

Donor 5590

Genetic Testing Summary

Fairfax Cryobank recommends reviewing this genetic testing summary with your healthcare provider to determine suitability.

Last Updated: 5/10/21

Donor Reported Ancestry: French, Latvian, German, Filipino Jewish Ancestry: No

| Genetic Test* | Result | Comments/Donor's Residual Risk** |
|---------------|--------|----------------------------------|
| | | |

| Chromosome analysis (karyotype) | Normal male karyotype | No evidence of clinically significant chromosome abnormalities | | |
|--|---|---|--|--|
| Hemoglobin evaluation | Normal hemoglobin fractionation and MCV/MCH results | Reduced risk to be a carrier for sickle cell anemia, beta thalassemia, alpha thalassemia trait (aa/ and a-/a-) and other hemoglobinopathies | | |
| Expanded Genetic Disease Carrier Screening Panel attached- 283 diseases by gene sequencing | Carrier: Cystic Fibrosis (CFTR) Carrier: Hermansky-Pudlak Syndrome Type 1 (HPS1) Carrier: Phenylalanine Hydroxylase Deficiency (PAH) Carrier: Spinal Muscular Atrophy (SMN1) Negative for other genes sequenced | Partner testing recommended before using this donor. | | |

^{*}No single test can screen for all genetic disorders. A negative screening result significantly reduces, but cannot eliminate, the risk for these conditions in a pregnancy.

^{**}Donor residual risk is the chance the donor is still a carrier after testing negative.





Patient Information

Name: Donor 5590

Date of Birth:

Sema4

Client ID:

Indication: Carrier Testing

Specimen Information

Specimen Type: Blood
Date Collected: 09/15/2020
Date Received: 09/16/2020
Final Report: 10/06/2020



Expanded Carrier Screen (283) Minus TSE

Number of genes tested: 283

SUMMARY OF RESULTS AND RECOMMENDATIONS

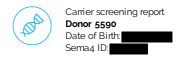
| ① Positive | ○ Negative |
|--|--|
| Carrier of Cystic Fibrosis (AR) Associated gene(s): CFTR Variant(s) Detected: c.1646G>A, p.S549N, Pathogenic, Heterozygous (one copy) | Negative for all other genes tested To view a full list of genes and diseases tested please see Table 1 in this report |
| Carrier of Hermansky-Pudlak Syndrome, Type 1 (AR) | |
| Associated gene(s): HPS1 | |
| Variant(s) Detected: c.1870C>T, p.Q624X, Likely Pathogenic, | |
| Heterozygous (one copy) | |
| Carrier of Phenylalanine Hydroxylase Deficiency (AR) | |
| Associated gene(s): PAH | |
| Variant(s) Detected: c.1222C>T, p.R408W, Pathogenic, Heterozygous | |
| (one copy) | |
| Carrier of Spinal Muscular Atrophy (AR) | |
| Associated gene(s): SMN1 | |
| Variant(s) Detected: Loss of one copy of <i>SMN1</i> | |

AR=Autosomal recessive; XL=X-linked

Recommendations

- Testing the partner for the above positive disorder(s) and genetic counseling are recommended.
- · Please note that for female carriers of X-linked diseases, follow-up testing of a male partner is not indicated.
- CGG repeat analysis of *FMR1* for fragile X syndrome is not performed on males as repeat expansion of premutation alleles is not expected in the male germline.
- Individuals of Asian, African, Hispanic and Mediterranean ancestry should also be screened for hemoglobinopathies by CBC and hemoglobin electrophoresis.
- Consideration of residual risk by ethnicity after a negative carrier screen is recommended for the other diseases on the panel, especially in the case of a positive family history for a specific disorder.





Interpretation of positive results

Cystic Fibrosis (AR)

Results and Interpretation

A heterozygous (one copy) pathogenic missense variant, c.1646G>A, p.S549N, was detected in the *CFTR* gene (NM_000492.3). When this variant is present in trans with a pathogenic variant, it is considered to be causative for cystic fibrosis. Therefore, this individual is expected to be at least a carrier for cystic fibrosis. Heterozygous carriers are not expected to exhibit symptoms of this disease.

What is Cystic Fibrosis?

Cystic fibrosis is an autosomal recessive disorder caused by pathogenic variants in the gene *CFTR*. It may be diagnosed in individuals worldwide, but has the highest prevalence in the Caucasian population, in individuals with Northern European ancestry. The clinical presentation includes thick mucus accumulation in the lungs leading to breathing difficulties and infection, poor digestion, and male infertility. The average life expectancy is in the 30s. Although some genotype/phenotype correlations exist, individuals with two classic pathogenic variants in *CFTR* are expected to present with a more severe disease phenotype. Non-classic variants in *CFTR* may lead to less severe forms of disease or specific phenotypes, such as male infertility as a result of congenital absence or hypoplasia of the vas deferens.

Hermansky-Pudlak Syndrome, Type 1 (AR)

Results and Interpretation

A heterozygous (one copy) likely pathogenic premature stop codon, c.1870C>T, p.Q624X, was detected in the *HPS1* gene (NM_000195.4). When this variant is present in trans with a pathogenic variant, it is considered to be causative for Hermansky-Pudlak syndrome, type 1. Therefore, this individual is expected to be at least a carrier for Hermansky-Pudlak syndrome, type 1. Heterozygous carriers are not expected to exhibit symptoms of this disease.

What is Hermansky-Pudlak Syndrome, Type 1?

Hermansky-Pudlak syndrome, type 1 is an autosomal recessive disorder that is caused by pathogenic variants in the gene *HPS1*. It is a rare disease worldwide, but is prevalent in Puerto Rican individuals due to the presence of a founder mutation. The disease is characterized by the presence of light-colored skin and hair (albinism) and reduced vision. Patients also have a susceptibility to prolonged bleeding caused by abnormalities in the platelets, which normally function in the clotting process. Most patients develop pulmonary fibrosis, which usually begins in the patient's early 30s and can be fatal within a decade. About 15% of patients will develop Crohn's disease. Life expectancy is usually in the patient's 40s or 50s due to progressive lung disease. It is not currently possible to predict the severity of symptoms based on the variants inherited.

Phenylalanine Hydroxylase Deficiency (AR)

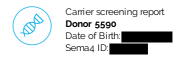
Results and Interpretation

A heterozygous (one copy) pathogenic missense variant, c.1222C>T, p.R408W, was detected in the *PAH* gene (NM_000277.1). When this variant is present in trans with a pathogenic variant, it is considered to be causative for phenylalanine hydroxylase deficiency. Therefore, this individual is expected to be at least a carrier for phenylalanine hydroxylase deficiency. Heterozygous carriers are not expected to exhibit symptoms of this disease.

What is Phenylalanine Hydroxylase Deficiency?

