

Patient Information: Partner Information: Not Tested DOB: Sex: M MR#: BFA0179 Patient#: Accession Accession: N/A Test# Specimen Type: Saliva Swab Collected: Nov 30,2022 FINAL RESULTS

Physician: Shaikly, Valerie ATTN: Shaikly, Valerie Fertility Genetics 1 Lanswood Park Elmstead Market, Essex CO7 7FD GB Dr. Hanlin (Harry) Gao Phone: 7711197938

Laboratory: **Fulgent Genetics** CAP#: 8042697 CLIA#: 05D2043189 Laboratory Director: Report Date: Dec 22,2022

TEST PERFORMED

Carrier for ONE genetic condition Genetic counseling is recommended.

176 Matched Fors Male (167 Gene Panel; gene sequencing

with deletion and duplication analysis)

Condition and Gene	Inheritance		Partner
Glycine encephalopathy, GLDC-related GLDC	AR	Carrier c.2316-1G>A (p.?)	N/A

INTERPRETATION:

Notes and Recommendations:

- Based on these results, this individual is positive for a carrier mutation in 1 gene. The risk estimates below are quantified based on general population carrier frequencies. Carrier screening for the reproductive partner is recommended to accurately assess this risk:
 - There is a 1/772 chance of having a child affected with Glycine encephalopathy, GLDC-related, a GLDC-related condition.
- Testing for copy number changes in the SMN1 gene was performed to screen for the carrier status of Spinal Muscular Atrophy. The results for this individual are within the normal range for non-carriers. See Limitations section for more information.
- This carrier screening test does not screen for all possible genetic conditions, nor for all possible mutations in every gene tested. Individuals with negative test results may still have up to a 3-4% risk to have a child with a birth defect due to genetic and/or environmental factors.
- Patients may wish to discuss any carrier results with blood relatives, as there is an increased chance that they are also carriers. These results should be interpreted in the context of this individual's clinical findings, biochemical profile, and family history.
- · Gene specific notes and limitations may be present. See below.
- This report does not include variants of uncertain significance.
- Genetic counseling is recommended. Contact your physician about the available options for genetic counseling.



GLYCINE ENCEPHALOPATHY, GLDC-RELATED

Patient		Partner
Result	 Carrier 	N/A
Variant Details	<i>GLDC</i> (NM_000170.3) c.2316-1G>A (p.?)	N/A

What is Glycine encephalopathy, GLDC-related?

Glycine encephalopathy is characterized by abnormally high levels of a molecule called glycine which also acts as a neurotransmitter. Glycine encephalopathy is caused by the shortage of an enzyme that normally breaks down glycine in the body. A lack of this enzyme allows excess glycine to build up in tissues and organs, particularly the brain, leading to serious medical problems. The most common form of glycine encephalopathy, called the classical type, appears shortly after birth. Affected infants experience lethargy, feeding difficulties, hypotonia, abnormal jerking movements, and life-threatening problems with breathing. Most children who survive these early signs and symptoms develop profound intellectual disability and seizures that are difficult to treat. For unknown reasons, affected males are more likely to survive and have less severe developmental problems than affected females.

What is my risk of having an affected child?

The risk of being a carrier of Glycine encephalopathy is about 1/193 in the general population. Individuals of Finnish descent have an increased carrier risk of 1/117. Individuals from British Columbia have reported a carrier risk of 1/125. If both partners are carriers for the disease allele, the risk for an affected child is 1 in 4 (25%).

What kind of medical management is available?

No definitive treatment exists for glycine encephalopathy. Most cases of glycine encephalopathy present as the neonatal form of disease and neonates may die without major medical intervention (e.g. intubation and ventilation). For the cases that survive past infancy, medical management is based on reducing serum glycine levels, seizure management; gastrostomy tube for feeding problems; therapy for gastroesophageal reflux; and physical therapy. Prognosis is poor for most of these patients.

What mutation was detected?

The detected heterozygous variant was NM_000170.3:c.2316-1G>A (p.?). This intronic variant alters the highly conserved splice acceptor site for exon 20 of this transcript and is predicted by all four splice site prediction tools queried to abolish canonical splice acceptor activity. This variant is expected to result in altered function of the GLDC gene product as a result of aberrant splicing. This variant has been reported in the homozygous and compound heterozygous state in several unrelated individuals with glycine encephalopathy (also known as nonketotic hyperglycinemia) (PubMed: 16601880, 27362913). The laboratory classifies this variant as pathogenic.



GENES TESTED:

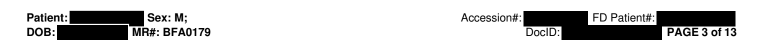
176 Matched Fors Male - 167 Genes

This analysis was run using the 176 Matched Fors Male gene list. 167 genes were tested with 98.99% of targets sequenced at >20x coverage. For more gene specific information and assistance with residual risk calculation, see the SUPPLEMENTAL TABLE.

ABCC8	ACADM	ACADS	ACADVL	ADA	AGA
AGL	AGXT	AIRE	ALDH3A2	ALDOB	ALG6
ALMS1	ALPL	AMT	ARG1	ARSA	ASL
ASPA	ASS1	ATM	ATP7B	BBS1	BBS10
BBS12	BBS2	BCKDHA	BCKDHB	BCS1L	BLM
BTD	CAPN3	CBS	CFTR	CLN3	CLN5
CLN6	CLN8	CLRN1	COL4A3	COL4A4	CPS1
CPT1A	CPT2	CRYL1	CTNS	CTSK	CYP11B1
CYP21A2	CYP27A1	DBT	DHCR7	DLD	DYSF
ELP1	ERCC6	ERCC8	EVC	EVC2	FAH
FANCA	FANCC	FKRP	FKTN	G6PC	GAA
GALC	GALK1	GALT	GBA	GCDH	GJB2
GJB6	GLB1	GLDC	GNE	GNPTAB	GNPTG
GRHPR	HADHA	HBA1	HBA2	HBB	HEXA
HEXB	HGSNAT	HLCS	HMGCL	HOGA1	HSD17B4
HYLS1	IDUA	IVD	KCNJ11	LAMA2	LAMA3
LAMB3	LAMC2	LIPA	LRPPRC	MAN2B1	MCOLN1
MEFV	MESP2	MKS1	MLC1	MMAA	MMAB
MMACHC	MPI	MUT	MYO7A	NAGLU	NBN
NEB	NPC1	NPC2	NPHS1	NPHS2	OPA3
PAH	PC	PCCA	PCCB	PCDH15	PEX1
PEX10	PEX12	PEX2	PEX6	PEX7	PKHD1
PMM2	POMGNT1	PPT1	PROP1	PTS	RMRP
RTEL1	SACS	SGCA	SGCB	SGCD	SGCG
SGSH	SLC12A6	SLC17A5	SLC22A5	SLC26A2	SLC26A4
SLC37A4	SMN1	SMPD1	STAR	TAT	TCIRG1
TGM1	ТН	TMEM216	TPP1	TTPA	USH1C
USH2A	VPS13B	XPA	XPC	ZFYVE26	

