

Foresight® Carrier Screen

RESULTS RECIPIENT **SEATTLE SPERM BANK** Attn: Jeffrey Olliffe 4915 25th Ave NE Ste 204w Seattle, WA 98105-5668 Phone: (206) 588-1484 Fax: (206) 466-4696 NPI: 1306838271 Report Date: 08/21/2019 MALE DONOR 12445 DOB Ethnicity: Northern European Sample Type: EDTA Blood Date of Collection: 08/15/2019 Date Received: 08/16/2019 Date Tested: 08/21/2019 Barcode: 11004212718545 Accession ID: CSL92M2GNCE4YK6 Indication: Egg or sperm donor

#### FEMALE N/A

## POSITIVE: CARRIER

#### 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 12445	Partner
Panel Information	Foresight Carrier Screen Universal Panel Fundamental Plus Panel Fundamental Panel <b>(175 conditions tested)</b>	N/A
POSITIVE: CARRIER		The reproductive risk presented
Galactosemia	NM_000155.3(GALT):c.563A>G (Q188R) heterozyaote	is based on a hypothetical pairing with a partner of the
Reproductive Risk: 1 in 350	· · · · · · · · · · · · · · · · · · ·	same ethnic group. Carrier
Inheritance: Autosomal Recessive		testing should be considered.
		See "Next Steps".

\*Carriers generally do not experience symptoms.

No disease-causing mutations were detected in any other gene tested. A complete list of all conditions tested can be found on page 6.

#### CLINICAL NOTES

• None

#### NEXT STEPS

- Carrier testing should be considered for the diseases specified above for the patient's partner, as both parents must be carriers before a child is at high risk of developing the disease.
- Genetic counseling is recommended and patients may wish to discuss any positive results with blood relatives, as there is an increased chance that they are also carriers.



MALE DONOR 12445 DOB: Ethnicity: Northern European Barcode: 11004212718545

FEMALE N/A

# positive: carrier Galactosemia

**Reproductive risk: 1 in 350** Risk before testing: 1 in 30,000

Gene: GALT | Inheritance Pattern: Autosomal Recessive

Patient	DONOR 12445	No partner tested
Result	Garrier	N/A
Variant(s)	NM_000155.3(GALT):c.563A>G(Q188R) heterozygote	N/A
Methodology	Sequencing with copy number analysis	N/A
Interpretation	This individual is a carrier of galactosemia. Carriers generally do not experience symptoms.	N/A
Detection rate	>99%	N/A
Exons tested	NM_000155:1-11.	N/A

### What Is Galactosemia?

Galactosemia is a treatable inherited condition that reduces the body's ability to metabolize galactose, a simple sugar found in milk. It is caused by mutations in the *GALT* gene, which result in a deficiency in an enzyme called galactose-1-phosphate uridyltransferase. The classic form of galactosemia can be fatal without prompt treatment and careful management. Because milk is a staple of an infant's diet, diagnosis and treatment within the first week of life is critical to avoiding intellectual disability and life-threatening complications.

#### CLASSIC FORM

Classic galactosemia, the most severe form of the disease, occurs when galactose-1-phosphate uridyltransferase activity is very low or absent. After only a few days of drinking milk, including breast milk, an infant with classic galactosemia will show symptoms including loss of appetite, jaundice, vomiting, lethargy, and convulsions. Without immediate and vigilant lifelong treatment, children with the condition will experience life-threatening complications such as severe infections, cirrhosis of the liver, and intellectual disability. Even with treatment, children can still develop cataracts, speech problems, stunted growth and motor function, and learning disabilities, and most females will eventually develop menstrual irregularities and go through premature menopause.

#### CLINICAL VARIANT FORM

Clinical variant galactosemia occurs when occurs when galactose-1-phosphate uridyltransferase activity is approximately 10% of the normal level. People with this form of galactosemia can have some of the symptoms of classic galactosemia, such as growth problems, severe infections, cirrhosis of the liver, cataracts, and mild intellectual disability. However, females do not develop menstrual irregularities or go through premature menopause.

#### **BIOCHEMICAL VARIANT FORM**

The biochemical variant form, also called Duarte galactosemia, is a much milder form of the disease in which a person has 14 to 25% of the normal amount of galactose-1-phosphate uridyltransferase. People with Duarte galactosemia generally do not suffer any of the symptoms of classic galactosemia.

Please note that galactosemia is not the same as lactose intolerance, a more-common and less-serious condition.



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### How Common Is Galactosemia?

Classic galactosemia affects 1 in 30,000 to 1 in 60,000 newborns, and it is more common in individuals of Irish ancestry. The prevalence of clinical variant galactosemia is estimated to be 1 in 20,000. The prevalence of Duarte galactosemia is approximately 1 in 4,000.

### How Is Galactosemia Treated?

People with classic galactosemia and clinical variant galactosemia must monitor their galactose-1-phosphate levels with regular blood tests and follow a lifelong diet free of milk, milk products, or other foods containing lactose. Infants should be fed with galactose-free formulas such as soy formula or Nutramigen, a hypoallergenic formula with no galactose, lactose, or soy. As children learn to feed themselves, parents must teach them how to read product labels so that they can avoid any food containing milk, dry milk, milk products, and other galactose-containing foods. Often they require calcium supplements to avoid calcium deficiency.

There is debate on whether people with Duarte galactosemia need to adhere to a galactose-free diet. Some medical professionals recommend modifying an affected person's diet, while others do not. The decision as to whether or not to treat a person with Duarte galactosemia may depend upon his or her level of enzyme activity.

People with galactosemia should work with a nutritionist to determine the best course of treatment.

### What Is the Prognosis for a Person with Galactosemia?

Most people who are diagnosed early with classic or clinical variant galactosemia and carefully follow a galactose-free diet can have a normal lifespan. However, they are still at risk for cataracts, speech defects, poor growth, poor intellectual function, neurologic deficits, and (in women with classic galactosemia) ovarian failure. If the treatment of classic or clinical variant galactosemia is not prompt and consistent, life-threatening complications and irreversible intellectual disability can result.

Duarte galactosemia has not been associated with any long-term health problems.



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# **Methods and Limitations**

DONOR 12445 [Foresight Carrier Screen]: Sequencing with copy number analysis, spinal muscular atrophy, and analysis of homologous regions.

## Sequencing with copy number analysis

High-throughput sequencing and read depth-based copy number analysis are used to analyze the listed exons, as well as selected intergenic and intronic regions, of the genes in the Conditions Tested section of the report. The region of interest (ROI) of the test comprises these regions, in addition to the 20 intronic bases flanking each exon. 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 high coverage and the sequences are compared to standards and references of normal variation. More than 99% of all bases in the ROI are sequenced at greater than the minimum read depth. Mutations may not be detected in areas of lower sequence coverage. Small insertions and deletions may not be as accurately determined as single nucleotide variants. Genes that have closely related pseudogenes may be addressed by a different method. *CFTR* and *DMD* testing includes analysis for both large (exon-level) deletions and duplications with an average sensitivity of 99%, while other genes are only analyzed for large deletions with a sensitivity of >75%. However, the sensitivity may be higher for selected founder deletions. The breakpoints of copy number variants and exons affected are estimated from probe positions. Only exons known to be included in the copy number variant are provided in the name. In some cases, the copy number variant may be larger or smaller than indicated. If *GJB2* is tested, two large upstream deletions which overlap *GJB6* and affect the expression of *GJB2*, del(*GJB6*-D13S1830) and del(*GJB6*-D13S1854), are also analyzed. Mosaicism or somatic variants present at low levels may not be detected. If detected, these may not be reported.

Detection rates are determined by using literature to estimate the fraction of disease alleles, weighted by frequency, that the methodology is unable to detect. Detection rates only account for analytical sensitivity and 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 mutations.

All variants that are a recognized cause of the disease will be 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". Likely pathogenic variants are evaluated and classified by assessing the nature of the variant and reviewing reports of allele frequencies in cases and controls, functional studies, variant annotation and effect prediction, and segregation studies. Exon level duplications 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 intended disease phenotype are not reported. Curation summaries of reported variants are available upon request.

# Spinal muscular atrophy

Targeted copy number analysis is used to determine the copy number of exon 7 of the *SMN1* gene relative to other genes. Other mutations may interfere with this analysis. Some individuals with two copies of *SMN1* are carriers with two *SMN1* genes on one chromosome and a *SMN1* deletion on the other chromosome. This is more likely in individuals who have 2 copies of the *SMN1* gene and are positive for the g.27134T>G SNP, 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 2 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 mutations in certain genes that have homology to other regions. The precise breakpoints of large deletions in these genes cannot be determined, but are estimated from copy number analysis. High numbers of pseudogene copies may interfere with this analysis.