Phenylalanine hydroxylase deficiency is an autosomal recessive disorder caused by pathogenic variants in the gene *PAH*. While it is found in many different ethnicities, it is particularly prevalent in Sephardic Jewish, Sicilian, Irish, and Turkish individuals, as well as Caucasians. Pathogenic *PAH* variants result in loss of function of the phenylalanine hydroxylase enzyme, which breaks down the amino acid phenylalanine. The most severe form of the disease is called phenylketonuria. If untreated, buildup of phenylalanine will result in irreversible brain damage and severe intellectual disability. Treatment involves the removal of phenylalanine from the diet. Even with strict adherence to the treatment, some neurologic deficiencies have been noticed in long-term survivors. Psychological problems, including anxiety, depression, phobias and panic attacks may occur in adults who do not comply well to their treatment. Some patients have a milder form of hyperphenylalaninemia and may





tolerate higher levels of phenylalanine in their diet. Depending on the genotype, patients may be responsive to BH4, which can direct their treatment. However, it is not always possible to predict the severity of the disease based on genotype.

Spinal Muscular Atrophy (AR)

Results and Interpretation

SMN1 copy number: 1 SMN2 copy number: >=3 c.*3+80T>G: Negative

Gene(s) analyzed: SMN1 (NM_000344.3) and SMN2 (NM_017411.3)

Inheritance: Autosomal Recessive

This patient is positive for loss of one copy of *SMN1* and is, therefore, a carrier for SMA. Complete loss of *SMN1* is causative in spinal muscular atrophy (SMA). One copy of *SMN1* was detected in this individual, which is consistent with being a carrier for SMA. This individual was found to be negative for c.*3*80T>G; however, given that this patient was found to be an SMA carrier by MLPA analysis, this finding does not modify residual risk.

What is spinal muscular atrophy?

Spinal muscular atrophy (SMA) is a pan-ethnic, autosomal recessive disease caused by loss of function of the *SMN1* gene. In over 95% of cases, patients are missing both copies of the *SMN1* gene. The disease is characterized by the degeneration of alpha motor neurons of the spinal cord anterior horn cells, leading to progressive symmetric weakness, atrophy of the proximal voluntary muscles and early death. Age of onset can be anywhere on a continuum from the prenatal period to adulthood.

- SMA o represents the most severe form. Infants are born with severe hypotonia and joint contractures; no motor milestones are achieved and patients die before 6 months of age.
- SMA I has an age of onset in the first six months of life. These cases are associated with death usually by age 2 and the lack of development of motor skills.
- SMA II has an age of onset between 3 and 15 months; patients may be able to sit independently. Intelligence is not affected. Life expectancy may vary from early childhood to early adulthood.
- SMA III has an age of onset after 18 months of age and as late as adolescence; patients may learn to stand and to walk short distances.

 These patients may have a normal lifespan.
- SMA IV is an adult-onset disorder of muscle weakness; life span is not shortened.

Most patients, regardless of the severity of disease, have a deletion of both *SMN1* copies. Patients with later-onset disease usually have three or more copies of *SMN2*, which encodes a small amount of residual protein and lessens the severity of the symptoms. However, other factors besides *SMN2* copy number may affect the phenotype, and therefore the severity of the disease may not be able to be accurately predicted in all patients based on genotype. New treatments may be available to infants and children to prevent development of symptoms and slow progression of the disease.

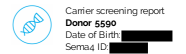
Test description

This patient was tested for a panel of diseases using a combination of sequencing, targeted genotyping and copy number analysis. Please note that negative results reduce but do not eliminate the possibility that this individual is a carrier for one or more of the disorders tested. Please see Table 1 for a list of genes and diseases tested, and **go.sema4.com/residualrisk** for specific detection rates and residual risk by ethnicity. With individuals of mixed ethnicity, it is recommended to use the highest residual risk estimate. Only variants determined to be pathogenic or likely pathogenic are reported in this carrier screening test.

Anastasia Larmore, Ph.D., Assistant Laboratory Director

Laboratory Medical Consultant: George A. Diaz, M.D., Ph.D.





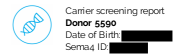
Genes and diseases tested

For specific detection rates and residual risk by ethnicity, please visit go.sema4.com/residualrisk

Table 1: List of genes and diseases tested with detailed results

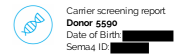
| | Disease | Gene | Inheritance Pattern | Status | Detailed Summary |
|----------|--|-------------|------------------------|----------------------------|--|
| ⊕ | Positive | | | | |
| | Cystic Fibrosis | CFTR | AR | Carrier | c.1646G>A, p.S549N, Pathogenic, Heterozygous (one copy) |
| | Hermansky-Pudlak Syndrome, Type 1 | HPS1 | AR | Carrier | c.1870C>T, p.Q624X, Likely Pathogenic, Heterozygous (one copy) |
| | Phenylalanine Hydroxylase Deficiency | PAH | AR | Carrier | c.1222C>T, p.R408W, Pathogenic, Heterozygous (one copy) |
| | Spinal Muscular Atrophy | SMN1 | AR | Carrier | SMN1 copy number: 1 SMN2 copy number: >=3 c.*3+80T>G: Negative |
| Θ | Negative | | | | |
| | 3-Beta-Hydroxysteroid Dehydrogenase Type II Deficiency | HSD3B2 | AR | Reduced Risk | |
| | 3-Methylcrotonyl-CoA Carboxylase Deficiency (MCCC1- Related) | MCCC1 | AR | Reduced Risk | |
| | 3-Methylcrotonyl-CoA Carboxylase Deficiency (MCCC2-Related) | MCCC2 | AR | Reduced Risk | |
| | 3-Methylglutaconic Aciduria, Type III | OPA3 | AR | Reduced Risk | |
| | 3-Phosphoglycerate Dehydrogenase Deficiency | PHGDH | AR | Reduced Risk | |
| | 6-Pyruvoyl-Tetrahydropterin Synthase Deficiency | PTS | AR | Reduced Risk | |
| | Abetalipoproteinemia | MTTP | AR | Reduced Risk | |
| | Achromatopsia (CNGB3-related) | CNGB3 | AR | Reduced Risk | |
| | Acrodermatitis Enteropathica | SLC39A4 | AR | Reduced Risk | |
| | Acute Infantile Liver Failure | TRMU | AR | Reduced Risk | |
| | Acyl-CoA Oxidase I Deficiency | ACOX1 | AR | Reduced Risk | |
| | Adenosine Deaminase Deficiency | ADA | AR | Reduced Risk | |
| | Adrenoleukodystrophy, X-Linked | ABCD1 | XL | Reduced Risk | |
| | Aicardi-Goutieres Syndrome (SAMHD1-Related) | SAMHD1 | AR | Reduced Risk | |
| | Alpha-Mannosidosis | MAN2B1 | AR | Reduced Risk | |
| | Alpha-Thalassemia | HBA1/HBA2 | AR | Reduced Risk | HBA1 Copy Number: 2 HBA2 Copy Number: 2 No pathogenic copy number variants detected HBA1/HBA2 Sequencing: Negative |
| | Alpha-Thalassemia Mental Retardation Syndrome | ATRX | XL | Reduced Risk | |
| | Alport Syndrome (COL4A3-Related) | COL4A3 | AR | Reduced Risk | |
| | Alport Syndrome (COL4A4-Related) | COL4A4 | AR | Reduced Risk | |
| | Alport Syndrome (COL4A5-Related) | COL4A5 | XL | Reduced Risk | |
| | Alstrom Syndrome | ALMS1 | AR | Reduced Risk | |
| | Andermann Syndrome | SLC12A6 | AR | Reduced Risk | |
| | Argininosuccinic Aciduria | ASL | AR | Reduced Risk | |
| | Aromatase Deficiency | CYP19A1 | AR | Reduced Risk | |
| | Arthrogryposis, Mental Retardation, and Seizures | SLC35A3 | AR | Reduced Risk | |
| | Asparagine Synthetase Deficiency | ASNS | AR | Reduced Risk | |
| | Aspartylglycosaminuria | AGA | AR | Reduced Risk | |
| | Ataxia With Isolated Vitamin E Deficiency | TTPA | AR | Reduced Risk | |
| - | Ataxia-Telangiectasia Autosomal Recessive Spastic Ataxia of Charlevoix- | ATM SACS | AR AR | Reduced Risk Reduced Risk | |
| | Saguenay | | | | |
| | Bardet-Biedl Syndrome (BBS10-Related) | BBS10 | AR | Reduced Risk | |