METHODS:

Genomic DNA was isolated from the submitted specimen indicated above (if cellular material was submitted). DNA was barcoded, and enriched for the coding exons of targeted genes using hybrid capture technology. Prepared DNA libraries were then sequenced using a Next Generation Sequencing technology. Following alignment to the human genome reference sequence (assembly GRCh37), variants were detected in regions of at least 10x coverage. For this specimen, 99.09% and 98.99% of coding regions and splicing junctions of genes listed had been sequenced with coverage of at least 10x and 20x, respectively, by NGS or by Sanger sequencing. The remaining regions did not have 10x coverage, and were not evaluated. Variants were interpreted manually using locus specific databases, literature searches, and other molecular biological principles. To minimize false positive results, any variants that do not meet internal quality standards are confirmed by Sanger sequencing. Variants classified as pathogenic, likely pathogenic, or risk allele which are located in the coding regions and nearby intronic regions (+/- 20bp) of the genes listed above are reported. Variants outside these intervals may be reported but are typically not guaranteed. When a single pathogenic or likely pathogenic variant is identified in a clinically relevant gene with autosomal recessive inheritance, the laboratory will attempt to ensure 100% coverage of coding sequences either through NGS or Sanger sequencing technologies ("fill-in"). All genes listed were evaluated for large deletions and/or duplications. However, single exon deletions or duplications will not be detected in this assay, nor will copy number alterations in regions of genes with significant pseudogenes. Putative deletions or duplications are analyzed using Fulgent Germline proprietary pipeline for this specimen. New York patients: diagnostic findings are confirmed by Sanger, MLPA, or gPCR; exception SNV variants in genes for which confirmation of NGS results has been performed >=10 times may not be confirmed if identified with high quality by NGS. Bioinformatics: The Fulgent Germline v2019.2 pipeline was used to analyze this specimen.





LIMITATIONS:

General Limitations

These test results and variant interpretation are based on the proper identification of the submitted specimen, accuracy of any stated familial relationships, and use of the correct human reference sequences at the queried loci. In very rare instances, errors may result due to mix-up or co-mingling of specimens. Positive results do not imply that there are no other contributors. genetic or otherwise, to future pregnancies, and negative results do not rule out the genetic risk to a pregnancy. Official gene names change over time. Fulgent uses the most up to date gene names based on HUGO Gene Nomenclature Committee (https://www.genenames.org) recommendations. If the gene name on report does not match that of ordered gene, please contact the laboratory and details can be provided. Result interpretation is based on the available clinical and family history information for this individual, collected published information, and Alamut annotation available at the time of reporting. This assay is not designed or validated for the detection of low-level mosaicism or somatic mutations. This assay will not detect certain types of genomic aberrations such as translocations, inversions, or repeat expansions other than specified genes. DNA alterations in regulatory regions or deep intronic regions (greater than 20bp from an exon) may not be detected by this test. Unless otherwise indicated, no additional assays have been performed to evaluate genetic changes in this specimen. There are technical limitations on the ability of DNA sequencing to detect small insertions and deletions. Our laboratory uses a sensitive detection algorithm, however these types of alterations are not detected as reliably as single nucleotide variants. Rarely, due to systematic chemical, computational, or human error, DNA variants may be missed. Although next generation seguencing technologies and our bioinformatics analysis significantly reduce the confounding contribution of pseudogene sequences or other highly-homologous sequences, sometimes these may still interfere with the technical ability of the assay to identify pathogenic alterations in both sequencing and deletion/duplication analyses. Deletion/duplication analysis can identify alterations of genomic regions which include one whole gene (buccal swab specimens and whole blood specimens) and are two or more contiguous exons in size (whole blood specimens only); single exon deletions or duplications may occasionally be identified, but are not routinely detected by this test. When novel DNA duplications are identified, it is not possible to discern the genomic location or orientation of the duplicated segment, hence the effect of the duplication cannot be predicted. Where deletions are detected, it is not always possible to determine whether the predicted product will remain in-frame or not. Unless otherwise indicated, deletion/duplication analysis has not been performed in regions that have been sequenced by Sanger.

Gene Specific Notes and Limitations

BTD: If detected, the variant NM_001370658.1:c.1270G>C (p.Asp424His) will not be reported as this variant is associated with low disease penetrance and is primarily associated with reduced enzyme activity when homozygous. CFTR: Analysis of the intron 8 polymorphic region (e.g. IVS8-5T allele) is only performed if the p.Arg117His (R117H) mutation is detected. Single exon deletion/duplication analysis is limited to deletions of previously reported exons: 1, 2, 3, 11, 19, 20, 21. CRYL1: As mutations in the CRYL1 gene are not known to be associated with any clinical condition, sequence variants in this gene are not analyzed. However, to increase copy number detection sensitivity for large deletions including this gene and a neighboring on gene on the panel (GJB6, also known as connexin 30), this gene was evaluated for copy number variation. CYP11B1: The current testing method is not able to reliably detect certain pathogenic variants in this gene due to the interference by highly homologous regions. This analysis is not designed to detect or rule-out copy-neutral chimeric CYP11B1/CYP11B2 gene. CYP21A2: Significant pseudogene interference and/or reciprocal exchanges between the CYP21A2 gene and its pseudogene, CYP21A1P, have been known to occur and may impact results. As such, the relevance of variants reported in this gene must be interpreted clinically in the context of the clinical findings, biochemical profile, and family history of each patient. The variants c.188A>T (p.His63Leu), c.844G>T (p.Val282Leu), c.1174G>A (p.Ala392Thr), and c.1360C>T (p.Pro454Ser) in CYP21A2 will not be routinely reported as these variants are primarily associated with non-classic congenital adrenal hyperplasia and low disease penetrance. Additionally, the variant c.955C>T (p.Gln319Ter) is in the region with pseudogene interference, and the probability of this variant occurring in the real gene is greater than 50%. When observed, this variant will be reported as a possible carrier without LR-PCR. The confirmation test is recommended if the second reproductive partner is tests positive for variants in CYP21A2. GALT: In general, the D2 "Duarte" allele is not reported if detected, but can be reported upon request. While this allele can cause positive newborn screening results, it is not known to cause clinical symptoms in any state (PubMed: 25473725, 30593450). GBA: The current testing method may not be able to reliably detect certain pathogenic variants in the GBA gene due to homologous recombination between the pseudogene and the functional gene. HBA1: The phase of heterozygous alterations in the HBA1 gene cannot be determined, but can be confirmed through parental testing. HBA2: The phase of heterozygous alterations in the HBA2 gene cannot be determined, but can be confirmed through parental testing. NEB: This gene contains a 32-kb triplicate region (exons 82-105) which is not amenable to sequencing and deletion/duplication analysis. <u>NPHS2:</u> If detected, the variant NM_014625.3:c.686G>A (p.Arg229GIn) will not be reported as this variant is not significantly associated with disease when homozygous or in the compound heterozygous state with

