593.3: p.Gln1787ArgfsTer5   | uncertain                                                 | definitely causal          | de novo heterozygous         | Chopra-Amiel-Gordon syndrome (619504) | new publication (includes this individual) |
| G469-1 M      | exome                       | BCL11B | NM_138576.4:c.726_727insCGCAGCAC | NP_612808.1: p.Thr243ArgfsTer41   | uncertain                                                 | probably causal            | heterozygous (inherited from mosaic father) | intellectual developmental disorder with dysmorphic facies, speech delay, and T cell abnormalities (618092) | new publication |
| G483-1 F      | exome                       | ALPL | NM_000478.5:c.407G>A             | NP_000469.3: p.Arg136His           | definitely causal                                         | uncertain                  | homozygous (maternally inherited chromosome 1 uniparental disomy) | hypophosphatasia, adult (146300) | referring physician’s interpretation based on patient phenotype |
| G494-1 M      | exome                       | EDEM3 | NM_025191.4:c.940A>T             | NP_09467.3: p.Arg314Ter            | uncertain                                                 | probably causal            | heterozygous (paternally inherited) | congenital disorder of glycosylation, type 2V (619493) | new publication (includes this individual) |
| G495-1 M      | exome                       | BRD4 | NM_001379291.1:c.1339C>T         | NP_00136220.1: p.Gln447Ter         | uncertain                                                 | probably causal            | de novo heterozygous         | BRD4-related disorder | newly described disorder |
| G498-1 F      | exome                       | TANC2 | NM_025185.3:c.5308A>G            | NP_079461.2: p.Arg1770Gly          | uncertain                                                 | probably causal            | de novo heterozygous         | intellectual developmental disorder with autistic features and language delay, with or without seizures (618906) | new publication |
| G504-1 F      | exome                       | TRRAP | NM_001375524.1:c.29C>T           | NP_001362453.1: p.Thr10Met        | uninformative                                             | probably causal            | VUS                        | developmental delay with or without dysmorphic facies and autism (618454) | new publication (includes this individual) |
| G536-1 M      | exome                       | NF1  | NM_001042492.3:c.5609+1G>T       | –                                  | uncertain                                                 | probably causal            | de novo heterozygous         | neurofibromatosis 1 (162200) | referring physician’s interpretation based on patient phenotype |
| G550-1 F      | exome                       | USP7 | NM_003470.2:c.963delC            | NP_003461.2: p.Lys322AsnfsTer16    | probably causal                                           | uncertain                  | de novo heterozygous         | Hao-Fountain syndrome (616863) | referring physician’s interpretation based on patient phenotype |
| G558-1 F      | exome                       | CDH2 | NM_001792.5:c.1879G>T           | NP_001783.2: p.Asp627Tyr          | uninformative                                             | definitely causal          | de novo heterozygous         | agenesis of corpus callosum, cardiac, ocular, and genital syndrome (618929) | new publication |

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Table 2. Continued

| CAUSES ID no. | Exome or genome sequencing | Gene | Variant (HGVS cDNA nomenclature) | Variant (HGVS protein nomenclature) | Diagnostic interpretation by multidisciplinary research team | ACMG variant classification | Mechanism | Disease (phenotype OMIM no.) | Reason for reinterpretation |
|---------------|---------------------------|------|---------------------------------|-----------------------------------|-------------------------------------------------------------|-----------------------------|-----------|-----------------------------|-----------------------------|
| G559-1        | F exome                  | MPZL2| NM_005797.4c:72del; NM_005797.4x:2,78A>T | NP_005788.1: p.Ile24MetTer22; NP_005788.1: p.Asp93Val | uninformative; probably causal | pathogenic; VUS | compound heterozygous | autosomal recessive deafness 111 (618145) | newly described disorder |
| G63-1         | F exome                  | MTHFS| NM_006441.4x:10,25dup          | NP_006432.1: p.Ala9GlyTer42       | uninformative; probably causal | unclassified | homozgyous | neurodevelopmental disorder with microcephaly, epilepsy, and hypomyelination | newly described disorder |
| G75-1         | M exome                  | COL5A1| NM_000093.5x:4,697C>−T       | NP_000084.3: p.Pro1566Leu        | uncertain; probably causal | VUS | heterozygous (maternally inherited) | Ehlers-Danlos syndrome, classic type 1 (130000) | referring physician's interpretation based on patient phenotype |
| G78-1         | F genome                 | SPG7 | NM_003119.3x:1226A>C          | NP_003110.1: p.Glu409Ala         | uncertain; probably causal | likely pathogenic | heterozygous (inherited from mosaic mother) | 'autosomal recessive' spastic paraplegia 7 (607259) | referring physician's interpretation based on patient phenotype |

*Variants that were interpreted as “uninformative” were not Sanger sequenced. Sanger sequencing and ACMG classification of variants were obtained when the individual was diagnosed by our multidisciplinary research team as having a genetic disease associated with a variant that was probably or definitely causal or of uncertain, but suspected, relationship to the phenotype.

