



DOB: Sam	ple type: Blood ple collection date: 06-JUN ple accession date: 07-JUN	N-2023 Invitae #:	15-JUN-2023 RQ5172543 Hallie Yoshimura Jeffrey Olliffe
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Reason for testing

Gamete donor

Test performed

Invitae Carrier Screen



RESULT: POSITIVE

This carrier test evaluated 514 gene(s) for genetic changes (variants) that are associated with an increased risk of having a child with a genetic condition. Knowledge of carrier status for one of these conditions may provide information that can be used to assist with family planning and/or preparation. Carrier screening is not intended for diagnostic purposes. To identify a potential genetic basis for a condition in the individual being tested, diagnostic testing for the gene(s) of interest is recommended.

This test shows the presence of clinically significant genetic change(s) in this individual in the gene(s) indicated below. No other clinically significant changes were identified in the remaining genes evaluated with this test.

RESULTS	GENE	VARIANT(S)	INHERITANCE	PARTNER TESTING RECOMMENDED
Carrier: Adenosine deaminase deficiency	ADA	c.715G>A (p.Gly239Ser)	Autosomal recessive	Yes
Carrier: Gitelman syndrome	SLC12A3	c.539C>A (p.Thr180Lys)	Autosomal recessive	Yes
Carrier: Oculocutaneous albinism type 2	OCA2	c.1160C>T (p.Thr387Met)	Autosomal recessive	Yes
Carrier: PCDH15-related conditions	PCDH15	c.4548_4551dup (p.Asp15181lefs*2)	Autosomal recessive	Yes
Carrier: Tyrosine hydroxylase deficiency	ТН	c.1495G>A (p.Val499Met)	Autosomal recessive	Yes

Next steps

- See the table above for recommendations regarding testing of this individual's reproductive partner.
- Even for genes that have a negative test result, there is always a small risk that an individual could still be a carrier. This is called "residual risk." See the Carrier detection rates and residual risks document.
- Discussion with a physician and/or genetic counselor is recommended to further review the implications of this test result and to understand these results in the context of any family history of a genetic condition.
- All patients, regardless of result, may wish to consider additional screening for hemoglobinopathies by complete blood count (CBC) and hemoglobin electrophoresis, if this has not already been completed.
- Individuals can register their tests at https://www.invitae.com/patients/ to access online results, educational resources, and next steps.





DOB

Patient name: DONOR 14527 Invitae #: RQ5172543

Clinical summary

RESULT: CARRIER

Adenosine deaminase deficiency

A single Likely Pathogenic variant, c.715G>A (p.Gly239Ser), was identified in ADA.

What is adenosine deaminase deficiency?

Adenosine deaminase (ADA) deficiency is a condition that affects the immune system and causes recurrent infections. Affected individuals have a reduced amount of the enzyme ADA, which leads to a deficiency of certain infection-fighting white blood cells (lymphocytes). The majority of individuals with ADA will develop severe combined immunodeficiency (SCID), which is typically diagnosed by 6 to 12 months of age. Due to their reduced immune system response, these individuals are prone to repeated infections, which can be life-threatening. Additional symptoms may include poor growth (failure to thrive), chronic diarrhea, and skin rashes. Children with SCID whose immune function is not restored rarely survive beyond 2 years of age. Less commonly, the onset of immune deficiency in individuals with ADA may be delayed or may occur in adulthood, with less severe symptoms. Partial ADA deficiency is associated with reduced ADA enzyme activity, but normal or near-normal immune function.

Treatment may include a bone marrow or stem cell transplant from an HLA-matched individual and/or enzyme replacement therapy. Follow-up depends on each affected individual's specific situation, and discussion with a healthcare provider should be considered.

Next steps

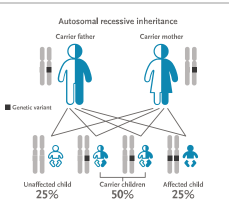
Carrier testing for the reproductive partner is recommended.

(+) If your partner tests positive:

In autosomal recessive inheritance, an individual must have disease-causing genetic changes in each copy of the ADA gene to be affected. Carriers, who have a disease-causing genetic change in only one copy of the gene, typically do not have symptoms. When both reproductive partners are carriers of an autosomal recessive condition, there is a 25% chance for each child to have the condition.

If your partner tests negative:

A negative carrier test result reduces, but does not eliminate, the chance that a person may be a carrier. The risk that a person could still be a carrier, even after a negative test result, is called a residual risk. See the table below for your partner's hypothetical



residual risk after testing negative for adenosine deaminase deficiency. These values are provided only as a guide, are based on the detection rate for the condition as tested at Invitae, and assume a negative family history, the absence of symptoms, and vary based on the ethnic background of an individual. For genes associated with both dominant and recessive inheritance, the numbers provided apply to the recessive condition(s) associated with the gene.

DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY BEFORE SCREENING	CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
Adenosine deaminase deficiency (AR) NM_000022.2	ADA	Pan-ethnic	1 in 224	1 in 2788





) INVITAE CARRIER SCREEN RESULTS

DOB:

Patient name: DONOR 14527 Invitae #: RQ5172543

RESULT: CARRIER

Gitelman syndrome

A single Pathogenic variant, c.539C>A (p.Thr180Lys), was identified in SLC12A3.

What is Gitelman syndrome?

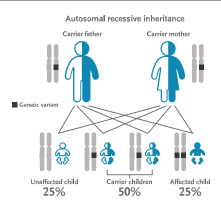
Gitelman syndrome is a condition that affects kidney function. Affected individuals have difficulty reabsorbing salt from urine back into the blood, causing low levels of potassium (hypokalemia), magnesium (hypomagnesemia), and calcium (hypocalciuria). Symptoms usually appear in late childhood or adolescence and may include intermittent muscle spasms (tetany), muscle weakness or cramps, joint pain caused by calcium crystals forming in the cartilage (chondrocalcinosis), fatigue, excessive thirst (polydipsia), excessive urination (polyuria), tingling or prickly sensation in the skin caused by pressure on, or damage to, peripheral nerves (paresthesia) and a sensation of rapid heart beats (palpitations). Less commonly, affected individuals have short stature, seizures, abnormally high or low thyroid hormone levels (hyperthyroidism or hypothyroidism, respectively), or an abnormal heart rhythm (arrhythmia). Gitelman syndrome is a highly variable condition; symptoms can vary widely, even among affected members of the same family, and in some cases there may only be mild symptoms or no overt symptoms of the condition (asymptomatic). With treatment and dietary management, affected individuals have a normal life expectancy.

Next steps

Carrier testing for the reproductive partner is recommended.

(+) If your partner tests positive:

In autosomal recessive inheritance, an individual must have disease-causing genetic changes in each copy of the SLC12A3 gene to be affected. Carriers, who have a disease-causing genetic change in only one copy of the gene, typically do not have symptoms. When both reproductive partners are carriers of an autosomal recessive condition, there is a 25% chance for each child to have the condition.



) If your partner tests negative:

A negative carrier test result reduces, but does not eliminate, the chance that a person may be a carrier. The risk that a person could still be a carrier, even after a negative test result, is called a residual risk. See the table below for your partner's hypothetical

residual risk after testing negative for Gitelman syndrome. These values are provided only as a guide, are based on the detection rate for the condition as tested at Invitae, and assume a negative family history, the absence of symptoms, and vary based on the ethnic background of an individual. For genes associated with both dominant and recessive inheritance, the numbers provided apply to the recessive condition(s) associated with the gene.

DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY BEFORE SCREENING	CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
Gitelman syndrome (AR) NM_000339.2	SLC12A3	Pan-ethnic	1 in 100	1 in 9900





DOB

Patient name: DONOR 14527 Invitae #: RQ5172543

RESULT: CARRIER

Oculocutaneous albinism type 2

A single Pathogenic variant, c.1160C>T (p.Thr387Met), was identified in OCA2.

What is oculocutaneous albinism type 2?

Oculocutaneous albinism (OCA) is a condition that causes decreased color (hypopigmentation) of the hair, skin, and eyes. Affected individuals produce a reduced amount of melanin, the pigment that gives skin, hair, and eyes their color, resulting in hypopigmentation. Additional symptoms of OCA include reduced visual acuity (farsightedness or nearsightedness), increased sensitivity to light (photophobia), involuntary eye movements (nystagmus), and eyes that do not look in the same direction (strabismus). Other eye findings, such as reduced pigmentation of the light-sensitive tissue that lines the back of the eye (retina) and misrouting of the nerves of the eye (optic nerves), are seen on ophthalmologic exam. Individuals with fair complexions have an increased risk for skin cancers. Intelligence is not typically affected. Treatment is aimed at correcting vision and providing visual aids, or other visual resources. Sun protection is essential due to the increased risk for skin cancer.

Next steps

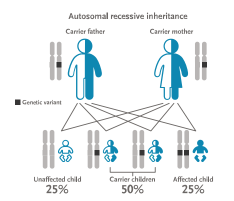
Carrier testing for the reproductive partner is recommended.

+) If your partner tests positive:

In autosomal recessive inheritance, an individual must have disease-causing genetic changes in each copy of the OCA2 gene to be affected. Carriers, who have a disease-causing genetic change in only one copy of the gene, typically do not have symptoms. When both reproductive partners are carriers of an autosomal recessive condition, there is a 25% chance for each child to have the condition.

(-) If your partner tests negative:

A negative carrier test result reduces, but does not eliminate, the chance that a person may be a carrier. The risk that a person could still be a carrier, even after a negative test result, is called a residual risk. See the table below for your partner's hypothetical



residual risk after testing negative for oculocutaneous albinism type 2. These values are provided only as a guide, are based on the detection rate for the condition as tested at Invitae, and assume a negative family history, the absence of symptoms, and vary based on the ethnic background of an individual. For genes associated with both dominant and recessive inheritance, the numbers provided apply to the recessive condition(s) associated with the gene.

DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY BEFORE SCREENING	CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
Oculocutaneous albinism type 2 (AR) NM_000275.2	OCA2	Pan-ethnic	1 in 95	1 in 9400





) INVITAE CARRIER SCREEN RESULTS

DOB:

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RESULT: CARRIER

PCDH15-related conditions

A single Pathogenic variant, c.4548_4551dup (p.Asp1518llefs*2), was identified in PCDH15.

What are PCDH15-related conditions?

PCDH15-related conditions include Usher syndrome type IF (USH1F) and autosomal recessive nonsyndromic deafness (DFNB23). Usher syndrome is a group of related conditions that causes deafness, progressive vision loss due to an eye disease called retinitis pigmentosa (RP), and, in certain forms, balance difficulties due to inner ear problems (vestibular dysfunction). Autosomal recessive nonsyndromic deafness is a group of related conditions that affects an individual's ability to hear.

Individuals with USH1F are usually born with severe to profound deafness. Balance issues may delay meeting developmental milestones such as independent sitting and walking. Progressive vision loss due to RP typically begins during childhood or adolescence; however, complete blindness is uncommon. Severity of symptoms can vary, even between family members with the same genetic change. Digenic inheritance, which occurs when an individual has a genetic change in two different Usher syndrome-associated genes, has been reported (PMID: 15537665); however, the evidence available at this time is insufficient to confirm this as a mode of inheritance.

Individuals with nonsyndromic deafness are born with mild to profound deafness that typically does not worsen over time. Nonsyndromic deafness does not affect any other part of the body. Severity of deafness may vary, even among members of the same family. Intellect and life span are not impacted.

For PCDH15-related conditions, early initiation of medical, educational, and social services is recommended to maximize outcomes.

Next steps

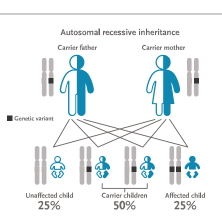
Carrier testing for the reproductive partner is recommended.

(+) If your partner tests positive:

In autosomal recessive inheritance, an individual must have disease-causing genetic changes in each copy of the PCDH15 gene to be affected. Carriers, who have a disease-causing genetic change in only one copy of the gene, typically do not have symptoms. When both reproductive partners are carriers of an autosomal recessive condition, there is a 25% chance for each child to have the condition.

If your partner tests negative:

A negative carrier test result reduces, but does not eliminate, the chance that a person may be a carrier. The risk that a person could still be a carrier, even after a negative test result, is called a residual risk. See the table below for your partner's hypothetical



residual risk after testing negative for PCDH15-related conditions. These values are provided only as a guide, are based on the detection rate for the condition as tested at Invitae, and assume a negative family history, the absence of symptoms, and vary based on the ethnic background of an individual. For genes associated with both dominant and recessive inheritance, the numbers provided apply to the recessive condition(s) associated with the gene.

DISORDER (INHERITANCE)	GENE	ETHNICITY		CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
PCDH15-related conditions (AR) NM_033056.3	PCDH15	Pan-ethnic	1 in 400	1 in 39900





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RESULT: CARRIER

Tyrosine hydroxylase deficiency

A single Pathogenic variant, c.1495G>A (p.Val499Met), was identified in TH.

What is tyrosine hydroxylase deficiency?

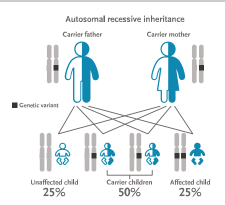
Tyrosine hydroxylase (TH) deficiency refers to a spectrum of conditions that affect the nervous system. TH deficiency, also known as autosomal recessive Segawa syndrome, has two main forms which are differentiated by the degree of severity of symptoms and response to treatment. Affected individuals have reduced levels of dopamine and other hormones that are involved in the functioning of the nervous system. The less severe type of TH deficiency is characterized by infantile or childhood onset progressive loss of muscle movement (hypokinesia), parkinsonian features such as muscle rigidity or tremors, and involuntary muscle tensing (dystonia). Individuals with the less severe form of the condition typically have good response to the medication levodopa. In the more severe type of TH deficiency, symptoms typically appear shortly after birth and include low muscle tone (hypotonia), complex brain dysfunction (encephalopathy) leading to developmental delay and intellectual disability, parkinsonian features, and disturbances of the autonomic nervous system, which controls involuntary body processes such as heart and breathing rates, blood pressure, digestion, and body temperature. These symptoms can be life-threatening. Individuals with the more severe form of TH deficiency have a lesser response to levodopa. Prognosis depends on the severity of symptoms. Follow-up depends on each affected individual's specific situation, and discussion with a healthcare provider should be considered.

Next steps

Carrier testing for the reproductive partner is recommended.

(+) If your partner tests positive:

In autosomal recessive inheritance, an individual must have disease-causing genetic changes in each copy of the TH gene to be affected. Carriers, who have a disease-causing genetic change in only one copy of the gene, typically do not have symptoms. When both reproductive partners are carriers of an autosomal recessive condition, there is a 25% chance for each child to have the condition.



•) If your partner tests negative:

A negative carrier test result reduces, but does not eliminate, the chance that a person may be a carrier. The risk that a person could still be a carrier, even after a negative test result, is called a residual risk. See the table below for your partner's hypothetical

residual risk after testing negative for tyrosine hydroxylase deficiency. These values are provided only as a guide, are based on the detection rate for the condition as tested at Invitae, and assume a negative family history, the absence of symptoms, and vary based on the ethnic background of an individual. For genes associated with both dominant and recessive inheritance, the numbers provided apply to the recessive condition(s) associated with the gene.

DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY BEFORE SCREENING	CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
Tyrosine hydroxylase deficiency (AR) NM_199292.2	тн	Pan-ethnic	≤1 in 500	Reduced



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Results to note

GALT

- c.-119_-116del (Non-coding) was identified in the GALT gene.
- This benign variant is not known to cause disease and does not impact this individual's risk to be a carrier for galactosemia (GALT-related). Carrier testing for the reproductive partner is not indicated based on this result. See Variant details for more information.

HBA1/HBA2

- Additional copy(ies) detected (HBA1 copy number=3). Result not associated with alpha-thalassemia.
- Additional copy(ies) of the alpha-globin gene (HBA1/HBA2) do not impact alpha-thalassemia risk, however, co-inheritance of this genetic change with beta-thalassemia may worsen the clinical and hematological features of the latter condition. There is also a possibility for hematological changes or clinical symptoms in beta-thalassemia carriers when this alpha-globin gene change is present. Carrier testing of the reproductive partner for beta-thalassemia (HBB) may be considered.

SMN1

Negative result. SMN1: 2 copies; c.*3+80T>G not detected.

Variant details

ADA, Exon 8, c.715G>A (p.Gly239Ser), heterozygous, Likely Pathogenic

- This sequence change replaces glycine, which is neutral and non-polar, with serine, which is neutral and polar, at codon 239 of the ADA protein (p.Gly239Ser).
- This variant is present in population databases (rs777820729, gnomAD 0.004%).
- This missense change has been observed in individual(s) with ADA-related conditions (PMID: 21410451).
- ClinVar contains an entry for this variant (Variation ID: 338506).
- Advanced modeling of protein sequence and biophysical properties (such as structural, functional, and spatial information, amino acid conservation, physicochemical variation, residue mobility, and thermodynamic stability) performed at Invitae indicates that this missense variant is expected to disrupt ADA protein function.
- Experimental studies have shown that this missense change affects ADA function (PMID: 11160213).
- This variant disrupts the p.Gly239 amino acid residue in ADA. Other variant(s) that disrupt this residue have been determined to be pathogenic (PMID: 26255240, 30858051). This suggests that this residue is clinically significant, and that variants that disrupt this residue are likely to be disease-causing.
- In summary, the currently available evidence indicates that the variant is pathogenic, but additional data are needed to prove that conclusively. Therefore, this variant has been classified as Likely Pathogenic.

GALT, Exon 1, c.-119_-116del (Non-coding), heterozygous, Benign (reportable variant)

- This variant occurs in a non-coding region of the GALT gene. It does not change the encoded amino acid sequence of the GALT protein. This variant is unique to the D2 allele and is also known as the Duarte variant.
- This variant is present in population databases (rs142496102, gnomAD 8%), including at least one homozygous and/or hemizygous individual. The c.-119_-116del variant is the most common galactosemia variant (PMID: 19904210).
- Compound heterozygosity for the Duarte allele and a pathogenic galactosemia variant (termed Duarte variant Galactosemia, DG) results in approximately 14-25% of normal GALT enzyme activity (PMID: 25473725, 25681083) and causes elevations of the metabolites found in galactosemia. DG may trigger a positive galactosemia newborn screening or abnormal biochemical test results but does not require dietary intervention (PMID: 30593450, 30593448) and does not cause the significant clinical consequences of classic galactosemia (PMID: 30593450,





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31160755). A homozygous c.-119_-116del variant alone (DD) can have mildly reduced GALT enzyme activity but is insufficient to cause metabolite accumulation and is considered clinically benign (PMID: 24718839, 25473725).