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variants in exons 1-6 of NPHS2. <u>SMN1</u>: The current testing method detects sequencing variants in exon 7 and copy number variations in exons 7-8 of the SMN1 gene (NM_022874.2). Sequencing and deletion/duplication analysis are not performed on any other region in this gene. About 5%-8% of the population have two copies of SMN1 on a single chromosome and a deletion on the other chromosome, known as a [2+0] configuration (PubMed: 20301526). The current testing method cannot directly detect carriers with a [2+0] SMN1 configuration, but can detect linkage between the silent carrier allele and certain population-specific single nucleotide changes. As a result, a negative result for carrier testing greatly reduces but does not eliminate the chance that a person is a carrier. Only abnormal results will be reported.

SIGNATURE:

a Nelonie A

Melanie Jones, Ph.D., CGMBS, FACMG on 12/22/2022 03:18 PM PST Electronically signed

DISCLAIMER:

This test was developed and its performance characteristics determined by **Fulgent Genetics**. It has not been cleared or approved by the FDA. The laboratory is regulated under CLIA as qualified to perform high-complexity testing. This test is used for clinical purposes. It should not be regarded as investigational or for research. Since genetic variation, as well as systematic and technical factors, can affect the accuracy of testing, the results of testing should always be interpreted in the context of clinical and familial data. For assistance with interpretation of these results, healthcare professionals may contact us directly at (626) 350-0537 or info@fulgentgenetics.com. It is recommended that patients receive appropriate genetic counseling to explain the implications of the test result, including its residual risks, uncertainties and reproductive or medical options.

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			emental Table	Carrier	Detection	Post-test	
Gene	Condition	Inheritance	Ethnicity	Rate	Rate	Carrier Probability*	Residual Risk*
ABCC8	Familial hyperinsulinism	AR	General Population Ashkenazi Jewish Population Finnish Population	1 in 112 1 in 44 1 in 25	98% 98% 98%	1 in 5,551 1 in 2,151 1 in 1,201	1 in 2,486,848 1 in 378,576 1 in 120,100
			Middle-Eastern Population	1 in 25	98%	1 in 1,201	1 in 120,100
ACADM	Medium-chain acyl-CoA dehydrogenase (MCAD) deficiency	AR	General Population Caucasian / European Population East Asian Population Native American Population	1 in 69 1 in 52 1 in 198 1 in 43	98% 99% 99% 96%	1 in 3,401 1 in 5,101 1 in 19,701 1 in 1,051	1 in 938,676 1 in 1,061,008 <1 in 10 million 1 in 180,772
ACADS	Short-chain acyl-coA dehydrogenase (SCAD) deficiency	AR	General Population African/African American Population Caucasian / European Population	1 in 85 1 in 52 1 in 76	99% 99% 99%	1 in 8,401 1 in 5,101 1 in 7,501	1 in 2,856,340 1 in 1,061,008 1 in 2,280,304
			Middle-Eastern Population South Asian/Indian Population	1 in 52 1 in 51	99% 99%	1 in 5,101 1 in 5,001	1 in 1,061,008 1 in 1,020,204
ACADVL	Very long-chain acyl-CoA dehydrogenase (VLCAD) deficiency	AR	General Population Middle-Eastern Population Native American Population South Asian/Indian Population	1 in 118 1 in 74 1 in 61 1 in 73	93% 93% 93% 93%	1 in 1,672 1 in 1,044 1 in 858 1 in 1,030	1 in 789,184 1 in 309,024 1 in 209,352 1 in 300,760
ADA	Adenosine deaminase deficiency	AR	General Population	1 in 224	93%	1 in 3,187	1 in 2,855,552
AGA	Aspartylglucosaminuria	AR	General Population Finnish Population	<1 in 500 1 in 71	98% 98%	1 in 24,951 1 in 3,501	<1 in 10 million 1 in 994,284
AGL	Glycogen storage disease type III	AR	General Population Faroese Population Inuit Population North African Jewish Population	1 in 158 1 in 28 1 in 25 1 in 37	95% 95% 95% 95%	1 in 3,141 1 in 541 1 in 481 1 in 721	1 in 1,985,112 1 in 60,592 1 in 48,100 1 in 106,708
AGXT	Primary hyperoxaluria type 1	AR	General Population Caucasian / European Population	1 in 120 1 in 173	99% 99%	1 in 11,901	1 in 5,712,480
AIRE	Autoimmune polyendocrinopathy syndrome type I	AR	General Population Finnish Population	1 in 150 1 in 79	98% 98%	1 in 7,451 1 in 3,901	1 in 4,470,600 1 in 1,232,716
ALDH3A2	Sjögren-Larsson syndrome	AR	General Population	1 in 250	98%	1 in 12,451	<1 in 10 millio
ALDOB	Hereditary fructose intolerance	AR	General Population African/African American Population Caucasian / European Population	1 in 122 1 in 250 1 in 67	99% 99% 99%	1 in 24,901 1 in 6,601	1 in 5,905,288 <1 in 10 million 1 in 1,769,068
ALG6	Congenital disorder of glycosylation type Ic	AR	Middle-Eastern Population General Population	1 in 97 <1 in 500	99% 98%	1 in 9,601	1 in 3,725,188
ALMS1	Alstrom syndrome	AR	General Population	1 in 500	98%		<1 in 10 millio
ALPL	Hypophosphatasia	AR	General Population Caucasian / European Population Mennonite Population	1 in 158 1 in 274 1 in 25	95% 95% 95%	1 in 3,141 1 in 5,461 1 in 481	1 in 1,985,112 1 in 5,985,256 1 in 48,100
AMT	Glycine encephalopathy	AR	General Population Finnish Population	1 in 373 1 in 117	98% 98%	1 in 18,601 1 in 5,801	<1 in 10 millio 1 in 2,714,868
ARG1	Arginase deficiency	AR	General Population	1 in 296	98%	1 in 14,751	<1 in 10 millio
ARSA	Metachromatic leukodystrophy	AR	General Population Caucasian / European Population Yemenite Jewish Population	1 in 100 1 in 78 1 in 75	99% 99% 99%	1 in 9,901 1 in 7,701 1 in 7,401	1 in 3,960,400 1 in 2,402,712 1 in 2,220,300
ASL	Argininosuccinate lyase deficiency	AR	General Population	1 in 132	90%	1 in 1,311	1 in 692,208
ASPA	Canavan disease	AR	General Population Ashkenazi Jewish Population	1 in 300 1 in 55	97% 96%	1 in 9,968 1 in 1,351	<1 in 10 millio 1 in 297,220
ASS1	Citrullinemia	AR	General Population East Asian Population	1 in 119 1 in 132	96% 96%	1 in 2,951 1 in 3,276	1 in 1,404,676 1 in 1,729,728
ATM	Ataxia-telangiectasia	AR	General Population	1 in 100	92%	1 in 1,239	1 in 495,600
ATP7B	Wilson disease	AR	General Population Caucasian / European Population Ashkenazi Jewish Population	1 in 87 1 in 42 1 in 70	98% 98% 98%	1 in 4,301 1 in 2,051 1 in 3,451	1 in 1,496,748 1 in 344,568 1 in 966,280
BBS1	Bardet-Biedl syndrome type 1	AR	General Population	1 in 367	99%		<1 in 10 millio
BBS10	Bardet-Biedl syndrome type 10	AR	General Population	1 in 395	99%		<1 in 10 millio
3BS12 3BS2	Bardet-Biedl syndrome type 12 Bardet-Biedl syndrome 2	AR AR	General Population General Population Ashkenazi Jewish Population	1 in 791 1 in 621 1 in 107	99% 99% 99%	1 in 62,001	<1 in 10 millio <1 in 10 millio
BBS2	Retinitis Pigmentosa 74	AR	General Population Ashkenazi Jewish Population	1 in 107 1 in 621 1 in 107	99% 99% 99%	1 in 62,001	1 in 4,537,228 <1 in 10 millio 1 in 4,537,228
BCKDHA	Maple syrup urine disease type la	AR	General Population Mennonite Population	1 in 321 1 in 10	98% 98%		<1 in 10 millio 1 in 18,040
	Maple syrup urine disease type Ib	AR	General Population	1 in 364	98%	1 in 18,151	<1 in 10 million