If *CYP21A2* is tested, patients who have one or more additional copies of the *CYP21A2* gene and a loss of function mutation may not actually be a carrier of 21-hydroxylasedeficient congenital adrenal hyperplasia (CAH). Because the true incidence of non-classic CAH is unknown, the residual carrier and reproductive risk numbers on the report are only based on published incidences 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 of overall chances for 21-hydroxylase-deficient CAH, especially in the aforementioned populations, as they do not account for non-classic CAH. If *HBA1/HBA2* are tested, some individuals with four alpha globin genes may be carriers, with three genes on one chromosome and a deletion on the other chromosome. This and similar, but rare, carrier states, where complementary changes exist in both the gene and a pseudogene, may not be detected by the assay.



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### Limitations

In an unknown number of cases, nearby genetic variants may interfere with mutation detection. Other possible sources of diagnostic error include sample mix-up, trace contamination, bone marrow transplantation, blood transfusions and technical errors. This test is designed to detect and report germline alterations. While somatic variants present at low levels may be detected, these 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. The test does not fully address all inherited forms of intellectual disability, birth defects and genetic disease. A family history of any of these conditions may warrant additional evaluation. Furthermore, not all mutations will be identified in the genes analyzed and additional testing may be beneficial for some patients. For example, individuals of African, Southeast Asian, and Mediterranean ancestry are at increased risk for being carriers for hemoglobin opathies, which can be identified by CBC and hemoglobin electrophoresis or HPLC (*ACOG Practice Bulletin No. 78. Obstet.Gynecol. 2007;109:229-37*).

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**.

#### Resources

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

Patients can share their reports via research registries such as Genome Connect, an online research registry working to build the knowledge base about genetics and health. 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

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Jack Ji, PhD, FACMG

Report content approved by Jack Ji, PhD, FACMG on Aug 22, 2019



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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:** Northern European 94%.

6-pyruvoyl-tetrahydropterin Synthase Deficiency - Gene: PTS. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000317:1-6. Detection Rate: Northern European >99%.

ABCC8-related Familial Hyperinsulinism - Gene: ABCC8. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000352:1-39. Detection Rate: Northern European >99%.

Adenosine Deaminase Deficiency - Gene: ADA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000022:1-12. Detection Rate: Northern European >99%.

Alpha Thalassemia - Genes: HBA1, HBA2. Autosomal Recessive. Analysis of homologous regions. Variants (13): -(alpha)20.5, --BRIT, --MEDI, --MEDI, --SEA, --THAI or --FIL, -alpha3.7, -alpha4.2, HBA1+HBA2 deletion, Hb Constant Spring, anti3.7, anti4.2, del HS-40. Detection Rate: Unknown due to rarity of disease. Alpha-mannosidosis - Gene: MAN2B1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000528:1-23. Detection Rate: Northern European >99%.

Alpha-sarcoglycanopathy - Gene: SGCA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000023:1-9. Detection Rate: Northern European >99%.

Alstrom Syndrome - Gene: ALMS1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_015120:1-23. Detection Rate: Northern European >99%.

**AMT-related Glycine Encephalopathy** - Gene: AMT. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000481:1-9. Detection Rate: Northern European >99%.

Andermann Syndrome - Gene: SLC12A6. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_133647:1-25. Detection Rate: Northern European >99%.

Argininemia - Gene: ARG1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000045:1-8. Detection Rate: Northern European 97%. Argininosuccinic Aciduria - Gene: ASL. Autosomal Recessive. Sequencing with copy

number analysis. Exons: NM\_001024943:1-16. Detection Rate: Northern European >99%.

Aspartylglucosaminuria - Gene: AGA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000027:1-9. Detection Rate: Northern European >99%.

Ataxia with Vitamin E Deficiency - Gene: TTPA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000370:1-5. Detection Rate: Northern European >99%.

Ataxia-telangiectasia - Gene: ATM. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000051:2-63. Detection Rate: Northern European 98%.

ATP7A-related Disorders - Gene: ATP7A. X-linked Recessive. Sequencing with copy number analysis. Exons: NM\_000052:2-23. Detection Rate: Northern European 96%.

Autoimmune Polyglandular Syndrome Type 1 - Gene: AIRE. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000383:1-14. Detection Rate: Northern European >99%.

Autosomal Recessive Osteopetrosis Type 1 - Gene: TCIRG1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_006019:2-20. Detection Rate: Northern European >99%.

Autosomal Recessive Polycystic Kidney Disease, PKHD1-related - Gene: PKHD1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_138694:2-67. Detection Rate: Northern European >99%. Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay - Gene: SACS. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_014363:2-10. Detection Rate: Northern European 99%.

Bardet-Biedl Syndrome, BBS1-related - Gene: BBS1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_024649:1-17. Detection Rate: Northern European >99%.

**Bardet-Biedl Syndrome, BBS10-related** - Gene: BBS10. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_024685:1-2. Detection Rate: Northern European >99%.

Bardet-Biedl Syndrome, BBS12-related - Gene: BBS12. Autosomal Recessive. Sequencing with copy number analysis. Exon: NM\_152618:2. Detection Rate: Northern European >99%.

Bardet-Biedl Syndrome, BBS2-related - Gene: BBS2. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_031885:1-17. Detection Rate: Northern European >99%.

**BCS1L-related Disorders** - Gene: BCS1L. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_004328:3-9. Detection Rate: Northern European >99%.

**Beta-sarcoglycanopathy** - Gene: SGCB. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000232:1-6. Detection Rate: Northern European >99%.

Biotinidase Deficiency - Gene: BTD. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000060:1-4. Detection Rate: Northern European >99%.

Bloom Syndrome - Gene: BLM. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000057:2-22. Detection Rate: Northern European >99%.

Calpainopathy - Gene: CAPN3. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000070:1-24. Detection Rate: Northern European >99%.

Canavan Disease - Gene: ASPA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000049:1-6. Detection Rate: Northern European 98%. Carbamoylphosphate Synthetase I Deficiency - Gene: CPS1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_001875:1-38. Detection Rate: Northern European >99%.

Carnitine Palmitoyltransferase IA Deficiency - Gene: CPT1A. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_001876:2-19. Detection Rate: Northern European >99%.

Carnitine Palmitoyltransferase II Deficiency - Gene: CPT2. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000098:1-5. Detection Rate: Northern European >99%.

Cartilage-hair Hypoplasia - Gene: RMRP. Autosomal Recessive. Sequencing with copy number analysis. Exon: NR\_003051:1. Detection Rate: Northern European >99%.

**Cerebrotendinous Xanthomatosis** - Gene: CYP27A1. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_000784:1-9. **Detection Rate:** Northern European >99%.

Citrullinemia Type 1 - Gene: ASS1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000050:3-16. Detection Rate: Northern European >99%.

CLN3-related Neuronal Ceroid Lipofuscinosis - Gene: CLN3. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_001042432:2-16. Detection Rate: Northern European >99%.

CLN5-related Neuronal Ceroid Lipofuscinosis - Gene: CLN5. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_006493:1-4. Detection Rate: Northern European >99%.

CLN6-related Neuronal Ceroid Lipofuscinosis - Gene: CLN6. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_017882:1-7. Detection Rate: Northern European >99%.



**CLN8-related Neuronal Ceroid Lipofuscinosis** - Gene: CLN8. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_018941:2-3. **Detection Rate:** Northern European >99%.

Cohen Syndrome - Gene: VPS13B. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_017890:2-62. Detection Rate: Northern European 97%.

**COL4A3**-related Alport Syndrome - Gene: COL4A3. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000091:1-52. Detection Rate: Northern European 97%.

**COL4A4-related Alport Syndrome** - Gene: COL4A4. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_000092:2-48. **Detection Rate:** Northern European 98%.

Combined Pituitary Hormone Deficiency, PROP1-related - Gene: PROP1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_006261:1-3. Detection Rate: Northern European >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, V281L, [I237N;V238E;M240K], c.293-13C>G. Detection Rate: Northern European 96%.

**Congenital Disorder of Glycosylation Type Ia** - Gene: PMM2. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_000303:1-8. **Detection Rate:** Northern European >99%.

**Congenital Disorder of Glycosylation Type Ic** - Gene: ALG6. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_013339:2-15. **Detection Rate:** Northern European >99%.

Congenital Disorder of Glycosylation, MPI-related - Gene: MPI. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_002435:1-8. Detection Rate: Northern European >99%.

**Costeff Optic Atrophy Syndrome** - Gene: OPA3. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_025136:1-2. **Detection Rate:** Northern European >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: Northern European >99%. Cystinosis - Gene: CTNS. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_004937:3-12. Detection Rate: Northern European >99%.

D-bifunctional Protein Deficiency - Gene: HSD17B4. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000414:1-24. Detection Rate: Northern European 98%.

**Delta-sarcoglycanopathy** - Gene: SGCD. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_000337:2-9. **Detection Rate:** Northern European 99%.

