

RESULTS RECIPIENT

SEATTLE SPERM BANK

Attn: Jeffrey Olliffe 4915 25th Ave NE Ste 204W Seattle, WA 98105

Phone: (206) 588-1484 Fax: (206) 466-4696 NPI: 1306838271 Report Date: 12/21/2020 MALE DONOR 12680

DOB: Ethnicity: Mixed or Other

Caucasian

Sample Type: EDTA Blood Date of Collection: 12/08/2020 Date Received: 12/10/2020 Date Tested: 12/18/2020 Barcode: 11004512731825 Accession ID: CSL4ZF4EXYFD9F6 Indication: Egg or sperm donor FEMALE N/A

Foresight® Carrier Screen

NEGATIVE

ABOUT THIS TEST

The **Myriad Foresight Carrier Screen** utilizes sequencing, maximizing coverage across all DNA regions tested, to help you learn about your chance to have a child with a genetic disease.

RESULTS SUMMARY

Risk Details	DONOR 12680	Partner
Panel Information	Foresight Carrier Screen Universal Panel Fundamental Plus Panel Fundamental Panel (175 conditions tested)	N/A
All conditions tested A complete list of all conditions tested can be found on page 5.	□ NEGATIVE No disease-causing mutations were detected.	N/A

CLINICAL NOTES

None

NEXT STEPS

• If necessary, patients can discuss residual risks with their physician or a genetic counselor.



MALE
DONOR 12680
DOB:

Ethnicity: Mixed or Other

Caucasian

Barcode: 11004512731825

FEMALE N/A

Methods and Limitations

DONOR 12680 [Foresight Carrier Screen]: Sequencing with copy number analysis, spinal muscular atrophy, analysis of homologous regions, and alpha thalassemia (HBA1/HBA2) sequencing with targeted copy number analysis (Assay(s): DTS v3.2).

Sequencing with copy number analysis

High-throughput sequencing and read-depth-based copy number analysis are used to analyze the genes listed in the Conditions Tested section of the report. Except where otherwise noted, the region of interest (ROI) comprises the indicated coding regions and 20 non-coding bases flanking each region. In a minority of cases where genomic features (e.g., long homopolymers) compromise calling fidelity, the affected non-coding bases are excluded from the ROI. The ROI is sequenced to a minimum acceptable read depth, and the sequences are compared to a reference genomic sequence (Genome Reference Consortium Human Build 37 [GRCh37]/hg19). On average, 99% of all bases in the ROI are sequenced at a read depth that is greater than the minimum read depth. Sequence variants may not be detected in areas of lower sequence coverage. Insertions and deletions may not be detected as accurately as single-nucleotide variants. Select genes or regions for which pseudogenes or other regions of homology impede reliable variant detection may be assayed using alternate technology, or they may be excluded from the ROI. *CFTR* and *DMD* testing includes analysis for exon-level deletions and duplications with an average sensitivity of ~99%. Only exon-level deletions are assayed for other genes on the panel and such deletions are detected with a sensitivity of ≥75%. Selected founder deletions may be detected at slightly higher sensitivity. Affected exons and/or breakpoints of copy number variants are estimated from junction reads, where available, or using the positions of affected probes. Only exons known to be included in the region affected by a copy number variant are provided in the variant nomenclature. In some cases, the copy number variant may be larger or smaller than indicated. If *GJB2* is tested, large upstream deletions involving the *GJB6* and/or *CRYL1* genes that may affect the expression of *GJB2* are also analyzed.

Spinal muscular atrophy

Targeted copy number analysis via high-throughput sequencing is used to determine the copy number of exon 7 of the *SMN1* gene. Other genetic variants may interfere with this analysis. Some individuals with two copies of *SMN1* are "silent" carriers with both *SMN1* genes on one chromosome and no copies of the gene on the other chromosome. This is more likely in individuals who have two copies of the *SMN1* gene and are positive for the g.27134T>G single-nucleotide polymorphism (SNP) (PMID: 9199562, 23788250, and 28676062), which affects the reported residual risk; Ashkenazi Jewish or Asian patients with this genotype have a high post-test likelihood of being carriers for SMA and are reported as carriers. The g.27134T>G SNP is only reported in individuals who have two copies of *SMN1*.

Analysis of homologous regions

A combination of high-throughput sequencing, read-depth-based copy number analysis, and targeted genotyping is used to determine the number of functional gene copies and/or the presence of selected loss-of-function variants in certain genes that have homology to other genomic regions. The precise breakpoints of large deletions in these genes cannot be determined but are instead estimated from copy number analysis. Pseudogenes may interfere with this analysis, especially when many pseudogene copies are present.

If CYP21A2 is tested, patients who have one or more additional copies of the CYP21A2 gene and a pathogenic variant may or may not be a carrier of 21-hydroxylase deficient CAH, depending on the chromosomal location of the variants (phase). Benign CYP21A2 gene duplications and/or triplications will only be reported in this context. Some individuals with two functional CYP21A2 gene copies may be "silent" carriers, with two gene copies resulting from a duplication on one chromosome and a gene deletion on the other chromosome. This and other similar rare carrier states, where complementary changes exist between the chromosomes, may not be detected by the assay. Given that the true incidence of non-classic CAH is unknown, the residual carrier and reproductive risk numbers on the report are based only on the published incidence for classic CAH. However, the published prevalence of non-classic CAH is highest in individuals of Ashkenazi Jewish, Hispanic, Italian, and Yugoslav descent. Therefore, the residual and reproductive risks are likely an underestimate for CAH, especially in the aforementioned populations, as they do not account for non-classic CAH.



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Alpha thalassemia (HBA1/HBA2) sequencing with targeted copy number analysis

High-throughput sequencing and read-depth-based copy number analysis are used to identify sequence variation and functional gene copies within the region of interest (ROI) of *HBA1* and *HBA2*, which includes the listed exons plus 20 intronic flanking bases. In a minority of cases where genomic features (e.g., long homopolymers) compromise calling fidelity, the affected intronic bases are not included in the ROI. The ROI is sequenced to a minimum acceptable read depth, and the sequences are compared to a reference genomic sequence (Genome Reference Consortium Human Build 37 [GRCh37]/hg19). On average, 99% of all bases in the ROI are sequenced at a read depth that is greater than the minimum read depth. Sequence variants may not be detected in areas of lower sequence coverage. Insertions and deletions may not be detected as accurately as single-nucleotide variants. For large deletions or duplications in these genes, the precise breakpoints cannot be determined but are instead estimated from copy number analysis. This assay has been validated to detect up to two additional copies of each alpha globin gene. In rare instances where assay results suggest greater than two additional copies are present, this will be noted but the specific number of gene copies observed will not be provided.

Extensive sequence homology exists between *HBA1* and *HBA2*. This sequence homology can prevent certain variants from being localized to one gene over the other. In these instances, variant nomenclature will be provided for both genes. If follow-up testing is indicated for patients with the nomenclature provided for both genes, both *HBA1* and *HBA2* should be tested. Some individuals with four functional alpha globin gene copies may be "silent" carriers, with three gene copies resulting from triplication on one chromosome and a single gene deletion on the other chromosome. This and other similar rare carrier states, where complementary changes exist between the chromosomes, may not be detected by the assay.

Interpretation of reported variants

Variants are classified according to internally defined criteria, which are compatible with the ACMG Standards and Guidelines for the Interpretation of Sequence Variants (PMID: 25741868). Variants that are considered disease-causing by Myriad Women's Health (MWH) are reported. In addition, variants that have not previously been established as a recognized cause of disease may be identified. In these cases, only variants classified as 'likely' pathogenic are reported. Likely pathogenic variants are described elsewhere in the report as 'likely to have a negative impact on gene function.' Variant pathogenicity is evaluated and classified by an assessment of the nature of the variant and reviews of reports of allele frequencies in cases and controls, functional studies, variant annotation and effect prediction, segregation studies, and other resources, where available. Exon-level duplications in the *DMD* and *CFTR* genes are assumed to be in tandem and are classified according to their predicted effect on the reading frame. Benign variants, variants of uncertain significance, and variants not directly associated with the specified disease phenotype(s) are not reported.

Limitations

The MWH Foresight Carrier Screen is designed to detect and report germline (constitutional) alterations. Mosaic (somatic) variation may not be detected, and if it is detected, it may not be reported. If more than one variant is detected in a gene, additional studies may be necessary to determine if those variants lie on the same chromosome or different chromosomes (phase). This test is not designed to detect sex-chromosome copy number variations. If present, sex-chromosome abnormalities may significantly reduce test sensitivity for X-linked conditions. Variant interpretation and residual and reproductive risk estimations assume a normal karyotype and may be different for individuals with abnormal karyotypes. The test does not fully address all inherited forms of intellectual disability, birth defects, or heritable diseases. Furthermore, not all forms of genetic variation are detected by this assay (i.e., duplications [except in specified genes], chromosomal rearrangements, structural abnormalities, etc.). Additional testing may be appropriate for some individuals. Pseudogenes and other regions of homology may interfere with this analysis. In an unknown number of cases, other genetic variation may interfere with variant detection. Rare carrier states where complementary changes exist between the chromosomes may not be detected by the assay. Other possible sources of diagnostic error include sample mix-up, trace contamination, bone marrow transplantation, blood transfusions, and technical or analytical errors.

Detection rates are determined using published scientific literature and/or reputable databases, when available, to estimate the fraction of disease alleles, weighted by frequency, that the methodology is predicted to be able or unable to detect. Detection rates are approximate and only account for analytical sensitivity. Certain variants that have been previously described in the literature may not be reported, if there is insufficient evidence for pathogenicity. Detection rates do not account for the disease specific rates of *de novo* variation.

This test was developed, and its performance characteristics determined by, Myriad Women's Health, Inc. It has not been cleared or approved by the US Food and Drug Administration (FDA). The FDA does not require this test to go through premarket review. This test is used for clinical purposes. It should not be regarded as investigational or for research. This laboratory is certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA) as qualified to perform high-complexity clinical testing. These results are adjunctive to the ordering physician's evaluation. CLIA Number: #05D1102604.

Incidental Findings



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Unless otherwise indicated, these results and interpretations are limited to the specific disease panel(s) requested by the ordering healthcare provider. In some cases, standard data analyses may identify genetic findings beyond the region(s) of interest specified by the test, and such findings may not be reported. These findings may include genomic abnormalities with major, minor, or no, clinical significance.

If you have questions or would like more information about any of the test methods or limitations, please contact (888) 268-6795.

Resources

GENOME CONNECT | http://www.genomeconnect.org

Patients can share their reports using research registries such as Genome Connect, an online research registry building a genetics and health knowledge base. Genome Connect provides patients, physicians, and researchers an opportunity to share genetic information to support the study of the impact of genetic variation on health conditions.

SENIOR LABORATORY DIRECTOR

Karla R. Bowles, PhD, FACMG, CGMB

Karla R Boules

Report content approved by Karla Bowles, PhD, FACMG, CGMB on Dec 21, 2020



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Conditions Tested

11-beta-hydroxylase-deficient Congenital Adrenal Hyperplasia - Gene: CYP11B1. Autosomal Recessive. Sequencing with copy number analysis. **Exons:**

NM_000497:1-9. Detection Rate: Mixed or Other Caucasian 94%.

6-pyruvoyl-tetrahydropterin Synthase Deficiency - Gene: PTS. Autosomal

Recessive. Sequencing with copy number analysis. Exons: NM_000317:1-6. Detection Rate: Mixed or Other Caucasian >99%.

ABCC8-related Familial Hyperinsulinism - Gene: ABCC8. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000352:1-39. Detection Rate: Mixed or Other Caucasian >99%.

Adenosine Deaminase Deficiency - Gene: ADA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000022:1-12. Detection Rate: Mixed or Other Caucasian >99%.