- ClinVar contains two entries for this variant (Variation ID: 140570, 25111).
- Algorithms developed to predict the effect of variants on protein structure and function are not available or were not evaluated for this variant.
- Experimental studies have shown that this variant affects GALT enzyme activity (PMID: 11286503, 11479743, 19224951).
- For these reasons, this variant has been classified as a Benign reportable variant.

OCA2, Exon 11, c.1160C>T (p.Thr387Met), heterozygous, PATHOGENIC

- This sequence change replaces threonine, which is neutral and polar, with methionine, which is neutral and non-polar, at codon 387 of the OCA2 protein (p.Thr387Met).
- This variant is present in population databases (rs150335311, gnomAD 0.08%).
- This missense change has been observed in individual(s) with clinical features of oculocutaneous albinism (PMID: 29345414; Invitae).
- ClinVar contains an entry for this variant (Variation ID: 888403).
- Advanced modeling of protein sequence and biophysical properties (such as structural, functional, and spatial information, amino acid conservation, physicochemical variation, residue mobility, and thermodynamic stability) has been performed at Invitae for this missense variant, however the output from this modeling did not meet the statistical confidence thresholds required to predict the impact of this variant on OCA2 protein function.
- For these reasons, this variant has been classified as Pathogenic.

PCDH15, Exon 33, c.4548_4551dup (p.Asp1518llefs*2), heterozygous, PATHOGENIC

- This sequence change creates a premature translational stop signal (p.Asp1518Ilefs*2) in the PCDH15 gene. While this is not anticipated to result in nonsense mediated decay, it is expected to disrupt the last 438 amino acid(s) of the PCDH15 protein.
- This variant is present in population databases (no rsID available, gnomAD 0.0009%).
- This variant has not been reported in the literature in individuals affected with PCDH15-related conditions.
- ClinVar contains an entry for this variant (Variation ID: 847128).
- This variant disrupts a region of the PCDH15 protein in which other variant(s) (p.Gln1576*) have been determined to be pathogenic (PMID: 28281779). This suggests that this is a clinically significant region of the protein, and that variants that disrupt it are likely to be disease-causing.
- For these reasons, this variant has been classified as Pathogenic.

SLC12A3, Exon 4, c.539C>A (p.Thr180Lys), heterozygous, PATHOGENIC

- This sequence change replaces threonine, which is neutral and polar, with lysine, which is basic and polar, at codon 180 of the SLC12A3 protein (p.Thr180Lys).
- This variant is present in population databases (rs146158333, gnomAD 0.4%).
- This missense change has been observed in individual(s) with Gitelman syndrome (PMID: 10616841, 21628937, 21757836, 26041598). In at least one individual the data is consistent with being in trans (on the opposite chromosome) from a pathogenic variant. It has also been observed to segregate with disease in related individuals.
- ClinVar contains an entry for this variant (Variation ID: 648571).
- Advanced modeling of protein sequence and biophysical properties (such as structural, functional, and spatial information, amino acid conservation, physicochemical variation, residue mobility, and thermodynamic stability) has been performed at Invitae for this missense variant, however the output from this modeling did not meet the statistical confidence thresholds required to predict the impact of this variant on SLC12A3 protein function.
- For these reasons, this variant has been classified as Pathogenic.

TH, Exon 14, c.1495G>A (p.Val499Met), heterozygous, PATHOGENIC

- This sequence change replaces valine, which is neutral and non-polar, with methionine, which is neutral and non-polar, at codon 499 of the TH protein (p.Val499Met).
- This variant is present in population databases (rs1800033, gnomAD 0.1%).





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- This missense change has been observed in individuals with clinical features of TH-related conditions and/or TH-deficient dopa-responsive dystonia (PMID: 29724574, 33072517, 34054692). It has also been observed to segregate with disease in related individuals.
- This variant is also known as p.Val468Met.
- ClinVar contains an entry for this variant (Variation ID: 412030).
- Advanced modeling of protein sequence and biophysical properties (such as structural, functional, and spatial information, amino acid conservation, physicochemical variation, residue mobility, and thermodynamic stability) has been performed at Invitae for this missense variant, however the output from this modeling did not meet the statistical confidence thresholds required to predict the impact of this variant on TH protein function.
- For these reasons, this variant has been classified as Pathogenic.

Residual risk

No carrier test can detect 100% of carriers. There still remains a small risk of being a carrier after a negative test (residual risk). Residual risk values assume a negative family history and are inferred from published carrier frequencies and estimated detection rates based on testing technologies used at Invitae. You can view Invitae's complete Carrier detection rates and residual risks document (containing all carrier genes) online at https://www.invitae.com/carrier-residual-risks/. Additionally, the order-specific information for this report is available to download in the portal (under this order's documents) or can be requested by contacting Invitae Client Services. The complete Carrier detection rates and residual risks document will not be applicable for any genes with specimen-specific limitations in sequencing and/or deletion/duplication coverage. Please see the final bullet point in the Limitations section of this report to view if this specimen had any gene-specific coverage gaps.





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Genes analyzed

This table represents a complete list of genes analyzed for this individual, including the relevant gene transcript(s). If more than one transcript is listed for a single gene, variants were reported using the first transcript listed unless otherwise indicated in the report. An asterisk (*) indicates that this gene has a limitation. Please see the Limitations section for details. Results are negative, unless otherwise indicated in the report.

GENE	TRANSCRIPT	GENE	TRANSCRIPT	GENE	TRANSCRIPT
AAAS	NM_015665.5	AP1S1	NM_001283.3	CBS	NM_000071.2
ABCA12	NM_173076.2	AQP2	NM_000486.5	CC2D1A	NM_017721.5
ABCA3	NM_001089.2	ARG1	NM_000045.3	CC2D2A	NM_001080522.2
ABCA4	NM_000350.2	ARL6	NM_177976.2	CCDC103	NM_213607.2
ABCB11	NM_003742.2	ARSA	NM_000487.5	CCDC39	NM_181426.1
ABCB4	NM_000443.3	ARSB	NM_000046.3	CCDC88C	NM_001080414.3
ABCC2*	NM_000392.4	ASL	NM_000048.3	CD3D	NM_000732.4
ABCC8	NM_000352.4	ASNS	NM_133436.3	CD3E	NM_000733.3
ACAD9	NM_014049.4	ASPA	NM_000049.2	CD40	NM_001250.5
ACADM	NM_000016.5	ASS1	NM_000050.4	CD59	NM_203330.2
ACADVL	NM_000018.3	ATM*	NM_000051.3	CDH23	NM_022124.5
ACAT1	NM_000019.3	ATP6V1B1	NM_001692.3	CEP152	NM_014985.3
ACOX1	NM_004035.6	ATP7B	NM_000053.3	CEP290	NM_025114.3
ACSF3	NM_174917.4	ATP8B1*	NM_005603.4	CERKL	NM_001030311.2
ADA	NM_000022.2	BBS1	NM_024649.4	CFTR*	NM_000492.3
ADAMTS2	NM_014244.4	BBS10	NM_024685.3	CHAT	NM_020549.4
ADAMTSL4	NM_019032.5	BBS12	NM_152618.2	CHRNE	NM_000080.3
ADGRG1	NM_005682.6	BBS2	NM_031885.3	CHRNG	NM_005199.4
ADGRV1	NM_032119.3	BBS4	NM_033028.4	CIITA	NM_000246.3
AGA	NM_000027.3	BBS5	NM_152384.2	CLCN1	NM_000083.2
AGL	NM_000642.2	BBS7	NM_176824.2	CLN3	NM_001042432.1
AGPS	NM_003659.3	BBS9*	NM_198428.2	CLN5	NM_006493.2
AGXT	NM_000030.2	BCKDHA	NM_000709.3	CLN6	NM_017882.2
AHII	NM_017651.4	BCKDHB	NM_183050.2	CLN8	NM_018941.3
AIPL1*	NM_014336.4	BCS1L	NM_004328.4	CLRN1	NM_174878.2
AIRE	NM_000383.3	BLM	NM_000057.3	CNGB3	NM_019098.4
ALDH3A2	NM_000382.2	BLOC1S3	NM_212550.4	COL11A2*	NM_080680.2
ALDH7A1	NM_001182.4	BLOC1S6	NM_012388.3	COL17A1	NM_000494.3
ALDOB	NM_000035.3	BMP1	NM_006129.4;NM_001199.3	COL27A1	NM_032888.3
ALG1	NM_019109.4	BRIP1	NM_032043.2	COL4A3	NM_000091.4
ALG6	NM_013339.3	BSND	NM_057176.2	COL4A4	NM_000092.4
ALMS1	NM_015120.4	BTD	NM_000060.3	COL7A1	NM_000094.3
ALPL	NM_000478.5	CAD	NM_004341.4	COX15	NM_004376.6
AMN*	NM_030943.3	CANT1	NM_138793.3	CPS1	NM_001875.4
AMT	NM_000481.3	CAPN3	NM_000070.2	CPT1A	NM_001876.3
ANO10*	NM_018075.3	CASQ2	NM_001232.3	CPT2	NM_000098.2





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GENE	TRANSCRIPT	GENE	TRANSCRIPT	GENE	TRANSCRIPT
CRB1	NM_201253.2	EIF2B1	NM_001414.3	FUCA1	NM_000147.4
CRTAP	NM_006371.4	EIF2B2	NM_014239.3	G6PC	NM_000151.3
CTNS	NM_004937.2	EIF2B3	NM_020365.4	G6PC3	NM_138387.3
CTSA	NM_000308.3	EIF2B4	NM_015636.3	GAA	NM_000152.3
стѕс	NM_001814.5	EIF2B5	NM_003907.2	GALC*	NM_000153.3
CTSD	NM_001909.4	ELP1	NM_003640.3	GALE*	NM_000403.3
стѕк	NM_000396.3	EPG5	NM_020964.2	GALK1	NM_000154.1
СҮВА	NM_000101.3	ERCC2	NM_000400.3	GALNS	NM_000512.4
CYP11A1	NM_000781.2	ERCC6	NM_000124.3	GALNT3	NM_004482.3
CYP11B1	NM_000497.3	ERCC8	NM_000082.3	GALT	NM_000155.3
CYP11B2	NM_000498.3	ESCO2	NM_001017420.2	GAMT	NM_000156.5
CYP17A1	NM_000102.3	ETFA	NM_000126.3	GATM	NM_001482.2
CYP19A1	NM_031226.2	ETFB	NM_001985.2	GBA*	NM_001005741.2
CYP1B1	NM_000104.3	ETFDH	NM_004453.3	GBE1	NM_000158.3
CYP21A2*	NM_000500.7	ETHE1	NM_014297.3	GCDH	NM_000159.3
CYP27A1	NM_000784.3	EVC	NM_153717.2	GCH1	NM_000161.2
CYP27B1	NM_000785.3	EVC2	NM_147127.4	GDF5	NM_000557.4
CYP7B1	NM_004820.3	EXOSC3	NM_016042.3	GFM1	NM_024996.5
DBT	NM_001918.3	EYS*	NM_001142800.1	GHR*	NM_000163.4
DCAF17	NM_025000.3	FAH*	NM_000137.2	GJB2	NM_004004.5
DCLRE1C	NM_001033855.2	FAM161A	NM_001201543.1	GLB1	NM_000404.2
DDX11*	NM_030653.3	FANCA	NM_000135.2	GLDC	NM_000170.2
DFNB59	NM_001042702.3	FANCC	NM_000136.2	GLE1	NM_001003722.1
DGAT1	NM_012079.5	FANCD2*	NM_033084.3	GNE*	NM_001128227.2
DGUOK	NM_080916.2	FANCE	NM_021922.2	GNPAT	NM_014236.3
DHCR7	NM_001360.2	FANCG	NM_004629.1	GNPTAB	NM_024312.4
DHDDS	NM_024887.3	FANCI	NM_001113378.1	GNPTG	NM_032520.4
DLD	NM_000108.4	FANCL*	NM_018062.3	GNS	NM_002076.3
DLL3	NM_016941.3	FBP1	NM_000507.3	GORAB	NM_152281.2
DNAH11	NM_001277115.1	FBXO7	NM_012179.3	GRHPR	NM_012203.1
DNAH5	NM_001369.2	FH*	NM_000143.3	GRIP1	NM_021150.3
DNAI1	NM_012144.3	FKBP10	NM_021939.3	GSS	NM_000178.2
DNAI2	NM_023036.4	FKRP	NM_024301.4	GUCY2D	NM_000180.3
DNMT3B	NM_006892.3	FKTN	NM_001079802.1	GUSB	NM_000181.3
DOK7	NM_173660.4	FMO3	NM_006894.6	HADH	NM_005327.4
DUOX2*	NM_014080.4	FOXN1	NM_003593.2	HADHA	NM_000182.4
DYNC2H1	NM_001080463.1	FOXRED1	NM_017547.3	HADHB	NM_000183.2
DYSF	NM_003494.3	FRAS1	NM_025074.6	НАМР	NM_021175.2
EIF2AK3	NM_004836.6	FREM2	NM_207361.5	HAX1	NM_006118.3





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GENE	TRANSCRIPT	GENE	TRANSCRIPT	GENE	TRANSCRIPT
HBA1*	NM_000558.4	LCA5	NM_181714.3	MTHFR*	NM_005957.4
HBA2	NM_000517.4	LDLR	NM_000527.4	MTR	NM_000254.2
НВВ	NM_000518.4	LDLRAP1	NM_015627.2	MTRR	NM_002454.2
HEXA	NM_000520.4	LHX3	NM_014564.4	MTTP	NM_000253.3
HEXB	NM_000521.3	LIFR*	NM_002310.5	MUSK	NM_005592.3
HGSNAT	NM_152419.2	LIG4	NM_002312.3	MUT	NM_000255.3
НЈ∨	NM_213653.3	LIPA	NM_000235.3	MVK	NM_000431.3
HLCS	NM_000411.6	LMBRD1	NM_018368.3	MYO15A	NM_016239.3
HMGCL	NM_000191.2	LOXHD1	NM_144612.6	MYO7A	NM_000260.3
HMOX1	NM_002133.2	LPL	NM_000237.2	NAGA	NM_000262.2
HOGA1	NM_138413.3	LRAT	NM_004744.4	NAGLU	NM_000263.3
HPD	NM_002150.2	LRP2	NM_004525.2	NAGS	NM_153006.2
HPS1	NM_000195.4	LRPPRC	NM_133259.3	NBN	NM_002485.4
HPS3	NM_032383.4	LYST	NM_000081.3	NCF2	NM_000433.3
HPS4	NM_022081.5	МАК	NM_001242957.2	NDRG1	NM_006096.3
HPS5	NM_181507.1	MAN2B1	NM_000528.3	NDUFAF2	NM_174889.4
HPS6	NM_024747.5	MANBA	NM_005908.3	NDUFAF5	NM_024120.4
HSD17B3	NM_000197.1	MCEE	NM_032601.3	NDUFS4	NM_002495.3
HSD17B4	NM_000414.3	MCOLN1	NM_020533.2	NDUFS6	NM_004553.4
HSD3B2	NM_000198.3	MCPH1	NM_024596.4	NDUFS7	NM_024407.4
HYAL1	NM_153281.1	MECR	NM_016011.3	NDUFV1	NM_007103.3
HYLS1	NM_145014.2	MED17	NM_004268.4	NEB*	NM_001271208.1
IDUA	NM_000203.4	MESP2	NM_001039958.1	NEU1	NM_000434.3
IGHMBP2	NM_002180.2	MFSD8	NM_152778.2	NGLY1	NM_018297.3
ІКВКВ	NM_001556.2	MKKS	NM_018848.3	NPC1	NM_000271.4
IL7R	NM_002185.3	MKS1	NM_017777.3	NPC2	NM_006432.3
INVS	NM_014425.3	MLC1*	NM_015166.3	NPHP1	NM_000272.3
ITGA6	NM_000210.3	MLYCD	NM_012213.2	NPHS1	NM_004646.3
ITGB3	NM_000212.2	MMAA	NM_172250.2	NPHS2	NM_014625.3
ITGB4	NM_001005731.2	MMAB	NM_052845.3	NR2E3	NM_014249.3
IVD	NM_002225.3	ММАСНС	NM_015506.2	NSMCE3	NM_138704.3
JAK3	NM_000215.3	MMADHC	NM_015702.2	NTRK1	NM_001012331.1
, KCNJ1	NM_000220.4	MOCS1	NM_001358530.2	OAT*	NM_000274.3
KCNJ11	NM_000525.3	MOCS2A	NM_176806.3	OCA2	NM_000275.2
LAMA2	NM_000426.3	MOCS2B	NM_004531.4	OPA3	NM_025136.3
LAMA3	NM_000227.4	MPI	NM_002435.2	OSTM1	NM_014028.3
LAMB3	NM_000228.2	MPL	NM_005373.2	OTOA*	NM_144672.3
LAMC2	NM_005562.2	MPV17	NM_002437.4	OTOF	NM_194248.2;NM_194323.2
LARGE1	NM_004737.4	MRE11	NM_005591.3	P3H1	NM_022356.3