Dihydrolipoamide Dehydrogenase Deficiency - Gene: DLD. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000108:1-14. Detection Rate: Northern European >99%.

Dysferlinopathy - Gene: DYSF. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_003494:1-55. Detection Rate: Northern European 98%. Dystrophinopathy (Including Duchenne/Becker Muscular Dystrophy) - Gene: DMD. X-linked Recessive. Sequencing with copy number analysis. Exons:

NM 004006:1-79. Detection Rate: Northern European >99%.

**ERCC6-related Disorders** - Gene: ERCC6. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_000124:2-21. **Detection Rate:** Northern European 99%.

**ERCC8-related Disorders** - Gene: ERCC8. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_000082:1-12. **Detection Rate:** Northern European 95%.

**EVC-related Ellis-van Creveld Syndrome** - Gene: EVC. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_153717:1-21. **Detection Rate:** Northern European 96%.

**EVC2-related Ellis-van Creveld Syndrome** - Gene: EVC2. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_147127:1-22. **Detection Rate:** Northern European >99%. MALE DONOR 12445 DOB Ethnicity: Northern European Barcode: 11004212718545 FEMALE N/A

 Fabry Disease - Gene: GLA. X-linked Recessive. Sequencing with copy number analysis. Exons: NM\_000169:1-7. Detection Rate: Northern European 98%.
 Familial Dysautonomia - Gene: IKBKAP. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_003640:2-37. Detection Rate: Northern European >99%.

Familial Mediterranean Fever - Gene: MEFV. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000243:1-10. Detection Rate: Northern European >99%.

Fanconi Anemia Complementation Group A - Gene: FANCA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000135:1-43. Detection Rate: Northern European 92%.

Fanconi Anemia, FANCC-related - Gene: FANCC. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000136:2-15. Detection Rate: Northern European >99%.

FKRP-related Disorders - Gene: FKRP. Autosomal Recessive. Sequencing with copy number analysis. Exon: NM\_024301:4. Detection Rate: Northern European >99%. FKTN-related Disorders - Gene: FKTN. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_001079802:3-11. Detection Rate: Northern European >99%.

Galactokinase Deficiency - Gene: GALK1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000154:1-8. Detection Rate: Northern European >99%.

Galactosemia - Gene: GALT. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000155:1-11. Detection Rate: Northern European >99%. Gamma-sarcoglycanopathy - Gene: SGCG. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000231:2-8. Detection Rate: Northern European 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: Northern European 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: Northern European >99%.

GLB1-related Disorders - Gene: GLB1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000404:1-16. Detection Rate: Northern European

>99%. GLDC-related Glycine Encephalopathy - Gene: GLDC. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000170:1-25. Detection Rate: Northern European 94%.

Glutaric Acidemia, GCDH-related - Gene: GCDH. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000159:2-12. Detection Rate: Northern European >99%.

**Glycogen Storage Disease Type la** - Gene: G6PC. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_000151:1-5. **Detection Rate:** Northern European >99%.

**Glycogen Storage Disease Type Ib** - Gene: SLC37A4. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_001164277:3-11. **Detection Rate:** Northern European >99%.

**Glycogen Storage Disease Type III** - **Gene:** AGL. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_000642:2-34. **Detection Rate:** Northern European >99%.

**GNE Myopathy** - Gene: GNE. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_001128227:1-12. Detection Rate: Northern European >99%. **GNPTAB-related Disorders** - Gene: GNPTAB. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_024312:1-21. Detection Rate: Northern European >99%.

HADHA-related Disorders - Gene: HADHA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000182:1-20. Detection Rate: Northern European >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: Northern European >99%.



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Hereditary Fructose Intolerance - Gene: ALDOB. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000035:2-9. Detection Rate: Northern European >99%.

Herlitz Junctional Epidermolysis Bullosa, LAMB3-related - Gene: LAMB3. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000228:2-23. Detection Rate: Northern European >99%.

Hexosaminidase A Deficiency (Including Tay-Sachs Disease) - Gene: HEXA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000520:1-14. Detection Rate: Northern European >99%.

HMG-CoA Lyase Deficiency - Gene: HMGCL. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000191:1-9. Detection Rate: Northern European 98%.

Holocarboxylase Synthetase Deficiency - Gene: HLCS. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000411:4-12. Detection Rate: Northern European >99%.

Homocystinuria, CBS-related - Gene: CBS. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000071:3-17. Detection Rate: Northern European >99%.

Hydrolethalus Syndrome - Gene: HYLS1. Autosomal Recessive. Sequencing with copy number analysis. Exon: NM\_145014:4. Detection Rate: Northern European >99%.

Hypophosphatasia - Gene: ALPL. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000478:2-12. Detection Rate: Northern European >99%.

**Isovaleric Acidemia** - Gene: IVD. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_002225:1-12. Detection Rate: Northern European >99%.

Joubert Syndrome 2 - Gene: TMEM216. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_001173990:1-5. Detection Rate: Northern European >99%.

Junctional Epidermolysis Bullosa, LAMA3-related - Gene: LAMA3. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000227:1-38. Detection Rate: Northern European >99%.

Junctional Epidermolysis Bullosa, LAMC2-related - Gene: LAMC2. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_005562:1-23. Detection Rate: Northern European >99%.

KCNJ11-related Familial Hyperinsulinism - Gene: KCNJ11. Autosomal Recessive. Sequencing with copy number analysis. Exon: NM\_000525:1. Detection Rate: Northern European >99%.

Krabbe Disease - Gene: GALC. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000153:1-17. Detection Rate: Northern European >99%.

LAMA2-related Muscular Dystrophy - Gene: LAMA2. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000426:1-65. Detection Rate: Northern European >99%.

Leigh Syndrome, French-Canadian Type - Gene: LRPPRC. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_133259:1-38. Detection Rate: Northern European >99%.

Lipoid Congenital Adrenal Hyperplasia - Gene: STAR. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000349:1-7. Detection Rate: Northern European >99%.

Lysosomal Acid Lipase Deficiency - Gene: LIPA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000235:2-10. Detection Rate: Northern European >99%.

Maple Syrup Urine Disease Type Ia - Gene: BCKDHA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000709:1-9. Detection Rate: Northern European >99%.

Maple Syrup Urine Disease Type Ib - Gene: BCKDHB. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_183050:1-10. Detection Rate: Northern European >99%.

Maple Syrup Urine Disease Type II - Gene: DBT. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_001918:1-11. Detection Rate: Northern European 96%.

Medium Chain Acyl-CoA Dehydrogenase Deficiency - Gene: ACADM. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000016:1-12. Detection Rate: Northern European >99%.

Megalencephalic Leukoencephalopathy with Subcortical Cysts - Gene: MLC1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_015166:2-12. Detection Rate: Northern European >99%.

Metachromatic Leukodystrophy - Gene: ARSA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000487:1-8. Detection Rate: Northern European >99%.

Methylmalonic Acidemia, cblA Type - Gene: MMAA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_172250:2-7. Detection Rate: Northern European >99%.

Methylmalonic Acidemia, cblB Type - Gene: MMAB. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_052845:1-9. Detection Rate: Northern European >99%.

Methylmalonic Aciduria and Homocystinuria, cblC Type - Gene: MMACHC. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM 015506:1-4. Detection Rate: Northern European >99%.

MKS1-related Disorders - Gene: MKS1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_017777:1-18. Detection Rate: Northern European >99%.

**Mucolipidosis III Gamma** - Gene: GNPTG. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_032520:1-11. Detection Rate: Northern European >99%.

Mucolipidosis IV - Gene: MCOLN1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_020533:1-14. Detection Rate: Northern European >99%.

**Mucopolysaccharidosis Type I** - Gene: IDUA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000203:1-14. Detection Rate: Northern European >99%.

**Mucopolysaccharidosis Type II** - Gene: IDS. X-linked Recessive. Sequencing with copy number analysis. Exons: NM\_000202:1-9. Detection Rate: Northern European 88%.

**Mucopolysaccharidosis Type IIIA** - Gene: SGSH. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000199:1-8. Detection Rate: Northern European >99%.

Mucopolysaccharidosis Type IIIB - Gene: NAGLU. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000263:1-6. Detection Rate: Northern European >99%.

**Mucopolysaccharidosis Type IIIC** - Gene: HGSNAT. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_152419:1-18. Detection Rate: Northern European >99%.

MUT-related Methylmalonic Acidemia - Gene: MUT. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000255:2-13. Detection Rate: Northern European >99%.

MYO7A-related Disorders - Gene: MYO7A. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000260:2-49. Detection Rate: Northern European >99%.

**NEB-related Nemaline Myopathy** - Gene: NEB. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_001271208:3-80,117-183. Detection Rate: Northern European 92%.