Alpha Thalassemia, HBA1/HBA2-related - Genes: HBA1, HBA2. Autosomal Recessive. Alpha thalassemia (HBA1/HBA2) sequencing with targeted copy number analysis. Exons: NM_000517:1-3; NM_000558:1-3. Variants (16): -(alpha)20.5, --BRIT, --MEDI, --MEDII, --SEA, --THAI or --FIL, -alpha3.7, -alpha4.2, HBA1+HBA2 deletion, Hb Constant Spring, Poly(A) AATAAA>AATA--, Poly(A) AATAAA>AATAAG, Poly(A) AATAAA>AATGAA, anti3.7, anti4.2, del HS-40. Detection Rate: Not calculated due to rarity of disease in this individual's reported ethnicity.

Alpha-mannosidosis - **Gene:** MAN2B1. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000528:1-23. **Detection Rate:** Mixed or Other Caucasian >99%.

Alpha-sarcoglycanopathy - **Gene:** SGCA. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000023:1-9. **Detection Rate:** Mixed or Other Caucasian >99%.

Alstrom Syndrome - Gene: ALMS1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_015120:1-23. Detection Rate: Mixed or Other Caucasian >99%.

AMT-related Glycine Encephalopathy - Gene: AMT. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000481:1-9. Detection Rate: Mixed or Other Caucasian >99%.

Andermann Syndrome - **Gene**: SLC12A6. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_133647:1-25. **Detection Rate**: Mixed or Other Caucasian >99%.

Argininemia - Gene: ARG1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000045:1-8. Detection Rate: Mixed or Other Caucasian 97%. Argininosuccinic Aciduria - Gene: ASL. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_001024943:1-16. Detection Rate: Mixed or Other Caucasian >99%.

Aspartylglucosaminuria - Gene: AGA. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000027:1-9. **Detection Rate:** Mixed or Other Caucasian >99%.

Ataxia with Vitamin E Deficiency - Gene: TTPA. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000370:1-5. **Detection Rate:** Mixed or Other Caucasian >99%.

Ataxia-telangiectasia - Gene: ATM. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000051:2-63. Detection Rate: Mixed or Other Caucasian 98%

ATP7A-related Disorders - Gene: ATP7A. X-linked Recessive. Sequencing with copy number analysis. Exons: NM_000052:2-23. Detection Rate: Mixed or Other Caucasian 96%.

Autoimmune Polyglandular Syndrome Type 1 - Gene: AIRE. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000383:1-14. **Detection Rate:** Mixed or Other Caucasian >99%.

Autosomal Recessive Osteopetrosis Type 1 - Gene: TCIRG1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_006019:2-20. **Detection Rate:** Mixed or Other Caucasian >99%.

Autosomal Recessive Polycystic Kidney Disease, PKHD1-related - Gene: PKHD1.

Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_138694 2-67. Detection Rate: Mixed or Other Caucasian >99%.

Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay - Gene: SACS.

Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_014363 2-10. Detection Rate: Mixed or Other Caucasian 99%.

Bardet-Biedl Syndrome, BBS1-related - Gene: BBS1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_024649:1-17. Detection Rate: Mixed or Other Caucasian >99%.

Bardet-Biedl Syndrome, BBS10-related - Gene: BBS10. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_024685:1-2. **Detection Rate:** Mixed or Other Caucasian >99%.

Bardet-Biedl Syndrome, BBS12-related - Gene: BBS12. Autosomal Recessive. Sequencing with copy number analysis. **Exon:** NM_152618:2. **Detection Rate:** Mixed or Other Caucasian >99%.

Bardet-Biedl Syndrome, BBS2-related - **Gene**: BBS2. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_031885:1-17. **Detection Rate**: Mixed or Other Caucasian >99%.

BCS1L-related Disorders - Gene: BCS1L. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_004328:3-9. Detection Rate: Mixed or Other Caucasian >99%.

Beta-sarcoglycanopathy - **Gene**: SGCB. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_000232:1-6. **Detection Rate**: Mixed or Other Caucasian >99%.

Biotinidase Deficiency - **Gene**: BTD. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_000060:1-4. **Detection Rate**: Mixed or Other Caucasian >99%.

Bloom Syndrome - Gene: BLM. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000057:2-22. **Detection Rate:** Mixed or Other Caucasian >99%

Calpainopathy - Gene: CAPN3. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000070:1-24. Detection Rate: Mixed or Other Caucasian >99%.

Canavan Disease - **Gene**: ASPA. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_000049:1-6. **Detection Rate**: Mixed or Other Caucasian 98%.

Carbamoylphosphate Synthetase I Deficiency - Gene: CPS1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_001875:1-38. Detection Rate: Mixed or Other Caucasian >99%.

Carnitine Palmitoyltransferase IA Deficiency - Gene: CPT1A. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_001876:2-19. Detection Rate: Mixed or Other Caucasian >99%.

Carnitine Palmitoyltransferase II Deficiency - Gene: CPT2. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000098:1-5. Detection Rate: Mixed or Other Caucasian >99%.

Cartilage-hair Hypoplasia - Gene: RMRP. Autosomal Recessive. Sequencing with copy number analysis. **Exon:** NR_003051:1. **Detection Rate:** Mixed or Other Caucasian >99%.

Cerebrotendinous Xanthomatosis - Gene: CYP27A1. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000784:1-9. **Detection Rate:** Mixed or Other Caucasian >99%.

Citrullinemia Type 1 - Gene: ASS1. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_000050:3-16. **Detection Rate**: Mixed or Other Caucasian >99%.

CLN3-related Neuronal Ceroid Lipofuscinosis - Gene: CLN3. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_001042432 2-16. Detection Rate: Mixed or Other Caucasian >99%.



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CLN5-related Neuronal Ceroid Lipofuscinosis - Gene: CLN5. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_006493:1-4. **Detection Rate:** Mixed or Other Caucasian >99%.

CLN6-related Neuronal Ceroid Lipofuscinosis - Gene: CLN6. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_017882:1-7. **Detection Rate:** Mixed or Other Caucasian >99%.

CLN8-related Neuronal Ceroid Lipofuscinosis - Gene: CLN8. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_018941:2-3. **Detection Rate:** Mixed or Other Caucasian >99%.

Cohen Syndrome - Gene: VPS13B. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_017890:2-62. **Detection Rate:** Mixed or Other Caucasian 97%.

COL4A3-related Alport Syndrome - Gene: COL4A3. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000091:1-52. Detection Rate: Mixed or Other Caucasian 97%.

COL4A4-related Alport Syndrome - Gene: COL4A4. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000092:2-48. Detection Rate: Mixed or Other Caucasian 98%.

Combined Pituitary Hormone Deficiency, PROP1-related - Gene: PROP1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_006261:1-3. Detection Rate: Mixed or Other Caucasian >99%.

Congenital Adrenal Hyperplasia, CYP21A2-related - Gene: CYP21A2. Autosomal Recessive. Analysis of homologous regions. Variants (13): CYP21A2 deletion, CYP21A2 duplication, CYP21A2 triplication, G111Vfs*21, I173N, L308Ffs*6, P31L, Q319*, Q319*+CYP21A2dup, R357W, V282L, [I237N;V238E;M240K], c.293-13C>G. Detection Rate: Mixed or Other Caucasian 96%.

Congenital Disorder of Glycosylation Type Ia - Gene: PMM2. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000303:1-8. Detection Rate: Mixed or Other Caucasian >99%.

Congenital Disorder of Glycosylation Type Ic - Gene: ALG6. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_013339:2-15. Detection Rate: Mixed or Other Caucasian >99%.

Congenital Disorder of Glycosylation, MPI-related - Gene: MPI. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_002435:1-8. Detection Rate: Mixed or Other Caucasian >99%.

Costeff Optic Atrophy Syndrome - Gene: OPA3. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_025136:1-2. **Detection Rate:** Mixed or Other Caucasian >99%.

Cystic Fibrosis - **Gene**: CFTR. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_000492:1-27. IVS8-5T allele analysis is only reported in the presence of the R117H mutation. **Detection Rate**: Mixed or Other Caucasian >99%.

Cystinosis - **Gene**: CTNS. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_004937:3-12. **Detection Rate**: Mixed or Other Caucasian >99%.

D-bifunctional Protein Deficiency - Gene: HSD17B4. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000414:1-24. **Detection Rate:** Mixed or Other Caucasian 98%.

Delta-sarcoglycanopathy - **Gene**: SGCD. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_000337:2-9. **Detection Rate**: Mixed or Other Caucasian 99%.

Dihydrolipoamide Dehydrogenase Deficiency - Gene: DLD. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_000108:1-14. **Detection Rate**: Mixed or Other Caucasian >99%.

Dysferlinopathy - **Gene**: DYSF. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_003494:1-55. **Detection Rate**: Mixed or Other Caucasian 98%.

Dystrophinopathy (Including Duchenne/Becker Muscular Dystrophy) - Gene: DMD. X-linked Recessive. Sequencing with copy number analysis. Exons: NM_004006:1-79. Detection Rate: Mixed or Other Caucasian >99%.

ERCC6-related Disorders - **Gene**: ERCC6. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_000124:2-21. **Detection Rate**: Mixed or Other Caucasian 99%.

ERCC8-related Disorders - Gene: ERCC8. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000082:1-12. **Detection Rate:** Mixed or Other Caucasian 95%.

EVC-related Ellis-van Creveld Syndrome - Gene: EVC. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_153717:1-21. **Detection Rate:** Mixed or Other Caucasian 96%.

EVC2-related Ellis-van Creveld Syndrome - Gene: EVC2. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_147127:1-22. **Detection Rate:** Mixed or Other Caucasian >99%.

Fabry Disease - Gene: GLA. X-linked Recessive. Sequencing with copy number analysis. Exons: NM_000169:1-7. Detection Rate: Mixed or Other Caucasian 98%. Familial Dysautonomia - Gene: IKBKAP. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_003640:2-37. Detection Rate: Mixed or Other Caucasian >99%.

Familial Mediterranean Fever - Gene: MEFV. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000243:1-10. Detection Rate: Mixed or Other Caucasian >99%.

Fanconi Anemia Complementation Group A - Gene: FANCA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000135:1-43. Detection Rate: Mixed or Other Caucasian 92%.

Fanconi Anemia, FANCC-related - Gene: FANCC. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000136:2-15. **Detection Rate:** Mixed or Other Caucasian >99%.

FKRP-related Disorders - Gene: FKRP. Autosomal Recessive. Sequencing with copy number analysis. **Exon:** NM_024301:4. **Detection Rate:** Mixed or Other Caucasian >99%

FKTN-related Disorders - Gene: FKTN. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_001079802:3-11. **Detection Rate**: Mixed or Other Caucasian >99%.

Free Sialic Acid Storage Disorders - Gene: SLC17A5. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_012434:1-11. Detection Rate: Mixed or Other Caucasian 98%.

Galactokinase Deficiency - **Gene:** GALK1. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000154:1-8. **Detection Rate:** Mixed or Other Caucasian >99%.

Galactosemia - **Gene:** GALT. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000155:1-11. **Detection Rate:** Mixed or Other Caucasian >99%.

Gamma-sarcoglycanopathy - **Gene:** SGCG. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000231:2-8. **Detection Rate:** Mixed or Other Caucasian 88%.

Gaucher Disease - **Gene:** GBA. Autosomal Recessive. Analysis of homologous regions. **Variants (10):** D409V, D448H, IVS2+1G>A, L444P, N370S, R463C, R463H, R496H, V394L, p.L29Afs*18. **Detection Rate:** Mixed or Other Caucasian 60%.