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GENE	TRANSCRIPT	GENE	TRANSCRIPT	GENE	TRANSCRIPT
РАН	NM_000277.1	POR	NM_000941.2	SGSH	NM_000199.3
PANK2	NM_153638.2	POU1F1	NM_000306.3	SKIV2L	NM_006929.4
PC	NM_000920.3	PPT1	NM_000310.3	SLC12A1	NM_000338.2
PCBD1	NM_000281.3	PRCD	NM_001077620.2	SLC12A3	NM_000339.2
PCCA	NM_000282.3	PRDM5	NM_018699.3	SLC12A6	NM_133647.1
PCCB	NM_000532.4	PRF1	NM_001083116.1	SLC17A5	NM_012434.4
PCDH15	NM_033056.3	PROP1	NM_006261.4	SLC19A2	NM_006996.2
PCNT	NM_006031.5	PSAP	NM_002778.3	SLC19A3	NM_025243.3
PDHB	NM_000925.3	PTPRC*	NM_002838.4	SLC1A4	NM_003038.4
PEPD	NM_000285.3	PTS	NM_000317.2	SLC22A5	NM_003060.3
PET100	NM_001171155.1	PUS1	NM_025215.5	SLC25A13	NM_014251.2
PEX1*	NM_000466.2	PYGM	NM_005609.3	SLC25A15	NM_014252.3
PEX10	NM_153818.1	QDPR	NM_000320.2	SLC25A20	NM_000387.5
PEX12	NM_000286.2	RAB23	NM_183227.2	SLC26A2	NM_000112.3
PEX13	NM_002618.3	RAG1	NM_000448.2	SLC26A3	NM_000111.2
PEX16	NM_004813.2	RAG2	NM_000536.3	SLC26A4	NM_000441.1
PEX2	NM_000318.2	RAPSN	NM_005055.4	SLC27A4	NM_005094.3
PEX26	NM_017929.5	RARS2	NM_020320.3	SLC35A3	NM_012243.2
PEX5	NM_001131025.1	RDH12	NM_152443.2	SLC37A4	NM_001164277.1
PEX6	NM_000287.3	RLBP1	NM_000326.4	SLC38A8	NM_001080442.2
PEX7	NM_000288.3	RMRP	NR_003051.3	SLC39A4	NM_130849.3
PFKM	NM_000289.5	RNASEH2A	NM_006397.2	SLC45A2	NM_016180.4
PGM3	NM_001199917.1	RNASEH2B	NM_024570.3	SLC4A11	NM_032034.3
PHGDH	NM_006623.3	RNASEH2C	NM_032193.3	SLC5A5	NM_000453.2
РНКВ	NM_000293.2;NM_00103183	RPE65	NM_000329.2	SLC7A7	NM_001126106.2
	5.2	RPGRIP1L	NM_015272.2	SMARCAL1	NM_014140.3
PHKG2	NM_000294.2	RTEL1	NM_001283009.1	SMN1*	NM_000344.3
РНҮН	NM_006214.3	RXYLT1	NM_014254.2	SMPD1	NM_000543.4
PIGN	NM_176787.4	RYR1	NM_000540.2	SNAP29	NM_004782.3
PKHD1*	NM_138694.3	SACS	NM_014363.5	SPG11	NM_025137.3
PLA2G6	NM_003560.2	SAMD9	NM_017654.3	SPR	NM_003124.4
PLEKHG5	NM_020631.4	SAMHD1	NM_015474.3	SRD5A2	NM_000348.3
PLOD1	NM_000302.3	SCO2	NM_005138.2	ST3GAL5	NM_003896.3
PMM2	NM_000303.2	SEC23B	NM_006363.4	STAR	NM_000349.2
PNPO	NM_018129.3	SEPSECS	NM_016955.3	STX11	NM_003764.3
POLG	NM_002693.2	SGCA	NM_000023.2	STXBP2	NM_006949.3
POLH	NM_006502.2	SGCB	NM_000232.4	SUMF1	NM_182760.3
POMGNT1	NM_017739.3	SGCD	NM_000337.5	SUOX	NM_000456.2
POMT1	NM_007171.3	SGCG	NM_000231.2	SURF1	NM_003172.3
POMT2	NM_013382.5				





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SYNE4 NM_001039876.2 TANGO2 NM_152906.6 TAT NM_000353.2 TBCD NM_005993.4 TBCE* NM_00619.3 TCN2 NM_00355.3 TECPR2 NM_014844.3 TERT NM_198253.2
TAT NM_000353.2 TBCD NM_005993.4 TBCE* NM_003193.4 TCIRG1 NM_006019.3 TCN2 NM_000355.3 TECPR2 NM_014844.3 TERT NM_198253.2
TBCD NM_005993.4 TBCE* NM_003193.4 TCIRG1 NM_006019.3 TCN2 NM_000355.3 TECPR2 NM_014844.3 TERT NM_198253.2
TBCE* NM_003193.4 TCIRG1 NM_006019.3 TCN2 NM_000355.3 TECPR2 NM_014844.3 TERT NM_198253.2
TCIRG1 NM_006019.3 TCN2 NM_000355.3 TECPR2 NM_014844.3 TERT NM_198253.2
TCN2 NM_000355.3 TECPR2 NM_014844.3 TERT NM_198253.2
TECPR2 NM_014844.3 TERT NM_198253.2
TERT NM_198253.2
TF NM_001063.3
TFR2 NM_003227.3
TG* NM_003235.4
TGM1 NM_000359.2
TH NM_199292.2
TK2 NM_004614.4
TMC1 NM_138691.2
TMEM216 NM_001173990.2
TMEM67 NM_153704.5
TMPRSS3 NM_024022.2
TPO NM_000547.5
TPP1 NM_000391.3
TREX1 NM_033629.4
TRIM32 NM_012210.3
TRIM37 NM_015294.4
TRMU NM_018006.4
TSEN54 NM_207346.2
TSFM* NM_001172696.1
TSHB NM_000549.4
TSHR NM_000369.2
TTC37 NM_014639.3
TTPA NM_000370.3
TULP1 NM_003322.4
TYMP NM_001953.4
TYR* NM_000372.4
TYRP1 NM_000550.2
UBR1 NM_174916.2
UNC13D NM_199242.2
USH1C* NM_005709.3
USH2A NM_206933.2

GENE	TRANSCRIPT
VDR	NM_001017535.1
VLDLR	NM_003383.4
VPS11	NM_021729.5
VPS13A*	NM_033305.2
VPS13B	NM_017890.4
VPS45	NM_007259.4
VPS53*	NM_001128159.2
VRK1	NM_003384.2
VSX2	NM_182894.2
WISP3	NM_003880.3
WNT10A	NM_025216.2
WRN*	NM_000553.4
XPA	NM_000380.3
XPC	NM_004628.4
ZBTB24	NM_014797.2
ZFYVE26	NM_015346.3
ZNF469	NM_001127464.2





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Invitae #: RQ5172543

Methods

■ Genomic DNA obtained from the submitted sample is enriched for targeted regions using a hybridization-based protocol, and sequenced using Illumina technology. Unless otherwise indicated, all targeted regions are sequenced with ≥50x depth or are supplemented with additional analysis. Reads are aligned to a reference sequence (GRCh37), and sequence changes are identified and interpreted in the context of a single clinically relevant transcript, indicated in the Genes Analyzed table. Enrichment and analysis focus on the coding sequence of the indicated transcripts, 20bp of flanking intronic sequence, and other specific genomic regions demonstrated to be causative of disease at the time of assay design. Promoters, untranslated regions, and other non-coding regions are not otherwise interrogated. Exonic deletions and duplications are called using an in-house algorithm that determines copy number at each target by comparing the read depth for each target in the proband sequence with both mean read-depth and read-depth distribution, obtained from a set of clinical samples. Markers across the X and Y chromosomes are analyzed for quality control purposes and may detect deviations from the expected sex chromosome complement. Such deviations may be included in the report in accordance with internal guidelines. Invitae utilizes a classification methodology to identify next-generation sequencing (NGS)-detected variants that require orthogonal confirmation (Lincoln, et al. J Mol Diagn. 2019 Mar;21(2):318-329). Confirmation of the presence and location of reportable variants is performed as needed based on stringent criteria using one of several validated orthogonal approaches (PubMed ID 30610921). Sequencing is performed by Invitae Corporation (1400 16th Street, San Francisco, CA 94103, #05D2040778).

The following additional analyses are performed if relevant to the requisition. For GBA the reference genome has been modified to mask the sites of polymorphic paralog sequence variants (PSVs) in both the gene and pseudogene. For CYP21A2 and GBA, if one or more reportable variants, gene conversion, or fusion event is identified via our NGS pipeline (see Limitations), these variants are confirmed by PacBio sequencing of an amplicon generated by long-range PCR and subsequent short-range PCR. In some cases, it may not be possible to disambiguate between the gene and pseudogene. For GJB2, the reportable range includes large upstream deletions overlapping GJB6. For HBA1/2, the reference genome has been modified to force some sequencing reads derived from HBA1 to align to HBA2, and variant calling algorithms are modified to support an expectation of 4 alleles in these regions. HBA1/2 copy number calling is performed by a custom hypothesis testing algorithm which generates diplotype calls. If sequence data for a sample does not support a unique high confidence match from among hypotheses tested, that sample is flagged for manual review. Copy number variation is only reported for coding sequence of HBA1 and HBA2 and the HS-40 region. This assay does not distinguish among the $-\alpha$ 3.7 subtypes, and all $-\alpha$ 3.7 variants are called as HBA1 deletions. This assay may not detect overlapping copy gain and copy loss events when the breakpoints of those events are similar. For FMR1, cytosine-guanine-guanine (CGG) triplet repeats in the 5' untranslated region (5' UTR) of the FMR1 gene are detected by triplet repeat-primed PCR (RP-PCR) with fluorescently labeled primers followed by capillary electrophoresis. Reference ranges: Normal: <45 CGG repeats, intermediate: 45-54 CGG repeats, premutation: 55-200 CGG repeats, full mutation: >200 CGG repeats. For alleles with 55-90 triplet repeats, the region surrounding the FMR1 repeat is amplified by PCR. The PCR amplicons are then processed through PacBio SMRTBell library prep and sequenced using PacBio long read technology. The number of AGG interruptions within the 55-90 triplet repeat is read directly from the resulting DNA sequences.

- This report only includes variants that have a clinically significant association with the conditions tested as of the report date. Variants of uncertain significance, benign variants, and likely benign variants are not included in this report. However, if additional evidence becomes available to indicate that the clinical significance of a variant has changed, Invitae may update this report and provide notification.
- A PMID is a unique identifier referring to a published, scientific paper. Search by PMID at http://www.ncbi.nlm.nih.gov/pubmed.
- An rsID is a unique identifier referring to a single genomic position, and is used to associate population frequency information with sequence changes at that position. Reported population frequencies are derived from a number of public sites that aggregate data from large-scale population sequencing projects, including ExAC (http://exac.broadinstitute.org), gnomAD (http://gnomad.broadinstitute.org), and dbSNP (http://ncbi.nlm.nih.gov/SNP).

Disclaimer

DNA studies do not constitute a definitive test for the selected condition(s) in all individuals. It should be realized that there are possible sources of error. Errors can result from trace contamination, rare technical errors, rare genetic variants that interfere with analysis, recent scientific developments, and alternative classification systems. This test should be one of many aspects used by the healthcare provider to help with a diagnosis and treatment plan, but it is not a diagnosis itself. This test was developed and its performance characteristics determined by Invitae. It has not been cleared or approved by





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the FDA. The laboratory is regulated under the Clinical Laboratory Improvement Act (CLIA) as qualified to perform high-complexity clinical tests (CLIA ID: 05D2040778). This test is used for clinical purposes. It should not be regarded as investigational or for research.

Limitations

- Based on validation study results, this assay achieves >99% analytical sensitivity and specificity for single nucleotide variants, insertions and deletions <15bp in length, and exon-level deletions and duplications. Invitae's methods also detect insertions and deletions larger than 15bp but smaller than a full exon but sensitivity for these may be marginally reduced. Invitae's deletion/duplication analysis determines copy number at a single exon resolution at virtually all targeted exons. However, in rare situations, single-exon copy number events may not be analyzed due to inherent sequence properties or isolated reduction in data quality. Certain types of variants, such as structural rearrangements (e.g. inversions, gene conversion events, translocations, etc.) or variants embedded in sequence with complex architecture (e.g. short tandem repeats or segmental duplications), may not be detected. Additionally, it may not be possible to fully resolve certain details about variants, such as mosaicism, phasing, or mapping ambiguity. Unless explicitly guaranteed, sequence changes in the promoter, non-coding exons, and other non-coding regions are not covered by this assay. Please consult the test definition on our website for details regarding regions or types of variants that are covered or excluded for this test. This report reflects the analysis of an extracted genomic DNA sample. While this test is intended to reflect the analysis of extracted genomic DNA from a referred patient, in very rare cases the analyzed DNA may not represent that individual's constitutional genome, such as in the case of a circulating hematolymphoid neoplasm, bone marrow transplant, blood transfusion, chimerism, culture artifact or maternal cell contamination.</p>
- DUOX2: Deletion/duplication and sequencing analysis is not offered for exons 6-7. PTPRC: Sequencing analysis is not offered for exons 3, 15. ABCC2: Deletion/duplication analysis is not offered for exons 24-25. OTOA: Deletion/duplication and sequencing analysis is not offered for exons 20-28. TBCE: Sequencing analysis for exons 2 includes only cds +/- 10 bp. GALE: Sequencing analysis for exons 10 includes only cds +/- 5 bp. DDX11: NM_030653.3:c.1763-1G>C variant only. PKHD1: Deletion/duplication analysis is not offered for exon 13. VPS13A: Deletion/duplication analysis is not offered for exons 2-3, 27-28. GNE: Sequencing analysis for exons 8 includes only cds +/- 10 bp. NEB: Deletion/duplication analysis is not offered for exons 82-105. NEB variants in this region with no evidence towards pathogenicity are not included in this report, but are available upon request. BBS9: Deletion/duplication analysis is not offered for exon 4. WRN: Deletion/duplication analysis is not offered for exons 10-11. Sequencing analysis for exons 8, 10-11 includes only cds +/- 10 bp. OAT: Deletion/duplication analysis is not offered for exon 2. GHR: Deletion/duplication and sequencing analysis is not offered for exon 3. CFTR: Sequencing analysis for exons 7 includes only cds +/- 10 bp. EYS: Sequencing analysis for exons 30 includes only cds +/- 0 bp. FH: Sequencing analysis for exons 9 includes only cds +/- 10 bp. ANO10: Sequencing analysis for exons 8 includes only cds +/- 0 bp. ATP8B1: Sequencing analysis for exons 19 includes only cds +/- 10 bp. FANCD2: Deletion/ duplication analysis is not offered for exons 14-17, 22 and sequencing analysis is not offered for exons 15-17. Sequencing analysis for exons 6, 14, 18, 20, 23, 25, 34 includes only cds +/- 10 bp. COL11A2: Deletion/duplication analysis is not offered for exon 36. TSFM: Sequencing analysis is not offered for exon 5. VPS53: Sequencing analysis for exons 14 includes only cds +/- 5 bp. HBA1/2: This assay is designed to detect deletions and duplications of HBA1 and/or HBA2, resulting from the -alpha20.5, --MED, --SEA, --FIL/--THAI, -alpha3.7, -alpha4.2, anti3.7 and anti4.2. Sensitivity to detect other copy number variants may be reduced. Detection of overlapping deletion and duplication events will be limited to combinations of events with significantly differing boundaries. In addition, deletion of the enhancer element HS-40 and the sequence variant, Constant Spring (NM_000517.4:c.427T>C), can be identified by this assay. MTHFR: The NM_005957.4:c.665C>T (p.Ala222Val) (aka 677C>T) and c.1286A>C (p.Glu429Ala) (aka 1298A>C) variants are not reported in our primary report. GBA: c.84dupG (p.Leu29Alafs*18), c.115+1G>A (Splice donor), c.222_224delTAC (p.Thr75del), c.475C>T (p.Arg159Trp), c.595_596delCT (p.Leu199Aspfs*62), c.680A>G (p.Asn227Ser), c.721G>A (p.Gly241Arg), c.754T>A (p.Phe252Ile), c.1226A>G (p.Asn409Ser), c.1246G>A (p.Gly416Ser), c.1263_1317del (p.Leu422Profs*4), c.1297G>T (p.Val433Leu), c.1342G>C (p.Asp448His), c.1343A>T (p.Asp448Val), c.1448T>C (p.Leu483Pro), c.1504C>T (p.Arg502Cys), c.1505G>A (p.Arg502His), c.1603C>T (p.Arg535Cys), c.1604G>A (p.Arg535His) variants only. Rarely, sensitivity to detect these variants may be reduced. When sensitivity is reduced, zygosity may be reported as "unknown". SMN1: Systematic exon numbering is used for all genes, including SMN1, and for this reason the exon typically referred to as exon 7 in the literature (PMID: 8838816) is referred to as exon 8 in this report. This assay unambiguously detects SMN1 exon 8 copy number. The presence of the g.27134T>G variant (also known as c.*3+80T>G) is reported if SMN1 copy number = 2. SMN1 or SMN2: NM_000344.3:c.*3+80T>G variant only. CYP21A2: Analysis includes the most common variants (c.92C>T(p.Pro31Leu), c.293-13C>G (intronic), c.332_339delGAGACTAC (p.Gly111Valfs*21), c.518T>A (p.Ile173Asn), c.710T>A (p.Ile237Asn), c.713T>A (p.Val238Glu), c.719T>A (p.Met240Lys), c.844G>T (p.Val282Leu), c.923dupT (p.Leu308Phefs*6), c.955C>T (p.Gln319*), c.1069C>T(p.Arg357Trp), c.1360C>T (p.Pro454Ser) and the 30Kb deletion) as well as select rare HGMD variants only (list available upon request). Full gene duplications are reported only in the presence of a pathogenic variant(s). When a duplication and a pathogenic variant(s) is identified, phase (cis/trans) cannot be determined. Full gene deletion analysis is not offered. Sensitivity to detect these variants, if they result from complex gene conversion/fusion events, may be reduced. AIPL1: Sequencing analysis for exons 2 includes only cds +/- 10 bp. LIFR: Sequencing analysis for exons 3 includes only





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cds +/- 5 bp. AMN: Deletion/duplication analysis is not offered for exon 1. PEX1: Sequencing analysis for exons 16 includes only cds +/- 0 bp. USH1C: Deletion/duplication analysis is not offered for exons 5-6. TYR: Deletion/duplication and sequencing analysis is not offered for exon 5. TG: Deletion/duplication analysis is not offered for exon 18. Sequencing analysis for exons 44 includes only cds +/- 0 bp. FANCL: Sequencing analysis for exons 4, 10 includes only cds +/- 10 bp. MLC1: Sequencing analysis for exons 11 includes only cds +/- 10 bp. ATM: Sequencing analysis for exons 6, 24, 43 includes only cds +/- 10 bp. FAH: Deletion/duplication analysis is not offered for exon 6.

This report has been reviewed and approved by:

Burth.

Arunkanth Ankala, Ph.D., FACMG Clinical Molecular Geneticist



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This table displays residual risks after a negative result for each of the genes and corresponding disorders. The values provided assume a negative family history and the absence of symptoms for each disorder. For genes associated with both dominant and recessive inheritance, the numbers in this table apply to the recessive condition(s) associated with the gene, unless otherwise noted. Residual risk values are provided for disorders when carrier frequency is greater than 1 in 500. For disorders with carrier frequency equal to, or less than, 1 in 500, residual risk is considered to be reduced substantially. When provided, residual risk values are inferred from published carrier frequencies, and estimated detection rates are based on testing technologies used at Invitae. Residual risks are provided only as a guide for assessing approximate risk given a negative result; values may vary based on the ethnic background(s) of an individual. For any genes marked with an asterisk*, refer to the Limitations section of the patient report for detailed coverage information. In the case of a sample-specific limitation, "N/A" indicates that a residual risk value could not be calculated. AR = autosomal recessive, XL = X-linked, AD = autosomal dominant.

DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY	DETECTION RATE	RISK TO BE A CARRIER AFTER NEGATIVE RESULT
3-hydroxy-3-methylglutaryl-CoA lyase deficiency (AR) NM_000191.2	HMGCL	Pan-ethnic	≤1 in 500	99%	Reduced
17-beta hydroxysteroid dehydrogenase 3 deficiency (AR) NM_000197.1	HSD17B3	Pan-ethnic	≤1 in 500	99%	Reduced
ABCA3-related conditions (AR) NM_001089.2	ABCA3	Pan-ethnic	1 in 277	99%	1 in 27600
ABCA4-related conditions (AR) NM_000350.2	ABCA4	Pan-ethnic	1 in 45	90%	1 in 441
ABCB4-related conditions (AR) NM_000443.3	ABCB4	Pan-ethnic	1 in 204	99%	1 in 20300
ABCB11-related conditions (AR) NM_003742.2	ABCB11	Pan-ethnic	1 in 100	99%	1 in 9900
ABCC8-related conditions (AR) NM_000352.4 When the mother is a noncarrier, but the father is a carrier, there is a residual risk for focal disease (1 in 540 for the Ashkenazi Jewish population; undetermined in other ethnic groups)	ABCC8	Pan-ethnic	1 in 177	99%	1 in 17600
Abetalipoproteinemia (AR) NM_000253.3	MTTP	Pan-ethnic	≤1 in 500	99%	Reduced
Achromatopsia (CNGB3-related) (AR) NM_019098.4	CNGB3	Pan-ethnic	1 in 93	99%	1 in 9200
ACOX1-related conditions (AR) NM_004035.6	ACOX1	Pan-ethnic	≤1 in 500	99%	Reduced
Acrodermatitis enteropathica (AR) NM_130849.3	SLC39A4	Pan-ethnic	1 in 354	99%	1 in 35300
ADGRV1-related conditions (AR) NM_032119.3	ADGRV1	Pan-ethnic	1 in 223	99%	1 in 22200
AHI1-related conditions (AR) NM_017651.4	AHI1	Pan-ethnic	1 in 447	99%	1 in 44600
Aicardi-Goutieres syndrome 2 (AR) NM_024570.3	RNASEH2B	Pan-ethnic	≤1 in 500	99%	Reduced
Aicardi-Goutieres syndrome 3 (AR) NM_032193.3	RNASEH2C	Pan-ethnic	≤1 in 500	99%	Reduced
Aicardi-Goutieres syndrome 4 (AR) NM_006397.2	RNASEH2A	Pan-ethnic	≤1 in 500	99%	Reduced
Aicardi-Goutieres syndrome 5 (AR) NM_015474.3	SAMHD1	Pan-ethnic	≤1 in 500	99%	Reduced
AIPL1-related conditions (AR) NM_014336.4	AIPL1 *	Pan-ethnic	1 in 408	99%	1 in 40700
Aldosterone synthase deficiency (AR) NM_000498.3	CYP11B2	Pan-ethnic	≤1 in 500	99%	Reduced
Alpha-mannosidosis (AR) NM_000528.3	MAN2B1	Pan-ethnic	1 in 354	99%	1 in 35300
Alpha-N-acetylgalactosaminidase deficiency (AR) NM_000262.2	NAGA	Pan-ethnic	≤1 in 500	99%	Reduced
		African-American	1 in 30	90%	1 in 291
Alpha-thalassemia (AR)	HBA1/	Asian	1 in 20	90%	1 in 191
NM_000558.4, NM_000517.4	HBA2 *	Caucasian	≤1 in 500	90%	Reduced
		Pan-ethnic	1 in 25	90%	1 in 241



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DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY	DETECTION RATE	RISK TO BE A CARRIER AFTER NEGATIVE RESULT
Alport syndrome (COL4A3-related) (AR) NM_000091.4	COL4A3	Pan-ethnic	1 in 354	99%	1 in 35300
Alport syndrome (COL4A4-related) (AR) NM_000092.4	COL4A4	Pan-ethnic	1 in 353	99%	1 in 35200
Alström syndrome (AR) NM_015120.4	ALMS1	Pan-ethnic	≤1 in 500	99%	Reduced
Arginase deficiency (AR) NM_000045.3	ARG1	Pan-ethnic	1 in 274	99%	1 in 27300
Argininosuccinate lyase deficiency (AR) NM_000048.3	ASL	Pan-ethnic	1 in 133	90%	1 in 1321
ARL6-related conditions (AR) NM_177976.2	ARL6	Pan-ethnic	≤1 in 500	99%	Reduced
Aromatase deficiency (AR) NM_031226.2	CYP19A1	Pan-ethnic	≤1 in 500	99%	Reduced
Asparagine synthetase deficiency (AR) NM_133436.3	ASNS	Pan-ethnic	≤1 in 500	99%	Reduced
Aspartylglucosaminuria (AR) NM_000027.3	AGA	Pan-ethnic	≤1 in 500	99%	Reduced
Ataxia with vitamin E deficiency (AR) NM_000370.3	ТТРА	Pan-ethnic	≤1 in 500	90%	Reduced
Ataxia-telangiectasia-like disorder (AR) NM_005591.3	MRE11	Pan-ethnic	≤1 in 500	99%	Reduced
ATM-related conditions (AR) NM_000051.3	ATM *	Pan-ethnic	1 in 100	99%	1 in 9900
ATP8B1-related conditions (AR) NM_005603.4	ATP8B1 *	Pan-ethnic	1 in 112	99%	1 in 11100
Atransferrinemia (AR) NM_001063.3	TF	Pan-ethnic	≤1 in 500	99%	Reduced
Autoimmune polyendocrinopathy with candidiasis and ectodermal dysplasia (AR) NM_000383.3	AIRE	Pan-ethnic	1 in 150	99%	1 in 14900
Autosomal recessive congenital ichthyosis (ABCA12-related) (AR) NM_173076.2	ABCA12	Pan-ethnic	≤1 in 500	99%	Reduced
Autosomal recessive congenital ichthyosis (TGM1-related) (AR) NM_000359.2	TGM1	Pan-ethnic	1 in 224	95%	1 in 4460
Autosomal recessive spastic ataxia of Charlevoix-Saguenay (AR) NM_014363.5	SACS	Pan-ethnic	≤1 in 500	99%	Reduced
Bardet-Biedl syndrome (BBS7-related) (AR) NM_176824.2	BBS7	Pan-ethnic	≤1 in 500	99%	Reduced
Bardet-Biedl syndrome (BBS9-related) (AR) NM_198428.2	BBS9 *	Pan-ethnic	≤1 in 500	99%	Reduced
Bardet-Biedl syndrome (BBS10-related) (AR) NM_024685.3	BBS10	Pan-ethnic	1 in 354	99%	1 in 35300
Bardet-Biedl syndrome (BBS12-related) (AR) NM_152618.2	BBS12	Pan-ethnic	≤1 in 500	99%	Reduced
Bartter syndrome type 1 (AR) NM_000338.2	SLC12A1	Pan-ethnic	1 in 224	99%	1 in 22300
Bartter syndrome type 2 (AR) NM_000220.4	KCNJ1	Pan-ethnic	≤1 in 500	99%	Reduced
BBS1-related conditions (AR) NM_024649.4	BBS1	Pan-ethnic	1 in 330	99%	1 in 32900
BBS2-related conditions (AR) NM_031885.3	BBS2	Pan-ethnic	≤1 in 500	99%	Reduced
BBS4-related conditions (AR) NM_033028.4	BBS4	Pan-ethnic	≤1 in 500	99%	Reduced
BBS5-related conditions (AR) NM_152384.2	BBS5	Pan-ethnic	≤1 in 500	99%	Reduced
BCS1L-related conditions (AR) NM_004328.4	BCS1L	Pan-ethnic	≤1 in 500	99%	Reduced

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DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY	DETECTION RATE	RISK TO BE A CARRIER AFTER NEGATIVE RESULT
Beta-ketothiolase deficiency (AR) NM_000019.3	ACAT1	Pan-ethnic	≤1 in 500	99%	Reduced
Beta-mannosidosis (AR) NM_005908.3	MANBA	Pan-ethnic	≤1 in 500	99%	Reduced
Biopterin-deficient hyperphenylalaninemia (PCBD1-related) (AR) NM_000281.3	PCBD1	Pan-ethnic	≤1 in 500	99%	Reduced
Biopterin-deficient hyperphenylalaninemia (PTS-related) (AR) NM_000317.2	PTS	Pan-ethnic	1 in 433	99%	1 in 43200
Biopterin-deficient hyperphenylalaninemia (QDPR-related) (AR) NM_000320.2	QDPR	Pan-ethnic	≤1 in 500	99%	Reduced
Biotin-responsive basal ganglia disease (AR) NM_025243.3	SLC19A3	Pan-ethnic	≤1 in 500	99%	Reduced
Biotinidase deficiency (AR) NM_000060.3	BTD	Pan-ethnic	1 in 125	99%	1 in 12400
Bloom syndrome (AR) NM_000057.3	BLM	Pan-ethnic	≤1 in 500	99%	Reduced
BRIP1-related conditions (AR) NM_032043.2	BRIP1	Pan-ethnic	≤1 in 500	99%	Reduced
Brittle cornea syndrome (PRDM5-related) (AR) NM_018699.3	PRDM5	Pan-ethnic	≤1 in 500	99%	Reduced
Brittle cornea syndrome (ZNF469-related) (AR) NM_001127464.2	ZNF469	Pan-ethnic	≤1 in 500	99%	Reduced
BSND-related conditions (AR) NM_057176.2	BSND	Pan-ethnic	≤1 in 500	99%	Reduced
Canavan disease (AR) NM_000049.2	ASPA	Pan-ethnic	1 in 159	99%	1 in 15800
Carbamoyl phosphate synthetase I deficiency (AR) NM_001875.4	CPS1	Pan-ethnic	≤1 in 500	99%	Reduced
Cardioencephalomyopathy (AR) NM_005138.2	SCO2	Pan-ethnic	1 in 387	99%	1 in 38600
Carnitine palmitoyltransferase I deficiency (AR) NM_001876.3	CPT1A	Pan-ethnic	≤1 in 500	99%	Reduced
Carnitine palmitoyltransferase II deficiency (AR) NM_000098.2	CPT2	Pan-ethnic	1 in 182	99%	1 in 18100
Carnitine-acylcarnitine translocase deficiency (AR) NM_000387.5	SLC25A20	Pan-ethnic	≤1 in 500	99%	Reduced
Carpenter syndrome (RAB23-related) (AR) NM_183227.2	RAB23	Pan-ethnic	≤1 in 500	99%	Reduced
Cartilage-hair hypoplasia-anauxetic dysplasia spectrum disorders (AR) NR_003051.3	RMRP	Pan-ethnic	≤1 in 500	99%	Reduced
Catecholaminergic polymorphic ventricular tachycardia (CASQ2-related) (AR) NM_001232.3	CASQ2	Pan-ethnic	1 in 224	99%	1 in 22300
CC2D2A-related conditions (AR) NM_001080522.2	CC2D2A	Pan-ethnic	1 in 426	99%	1 in 42500
CDH23-related conditions (AR) NM_022124.5	CDH23	Pan-ethnic	1 in 202	95%	1 in 4020
CEP290-related conditions (AR) NM_025114.3	CEP290	Pan-ethnic	1 in 185	99%	1 in 18400
Cerebellar ataxia, intellectual disability, and dysequilibrium syndrome 1 (AR) NM_003383.4	VLDLR	Pan-ethnic	≤1 in 500	99%	Reduced
Cerebral dysgenesis, neuropathy, ichthyosis, and keratoderma (AR) NM_004782.3	SNAP29	Pan-ethnic	≤1 in 500	99%	Reduced
Cerebrotendinous xanthomatosis (AR) NM_000784.3	CYP27A1	Pan-ethnic	1 in 112	98%	1 in 5550
CERKL-related conditions (AR) NM_001030311.2	CERKL	Pan-ethnic	1 in 137	99%	1 in 13600



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DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY	DETECTION RATE	RISK TO BE A CARRIER AFTER NEGATIVE RESULT
CETD related conditions (AD)		Pan-ethnic - classic CF	1 in 45	99%	1 in 4400
CFTR-related conditions (AR) NM_000492.3	CFTR *	Pan-ethnic - classic CF and CFTR-related disorders	1 in 9	99%	1 in 800
Charcot-Marie-Tooth disease type 4D (AR) NM_006096.3	NDRG1	Pan-ethnic	≤1 in 500	99%	Reduced
Chediak-Higashi syndrome (AR) NM_000081.3	LYST	Pan-ethnic	≤1 in 500	99%	Reduced
Childhood-onset dystonia with optic atrophy and basal ganglia abnormalities (AR) NM_016011.3	MECR	Pan-ethnic	≤1 in 500	99%	Reduced
Chorea-acanthocytosis (AR) NM_033305.2	VPS13A *	Pan-ethnic	≤1 in 500	97%	Reduced
Chronic granulomatous disease (CYBA-related) (AR) NM_000101.3	СҮВА	Pan-ethnic	≤1 in 500	99%	Reduced
Chronic granulomatous disease (NCF2-related) (AR) NM_000433.3	NCF2	Pan-ethnic	≤1 in 500	99%	Reduced
Citrin deficiency (AR) NM_014251.2	SLC25A13	Pan-ethnic	1 in 313	99%	1 in 31200
Citrullinemia type 1 (AR) NM_000050.4	ASS1	Pan-ethnic	1 in 120	96%	1 in 2975
CLN3-related conditions (AR) NM_001042432.1	CLN3	Pan-ethnic	1 in 230	99%	1 in 22900
CLRN1-related conditions (AR) NM_174878.2	CLRN1	Pan-ethnic	≤1 in 500	99%	Reduced
Cobalamin C deficiency (AR) NM_015506.2	MMACHC	Pan-ethnic	1 in 123	99%	1 in 12200
Cobalamin D deficiency (AR) NM_015702.2	MMADHC	Pan-ethnic	≤1 in 500	99%	Reduced
Cobalamin F deficiency (AR) NM_018368.3	LMBRD1	Pan-ethnic	≤1 in 500	99%	Reduced
Cockayne syndrome A (AR) NM_000082.3	ERCC8	Pan-ethnic	≤1 in 500	99%	Reduced
Cockayne syndrome B (AR) NM_000124.3	ERCC6	Pan-ethnic	1 in 377	99%	1 in 37600
Cohen syndrome (AR) NM_017890.4	VPS13B	Pan-ethnic	≤1 in 500	99%	Reduced
COL11A2-related conditions (AR) NM_080680.2	COL11A2 *	Pan-ethnic	≤1 in 500	99%	Reduced
COL17A1-related conditions (AR) NM_000494.3	COL17A1	Pan-ethnic	≤1 in 500	99%	Reduced
Combined malonic and methylmalonic aciduria (AR) NM_174917.4	ACSF3	Pan-ethnic	1 in 87	99%	1 in 8600
Combined oxidative phosphorylation deficiency 1 (AR) NM_024996.5	GFM1	Pan-ethnic	≤1 in 500	99%	Reduced
Combined oxidative phosphorylation deficiency 3 (AR) NM_001172696.1	TSFM *	Pan-ethnic	≤1 in 500	93%	Reduced
Combined pituitary hormone deficiency (LHX3-related) (AR) NM_014564.4	LHX3	Pan-ethnic	≤1 in 500	99%	Reduced
Combined pituitary hormone deficiency (POU1F1-related) (AR) NM_000306.3	POU1F1	Pan-ethnic	≤1 in 500	99%	Reduced
Combined pituitary hormone deficiency (PROP1-related) (AR) NM_006261.4	PROP1	Pan-ethnic	1 in 45	98%	1 in 2200
Congenital adrenal hyperplasia due to 3-beta- hydroxysteroid dehydrogenase deficiency (AR) NM_000198.3	HSD3B2	Pan-ethnic	≤1 in 500	99%	Reduced
Congenital adrenal hyperplasia due to 21-hydroxylase deficiency (AR) NM_000500.7	CYP21A2 *	Pan-ethnic	1 in 61	92%	1 in 751
Congenital adrenal insufficiency (AR) NM_000781.2	CYP11A1	Pan-ethnic	≤1 in 500	99%	Reduced

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DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY	DETECTION RATE	RISK TO BE A CARRIER AFTER NEGATIVE RESULT
Congenital chronic diarrhea (DGAT1-related) (AR) NM_012079.5	DGAT1	Pan-ethnic	≤1 in 500	99%	Reduced
Congenital disorder of glycosylation (SLC35A3-related) (AR) NM_012243.2	SLC35A3	Pan-ethnic	≤1 in 500	99%	Reduced
Congenital disorder of glycosylation type Ia (AR) NM_000303.2	PMM2	Pan-ethnic	1 in 190	99%	1 in 18900
Congenital disorder of glycosylation type Ib (AR) NM_002435.2	MPI	Pan-ethnic	≤1 in 500	99%	Reduced
Congenital disorder of glycosylation type Ic (AR) NM_013339.3	ALG6	Pan-ethnic	≤1 in 500	99%	Reduced
Congenital disorder of glycosylation type Ik (AR) NM_019109.4	ALG1	Pan-ethnic	≤1 in 500	99%	Reduced
Congenital disorder of glycosylation type Iv (AR) NM_018297.3	NGLY1	Pan-ethnic	≤1 in 500	99%	Reduced
Congenital dyserythropoietic anemia type II (AR) NM_006363.4	SEC23B	Pan-ethnic	≤1 in 500	99%	Reduced
Congenital hydrocephalus-1 (AR) NM_001080414.3	CCDC88C	Pan-ethnic	≤1 in 500	99%	Reduced
Congenital hypothyroidism (TSHB-related) (AR) NM_000549.4	тѕнв	Pan-ethnic	≤1 in 500	99%	Reduced
Congenital insensitivity to pain with anhidrosis (AR) NM_001012331.1	NTRK1	Pan-ethnic	≤1 in 500	99%	Reduced
Congenital myasthenic syndrome (CHAT-related) (AR) NM_020549.4	СНАТ	Pan-ethnic	≤1 in 500	99%	Reduced
Congenital myasthenic syndrome (CHRNE-related) (AR) NM_000080.3	CHRNE	Pan-ethnic	1 in 200	99%	1 in 19900
Congenital nephrotic syndrome type 1 (AR) NM_004646.3	NPHS1	Pan-ethnic	≤1 in 500	99%	Reduced
Congenital nephrotic syndrome type 2 (AR) NM_014625.3	NPHS2	Pan-ethnic	≤1 in 500	99%	Reduced
Congenital secretory chloride diarrhea (AR) NM_000111.2	SLC26A3	Pan-ethnic	≤1 in 500	99%	Reduced
Corneal dystrophy and perceptive deafness (AR) NM_032034.3	SLC4A11	Pan-ethnic	≤1 in 500	99%	Reduced
CRB1-related conditions (AR) NM_201253.2	CRB1	Pan-ethnic	1 in 112	99%	1 in 11100
CTSC-related conditions (AR) NM_001814.5	стѕс	Pan-ethnic	1 in 250	99%	1 in 24900
CYP1B1-related conditions (AR) NM_000104.3	CYP1B1	Pan-ethnic	1 in 79	99%	1 in 7800
CYP7B1-related conditions (AR) NM_004820.3	CYP7B1	Pan-ethnic	≤1 in 500	99%	Reduced
CYP11B1-related conditions (AR) NM_000497.3	CYP11B1	Pan-ethnic	1 in 194	99%	1 in 19300
CYP17A1-related conditions (AR) NM_000102.3	CYP17A1	Pan-ethnic	≤1 in 500	99%	Reduced
Cystinosis (AR) NM_004937.2	CTNS	Pan-ethnic	1 in 158	99%	1 in 15700
Cytochrome P450 oxidoreductase deficiency (AR) NM_000941.2	POR	Pan-ethnic	1 in 158	99%	1 in 15700
Desbuquois dysplasia type 1 (AR) NM_138793.3	CANT1	Pan-ethnic	≤1 in 500	99%	Reduced
Developmental and epileptic encephalopathy (CAD-related) (AR) NM_004341.4	CAD	Pan-ethnic	≤1 in 500	99%	Reduced
DGUOK-related conditions (AR) NM_080916.2	DGUOK	Pan-ethnic	≤1 in 500	99%	Reduced
DHDDS-related conditions (AR) NM_024887.3	DHDDS	Pan-ethnic	≤1 in 500	99%	Reduced
Dihydrolipoamide dehydrogenase deficiency (AR) NM_000108.4	DLD	Pan-ethnic	≤1 in 500	99%	Reduced