**Nephrotic Syndrome, NPHS1-related** - Gene: NPHS1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_004646:1-29. Detection Rate: Northern European >99%.

Nephrotic Syndrome, NPHS2-related - Gene: NPHS2. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_014625:1-8. Detection Rate: Northern European >99%.

Niemann-Pick Disease Type C1 - Gene: NPC1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000271:1-25. Detection Rate: Northern European >99%.

Niemann-Pick Disease Type C2 - Gene: NPC2. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_006432:1-5. Detection Rate: Northern European >99%.



Niemann-Pick Disease, SMPD1-related - Gene: SMPD1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000543:1-6. Detection Rate: Northern European >99%.

Nijmegen Breakage Syndrome - Gene: NBN. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_002485:1-16. Detection Rate: Northern European >99%.

**Ornithine Transcarbamylase Deficiency** - Gene: OTC. X-linked Recessive. Sequencing with copy number analysis. **Exons:** NM\_000531:1-10. **Detection Rate:** Northern European 97%.

PCCA-related Propionic Acidemia - Gene: PCCA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000282:1-24. Detection Rate: Northern European 95%.

PCCB-related Propionic Acidemia - Gene: PCCB. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000532:1-15. Detection Rate: Northern European >99%.

PCDH15-related Disorders - Gene: PCDH15. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_033056:2-33. Detection Rate: Northern European 93%.

**Pendred Syndrome** - Gene: SLC26A4. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000441:2-21. Detection Rate: Northern European >99%.

**Peroxisome Biogenesis Disorder Type 1** - Gene: PEX1. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_000466:1-24. **Detection Rate:** Northern European >99%.

**Peroxisome Biogenesis Disorder Type 3** - Gene: PEX12. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_000286:1-3. **Detection Rate:** Northern European >99%.

**Peroxisome Biogenesis Disorder Type 4** - Gene: PEX6. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_000287:1-17. **Detection Rate:** Northern European 97%.

**Peroxisome Biogenesis Disorder Type 5** - Gene: PEX2. Autosomal Recessive. Sequencing with copy number analysis. **Exon:** NM\_000318:4. **Detection Rate:** Northern European >99%.

**Peroxisome Biogenesis Disorder Type 6** - Gene: PEX10. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_153818:1-6. **Detection Rate:** Northern European >99%.

Phenylalanine Hydroxylase Deficiency - Gene: PAH. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000277:1-13. Detection Rate: Northern European >99%.

**POMGNT-related Disorders** - Gene: POMGNT1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_017739:2-22. Detection Rate: Northern European 96%.

Pompe Disease - Gene: GAA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000152:2-20. Detection Rate: Northern European 98%.

**PPT1-related Neuronal Ceroid Lipofuscinosis** - Gene: PPT1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000310:1-9. Detection Rate: Northern European >99%.

**Primary Carnitine Deficiency** - Gene: SLC22A5. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_003060:1-10. **Detection Rate:** Northern European >99%.

**Primary Hyperoxaluria Type 1** - Gene: AGXT. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_000030:1-11. **Detection Rate:** Northern European >99%.

**Primary Hyperoxaluria Type 2** - Gene: GRHPR. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_012203:1-9. Detection Rate: Northern European >99%.

**Primary Hyperoxaluria Type 3** - Gene: HOGA1. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_138413:1-7. **Detection Rate:** Northern European >99%.

**Pycnodysostosis** - Gene: CTSK. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000396:2-8. Detection Rate: Northern European >99%.

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**Pyruvate Carboxylase Deficiency** - Gene: PC. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_000920:3-22. **Detection Rate:** Northern European >99%.

Rhizomelic Chondrodysplasia Punctata Type 1 - Gene: PEX7. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000288:1-10. Detection Rate: Northern European >99%.

**RTEL1-related Disorders** - Gene: RTEL1. Autosomal Recessive. Sequencing with copy number analysis. **Exons:** NM\_032957:2-35. **Detection Rate:** Northern European >99%.

Salla Disease - Gene: SLC17A5. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_012434:1-11. Detection Rate: Northern European 98%.

Sandhoff Disease - Gene: HEXB. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000521:1-14. Detection Rate: Northern European >99%.

Short-chain Acyl-CoA Dehydrogenase Deficiency - Gene: ACADS. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000017:1-10. Detection Rate: Northern European >99%.

Sjogren-Larsson Syndrome - Gene: ALDH3A2. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000382:1-10. Detection Rate: Northern European 96%.

SLC26A2-related Disorders - Gene: SLC26A2. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000112:2-3. Detection Rate: Northern European >99%.

Smith-Lemli-Opitz Syndrome - Gene: DHCR7. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_001360:3-9. Detection Rate: Northern European >99%.

Spastic Paraplegia Type 15 - Gene: ZFYVE26. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_015346:2-42. Detection Rate: Northern European >99%.

Spinal Muscular Atrophy - Gene: SMN1. Autosomal Recessive. Spinal muscular atrophy. Variant (1): SMN1 copy number. Detection Rate: Northern European 95%. Spondylothoracic Dysostosis - Gene: MESP2. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_001039958:1-2. Detection Rate: Northern European >99%.

TGM1-related Autosomal Recessive Congenital Ichthyosis - Gene: TGM1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000359:2-15. Detection Rate: Northern European >99%.

**TPP1-related Neuronal Ceroid Lipofuscinosis** - Gene: TPP1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000391:1-13. Detection Rate: Northern European >99%.

Tyrosine Hydroxylase Deficiency - Gene: TH. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_199292:1-14. Detection Rate: Northern European >99%.

Tyrosinemia Type I - Gene: FAH. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000137:1-14. Detection Rate: Northern European >99%.

Tyrosinemia Type II - Gene: TAT. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000353:2-12. Detection Rate: Northern European >99%.

**USH1C-related Disorders** - Gene: USH1C. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_005709:1-21. Detection Rate: Northern European >99%.

USH2A-related Disorders - Gene: USH2A. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_206933:2-72. Detection Rate: Northern European 94%.

Usher Syndrome Type 3 - Gene: CLRN1. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_174878:1-3. Detection Rate: Northern European >99%.

Very-long-chain Acyl-CoA Dehydrogenase Deficiency - Gene: ACADVL. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000018:1-20. Detection Rate: Northern European >99%.



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X-linked Myotubular Myopathy - Gene: MTM1. X-linked Recessive. Sequencing with copy number analysis. Exons: NM\_000252:2-15. Detection Rate: Northern European 98%.

X-linked Severe Combined Immunodeficiency - Gene: IL2RG. X-linked Recessive. Sequencing with copy number analysis. Exons: NM\_000206:1-8. Detection Rate: Northern European >99%.

Xeroderma Pigmentosum Group A - Gene: XPA. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000380:1-6. Detection Rate: Northern European >99%.

Xeroderma Pigmentosum Group C - Gene: XPC. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_004628:1-16. Detection Rate: Northern European 97%.

Wilson Disease - Gene: ATP7B. Autosomal Recessive. Sequencing with copy number analysis. Exons: NM\_000053:1-21. Detection Rate: Northern European >99%.

X-linked Adrenoleukodystrophy - Gene: ABCD1. X-linked Recessive. Sequencing with copy number analysis. Exons: NM\_000033:1-6. Detection Rate: Northern European 77%.

X-linked Alport Syndrome - Gene: COL4A5. X-linked Recessive. Sequencing with copy number analysis. Exons: NM\_000495:1-51. Detection Rate: Northern European 95%.

X-linked Congenital Adrenal Hypoplasia - Gene: NR0B1. X-linked Recessive. Sequencing with copy number analysis. Exons: NM\_000475:1-2. Detection Rate: Northern European 99%.

X-linked Juvenile Retinoschisis - Gene: RS1. X-linked Recessive. Sequencing with copy number analysis. Exons: NM\_000330:1-6. Detection Rate: Northern European 98%.



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FEMALE N/A

# **Risk Calculations**

Below are the risk calculations for all conditions tested. Since negative results do not completely rule out the possibility of being a carrier, the **residual risk** represents the patient's post-test likelihood of being a carrier and the **reproductive risk** represents the likelihood the patient's 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 after a negative test result. Inaccurate reporting of ethnicity may cause errors in risk calculation. The reproductive risk presented is based on a hypothetical pairing with a partner of the same ethnic group.

†Indicates a positive result. See the full clinical report for interpretation and details.