| Bardet-Biedl Syndrome (BBS12-Related) | DDC42 | A.D. | Reduced Risk | |
|--|----------|----------|---------------------------|---|
| Bardet-Biedl Syndrome (BBS12-Related) | BBS12 | AR | Reduced Risk | |
| <u> </u> | BBS1 | AR | | |
| Bardet-Biedl Syndrome (BBS2-Related) | BBS2 | AR AR | Reduced Risk | |
| Bare Lymphocyte Syndrome, Type II | CIITA | | Reduced Risk Reduced Risk | |
| Bartter Syndrome, Type 4A | BSND | AR | | |
| Bernard-Soulier Syndrome, Type A1 | GP1BA | AR | Reduced Risk | |
| Bernard-Soulier Syndrome, Type C | GP9 | AR | Reduced Risk | |
| Beta-Globin-Related Hemoglobinopathies | HBB | AR | Reduced Risk | |
| Beta-Ketothiolase Deficiency | ACAT1 | AR | Reduced Risk | |
| Bilateral Frontoparietal Polymicrogyria | GPR56 | AR | Reduced Risk | |
| Biotinidase Deficiency | BTD | AR | Reduced Risk | |
| Bloom Syndrome | BLM | AR | Reduced Risk | |
| Canavan Disease | ASPA | AR | Reduced Risk | |
| Carbamoylphosphate Synthetase I Deficiency | CPS1 | AR | Reduced Risk | |
| Carnitine Palmitoyltransferase IA Deficiency | CPT1A | AR | Reduced Risk | |
| Carnitine Palmitoyltransferase II Deficiency | CPT2 | AR | Reduced Risk | |
| Carpenter Syndrome | RAB23 | AR | Reduced Risk | |
| Cartilage-Hair Hypoplasia | RMRP | AR | Reduced Risk | |
| Cerebral Creatine Deficiency Syndrome 1 | SLC6A8 | XL | Reduced Risk | |
| Cerebral Creatine Deficiency Syndrome 2 | GAMT | AR | Reduced Risk | |
| Cerebrotendinous Xanthomatosis | CYP27A1 | AR | Reduced Risk | |
| Charcot-Marie-Tooth Disease, Type 4D | NDRG1 | AR | Reduced Risk | |
| Charcot-Marie-Tooth Disease, Type 5 / Arts Syndrome | PRPS1 | XL | Reduced Risk | |
| Charcot-Marie-Tooth Disease, X-Linked | GJB1 | XL | Reduced Risk | |
| Choreoacanthocytosis | VPS13A | AR | Reduced Risk | |
| Choroideremia | СНМ | XL | Reduced Risk | |
| Chronic Granulomatous Disease (CYBA-Related) | CYBA | AR | Reduced Risk | |
| Chronic Granulomatous Disease (CYBB-Related) | CYBB | XL | Reduced Risk | |
| Citrin Deficiency | SLC25A13 | AR | Reduced Risk | |
| Citrullinemia, Type 1 | ASS1 | AR | Reduced Risk | |
| Cohen Syndrome | VPS13B | AR | Reduced Risk | |
| Combined Malonic and Methylmalonic Aciduria | ACSF3 | AR | Reduced Risk | |
| Combined Oxidative Phosphorylation Deficiency 1 | GFM1 | AR | Reduced Risk | |
| Combined Oxidative Phosphorylation Deficiency 3 | TSFM | AR | Reduced Risk | |
| Combined Pituitary Hormone Deficiency 2 | PROP1 | AR | Reduced Risk | |
| Combined Pituitary Hormone Deficiency 3 | LHX3 | AR | Reduced Risk | |
| Combined SAP Deficiency | PSAP | AR | Reduced Risk | |
| Congenital Adrenal Hyperplasia due to 17-Alpha- | , 0, | 7.11. | Troubou Trior | |
| Hydroxylase Deficiency | CYP17A1 | AR | Reduced Risk | |
| Congenital Adrenal Hyperplasia due to 21-Hydroxylase | | | | CYP21A2 copy number: 2 |
| Deficiency | CYP21A2 | AR | Reduced Risk | CYP21A2 sequencing: Negative |
| Congenital Amegakaryocytic Thrombocytopenia | MPL | AR | Reduced Risk | 277 <u>22 2 334451.511.9</u> 17 1394477 |
| Congenital Disorder of Glycosylation, Type la | PMM2 | AR | Reduced Risk | |
| Congenital Disorder of Glycosylation, Type Ib | MPI | AR | Reduced Risk | |
| Congenital Disorder of Glycosylation, Type Ib | ALG6 | AR | Reduced Risk | |
| Congenital Insensitivity to Pain with Anhidrosis | NTRK1 | AR | Reduced Risk | |
| Congenital Myasthenic Syndrome (CHRNE-Related) | | | Reduced Risk | |
| | CHRNE | AR | | |
| Congenital Neutropopia (HAY) Polated) | RAPSN | AR | Reduced Risk | |
| Congenital Neutropenia (HAX1-Related) | HAX1 | AR | Reduced Risk | |
| Congenital Neutropenia (VPS45-Related) | VPS45 | AR | Reduced Risk | |
| Corneal Dystrophy and Perceptive Deafness | SLC4A11 | AR | Reduced Risk | |
| Corticosterone Methyloxidase Deficiency | CYP11B2 | AR | Reduced Risk | |
| Cystinosis | CTNS | AR | Reduced Risk | |
| D-Bifunctional Protein Deficiency | HSD17B4 | AR | Reduced Risk | |
| Deafness, Autosomal Recessive 77 | LOXHD1 | AR | Reduced Risk | |
| Duchenne Muscular Dystrophy / Becker Muscular | DMD | XL | Reduced Risk | |
| Dystrophy | | | | |
| Dyskeratosis Congenita (RTEL1-Related) | RTEL1 | AR | Reduced Risk | |
| Dystrophic Epidermolysis Bullosa | COL7A1 | AR | Reduced Risk | |
| | ADAMTS2 | AR | | |