GJB2-related DFNB1 Nonsyndromic Hearing Loss and Deafness - Gene: GJB2. Autosomal Recessive. Sequencing with copy number analysis. Exons:

NM_004004:1-2. Detection Rate: Mixed or Other Caucasian >99%.

GLB1-related Disorders - Gene: GLB1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000404:1-16. Detection Rate: Mixed or Other Caucasian >99%

GLDC-related Glycine Encephalopathy - **Gene:** GLDC. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000170:1-25. **Detection Rate:** Mixed or Other Caucasian 94%.

Glutaric Acidemia, GCDH-related - Gene: GCDH. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000159:2-12. **Detection Rate:** Mixed or Other Caucasian >99%.

Glycogen Storage Disease Type la - Gene: G6PC. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000151:1-5. **Detection Rate:** Mixed or Other Caucasian >99%.

Glycogen Storage Disease Type Ib - Gene: SLC37A4. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_001164277 3-11. **Detection Rate:** Mixed or Other Caucasian >99%.



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Glycogen Storage Disease Type III - Gene: AGL. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000642:2-34. **Detection Rate:** Mixed or Other Caucasian >99%.

GNE Myopathy - **Gene**: GNE. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_001128227:1-12. **Detection Rate**: Mixed or Other Caucasian >99%.

GNPTAB-related Disorders - Gene: GNPTAB. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_024312:1-21. Detection Rate: Mixed or Other Caucasian >99%.

HADHA-related Disorders - Gene: HADHA. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_000182:1-20. **Detection Rate**: Mixed or Other Caucasian >99%.

Hb Beta Chain-related Hemoglobinopathy (Including Beta Thalassemia and Sickle Cell Disease) - Gene: HBB. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000518:1-3. Detection Rate: Mixed or Other Caucasian >99%. Hereditary Fructose Intolerance - Gene: ALDOB. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000035:2-9. Detection Rate: Mixed or Other Caucasian >99%.

Herlitz Junctional Epidermolysis Bullosa, LAMB3-related - Gene: LAMB3. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000228 2-23. Detection Rate: Mixed or Other Caucasian >99%.

Hexosaminidase A Deficiency (Including Tay-Sachs Disease) - Gene: HEXA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000520:1-14. Detection Rate: Mixed or Other Caucasian >99%.

HMG-CoA Lyase Deficiency - Gene: HMGCL. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000191:1-9. **Detection Rate:** Mixed or Other Caucasian 98%.

Holocarboxylase Synthetase Deficiency - Gene: HLCS. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000411:4-12. **Detection Rate:** Mixed or Other Caucasian >99%.

Homocystinuria, CBS-related - Gene: CBS. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000071:3-17. Detection Rate: Mixed or Other Caucasian >99%

Hydrolethalus Syndrome - **Gene**: HYLS1. Autosomal Recessive. Sequencing with copy number analysis. **Exon**: NM_145014:4. **Detection Rate**: Mixed or Other Caucasian >99%.

Hypophosphatasia - Gene: ALPL. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000478:2-12. **Detection Rate:** Mixed or Other Caucasian >99%.

Isovaleric Acidemia - **Gene**: IVD. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_002225:1-12. **Detection Rate**: Mixed or Other Caucasian > 90%

Joubert Syndrome 2 - Gene: TMEM216. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_001173990:1-5. **Detection Rate:** Mixed or Other Caucasian >99%.

Junctional Epidermolysis Bullosa, LAMA3-related - Gene: LAMA3. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000227:1-38. Detection Rate: Mixed or Other Caucasian >99%.

Junctional Epidermolysis Bullosa, LAMC2-related - Gene: LAMC2. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_005562:1-23. Detection Rate: Mixed or Other Caucasian >99%.

KCNJ11-related Familial Hyperinsulinism - Gene: KCNJ11. Autosomal Recessive. Sequencing with copy number analysis. Exon: NM_000525:1. Detection Rate: Mixed or Other Caucasian >99%.

Krabbe Disease - **Gene:** GALC. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000153:1-17. **Detection Rate:** Mixed or Other Caucasian >99%.

LAMA2-related Muscular Dystrophy - Gene: LAMA2. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000426:1-43,45-65. Detection Rate: Mixed or Other Caucasian >99%.

Leigh Syndrome, French-Canadian Type - Gene: LRPPRC. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_133259:1-38. **Detection Rate:** Mixed or Other Caucasian >99%.

Lipoid Congenital Adrenal Hyperplasia - Gene: STAR. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000349:1-7. **Detection Rate:** Mixed or Other Caucasian >99%.

Lysosomal Acid Lipase Deficiency - Gene: LIPA. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000235:2-10. **Detection Rate:** Mixed or Other Caucasian >99%.

Maple Syrup Urine Disease Type Ia - Gene: BCKDHA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000709:1-9. Detection Rate: Mixed or Other Caucasian >99%.

Maple Syrup Urine Disease Type Ib - Gene: BCKDHB. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_183050:1-10. Detection Rate: Mixed or Other Caucasian >99%.

Maple Syrup Urine Disease Type II - Gene: DBT. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_001918:1-11. Detection Rate: Mixed or Other Caucasian 96%.

Medium Chain Acyl-CoA Dehydrogenase Deficiency - Gene: ACADM. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000016:1-12. Detection Rate: Mixed or Other Caucasian >99%.

Megalencephalic Leukoencephalopathy with Subcortical Cysts - Gene: MLC1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_015166 2-12. Detection Rate: Mixed or Other Caucasian >99%.

Metachromatic Leukodystrophy - **Gene**: ARSA. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_000487:1-8. **Detection Rate**: Mixed or Other Caucasian >99%.

Methylmalonic Acidemia, cblA Type - Gene: MMAA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_172250:2-7. Detection Rate: Mixed or Other Caucasian >99%.

Methylmalonic Acidemia, cblB Type - Gene: MMAB. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_052845:1-9. Detection Rate: Mixed or Other Caucasian >99%.

Methylmalonic Aciduria and Homocystinuria, cblC Type - Gene: MMACHC. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_015506:1-4. Detection Rate: Mixed or Other Caucasian >99%.

MKS1-related Disorders - Gene: MKS1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_017777:1-18. Detection Rate: Mixed or Other Caucasian >99%.

Mucolipidosis III Gamma - **Gene**: GNPTG. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_032520:1-11. **Detection Rate**: Mixed or Other Caucasian >99%.

Mucolipidosis IV - Gene: MCOLN1. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_020533:1-14. **Detection Rate:** Mixed or Other Caucasian >99%.

Mucopolysaccharidosis Type I - **Gene**: IDUA. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_000203:1-14. **Detection Rate**: Mixed or Other Caucasian >99%.

Mucopolysaccharidosis Type II - Gene: IDS. X-linked Recessive. Sequencing with copy number analysis. **Exons:** NM_000202:1-9. **Detection Rate:** Mixed or Other Caucasian 88%.

Mucopolysaccharidosis Type IIIA - Gene: SGSH. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000199:1-8. **Detection Rate:** Mixed or Other Caucasian >99%.

Mucopolysaccharidosis Type IIIB - Gene: NAGLU. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000263:1-6. **Detection Rate:** Mixed or Other Caucasian >99%.

Mucopolysaccharidosis Type IIIC - Gene: HGSNAT. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_152419:1-18. Detection Rate: Mixed or Other Caucasian >99%.

MUT-related Methylmalonic Acidemia - Gene: MUT. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000255:2-13. Detection Rate: Mixed or Other Caucasian >99%.

MYO7A-related Disorders - Gene: MYO7A. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_000260:2-49. **Detection Rate**: Mixed or Other Caucasian >99%.



MALE DONOR 12680

DOB: Ethnicity: Mixed or Other

Caucasian

Barcode: 11004512731825

FEMALE N/A

Primary Carnitine Deficiency - Gene: SLC22A5. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_003060:1-10. Detection Rate: Mixed or Other Caucasian >99%.

Primary Hyperoxaluria Type 1 - Gene: AGXT. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000030:1-11. **Detection Rate:** Mixed or Other Caucasian >99%.

Primary Hyperoxaluria Type 2 - Gene: GRHPR. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_012203:1-9. Detection Rate: Mixed or Other Caucasian >99%.

Primary Hyperoxaluria Type 3 - Gene: HOGA1. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_138413:1-7. **Detection Rate:** Mixed or Other Caucasian >99%.

Pycnodysostosis - **Gene:** CTSK. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000396:2-8. **Detection Rate:** Mixed or Other Caucasian >99%.

Pyruvate Carboxylase Deficiency - Gene: PC. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_000920:3-22. **Detection Rate**: Mixed or Other Caucasian >99%.

Rhizomelic Chondrodysplasia Punctata Type 1 - Gene: PEX7. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000288:1-10. Detection Rate: Mixed or Other Caucasian >99%.

RTEL1-related Disorders - Gene: RTEL1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_032957:2-35. Detection Rate: Mixed or Other Caucasian >99%.

Sandhoff Disease - Gene: HEXB. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000521:1-14. Detection Rate: Mixed or Other Caucasian >99%.

Short-chain Acyl-CoA Dehydrogenase Deficiency - Gene: ACADS. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000017:1-10. Detection Rate: Mixed or Other Caucasian >99%.

Sjogren-Larsson Syndrome - Gene: ALDH3A2. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000382:1-10. **Detection Rate:** Mixed or Other Caucasian 96%.

SLC26A2-related Disorders - Gene: SLC26A2. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000112:2-3. Detection Rate: Mixed or Other Caucasian > 99%.

Smith-Lemli-Opitz Syndrome - Gene: DHCR7. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_001360:3-9. Detection Rate: Mixed or Other Caucasian >99%.

Spastic Paraplegia Type 15 - Gene: ZFYVE26. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_015346:2-42. **Detection Rate**: Mixed or Other Caucasian >99%.

Spinal Muscular Atrophy - Gene: SMN1. Autosomal Recessive. Spinal muscular atrophy. **Variant** (1): SMN1 copy number. **Detection Rate:** Mixed or Other Caucasian 95%.

Spondylothoracic Dysostosis - Gene: MESP2. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_001039958:1-2. Detection Rate: Mixed or Other Caucasian >99%.

TGM1-related Autosomal Recessive Congenital Ichthyosis - Gene: TGM1.
Autosomal Recessive. Sequencing with copy number analysis. Exons:

NM 000359 2-15. Detection Rate: Mixed or Other Caucasian >99%.

TPP1-related Neuronal Ceroid Lipofuscinosis - Gene: TPP1. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000391:1-13. **Detection Rate:** Mixed or Other Caucasian >99%.

Tyrosine Hydroxylase Deficiency - Gene: TH. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_199292:1-14. **Detection Rate:** Mixed or Other Caucasian >99%.

Tyrosinemia Type I - Gene: FAH. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000137:1-14. **Detection Rate:** Mixed or Other Caucasian >99%.

Tyrosinemia Type II - Gene: TAT. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000353:2-12. **Detection Rate:** Mixed or Other Caucasian >99%.

NEB-related Nemaline Myopathy - Gene: NEB. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_001271208:3-80,117-183. **Detection Rate:** Mixed or Other Caucasian 92%.

Nephrotic Syndrome, NPHS1-related - Gene: NPHS1. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_004646:1-29. **Detection Rate:** Mixed or Other Caucasian >99%.

Nephrotic Syndrome, NPHS2-related - Gene: NPHS2. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_014625:1-8. **Detection Rate:** Mixed or Other Caucasian >99%.

Niemann-Pick Disease Type C1 - Gene: NPC1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000271:1-25. Detection Rate: Mixed or Other Caucasian >99%.

Niemann-Pick Disease Type C2 - Gene: NPC2. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_006432:1-5. Detection Rate: Mixed or Other Caucasian >99%.

Niemann-Pick Disease, SMPD1-related - Gene: SMPD1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000543:1-6. Detection Rate: Mixed or Other Caucasian >99%.