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Distal renal tubular acidosis with deafness (ATP6V1B1-related) (AR) NM_001692.3	ATP6V1B1	Pan-ethnic	≤1 in 500	99%	Reduced
DOK7-related conditions (AR) NM_173660.4	DOK7	Pan-ethnic	1 in 115	99%	1 in 11400
Donnai-Barrow syndrome (AR) NM_004525.2	LRP2	Pan-ethnic	≤1 in 500	99%	Reduced
Dubin-Johnson syndrome (AR) NM_000392.4	ABCC2 *	Pan-ethnic	≤1 in 500	99%	Reduced
DUOX2-related conditions (AR) NM_014080.4	DUOX2 *	Pan-ethnic	1 in 58	91%	1 in 634
DYNC2H1-related conditions (AR) NM_001080463.1	DYNC2H1	Pan-ethnic	1 in 224	99%	1 in 22300
DYSF-related conditions (AR) NM_003494.3	DYSF	Pan-ethnic	1 in 311	99%	1 in 31000
Dyskeratosis congenita spectrum disorders (RTEL1-related) (AR) NM_001283009.1	RTEL1	Pan-ethnic	≤1 in 500	99%	Reduced
Dyskeratosis congenita spectrum disorders (TERT-related) (AR) NM_198253.2	TERT	Pan-ethnic	≤1 in 500	99%	Reduced
Dystrophic epidermolysis bullosa (AR) NM_000094.3	COL7A1	Pan-ethnic	1 in 370	97%	1 in 12300
Ehlers-Danlos syndrome, dermatosparaxis type (AR) NM_014244.4	ADAMTS2	Pan-ethnic	≤1 in 500	99%	Reduced
Ehlers-Danlos syndrome, kyphoscoliotic type (AR) NM_000302.3	PLOD1	Pan-ethnic	1 in 150	99%	1 in 14900
Ellis-van Creveld syndrome (EVC-related) (AR) NM_153717.2	EVC	Pan-ethnic	1 in 220	99%	1 in 21900
Epidermolysis bullosa with pyloric atresia (ITGB4-related) (AR) NM_001005731.2	ITGB4	Pan-ethnic	1 in 393	99%	1 in 39200
Epimerase deficiency galactosemia (AR) NM_000403.3	GALE *	Pan-ethnic	≤1 in 500	99%	Reduced
ERCC2-related conditions (AR) NM_000400.3	ERCC2	Pan-ethnic	≤1 in 500	99%	Reduced
Ethylmalonic encephalopathy (AR) NM_014297.3	ETHE1	Pan-ethnic	≤1 in 500	99%	Reduced
EVC2-related conditions (AR) NM_147127.4	EVC2	Pan-ethnic	1 in 199	99%	1 in 19800
Familial chylomicronemia syndrome (AR) NM_000237.2	LPL	Pan-ethnic	≤1 in 500	99%	Reduced
Familial dysautonomia (AR) NM_003640.3	ELP1	Pan-ethnic	≤1 in 500	99%	Reduced
Familial hemophagocytic lymphohistiocytosis type 2 (AR) NM_001083116.1	PRF1	Pan-ethnic	1 in 177	99%	1 in 17600
Familial hemophagocytic lymphohistiocytosis type 3 (AR) NM_199242.2	UNC13D	Pan-ethnic	1 in 177	93%	1 in 2515
Familial hemophagocytic lymphohistiocytosis type 4 (AR) NM_003764.3	STX11	Pan-ethnic	1 in 224	99%	1 in 22300
Familial hemophagocytic lymphohistiocytosis type 5 (AR) NM_006949.3	STXBP2	Pan-ethnic	1 in 224	99%	1 in 22300
Familial hypercholesterolemia (LDLR-related) (AD) NM_000527.4	LDLR	Pan-ethnic	1 in 250	99%	1 in 24900
Familial hypercholesterolemia (LDLRAP1-related) (AR) NM_015627.2	LDLRAP1	Pan-ethnic	≤1 in 500	99%	Reduced
Fanconi anemia type A (AR) NM_000135.2	FANCA	Pan-ethnic	1 in 345	99%	1 in 34400
Fanconi anemia type C (AR) NM_000136.2	FANCC	Pan-ethnic	1 in 417	99%	1 in 41600
Fanconi anemia type D2 (AR) NM_033084.3	FANCD2 *	Pan-ethnic	≤1 in 500	94%	Reduced

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Fanconi anemia type E (AR) NM_021922.2	FANCE	Pan-ethnic	≤1 in 500	99%	Reduced
Fanconi anemia type G (AR) NM_004629.1	FANCG	Pan-ethnic	≤1 in 500	99%	Reduced
Fanconi anemia type I (AR) NM_001113378.1	FANCI	Pan-ethnic	≤1 in 500	99%	Reduced
Fanconi anemia type L (AR) NM_018062.3	FANCL *	Pan-ethnic	≤1 in 500	99%	Reduced
FH-related conditions (AR) NM_000143.3	FH *	Pan-ethnic	≤1 in 500	99%	Reduced
FKBP10-related conditions (AR) NM_021939.3	FKBP10	Pan-ethnic	≤1 in 500	99%	Reduced
Foveal hypoplasia (SLC38A8-related) (AR) NM_001080442.2	SLC38A8	Pan-ethnic	≤1 in 500	99%	Reduced
FOXN1-related conditions (AR) NM_003593.2	FOXN1	Pan-ethnic	≤1 in 500	99%	Reduced
Fraser syndrome (FRAS1-related) (AR) NM_025074.6	FRAS1	Pan-ethnic	1 in 316	99%	1 in 31500
Fraser syndrome (FREM2-related) (AR) NM_207361.5	FREM2	Pan-ethnic	≤1 in 500	99%	Reduced
Fraser syndrome (GRIP1-related) (AR) NM_021150.3	GRIP1	Pan-ethnic	1 in 447	99%	1 in 44600
Fructose-1,6-bisphosphatase deficiency (AR) NM_000507.3	FBP1	Pan-ethnic	≤1 in 500	99%	Reduced
Fucosidosis (AR) NM_000147.4	FUCA1	Pan-ethnic	≤1 in 500	99%	Reduced
Galactokinase deficiency galactosemia (AR) NM_000154.1	GALK1	Pan-ethnic	1 in 122	99%	1 in 12100
Galactosemia (GALT-related) (AR) NM_000155.3	GALT	Pan-ethnic	1 in 100	99%	1 in 9900
Galactosialidosis (AR) NM_000308.3	CTSA	Pan-ethnic	≤1 in 500	99%	Reduced
GATM-related conditions (AR) NM_001482.2	GATM	Pan-ethnic	≤1 in 500	99%	Reduced
GBA-related conditions including Gaucher disease (AR) NM_001005741.2	GBA *	Ashkenazi Jewish Pan-ethnic	1 in 15 1 in 158	94% 72%	1 in 234 1 in 561
GET-related conditions (AR) NM_000158.3	GBE1	Pan-ethnic	1 in 387	99%	1 in 38600
GCH1-related conditions (AR) NM_000161.2	GCH1	Pan-ethnic	≤1 in 500	99%	Reduced
GDF5-related conditions (AR) NM_000557.4	GDF5	Pan-ethnic	≤1 in 500	99%	Reduced
Geroderma osteodysplastica (AR) NM_152281.2	GORAB	Pan-ethnic	≤1 in 500	99%	Reduced
GHR-related conditions (AR) NM_000163.4	GHR *	Pan-ethnic	≤1 in 500	98%	Reduced
GJB2-related conditions (AR) NM 004004.5	GJB2	Pan-ethnic	1 in 50	99%	1 in 4900
GLB1-related conditions (AR) NM_000404.2	GLB1	Pan-ethnic	1 in 158	99%	1 in 15700
GLE1-related conditions (AR) NM_001003722.1	GLE1	Pan-ethnic	≤1 in 500	99%	Reduced
Glutaric acidemia type I (AR) NM_000159.3	GCDH	Pan-ethnic	1 in 87	99%	1 in 8600
Glutaric acidemia type IIA (AR) NM_000126.3	ETFA	Pan-ethnic	≤1 in 500	99%	Reduced
Glutaric acidemia type IIB (AR) NM_001985.2	ETFB	Pan-ethnic	≤1 in 500	99%	Reduced
Glutaric acidemia type IIC (AR) NM_004453.3	ETFDH	Pan-ethnic	1 in 250	99%	1 in 24900
Glutathione synthetase deficiency (AR) NM_000178.2	GSS	Pan-ethnic	≤1 in 500	99%	Reduced

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Glycine encephalopathy (AMT-related) (AR) NM_000481.3	AMT	Pan-ethnic	1 in 325	99%	1 in 32400
Glycine encephalopathy (GLDC-related) (AR) NM_000170.2	GLDC	Pan-ethnic	1 in 165	99%	1 in 16400
Glycogen storage disease type Ia (AR) NM_000151.3	G6PC	Pan-ethnic	1 in 177	95%	1 in 3520
Glycogen storage disease type II (Pompe disease) (AR) NM_000152.3	GAA	Pan-ethnic	1 in 100	99%	1 in 9900
Glycogen storage disease type III (AR) NM_000642.2	AGL	Pan-ethnic	1 in 159	95%	1 in 3160
Glycogen storage disease type IXb (AR) NM_000293.2	РНКВ	Pan-ethnic	≤1 in 500	99%	Reduced
Glycogen storage disease type IXc (AR) NM_000294.2	PHKG2	Pan-ethnic	≤1 in 500	99%	Reduced
Glycogen storage disease type V (AR) NM_005609.3	PYGM	Pan-ethnic	1 in 171	99%	1 in 17000
Glycogen storage disease type VII (AR) NM_000289.5	PFKM	Pan-ethnic	≤1 in 500	99%	Reduced
GM3 synthase deficiency (AR) NM_003896.3	ST3GAL5	Pan-ethnic	≤1 in 500	99%	Reduced
GNE-related conditions (AR) NM_001128227.2	GNE *	Pan-ethnic	1 in 179	99%	1 in 17800
GNPTAB-related conditions (AR) NM_024312.4	GNPTAB	Pan-ethnic	1 in 200	99%	1 in 19900
Guanidinoacetate methyltransferase deficiency (AR) NM_000156.5	GAMT	Pan-ethnic	≤1 in 500	99%	Reduced
GUCY2D-related conditions (AR) NM_000180.3	GUCY2D	Pan-ethnic	1 in 204	99%	1 in 20300
Gyrate atrophy of the choroid and retina (AR) NM_000274.3	OAT *	Pan-ethnic	≤1 in 500	99%	Reduced
HADHA-related conditions (AR) NM_000182.4	HADHA	Pan-ethnic	1 in 350	99%	1 in 34900
HBB-related hemoglobinopathies (AR) NM_000518.4	НВВ	Pan-ethnic	1 in 49	99%	1 in 4800
Heme oxygenase 1 deficiency (AR) NM_002133.2	HMOX1	Pan-ethnic	≤1 in 500	99%	Reduced
Hemolytic anemia, CD59-mediated (AR) NM_203330.2	CD59	Pan-ethnic	≤1 in 500	99%	Reduced
Hereditary fructose intolerance (AR) NM_000035.3	ALDOB	Pan-ethnic	1 in 122	99%	1 in 12100
Hereditary hemochromatosis type 2 (HAMP-related) (AR) NM_021175.2	НАМР	Pan-ethnic	≤1 in 500	99%	Reduced
Hereditary hemochromatosis type 2 (HJV-related) (AR) NM_213653.3	HJV	Pan-ethnic	≤1 in 500	99%	Reduced
Hereditary hemochromatosis type 3 (AR) NM_003227.3	TFR2	Pan-ethnic	≤1 in 500	99%	Reduced
Hermansky-Pudlak syndrome type 1 (AR) NM_000195.4	HPS1	Pan-ethnic	≤1 in 500	99%	Reduced
Hermansky-Pudlak syndrome type 3 (AR) NM_032383.4	HPS3	Pan-ethnic	≤1 in 500	99%	Reduced
Hermansky-Pudlak syndrome type 4 (AR) NM_022081.5	HPS4	Pan-ethnic	≤1 in 500	99%	Reduced
Hermansky-Pudlak syndrome type 5 (AR) NM_181507.1	HPS5	Pan-ethnic	≤1 in 500	99%	Reduced
Hermansky-Pudlak syndrome type 6 (AR) NM_024747.5	HPS6	Pan-ethnic	≤1 in 500	99%	Reduced
Hermansky-Pudlak syndrome type 8 (AR) NM_212550.4	BLOC1S3	Pan-ethnic	≤1 in 500	99%	Reduced
Hermansky-Pudlak syndrome type 9 (AR) NM_012388.3	BLOC1S6	Pan-ethnic	≤1 in 500	99%	Reduced
HGSNAT-related conditions (AR) NM_152419.2	HGSNAT	Pan-ethnic	≤1 in 500	99%	Reduced



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Holocarboxylase synthetase deficiency (AR) NM_000411.6	HLCS	Pan-ethnic	1 in 224	99%	1 in 22300
Homocystinuria due to cobalamin E deficiency (AR) NM_002454.2	MTRR	Pan-ethnic	≤1 in 500	99%	Reduced
Homocystinuria due to cobalamin G deficiency (AR) NM_000254.2	MTR	Pan-ethnic	≤1 in 500	99%	Reduced
Homocystinuria due to cystathionine beta-synthase deficiency (AR) NM_000071.2	CBS	Pan-ethnic	1 in 224	99%	1 in 22300
Homocystinuria due to MTHFR deficiency (AR) NM_005957.4	MTHFR *	Pan-ethnic	≤1 in 500	99%	Reduced
HSD17B4-related conditions (AR) NM_000414.3	HSD17B4	Pan-ethnic	1 in 158	99%	1 in 15700
Hydrolethalus syndrome type 1 (AR) NM_145014.2	HYLS1	Pan-ethnic	≤1 in 500	99%	Reduced
Hyper-IgM immunodeficiency (CD40-related) (AR) NM_001250.5	CD40	Pan-ethnic	≤1 in 500	99%	Reduced
Hyperornithinemia-hyperammonemia-homocitrullinuria syndrome (AR) NM_014252.3	SLC25A15	Pan-ethnic	≤1 in 500	99%	Reduced
Hyperphosphatemic familial tumoral calcinosis (GALNT3-related) (AR) NM_004482.3	GALNT3	Pan-ethnic	≤1 in 500	99%	Reduced
Hypomyelinating leukodystrophy-12 (AR) NM_021729.5	VPS11	Pan-ethnic	≤1 in 500	99%	Reduced
Hypophosphatasia (AR) NM_000478.5	ALPL	Pan-ethnic	1 in 150	95%	1 in 2980
Ichthyosis prematurity syndrome (AR) NM_005094.3	SLC27A4	Pan-ethnic	≤1 in 500	99%	Reduced
IGHMBP2-related conditions (AR) NM_002180.2	IGHMBP2	Pan-ethnic	≤1 in 500	99%	Reduced
IKBKB-related conditions (AR) NM_001556.2	ІКВКВ	Pan-ethnic	≤1 in 500	99%	Reduced
Imerslund-Gräsbeck syndrome (AR) NM_030943.3	AMN *	Pan-ethnic	≤1 in 500	99%	Reduced
Immunodeficiency-centromeric instability-facial anomalies syndrome 1 (AR) NM_006892.3	DNMT3B	Pan-ethnic	≤1 in 500	99%	Reduced
Immunodeficiency-centromeric instability-facial anomalies syndrome 2 (AR) NM_014797.2	ZBTB24	Pan-ethnic	≤1 in 500	99%	Reduced
Isolated ectopia lentis (AR) NM_019032.5	ADAMTSL4	Pan-ethnic	≤1 in 500	99%	Reduced
Isovaleric acidemia (AR) NM_002225.3	IVD	Pan-ethnic	1 in 250	99%	1 in 24900
ITGB3-related conditions (AR) NM_000212.2	ITGB3	Pan-ethnic	≤1 in 500	99%	Reduced
Johanson-Blizzard syndrome (AR) NM_174916.2	UBR1	Pan-ethnic	1 in 250	99%	1 in 24900
Joubert syndrome and related disorders (MKS1-related) (AR) NM_017777.3	MKS1	Pan-ethnic	1 in 260	95%	1 in 5180
Joubert syndrome and related disorders (RPGRIP1L- related) (AR) NM_015272.2	RPGRIP1L	Pan-ethnic	1 in 259	95%	1 in 5160
Joubert syndrome and related disorders (TMEM216-related) (AR) NM_001173990.2	TMEM216	Pan-ethnic	≤1 in 500	99%	Reduced
Junctional epidermolysis bullosa (LAMC2-related) (AR) NM_005562.2	LAMC2	Pan-ethnic	≤1 in 500	99%	Reduced
Junctional epidermolysis bullosa with pyloric atresia (ITGA6-related) (AR) NM_000210.3	ITGA6	Pan-ethnic	≤1 in 500	99%	Reduced