In seven families who underwent trio ES, a diagnosis was not established by other testing initiated in the CAUSES study (Table 3). In three families, additional clinical exome analysis in the CAUSES study and Sanger sequencing of the third allele of a recessive locus that appeared to have a disease-causing variant identified by research GS. In one individual, a deep intronic variant was found in the second allele of a recessive locus that appeared to have a disease-causing variant identified by research GS. In the third family, a deep intronic variant was found in the second allele of a recessive locus that appeared to have a disease-causing variant identified by research GS. In the fourth family, a deep intronic variant was found in the second allele of a recessive locus that appeared to have a disease-causing variant identified by research GS. In the fifth family, a deep intronic variant was found in the second allele of a recessive locus that appeared to have a disease-causing variant identified by research GS. In the sixth family, a deep intronic variant was found in the second allele of a recessive locus that appeared to have a disease-causing variant identified by research GS. In the seventh family, a deep intronic variant was found in the second allele of a recessive locus that appeared to have a disease-causing variant identified by research GS.
Comparison of ACMG variant classification to multidisciplinary diagnosis of individuals

Over the course of this study, our multidisciplinary research team diagnosed 261 families with one or more genetic diseases caused by variants discovered on CAUSES GWS. If identification of genetic disease in the CAUSES families had been based on the ACMG classification alone (i.e., one pathogenic or likely pathogenic variant allele of an autosomal dominant or X-linked disease locus in a hemizygous male or two pathogenic/likely pathogenic alleles or one pathogenic/likely pathogenic allele and one VUS/unclassified allele for an autosomal recessive or X-linked disease in a female), a genetic disease rate would have been recognized in 33.6% instead of the 52.2% of families diagnosed by our multidisciplinary team.**

Compelling research candidates

Variants in compelling research candidate genes were identified in 113 families (Table S5), with four sets of siblings sharing candidate variants. In 14 cases, a variant in another gene was identified as probably or definitely disease causing by our multidisciplinary research team, raising the possibility of dual diagnoses in these individuals.

Discussion

The heterogeneous ancestry of the pediatric individuals referred to the CAUSES study reflects the marked diversity of the population of British Columbia. Families of European descent were most frequent, but made up a little less than half of the total. South and East Asian families constituted almost 1/3 of the total, with Middle Eastern, First Nations, and other groups accounting for the rest. A Canadian GWS study done in Toronto reported European ancestry in 61% of individuals enrolled. Apart from English, the most frequently spoken languages in British Columbia include Punjabi, Cantonese, and Mandarin (www.statcan.gc.ca). Most of the families seen in the CAUSES study were fluent in English, but translator services were used for non-English-speaking families. The ethnic diversity represented in the CAUSES cohort poses challenges for variant interpretation owing to the lack of adequate representation of non-European ethnic groups, particularly Indigenous populations, in reference databases. The frequency of de novo autosomal dominant variants identified in our cohort supports a trio-based approach to GWS, especially for individuals whose ethnicity is poorly represented in reference databases.

Neurology, medical genetics, and biochemical diseases were the clinical services that referred most individuals to the CAUSES study and are the most frequent users of GWS in the province. The largest cohort by indication was individuals with syndromic ID, who may be followed by medical specialists in any (or all) of these three clinical services.