11.beta-hydroxylase-deficient Congenital Adrenal Hyperplasia         1 in 3,800         <1 in 1,000,000           6-privosyl-tetrahydropterin Synthase Deficiency         <1 in 1,000,000         <1 in 1,000,000           ABCC3R-related Familial Hyperplasia         1 in 17,000         <1 in 1,000,000           ABCC3R-related Familial Hyperinsulnism         1 in 17,000         <1 in 1,000,000           Abpha-stassesmia         Alpha globin status: sa/aa.         Not calculated           Alpha-stassegylcxenopathy         1 in 35,000         <1 in 1,000,000           Alpha-stacgylcxenopathy         1 in 50,000         <1 in 1,000,000           Andreman Syndrome         <1 in 50,000         <1 in 1,000,000           Andreman Syndrome         <1 in 50,000         <1 in 1,000,000           Argininosuccinic Aciduria         1 in 1,000,000         <1 in 1,000,000           Argininosuccinic Aciduria         1 in 1,000,000         <1 in 1,000,000           Ataxia with Vtamin E Deficiency         <1 in 50,000         <1 in 1,000,000           Ataxia with Vtamin E Deficiency         <1 in 10,000,000         <1 in 1,000,000           Ataxia with Vtamin E Deficiency         <1 in 1,000,000         <1 in 1,000,000           Ataxia with Vtamin E Deficiency         <1 in 1,000,000         <1 in 1,000,000           Ataxia with Vtamin E Deficiency         <	Disease	DONOR 12445 Residual Risk	Reproductive Risk
G-pyrusyl-tetrahydropterin Synthase Deficiency         < 1 in 100,000         < 1 in 100,000           ABCC8-related Familial Hyperinsulinism         1 in 17,000         < 1 in 1,000,000         1 in 0,00,000			
ABCCB-related Familial Hyperinsulinism         1 in 17,000         < 1 in 1000,000           Adenosine Derinicacy         1 in 22,000         < 1 in 1000,000           Alpha Thalassemia         Alpha globin status: aa/aa.         Not calculated           Alpha-smooidosis         1 in 35,000         < 1 in 1,000,000           Alpha-sarcoglycanopathy         1 in 50,000         < 1 in 1,000,000           AltTrestated Glycine Encephalopathy         1 in 50,000         < 1 in 1,000,000           Andresited Glycine Encephalopathy         1 in 17,000         < 1 in 1,000,000           Andresited Glycine Encephalopathy         1 in 17,000         < 1 in 1,000,000           Argininasuccinic Aciduria         1 in 17,000         < 1 in 1,000,000           Argininasuccinic Aciduria         1 in 13,000         < 1 in 1,000,000           Ataxia with Vitamin E Deficiency         1 in 50,000         < 1 in 1,000,000           Ataxia with Vitamin E Deficiency         1 in 1,000,000         < 1 in 1,000,000           Autosomal Recessive Oblycytic Kidney Disease, PKHD1-related         1 in 35,000         < 1 in 1,000,000           Autosomal Recessive Polycytic Kidney Disease, PKHD1-related         1 in 32,000         < 1 in 1,000,000           Bardet-Bield Syndrome, BBS1-related         1 in 50,000         < 1 in 1,000,000           Bardet-Bield Syndrome, BBS			
Adenosine Dearninase Deficiency         1 in 22,000         < 1 in 1,000,000           Alpha Thalassemia         Alpha globin status: aa/aa.         Not calculated           Alpha-manosidosis         1 in 35,000         < 1 in 1,000,000           Alpha-sarcoglycanopathy         1 in 45,000         < 1 in 1,000,000           Alpha-sarcoglycanopathy         1 in 25,000         < 1 in 1,000,000           AMT-related Glycine Encephalopathy         1 in 20,000         < 1 in 1,000,000           Ardgrinnemia         < 1 in 1,000,000         < 1 in 1,000,000           Argininemia         < 1 in 1,000,000         < 1 in 1,000,000           Argininegicasinuria         1 in 30,000         < 1 in 1,000,000           Aspartylgicosaminuria         1 in 50,000         < 1 in 1,000,000           Ataxia -telangicetasia         1 in 1,000,000         < 1 in 1,000,000           Ataxia -telangicetasia         1 in 1,000,000         < 1 in 1,000,000           Ataxia -telangicetasia         1 in 1,000,000         < 1 in 1,000,000           Autosomal Recessive Polycystic Kidney Disease, PKHD1-related         1 in 35,000         < 1 in 1,000,000           Autosomal Recessive Polycystic Kidney Disease, PKHD1-related         1 in 42,000         < 1 in 1,000,000           Bardet-Bield Syndrome, BBS1-related         1 in 32,000         < 1 in 1,000,0			
Alpha         Alpha globin staus: aa/aa.         Not calculated           Alpha-mannosidosis         1 in 35,000         <1 in 1,000,000           Alpha-sarcoglycanopathy         1 in 45,000         <1 in 1,000,000           Alpha-sarcoglycanopathy         1 in 50,000         <1 in 1,000,000           AltTrelated Glycine Encephalopathy         1 in 20,000         <1 in 1,000,000           Andermann Syndrome         <1 in 50,000         <1 in 1,000,000           Argininosucchir Aciduria         1 in 13,000         <1 in 1,000,000           Argininosucchir Aciduria         1 in 13,000         <1 in 1,000,000           Ataxia with Vitamin E Deficiency         <1 in 50,000         <1 in 1,000,000           Ataxia with Vitamin E Deficiency         <1 in 1,000,000         <1 in 1,000,000           Ataxia with Vitamin E Deficiency         <1 in 1,000,000         <1 in 1,000,000           Autoinmune Polyglandular Syndrome Type 1         1 in 1,500,00         <1 in 1,000,000           Autoismune Recessive Ostopoetrosis Type 1         1 in 8,100         <1 in 1,000,000           Autoismune Recessive Spastic Ataxia of Charlevoix-Saguenay         <1 in 4,000         <1 in 1,000,000           Bardet-Bield Syndrome, BBS1-related         1 in 2,000         <1 in 1,000,000         <1 in 1,000,000         <1 in 1,000,000         <1 in 1,000,000			
Alpha-mannosidosis         1 in 5,000         <1 in 1,000,000           Alpha-sarcoglycanopathy         1 in 45,000         <1 in 1,000,000           Alstrom Syndrome         <1 in 5,000         <1 in 1,000,000           Andermann Syndrome         1 in 5,000         <1 in 1,000,000           Andermann Syndrome         <1 in 5,000         <1 in 1,000,000           Argininemia         <1 in 7,000         <1 in 1,000,000           Argininemia         <1 in 1,000,000         <1 in 1,000,000           Argininemia         <1 in 5,000         <1 in 1,000,000           Argininesucini Aciduria         1 in 5,000         <1 in 1,000,000           Atxaia-telangiectasia         1 in 1,000,000         <1 in 1,000,000           Atxaia-telangiectasia         1 in 1,50,000         <1 in 1,000,000           Autosomal Recessive Polycystic Kidney Disease, PKHD1-related         1 in 8,000         <1 in 1,000,000           Autosomal Recessive Polycystic Kidney Disease, PKHD1-related         1 in 4,000         <1 in 1,000,000           Autosomal Recessive Polycystic Kidney Disease, PKHD1-related         1 in 4,000         <1 in 1,000,000           Bardet-Bield Syndrome, BBS1-related         1 in 5,000         <1 in 1,000,000         <1 in 1,000,000           Bardet-Bield Syndrome, BBS1-related         1 in 5,0000         <1 in 1,000,000	•		
Alpha-sarcoglycanopathy         1 in 45,000         <1 in 1,000,000           Alstrom Syndrome         <1 in 50,000         <1 in 1,000,000           AndTr-related Glycine Encephalopathy         1 in 22,000         <1 in 1,000,000           AndErrelated Glycine Encephalopathy         1 in 22,000         <1 in 1,000,000           Argininemia         <1 in 50,000         <1 in 1,000,000           Argininemia         1 in 13,000         <1 in 1,000,000           Atsia with Vitamin E Deficiency         <1 in 50,000         <1 in 1,000,000           Ataxia with Vitamin E Deficiency         <1 in 50,000         <1 in 1,000,000           Ataxia with Vitamin E Deficiency         <1 in 50,000         <1 in 1,000,000           Attaxia with Vitamin E Deficiency         <1 in 50,000         <1 in 1,000,000           Attaxia with Vitamin E Deficiency         <1 in 1,000,000         <1 in 1,000,000           Autoimune Polyglandular Syndrome Type 1         1 in 15,000         <1 in 1,000,000           Autoimune Polyglandular Syndrome Type 1         1 in 32,000         <1 in 1,000,000           Autoimune Besti Syndrome, BBS1-related         1 in 32,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS1-related         1 in 50,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS1-related         1 in 50,000         <1	•	· · ·	
Alstrom Syndrome         <1 in 50,000         <1 in 1,000,000           AMT-related Glycine Encephalopathy         1 in 50,000         <1 in 1,000,000           Anderman Syndrome         <1 in 50,000         <1 in 1,000,000           Argininosuccinic Aciduria         1 in 13,000         <1 in 1,000,000           Argininosuccinic Aciduria         1 in 13,000         <1 in 1,000,000           Aspartyfiglucosaminuria         <1 in 50,000         <1 in 1,000,000           Ataxia vith Vitamin E Deficiency         <1 in 50,000         <1 in 1,000,000           Ataxia-telangiettasia         1 in 1,000,000         <1 in 0,000,000           Ataxia-telangiettasia         1 in 1,000,000         <1 in 0,000,000           Autosomal Recessive Polycystic Kidney Disease, PKHD1-related         1 in 8,100         <1 in 1,000,000           Autosomal Recessive Polycystic Kidney Disease, PKHD1-related         1 in 4,000         <1 in 1,000,000           Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay         <1 in 50,000         <1 in 1,000,000           Bardet-Bield Syndrome, BBS10-related         1 in 42,000         <1 in 1,000,000           Bardet-Bield Syndrome, BBS12-related         1 in 50,000         <1 in 1,000,000           Bardet-Bield Syndrome, BBS12-related         1 in 50,000         <1 in 1,000,000           Bardet-Bield Syndrome, BBS12-r	•		
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 1,000,000         <1 in 1,000,000           Argininemia         <1 in 1,000,000         <1 in 1,000,000           Argininesuccinic Aciduria         1 in 1,000,000         <1 in 1,000,000           Ataxia with Vitamin E Deficiency         <1 in 50,000         <1 in 1,000,000           Ataxia with Vitamin E Deficiency         <1 in 50,000         <1 in 1,000,000           Ataxia with Vitamin E Deficiency         <1 in 1,000,000         <1 in 600,000           Autoimune Polyglandular Syndrome Type 1         1 in 1,500,000         <1 in 1,000,000           Autosomal Recessive Osteopetrosis Type 1         1 in 8,5000         <1 in 1,000,000           Autosomal Recessive Polycystic Kidney Disease, PKHD1-related         1 in 8,100         <1 in 1,000,000           Autosomal Recessive Polycystic Kidney Disease, PKHD1-related         1 in 4,000         <1 in 1,000,000           Bardet-Bield Syndrome, BBS1-related         1 in 50,000         <1 in 1,000,000         <1 in 50,000         <1 in 1,000,000         <1 in 1,000,000 <td< th=""><th></th><th></th><th></th></td<>			