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		Suppl	emental Table				
Gene	Condition	Inheritance	Ethnicity	Carrier Rate	Detection Rate	Post-test Carrier Probability*	Residual Risk*
BOOM	Differentia di anna dia ma		Ormanal Dansslation	1 - 500	000/	1 - 04 051	1 := 10 == :!!! = =
BCS1L BCS1L	Björnstad syndrome GRACILE syndrome	AR AR	General Population General Population	<1 in 500 <1 in 500			<1 in 10 million <1 in 10 million
BCS1L	Mitochondrial complex III deficiency	AR	General Population	<1 in 500		,	<1 in 10 million
BLM	Bloom syndrome	AR	General Population	1 in 800	87%	1 in 6,147	<1 in 10 million
			Ashkenazi Jewish Population	1 in 134	99%		1 in 7,129,336
BTD	Biotinidase deficiency	AR	General Population Caucasian / European Population Latino Population Middle-Eastern Population	1 in 124 1 in 71 1 in 136 1 in 55	99% 99% 99% 99%	1 in 7,001	1 in 6,101,296 1 in 1,988,284 1 in 7,344,544 1 in 1,188,220
CAPN3	Limb-girdle muscular dystrophy type 2A	AR	General Population Caucasian / European Population	<1 in 500 1 in 103	98% 98%	1 in 24,951 1 in 5,101	<1 in 10 million 1 in 2,101,612
CBS	Homocystinuria due to cystathionine beta-synthase deficiency	AR	General Population Caucasian / European Population	1 in 224 1 in 86	99% 99%	1 in 22,301 1 in 8,501	<1 in 10 million 1 in 2,924,344
CFTR	Cystic Fibrosis	AR	Middle-Eastern Population General Population	1 in 21 1 in 32	99% 99%	1 in 2,001 1 in 3,101	1 in 168,084 1 in 396,928
CFIR	Cystic Fibrosis	Ап	African/African American Population Ashkenazi Jewish Population Caucasian / European Population East Asian Population Latino Population	1 in 32 1 in 61 1 in 24 1 in 25 1 in 94 1 in 58	99% 99% 99% 99% 99%	1 in 3,101 1 in 6,001 1 in 2,301 1 in 2,401 1 in 9,301 1 in 5,701	1 in 1,464,244 1 in 220,896 1 in 240,100 1 in 3,497,176 1 in 1,322,632
CLN3	Neuronal ceroid lipofuscinosis	AR	General Population Finnish Population	1 in 230 1 in 72	98% 98%	1 in 11,451 1 in 3,551	<1 in 10 million 1 in 1,022,688
CLN5	Neuronal ceroid lipofuscinosis 5	AR	General Population Finnish Population	<1 in 500 1 in 115	95% 95%	1 in 9,981 1 in 2,281	<1 in 10 million 1 in 1,049,260
CLN6	Neuronal ceroid lipofuscinosis, CLN6-related	AR	General Population	<1 in 500	92%	1 in 6,239	<1 in 10 million
CLN8	Neuronal ceroid lipofuscinosis, CLN8-related	AR	General Population Finnish Population	<1 in 500 1 in 135	95% 95%	1 in 9,981 1 in 2,681	<1 in 10 million 1 in 1,447,740
CLRN1	Usher syndrome, type 3A	AR	General Population Ashkenazi Jewish Population Finnish Population	1 in 500 1 in 120 1 in 70	98% 98% 98%	1 in 24,951 1 in 5,951 1 in 3,451	<1 in 10 million 1 in 2,856,480 1 in 966,280
COL4A3	Alport syndrome, COL4A3-related	AR	General Population Ashkenazi Jewish Population	1 in 267 1 in 188	98% 98%	1 in 13,301 1 in 9,351	<1 in 10 million 1 in 7,031,952
COL4A4	Alport syndrome, COL4A4-related	AR	General Population	1 in 267	98%		<1 in 10 million
CPS1	Carbamoylphosphate synthetase I deficiency	AR	General Population	1 in 570	98%		<1 in 10 million
CPT1A	Carnitine palmitoyltransferase IA deficiency	AR	General Population Hutterite Population	1 in 354 1 in 16	90% 90%	1 in 3,531 1 in 151	1 in 4,999,896 1 in 9,664
CPT2	Carnitine palmitoyltransferase II deficiency	AR	General Population Ashkenazi Jewish Population	<1 in 500 1 in 51	95% 95%	1 in 9,981 1 in 1,001	<1 in 10 million 1 in 204,204
CRYL1	GJB6-CRYL1 related nonsyndromic hearing loss	UK	General Population	1 in 423	99%		<1 in 10 million
CTNS	Cystinosis	AR	General Population British Population	1 in 158 1 in 81	99% 99%	1 in 8,001	1 in 9,923,032 1 in 2,592,324
CTSK	Pycnodysostosis	AR	Moroccan Jewish Population General Population	1 in 100 <1 in 500			1 in 3,960,400 <1 in 10 million
CYP11B1	Congenital adrenal hyperplasia due to 11-beta-	AR	General Population	1 in 158	98%	1 in 7,851	1 in 4,961,832
CYP21A2	hydroxylase deficiency Congenital adrenal hyperplasia due to 21-hydroxylase deficiency	AR	Morrocan Jewish Population General Population Inuit Population Middle-Eastern Population	1 in 35 1 in 61 1 in 9 1 in 35	98% 99% 99%	1 in 1,701 1 in 6,001 1 in 801 1 in 2,401	1 in 238,140 1 in 1,464,244 1 in 28,836 1 in 476,140
CYP27A1	Cerebrotendinous xanthomatosis	AR	General Population Morrocan Jewish Population	1 in 35 1 in 500 1 in 5	99% 98% 98%	1 in 3,401 1 in 24,951 1 in 201	1 in 476,140 <1 in 10 million 1 in 4,020
DBT	Maple syrup urine disease, type II	AR	General Population	1 in 481	98%		<1 in 10 million
DHCR7	Smith-Lemli-Opitz syndrome	AR	General Population African/African American Population Ashkenazi Jewish Population	1 in 30 1 in 138 1 in 36	96% 96% 96%	1 in 726 1 in 3,426 1 in 876	1 in 87,120 1 in 1,891,152 1 in 126,144
DLD	Dihydrolipoamide dehydrogenase deficiency	AR	General Population Ashkenazi Jewish Population	1 in 500 1 in 107	98% 98%		<1 in 10 million 1 in 2,268,828
DYSF	Limb-girdle muscular dystrophy type 2B	AR	General Population Japanese Population Libyan Jewish Population	<1 in 500 1 in 332 1 in 18		1 in 9,981 1 in 6,621 1 in 341	<1 in 10 million 1 in 8,792,688 1 in 24,552
ELP1	Familial Dysautonomia	AR	General Population Ashkenazi Jewish Population	1 in 300 1 in 31	99% 99%	1 in 29,901 1 in 3,001	<1 in 10 million 1 in 372,124