The overall diagnostic rate of 52.2% in the CAUSES study is higher than that reported in many other series, but we used the term “diagnosis” to mean clinical identification of a genetic disease in an affected individual rather than the surrogate “molecular diagnosis” used in many published reports. “Molecular diagnosis” is usually based only on standard ACMG variant classification, zygosity, allelism, and the limited phenotypic information provided...
| CAUSES ID no. | Sex | Age (years) | Phenotype | CAUSES finding | Follow-up investigations | Gene | Variant (HGVS cDNA nomenclature) | Variant (HGVS protein nomenclature) | Mechanism | Disease (phenotype OMIM no.) | Comment |
|--------------|-----|-------------|-----------|----------------|------------------------|------|------------------------------|--------------------------------|-----------|----------------------------|---------|
| G012-1       | M   | 12          | moderate intellectual disability; mild dysmorphic facial features; mild webbed neck; mild brachydactyly with short distal phalanx of the finger; right lower limb vascular skin abnormality | uninformative exome | clinical exome sequencing with deletion/duplication analysis identified a heterozygous 25 kb likely pathogenic duplication involving exons 4–20 of CUL4B | CUL4B | NC_000023.10: (c.610+1_611/C0) (2493+1_2494−1)dup | − | hemizygous (inheritance unknown) | X-linked syndromic mental retardation 15 (Cabezas type) (300354) | CAUSES exome sequencing analysis did not include assessment of copy number |
| G013-1       | F   | 9           | short stature (−5 SD); craniofacial dysmorphism; bilateral hip deformities; feeding difficulties; abnormal growth hormone level; congenital adrenal hypoplasia; nephrocalcinosis; osteopenia | uninformative exome | targeted research sequencing found biallelic variants in POLE | POLE | NM_006231.4:c.3265G>C; NM_006231.3:c.1682+32C>G | NP_006222.2:p.Val1089Leu; NP_006222.2:p.Asn563Valfs*16 | compound heterozygous | IMAGe syndrome (618336) | second variant is deep intronic and was not captured in exome sequencing |
| G139-1       | M   | 10          | developmental delay; microcephaly; cataracts; myopia; hearing loss; renal cysts; cysts of spleen | uninformative exome | clinical exome sequencing with deletion/duplication analysis identified a heterozygous 8,928 bp likely pathogenic deletion involving exons 13–18 of COL11A1 | COL11A1 | NC_000001.10:g.103471300−103480228del | − | heterozygous (inheritance unknown) | Stickler syndrome, type II (604841) | CAUSES exome sequencing analysis did not include assessment of copy number |
| G199-1       | F   | 15          | learning disability; astrocytoma; neurofibroma in a muscle | uninformative exome | research genome sequencing identified a heterozygous 92 kb pathogenic deletion involving all of CDNKA and part of CDNKB | CDNKA/CDNK2B | NC_000009.11:g.21915312_22006909del | − | de novo heterozygous | melanoma and neural system tumor syndrome (155755) | CAUSES exome sequencing analysis did not include assessment of copy number |

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| CAUSES ID no. | Sex | Age (years) | Phenotype                                      | CAUSES finding | Follow-up investigations | Gene                          | Variant (HGVS cDNA nomenclature) | Variant (HGVS protein nomenclature) | Mechanism   | Disease (phenotype OMIM no.) | Comment                                                                 |
|---------------|-----|-------------|-----------------------------------------------|----------------|--------------------------|-------------------------------|----------------------------------|----------------------------------|-------------|--------------------------|--------------------------------------------------------------------------------|
| G222-1        | F   | 4           | global developmental delay; cerebellar hypoplasia; microcephaly; cleft palate; Angelman syndrome phenotype with normal UBE3A methylation | uninformative exome | research genome sequencing identified a heterozygous 13.9 kb tandem duplication involving exons 11–12 of CASK; the deletion was classified as a VUS by ACMG criteria but was considered to be disease causing by the patient’s clinicians | CASK                          | NC_000023.10:g.41468838-41482710dup1.8 | –                       | de novo heterozygous         | mental retardation and microcephaly with pontine and cerebellar hypoplasia (300749) | CAUSES exome sequencing analysis did not include assessment of copy number |
| G351-1        | M   | 12          | myotonic dystrophy; mild intellectual disability; dysmorphic facial features; strabismus; joint hypermobility; inguinal hernias | compound heterozygous variants of PYCR1 considered to be definitely causal of autosomal recessive cutis laxa type IIb | clinical short tandem repeat analysis of DMPK prior to enrollment in CAUSES identified a pathogenic 150 repeat expansion in the proband and a 430 repeat expansion in the mother, both of whom were diagnosed with myotonic dystrophy | DMPK                          | NM_004409.5:c.*224_*226CTG150 | –                       | heterozygous (maternally inherited) | myotonic dystrophy type 1 (160900) | CAUSES exome sequencing analysis did not include assessment for expansions of short tandem repeats |
| G410-1        | F   | 12          | facial muscle weakness; velopharyngeal insufficiency; muscular hypotonia of the trunk; proximal muscle weakness; failure to thrive | uninformative exome | clinical testing with a multigene muscle disorder panel; two pathogenic variants of SEPN1 | SEPN1                         | SEPN1                          | SEPN1-related myopathy (602771) | compound heterozygous           | -           | -                        | second variant is a 10 bp insertion in exon 1; SEPN1 exon 1 was poorly covered in CAUSES exome dataset |
on the test requisition. This siloing of the data available to the laboratory from the rest of the information the clinical team knows about the affected individual makes it difficult or impossible for the laboratory to assess the genotype-phenotype correlation fully, and genotype-phenotype correlation is essential to genetic disease diagnosis. Our use of “diagnosis” to mean interpretation of the GWS laboratory report in the context of all available clinical information about the affected individual and family is consistent with ACMG and UK practice guidelines1,6,44 and the recommendations of other expert groups.7,45 High rates of genetic disease diagnosis have also been reported in other GWS studies that have based genetic disease diagnosis on multidisciplinary interpretation and access to patients’ complete medical records.8-10,17,26