Nijmegen Breakage Syndrome - Gene: NBN. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_002485:1-16. Detection Rate: Mixed or Other Caucasian >99%.

Ornithine Transcarbamylase Deficiency - Gene: OTC. X-linked Recessive. Sequencing with copy number analysis. Exons: NM_000531:1-10. Detection Rate: Mixed or Other Caucasian 97%.

PCCA-related Propionic Acidemia - Gene: PCCA. Autosomal Recessive.

Sequencing with copy number analysis. Exons: NM_000282:1-24. Detection Rate: Mixed or Other Caucasian 95%.

PCCB-related Propionic Acidemia - Gene: PCCB. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000532:1-15. Detection Rate: Mixed or Other Caucasian >99%.

PCDH15-related Disorders - Gene: PCDH15. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_033056:2-33. Detection Rate: Mixed or Other Caucasian 93%.

Pendred Syndrome - Gene: SLC26A4. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000441:2-21. **Detection Rate:** Mixed or Other Caucasian >99%.

Peroxisome Biogenesis Disorder Type 1 - Gene: PEX1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000466:1-24. Detection Rate: Mixed or Other Caucasian >99%.

Peroxisome Biogenesis Disorder Type 3 - Gene: PEX12. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000286:1-3. **Detection Rate:** Mixed or Other Caucasian >99%.

Peroxisome Biogenesis Disorder Type 4 - Gene: PEX6. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000287:1-17. **Detection Rate:** Mixed or Other Caucasian 97%.

Peroxisome Biogenesis Disorder Type 5 - Gene: PEX2. Autosomal Recessive. Sequencing with copy number analysis. Exon: NM_000318:4. Detection Rate: Mixed or Other Caucasian >99%.

Peroxisome Biogenesis Disorder Type 6 - Gene: PEX10. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_153818:1-6. **Detection Rate:** Mixed or Other Caucasian >99%.

Phenylalanine Hydroxylase Deficiency - Gene: PAH. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000277:1-13. Detection Rate: Mixed or Other Caucasian >99%.

POMGNT-related Disorders - Gene: POMGNT1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_017739:2-22. Detection Rate: Mixed or Other Caucasian 96%.

Pompe Disease - Gene: GAA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000152:2-20. Detection Rate: Mixed or Other Caucasian 98%. PPT1-related Neuronal Ceroid Lipofuscinosis - Gene: PPT1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_000310:1-9. Detection Rate: Mixed or Other Caucasian >99%.



MALE DONOR 12680

DOB: Ethnicity: Mixed or Other

Caucasian

Barcode: 11004512731825

FEMALE N/A

USH1C-related Disorders - Gene: USH1C. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_005709:1-21. **Detection Rate:** Mixed or Other Caucasian >99%.

USH2A-related Disorders - **Gene:** USH2A. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_206933:2-72. **Detection Rate:** Mixed or Other Caucasian 94%.

Usher Syndrome Type 3 - Gene: CLRN1. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_174878:1-3. **Detection Rate:** Mixed or Other Caucasian >99%.

Very-long-chain Acyl-CoA Dehydrogenase Deficiency - Gene: ACADVL. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000018:1-20. **Detection Rate:** Mixed or Other Caucasian >99%.

Wilson Disease - Gene: ATP7B. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM_000053:1-21. **Detection Rate:** Mixed or Other Caucasian >99%.

X-linked Adrenoleukodystrophy - Gene: ABCD1. X-linked Recessive. Sequencing with copy number analysis. Exons: NM_000033:1-6. Detection Rate: Mixed or Other Caucasian 77%.

X-linked Alport Syndrome - Gene: COL4A5. X-linked Recessive. Sequencing with copy number analysis. Exons: NM_000495:1-51. Detection Rate: Mixed or Other Caucasian 95%.

X-linked Congenital Adrenal Hypoplasia - Gene: NR0B1. X-linked Recessive. Sequencing with copy number analysis. Exons: NM_000475:1-2. Detection Rate: Mixed or Other Caucasian 99%.

X-linked Juvenile Retinoschisis - Gene: RS1. X-linked Recessive. Sequencing with copy number analysis. Exons: NM_000330:1-6. Detection Rate: Mixed or Other Caucasian 98%.

X-linked Myotubular Myopathy - Gene: MTM1. X-linked Recessive. Sequencing with copy number analysis. Exons: NM_000252:2-15. Detection Rate: Mixed or Other Caucasian 98%.

X-linked Severe Combined Immunodeficiency - Gene: IL2RG. X-linked Recessive. Sequencing with copy number analysis. Exons: NM_000206:1-8. Detection Rate: Mixed or Other Caucasian >99%.

Xeroderma Pigmentosum Group A - **Gene**: XPA. Autosomal Recessive. Sequencing with copy number analysis. **Exons**: NM_000380:1-6. **Detection Rate**: Mixed or Other Caucasian >99%.

Xeroderma Pigmentosum Group C - Gene: XPC. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM_004628:1-16. Detection Rate: Mixed or Other Caucasian 97%.



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DOB: Ethnicity: Mixed or Other

Caucasian

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FEMALE N/A

Risk Calculations

Below are the risk calculations for all conditions tested. Negative results do not rule out the possibility of being a carrier. Residual risk is an estimate of each patient's post-test likelihood of being a carrier, while the reproductive risk represents an estimated likelihood that the patients' future children could inherit each disease. These risks are inherent to all carrier-screening tests, may vary by ethnicity, are predicated on a negative family history, and are present even given a negative test result. Inaccurate reporting of ethnicity may cause errors in risk calculation. In addition, average carrier rates are estimated using incidence or prevalence data from published scientific literature and/or reputable databases, where available, and are incorporated into residual risk calculations for each population/ethnicity. When population-specific data is not available for a condition, average worldwide incidence or prevalence is used. Further, incidence and prevalence data are only collected for the specified phenotypes (which include primarily the classic or severe forms of disease) and may not include alternate or milder disease manifestations associated with the gene. Actual incidence rates, prevalence rates, and carrier rates, and therefore actual residual risks, may be higher or lower than the estimates provided. Carrier rates, incidence/prevalence, and/or residual risks are not provided for some genes with biological or heritable properties that would make these estimates inaccurate. A '†' symbol indicates a positive result. See the full clinical report for interpretation and details. The reproductive risk presented is based on a hypothetical pairing with a partner of the same ethnic group.

Disease	DONOR 12680 Residual Risk	Reproductive Risk
11-beta-hydroxylase-deficient Congenital Adrenal Hyperplasia	1 in 3,800	< 1 in 1,000,000
6-pyruvoyl-tetrahydropterin Synthase Deficiency	< 1 in 50,000	< 1 in 1,000,000
ABCC8-related Familial Hyperinsulinism	1 in 17,000	< 1 in 1,000,000
Adenosine Deaminase Deficiency	1 in 22,000	< 1 in 1,000,000
Alpha Thalassemia, HBA1/HBA2-related	Alpha globin status: aa/aa.	Not calculated
Alpha-mannosidosis	1 in 35,000	< 1 in 1,000,000
Alpha-sarcoglycanopathy	< 1 in 50,000	< 1 in 1,000,000
Alstrom Syndrome	< 1 in 50,000	< 1 in 1,000,000
AMT-related Glycine Encephalopathy	1 in 22,000	< 1 in 1,000,000
Andermann Syndrome	< 1 in 50,000	< 1 in 1,000,000
Argininemia	< 1 in 17,000	< 1 in 1,000,000
Argininosuccinic Aciduria	1 in 13,000	< 1 in 1,000,000
Aspartylglucosaminuria	< 1 in 50,000	< 1 in 1,000,000
Ataxia with Vitamin E Deficiency	< 1 in 50,000	< 1 in 1,000,000
Ataxia-telangiectasia	1 in 11,000	< 1 in 1,000,000
ATP7A-related Disorders	< 1 in 1,000,000	1 in 600,000
Autoimmune Polyglandular Syndrome Type 1	1 in 15,000	< 1 in 1,000,000
Autosomal Recessive Osteopetrosis Type 1	1 in 35,000	< 1 in 1,000,000
Autosomal Recessive Polycystic Kidney Disease, PKHD1-related	1 in 8,100	< 1 in 1,000,000
Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay	< 1 in 44,000	< 1 in 1,000,000
Bardet-Biedl Syndrome, BBS1-related	1 in 32,000	< 1 in 1,000,000
Bardet-Biedl Syndrome, BBS10-related	1 in 42,000	< 1 in 1,000,000
Bardet-Biedl Syndrome, BBS12-related	< 1 in 50,000	< 1 in 1,000,000
Bardet-Biedl Syndrome, BBS2-related	< 1 in 50,000	< 1 in 1,000,000
BCS1L-related Disorders	< 1 in 50,000	< 1 in 1,000,000
Beta-sarcoglycanopathy	1 in 39,000	< 1 in 1,000,000
Biotinidase Deficiency	1 in 13,000	1 in 650,000
Bloom Syndrome	< 1 in 50,000	< 1 in 1,000,000
Calpainopathy	1 in 13,000	< 1 in 1,000,000
Canavan Disease	1 in 9,700	< 1 in 1,000,000
Carbamoylphosphate Synthetase I Deficiency	< 1 in 57,000	< 1 in 1,000,000
Carnitine Palmitoyltransferase IA Deficiency	< 1 in 50,000	< 1 in 1,000,000
Carnitine Palmitoyltransferase II Deficiency	1 in 25,000	< 1 in 1,000,000
Cartilage-hair Hypoplasia	< 1 in 50,000	< 1 in 1,000,000
Cerebrotendinous Xanthomatosis	1 in 11,000	< 1 in 1,000,000
Citrullinemia Type 1	1 in 14,000	< 1 in 1,000,000
CLN3-related Neuronal Ceroid Lipofuscinosis	1 in 8,600	< 1 in 1,000,000
CLN5-related Neuronal Ceroid Lipofuscinosis CLN5-related Neuronal Ceroid Lipofuscinosis	< 1 in 50,000	< 1 in 1,000,000
CLN5-related Neuronal Ceroid Lipotuscinosis CLN6-related Neuronal Ceroid Lipofuscinosis	1 in 43,000	< 1 in 1,000,000 < 1 in 1,000,000
CLN8-related Neuronal Ceroid Lipofuscinosis CLN8-related Neuronal Ceroid Lipofuscinosis	< 1 in 50,000	< 1 in 1,000,000
Cohen Syndrome	< 1 in 50,000 < 1 in 15,000	
	•	< 1 in 1,000,000
COLAAA related Alport Syndrome	1 in 6,200	< 1 in 1,000,000
COL4A4-related Alport Syndrome	1 in 12,000	< 1 in 1,000,000
Combined Pituitary Hormone Deficiency, PROP1-related	1 in 6,100	< 1 in 1,000,000
Congenital Adrenal Hyperplasia, CYP21A2-related	1 in 1,300	1 in 280,000



MALE DONOR 12680

DOB:

Ethnicity: Mixed or Other

Caucasian

Barcode: 11004512731825

FEMALE N/A

Companies Disorder of Glycopystation Type Is Companies Com	Disease	DONOR 12680	Pannaduativa Biole
Congenital Disorder of Glycopistation Type Is		Residual Risk	Reproductive Risk
Congential Disorder of Glycospikation, MPI-related			
Caster Optic Atrosphy Syndrome			
Cystle Fibrosis			
Cysthosis 1 n 2,2000 4 l n 1,000,000 Delts-accoplycanopathy 1 n 4,000 4 l n 1,000,000 Delts-accoplycanopathy 4 l n 4,000 4 l n 1,000,000 Dyferfinopathy 1 n 1,000,000 4 l n 1,000,000 Dyferfonpathy (Including Duchenne/Backer Muscular Dystrophy) 1 n 1,000 4 l n 1,000,000 ERCCE-related Disorders 1 n 2,000 4 l n 1,000,000 ERCCE-related Disorders 1 n 2,000 4 l n 1,000,000 EVC-related Ellis-van Creveld Syndrome 1 n 7,500 4 l n 1,000,000 EVC-related Syndrome 1 n 7,500 1 l n 1,000,000 Fabry Disease 4 l n 1,000,000 1 l n 1,000,000 Fabry Disease 4 l n 1,000,000 1 l n 1,000,000 Familial Poyautonomia 4 l n 2,000 4 l n 1,000,000 Familial Poyautonomia 4 l n 2,000 4 l n 1,000,000 Familial Poyautonomia 4 l n 2,000 4 l n 1,000,000 Familial Poyautonomia 4 l n 2,000 4 l n 1,000,000 Familial Poyautonomia 4 l n 2,000 4 l n 1,000,000 Familial Poyautonomia 4 l n 2,000 4 l n 1,000			
Delifunctional Protein Deficiency	•	•	
Delta-sarcalykamopathy	•		
Dihydrolipoamide Dehydrogenase Deficiency	· · · · · · · · · · · · · · · · · · ·		
Dysferchinopathy			
Dystrophinopathy (Including Duchenne/Becker Muscular Dystrophy)	·		
ERCC6-elated Disorders	• • •	•	
ERCCF-elated Disorders			
EVC-related Ellis-van Creveld Syndrome			
EVC2-related Ellis-van Creveld Syndrome			
Fabry Disease	· · · · · · · · · · · · · · · · · · ·		
Familial Moditarranean Fewer	· · · · · · · · · · · · · · · · · · ·		
Familial Mediterranean Fever	· · · · · · · · · · · · · · · · · · ·		
Fanconi Anemia Camplementation Group A	•		
Fancon Anemia, FANCC-related			
FKRP-related Disorders	·		
FKTN-related Disorders			
Free Salici Acid Storage Disorders 1 in 3,0000 Calactosemia 1 in 10,00000 Calactosemia 1 in 10,00000 Calactosemia 1 in 10,00000 Calactosemia 1 in 10,00000 Camma-sarcoglycanopathy 1 in 3,0000 Calactosemia 1 in 10,00000 Calactosemia			
Salactokinase Deficiency			
Salactosemia	·		
Gamma-sarcoglycanopathy 1 in 3,000 < 1 in 1,000,000 Gaucher Disease 1 in 260 1 in 110,000 GB2-related Disorders 1 in 2,500 < 1 in 2,600 GLB1-related Glycine Encephalopathy 1 in 1,2800 < 1 in 1,000,000 GLDC-related Glycine Encephalopathy 1 in 1,2800 < 1 in 1,000,000 Glutaric Acidemia, GCDH-related 1 in 1,000 < 1 in 1,000,000 Glycogen Storage Disease Type IB 1 in 1,000 < 1 in 1,000,000 Glycogen Storage Disease Type IB 1 in 16,000 < 1 in 1,000,000 GNE Myopathy 1 in 23,000 < 1 in 1,000,000 GNE Myopathy 1 in 23,000 < 1 in 1,000,000 GNE Myopathy 1 in 20,000 < 1 in 1,000,000 GNE Myopathy 1 in 20,000 < 1 in 1,000,000 HB Beta Chain-related Disorders 1 in 32,000 < 1 in 1,000,000 HB Beta Chain-related Hemoglobinopathy (Including Beta Thalassemia and Sickle Cell Disease) 1 in 3,100 1 in 3,000 Hereditary Fructose Intolerance 1 in 7,900 < 1 in 1,000,000 Hereditary Fructose Intolerance 1 in 7,000 < 1 in 1,000,000	·		
Gauche Disease 1 in 260			
GLB2-related DFNB1 Nonsyndromic Hearing Loss and Deafness			
GLB1-elated Disorders 1 in 1,000,000 GLDC-related Glycine Encephalopathy 1 in 2,800 <1 in 1,000,000 Glutaric Acidemia, GCDH-related 1 in 16,000 <1 in 1,000,000 Glycogen Storage Disease Type Ia 1 in 18,000 <1 in 1,000,000 Glycogen Storage Disease Type Ib 1 in 16,000 <1 in 1,000,000 GNE Myopathy 1 in 12,000 <1 in 1,000,000 GNE Myopathy 1 in 32,000 <1 in 1,000,000 GNE TAB-related Disorders 1 in 20,000 <1 in 1,000,000 HADHA-related Disorders 1 in 20,000 <1 in 1,000,000 HB Beta Chain-related Hemoglobinopathy (Including Beta Thalassemia and Sickle Cell) 1 in 3,100 1 in 1,000,000 Herditary Fructose Intolerance 1 in 7,900 <1 in 1,000,000 Herditary Fructose Intolerance 1 in 50,000 <1 in 1,000,000 Herditary Fructose Intolerance 1 in 3,000 <1 in 1,000,000 Herditary Fructose Intolerance 1 in 50,000 <1 in 1,000,000 Herditary Fructose Intolerance 1 in 1,000,000 <1 in 1,000,000 Herditary Fructose Intolerance 1 in 5,000 <1 in 1,000,000 <tr< th=""><th></th><th></th><th></th></tr<>			
GLDC-related Glycine Encephalopathy	· · · · · · · · · · · · · · · · · · ·		
Glutaric Acidemia, GCDH-related			
Glycogen Storage Disease Type Ia	· · · · · · · · · · · · · · · · · · ·		
Glycogen Storage Disease Type Ib			
Signature Sign			
Section			
GNPTAB-related Disorders 1 in 32,000 < 1 in 1,000,000 HADHA-related Disorders 1 in 20,000 1 in 1,000,000 HB Beta Chain-related Hemoglobinopathy (Including Beta Thalassemia and Sickle Cell Disease) 1 in 370,000 1 in 390,000 Hereditary Fructose Intolerance 1 in 7,900 < 1 in 1,000,000 Herlitz Junctional Epidermolysis Bullosa, LAMB3-related 1 in 50,000 < 1 in 1,000,000 Hexosaminidase A Deficiency (Including Tay-Sachs Disease) 1 in 30,000 < 1 in 1,000,000 HMG-CoA Lyase Deficiency 1 in 30,000 < 1 in 1,000,000 Holocarboxylase Synthetase Deficiency 1 in 15,000 < 1 in 1,000,000 Homocystinuria, CBS-related 1 in 9,400 < 1 in 1,000,000 Hypophosphatasia 1 in 50,000 < 1 in 1,000,000 Hypophosphatasia 1 in 32,000 < 1 in 1,000,000 Isovaleric Acidemia 1 in 32,000 < 1 in 1,000,000 Jounctional Epidermolysis Bullosa, LAMA3-related 1 in 50,000 < 1 in 1,000,000 Morth Syndrome 1 in 50,000 < 1 in 1,000,000 KCDJ11-related Familial Hyperinsulinism 1 in 50,000 < 1 in 1,000,000 Krabbe Di			
Hb Beta Chain-related Hemoglobinopathy (Including Beta Thalassemia and Sickle Cell Disease) 1 in 3,100 1 in 300,000 Hereditary Fructose Intolerance 1 in 7,900 < 1 in 1,000,000		1 in 32,000	< 1 in 1,000,000
Disease) 1 in 3,000 Hereditary Fructose Intolerance 1 in 7,900 <1 in 1,000,000 Herlitz Junctional Epidermolysis Bullosa, LAMB3-related <1 in 50,000 <1 in 1,000,000 Hexosaminidase A Deficiency (Including Tay-Sachs Disease) 1 in 30,000 <1 in 1,000,000 HMG-CoA Lyase Deficiency <1 in 33,000 <1 in 1,000,000 Holocarboxylase Synthetase Deficiency 1 in 15,000 <1 in 1,000,000 Homocystinuria, CBS-related 1 in 9,400 <1 in 1,000,000 Hydrolethalus Syndrome <1 in 50,000 <1 in 1,000,000 Junctional Epidermolysis Bullosa, LAMA3-related <1 in 50,000 <1 in 1,000,000 Junctional Epidermolysis Bullosa, LAMA2-related <1 in 50,000 <1 in 1,000,000 KCNJ11-related Familial Hyperinsulinism <1 in 15,000 <1 in 1,000	HADHA-related Disorders	1 in 20,000	< 1 in 1,000,000
Disease	Hb Beta Chain-related Hemoglobinopathy (Including Beta Thalassemia and Si	ckle Cell	
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Hexosaminidase A Deficiency (Including Tay-Sachs Disease) 1 in 30,000 <1 in 1,000,000 HMG-CoA Lyase Deficiency 1 in 33,000 <1 in 1,000,000 Holocarboxylase Synthetase Deficiency 1 in 15,000 <1 in 1,000,000 Homocystinuria, CBS-related 1 in 9,400 <1 in 1,000,000 Hydrolethalus Syndrome <1 in 50,000 <1 in 1,000,000 Hypophosphatasia 1 in 27,000 <1 in 1,000,000 Isovaleric Acidemia 1 in 32,000 <1 in 1,000,000 Joubert Syndrome 2 <1 in 50,000 <1 in 1,000,000 Junctional Epidermolysis Bullosa, LAMA3-related <1 in 50,000 <1 in 1,000,000 KCNJ11-related Familial Hyperinsulinism <1 in 50,000 <1 in 1,000,000 KCNJ11-related Familial Hyperinsulinism <1 in 50,000 <1 in 1,000,000 Krabbe Disease 1 in 14,000 <1 in 1,000,000 Leigh Syndrome, French-Canadian Type <1 in 50,000 <1 in 1,000,000 Leigh Syndrome, French-Canadian Type <1 in 50,000 <1 in 1,000,000 Lysosomal Acid Lipase Deficiency 1 in 18,000 <1 in 1,000,000 Lysosomal Acid Lipase Deficiency 1 in 1,000,000	Hereditary Fructose Intolerance	1 in 7,900	< 1 in 1,000,000
HMG-CoA Lyase Deficiency < 1 in 33,000 < 1 in 1,000,000 Holocarboxylase Synthetase Deficiency 1 in 15,000 < 1 in 1,000,000 Homocystinuria, CBS-related 1 in 9,400 < 1 in 1,000,000 Hydrolethalus Syndrome < 1 in 50,000 < 1 in 1,000,000 Hypophosphatasia 1 in 27,000 < 1 in 1,000,000 Isovaleric Acidemia 1 in 32,000 < 1 in 1,000,000 Joubert Syndrome 2 < 1 in 50,000 < 1 in 1,000,000 Junctional Epidermolysis Bullosa, LAMA3-related < 1 in 50,000 < 1 in 1,000,000 Junctional Epidermolysis Bullosa, LAMC2-related < 1 in 50,000 < 1 in 1,000,000 KCNJ11-related Familial Hyperinsulinism < 1 in 50,000 < 1 in 1,000,000 KRONJ11-related Familial Hyperinsulinism < 1 in 14,000 < 1 in 1,000,000 LAMA2-related Muscular Dystrophy 1 in 34,000 < 1 in 1,000,000 Leigh Syndrome, French-Canadian Type < 1 in 50,000 < 1 in 1,000,000 Lipoid Congenital Adrenal Hyperplasia < 1 in 50,000 < 1 in 1,000,000 Lysosomal Acid Lipase Deficiency 1 in 1,000,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type	Herlitz Junctional Epidermolysis Bullosa, LAMB3-related	< 1 in 50,000	< 1 in 1,000,000