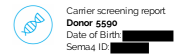
| Ellis-van Creveld Syndrome (EVC-Related) | EVC | AR | Reduced Risk | |
|---|----------|-----|--------------|---|
| Emery-Dreifuss Myopathy 1 | EMD | XL | Reduced Risk | |
| Enhanced S-Cone Syndrome | NR2E3 | AR | Reduced Risk | |
| Ethylmalonic Encephalopathy | ETHE1 | AR | Reduced Risk | |
| Fabry Disease | GLA | XL | Reduced Risk | |
| Factor IX Deficiency | F9 | XL | Reduced Risk | |
| Factor XI Deficiency | F11 | AR | Reduced Risk | |
| Familial Autosomal Recessive Hypercholesterolemia | LDLRAP1 | AR | Reduced Risk | |
| Familial Dysautonomia | IKBKAP | AR | Reduced Risk | |
| Familial Hypercholesterolemia | LDLR | AR | Reduced Risk | |
| Familial Hyperinsulinism (ABCC8-Related) | ABCC8 | AR | Reduced Risk | |
| Familial Hyperinsulinism (KCNJ11-Related) | KCNJ11 | AR | Reduced Risk | |
| Familial Mediterranean Fever | MEFV | AR | Reduced Risk | |
| Fanconi Anemia, Group A | FANCA | AR | Reduced Risk | |
| Fanconi Anemia, Group C | FANCC | AR | Reduced Risk | |
| Fanconi Anemia, Group G | FANCG | AR | Reduced Risk | |
| Tarkori Arieriia, Group G | TAIVCG | AIN | Neduced Nisk | FMR1 CGG repeat sizes: Not Performed |
| | | | | FMR1 Cadrepeat sizes. Not Performed FMR1 Sequencing: Negative |
| Fragila V S andrama | FMR1 | XL | Reduced Risk | Fragile X CGG triplet repeat expansion testing v |
| Fragile X Syndrome | FMKI | XL | Reduced RISK | |
| | | | | not performed at this time, as the patient has either particularly tested exists make |
| E | | 4.5 | D 1 10'1 | been previously tested or is a male. |
| Furnarase Deficiency | FH | AR | Reduced Risk | |
| GRACILE Syndrome and Other BCS1L-Related | BCS1L | AR | Reduced Risk | |
| Disorders | | | | |
| Galactokinase Deficiency | GALK1 | AR | Reduced Risk | |
| Galactosemia | GALT | AR | Reduced Risk | |
| Gaucher Disease | GBA | AR | Reduced Risk | |
| Gitelman Syndrome | SLC12A3 | AR | Reduced Risk | |
| Glutaric Acidemia, Type I | GCDH | AR | Reduced Risk | |
| Glutaric Acidemia, Type Ila | ETFA | AR | Reduced Risk | |
| Glutaric Acidemia, Type IIc | ETFDH | AR | Reduced Risk | |
| Glycine Encephalopathy (AMT-Related) | AMT | AR | Reduced Risk | |
| Glycine Encephalopathy (GLDC-Related) | GLDC | AR | Reduced Risk | |
| Glycogen Storage Disease, Type II | GAA | AR | Reduced Risk | |
| Glycogen Storage Disease, Type III | AGL | AR | Reduced Risk | |
| Glycogen Storage Disease, Type IV / Adult | | | | |
| Polyglucosan Body Disease | GBE1 | AR | Reduced Risk | |
| Glycogen Storage Disease, Type Ia | G6PC | AR | Reduced Risk | |
| Glycogen Storage Disease, Type Ib | SLC37A4 | AR | Reduced Risk | |
| Glycogen Storage Disease, Type ID | PYGM | AR | Reduced Risk | |
| Glycogen Storage Disease, Type VII | PFKM | AR | Reduced Risk | |
| | | | | |
| HMG-CoA Lyase Deficiency | HMGCL | AR | Reduced Risk | |
| Hemochromatosis, Type 2A | HFE2 | AR | Reduced Risk | |
| Hemochromatosis, Type 3 | TFR2 | AR | Reduced Risk | |
| Hereditary Fructose Intolerance | ALDOB | AR | Reduced Risk | |
| Hereditary Spastic Paraparesis 49 | TECPR2 | AR | Reduced Risk | |
| Hermansky-Pudlak Syndrome, Type 3 | HPS3 | AR | Reduced Risk | |
| Holocarboxylase Synthetase Deficiency | HLCS | AR | Reduced Risk | |
| Homocystinuria (CBS-Related) | CBS | AR | Reduced Risk | |
| Homocystinuria due to MTHFR Deficiency | MTHFR | AR | Reduced Risk | |
| Homocystinuria, cblEType | MTRR | AR | Reduced Risk | |
| Hydrolethalus Syndrome | HYLS1 | AR | Reduced Risk | |
| Hyperomithinemia-Hyperammonemia- | | | 5 / | |
| Homocitrullinuria Syndrome | SLC25A15 | AR | Reduced Risk | |
| Hypohidrotic Ectodermal Dysplasia 1 | EDA | XL | Reduced Risk | |
| Hypophosphatasia | ALPL | AR | Reduced Risk | |
| Inclusion Body Myopathy 2 | GNE | AR | Reduced Risk | |
| | | | | |
| Infantile Cerebral and Cerebellar Atrophy | MED17 | AR | Reduced Risk | |
| Isovaleric Acidemia | IVD | AR | Reduced Risk | |
| Joubert Syndrome 2 | TMEM216 | AR | Reduced Risk | |