Discordance between ACMG variant classification and clinical diagnosis of genetic disease has been noted in studies comparing ACMG classifications of disease genes to ClinVar46 or the Human Gene Mutation Database.17 Some of these differences are probably attributable to incomplete penetrance of many genetic diseases, a problem that is recognized in the ACMG guidelines.1 Late-onset and variable expressivity of genetic disease also confound “molecular diagnosis” based solely on ACMG variant classification.

We found no difference in the diagnostic rate between 415 families studied with ES (52.3%) and 85 families studied with GS (51.8%) in the CAUSES study. Most previous studies have found similar diagnostic rates in patients who received GS and those who received ES,2,18 but there is also evidence that GS identifies some disease-causing variants that are not found by ES.11,17 The high diagnostic rate in our study may also reflect patient selection and our use of trio-based, rather than singleton, GWS, as well as our use of a multidisciplinary research team to make diagnoses in all cases, including those in which variants were unclassified or classified as ACMG VUS.

CAUSES participants were followed clinically after initial interpretation of their GWS results and with periodic bioinformatic reanalysis and reinterpretation of variants as indicated. This follow-up and reclassification increased our diagnostic rate from 43.0% at the time of initial interpretation to 52.2% at the end of the study. Reinterpretation by the CAUSES team resulted in an average increase in diagnosis of 4.8% per year in families initially interpreted as uninformative or uncertain. This rate of additional diagnoses is consistent with other studies that reanalyzed GWS data after a shorter period of time.48,49 We found that additional genetic diagnoses continued to be made at a similar rate for at least 5 years after the test was done, with no sign of decreasing over this period (Figure 1).

Dual diagnoses
The rate of dual diagnosis in the CAUSES study (eight families; 1.6%) is lower than the ~5%-7% reported in most other studies.50,51 We did not count variants that had an uncertain relationship to the disease among individuals with a diagnosis, and we did not routinely reanalyze bioinformatic datasets once one diagnosis had been established for an affected individual. However, there are 14 individuals in the CAUSES study who were diagnosed with a genetic disease and also have compelling research variants (Table S5).

“False negatives” and false positives
Seven (2.9%) of the 241 CAUSES families who underwent trio ES and had a result that was interpreted as either uninformative (n = 226) or uncertain (n = 15) were subsequently diagnosed with a specific genetic disorder after a disease-causing genetic variant was found by another clinical or research test (Table 3). Our failure to identify these variants in the CAUSES study reflected either technical limitations of the ES platform or the fact that we did not test for copy number variants in ES data.

It is interesting to note that in six of the seven families in which a “false negative” CAUSES ES result occurred, a disease-causing variant either was found on research GS (G199 and G222) or probably would have been found had GS been done on the family (G012, G013, G139, G410). There were no “false negatives” among individuals who underwent GS, but trio GS was performed in only 85 of the 500 families included in the CAUSES study.