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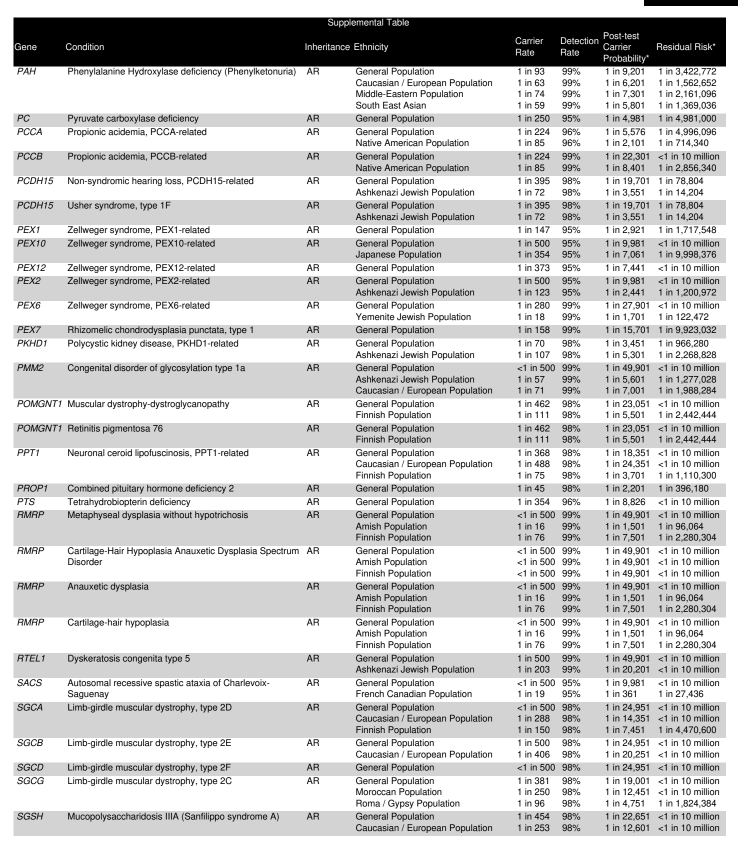
Supplemental Table							
Gene	Condition	Inheritance	Ethnicity	Carrier Rate	Detection Rate	Post-test Carrier Probability*	Residual Risk*
GLDC	Glycine encephalopathy, GLDC-related	AR	General Population British Columbia Canadian Population Finnish Population	1 in 193 1 in 125 1 in 117	98% 99% 99%		1 in 7,411,972 1 in 6,200,500 1 in 5,429,268
GNE	Inclusion body myopathy type 2 (Nonaka myopathy)	AR	General Population Iranian Jewish Population	<1 in 500 1 in 11	99% 99%	1 in 49,901 1 in 1,001	1 in 99,802,000 1 in 44,044
GNPTAB	Mucolipidosis II alpha/beta	AR	General Population	<1 in 500	95%	1 in 9,981	<1 in 10 million
GNPTAB	Mucolipidosis III alpha/beta	AR	General Population	<1 in 500	95%	1 in 9,981	<1 in 10 million
GNPTG	Mucolipidosis III gamma	AR	General Population	<1 in 500	95%	1 in 9,981	<1 in 10 million
GRHPR	Primary hyperoxaluria type II	AR	General Population	<1 in 500	99%	1 in 49,901	<1 in 10 million
HADHA	Trifunctional protein deficiency	AR	General Population Finnish Population	<1 in 500 1 in 124	98% 98%	1 in 24,951 1 in 6,151	<1 in 10 million 1 in 3,050,896
HADHA	Long-chain 3-hydroxyacyl-CoA dehydrogenase (LCHAD) deficiency	AR	General Population Finnish Population	<1 in 500 1 in 124	98% 98%	1 in 24,951 1 in 6,151	<1 in 10 million 1 in 3,050,896
HBA1	Alpha thalassemia	AR	General Population General Population† Southeast Asian Population Southeast Asian Population† Mediterranean Population† African/African American Population	1 in 18 1 in 1000 ≤1 in 7 ≤1 in 14 ≤1 in 6 1 in 500 1 in 30	98% 98% 98% 98% 98% 98%	1 in 860 1 in 860 ≤1 in 305 ≤1 in 305 ≤1 in 229 ≤1 in 229 1 in 1,451	1 in 3,440,364 1 in 3,440,364 ≤1 in 17,228 ≤1 in 17,228 ≤1 in 457,556 ≤1 in 457,556 1 in 5,804,000
HBA2	Alpha thalassemia	AR	General Population General Population† Southeast Asian Population Southeast Asian Population† Mediterranean Population† Mediterranean Population† African/African American Population	1 in 18 1 in 1000 ≤1 in 7 ≤1 in 14 ≤1 in 6 1 in 500 1 in 30	98% 98% 98% 98% 98% 98%	1 in 860 1 in 860 ≤1 in 305 ≤1 in 305 ≤1 in 229 ≤1 in 229 1 in 1,451	1 in 3,440,364 1 in 3,440,364 ≤1 in 17,228 ≤1 in 17,228 ≤1 in 457,556 ≤1 in 457,556 1 in 5,804,000