Andermann Syndrome         <1 in 50,000         <1 in 1000,000           Argininemia         <1 in 17,000         <1 in 1000,000           Argininemia         1 in 17,000         <1 in 1000,000           Argininemia         1 in 13,000         <1 in 1000,000           Aspartylglucosaminuria         <1 in 50,000         <1 in 1000,000           Ataxia with Vitamin E Deficiency         <1 in 50,000         <1 in 1,000,000           Ataxia-telangiectasia         1 in 1,000,000         <1 in 1,000,000           Attaxia-telangiectasia         <1 in 1,000,000         <1 in 1,000,000           Autoismune Polyglandlular Syndrome Type 1         1 in 15,000         <1 in 1,000,000           Autosomal Recessive Polycystic Kidney Disease, PKHD1-related         1 in 8,100         <1 in 1,000,000           Autosomal Recessive Polycystic Kidney Disease, PKHD1-related         1 in 32,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS1-related         1 in 32,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS1-related         1 in 50,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS1-related         1 in 50,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS1-related         1 in 50,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS12-related         <1 in 50,000	•		
Argininemia         <1 in 17,000         <1 in 1,000,000           Argininosuccinic Aciduria         1 in 13,000         <1 in 1,000,000           Aspartylgluccosaminuria         <1 in 50,000         <1 in 1,000,000           Ataxia with Vitamin E Deficiency         <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 1,000,000         <1 in 0,000,000           Attra-telangiectasia         1 in 1,000,000         <1 in 0,000,000           Autosmal Recessive Ostoepetrosis Type 1         1 in 5,000         <1 in 1,000,000           Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay         <1 in 4,000         <1 in 1,000,000           Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay         <1 in 44,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS10-related         1 in 50,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS10-related         <1 in 50,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS10-related         <1 in 50,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS10-related         <1 in 50,000         <1 in 1,000,000           Bord Syndrome, BBS10-related         <1 in 50,000         <1 in 1,000,000           Bord Syndrome, BBS10-related	· · · · ·		
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 with Vitamin E Deficiency         <1 in 50,000         <1 in 1,000,000           Ataxia with Vitamin E Deficiency         <1 in 1,000,000         1 in 1,000,000           Ataxia with Vitamin E Deficiency         <1 in 1,000,000         1 in 600,000           Autosomal Recessive Osteopetrosis Type 1         1 in 15,000         <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           Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay         <1 in 42,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BB51-related         1 in 50,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BB51-related         <1 in 50,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BB51-related         <1 in 50,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BB52-related         <1 in 50,000         <1 in 1,000,000           Border Biedl Syndrome, BB52-related         <1 in 50,000         <1 in 1,000,00	•		
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 vita mith Vitamin E Deficiency         <1 in 50,000         <1 in 1,000,000           Ataxia vita mith Vitamin E Deficiency         <1 in 1,000,000         <1 in 1,000,000           Ataxia vita mith Polyglandular Syndrome Type 1         1 in 15,000         <1 in 1,000,000           Autosomal Recessive Ostopetrosis Type 1         1 in 35,000         <1 in 1,000,000           Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay         <1 in 44,000         <1 in 1,000,000           Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay         <1 in 42,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS1-related         1 in 32,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS2-related         <1 in 50,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS2-related         <1 in 50,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS2-related         <1 in 30,000         <1 in 1,000,000           Bottonicidase Deficiency         1 in 1,30,000         <1 in 1,000,000           Calpanopathy         1 in 30,000         <1 in 1,000,000           Garavan Disease         1 in 7,000,000         <1 in 1,000,000           Car			
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           Attra-telangiectasia         1 in 1,000,000         1 in 600,000           Autoimuune 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 44,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS1-related         1 in 32,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS1-related         1 in 50,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS2-related         <1 in 50,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS2-related         <1 in 50,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BBS2-related         <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 1,000,000           Biotinidase Deficiency         1 in 1,000,000         <1 in 1,000,000           Calpainopathy         1 in 3,000         <1 in 1,000,000         <1 in 1,000,000           Carbanoylphosphate Synthetase I			
Ataxia-telangiectasia         1 in 11,000         <1 in 1,000,000           ATP3A-related Disorders         <1 in 1,000,000         1 in 600,000           Autoimmune Polyglandular Syntome 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 Osteopetrosis Type 1         1 in 35,000         <1 in 1,000,000           Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay         <1 in 4,000         <1 in 1,000,000           Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay         <1 in 4,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BB51-related         1 in 32,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BB51-related         1 in 50,000         <1 in 1,000,000           Bardet-Biedl Syndrome, BB52-related         <1 in 50,000         <1 in 1,000,000           BCS11-related Disorders         <1 in 50,000         <1 in 1,000,000           BCS11-related Disorders         <1 in 50,000         <1 in 1,000,000           Bloom Syndrome         <1 in 50,000         <1 in 1,000,000           Canavan Disease         1 in 3,000         <1 in 1,000,000           Carabinoylphosphate Synthetase I Deficiency         1 in 50,000         <1 in 1,000,000           Caraitine Palmitoyltransferase IA	, , , , , , , , , , , , , , , , , , , ,		
ATP7A-related Disorders         < 1 in 1,000,000         1 in 600,000           Autosomal Recessive Osteopetrosis 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 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           Bardet-Biedl Syndrome, BBS1-related         1 in 32,000         < 1 in 1,000,000           Bardet-Biedl Syndrome, BBS12-related         1 in 50,000         < 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           BcS11-related Disorders         < 1 in 50,000         < 1 in 1,000,000           BcS12-related Disorders         < 1 in 50,000         < 1 in 1,000,000           BcS11-related Disorders         < 1 in 50,000         < 1 in 1,000,000           BcS11-related Disorders         < 1 in 50,000         < 1 in 1,000,000           Gcahamoylphosphate         1 in 1,000,000         < 1 in 1,000,000         < 1 in 1,000,000            Ganavan Disease         1 in 9,700         < 1 in 1,000,000         < 1 in	•		
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 Spatic Ataxia of Charlevoix-Saguenay         < 1 in 44,000         < 1 in 0,000,000           Bardet-Biedl Syndrome, BBS1-related         1 in 32,000         < 1 in 1,000,000           Bardet-Biedl Syndrome, BBS1-related         1 in 42,000         < 1 in 1,000,000           Bardet-Biedl Syndrome, BBS1-related         1 in 42,000         < 1 in 1,000,000           Bardet-Biedl Syndrome, BBS1-related         1 in 50,000         < 1 in 1,000,000           Bardet-Biedl Syndrome, BBS1-related         < 1 in 50,000         < 1 in 1,000,000           Bardet-Biedl Syndrome, BBS2-related         < 1 in 50,000         < 1 in 1,000,000           Bardet-Biedl Syndrome, BBS2-related         < 1 in 50,000         < 1 in 1,000,000           Bardet-Biedl Syndrome, BBS2-related         < 1 in 50,000         < 1 in 1,000,000           Bardet-Biedl Syndrome, BBS2-related         < 1 in 50,000         < 1 in 1,000,000           Bardet-Biedl Syndrome, BBS2-related         < 1 in 50,000         < 1 in 1,000,000           Castama Disease         < 1 in 50,000         < 1 in 1,000,000           Carbamoylphosphate Synthetase I Deficiency         < 1 in 50,000         < 1 in 1,000,00	•		
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, BBS12-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           BcS1L-related Disorders         < 1 in 50,000         < 1 in 1,000,000           BcS1L-related Disorders         < 1 in 50,000         < 1 in 1,000,000           Biotinidase Deficiency         1 in 13,000         < 1 in 1,000,000           Calpainopathy         1 in 13,000         < 1 in 1,000,000           Caravan Disease         1 in 57,000         < 1 in 1,000,000           Caramoylphosphate Synthetase I Deficiency         < 1 in 50,000         < 1 in 1,000,000           Carnitine Palmitoyltransferase I Deficiency         1 in 50,000         < 1 in 1,000,000           Carnitin			