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KCNJ11-related conditions (AR) NM_000525.3	KCNJ11	Pan-ethnic	≤1 in 500	99%	Reduced
Krabbe disease (AR) NM_000153.3	GALC *	Pan-ethnic	1 in 158	99%	1 in 15700
LAMA2-related muscular dystrophy (AR) NM_000426.3	LAMA2	Pan-ethnic	1 in 87	99%	1 in 8600
LAMA3-related conditions (AR) NM_000227.4	LAMA3	Pan-ethnic	≤1 in 500	99%	Reduced
LAMB3-related conditions (AR) NM_000228.2	LAMB3	Pan-ethnic	1 in 317	99%	1 in 31600
Leber congenital amaurosis 5 (AR) NM_181714.3	LCA5	Pan-ethnic	≤1 in 500	97%	Reduced
Leukoencephalopathy with vanishing white matter (EIF2B1-related) (AR) NM_001414.3	EIF2B1	Pan-ethnic	≤1 in 500	99%	Reduced
Leukoencephalopathy with vanishing white matter (EIF2B2-related) (AR) NM_014239.3	EIF2B2	Pan-ethnic	≤1 in 500	99%	Reduced
Leukoencephalopathy with vanishing white matter (EIF2B3-related) (AR) NM_020365.4	EIF2B3	Pan-ethnic	≤1 in 500	99%	Reduced
Leukoencephalopathy with vanishing white matter (EIF2B4-related) (AR) NM_015636.3	EIF2B4	Pan-ethnic	≤1 in 500	99%	Reduced
Leukoencephalopathy with vanishing white matter (EIF2B5-related) (AR) NM_003907.2	EIF2B5	Pan-ethnic	≤1 in 500	99%	Reduced
LIG4 syndrome (AR) NM_002312.3	LIG4	Pan-ethnic	≤1 in 500	99%	Reduced
Limb-girdle muscular dystrophy (CAPN3-related) (AR) NM_000070.2	CAPN3	Pan-ethnic	1 in 134	99%	1 in 13300
Limb-girdle muscular dystrophy type 2C (AR) NM_000231.2	SGCG	Pan-ethnic	≤1 in 500	99%	Reduced
Limb-girdle muscular dystrophy type 2D (AR) NM_000023.2	SGCA	Pan-ethnic	≤1 in 500	99%	Reduced
Limb-girdle muscular dystrophy type 2E (AR) NM_000232.4	SGCB	Pan-ethnic	≤1 in 500	92%	Reduced
Limb-girdle muscular dystrophy type 2F (AR) NM_000337.5	SGCD	Pan-ethnic	≤1 in 500	99%	Reduced
Lipoid congenital adrenal hyperplasia (AR) NM_000349.2	STAR	Pan-ethnic	≤1 in 500	99%	Reduced
LRAT-related conditions (AR) NM_004744.4	LRAT	Pan-ethnic	1 in 296	99%	1 in 29500
Lysinuric protein intolerance (AR) NM_001126106.2	SLC7A7	Pan-ethnic	≤1 in 500	99%	Reduced
Lysosomal acid lipase deficiency (AR) NM_000235.3	LIPA	Pan-ethnic	1 in 359	94%	1 in 5967
Major histocompatibility complex class II deficiency (CIITA- related) (AR) NM_000246.3	CIITA	Pan-ethnic	≤1 in 500	99%	Reduced
Malonyl-CoA decarboxylase deficiency (AR) NM_012213.2	MLYCD	Pan-ethnic	≤1 in 500	99%	Reduced
Maple syrup urine disease type 1A (AR) NM_000709.3	BCKDHA	Pan-ethnic	1 in 373	99%	1 in 37200
Maple syrup urine disease type 1B (AR) NM_183050.2	BCKDHB	Pan-ethnic	1 in 346	99%	1 in 34500
Maple syrup urine disease type 2 (AR) NM_001918.3	DBT	Pan-ethnic	≤1 in 500	99%	Reduced
Medium-chain acyl-CoA dehydrogenase deficiency (AR) NM_000016.5	ACADM	Pan-ethnic	1 in 66	99%	1 in 6500
Medium/short-chain 3-hydroxyacyl-CoA dehydrogenase deficiency (AR) NM_005327.4	HADH	Pan-ethnic	≤1 in 500	99%	Reduced



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MEDNIK syndrome (AR) NM_001283.3	AP1S1	Pan-ethnic	≤1 in 500	99%	Reduced
Megalencephalic leukoencephalopathy with subcortical cysts 1 (AR) NM_015166.3	MLC1 *	Pan-ethnic	≤1 in 500	99%	Reduced
Metabolic crises with rhabdomyolysis, cardiac arrhythmias and neurodegeneration (AR) NM_152906.6	TANGO2	Pan-ethnic	≤1 in 500	99%	Reduced
Metachromatic leukodystrophy (ARSA-related) (AR) NM_000487.5	ARSA	Pan-ethnic	1 in 100	95%	1 in 1980
Methylmalonic acidemia (MCEE-related) (AR) NM_032601.3	MCEE	Pan-ethnic	≤1 in 500	99%	Reduced
Methylmalonic acidemia (MMAA-related) (AR) NM_172250.2	MMAA	Pan-ethnic	1 in 316	97%	1 in 10500
Methylmalonic acidemia (MMAB-related) (AR) NM_052845.3	ММАВ	Pan-ethnic	1 in 456	98%	1 in 22750
Methylmalonic acidemia (MUT-related) (AR) NM_000255.3	MUT	Pan-ethnic	1 in 204	96%	1 in 5075
MFSD8-related conditions (AR) NM_152778.2	MFSD8	Pan-ethnic	≤1 in 500	99%	Reduced
Microcephalic osteodysplastic primordial dwarfism type II (AR) NM_006031.5	PCNT	Pan-ethnic	≤1 in 500	99%	Reduced
Microcephaly, postnatal progressive, with seizures and brain atrophy (AR) NM_004268.4	MED17	Pan-ethnic	≤1 in 500	99%	Reduced
Mitochondrial complex I deficiency 1 (AR) NM_002495.3	NDUFS4	Pan-ethnic	≤1 in 500	99%	Reduced
Mitochondrial complex I deficiency 3 (AR) NM_024407.4	NDUFS7	Pan-ethnic	1 in 387	99%	1 in 38600
Mitochondrial complex I deficiency 4 (AR) NM_007103.3	NDUFV1	Pan-ethnic	1 in 387	99%	1 in 38600
Mitochondrial complex I deficiency 9 (AR) NM_004553.4	NDUFS6	Pan-ethnic	≤1 in 500	99%	Reduced
Mitochondrial complex I deficiency 10 (AR) NM_174889.4	NDUFAF2	Pan-ethnic	1 in 387	99%	1 in 38600
Mitochondrial complex I deficiency 16 (AR) NM_024120.4	NDUFAF5	Pan-ethnic	≤1 in 500	99%	Reduced
Mitochondrial complex I deficiency 19 (AR) NM_017547.3	FOXRED1	Pan-ethnic	1 in 376	99%	1 in 37500
Mitochondrial complex I deficiency 20/ACAD9 deficiency (AR) NM_014049.4	ACAD9	Pan-ethnic	≤1 in 500	99%	Reduced
Mitochondrial complex IV deficiency 6 (AR) NM_004376.6	COX15	Pan-ethnic	≤1 in 500	99%	Reduced
Mitochondrial complex IV deficiency 12 (AR) NM_001171155.1	PET100	Pan-ethnic	1 in 387	99%	1 in 38600
Mitochondrial complex IV deficiency / Leigh syndrome, French Canadian type (AR) NM_133259.3	LRPPRC	Pan-ethnic	≤1 in 500	99%	Reduced
Mitochondrial DNA depletion syndrome-2 (AR) NM_004614.4	TK2	Pan-ethnic	≤1 in 500	99%	Reduced
Mitochondrial neurogastrointestinal encephalomyopathy (AR) NM_001953.4	ТҮМР	Pan-ethnic	≤1 in 500	99%	Reduced
Mitochondrial trifunctional protein deficiency (HADHB- related) (AR) NM_000183.2	HADHB	Pan-ethnic	≤1 in 500	99%	Reduced
MKKS-related conditions (AR) NM_018848.3	MKKS	Pan-ethnic	≤1 in 500	99%	Reduced
Molybdenum cofactor deficiency (MOCS1-related) (AR) NM_001358530.2	MOCS1	Pan-ethnic	1 in 226	99%	1 in 22500
Molybdenum cofactor deficiency (MOCS2-related) (AR) NM_004531.4	MOCS2B	Pan-ethnic	≤1 in 500	99%	Reduced



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DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY	DETECTION RATE	RISK TO BE A CARRIER AFTER NEGATIVE RESULT
Molybdenum cofactor deficiency (MOCS2-related) (AR) NM_176806.3	MOCS2A	Pan-ethnic	≤1 in 500	99%	Reduced
MPL-related conditions (AR) NM_005373.2	MPL	Pan-ethnic	≤1 in 500	99%	Reduced
MPV17-related conditions (AR) NM_002437.4	MPV17	Pan-ethnic	≤1 in 500	99%	Reduced
Mucolipidosis type III gamma (AR) NM_032520.4	GNPTG	Pan-ethnic	≤1 in 500	99%	Reduced
Mucolipidosis type IV (AR) NM_020533.2	MCOLN1	Pan-ethnic	≤1 in 500	99%	Reduced
Mucopolysaccharidosis type I (AR) NM_000203.4	IDUA	Pan-ethnic	1 in 148	97%	1 in 4900
Mucopolysaccharidosis type IIIA (AR) NM_000199.3	SGSH	Pan-ethnic	1 in 215	99%	1 in 21400
Mucopolysaccharidosis type IIIB (AR) NM_000263.3	NAGLU	Pan-ethnic	1 in 224	99%	1 in 22300
Mucopolysaccharidosis type IIID (AR) NM_002076.3	GNS	Pan-ethnic	≤1 in 500	99%	Reduced
Mucopolysaccharidosis type IVA (AR) NM_000512.4	GALNS	Pan-ethnic	1 in 224	99%	1 in 22300
Mucopolysaccharidosis type IX (AR) NM_153281.1	HYAL1	Pan-ethnic	≤1 in 500	99%	Reduced
Mucopolysaccharidosis type VI (AR) NM_000046.3	ARSB	Pan-ethnic	1 in 250	99%	1 in 24900
Mucopolysaccharidosis type VII (AR) NM_000181.3	GUSB	Pan-ethnic	1 in 250	99%	1 in 24900
Mulibrey nanism (AR) NM_015294.4	TRIM37	Pan-ethnic	≤1 in 500	99%	Reduced
Multiple pterygium syndrome (AR) NM_005199.4	CHRNG	Pan-ethnic	≤1 in 500	99%	Reduced
Multiple sulfatase deficiency (AR) NM_182760.3	SUMF1	Pan-ethnic	≤1 in 500	99%	Reduced
Muscular dystrophy-dystroglycanopathy (FKRP-related) (AR) NM_024301.4	FKRP	Pan-ethnic	1 in 158	99%	1 in 15700
Muscular dystrophy-dystroglycanopathy (FKTN-related) (AR) NM_001079802.1	FKTN	Pan-ethnic	≤1 in 500	99%	Reduced
Muscular dystrophy-dystroglycanopathy (LARGE1-related) (AR) NM_004737.4	LARGE1	Pan-ethnic	≤1 in 500	99%	Reduced
Muscular dystrophy-dystroglycanopathy (POMT1-related) (AR) NM_007171.3	POMT1	Pan-ethnic	1 in 268	99%	1 in 26700
Muscular dystrophy-dystroglycanopathy (POMT2-related) (AR) NM_013382.5	POMT2	Pan-ethnic	1 in 371	99%	1 in 37000
Muscular dystrophy-dystroglycanopathy (RXYLT1-related) (AR) NM_014254.2	RXYLT1	Pan-ethnic	≤1 in 500	99%	Reduced
MUSK-related conditions (AR) NM_005592.3	MUSK	Pan-ethnic	1 in 447	99%	1 in 44600
MVK-related conditions (AR) NM_000431.3	MVK	Pan-ethnic	≤1 in 500	99%	Reduced
MYO7A-related conditions (AR) NM_000260.3	MYO7A	Pan-ethnic	1 in 200	95%	1 in 3980
Myopathy, lactic acidosis, and sideroblastic anemia 1 (AR) NM_025215.5	PUS1	Pan-ethnic	≤1 in 500	99%	Reduced
Myotonia congenita (AR) NM_000083.2	CLCN1	Pan-ethnic	1 in 112	99%	1 in 11100
N-acetylglutamate synthase deficiency (AR) NM_153006.2	NAGS	Pan-ethnic	≤1 in 500	99%	Reduced

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Nemaline myopathy 2 (AR) NM_001271208.1	NEB *	Pan-ethnic	1 in 158	95%	1 in 3140
Nephrogenic diabetes insipidus (AQP2-related) (AR) NM_000486.5	AQP2	Pan-ethnic	≤1 in 500	99%	Reduced
Nephronophthisis (INVS-related) (AR) NM_014425.3	INVS	Pan-ethnic	≤1 in 500	99%	Reduced
Nephronophthisis (NPHP1-related) (AR) NM_000272.3	NPHP1	Pan-ethnic	≤1 in 500	99%	Reduced
Neuronal ceroid lipofuscinosis type 1 (AR) NM_000310.3	PPT1	Pan-ethnic	1 in 199	98%	1 in 9900
Neuronal ceroid lipofuscinosis type 2 (AR) NM_000391.3	TPP1	Pan-ethnic	1 in 250	97%	1 in 8300
Neuronal ceroid lipofuscinosis type 5 (AR) NM_006493.2	CLN5	Pan-ethnic	≤1 in 500	99%	Reduced
Neuronal ceroid lipofuscinosis type 6 (AR) NM_017882.2	CLN6	Pan-ethnic	≤1 in 500	99%	Reduced
Neuronal ceroid lipofuscinosis type 8 (AR) NM_018941.3	CLN8	Pan-ethnic	≤1 in 500	99%	Reduced
Neuronal ceroid lipofuscinosis type 10 (AR) NM_001909.4	CTSD	Pan-ethnic	≤1 in 500	99%	Reduced
Niemann-Pick disease type C (NPC1-related) (AR) NM_000271.4	NPC1	Pan-ethnic	1 in 183	99%	1 in 18200
Niemann-Pick disease type C (NPC2-related) (AR) NM_006432.3	NPC2	Pan-ethnic	≤1 in 500	99%	Reduced
Niemann-Pick disease types A and B (AR) NM_000543.4	SMPD1	Pan-ethnic	1 in 250	95%	1 in 4980
Nijmegen breakage syndrome (AR) NM_002485.4	NBN	Pan-ethnic	≤1 in 500	99%	Reduced
Nonsyndromic deafness (LOXHD1-related) (AR) NM_144612.6	LOXHD1	Pan-ethnic	≤1 in 500	99%	Reduced
Nonsyndromic deafness (MYO15A-related) (AR) NM_016239.3	MYO15A	Pan-ethnic	≤1 in 500	99%	Reduced
Nonsyndromic deafness (OTOA-related) (AR) NM_144672.3	OTOA *	Pan-ethnic	≤1 in 500	88%	Reduced
Nonsyndromic deafness (SYNE4-related) (AR) NM_001039876.2	SYNE4	Pan-ethnic	≤1 in 500	99%	Reduced
Nonsyndromic deafness (TMC1-related) (AR) NM_138691.2	TMC1	Pan-ethnic	≤1 in 500	99%	Reduced
Nonsyndromic deafness (TMPRSS3-related) (AR) NM_024022.2	TMPRSS3	Pan-ethnic	≤1 in 500	99%	Reduced
Nonsyndromic intellectual disability (CC2D1A-related) (AR) NM_017721.5	CC2D1A	Pan-ethnic	≤1 in 500	99%	Reduced
NR2E3-related conditions (AR) NM_014249.3	NR2E3	Pan-ethnic	≤1 in 500	99%	Reduced
NSMCE3 deficiency (AR) NM_138704.3	NSMCE3	Pan-ethnic	≤1 in 500	99%	Reduced
Oculocutaneous albinism type 3 (AR) NM_000550.2	TYRP1	Pan-ethnic	≤1 in 500	99%	Reduced
Oculocutaneous albinism type 4 (AR) NM_016180.4	SLC45A2	Pan-ethnic	1 in 158	99%	1 in 15700
Oculocutaneous albinism types 1A and 1B (AR) NM_000372.4	TYR *	Pan-ethnic	1 in 100	97%	1 in 3300
OPA3-related conditions (AR) NM_025136.3	OPA3	Pan-ethnic	≤1 in 500	99%	Reduced
Osteogenesis imperfecta (BMP1-related) (AR) NM_006129.4	BMP1	Pan-ethnic	≤1 in 500	99%	Reduced
Osteogenesis imperfecta (CRTAP-related) (AR) NM_006371.4	CRTAP	Pan-ethnic	≤1 in 500	99%	Reduced
Osteogenesis imperfecta (P3H1-related) (AR) NM_022356.3	P3H1	Pan-ethnic	≤1 in 500	99%	Reduced
Osteopetrosis (TCIRG1-related) (AR) NM_006019.3	TCIRG1	Pan-ethnic	1 in 317	99%	1 in 31600



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OSTM1 deficiency associated osteopetrosis (AR) NM_014028.3	OSTM1	Pan-ethnic	≤1 in 500	99%	Reduced
OTOF-related conditions (AR) NM_194248.2	OTOF	Pan-ethnic	≤1 in 500	99%	Reduced
Pantothenate kinase-associated neurodegeneration (AR) NM_153638.2	PANK2	Pan-ethnic	1 in 289	99%	1 in 28800
Parkinson disease 15 (AR) NM_012179.3	FBXO7	Pan-ethnic	≤1 in 500	99%	Reduced
PEX5-related conditions (AR) NM_001131025.1	PEX5	Pan-ethnic	≤1 in 500	99%	Reduced
PEX7-related conditions (AR) NM_000288.3	PEX7	Pan-ethnic	1 in 157	99%	1 in 15600
PGM3-congenital disorder of glycosylation (AR) NM_001199917.1	PGM3	Pan-ethnic	≤1 in 500	99%	Reduced
Phenylalanine hydroxylase deficiency (AR) NM_000277.1	РАН	Pan-ethnic	1 in 58	99%	1 in 5700
Phosphoglycerate dehydrogenase deficiency (AR) NM_006623.3	PHGDH	Pan-ethnic	≤1 in 500	99%	Reduced
PIGN-congenital disorder of glycosylation (AR) NM_176787.4	PIGN	Pan-ethnic	≤1 in 500	99%	Reduced
PJVK-related conditions (AR) NM_001042702.3	DFNB59	Pan-ethnic	≤1 in 500	99%	Reduced
PLA2G6-related conditions (AR) NM_003560.2	PLA2G6	Pan-ethnic	≤1 in 500	99%	Reduced
PLEKHG5-related conditions (AR) NM_020631.4	PLEKHG5	Pan-ethnic	≤1 in 500	99%	Reduced
POLG-related conditions (AR) NM_002693.2	POLG	Pan-ethnic	1 in 113	95%	1 in 2240
Polycystic kidney disease (PKHD1-related) (AR) NM_138694.3	PKHD1 *	Pan-ethnic	1 in 70	99%	1 in 6900
Polymicrogyria (ADGRG1-related) (AR) NM_005682.6	ADGRG1	Pan-ethnic	≤1 in 500	99%	Reduced
POMGNT1-related conditions (AR) NM_017739.3	POMGNT1	Pan-ethnic	≤1 in 500	99%	Reduced
Pontocerebellar hypoplasia (TSEN54-related) (AR) NM_207346.2	TSEN54	Pan-ethnic	≤1 in 500	99%	Reduced
Pontocerebellar hypoplasia type 1B (AR) NM_016042.3	EXOSC3	Pan-ethnic	≤1 in 500	99%	Reduced
Pontocerebellar hypoplasia type 2D (AR) NM_016955.3	SEPSECS	Pan-ethnic	≤1 in 500	99%	Reduced
Pontocerebellar hypoplasia type 6 (AR) NM_020320.3	RARS2	Pan-ethnic	≤1 in 500	99%	Reduced
Primary carnitine deficiency (AR) NM_003060.3	SLC22A5	Pan-ethnic	1 in 71	99%	1 in 7000
Primary ciliary dyskinesia (CCDC39-related) (AR) NM_181426.1	CCDC39	Pan-ethnic	1 in 211	99%	1 in 21000
Primary ciliary dyskinesia (CCDC103-related) (AR) NM_213607.2	CCDC103	Pan-ethnic	1 in 316	99%	1 in 31500
Primary ciliary dyskinesia (DNAH5-related) (AR) NM_001369.2	DNAH5	Pan-ethnic	1 in 109	99%	1 in 10800
Primary ciliary dyskinesia (DNAH11-related) (AR) NM_001277115.1	DNAH11	Pan-ethnic	1 in 211	99%	1 in 21000
Primary ciliary dyskinesia (DNAI1-related) (AR) NM_012144.3	DNAI1	Pan-ethnic	1 in 250	99%	1 in 24900
Primary ciliary dyskinesia (DNAI2-related) (AR) NM_023036.4	DNAI2	Pan-ethnic	1 in 354	99%	1 in 35300
Primary hyperoxaluria type 1 (AR) NM_000030.2	AGXT	Pan-ethnic	1 in 135	99%	1 in 13400
Primary hyperoxaluria type 2 (AR) NM_012203.1	GRHPR	Pan-ethnic	≤1 in 500	99%	Reduced
Primary hyperoxaluria type 3 (AR) NM_138413.3	HOGA1	Pan-ethnic	1 in 354	99%	1 in 35300