HBB	Sickle cell disease	AR	General Population African/African American Population East Asian Population Latino Population Mediterranean Population South Asian/Indian Population	1 in 158 1 in 10 1 in 50 1 in 128 1 in 3 1 in 25	95% 95% 95% 95% 95%	1 in 3,141 1 in 181 1 in 981 1 in 2,541 1 in 41 1 in 481	1 in 1,985,112 1 in 7,240 1 in 196,200 1 in 1,300,992 1 in 492 1 in 48,100
HBB	Hemoglobin C disease	AR	General Population African/African American Population East Asian Population Latino Population Mediterranean Population South Asian/Indian Population	1 in 158 1 in 10 1 in 50 1 in 128 1 in 3 1 in 25	95% 95% 95% 95% 95%	1 in 3,141 1 in 181 1 in 981 1 in 2,541 1 in 41 1 in 481	1 in 1,985,112 1 in 7,240 1 in 196,200 1 in 1,300,992 1 in 492 1 in 48,100
HBB	Beta thalassemia	AR	General Population African/African American Population East Asian Population Latino Population Mediterranean Population South Asian/Indian Population	1 in 158 1 in 10 1 in 50 1 in 128 1 in 3 1 in 25	95% 95% 95% 95% 95% 95%	1 in 3,141 1 in 181 1 in 981 1 in 2,541 1 in 41 1 in 481	1 in 1,985,112 1 in 7,240 1 in 196,200 1 in 1,300,992 1 in 492 1 in 48,100
HEXA	Tay-Sachs disease	AR	General Population Ashkenazi Jewish Population Moroccan Jewish Population	1 in 300 1 in 27 1 in 110	99% 99% 99%	1 in 2,601	<1 in 10 million 1 in 280,908 1 in 4,796,440
HEXB	Sandhoff disease	AR	General Population	1 in 600	98%		<1 in 10 million
HGSNAT	Mucopolysaccharidosis type IIIC (Sanfilippo syndrome C)	AR	General Population Caucasian / European Population	1 in 434 1 in 345	98% 98%	1 in 21,651	<1 in 10 million <1 in 10 million
HLCS	Holocarboxylase synthetase deficiency	AR	General Population	1 in 500	98%		<1 in 10 million
HMGCL	3-hydroxy-3-methylglutaryl-CoA lyase deficiency	AR	General Population	<1 in 500	98%		<1 in 10 million
HOGA1	Primary hyperoxaluria type III	AR	General Population	1 in 184	99%		<1 in 10 million
HSD17B4	D-bifunctional protein deficiency	AR	General Population	1 in 158	98%	1 in 7,851	1 in 4,961,832
HYLS1	Hydrolethalus syndrome	AR	General Population Finnish Population	<1 in 500 1 in 50		1 in 24,951 1 in 2,451	<1 in 10 million 1 in 490,200
IDUA	Mucopolysaccharidosis, type I (Hurler syndrome)	AR	General Population Caucasian / European Population	<1 in 500 1 in 153		1 in 9,981 1 in 3,041	<1 in 10 million 1 in 1,861,092
IVD	Isovaleric Acidemia	AR	General Population African/African American Population Caucasian / European Population East Asian Population	1 in 167 1 in 100 1 in 115 1 in 407	90% 90% 90% 90%	1 in 1,661 1 in 991 1 in 1,141 1 in 4,061	1 in 1,109,548 1 in 396,400 1 in 524,860 1 in 6,611,308