Holocarboxylase Synthetase Deficiency 1 in 15,000 <1 in 1,000,000 Homocystinuria, CBS-related 1 in 9,400 <1 in 1,000,000 Hydrolethalus Syndrome <1 in 50,000 <1 in 1,000,000 Hypophosphatasia 1 in 27,000 <1 in 1,000,000 Isovaleric Acidemia 1 in 32,000 <1 in 1,000,000 Joubert Syndrome 2 <1 in 50,000 <1 in 1,000,000 Junctional Epidermolysis Bullosa, LAMA3-related <1 in 50,000 <1 in 1,000,000 KCNJ11-related Familial Hyperinsulinism <1 in 50,000 <1 in 1,000,000 KCRbbe Disease 1 in 14,000 <1 in 1,000,000 LAMA2-related Muscular Dystrophy 1 in 34,000 <1 in 1,000,000 Leigh Syndrome, French-Canadian Type <1 in 50,000 <1 in 1,000,000 Lipoid Congenital Adrenal Hyperplasia <1 in 50,000 <1 in 1,000,000 Lysosomal Acid Lipase Deficiency 1 in 18,000 <1 in 1,000,000 Maple Syrup Urine Disease Type Ia 1 in 42,000 <1 in 1,000,000 Maple Syrup Urine Disease Type Ib 1 in 13,000 <1 in 1,000,000 Medium Chain Acyl-CoA Dehydrogenase Deficiency 1 in 4,400 <td< th=""><th>Hexosaminidase A Deficiency (Including Tay-Sachs Disease)</th><th>1 in 30,000</th><th>< 1 in 1,000,000</th></td<>	Hexosaminidase A Deficiency (Including Tay-Sachs Disease)	1 in 30,000	< 1 in 1,000,000
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Hypophosphatasia 1 in 27,000 < 1 in 1,000,000 Isovaleric Acidemia 1 in 32,000 < 1 in 1,000,000 Joubert Syndrome 2 < 1 in 50,000 < 1 in 1,000,000 Junctional Epidermolysis Bullosa, LAMA3-related < 1 in 50,000 < 1 in 1,000,000 Junctional Epidermolysis Bullosa, LAMC2-related < 1 in 50,000 < 1 in 1,000,000 KCNJ11-related Familial Hyperinsulinism < 1 in 50,000 < 1 in 1,000,000 Krabbe Disease 1 in 14,000 < 1 in 1,000,000 LAMA2-related Muscular Dystrophy 1 in 34,000 < 1 in 1,000,000 Leigh Syndrome, French-Canadian Type < 1 in 50,000 < 1 in 1,000,000 Lipoid Congenital Adrenal Hyperplasia < 1 in 50,000 < 1 in 1,000,000 Lysosomal Acid Lipase Deficiency 1 in 18,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type Ia 1 in 42,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type Ib 1 in 39,000 < 1 in 1,000,000 Medium Chain Acyl-CoA Dehydrogenase Deficiency 1 in 4,400 1 in 790,000	Homocystinuria, CBS-related	1 in 9,400	< 1 in 1,000,000
Isovaleric Acidemia 1 in 32,000 < 1 in 1,000,000 Joubert Syndrome 2 < 1 in 50,000 < 1 in 1,000,000 Junctional Epidermolysis Bullosa, LAMA3-related < 1 in 50,000 < 1 in 1,000,000 Junctional Epidermolysis Bullosa, LAMC2-related < 1 in 50,000 < 1 in 1,000,000 KCNJ11-related Familial Hyperinsulinism < 1 in 50,000 < 1 in 1,000,000 Krabbe Disease 1 in 14,000 < 1 in 1,000,000 LAMA2-related Muscular Dystrophy 1 in 34,000 < 1 in 1,000,000 Leigh Syndrome, French-Canadian Type < 1 in 50,000 < 1 in 1,000,000 Lipoid Congenital Adrenal Hyperplasia < 1 in 50,000 < 1 in 1,000,000 Lysosomal Acid Lipase Deficiency 1 in 18,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type Ia 1 in 42,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type Ib 1 in 39,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type II 1 in 13,000 < 1 in 1,000,000 Medium Chain Acyl-CoA Dehydrogenase Deficiency 1 in 4,400 1 in 790,000	Hydrolethalus Syndrome	< 1 in 50,000	< 1 in 1,000,000
Joubert Syndrome 2 <1 in 50,000 <1 in 1,000,000 Junctional Epidermolysis Bullosa, LAMA3-related <1 in 50,000 <1 in 1,000,000 Junctional Epidermolysis Bullosa, LAMC2-related <1 in 50,000 <1 in 1,000,000 KCNJ11-related Familial Hyperinsulinism <1 in 50,000 <1 in 1,000,000 Krabbe Disease 1 in 14,000 <1 in 1,000,000 LAMA2-related Muscular Dystrophy 1 in 34,000 <1 in 1,000,000 Leigh Syndrome, French-Canadian Type <1 in 50,000 <1 in 1,000,000 Lipoid Congenital Adrenal Hyperplasia <1 in 50,000 <1 in 1,000,000 Lysosomal Acid Lipase Deficiency 1 in 18,000 <1 in 1,000,000 Maple Syrup Urine Disease Type Ia 1 in 42,000 <1 in 1,000,000 Maple Syrup Urine Disease Type Ib 1 in 39,000 <1 in 1,000,000 Maple Syrup Urine Disease Type II 1 in 13,000 <1 in 1,000,000 Medium Chain Acyl-CoA Dehydrogenase Deficiency 1 in 4,400 1 in 790,000	Hypophosphatasia	1 in 27,000	< 1 in 1,000,000
Junctional Epidermolysis Bullosa, LAMA3-related < 1 in 50,000 < 1 in 1,000,000 Junctional Epidermolysis Bullosa, LAMC2-related < 1 in 50,000 < 1 in 1,000,000 KCNJ11-related Familial Hyperinsulinism < 1 in 50,000 < 1 in 1,000,000 Krabbe Disease 1 in 14,000 < 1 in 1,000,000 LAMA2-related Muscular Dystrophy 1 in 34,000 < 1 in 1,000,000 Leigh Syndrome, French-Canadian Type < 1 in 50,000 < 1 in 1,000,000 Lipoid Congenital Adrenal Hyperplasia < 1 in 50,000 < 1 in 1,000,000 Lysosomal Acid Lipase Deficiency 1 in 18,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type Ia 1 in 42,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type Ib 1 in 39,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type II 1 in 13,000 < 1 in 1,000,000 Medium Chain Acyl-CoA Dehydrogenase Deficiency 1 in 4,400 1 in 790,000	Isovaleric Acidemia	1 in 32,000	< 1 in 1,000,000
Junctional Epidermolysis Bullosa, LAMC2-related < 1 in 50,000 < 1 in 1,000,000 KCNJ11-related Familial Hyperinsulinism < 1 in 50,000 < 1 in 1,000,000 Krabbe Disease 1 in 14,000 < 1 in 1,000,000 LAMA2-related Muscular Dystrophy 1 in 34,000 < 1 in 1,000,000 Leigh Syndrome, French-Canadian Type < 1 in 50,000 < 1 in 1,000,000 Lipoid Congenital Adrenal Hyperplasia < 1 in 50,000 < 1 in 1,000,000 Lysosomal Acid Lipase Deficiency 1 in 18,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type Ia 1 in 42,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type Ib 1 in 39,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type II 1 in 13,000 < 1 in 1,000,000 Medium Chain Acyl-CoA Dehydrogenase Deficiency 1 in 4,400 1 in 790,000	Joubert Syndrome 2	< 1 in 50,000	< 1 in 1,000,000
KCNJ11-related Familial Hyperinsulinism <1 in 50,000 <1 in 1,000,000 Krabbe Disease 1 in 14,000 <1 in 1,000,000 LAMA2-related Muscular Dystrophy 1 in 34,000 <1 in 1,000,000 Leigh Syndrome, French-Canadian Type <1 in 50,000 <1 in 1,000,000 Lipoid Congenital Adrenal Hyperplasia <1 in 50,000 <1 in 1,000,000 Lysosomal Acid Lipase Deficiency 1 in 18,000 <1 in 1,000,000 Maple Syrup Urine Disease Type Ia 1 in 42,000 <1 in 1,000,000 Maple Syrup Urine Disease Type Ib 1 in 39,000 <1 in 1,000,000 Maple Syrup Urine Disease Type II 1 in 13,000 <1 in 1,000,000 Medium Chain Acyl-CoA Dehydrogenase Deficiency 1 in 4,400 1 in 790,000	Junctional Epidermolysis Bullosa, LAMA3-related	< 1 in 50,000	< 1 in 1,000,000
Krabbe Disease 1 in 14,000 < 1 in 1,000,000 LAMA2-related Muscular Dystrophy 1 in 34,000 < 1 in 1,000,000 Leigh Syndrome, French-Canadian Type < 1 in 50,000 < 1 in 1,000,000 Lipoid Congenital Adrenal Hyperplasia < 1 in 50,000 < 1 in 1,000,000 Lysosomal Acid Lipase Deficiency 1 in 18,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type Ia 1 in 42,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type Ib 1 in 39,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type II 1 in 13,000 < 1 in 1,000,000 Medium Chain Acyl-CoA Dehydrogenase Deficiency 1 in 4,400 1 in 790,000	Junctional Epidermolysis Bullosa, LAMC2-related	< 1 in 50,000	< 1 in 1,000,000
LAMA2-related Muscular Dystrophy 1 in 34,000 < 1 in 1,000,000 Leigh Syndrome, French-Canadian Type < 1 in 50,000 < 1 in 1,000,000 Lipoid Congenital Adrenal Hyperplasia < 1 in 50,000 < 1 in 1,000,000 Lysosomal Acid Lipase Deficiency 1 in 18,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type Ia 1 in 42,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type Ib 1 in 39,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type II 1 in 13,000 < 1 in 1,000,000 Medium Chain Acyl-CoA Dehydrogenase Deficiency 1 in 4,400 1 in 790,000	KCNJ11-related Familial Hyperinsulinism	< 1 in 50,000	< 1 in 1,000,000
Leigh Syndrome, French-Canadian Type < 1 in 50,000 < 1 in 1,000,000 Lipoid Congenital Adrenal Hyperplasia < 1 in 50,000 < 1 in 1,000,000 Lysosomal Acid Lipase Deficiency 1 in 18,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type Ia 1 in 42,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type Ib 1 in 39,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type II 1 in 13,000 < 1 in 1,000,000 Medium Chain Acyl-CoA Dehydrogenase Deficiency 1 in 4,400 1 in 790,000	Krabbe Disease	1 in 14,000	
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Lysosomal Acid Lipase Deficiency 1 in 18,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type Ia 1 in 42,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type Ib 1 in 39,000 < 1 in 1,000,000 Maple Syrup Urine Disease Type II 1 in 13,000 < 1 in 1,000,000 Medium Chain Acyl-CoA Dehydrogenase Deficiency 1 in 4,400 1 in 790,000	· · · · · · · · · · · · · · · · · · ·		
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Maple Syrup Urine Disease Type II 1 in 13,000 < 1 in 1,000,000 Medium Chain Acyl-CoA Dehydrogenase Deficiency 1 in 4,400 1 in 790,000			
Medium Chain Acyl-CoA Dehydrogenase Deficiency1 in 4,4001 in 790,000			
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Megalencephalic Leukoencephalopathy with Subcortical Cysts < 1 in 50,000 < 1 in 1,000,000	·		
	Megalencephalic Leukoencephalopathy with Subcortical Cysts	< 1 in 50,000	< 1 in 1,000,000