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, BBS1-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           Beta-sarcoglycanopathy         1 in 3,000         < 1 in 1,000,000           Botonidase Deficiency         1 in 13,000         < 1 in 1,000,000           Biotinidase Deficiency         1 in 13,000         < 1 in 1,000,000           Calpainopathy         1 in 3,000         < 1 in 1,000,000           Carbamoylphosphate Synthetase I Deficiency         1 in 9,700         < 1 in 1,000,000           Carritine Palmitoyltransferase II Deficiency         1 in 57,000         < 1 in 1,000,000           Cartilage-hair Hypoplasia         < 1 in 50,000         < 1 in 1,000,000           Cartilage-hair Hypoplasia         < 1 in 50,000         < 1 in 1,000,000           Cartilage-hair Hypoplasia         < 1 in 50,000         < 1 in 1,000,000           Cartilage-hair Hyp			
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           BcS1L-related Disorders         < 1 in 50,000         < 1 in 1,000,000           Btom Syndrome, BBS2-related         < 1 in 50,000         < 1 in 1,000,000           BcS1L-related Disorders         < 1 in 50,000         < 1 in 1,000,000           Btom Syndrome         < 1 in 1,000,000         < 1 in 1,000,000           Galpainopathy         1 in 13,000         < 1 in 1,000,000           Carbamoylphosphate Synthetase I Deficiency         < 1 in 50,000         < 1 in 1,000,000           Carnitine Palmitoyltransferase IA Deficiency         < 1 in 50,000         < 1 in 1,000,000           Carnitine Palmitoyltransferase ID Deficiency         < 1 in 50,000         < 1 in 1,000,000           Carnitine Palmitoyltransferase I Deficiency         1 in 14,000         < 1 in 1,000,000           Carn			< 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           BCS11-related Disorders         < 1 in 50,000         < 1 in 1,000,000           BcS11-related Disorders         < 1 in 50,000         < 1 in 1,000,000           Bta-sarcoglycanopathy         1 in 39,000         < 1 in 1,000,000           Biom Syndrome         < 1 in 50,000         < 1 in 1,000,000           Calpainopathy         1 in 13,000         < 1 in 1,000,000           Carbamoylphosphate Synthetase I Deficiency         1 in 9,700         < 1 in 1,000,000           Carbamoylphosphate Synthetase I Deficiency         < 1 in 50,000         < 1 in 1,000,000           Carbamoylphosphate Synthetase I Deficiency         < 1 in 50,000         < 1 in 1,000,000           Carbamoylphosphate Synthetase I Deficiency         < 1 in 50,000         < 1 in 1,000,000           Carbamoylphosphate Synthetase I Deficiency         < 1 in 50,000         < 1 in 1,000,000           Carbamoylphosphate Synthetase I Deficiency         < 1 in 50,000         < 1 in 1,000,000           Carbamoylphosphat	Autosomal Recessive Polycystic Kidney Disease, PKHD1-related	1 in 8,100	< 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           BcS11-related Disorders         < 1 in 50,000         < 1 in 1,000,000           BcS11-related Disorders         < 1 in 50,000         < 1 in 1,000,000           BcS11-related Disorders         < 1 in 50,000         < 1 in 1,000,000           Botinidase 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           Calpainopathy         1 in 1,000,000         < 1 in 1,000,000           Carbamoylphosphate Synthetase I Deficiency         1 in 50,000         < 1 in 1,000,000           Carnitine Palmitoyltransferase IA Deficiency         < 1 in 57,000         < 1 in 1,000,000           Carnitine Palmitoyltransferase II Deficiency         < 1 in 50,000         < 1 in 1,000,000           Carnitine Palmitoyltransferase II Deficiency         < 1 in 50,000         < 1 in 1,000,000           Carnitine Palmitoyltransferase II Deficiency         < 1 in 50,000         < 1 in 1,000,000           Carnitine Palmitoyltransferase II Deficiency	Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay	< 1 in 44,000	< 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           Carbamoylphosphate Synthetase I Deficiency         < 1 in 50,000         < 1 in 1,000,000           Carbamoylphosphate Synthetase I Deficiency         < 1 in 50,000         < 1 in 1,000,000           Carbinous Xanthomatosis         1 in 1,000,000         < 1 in 1,000,000         < 1 in 1,000,000           Cartilage-hair Hypoplasia         < 1 in 50,000         < 1 in 1,000,000         < 1 in 1,000,000         < 1 in 1,000,000           Citrullinemia Type 1         1 in 14,000         < 1 in 1,000,000	Bardet-Biedl Syndrome, BBS1-related	1 in 32,000	< 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           Biotom Syndrome         < 1 in 50,000         < 1 in 1,000,000           Calpainopathy         1 in 13,000         < 1 in 1,000,000           Calpainopathy         1 in 1,000         < 1 in 1,000,000           Carbamoylphosphate Synthetase I Deficiency         < 1 in 57,000         < 1 in 1,000,000           Carritine Palmitoyltransferase IA Deficiency         < 1 in 50,000         < 1 in 1,000,000           Cartilage-hair Hypoplasia         < 1 in 50,000         < 1 in 1,000,000           Cartilage-hair Hypoplasia         < 1 in 50,000         < 1 in 1,000,000           Citrullinemia Type 1         1 in 11,000         < 1 in 1,000,000           CLN3-related Neuronal Ceroid Lipofuscinosis         1 in 8,600         < 1 in 1,000,000           CLN5-related Neuronal Ceroid Lipofuscinosis         < 1 in 50,000         < 1 in 1,000,000           CLN5-related Neuronal Ceroid Lipofuscinosis         < 1 in 50,000         < 1 in 1,000,000           CLN5-related Neuronal Ceroid Lipofuscinosis         < 1 in 50,000	Bardet-Biedl Syndrome, BBS10-related	1 in 42,000	< 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           Calpainopathy         1 in 13,000         < 1 in 1,000,000           Caravan Disease         1 in 9,700         < 1 in 1,000,000           Carbamoylphosphate Synthetase I Deficiency         < 1 in 57,000         < 1 in 1,000,000           Carritine Palmitoyltransferase IA Deficiency         < 1 in 50,000         < 1 in 1,000,000           Cartilage-hair Hypoplasia         < 1 in 50,000         < 1 in 1,000,000           Cartilage-hair Hypoplasia         < 1 in 50,000         < 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         1 in 43,000         < 1 in 1,000,000           CLN6-related Neuronal Ceroid Lipofuscinosis         1 in 43,000         < 1 in 1,000,000           CLN6-related Neuronal Ceroid Lipofuscinosis         1 in 50,000         < 1 in 1,000,000 <th>Bardet-Biedl Syndrome, BBS12-related</th> <th>&lt; 1 in 50,000</th> <th>&lt; 1 in 1,000,000</th>	Bardet-Biedl Syndrome, BBS12-related	< 1 in 50,000	< 1 in 1,000,000
Beta-sarcoglycanopathy         1 in 39,000         < 1 in 1,000,000	Bardet-Biedl Syndrome, BBS2-related	< 1 in 50,000	< 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           Carbamoylphosphate Synthetase I Deficiency         < 1 in 50,000         < 1 in 1,000,000           Carnitine Palmitoyltransferase IA Deficiency         < 1 in 50,000         < 1 in 1,000,000           Carnitine Palmitoyltransferase II Deficiency         < 1 in 50,000         < 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 50,000         < 1 in 1,000,000           CLN5-related Neuronal Ceroid Lipofuscinosis         < 1 in 50,000         < 1 in 1,000,000           CLN6-related Neuronal Ceroid Lipofuscinosis         < 1 in 50,000         < 1 in 1,000,000           CLN8-related Neuronal Ceroid Lipofuscinosis         < 1 in 50,000         < 1 in 1,000,000	BCS1L-related Disorders	< 1 in 50,000	< 1 in 1,000,000
Bloom Syndrome         < 1 in 50,000	Beta-sarcoglycanopathy	1 in 39,000	< 1 in 1,000,000
Calpainopathy         1 in 13,000         < 1 in 1,000,000	Biotinidase Deficiency	1 in 13,000	1 in 650,000
Canavan Disease         1 in 9,700         < 1 in 1,000,000	Bloom Syndrome	< 1 in 50,000	< 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           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 50,000         < 1 in 1,000,000           CLN5-related Neuronal Ceroid Lipofuscinosis         < 1 in 50,000         < 1 in 1,000,000           CLN6-related Neuronal Ceroid Lipofuscinosis         1 in 43,000         < 1 in 1,000,000           CLN8-related Neuronal Ceroid Lipofuscinosis         < 1 in 50,000         < 1 in 1,000,000	Calpainopathy	1 in 13,000	< 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 50,000         < 1 in 1,000,000           CLN5-related Neuronal Ceroid Lipofuscinosis         < 1 in 50,000         < 1 in 1,000,000           CLN6-related Neuronal Ceroid Lipofuscinosis         1 in 43,000         < 1 in 1,000,000           CLN8-related Neuronal Ceroid Lipofuscinosis         1 in 50,000         < 1 in 1,000,000	Canavan Disease	1 in 9,700	< 1 in 1,000,000
Carnitine Palmitoyltransferase II Deficiency         1 in 25,000         < 1 in 1,000,000	Carbamoylphosphate Synthetase I Deficiency	< 1 in 57,000	< 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         < 1 in 50,000         < 1 in 1,000,000           CLN6-related Neuronal Ceroid Lipofuscinosis         1 in 43,000         < 1 in 1,000,000           CLN8-related Neuronal Ceroid Lipofuscinosis         1 in 50,000         < 1 in 1,000,000	Carnitine Palmitoyltransferase IA Deficiency	< 1 in 50,000	< 1 in 1,000,000
Cerebrotendinous Xanthomatosis         1 in 11,000         < 1 in 1,000,000	Carnitine Palmitoyltransferase II Deficiency	1 in 25,000	< 1 in 1,000,000
Citrullinemia Type 1         1 in 14,000         < 1 in 1,000,000	Cartilage-hair Hypoplasia	< 1 in 50,000	< 1 in 1,000,000
CLN3-related Neuronal Ceroid Lipofuscinosis         1 in 8,600         < 1 in 1,000,000	Cerebrotendinous Xanthomatosis	1 in 11,000	< 1 in 1,000,000
CLN5-related Neuronal Ceroid Lipofuscinosis         < 1 in 50,000	Citrullinemia Type 1	1 in 14,000	< 1 in 1,000,000
CLN6-related Neuronal Ceroid Lipofuscinosis         1 in 43,000         < 1 in 1,000,000	CLN3-related Neuronal Ceroid Lipofuscinosis	1 in 8,600	< 1 in 1,000,000
CLN8-related Neuronal Ceroid Lipofuscinosis < 1 in 50,000 < 1 in 1,000,000	CLN5-related Neuronal Ceroid Lipofuscinosis	< 1 in 50,000	< 1 in 1,000,000
	CLN6-related Neuronal Ceroid Lipofuscinosis	1 in 43,000	< 1 in 1,000,000
Cohen Syndrome < 1 in 15 000 < 1 in 1 000 000	CLN8-related Neuronal Ceroid Lipofuscinosis	< 1 in 50,000	< 1 in 1,000,000
	Cohen Syndrome	< 1 in 15,000	< 1 in 1,000,000
COL4A3-related Alport Syndrome         1 in 6,200         < 1 in 1,000,000	COL4A3-related Alport Syndrome	1 in 6,200	< 1 in 1,000,000
		1 in 12,000	< 1 in 1,000,000
Combined Pituitary Hormone Deficiency, PROP1-related 1 in 6,100 < 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	• •		
			< 1 in 1,000,000
			< 1 in 1,000,000
			< 1 in 1,000,000
	• • •		< 1 in 1,000,000