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		Suppl	emental Table				
Gene	Condition	Inheritance	Ethnicity	Carrier Rate	Detection Rate	Post-test Carrier Probability*	Residual Risk*
KCNJ11	Congenital hyperinsulinism	AR	General Population Caucasian / European Population	1 in 423 1 in 232	99% 99%		<1 in 10 million <1 in 10 million
KCNJ11	Permanent neonatal diabetes mellitus	AR	General Population Caucasian / European Population	1 in 423 1 in 232	99% 99%	,	<1 in 10 million <1 in 10 million
LAMA2	Muscular dystrophy, LAMA2-related	AR	General Population Caucasian / European Population	<1 in 500 1 in 125	99% 99%		<1 in 10 million 1 in 6,200,500
LAMA3	Junctional epidermolysis bullosa, LAMA3-related	AR	General Population	1 in 781	98%		<1 in 10 million
LAMA3	Laryngo-onycho-cutaneous syndrome	AR	General Population	1 in 781	98%	,	<1 in 10 million
LAMB3	Junctional epidermolysis bullosa, LAMB3-related	AR	General Population	1 in 781	98%		<1 in 10 million
LAMC2	Junctional epidermolysis bullosa, LAMC2-related	AR	General Population	1 in 781	98%		<1 in 10 million
LIPA	Lysosomal acid lipase deficiency	AR	General Population Caucasian / European Population Iranian Jewish Population	<1 in 500 1 in 112 1 in 26	99% 99% 99%	,	<1 in 10 million 1 in 4,973,248 1 in 260,104
LRPPRC	Leigh syndrome with Complex IV deficiency	AR	General Population Faroese Population French Canadian Population	1 in 447 1 in 21 1 in 22	98% 98% 98%	1 in 22,301 1 in 1,001 1 in 1,051	<1 in 10 million 1 in 84,084 1 in 92,488
MAN2B1	Alpha-Mannosidosis	AR	General Population Caucasian / European Population	1 in 354 1 in 274	99% 99%	1 in 35,301 1 in 27,301	<1 in 10 million <1 in 10 million
MCOLN1	Mucolipidosis IV	AR	General Population Ashkenazi Jewish Population	1 in 300 1 in 100	99% 99%	1 in 29,901 1 in 9,901	<1 in 10 million 1 in 3,960,400
MEFV	Familial Mediterranean fever	AR	General Population Mediterranean Population	1 in 20 1 in 7	99% 90%	1 in 1,901 1 in 61	1 in 152,080 1 in 1,708
MESP2	Spondylocostal dysostosis	AR	General Population	<1 in 500	98%	1 in 24,951	<1 in 10 million
MKS1	Bardet-Biedl syndrome 13	AR	General Population Finnish Population	1 in 260 1 in 47	98% 98%	1 in 12,951 1 in 2,301	<1 in 10 million 1 in 432,588
MKS1	Joubert syndrome 28	AR	General Population Finnish Population	1 in 260 1 in 47	98% 98%	1 in 12,951 1 in 2,301	<1 in 10 million 1 in 432,588
MKS1	Meckel syndrome 1	AR	General Population Finnish Population	1 in 260 1 in 47	98% 98%	1 in 12,951 1 in 2,301	<1 in 10 million 1 in 432,588
MLC1	Megalencephalic leukoencephalopathy with subcortical cysts	AR	General Population Libyan Jewish Population	<1 in 500 1 in 40	99% 99%	1 in 49,901 1 in 3,901	<1 in 10 million 1 in 624,160
MMAA	Methylmalonic aciduria, cbIA type	AR	General Population	1 in 301	97%	1 in 10,001	<1 in 10 million
MMAB	Methylmalonic aciduria, cblB type	AR	General Population	1 in 435	98%	1 in 21,701	<1 in 10 million
MMACHC	Methylmalonic aciduria and homocystinuria, cbIC type	AR	General Population	1 in 134	90%	1 in 1,331	1 in 713,416
MPI	Congenital disorder of glycosylation type lb	AR	General Population	<1 in 500			<1 in 10 million
MUT	Methylmalonic acidemia, MUT-related	AR	General Population East Asian Population Middle-Eastern Population	1 in 195 1 in 53 1 in 52	96% 96% 96%	1 in 4,851 1 in 1,301 1 in 1,276	1 in 3,783,780 1 in 275,812 1 in 265,408
MUT	Methylmalonic aciduria-methylmalonyl-CoA mutase deficiency	AR	General Population	1 in 100	99%	1 in 9,901	1 in 3,960,400
MYO7A	Usher syndrome, type 1B	AR	General Population East Asian Population	1 in 206 1 in 62	98% 98%	1 in 10,251 1 in 3,051	1 in 8,446,824 1 in 756,648
MYO7A	Non-syndromic hearing loss, MYO7A-related	AR	General Population East Asian Population	1 in 206 1 in 62	98% 98%	1 in 10,251 1 in 3,051	1 in 8,446,824 1 in 756,648
NAGLU	Mucopolysaccharidosis type IIIB (Sanfilippo syndrome B)	AR	General Population Caucasian / European Population East Asian Population	<1 in 500 1 in 346 1 in 298	99% 99% 99%	1 in 34,501	<1 in 10 million <1 in 10 million <1 in 10 million
NBN	Nijmegen breakage syndrome	AR	General Population	1 in 158	99%	1 in 15,701	1 in 9,923,032
NEB	Nemaline myopathy	AR	General Population Amish Population Ashkenazi Jewish Population Finnish Population	1 in 112 1 in 11 1 in 108 1 in 112	98% 98% 98% 98%	1 in 5,551 1 in 501 1 in 5,351 1 in 5,551	1 in 2,486,848 1 in 22,044 1 in 2,311,632 1 in 2,486,848
NPC1	Niemann-Pick disease, type C1	AR	General Population	1 in 194	90%	1 in 1,931	1 in 1,498,456
NPC2	Niemann-Pick disease, type C2	AR	General Population	1 in 194	99%	1 in 19,301	<1 in 10 million
NPHS1	Congenital nephrotic syndrome, type 1	AR	General Population Finnish Population	1 in 289 1 in 50	98% 98%	1 in 14,401 1 in 2,451	<1 in 10 million 1 in 490,200
NPHS2	Congenital nephrotic syndrome, type 2	AR	General Population Finnish Population	1 in 289 1 in 50	98% 98%	1 in 14,401 1 in 2,451	<1 in 10 million 1 in 490,200
OPA3	Costeff syndrome	AR	General Population Iraqi Jewish Population	<1 in 500 1 in 50	98% 98%	1 in 24,951 1 in 2,451	<1 in 10 million 1 in 490,200