MALE DONOR 12680

DOB: Ethnicity: Mixed or Other

Caucasian Barcode: 11004512731825 FEMALE N/A

Disease	DONOR 12680 Residual Risk	Papradustiva Pisk
		Reproductive Risk
Metachromatic Leukodystrophy	1 in 16,000	< 1 in 1,000,000 < 1 in 1,000,000
Methylmalonic Acidemia, cblA Type	< 1 in 50,000	
Methylmalonic Acidemia, cblB Type	1 in 48,000 1 in 16,000	< 1 in 1,000,000
Methylmalonic Aciduria and Homocystinuria, cblC Type	·	< 1 in 1,000,000
MKS1-related Disorders Mucolipidosis III Gamma	< 1 in 50,000 < 1 in 50,000	< 1 in 1,000,000
Mucolipidosis IV	< 1 in 50,000	< 1 in 1,000,000 < 1 in 1,000,000
Mucopolysaccharidosis Type I	1 in 16,000	< 1 in 1,000,000
Mucopolysaccharidosis Type II	1 in 600,000	1 in 150,000
Mucopolysaccharidosis Type IIIA	1 in 12,000	< 1 in 1,000,000
Mucopolysaccharidosis Type IIIB	1 in 25,000	< 1 in 1,000,000
Mucopolysaccharidosis Type IIIC	1 in 37,000	< 1 in 1,000,000
MUT-related Methylmalonic Acidemia	1 in 26,000	< 1 in 1,000,000
MYO7A-related Disorders	1 in 15,000	< 1 in 1,000,000
NEB-related Nemaline Myopathy	1 in 1,200	1 in 400,000
Nephrotic Syndrome, NPHS1-related	< 1 in 50,000	< 1 in 1,000,000
Nephrotic Syndrome, NPHS2-related	1 in 35,000	< 1 in 1,000,000
Niemann-Pick Disease Type C1	1 in 19,000	< 1 in 1,000,000
Niemann-Pick Disease Type C2	< 1 in 50,000	< 1 in 1,000,000
Niemann-Pick Disease, SMPD1-related	1 in 25,000	< 1 in 1,000,000
Nijmegen Breakage Syndrome	1 in 15,000	< 1 in 1,000,000
Ornithine Transcarbamylase Deficiency	< 1 in 1,000,000	1 in 140,000
PCCA-related Propionic Acidemia	1 in 4,200	< 1 in 1,000,000
PCCB-related Propionic Acidemia	1 in 22,000	< 1 in 1,000,000
PCDH15-related Disorders	1 in 3,300	< 1 in 1,000,000
Pendred Syndrome	1 in 8,200	< 1 in 1,000,000
Peroxisome Biogenesis Disorder Type 1	1 in 16,000	< 1 in 1,000,000
Peroxisome Biogenesis Disorder Type 3	1 in 44,000	< 1 in 1,000,000
Peroxisome Biogenesis Disorder Type 4	1 in 9,300	< 1 in 1,000,000
Peroxisome Biogenesis Disorder Type 5	< 1 in 71,000	< 1 in 1,000,000
Peroxisome Biogenesis Disorder Type 6	< 1 in 50,000	< 1 in 1,000,000
Phenylalanine Hydroxylase Deficiency	1 in 4,800	1 in 940,000
POMGNT-related Disorders	< 1 in 12,000	< 1 in 1,000,000
Pompe Disease	1 in 4,000	< 1 in 1,000,000
PPT1-related Neuronal Ceroid Lipofuscinosis	1 in 7,700	< 1 in 1,000,000
Primary Carnitine Deficiency	1 in 11,000	< 1 in 1,000,000
Primary Hyperoxaluria Type 1	1 in 17,000	< 1 in 1,000,000
Primary Hyperoxaluria Type 2	< 1 in 50,000	< 1 in 1,000,000
Primary Hyperoxaluria Type 3	1 in 13,000	< 1 in 1,000,000
Pycnodysostosis	1 in 43,000	< 1 in 1,000,000
Pyruvate Carboxylase Deficiency	1 in 25,000	< 1 in 1,000,000
Rhizomelic Chondrodysplasia Punctata Type 1	1 in 16,000	< 1 in 1,000,000
RTEL1-related Disorders	< 1 in 50,000	< 1 in 1,000,000
Sandhoff Disease	1 in 32,000	< 1 in 1,000,000
Short-chain Acyl-CoA Dehydrogenase Deficiency	1 in 11,000	< 1 in 1,000,000
Sjogren-Larsson Syndrome	< 1 in 12,000	< 1 in 1,000,000
SLC26A2-related Disorders	1 in 16,000	< 1 in 1,000,000
Smith-Lemli-Opitz Syndrome	1 in 9,400	< 1 in 1,000,000
Spastic Paraplegia Type 15	< 1 in 50,000	< 1 in 1,000,000
Spinal Muscular Atrophy	Negative for g.27134T>G SNP SMN1: 2 copies	1 in 110,000
	1 in 770	
Spondylothoracic Dysostosis	< 1 in 50,000	< 1 in 1,000,000
TGM1-related Autosomal Recessive Congenital Ichthyosis	1 in 22,000	< 1 in 1,000,000
TPP1-related Neuronal Ceroid Lipofuscinosis	1 in 30,000	< 1 in 1,000,000
Tyrosine Hydroxylase Deficiency	< 1 in 50,000	< 1 in 1,000,000
Tyrosinemia Type I	1 in 16,000	< 1 in 1,000,000
Tyrosinemia Type II	1 in 25,000	< 1 in 1,000,000
USH1C-related Disorders	1 in 35,000	< 1 in 1,000,000
USH2A-related Disorders	1 in 2,200	< 1 in 1,000,000
Usher Syndrome Type 3	1 in 41,000	< 1 in 1,000,000



MALE DONOR 12680

DOB: Ethnicity: Mixed or Other

Caucasian

Barcode: 11004512731825

FEMALE N/A

Disease	DONOR 12680 Residual Risk	Reproductive Risk < 1 in 1,000,000	
Wilson Disease	1 in 6,500		
X-linked Adrenoleukodystrophy	1 in 90,000	1 in 42,000	
X-linked Alport Syndrome	Not calculated	Not calculated < 1 in 1,000,000	
X-linked Congenital Adrenal Hypoplasia	< 1 in 1,000,000		
X-linked Juvenile Retinoschisis	< 1 in 1,000,000	1 in 40,000	
X-linked Myotubular Myopathy	Not calculated	Not calculated	
X-linked Severe Combined Immunodeficiency	< 1 in 1,000,000	1 in 200,000	
Xeroderma Pigmentosum Group A	< 1 in 50,000	< 1 in 1,000,000	
Xeroderma Pigmentosum Group C	1 in 7,300	< 1 in 1,000,000	



Client/Sending Facility: Seattle Sperm Bank

4915 25th Ave Ne Ste 204 SEATTLE, WA 98105 Ph: (206)588-1484

Fax: (206) 466-4696 WAB-55

LCLS Specimen Number: 343-129-1193-0 Account Number: 46857540

Patient Name: 12680, DONOR Ordering Physician: J OLLIFFE

Date of Birth: Specimen Type: BLOOD

Gender: M Client Reference:

Patient ID: Date Collected: 12/08/2020
Lab Number: DCGWA20-4098 L Date Received: 12/09/2020

Indications: DONOR Date Reported: 12/17/2020

Test: Chromosome, Blood, Routine

Cells Counted: 15 Cells Karyotyped: 4
Cells Analyzed: 5 Band Resolution: 550

CYTOGENETIC RESULT: 46,XY

INTERPRETATION: NORMAL MALE KARYOTYPE

Cytogenetic analysis of PHA stimulated cultures has revealed a MALE karyotype with an apparently normal GTW banding pattern in all cells observed.

This result does not exclude the possibility of subtle rearrangements below the resolution of cytogenetics or congenital anomalies due to other etiologies.



Client/Sending Facility: Seattle Sperm Bank

4915 25th Ave Ne Ste 204 SEATTLE, WA 98105 Ph: (206)588-1484

Fax: (206) 466-4696 WAB-55

LCLS Specimen Number: 343-129-1193-0

Patient Name: 12680, DONOR

Date of Birth:

Gender: M

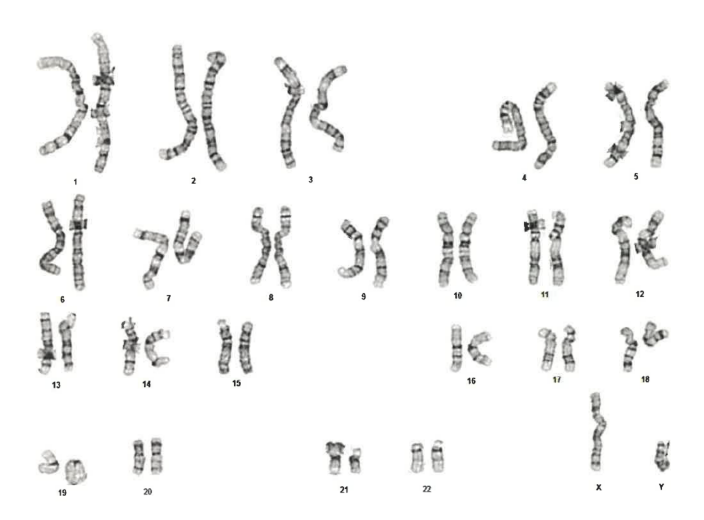
Patient ID:

Lab Number: DCGWA20-4098 L

Account Number: 46857540
Ordering Physician: J OLLIFFE
Specimen Type: BLOOD

Client Reference:

Date Collected: 12/08/2020 Date Received: 12/09/2020





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Specimen Type: BLOOD

Client Reference:

Date Collected: 12/08/2020

Date Received: 12/09/2020

Hiba Risheg, PhD., FACMG

Daniel Toweill, MD Medical Director Stuart Schwartz, PhD

National Director of Cytogenetics

Technical component performed by Laboratory Corporation of America Holdings, 550 17th Ave Ste 300, Seattle, WA, 98122-5789 (206) 861-7000

Professional Component performed by LabCorp CLIA 50D0630157, 550 17th Ave. Suite 200, Seattle WA 98122-5789. Medical Director, Daniel Toweill, MD

This document contains private and confidential health information protected by state and federal law.





Patient Information

Name: Donor 12680

Date of Birth:
Sema4 ID: 22130044CS

Client ID: SEATSB-S458158631

Indication: Carrier Screening

Specimen Information

Specimen Type: Blood
Date Collected: 06/29/2022
Date Received: 06/30/2022
Final Report: 07/10/2022

Referring Provider

Jeffrey Olliffe, M.D. Seattle Sperm Bank 4915 25th Avenue NE Suite 204W Seattle, WA, 98105 Fax: 206-466-4696

Custom Carrier Screen (1 gene)

with Personalized Residual Risk

SUMMARY OF RESULTS AND RECOMMENDATIONS

Negative

Negative for all genes tested: DUOX2

To view a full list of genes and diseases tested please see Table 1 in this report

AR=Autosomal recessive: XL=X-linked

Recommendations

• 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. Please note that residual risks for X-linked diseases (including full repeat expansions for Fragile X syndrome) may not be accurate for males and the actual residual risk is likely to be lower.

Test description

This patient was tested for the genes listed above using one or more of the following methodologies: target capture and short-read sequencing, long-range PCR followed by short-read sequencing, targeted genotyping, and/or 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 with the patient's personalized residual risk. If personalized residual risk is not provided, please see the complete residual risk table at **go.sema4.com/residualrisk**. Only known pathogenic or likely pathogenic variants are reported. This carrier screening test does not report likely benign variants and variants of uncertain significance (VUS). If reporting of likely benign variants and VUS are desired in this patient, please contact the laboratory at 800-298-6470, option 2 to request an amended report.

Anastasia Larmore, Ph.D., Associate Laboratory Director

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





Genes and diseases tested

The personalized residual risks listed below are specific to this individual. The complete residual risk table is available at go.sema4.com/residualrisk

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

	Disease	Gene	Inheritance Pattern	Status	Detailed Summary
Θ	Negative				
	Thyroid Dyshormonogenesis 6	DUOX2	AR	Reduced Risk	Personalized Residual Risk: 1 in 190

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%)

 $MLPA^{\circledR}$ 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 gg%.