MALE DONOR 12445 DOB: Ethnicity: Northern European Barcode: 11004212718545 FEMALE N/A

Disease	DONOR 12445 Residual Risk	Reproductive Risk
Cystic Fibrosis	1 in 3,000	1 in 360,000
Cystinosis	1 in 22,000	< 1 in 1,000,000
D-bifunctional Protein Deficiency	1 in 9,000	< 1 in 1,000,000
Delta-sarcoglycanopathy	< 1 in 40,000	< 1 in 1,000,000
Dihydrolipoamide Dehydrogenase Deficiency	< 1 in 50,000	< 1 in 1,000,000
	1 in 11,000	< 1 in 1,000,000
Dysferlinopathy		
Dystrophinopathy (Including Duchenne/Becker Muscular Dystrophy)	Not calculated	Not calculated
ERCC6-related Disorders	1 in 26,000	< 1 in 1,000,000
ERCC8-related Disorders	< 1 in 9,900	< 1 in 1,000,000
EVC-related Ellis-van Creveld Syndrome	1 in 7,500	< 1 in 1,000,000
EVC2-related Ellis-van Creveld Syndrome	< 1 in 50,000	< 1 in 1,000,000
Fabry Disease	< 1 in 1,000,000	1 in 80,000
Familial Dysautonomia	< 1 in 50,000	< 1 in 1,000,000
Familial Mediterranean Fever	< 1 in 50,000	< 1 in 1,000,000
Fanconi Anemia Complementation Group A	1 in 2,800	< 1 in 1,000,000
Fanconi Anemia, FANCC-related	< 1 in 50,000	< 1 in 1,000,000
FKRP-related Disorders	1 in 16,000	< 1 in 1,000,000
FKTN-related Disorders	< 1 in 50,000	
		< 1 in 1,000,000
Galactokinase Deficiency	1 in 10,000	< 1 in 1,000,000
Galactosemia	Q188R heterozygote <sup>†</sup>	1 in 350
Gamma-sarcoglycanopathy	1 in 3,000	< 1 in 1,000,000
Gaucher Disease	1 in 260	1 in 110,000
GJB2-related DFNB1 Nonsyndromic Hearing Loss and Deafness	1 in 2,500	1 in 260,000
GLB1-related Disorders	1 in 19,000	< 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 35,000	< 1 in 1,000,000
Glycogen Storage Disease Type III	1 in 16,000	< 1 in 1,000,000
GNE Myopathy	1 in 23,000	< 1 in 1,000,000
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 Si	ckle Cell 1 in 3,100	1 in 390,000
Disease)		1 070,000
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
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 < 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
Megalencephalic Leukoencephalopathy with Subcortical Cysts	< 1 in 50,000	< 1 in 1,000,000
Metachromatic Leukodystrophy	1 in 16,000	< 1 in 1,000,000
Methylmalonic Acidemia, cblA Type	< 1 in 50,000	< 1 in 1,000,000
Methylmalonic Acidemia, cblB Type	1 in 48,000	< 1 in 1,000,000
Methylmalonic Aciduria and Homocystinuria, cblC Type	1 in 16,000	< 1 in 1,000,000
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MALE DONOR 12445 DOB: Ethnicity: Northern European Barcode: 11004212718545 FEMALE N/A

	DONOR 12445	
Disease	Residual Risk	Reproductive Risk
Mucolipidosis III Gamma	< 1 in 50,000	< 1 in 1,000,000
Mucolipidosis IV	< 1 in 50,000	< 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 16,000	< 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
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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
Salla Disease	< 1 in 30,000	< 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
Siggren-Larsson Syndrome		
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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
	Negative for g.27134T>G SNP	
Spinal Muscular Atrophy	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
Very-long-chain Acyl-CoA Dehydrogenase Deficiency	1 in 18,000	< 1 in 1,000,000
Wilson Disease	1 in 8,600	< 1 in 1,000,000
X-linked Adrenoleukodystrophy	1 in 90,000	1 in 42,000
X-linked Alport Syndrome	Not calculated	Not calculated



MALE DONOR 12445 DOB: Ethnicity: Northern European Barcode: 11004212718545 FEMALE N/A

DONOR 12445 Residual Risk	Reproductive Risk
< 1 in 1,000,000	1 in 40,000
Not calculated	Not calculated
< 1 in 1,000,000	1 in 200,000
< 1 in 50,000	< 1 in 1,000,000
1 in 7,300	< 1 in 1,000,000
	Residual Risk           < 1 in 1,000,000