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Supplemental Table							
Gene	Condition	Inheritance	Ethnicity	Carrier Rate	Detection Rate	Post-test Carrier Probability*	Residual Risk*
SLC12A6	Andermann syndrome	AR	General Population French Canadian Population	<1 in 500 1 in 23	98% 99%	1 in 24,951 1 in 2,201	<1 in 10 million 1 in 202,492
SLC17A5	Sialic acid storage disorder	AR	General Population Finnish Population	<1 in 500 1 in 100	91% 91%	1 in 5,545 1 in 1,101	<1 in 10 million 1 in 440,400
SLC22A5	Systemic primary carnitine deficiency	AR	General Population African/African American Population East Asian Population Faroese Population Pacific Islander Population South Asian/Indian Population	1 in 129 1 in 86 1 in 77 1 in 9 1 in 37 1 in 51	99% 99% 99% 99% 99%	1 in 12,801 1 in 8,501 1 in 7,601 1 in 801 1 in 3,601 1 in 5,001	1 in 6,605,316 1 in 2,924,344 1 in 2,341,108 1 in 28,836 1 in 532,948 1 in 1,020,204
SLC26A2	Diastrophic dysplasia	AR	General Population Finnish Population	1 in 158 1 in 50	90% 90%	1 in 1,571 1 in 491	1 in 992,872 1 in 98,200
SLC26A2	Achondrogenesis, type IB	AR	General Population Finnish Population	1 in 158 1 in 50	90% 90%	1 in 1,571 1 in 491	1 in 992,872 1 in 98,200
SLC26A2	Multiple epiphyseal dysplasia	AR	General Population Finnish Population	1 in 158 1 in 50	90% 90%	1 in 1,571 1 in 491	1 in 992,872 1 in 98,200
SLC26A2	Atelosteogenesis II	AR	General Population Finnish Population	1 in 158 1 in 50	90% 90%	1 in 1,571 1 in 491	1 in 992,872 1 in 98,200
SLC26A4	Pendred syndrome	AR	General Population African/African American Population Caucasian / European Population East Asian Population	1 in 80 1 in 76 1 in 88 1 in 74	98% 98% 98% 98%	1 in 3,951 1 in 3,751 1 in 4,351 1 in 3,651	1 in 1,264,320 1 in 1,140,304 1 in 1,531,552 1 in 1,080,696
SLC37A4	Glycogen storage disease, type lb	AR	General Population Ashkenazi Jewish Population	1 in 158 1 in 71	95% 95%	1 in 3,141 1 in 1,401	1 in 1,985,112 1 in 397,884
SMN1	Spinal muscular atrophy	AR	General Population African/African American Population Ashkenazi Jewish Population Caucasian / European Population East Asian Population Latino Population Sephardic Jewish Population	1 in 54 1 in 72 1 in 67 1 in 47 1 in 59 1 in 68 1 in 34	91% 71% 95% 93% 90% 96%	1 in 590 1 in 246 1 in 734 1 in 921 1 in 830 1 in 671 1 in 826	1 in 127,440 1 in 70,848 1 in 196,712 1 in 173,148 1 in 195,880 1 in 182,512 1 in 112,336
SMPD1	Niemann-Pick disease, type A/B	AR	General Population Ashkenazi Jewish Population Latino Population	1 in 250 1 in 115 1 in 106	95% 95% 95%	1 in 4,981 1 in 2,281 1 in 2,101	1 in 4,981,000 1 in 1,049,260 1 in 890,824
STAR	Lipoid congenital adrenal hyperplasia	AR	General Population	<1 in 500	98%	1 in 24,951	<1 in 10 million
TAT	Tyrosinemia, type II	AR	General Population	1 in 250	98%	1 in 12,451	<1 in 10 million
TCIRG1	Osteopetrosis, TCIRG1-related	AR	General Population	1 in 250	98%	1 in 12,451	<1 in 10 million
TGM1	Congenital ichthyosis	AR	General Population	1 in 224	95%	1 in 4,461	1 in 3,997,056
TH	Segawa syndrome	AR	General Population	1 in 224	98%		1 in 9,991,296
TMEM216	Joubert syndrome 2	AR	General Population Ashkenazi Jewish Population	1 in 141 1 in 92	98% 98%	1 in 7,001 1 in 4,551	1 in 3,948,564 1 in 1,674,768
TMEM216	Meckel syndrome 2	AR	General Population Ashkenazi Jewish Population	1 in 141 1 in 92	98% 98%	1 in 7,001 1 in 4,551	1 in 3,948,564 1 in 1,674,768
TPP1	Neuronal ceroid lipofuscinosis, TPP1-related	AR	General Population French Canadian Population	1 in 252 1 in 53	97% 97%	1 in 8,368 1 in 1,734	1 in 8,434,944 1 in 367,608
TTPA	Ataxia with isolated vitamin E deficiency	AR	General Population Caucasian / European Population	<1 in 500 1 in 267	98% 90%	1 in 24,951 1 in 2,661	<1 in 10 million 1 in 2,841,948
USH1C	Usher syndrome, type IC	AR	General Population French Canadian Population	1 in 353 1 in 227	90% 90%	1 in 3,521 1 in 2,261	1 in 4,971,652 1 in 2,052,988
USH1C	Non-syndromic hearing loss, USH1C-related	AR	General Population French Canadian Population	1 in 353 1 in 227	90% 90%	1 in 3,521 1 in 2,261	1 in 4,971,652 1 in 2,052,988
USH2A	Usher syndrome, type 2A	AR	General Population Caucasian / European Population Ashkenazi Jewish Population Iranian Jewish Population	1 in 126 1 in 73 1 in 35 1 in 60	96% 96% 99% 99%	1 in 3,126 1 in 1,801 1 in 3,401 1 in 5,901	1 in 1,575,504 1 in 525,892 1 in 476,140 1 in 1,416,240
VPS13B	Cohen syndrome	AR	General Population	<1 in 500	98%	1 in 24,951	<1 in 10 million
XPA	Xeroderma pigmentosum, group A	AR	General Population Japanese Population	1 in 500 1 in 74	99% 99%	1 in 49,901 1 in 7,301	<1 in 10 million 1 in 2,161,096
XPC	Xeroderma pigmentosum, group C	AR	General Population	1 in 500	99%	1 in 49,901	<1 in 10 million
ZFYVE26	Spastic paraplegia 15	AR	General Population	<1 in 500	98%	1 in 24,951	<1 in 10 million

* For genes that have tested negative

† The carrier frequency for alpha thalassemia trait cis is described in rows marked with a dagger symbol.

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Abbreviations: AR, autosomal recessive; XL, X-linked