| Joubert Syndrome 7 / Meckel Syndrome 5 / COACH | RPGRIP1L | AR | Reduced Risk |
|---|----------------------------------|----------------------------|---|
| Syndrome | 7.0 67.07 12 | | |
| Junctional Epidermolysis Bullosa (LAMA3-Related) | LAMA3 | AR | Reduced Risk |
| Junctional Epidermolysis Bullosa (<i>LAMB3</i> -Related) | LAMB3 | AR | Reduced Risk |
| Junctional Epidermolysis Bullosa (<i>LAMC2</i> -Related) | LAMC2 | AR | Reduced Risk |
| Krabbe Disease | GALC | AR | Reduced Risk |
| Lamellar Ichthyosis, Type 1 | TGM1 | AR | Reduced Risk |
| Leber Congenital Amaurosis 10 and Other CEP290- | CEP290 | AR | Reduced Risk |
| Related Ciliopathies | CLF290 | AIN | Neduced Mak |
| Leber Congenital Amaurosis 13 | RDH12 | AR | Reduced Risk |
| Leber Congenital Amaurosis 2 / Retinitis Pigmentosa 20 | RPE65 | AR | Reduced Risk |
| Leber Congenital Amaurosis 5 | LCA5 | AR | Reduced Risk |
| Leber Congenital Amaurosis 8 / Retinitis Pigmentosa 12 | CRB1 | AR | Reduced Risk |
| / Pigmented Paravenous Chorioretinal Atrophy | CRDI | AK | Reduced Risk |
| Leigh Syndrome, French-Canadian Type | LRPPRC | AR | Reduced Risk |
| Lethal Congenital Contracture Syndrome 1 / Lethal | GLE1 | AR | Reduced Risk |
| Arthrogryposis with Anterior Horn Cell Disease | GLEI | AK | Reduced Risk |
| Leukoencephalopathy with Vanishing White Matter | EIF2B5 | AR | Reduced Risk |
| Limb-Girdle Muscular Dystrophy, Type 2A | CAPN3 | AR | Reduced Risk |
| Limb-Girdle Muscular Dystrophy, Type 2B | DYSF | AR | Reduced Risk |
| Limb-Girdle Muscular Dystrophy, Type 2C | SGCG | AR | Reduced Risk |
| Limb-Girdle Muscular Dystrophy, Type 2D | SGCA | AR | Reduced Risk |
| Limb-Girdle Muscular Dystrophy, Type 2E | SGCB | AR | Reduced Risk |
| Limb-Girdle Muscular Dystrophy, Type 21 | FKRP | AR | Reduced Risk |
| Lipoamide Dehydrogenase Deficiency | DLD | AR | Reduced Risk |
| Lipoid Adrenal Hyperplasia | STAR | AR | Reduced Risk |
| Lipoprotein Lipase Deficiency | LPL | AR | Reduced Risk |
| Long-Chain 3-Hydroxyacyl-CoA Dehydrogenase | LIL | 7111 | Neddeed Hish |
| Deficiency | HADHA | AR | Reduced Risk |
| Lysinuric Protein Intolerance | SLC7A7 | AR | Reduced Risk |
| Maple Syrup Urine Disease, Type 1a | BCKDHA | AR | Reduced Risk |
| Maple Syrup Urine Disease, Type 1b | BCKDHB | AR | Reduced Risk |
| Meckel 1 / Bardet-Biedl Syndrome 13 | MKS1 | AR | Reduced Risk |
| Medium Chain Acyl-CoA Dehydrogenase Deficiency | ACADM | AR | Reduced Risk |
| Megalencephalic Leukoencephalopathy with | 710/12/17 | 7 11 1 | reduced Hist |
| Subcortical Cysts | MLC1 | AR | Reduced Risk |
| Menkes Disease | ATP7A | XL | Reduced Risk |
| Metachromatic Leukodystrophy | ARSA | AR | Reduced Risk |
| Methylmalonic Acidemia (<i>MMAA</i> -Related) | MMAA | AR | Reduced Risk |
| Methylmalonic Acidemia (<i>MMAB</i> -Related) | MMAB | AR | Reduced Risk |
| Methylmalonic Acidemia (<i>MUT</i> -Related) | MUT | AR | Reduced Risk |
| Methylmalonic Aciduria and Homocystinuria, | | | |
| Cobalamin C Type | MMACHC | AR | Reduced Risk |
| Methylmalonic Aciduria and Homocystinuria, | MMADHC | AR | Reduced Risk |
| Cobalamin D Type | MIMADAC | AK | Neduced risk |
| Microphthalmia / Anophthalmia | VSX2 | AR | Reduced Risk |
| Mitochondrial Complex I Deficiency (ACADg-Related) | ACAD9 | AR | Reduced Risk |
| Mitochondrial Complex Deficiency (NDUFAF5-Related) | NDUFAF5 | AR | Reduced Risk |
| Mitochondrial Complex Deficiency (NDUFS6-Related) | NDUFS6 | AR | Reduced Risk |
| Mitochondrial DNA Depletion Syndrome 6 / Navajo | MD14= | 4.0 | Dadwood Did |
| Neurohepatopathy | MPV17 | AR | Reduced Risk |
| Mitochondrial Myopathy and Sideroblastic Anemia 1 | PUS1 | AR | Reduced Risk |
| | | AR | Reduced Risk |
| Mucolipidosis II / IIIA | GNPTAB | | |
| Mucolipidosis II / IIIA Mucolipidosis III Gamma | GNPTAB GNPTG | AR | Reduced Risk |
| Mucolipidosis III Gamma | | AR AR | Reduced Risk Reduced Risk |
| Mucolipidosis III Gamma Mucolipidosis IV | GNPTG MCOLN1 | AR | Reduced Risk |
| Mucolipidosis III Gamma Mucolipidosis IV Mucopolysaccharidosis Type I | GNPTG MCOLN1 IDUA | AR AR | Reduced Risk Reduced Risk |
| Mucolipidosis III Gamma Mucolipidosis IV Mucopolysaccharidosis Type I Mucopolysaccharidosis Type II | GNPTG MCOLN1 IDUA IDS | AR AR XL | Reduced Risk Reduced Risk Reduced Risk |
| Mucolipidosis III Gamma Mucolipidosis IV Mucopolysaccharidosis Type I Mucopolysaccharidosis Type II Mucopolysaccharidosis Type IIIA | GNPTG MCOLN1 IDUA IDS SGSH | AR AR XL AR | Reduced Risk Reduced Risk Reduced Risk Reduced Risk |
| Mucolipidosis III Gamma Mucolipidosis IV Mucopolysaccharidosis Type I Mucopolysaccharidosis Type III Mucopolysaccharidosis Type IIIIA Mucopolysaccharidosis Type IIIIB | GNPTG MCOLN1 IDUA IDS SGSH NAGLU | AR AR XL AR AR | Reduced Risk Reduced Risk Reduced Risk Reduced Risk Reduced Risk Reduced Risk |
| Mucolipidosis III Gamma Mucolipidosis IV Mucopolysaccharidosis Type I Mucopolysaccharidosis Type II Mucopolysaccharidosis Type IIIA | GNPTG MCOLN1 IDUA IDS SGSH | AR AR XL AR | Reduced Risk Reduced Risk Reduced Risk Reduced Risk |