Although rescinding the clinical diagnosis of a genetic disease was frequent prior to the advent of routine genetic testing and often occurs as a result of genetic testing, few studies deal with the occurrence of false positive diagnoses made on the basis of GWS results. We had four such cases (G103, G369, G483, and G550) in our study, all resulting from lack of clinical concordance with the phenotype expected for the observed genotype. Our finding of an occasional false positive clinical diagnosis after GWS is consistent with observations in the DDD,52 UK 100,000 Genomes Pilot studies,17 and a recent review of medical records on 130 patients for whom the laboratory and clinical interpretations of sequencing test results were compared.53

The difference between variant classification and clinical diagnosis
Genotype-phenotype correlation is the core principle of genetic disease diagnosis. Physicians diagnose genetic disease on the basis of all of the available information about an affected individual, including the medical history and disease course over time, family history, physical examination, specialist consultations, imaging studies, and all of the laboratory results, including reports of ES or GS. The UK practice guideline for variant interpretation43 advocates use of a genomic multidisciplinary team “to assess the gene variant(s) identified in the context of the patient’s phenotype data and ascertain their contribution to the clinical presentation,” but genotype-phenotype correlation plays only a minor role (as the phenotypic specificity criterion, PP4) in the ACMG classification. It is, therefore, not surprising that the ACMG classification of an affected...
individual's variant(s) as “likely pathogenic” or of “uncertain significance” and the clinician’s interpretation of that report in the context of the affected individual's overall clinical picture may differ. The ACMG classification and the clinician are describing two different things: genetic variants on one hand and patients on the other.

The ACMG variant classification alone is not sufficient to diagnose a genetic disease. The information about a genetic variant identified by GWS and categorized by the ACMG guidelines must be interpreted in the context of an affected individual’s complete medical history, disease course, family history, physical examination findings, specialist consultations, imaging studies, and other laboratory test results. Having an ACMG pathogenic or likely pathogenic variant does not necessarily mean that the variant is causing an affected individual's genetic disease. For example, heterozygous carriers of an autosomal recessive condition or non-penetrant carrier of an autosomal dominant disorder have pathogenic variants that do not affect their own health. The ACMG pathogenic classification means that a variant is capable of causing genetic disease in some circumstances. The clinician who ordered the test must decide whether those circumstances exist in a particular affected individual; it is the physician (or multidisciplinary research team in CAUSES) who makes the diagnosis.

Many clinicians diagnose autosomal recessive genetic diseases in patients with characteristic phenotypes and biochemistry results and one ACMG pathogenic or likely pathogenic allele and a VUS of the second allele if the phenotype and other laboratory results are characteristic of the disease. Similarly, autosomal dominant genetic disease may be diagnosed in patients with an ACMG VUS of the associated gene, a classical phenotype, and ancillary supporting data. Discordance between ACMG variant classification and clinical diagnosis of genetic disease has also been observed in studies of large reference databases.

The CAUSES study was a translational research project that provided trio-based ES or GS to 500 families of children with suspected genetic diseases. We diagnosed a specific genetic condition in 52.2% of the individuals enrolled in this study, a high diagnostic rate that we attribute largely to (1) close collaboration between clinical geneticists, genetic counselors, laboratory geneticists, and clinical bioinformaticians on our research team and the affected individual's clinical team in interpreting the variants found and (2) continuing follow-up of GWS results, with reanalysis and reinterpretation pursued over many years to take advantage of technical improvements and new knowledge that have accumulated.

We learned that pre-test genetic counseling involves much more than just “consenting” the family, that “false negative” and false positive results occasionally do occur with clinical GWS, that genetic counseling is valuable in preparing families for possible changes in interpretation that may take place over time, and that follow-up is important for families with uninformative as well as those with positive GWS results. Finally, we were reminded that patient-oriented research is essential to the provision of high-quality genetic health care.

### Data and code availability
The scripts used for this analysis are available at https://github.com/FriedmanLab/StructuralVariantAnalysis. Additional clinical data may be available upon request from the corresponding author, subject to privacy or ethical restrictions. The variants reported in Table S1 have been deposited in the ClinVar database.

### Supplemental information
Supplemental information can be found online at https://doi.org/10.1016/j.xhgg.2022.100108.

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### Author contributions
A.M.E., S.A., C.D.S., A.L., T.N., C.V.K., and J.M.F. contributed to the conception and study design. All authors contributed to data acquisition. A.M.E. and J.M.F. drafted the manuscript and all authors contributed to critical review and editing of the manuscript. A.M.E., S.A., C.D.S., A.L., T.N., and J.M.F. accessed and verified the data.

### Declaration of interests
The authors declare no competing interests.

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### Web resources
ClinVar, [https://www.ncbi.nlm.nih.gov/clinvar/](https://www.ncbi.nlm.nih.gov/clinvar/)
DECIPHER, [https://www.deciphergenomics.org/](https://www.deciphergenomics.org/)
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