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Primary microcephaly (MCPH1-related) (AR) NM_024596.4	МСРН1	Pan-ethnic	≤1 in 500	99%	Reduced
Progressive early-onset encepahlopathy with brain atrophy and thin corpus callosum (PEBAT) (AR) NM_005993.4	TBCD	Pan-ethnic	≤1 in 500	99%	Reduced
Progressive pseudorheumatoid dysplasia (AR) NM_003880.3	WISP3	Pan-ethnic	≤1 in 500	99%	Reduced
Prolidase deficiency (AR) NM_000285.3	PEPD	Pan-ethnic	≤1 in 500	99%	Reduced
Propionic acidemia (PCCA-related) (AR) NM_000282.3	PCCA	Pan-ethnic	1 in 224	96%	1 in 5575
Propionic acidemia (PCCB-related) (AR) NM_000532.4	РССВ	Pan-ethnic	1 in 224	99%	1 in 22300
PSAP-related conditions (AR) NM_002778.3	PSAP	Pan-ethnic	≤1 in 500	99%	Reduced
Pycnodysostosis (AR) NM_000396.3	СТЅК	Pan-ethnic	1 in 438	99%	1 in 43700
Pyridoxal 5'-phosphate-dependent epilepsy (AR) NM_018129.3	PNPO	Pan-ethnic	≤1 in 500	99%	Reduced
Pyridoxine-dependent epilepsy (ALDH7A1-related) (AR) NM_001182.4	ALDH7A1	Pan-ethnic	1 in 127	99%	1 in 12600
Pyruvate carboxylase deficiency (AR) NM_000920.3	PC	Pan-ethnic	1 in 250	95%	1 in 4980
Pyruvate dehydrogenase complex deficiency (PDHB- related) (AR) NM_000925.3	PDHB	Pan-ethnic	≤1 in 500	99%	Reduced
RAPSN-related conditions (AR) NM_005055.4	RAPSN	Pan-ethnic	1 in 283	99%	1 in 28200
RDH12-related conditions (AR) NM_152443.2	RDH12	Pan-ethnic	1 in 460	99%	1 in 45900
Refsum disease (PHYH-related) (AR) NM_006214.3	РНҮН	Pan-ethnic	≤1 in 500	99%	Reduced
Retinitis pigmentosa 25 (AR) NM_001142800.1	EYS *	Pan-ethnic	1 in 129	99%	1 in 12800
Retinitis pigmentosa 28 (AR) NM_001201543.1	FAM161A	Pan-ethnic	1 in 289	99%	1 in 28800
Retinitis pigmentosa 36 (AR) NM_001077620.2	PRCD	Pan-ethnic	1 in 296	99%	1 in 29500
Retinitis pigmentosa 62 (AR) NM_001242957.2	МАК	Pan-ethnic	1 in 274	99%	1 in 27300
Rhizomelic chondrodysplasia punctata type 2 (AR) NM_014236.3	GNPAT	Pan-ethnic	≤1 in 500	99%	Reduced
Rhizomelic chondrodysplasia punctata type 3 (AR) NM_003659.3	AGPS	Pan-ethnic	≤1 in 500	99%	Reduced
RLBP1-related conditions (AR) NM_000326.4	RLBP1	Pan-ethnic	1 in 296	99%	1 in 29500
Roberts syndrome (AR) NM_001017420.2	ESCO2	Pan-ethnic	≤1 in 500	99%	Reduced
RPE65-related conditions (AR) NM_000329.2	RPE65	Pan-ethnic	1 in 228	99%	1 in 22700
RYR1-related conditions (AR) NM_000540.2	RYR1	Pan-ethnic	≤1 in 500	99%	Reduced
SAMD9-related conditions (AR) NM_017654.3	SAMD9	Pan-ethnic	≤1 in 500	99%	Reduced
Sandhoff disease (AR) NM_000521.3	НЕХВ	Pan-ethnic	1 in 180	99%	1 in 17900
Schimke immuno-osseous dysplasia (AR) NM_014140.3	SMARCAL1	Pan-ethnic	≤1 in 500	99%	Reduced
Seckel syndrome (CEP152-related) (AR) NM_014985.3	CEP152	Pan-ethnic	≤1 in 500	99%	Reduced
Sepiapterin reductase deficiency (AR) NM_003124.4	SPR	Pan-ethnic	≤1 in 500	99%	Reduced



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Severe combined immunodeficiency due to CD3-delta deficiency (AR) NM_000732.4	CD3D	Pan-ethnic	≤1 in 500	99%	Reduced
Severe combined immunodeficiency due to CD3-epsilon deficiency (AR) NM_000733.3	CD3E	Pan-ethnic	≤1 in 500	99%	Reduced
Severe combined immunodeficiency due to CD45 deficiency (AR) NM_002838.4	PTPRC *	Pan-ethnic	≤1 in 500	99%	Reduced
Severe combined immunodeficiency due to DCLRE1C (Artemis) deficiency (AR) NM_001033855.2	DCLRE1C	Pan-ethnic	≤1 in 500	99%	Reduced
Severe combined immunodeficiency due to IL7R-alpha deficiency (AR) NM_002185.3	IL7R	Pan-ethnic	1 in 348	99%	1 in 34700
Severe combined immunodeficiency due to JAK3 deficiency (AR) NM_000215.3	JAK3	Pan-ethnic	1 in 455	99%	1 in 45400
Severe combined immunodeficiency due to RAG1 deficiency (AR) NM_000448.2	RAG1	Pan-ethnic	1 in 301	99%	1 in 30000
Severe combined immunodeficiency due to RAG2 deficiency (AR) NM_000536.3	RAG2	Pan-ethnic	≤1 in 500	99%	Reduced
Severe congenital neutropenia due to G6PC3 deficiency (AR) NM_138387.3	G6PC3	Pan-ethnic	≤1 in 500	99%	Reduced
Severe congenital neutropenia due to HAX1 deficiency (AR) NM_006118.3	HAX1	Pan-ethnic	≤1 in 500	99%	Reduced
Severe congenital neutropenia due to VPS45 deficiency (AR) NM_007259.4	VPS45	Pan-ethnic	≤1 in 500	99%	Reduced
Sialic acid storage diseases (AR) NM_012434.4	SLC17A5	Pan-ethnic	≤1 in 500	99%	Reduced
Sialidosis (AR) NM_000434.3	NEU1	Pan-ethnic	≤1 in 500	99%	Reduced
Sjögren-Larsson syndrome (AR) NM_000382.2	ALDH3A2	Pan-ethnic	≤1 in 500	99%	Reduced
SLC12A6-related conditions (AR) NM_133647.1	SLC12A6	Pan-ethnic	≤1 in 500	99%	Reduced
SLC26A2-related conditions (AR) NM_000112.3	SLC26A2	Pan-ethnic	1 in 158	95%	1 in 3140
SLC26A4-related conditions (AR) NM_000441.1	SLC26A4	Pan-ethnic	1 in 80	99%	1 in 7900
SLC37A4-related conditions (AR) NM_001164277.1	SLC37A4	Pan-ethnic	1 in 354	95%	1 in 7060
Smith-Lemli-Opitz syndrome (AR) NM_001360.2	DHCR7	Pan-ethnic	1 in 71	99%	1 in 7000
Spastic paraplegia type 15 (AR) NM_015346.3	ZFYVE26	Pan-ethnic	≤1 in 500	99%	Reduced
Spastic paraplegia type 49 (AR) NM_014844.3	TECPR2	Pan-ethnic	≤1 in 500	99%	Reduced
Spastic tetraplegia, thin corpus callosum, and progressive microcephaly (AR) NM_003038.4	SLC1A4	Pan-ethnic	≤1 in 500	99%	Reduced
SPG11-related conditions (AR) NM_025137.3	SPG11	Pan-ethnic	1 in 141	99%	1 in 14000
		African-American	1 in 59	83%	1 in 342
Spinal muscular atrophy (AR)		Ashkenazi Jewish	1 in 62	94%	1 in 1017
NM_000344.3	SMN1 *	Asian	1 in 50	93%	1 in 701
Carrier residual risks listed are for 2 copy SMN1 results. Carrier residual risk for >2 copies are 5- to 10-fold lower.		Caucasian	1 in 45	95%	1 in 880
		Hispanic Ban othnic	1 in 48	94%	1 in 784
		Pan-ethnic	1 in 49	94%	1 in 800

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Spinocerebellar ataxia (ANO10-related) (AR) NM_018075.3	ANO10 *	Pan-ethnic	≤1 in 500	99%	Reduced
Spondylocostal dysostosis (DLL3-related) (AR) NM_016941.3	DLL3	Pan-ethnic	1 in 350	99%	1 in 34900
Spondylocostal dysostosis (MESP2-related) (AR) NM_001039958.1	MESP2	Pan-ethnic	1 in 224	99%	1 in 22300
Steel syndrome (AR) NM_032888.3	COL27A1	Pan-ethnic	≤1 in 500	99%	Reduced
Steroid 5-alpha-reductase deficiency (AR) NM_000348.3	SRD5A2	Pan-ethnic	≤1 in 500	99%	Reduced
Stüve-Wiedemann syndrome (AR) NM_002310.5	LIFR *	Pan-ethnic	≤1 in 500	99%	Reduced
Sulfite oxidase deficiency (AR) NM_000456.2	SUOX	Pan-ethnic	≤1 in 500	99%	Reduced
SURF1-related conditions (AR) NM_003172.3	SURF1	Pan-ethnic	1 in 128	99%	1 in 12700
Tay-Sachs disease (AR) NM_000520.4	HEXA	Pan-ethnic	1 in 250	99%	1 in 24900
TBCE-related conditions (AR) NM_003193.4	TBCE *	Pan-ethnic	≤1 in 500	99%	Reduced
Thiamine-responsive megaloblastic anemia (AR) NM_006996.2	SLC19A2	Pan-ethnic	≤1 in 500	99%	Reduced
Thyroid dyshormonogenesis (SLC5A5-related) (AR) NM_000453.2	SLC5A5	Pan-ethnic	≤1 in 500	99%	Reduced
Thyroid dyshormonogenesis (TG-related) (AR) NM_003235.4	TG *	Pan-ethnic	≤1 in 500	99%	Reduced
Thyroid dyshormonogenesis (TPO-related) (AR) NM_000547.5	ТРО	Pan-ethnic	1 in 129	99%	1 in 12800
TMEM67-related conditions (AR) NM_153704.5	TMEM67	Pan-ethnic	1 in 316	99%	1 in 31500
Transcobalamin II deficiency (AR) NM_000355.3	TCN2	Pan-ethnic	≤1 in 500	99%	Reduced
Transient infantile liver failure (AR) NM_018006.4	TRMU	Pan-ethnic	≤1 in 500	99%	Reduced
TREX1-related conditions (AR) NM_033629.4	TREX1	Pan-ethnic	≤1 in 500	99%	Reduced
Trichohepatoenteric syndrome (SKIV2L-related) (AR) NM_006929.4	SKIV2L	Pan-ethnic	≤1 in 500	99%	Reduced
Trichohepatoenteric syndrome (TTC37-related) (AR) NM_014639.3	TTC37	Pan-ethnic	≤1 in 500	99%	Reduced
TRIM32-related conditions (AR) NM_012210.3	TRIM32	Pan-ethnic	1 in 408	99%	1 in 40700
Trimethylaminuria (AR) NM_006894.6	FMO3	Pan-ethnic	≤1 in 500	99%	Reduced
Triple A syndrome (AR) NM_015665.5	AAAS	Pan-ethnic	≤1 in 500	99%	Reduced
TSHR-related conditions (AR) NM_000369.2	TSHR	Pan-ethnic	1 in 158	99%	1 in 15700
TULP1-related conditions (AR) NM_003322.4	TULP1	Pan-ethnic	1 in 296	99%	1 in 29500
Tyrosinemia type I (AR) NM_000137.2	FAH *	Pan-ethnic	1 in 125	95%	1 in 2480
Tyrosinemia type II (AR) NM_000353.2	TAT	Pan-ethnic	1 in 250	99%	1 in 24900
Tyrosinemia type III (AR) NM_002150.2	HPD	Pan-ethnic	≤1 in 500	99%	Reduced
USH1C-related conditions (AR) NM_005709.3	USH1C *	Pan-ethnic	1 in 353	90%	1 in 3521
USH2A-related conditions (AR) NM_206933.2	USH2A	Pan-ethnic	1 in 112	99%	1 in 11100
Very long-chain acyl-CoA dehydrogenase deficiency (AR) NM_000018.3	ACADVL	Pan-ethnic	1 in 100	99%	1 in 9900



This table is relevant to patient report RQ5172543 Issue date: 06/15/2023

DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY	DETECTION RATE	RISK TO BE A CARRIER AFTER NEGATIVE RESULT
Vici syndrome (AR) NM_020964.2	EPG5	Pan-ethnic	≤1 in 500	99%	Reduced
Vitamin D-dependent rickets type 1A (AR) NM_000785.3	CYP27B1	Pan-ethnic	≤1 in 500	99%	Reduced
Vitamin D-dependent rickets type 2A (AR) NM_001017535.1	VDR	Pan-ethnic	≤1 in 500	99%	Reduced
VPS53-related conditions (AR) NM_001128159.2	VPS53 *	Pan-ethnic	≤1 in 500	99%	Reduced
VRK1-related conditions (AR) NM_003384.2	VRK1	Pan-ethnic	≤1 in 500	99%	Reduced
VSX2-related conditions (AR) NM_182894.2	VSX2	Pan-ethnic	≤1 in 500	99%	Reduced
Warsaw syndrome (AR) NM_030653.3	DDX11 *	Pan-ethnic	≤1 in 500	15%	Reduced
Werner syndrome (AR) NM_000553.4	WRN *	Pan-ethnic	1 in 224	99%	1 in 22300
Wilson disease (AR) NM_000053.3	ATP7B	Pan-ethnic	1 in 90	98%	1 in 4450
WNT10A-related conditions (AR) NM_025216.2	WNT10A	Pan-ethnic	1 in 305	99%	1 in 30400
Wolcott-Rallison syndrome (AR) NM_004836.6	EIF2AK3	Pan-ethnic	≤1 in 500	99%	Reduced
Woodhouse-Sakati syndrome (AR) NM_025000.3	DCAF17	Pan-ethnic	≤1 in 500	99%	Reduced
Xeroderma pigmentosum complementation group A (AR) NM_000380.3	ХРА	Pan-ethnic	≤1 in 500	99%	Reduced
Xeroderma pigmentosum complementation group C (AR) NM_004628.4	XPC	Pan-ethnic	≤1 in 500	99%	Reduced
Xeroderma pigmentosum, variant type (AR) NM_006502.2	POLH	Pan-ethnic	≤1 in 500	99%	Reduced
Zellweger spectrum disorder (PEX1-related) (AR) NM_000466.2	PEX1 *	Pan-ethnic	1 in 144	99%	1 in 14300
Zellweger spectrum disorder (PEX2-related) (AR) NM_000318.2	PEX2	Pan-ethnic	≤1 in 500	99%	Reduced
Zellweger spectrum disorder (PEX6-related) (AR) NM_000287.3	PEX6	Pan-ethnic	1 in 294	99%	1 in 29300
Zellweger spectrum disorder (PEX10-related) (AR) NM_153818.1	PEX10	Pan-ethnic	≤1 in 500	94%	Reduced
Zellweger spectrum disorder (PEX12-related) (AR) NM_000286.2	PEX12	Pan-ethnic	1 in 409	99%	1 in 40800
Zellweger spectrum disorder (PEX13-related) (AR) NM_002618.3	PEX13	Pan-ethnic	≤1 in 500	99%	Reduced
Zellweger spectrum disorder (PEX16-related) (AR) NM_004813.2	PEX16	Pan-ethnic	≤1 in 500	99%	Reduced
Zellweger spectrum disorder (PEX26-related) (AR) NM_017929.5	PEX26	Pan-ethnic	≤1 in 500	99%	Reduced

Patient Information	n	Test Information		
Patient Name:	DONOR 14527	Ordering Physician:	Dr. James Kuan	horizon"
		Clinic Information:	Seattle Sperm Bank	natera carrier screen
Date Of Birth:				
Gender:	Male			CARRIER SCREENING REPORT
Ethnicity:	Other	Phone:	(206) 588-1484	
Patient ID:	N/A	Report Date:	04/16/2025	ABOUT THIS SCREEN: Horizon [™] is a carrier
Medical Record #:	N/A	Sample Collected:	04/07/2025	screen for specific autosomal recessive and X- linked diseases. This information can help
Collection Kit:	44378897-2-C	Sample Received:	04/08/2025	patients learn their risk of having a child with
Accession ID:	N/A	Sample Type:	Blood	specific genetic conditions.
Case File ID:	16381364			
				ORDER SELECTED: The Horizon Custom panel was ordered for this patient. Males are not screened for X-linked diseases

FINAL RESULTS SUMMARY:

Negative for 1 out of 1 diseases

No Pathogenic variants were detected in the genes that were screened. The patient's remaining carrier risk after the negative screening results is listed for each disease/gene on the Horizon website at https://www.natera.com/panel-option/h-all/. Please see the following pages of this report for a comprehensive list of all conditions included on this individual's screen.