| Mucopolysaccharidosis Type IVb / GM1 Gangliosidosis | GLB1 | AR | Reduced Risk | |
|---|----------------|------|----------------------------|---|
| Mucopolysaccharidosis type IX | HYAL1 | AR | Reduced Risk | — |
| Mucopolysaccharidosis type VI | ARSB | AR | Reduced Risk Reduced Risk | |
| | SUMF1 | AR | Reduced Risk Reduced Risk | |
| Multiple Sulfatase Deficiency | 30MF1 | AR | Reduced Risk | |
| Muscle-Eye-Brain Disease and Other POMGNT1- | DOMONTA | AR | Reduced Risk | |
| Related Congenital Muscular Dystrophy- | POMGNT1 | AK | Reduced Risk | |
| Dystroglycanopathies | TYMP | A.D. | Reduced Risk | |
| Myoneurogastrointestinal Encephalopathy | | AR | | |
| Myotubular Myopathy 1 | MTM1 | XL | Reduced Risk | |
| N-Acetylglutamate Synthase Deficiency | NAGS | AR | Reduced Risk | |
| Nemaline Myopathy 2 | NEB | AR | Reduced Risk | |
| Nephrogenic Diabetes Insipidus, Type II | AQP2 | AR | Reduced Risk | |
| Nephrotic Syndrome (NPHS1-Related) / Congenital | NPHS1 | AR | Reduced Risk | |
| Finnish Nephrosis | | | | |
| Nephrotic Syndrome (NPHS2-Related) / Steroid- | NPHS2 | AR | Reduced Risk | |
| Resistant Nephrotic Syndrome | | | | |
| Neuronal Ceroid-Lipofuscinosis (CLN3-Related) | CLN3 | AR | Reduced Risk | |
| Neuronal Ceroid-Lipofuscinosis (CLN5-Related) | CLN5 | AR | Reduced Risk | |
| Neuronal Ceroid-Lipofuscinosis (CLN6-Related) | CLN6 | AR | Reduced Risk | |
| Neuronal Ceroid-Lipofuscinosis (CLN8-Related) | CLN8 | AR | Reduced Risk | |
| Neuronal Ceroid-Lipofuscinosis (MFSD8-Related) | MFSD8 | AR | Reduced Risk | |
| Neuronal Ceroid-Lipofuscinosis (PPT1-Related) | PPT1 | AR | Reduced Risk | |
| Neuronal Ceroid-Lipofuscinosis (TPP1-Related) | TPP1 | AR | Reduced Risk | |
| Niemann-Pick Disease (SMPD1-Related) | SMPD1 | AR | Reduced Risk | |
| Niemann-Pick Disease, Type C (NPC1-Related) | NPC1 | AR | Reduced Risk | |
| Niemann-Pick Disease, Type C (NPC2-Related) | NPC2 | AR | Reduced Risk | |
| Nijmegen Breakage Syndrome | NBN | AR | Reduced Risk | |
| Non-Syndromic Hearing Loss (GJB2-Related) | GJB2 | AR | Reduced Risk | |
| Odonto-Onycho-Dermal Dysplasia / Schopf-Schulz- | IV/A/Tr o A | 4.0 | Dadward Bid | |
| Passarge Syndrome | WNT10A | AR | Reduced Risk | |
| Omenn Syndrome (RAG2-Related) | RAG2 | AR | Reduced Risk | |
| Omenn Syndrome / Severe Combined | DCI DE4C | ۸D | Dadward Did | |
| Immunodeficiency, Athabaskan-Type | DCLRE1C | AR | Reduced Risk | |
| Ornithine Aminotransferase Deficiency | OAT | AR | Reduced Risk | |
| Ornithine Transcarbamylase Deficiency | OTC | XL | Reduced Risk | |
| Osteopetrosis 1 | TCIRG1 | AR | Reduced Risk | |
| Pendred Syndrome | SLC26A4 | AR | Reduced Risk | |
| Polycystic Kidney Disease, Autosomal Recessive | PKHD1 | AR | Reduced Risk | |
| Polyglandular Autoimmune Syndrome, Type 1 | AIRE | AR | Reduced Risk | |
| Pontocerebellar Hypoplasia, Type 1A | VRK1 | AR | Reduced Risk | |
| Pontocerebellar Hypoplasia, Type 6 | RARS2 | AR | Reduced Risk | |
| Primary Carnitine Deficiency | SLC22A5 | AR | Reduced Risk | |
| Primary Ciliary Dyskinesia (<i>DNAH5</i> -Related) | DNAH5 | AR | Reduced Risk | |
| Primary Citiary Dyskinesia (DNA/1-Related) | DNAI1 | AR | Reduced Risk | |
| Primary Citiary Dyskinesia (DNAI2-Related) | DNAI1 DNAI2 | AR | Reduced Risk | |
| Primary Hyperoxaluria, Type 1 | AGXT | AR | Reduced Risk Reduced Risk | |
| Primary Hyperoxaluria, Type 2 | GRHPR | AR | Reduced Risk | |
| | | | Reduced Risk | |
| Primary Hyperoxaluria, Type 3 | HOGA1 | AR | | |
| Progressive Cerebello-Cerebral Atrophy | SEPSECS | AR | Reduced Risk | |
| Progressive Familial Intrahepatic Cholestasis, Type 2 | ABCB11 | AR | Reduced Risk | |
| Propionic Acidemia (<i>PCCA</i> -Related) | PCCA | AR | Reduced Risk | |
| Propionic Acidemia (PCCB-Related) | PCCB | AR | Reduced Risk | |
| Pycnodysostosis | CTSK | AR | Reduced Risk | |
| Pyruvate Dehydrogenase E1-Alpha Deficiency | PDHA1 | XL | Reduced Risk | |
| Pyruvate Dehydrogenase E1-Beta Deficiency | PDHB | AR | Reduced Risk | |
| Renal Tubular Acidosis and Deafness | ATP6V1B1 | AR | Reduced Risk | |
| Retinitis Pigmentosa 25 | EYS | AR | Reduced Risk | |
| Retinitis Pigmentosa 26 | CERKL | AR | Reduced Risk | |
| Retinitis Pigmentosa 28 | FAM161A | AR | Reduced Risk | |
| | DHDDS | AR | | |





| Rhizomelic Chondrodysplasia Punctata, Type 1 | PEX7 | AR | Reduced Risk |
|---|----------|----|--------------|
| Rhizomelic Chondrodysplasia Punctata, Type 3 | AGPS | AR | Reduced Risk |
| Roberts Syndrome | ESCO2 | AR | Reduced Risk |
| Salla Disease | SLC17A5 | AR | Reduced Risk |
| Sandhoff Disease | HEXB | AR | Reduced Risk |
| Schimke Immunoosseous Dysplasia | SMARCAL1 | AR | Reduced Risk |
| Segawa Syndrome | TH | AR | Reduced Risk |
| Sjogren-Larsson Syndrome | ALDH3A2 | AR | Reduced Risk |
| Smith-Lemli-Opitz Syndrome | DHCR7 | AR | Reduced Risk |
| Spondylothoracic Dysostosis | MESP2 | AR | Reduced Risk |
| Steel Syndrome | COL27A1 | AR | Reduced Risk |
| Stuve-Wiedemann Syndrome | LIFR | AR | Reduced Risk |
| Sulfate Transporter-Related Osteochondrodysplasia | SLC26A2 | AR | Reduced Risk |
| Tay-Sachs Disease | HEXA | AR | Reduced Risk |
| Tyrosinemia, Type I | FAH | AR | Reduced Risk |
| Usher Syndrome, Type IB | MY07A | AR | Reduced Risk |
| Usher Syndrome, Type IC | USH1C | AR | Reduced Risk |
| Usher Syndrome, Type ID | CDH23 | AR | Reduced Risk |
| Usher Syndrome, Type IF | PCDH15 | AR | Reduced Risk |
| Usher Syndrome, Type IIA | USH2A | AR | Reduced Risk |
| Usher Syndrome, Type III | CLRN1 | AR | Reduced Risk |
| Very Long Chain Acyl-CoA Dehydrogenase Deficiency | ACADVL | AR | Reduced Risk |
| Walker-Warburg Syndrome and Other FKTN-Related Dystrophies | FKTN | AR | Reduced Risk |
| Wilson Disease | ATP7B | AR | Reduced Risk |
| Wolman Disease / Cholesteryl Ester Storage Disease | LIPA | AR | Reduced Risk |
| X-Linked Juvenile Retinoschisis | RS1 | XL | Reduced Risk |
| X-Linked Severe Combined Immunodeficiency | IL2RG | XL | Reduced Risk |
| Zellweger Syndrome Spectrum (<i>PEX10</i> -Related) | PEX10 | AR | Reduced Risk |
| Zellweger Syndrome Spectrum (<i>PEX1</i> -Related) | PEX1 | AR | Reduced Risk |
| Zellweger Syndrome Spectrum (<i>PEX2</i> -Related) | PEX2 | AR | Reduced Risk |
| Zellweger Syndrome Spectrum (PEX6-Related) | PEX6 | AR | Reduced Risk |

AR=Autosomal recessive; XL=X-linked

Test methods and comments

Genomic DNA isolated from this patient was analyzed by one or more of the following methodologies, as applicable:

Fragile X CGG Repeat Analysis (Analytical Detection Rate >99%)

PCR amplification using Asuragen, Inc. AmplideX[®] FMR1 PCR reagents followed by capillary electrophoresis for allele sizing was performed. Samples positive for FMR1 CGG repeats in the premutation and full mutation size range were further analyzed by Southern blot analysis to assess the size and methylation status of the FMR1 CGG repeat.