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; ACADSB (NM_001609.3) chr10:124,810,695-124,810,707 (partial exon 9); ADA (NM_0000222) exon 1; ADAMTS2 (NM_014244 4) exon 1; AGPS (NM_003659.3) chrz:178,257,512-178,257,649 (partial exon 1); ALDH7A1 (NM_001182.4) chr5:125,911,150-125,911,163 (partial exon 7) and chr5:125,896,807-125,896,821 (partial exon 10); ALMS1 (NM_015120.4) chr2:73,612,990-73,613,041 (partial exon 1): APOPTI (NM_032374.4) chr14:104.040.437-104.040.455 (partial exon 3): CDAN1 (NM_138477.2) exon 2: CEP152 (NM_014985.3) chr15;49,061,146-49,061,165 (partial exon 14) and exon 22; CEP2go (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_0000924) chr2:227,942,604-227,942,619 (partial exon 25); COX10 (NM_001303.3) exon 6; CYP11B1 (NM_000497.3) exons 3-7; CYP11B2 (NM_000498.3) exons 3-7; DNAI2 (NM_023036.4) chr17;72,308,136-72,308,147 (partial exon 12); DOK7 (NM_173660.4) chr4;3,465,131-3,465,161 (partial exon 1) and exon 2; DUOX2 (NM_014080.4) exons 6-8; EIF2AK3 (NM_004836.5 exon 8: EVC (NM_1537172) exon 1: F5 (NM_000130 4) chr1:169.551.662-169.551.679 (partial exon 2): 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; GHR (NM_0001634) exon 3; GYS2 (NM_021957.3) chr12:21,699,370-21,699,409 (partial exon 12); HGSNAT (NM_152419.2) exon 1; IDS (NM_000202.6 exon 3; ITGB4 (NM_000213.4) chr17:73,749,976-73,750,060 (partial exon 33); JAK3 (NM_000215.3) chr19:17,950,462-17,950,483 (partial exon 10); LIFR (NM_002310.5 exon 19; LMBRD1 (NM_018368.3) chr6:70,459,226-70,459,257 (partial exon 5), chr6:70,447,828-70,447,836 (partial exon 7) and exon 12; LYST (NM_000081.3) chr1:235,944,158-235,944,176 (partial exon 16) and chr1:235,875,350-235,875,362 (partial exon 43); MLYCD (NM_012213.2) chr16:83,933,242-83,933,282 (partial exon 1); MTR (NM_000254.2) chr1 237,024,418-237,024,439 (partial exon 20) and chr1:237,038,019-237,038,029 (partial exon 24); NBEAL2 (NM_015175 2) chr3 47,021,385-47,021,407 (partial exon 1); NEB (NM_001271208.1 exons 82-105; NPC1 (NM_0002714) chr18:21,123,519-21,123,538 (partial exon 14); NPHP1 (NM_000272.3) chr2:110,937,251-110,937,263 (partial exon 3); OCRL (NM_000276.3) chrX:128,674,450-128,674,460 (partial exon 1); PHKB (NM_0002932) exon 1 and chr16:47,732,498-47,732,504 (partial exon 30); PIGN (NM_176787.4) chr18:59,815,547-59,815,576 (partial exon 8); PIP5K1C (NM_012398.2) exon 1 and chr19;3637602-3637616 (partial exon 17); POU1F1 (NM_000306.3) exon 5; PTPRC (NM_0028384) exons 11 and 23; PUS1 (NM_025215.5 chr12:132,414,446-132,414,532 (partial exon 2); RPGRIP1L (NM_0152722) exon 23; SGSH (NM_000199.3) chr17;78,194,022-78,194,072 (partial exon 1); SLC6A8 (NM_005629.3) exons 3 and 4; ST3GAL5 (NM_003896.3) exon 1; SURF1 (NM_003172.3) chrg:136,223,269-136,223,307 (partial exon 1); TRPM6 (NM_0176624) chrg:77,362,800-77,362,811 (partial exon 31); TSEN54 (NM_207346.2) exon 1; TYR (NM_0003724) exon 5; VWF (NM_000552.3) exons 24-26, chr12:6.125,675-6.125,684 (partial exon 30), chr12:6.121,244-6,121,265 (partial exon 33), and exon 34.

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%)

Th 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 through 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.

Tay-Sachs Disease (TSD) Enzyme Analysis (Analytical Detection Rate >98%)

Hexosaminidase activity and Hex A% activity were measured by a standard heat-inactivation, fluorometric method using artificial 4-MU-β-N-acetyl glucosaminide (4-MUG) substrate. This assay is highly sensitive and accurate in detecting Tay-Sachs carriers and individuals affected with TSD. Normal ranges of Hex A% activity are 55.0-72.0 for white blood cells and 58.0-72.0 for plasma. It is estimated that less than 0.5% of Tay-Sachs carriers have non-carrier levels of percent Hex A activity, and therefore may not be identified by this assay. In addition, this assay may detect individuals that are carriers of or are affected with Sandhoff disease. False positive results may occur if benign variants, such as pseudodeficiency alleles, interfere with the enzymatic assay. False negative results may occur if both HEXA and HEXB pathogenic or pseudodeficiency variants are present in the same individual.

Please note these tests were developed and their performance characteristics were determined by Sema4 Opco, 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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Carrier Screening

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Fragile X syndrome

Chen L et al. An information-rich CGG repeat primed PCR that detects the full range of Fragile X expanded alleles and minimizes the need for Southern blot analysis. *J Mol Diag* 2010 12:589-600.

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Luo M et al. An Ashkenazi Jewish SMN1 haplotype specific to duplication alleles improves pan-ethnic carrier screening for spinal muscular atrophy. *Genet Med.* 2014 16:149-56.

Ashkenazi Jewish Disorders:

Scott SA et al. Experience with carrier screening and prenatal diagnosis for sixteen Ashkenazi Jewish Genetic Diseases. *Hum. Mutat.* 2010 31:1-11.

Duchenne Muscular Dystrophy:

Flanigan KM et al. Mutational spectrum of DMD mutations in dystrophinopathy patients: application of modern diagnostic techniques to a large cohort. *Hum Mutat.* 2009 30:1657-66.

Variant Classification:

Richards S et al. Standards and guidelines for the interpretation of sequence variants: a joint consensus recommendation of the American College of Medical Genetics and Genomics and the Association for Molecular Pathology. *Genet Med.* 2015 May;17(5):405-24 Additional disease-specific references available upon request.

Specimen ID: 343-129-1194-0 **Control ID:** B0109683339

Acct #: 46857540

Phone: (206) 588-1484

Rte: 00

12680, DONOR

Seattle Sperm Bank 4915 25th Ave Ne Ste 204 SEATTLE WA 98105

<u>ընինուկնիինիգուսոր</u>ականիկրեկիիիի

Patient Details

DOB:

Age(y/m/d):

Gender: M

Patient ID:

Specimen Details

Date collected: 12/08/2020 1010 Local

Date received: 12/09/2020 Date entered: 12/08/2020

Date reported: 12/14/2020 1706 ET

Physician Details Ordering: J OLLIFFE

Referring:

ID:

NPI: 1306838271

General Comments & Additional Information Alternate Control Number: B0109683339

Alternate Patient ID: Not Provided

Ordered Items

LP+12AC+CBC/D/Plt+UA+Rh+ABO...

TESTS	RESULT	FLAG	UNITS RE	FERENCE INTERVAL	LAB
LP+12AC+CBC/D/Plt+UA+Rh+ABO.					
Glucose	110	High `	mg/dL	65-99	01
Uric Acid	6.9		mg/dL	3.8-8.4	01
	Therapeut **Plea	tic targ ase note	et for gout p reference in	oatients: <6.0 nterval change**	
BUN	13		mg/dL	6-20	01
Creatinine	1.12		mg/dL	0.76-1.27	01
eGFR If NonAfricn Am	91		mL/min/1.73		
eGFR If Africn Am	105		mL/min/1.73		
Calcium	9.3		mg/dL	8.7-10.2	01
Protein, Total	6.7		g/dL	6.0-8.5	01
Albumin	4.4		g/dL	4.1-5.2	01
Bilirubin, Total	0.6		mg/dL	0.0-1.2	01
Alkaline Phosphatase	66		IU/L	39-117	01
LDH	161		IU/L	121-224	01
AST (SGOT)	34		IU/L	0-40	01
ALT (SGPT)	57	High	IU/L	0-44	01
Cholesterol, Total	167		mg/dL	100-199	01
Triglycerides	58		mg/dL	0-149	01
HDL Cholesterol	42		mg/dL	>39	01
LDL Chol Calc (NIH)	114	High /	mg/dL	0-99	
LDL/HDL Ratio	2.7		ratio	0.0-3.6	
Please Note:					01
			LDL/	HDL Ratio Men Women	
			1/2 Avg.Ri		
el m	- \ .		Avg.Ri		
GAG /	TOCZ	ع ا	2X Avg.Ri		
90 09	, ,		3X Avg.Ri	lsk 8.0 6.1	
Ce					01
Hemoglobin Fractionation					01
Hgb F A	0.0		%	0.0-2.0	02



Patient: 12680, DONOR DOB:

Patient ID:

Control ID: B0109683339

Specimen ID: 343-129-1194-0 Date collected: 12/08/2020 1010 Local

TESTS	RESULT	FLAG	UNITS	REFERENCE INTERVAL	
Hgb A A	97.6		%	96.4-98.8	02
Hgb S A	0.0		%	0.0	02
Hgb C A	0.0		%	0.0	02
Hgb A2 ^A	2.4		ે	1.8-3.2	02
Hgb Variant A	0.0		ક	0.0	02
Interpretation					
Normal adult hemoglobin	_				02
ABO Grouping	0				01
Rh Factor	Positive		- 3DO / Di	one not	01
Please note: Prior reco	rds for this	s patient'	s ABO / R	n type are not	
available for additiona	I verificat.	1011.			01
					01
CBC, Platelet Ct, and Diff	4 0		x10E3/uL	3.4-10.8	01
WBC	4.0		x10E3/uL		01
RBC	5.22			13.0-17.7	01
Hemoglobin	14.8		g/dL		01
Hematocrit	44.7		%	37.5-51.0	01
MCV	86		fL	79-97	
MCH	28.4		pg	26.6-33.0	01
MCHC	33.1		g/dL	31.5-35.7	01
RDW	14.5		ે	11.6-15.4	01
Platelets	239		x10E3/uL		01
Neutrophils	61		8	Not Estab.	01
Lymphs	28		%	Not Estab.	01
Monocytes	6		%	Not Estab.	01
Eos	3		%	Not Estab.	01
Basos	1		8	Not Estab.	01
Neutrophils (Absolute)	2.5		x10E3/uL	1.4-7.0	01
Lymphs (Absolute)	1.1		x10E3/uL	0.7-3.1	01
Monocytes (Absolute)	0.3		x10E3/uL	0.1-0.9	01
Eos (Absolute)	0.1		x10E3/uL	0.0-0.4	01
Baso (Absolute)	0.0		x10E3/uL	0.0-0.2	01
Immature Granulocytes	1		%	Not Estab.	01
Immature Grans (Abs)	0.0		x10E3/uL	0.0-0.1	01
	*				01
Urinalysis Gross Exam					01
Specific Gravity	1.010			1.005-1.030	01
рн	7.5	20	17 De	5.0-7.5	01
Urine-Color	Yellow	APA	. 1 200	Yellow	01
Appearance	Clear	(000)		Clear	01
WBC Esterase	Negative			Negative	01
Protein	Negative			Negative/Trace	01
Glucose	Negative			Negative	01
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