Genotyping (Analytical Detection Rate >99%)

Multiplex PCR amplification and allele specific primer extension analyses using the MassARRAY® System were used to identify certain recurrent variants that are complex in nature or are present in low copy repeats. Rare sequence variants may interfere with assay performance.

Multiplex Ligation-Dependent Probe Amplification (MLPA) (Analytical Detection Rate >99%)

MLPA[®] probe sets and reagents from MRC-Holland were used for copy number analysis of specific targets versus known control samples. False positive or negative results may occur due to rare sequence variants in target regions detected by MLPA probes. Analytical sensitivity and specificity of the MLPA method are both 99%.

For alpha thalassemia, the copy numbers of the *HBA1* and *HBA2* genes were analyzed. Alpha-globin gene deletions, triplications, and the Constant Spring (CS) mutation are assessed. This test is expected to detect approximately 90% of all alpha-thalassemia mutations, varying by





ethnicity. Carriers of alpha-thalassemia with three or more *HBA* copies on one chromosome, and one or no copies on the other chromosome, may not be detected. With the exception of triplications, other benign alpha-globin gene polymorphisms will not be reported. Analyses of *HBA1* and *HBA2* are performed in association with long-range PCR of the coding regions followed by short-read sequencing.

For Duchenne muscular dystrophy, the copy numbers of all *DMD* exons were analyzed. Potentially pathogenic single exon deletions and duplications are confirmed by a second method. Analysis of *DMD* is performed in association with sequencing of the coding regions.

For congenital adrenal hyperplasia, the copy number of the *CYP21A2* gene was analyzed. This analysis can detect large deletions typically due to unequal meiotic crossing-over between *CYP21A2* and the pseudogene *CYP21A1P*. Classic 30-kb deletions make up approximately 20% of *CYP21A2* pathogenic alleles. This test may also identify certain point mutations in *CYP21A2* caused by gene conversion events between *CYP21A2* and *CYP21A1P*. Some carriers may not be identified by dosage sensitive methods as this testing cannot detect individuals with two copies (duplication) of the *CYP21A2* gene on one chromosome and loss of *CYP21A2* (deletion) on the other chromosome. Analysis of *CYP21A2* is performed in association with long-range PCR of the coding regions followed by short-read sequencing.

For spinal muscular atrophy (SMA), the copy numbers of the *SMN1* and *SMN2* genes were analyzed. The individual dosage of exons 7 and 8 as well as the combined dosage of exons 1, 4, 6 and 8 of *SMN1* and *SMN2* were assessed. Copy number gains and losses can be detected with this assay. Depending on ethnicity, 6 - 29 % of carriers will not be identified by dosage sensitive methods as this testing cannot detect individuals with two copies (duplication) of the *SMN1* gene on one chromosome and loss of *SMN1* (deletion) on the other chromosome (silent 2+0 carrier) or individuals that carry an intragenic mutation in *SMN1*. Please also note that 2% of individuals diagnosed with SMA have a causative *SMN1* variant that occurred *de novo*, and therefore cannot be picked up by carrier screening in the parents. Analysis of *SMN1* is performed in association with short-read sequencing of exons 2a-7, followed by confirmation using long-range PCR (described below).

The presence of the c.*3+80T>G (chr5:70,247,901T>G) variant allele in an individual with Ashkenazi Jewish or Asian ancestry is typically indicative of a duplication of *SMN1*. When present in an Ashkenazi Jewish or Asian individual with two copies of *SMN1*, c.*3+80T>G is likely indicative of a silent (2+0) carrier. In individuals with two copies of *SMN1* with African American, Hispanic or Caucasian ancestry, the presence or absence of c.*3+80T>G significantly increases or decreases, respectively, the likelihood of being a silent 2+0 silent carrier.

MLPA for Gaucher disease (*GBA*), cystic fibrosis (*CFTR*), and non-syndromic hearing loss (*GJB2/GJB6*) will only be performed if indicated for confirmation of detected CNVs. If *GBA* analysis was performed, the copy numbers of exons 1, 3, 4, and 6 - 10 of the *GBA* gene (of 11 exons total) were analyzed. If *CFTR* analysis was performed, the copy numbers of all 27 *CFTR* exons were analyzed. If *GJB2/GJB6* analysis was performed, the copy number of the two *GJB2* exons were analyzed, as well as the presence or absence of the two upstream deletions of the *GJB2* regulatory region, del(*GJB6*-D13S1830) and del(*GJB6*-D13S1854).

Next Generation Sequencing (NGS) (Analytical Detection Rate >95%)

NGS was performed on a panel of genes for the purpose of identifying pathogenic or likely pathogenic variants.

Agilent SureSelectTMXT Low Input technology was used with a custom capture library to target the exonic regions and intron/exon splice junctions of the relevant genes, as well as a number of UTR, intronic or promoter regions that contain previously reported mutations. Libraries were pooled and sequenced on the Illumina NovaSeq 9000 platform, using paired-end 100 bp reads. The sequencing data was analyzed using a custom bioinformatics algorithm designed and validated in house.

The coding exons and splice junctions of the known protein-coding RefSeq genes were assessed for the average depth of coverage (minimum of 20X) and data quality threshold values. Most exons not meeting a minimum of >20X read depth across the exon are further analyzed by Sanger sequencing. Please note that several genomic regions present difficulties in mapping or obtaining read depth >20X. These regions, which are described below, will not be reflexed to Sanger sequencing if the mapping quality or coverage is poor. Any variants identified during testing in these regions are confirmed by a second method and reported if determined to be pathogenic or likely pathogenic. However, as there is a possibility of false negative results within these regions, detection rates and residual risks for these genes have been calculated with the presumption that variants in these exons will not be detected, unless included in the MassARRAY® genotyping platform.

Exceptions: ABCD1 (NM_000033.3) exons 8 and 9; ADA (NM_000022.2) exon 1; ADAMTS2 (NM_014244.4) exon 1; AGPS (NM_003659.3) chr2:178,257.512 - 178,257.649 (partial exon 1); ALMS1 (NM_015120.4) chr2:73,612,990 - 73,613,041 (partial exon 1); CEP290 (NM_025114.3) exon 5, exon 7, chr12:88,519,017 - 88,519,039 (partial exon 13), chr12:88,514,049 - 88,514,058 (partial exon 15), chr12:88,502,837 - 88,502,841 (partial





exon 23), chr12:88,481,551 - 88,481,589 (partial exon 32), chr12:88,471,605 - 88,471,700 (partial exon 40); *CFTR* (NM_000492.3) exon 10; *COL4A4* (NM_000092.4) chr2:227,942,604 - 227,942,619 (partial exon 25); *CYP11B2* (NM_000498.3) exons 3 - 7; *DNAI2* (NM_023036.4) chr17:72,308,136 - 72,308,147 (partial exon 12); *EVC* (NM_153717.2) exon 1; *FH* (NM_000143.3) exon 1; *GAMT* (NM_000156.5 exon 1; *GLDC* (NM_000170.2) exon 1; *GNPTAB* (NM_024312.4) chr17:4,837,000 - 4,837,400 (partial exon 2); *GNPTG* (NM_032520.4) exon 1; *HGSNAT* (NM_152419.2) exon 1; *IDS* (NM_000202.6) exon 3; *LIFR* (NM_002310.5) exon 19; *NEB* (NM_001271208.1) exons 82 - 105; *NPC1* (NM_000271.4) chr18:21,123,519 - 21,123,538 (partial exon 14); *PUS1* (NM_025215.5); chr12:132,414,446 - 132,414,532 (partial exon 2); *RPGRIP1L* (NM_015272.2) exon 23; *SGSH* (NM_000199.3) chr17:78,194,022 - 78,194,072 (partial exon 1); *SLC6A8* (NM_005629.3) exons 3 and 4.

This test will detect variants within the exons and the intron-exon boundaries of the target regions. Variants outside these regions may not be detected, including, but not limited to, UTRs, promoters, and deep intronic areas, or regions that fall into the Exceptions mentioned above. This technology may not detect all small insertion/deletions and is not diagnostic for repeat expansions and structural genomic variation. In addition, a mutation(s) in a gene not included on the panel could be present in this patient.

Variant interpretation and classification was performed based on the American College of Medical Genetics Standards and Guidelines for the Interpretation of Sequence Variants (Richards et al. 2015). All potentially pathogenic variants may be confirmed by either a specific genotyping assay or Sanger sequencing, if indicated. Any benign variants, likely benign variants or variants of uncertain significance identified during this analysis will not be reported.

Next Generation Sequencing for SMN1

Exonic regions and intron/exon splice junctions of *SMN1* and *SMN2* were captured, sequenced, and analyzed as described above. Any variants located within exons 2a-7 and classified as pathogenic or likely pathogenic were confirmed to be in either *SMN1* or *SMN2* using gene-specific long-range PCR analysis followed by Sanger sequencing. Variants located in exon 1 cannot be accurately assigned to either *SMN1* or *SMN2* using our current methodology, and so these variants are considered to be of uncertain significance and are not reported.

Copy Number Variant Analysis (Analytical Detection Rate >95%)

Large duplications and deletions were called from the relative read depths on an exon-by-exon basis using a custom exome hidden Markov model (XHMM) algorithm. Deletions or duplications determined to be pathogenic or likely pathogenic were confirmed by either a custom arrayCGH platform, quantitative PCR, or MLPA (depending on CNV size and gene content). While this algorithm is designed to pick up deletions and duplications of 2 or more exons in length, potentially pathogenic single-exon CNVs will be confirmed and reported, if detected.

Exon Array (Confirmation method) (Accuracy >99%)

The customized oligonucleotide microarray (Oxford Gene Technology) is a highly-targeted exon-focused array capable of detecting medically relevant microdeletions and microduplications at a much higher resolution than traditional aCGH methods. Each array matrix has approximately 180,000 60-mer oligonucleotide probes that cover the entire genome. This platform is designed based on human genome NCBI Build 37 (hg19) and the CGH probes are enriched to target the exonic regions of the genes in this panel.

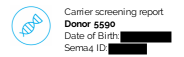
Quantitative PCR (Confirmation method) (Accuracy >99%)

The relative quantification PCR is utilized on a Roche Universal Library Probe (UPL) system, which relates the PCR signal of the target region in one group to another. To test for genomic imbalances, both sample DNA and reference DNA is amplified with primer/probe sets that specific to the target region and a control region with known genomic copy number. Relative genomic copy numbers are calculated based on the standard $\Delta\Delta$ Ct formula.

Long-Range PCR (Analytical Detection Rate >99%)

Long-range PCR was performed to generate locus-specific amplicons for *CYP21A2*, *HBA1* and *HBA2* and *GBA*. The PCR products were then prepared for short-read NGS sequencing and sequenced. Sequenced reads were mapped back to the original genomic locus and run through the bioinformatics pipeline. If indicated, copy number from MLPA was correlated with the sequencing output to analyze the results. For *CYP21A2*, a certain percentage of healthy individuals carry a duplication of the *CYP21A2* gene, which has no clinical consequences. In cases where two copies of a gene are located on the same chromosome in tandem, only the second copy will be amplified and assessed for potentially pathogenic variants, due to size limitations of the PCR reaction. However, because these alleles contain at least two copies of the *CYP21A2* gene in tandem, it is expected that this patient has at least one functional gene in the tandem allele and this patient is therefore less likely to be a carrier. When an individual carries both a duplication allele and a pathogenic variant, or multiple pathogenic variants, the current





analysis may not be able to determine the phase (cis/trans configuration) of the *CYP21A2* alleles identified. Family studies may be required in certain scenarios where phasing is required to determine the carrier status.

Residual Risk Calculations

Carrier frequencies and detection rates for each ethnicity were calculated trough the combination of internal curations of >30,000 variants and genomic frequency data from >138,000 individuals across seven ethnic groups in the gnomAD database. Additional variants in HGMD and novel deleterious variants were also incorporated into the calculation. Residual risk values are calculated using a Bayesian analysis combining the *a priori* risk of being a pathogenic mutation carrier (carrier frequency) and the detection rate. They are provided only as a guide for assessing approximate risk given a negative result, and values will vary based on the exact ethnic background of an individual. This report does not represent medical advice but should be interpreted by a genetic counselor, medical geneticist or physician skilled in genetic result interpretation and the relevant medical literature.

Personalized Residual Risk Calculations

Agilent SureSelectTMXT Low-Input technology was utilized in order to create whole-genome libraries for each patient sample. Libraries were then pooled and sequenced on the Illumina NovaSeq platform. Each sequencing lane was multiplexed to achieve 0.4-2x genome coverage, using paired-end 100 bp reads. The sequencing data underwent ancestral analysis using a customized, licensed bioinformatics algorithm that was validated in house. Identified sub-ethnic groupings were binned into one of 7 continental-level groups (African, East Asian, South Asian, Non-Finnish European, Finnish, Native American, and Ashkenazi Jewish) or, for those ethnicities that matched poorly to the continental-level groups, an 8th "unassigned" group, which were then used to select residual risk values for each gene. For individuals belonging to multiple high-level ethnic groupings, a weighting strategy was used to select the most appropriate residual risk. For genes that had insufficient data to calculate ethnic-specific residual risk values, or for sub-ethnic groupings that fell into the "unassigned" group, a "worldwide" residual risk was used. This "worldwide" residual risk was calculated using data from all available continental-level groups.

Sanger Sequencing (Confirmation method) (Accuracy >99%)

Sanger sequencing, as indicated, was performed using BigDye Terminator chemistry with the ABI 3730 DNA analyzer with target specific amplicons. It also may be used to supplement specific guaranteed target regions that fail NGS sequencing due to poor quality or low depth of coverage (<20 reads) or as a confirmatory method for NGS positive results. False negative results may occur if rare variants interfere with amplification or annealing.

Please note these tests were developed and their performance characteristics were determined by Mount Sinai Genomics, Inc. They have not been cleared or approved by the FDA. These analyses generally provide highly accurate information regarding the patient's carrier or affected status. Despite this high level of accuracy, it should be kept in mind that there are many potential sources of diagnostic error, including misidentification of samples, polymorphisms, or other rare genetic variants that interfere with analysis. Families should understand that rare diagnostic errors may occur for these reasons.

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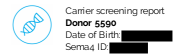
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Additional disease-specific references available upon request.