Carrier screening is not diagnostic and may not detect all possible pathogenic variants in a given gene.

RECOMMENDATIONS

Individuals who would like to review their Horizon report with a Natera Laboratory Genetic Counselor may schedule a telephone genetic information session by calling 650-249-9090 or visiting naterasession.com. Clinicians with questions may contact Natera at 650-249-9090 or email support@natera.com.

Dianne Keen-Kim, Ph.D., FACMGO

The pre-analytic and post-analytic phases of this test were performed by NSTX, Inc., 13011 McCallen Pass, Building A Suite 110, Austin, TX 78753 (CLIA ID 45D2093704). This test was performed by Baylor Miraca Genetics, DBA Baylor Genetics, D



Patient Information

DONOR 14527 Patient Name:

Test Information Ordering Physician:

Clinic Information:

Dr. James Kuan

Seattle Sperm Bank



Date Of Birth: Case File ID:

16381364

Report Date:

04/16/2025

DISEASES SCREENED

Below is a list of all diseases screened and the result. Certain conditions have unique patient-specific numerical values, therefore, results for those conditions are formatted differently.

Autosomal Recessive

C CONGENITAL ADRENAL HYPERPLASIA, 21-HYDROXYLASE DEFICIENCY (CYP21A2) negative



Patient Information Patient Name:	DONOR 14527	Test Information Ordering Physician:	Dr. James Kuan
Date Of Birth:		Clinic Information:	Seattle Sperm Bank
Case File ID:	16381364	Report Date:	04/16/2025

Testing Methodology, Limitations, and Comments:

Next-generation sequencing (NGS)

Sequencing library prepared from genomic DNA isolated from a patient sample is enriched for targets of interest using standard hybridization capture protocols and PCR amplification (for targets specified below). NGS is then performed to achieve the standards of quality control metrics, including a minimum coverage of 99% of targeted regions at 20X sequencing depth. Sequencing data is aligned to human reference sequence, followed by deduplication, metric collection and variant calling (coding region +/- 20bp). Variants are then classified according to ACMGG/AMP standards of interpretation using publicly available databases including but not limited to ENSEMBL, HGMD Pro, ClinGen, ClinVar, 1000G, ESP and gnomAD. Variants predicted to be pathogenic or likely pathogenic for the specified diseases are reported. It should be noted that the data interpretation is based on our current understanding of the genes and variants at the time of reporting. Putative positive sequencing variants that do not meet internal quality standards or are within highly homologous regions are confirmed by Sanger sequencing or gene-specific long-range PCR as needed prior to reporting.

Copy Number Variant (CNV) analysis is limited to deletions involving two or more exons for all genes on the panel, in addition to specific known recurrent singleexon deletions. CNVs of small size may have reduced detection rate. This method does not detect gene inversions, single-exonic and sub-exonic deletions (unless otherwise specified), and duplications of all sizes (unless otherwise specified). Additionally, this method does not define the exact breakpoints of detected CNV events. Confirmation testing for copy number variation is performed by specific PCR, Multiplex Ligation-dependent Probe Amplification (MLPA), next generation sequencing, or other methodology.

This test may not detect certain variants due to local sequence characteristics, high/low genomic complexity, homologous sequence, or allele dropout (PCR-based assays). Variants within noncoding regions (promoter, 5'UTR, 3'UTR, deep intronic regions, unless otherwise specified), small deletions or insertions larger than 25bp, low-level mosaic variants, structural variants such as inversions, and/or balanced translocations may not be detected with this technology.

SPECIAL NOTES

For CYP21A2, targets were enriched using long-range PCR amplification, followed by next generation sequencing. Duplication analysis will only be performed and reported when c.955C>T (p.Q319*) is detected. Sequencing and CNV analysis may have reduced sensitivity, if variants result from complex rearrangements, in trans with a gene deletion, or CYP21A2 gene duplication on one chromosome and deletion on the other chromosome. This analysis cannot detect sequencing variants located on the CYP21A2 duplicated copy.

Variant Classification

Only pathogenic or likely pathogenic variants are reported. Other variants including benign variants, likely benign variants, variants of uncertain significance, or inconclusive variants identified during this analysis may be reported in certain circumstances. Our laboratory's variant classification criteria are based on the ACMG and internal guidelines and our current understanding of the specific genes. This interpretation may change over time as more information about a gene and/or variant becomes available. Natera and its lab partner(s) may reclassify variants at certain intervals but may not release updated reports without a specific request made to Natera by the ordering provider. Natera may disclose incidental findings if deemed clinically pertinent to the test performed.

Negative Results

A negative carrier screening result reduces the risk for a patient to be a carrier of a specific disease but does not completely rule out carrier status. Please visit https://www.natera.com/panel-option/h-all/ for a table of carrier rates, detection rates, residual risks and promised variants/exons per gene. Carrier rates before and after testing vary by ethnicity and assume a negative family history for each disease screened and the absence of clinical symptoms in the patient. Any patient with a family history for a specific genetic disease will have a higher carrier risk prior to testing and, if the disease-causing mutation in their family is not included on the test, their carrier risk would remain unchanged. Genetic counseling is recommended for patients with a family history of genetic disease so that risk figures based on actual family history can be determined and discussed along with potential implications for reproduction. Horizon carrier screening has been developed to identify the reproductive risks for monogenic inherited conditions. Even when one or both members of a couple screen negative for pathogenic variants in a specific gene, the disease risk for their offspring is not zero. There is still a low risk for the condition in their offspring due to a number of different mechanisms that are not detected by Horizon including, but not limited to, pathogenic variant(s) in the tested gene or in a different gene not included on Horizon, pathogenic variant(s) in an upstream regulator, uniparental disomy, de novo mutation(s), or digenic or polygenic inheritance.

Additional Comments

These analyses generally provide highly accurate information regarding the patient's carrier status. Despite this high level of accuracy, it should be kept in mind that there are many potential sources of diagnostic error, including misidentification of samples, polymorphisms, or other rare genetic variants that interfere with analysis. Families should understand that rare diagnostic errors may occur for these reasons.



15 Crawford St., STE 100 Needham, MA 02494 (p) 626-350-0537 (f) 626-454-1667 Lab Director: Arash Radfar M.D. CLIA: 22D0957540







Patient Information: 14527, Donor DOB: Sex: M MR#: Patient#: FT-PT8863560

Accession: FT-7218531 Test#: FT-TS14970682 Specimen Type: Blood (EDTA) Collected: Not Provided

FINAL RESULTS

<u>Accession:</u> N/A

Not Tested

Partner Information:

Physician: Kuan, James ATTN: SSB Genetics, Dept San Diego Sperm Bank 4915 25th Avenue NE, Ste 204W Seattle, WA 98105 Phone: (206) 588-1484 Laboratory: Fulgent Therapeutics LLC CAP#: 8042697 CLIA#: 05D2043189 Laboratory Director: Lawrence M. Weiss, MD Report Date: Oct 15,2024

TEST PERFORMED

No carrier mutations identified

Single Gene Carrier Screening: TNXB (1 Gene Panel: *TNXB*; gene sequencing with deletion and

(1 Gene Panel: *TNXB*; gene sequencing with deletion and duplication analysis)

INTERPRETATION:

Notes and Recommendations:

- No carrier mutations were identified in the submitted specimen. A negative result does not rule out the possibility of a genetic predisposition nor does it rule out any pathogenic mutations in areas not assessed by this test or in regions that were covered at a level too low to reliably assess. Also, it does not rule out mutations that are of the sort not queried by this test; see Methods and Limitations for more information. A negative result reduces, but does not eliminate, the chance to be a carrier for any condition included in this screen. Please see the supplemental table for details.
- This carrier screening test does not screen for all possible genetic conditions, nor for all possible mutations in every gene tested. This report does not include variants of uncertain significance; only variants classified as pathogenic or likely pathogenic at the time of testing, and considered relevant for reproductive carrier screening, are reported. Please see the gene specific notes for details. Please note that the classification of variants can change over time.
- 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.
- Genetic counseling is recommended. Available genetic counselors and additional resources can be found at the National Society of Genetic Counselors (NSGC; https://www.nsgc.org)







GENES TESTED:

Custom Beacon Preconception Carrier Screening Panel - Gene

This analysis was run using the Custom Beacon Preconception Carrier Screening Panel gene list. 1 genes were tested with 100.0% of targets sequenced at >20x coverage. For more gene-specific information and assistance with residual risk calculation, see the SUPPLEMENTAL TABLE.

TNXB

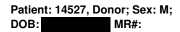
METHODS:

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, 100.00% and 100.00% 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. 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 sequencing 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.



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Gene Specific Notes and Limitations

<u>TNXB</u>: This gene is susceptible to significant pseudogene interference, particularly for exons 32-44 (NM_019105.6). Among these exons, copy number analysis is available for only exon 35. SNV analysis is available for only exons 35 and 40.

SIGNATURE:

- Gao

Dr. Harry Gao, DABMG, FACMG on 10/15/2024 Laboratory Director, Fulgent

DISCLAIMER:

This test was developed and its performance characteristics determined by Fulgent Therapeutics LLC CAP #8042697 CLIA #05D2043189; 4399 Santa Anita Ave., El Monte, CA, 91731. 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 by email at **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.

To view the supplemental table describing the carrier frequencies, detection rates, and residual risks associated with the genes on this test please visit the following link:

Beacon Expanded Carrier Screening Supplemental Table







Lab:EZ

14527, DONORSpecimen: EN361035R Requisition: 0001505Client #: 92037397 MAIL000 OLLIFFE, JEFFREY F SEATTLE SPERM BANKDOB:AGE:Collected:06/06/2023 / 11:10 PDT Received:SEATTLE SPERM BANKGender:MFasting: UCollected:06/06/2023 / 11:10 PDT Received:Attn: STE B214Phone:858.732.8500Received:06/07/2023 / 02:56 PDT Reported:Attn: STE B214Patient ID: 14527Reported:06/19/2023 / 13:45 PDTAttn: Attn: A	Patient Information	Specimen Information	Client Information
Health ID: X5730325316/18/17X	14527, DONOR DOB: AGE: Gender: M Fasting: U Phone: 858.732.8500	Specimen: EN361035R Requisition: 0001505 Collected: 06/06/2023 / 11:10 PDT Received: 06/07/2023 / 02:56 PDT	Client #: 92037397 MAIL000 OLLIFFE, JEFFREY F SEATTLE SPERM BANK Attn: STE B214 8950 VILLA LA JOLLA DR

COMMENTS: FASTING:UNKNOWN

Cytogenetic Report

CHROMOSOME ANALYSIS, BLOOD - 14596 CHROMOSOME ANALYSIS, BLOOD

Order ID: 23-253486 Specimen Type: Blood Clinical Indication: SDSB

RESULT: NORMAL MALE KARYOTYPE

INTERPRETATION:

Chromosome analysis revealed normal G-band patterns within the limits of standard cytogenetic analysis.

Please expect the results of any other concurrent study in a separate report.

NOMENCLATURE:

46,XY

ASSAY INFORMATION:

Method:	G-Band (Digital Analysis: MetaSyst
Cells Counted:	20
Band Level:	450
Cells Analyzed:	5
Cells Karyotyped:	3

This test does not address genetic disorders that cannot be detected by standard cytogenetic methods or rare events such as low level mosaicism or subtle rearrangements.

Morteza Hemmat, PhD, FACMG (800) NICHOLS-4307

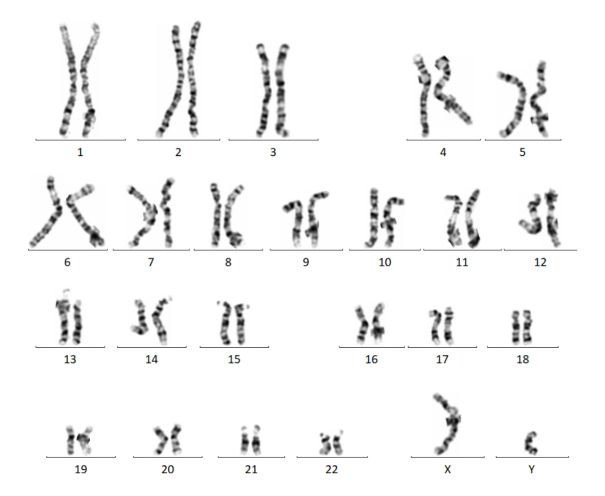
Electronic Signature: 6/19/2023 3:56 PM

CLIENT SERVICES: 866.697.8378





Patient Information	Specimen Information	Client Information
14527, DONOR	Specimen: EN361035R Collected: 06/06/2023 / 11:10 PDT	Client #: 92037397 OLLIFFE, JEFFREY F
DOB:AGE:Gender:MFasting: UPatient ID:14527Health ID:8573032531648478	Received: 06/07/2023 / 02:56 PDT Reported: 06/19/2023 / 13:45 PDT	



PERFORMING SITE: EZ QUEST DIAGNOSTICS/NICHOLS SJC, 33608 ORTEGA HWY, SAN JUAN CAPISTRANO, CA 92675-2042 Laboratory Director: IRINA MARAMICA,MD,PHD,MBA, CLIA: 05D0643352

14527, Donor	DOB:	Patient Report	labcor
Patient ID: Specimen ID: 157-253-3185-0	Age: Sex: Male	Account Number: 04087470 Ordering Physician: J OLLIFFE	
Date Collected: 06/06/2023	Date Received: 06/07/2023	Date Reported: 06/09/2023	Fasting: Not Given

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

Date Collected: 06/06/2023

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

Test	Current Resu	lt and Flag	Previous Result and Date	Units	Reference Interval
Glucose ⁰¹	102	High		mg/dL	70-99
Uric Acid ⁰¹	7.5			mg/dL	3.8-8.4
		The	erapeutic target for gout pa		
			Please note reference int	erval change	
BUN ⁰¹	15			mg/dL	6-20
Creatinine ⁰¹	1.03			mg/dL	0.76-1.27
eGFR	107			mL/min/1.73	>59
Calcium ⁰¹	10.6	High		mg/dL	8.7-10.2
Protein, Total ⁰¹	8.2			g/dL	6.0-8.5
Albumin ⁰¹	5.6	High		g/dL	4.1-5.2
Bilirubin, Total ⁰¹	1.2			mg/dL	0.0-1.2
Alkaline Phosphatase ⁰¹	97			IU/L	51-125
LDH ⁰¹	158			IU/L	121-224
AST (SGOT) ⁰¹	21			IU/L	0-40
ALT (SGPT) 01	11			IU/L	0-44
Cholesterol, Total ⁰¹	201	High		mg/dL	100-169
Triglycerides ⁰¹	128	High		mg/dL	0-89
HDL Cholesterol ⁰¹	61			mg/dL	>39
LDL Chol Calc (NIH)	117	High		mg/dL	0-109
LDL/HDL Ratio	1.9			ratio	0.0-3.6
Please Note: ⁰¹					
			LDL/H	DL Ratio	
				Men Women	

	LDL/HDL	Ratio	D
		Men	Women
1/2	Avg.Risk	1.0	1.5
	Avg.Risk	3.6	3.2
2X	Avg.Risk	6.2	5.0
3X	Avg.Risk	8.0	6.1

Hgb Fractionation by CE:02				
Hgb F ⁰²	0.0	%	0.0-2.0	
Hgb A ⁰²	97.0	%	96.4-98.8	
Hgb A2 02	3.0	%	1.8-3.2	
Hgb S ⁰²	0.0	%	0.0	
Interpretation: ⁰²	Normal hemoglobin present; no hemoglobin variant or beta thalassemia identified. Note: Alpha thalassemia may not be detected by the Hgb Fractionation Cascade panel. If alpha thalassemia is suspected, Labcorp offers Alpha-Thalassemia DNA Analysis (#511172).			
ABO Grouping ⁰¹	0			
Rh Factor ⁰¹	Positive Please note: Prior records f	or this patient's ABO / Rh type are not		

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Date Collected: 06/06/2023

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

available for additional verification.

.01					
CBC, Platelet Ct, and Diff ⁰¹					
WBC ⁰¹	7.1			x10E3/uL	3.4-10.8
RBC ⁰¹	5.71			x10E6/uL	4.14-5.80
Hemoglobin ⁰¹	17.0			g/dL	13.0-17.7
Hematocrit ⁰¹	50.3			%	37.5-51.0
MCV ⁰¹	88			fL	79-97
MCH ⁰¹	29.8			pg	26.6-33.0
MCHC ⁰¹	33.8			g/dL	31.5-35.7
RDW ⁰¹	12.5			%	11.6-15.4
Platelets ⁰¹	316			x10E3/uL	150-450
Neutrophils ⁰¹	64			%	Not Estab.
Lymphs ⁰¹	23			%	Not Estab.
Monocytes ⁰¹	9			%	Not Estab.
Eos ⁰¹	3			%	Not Estab.
Basos ⁰¹	1			%	Not Estab.
Neutrophils (Absolute) ⁰¹	4.5			x10E3/uL	1.4-7.0
Lymphs (Absolute) ⁰¹	1.6			x10E3/uL	0.7-3.1
Monocytes(Absolute) ⁰¹	0.6			x10E3/uL	0.1-0.9
Eos (Absolute) ⁰¹	0.2			x10E3/uL	0.0-0.4
Baso (Absolute) ⁰¹	0.1			x10E3/uL	0.0-0.2
Immature Granulocytes ⁰¹	0			%	Not Estab.
Immature Grans (Abs) ⁰¹	0.0			x10E3/uL	0.0-0.1
.01				-	
Urinalysis Gross Exam ⁰¹					
Specific Gravity ⁰¹	1.029				1.005-1.030
pH ⁰¹	5.5				5.0-7.5
Urine-Color ⁰¹	Yellow				Yellow
Appearance ⁰¹	Turbid	Abnormal			Clear
WBC Esterase ⁰¹	Negative				Negative
Protein ⁰¹	1+	Abnormal			Negative/Trac
Glucose ⁰¹	Negative				Negative
Ketones ⁰¹	Negative				Negative
Occult Blood ⁰¹	Negative				Negative
Bilirubin ⁰¹	Negative				Negative
Urobilinogen,Semi-Qn ⁰¹	0.2			mg/dL	0.2-1.0
Nitrite, Urine ⁰¹	Negative				Negative
Microscopic Examination ⁰¹	See below:				
		indicated an	d was performed.		
WBC ⁰¹	0-5			/hpf	0 - 5
RBC ⁰¹	None seen			/hpf	0 - 2
Epithelial Cells (non renal) ⁰¹	None seen			/hpf	0 - 10
Casts ⁰¹	None seen			/lpf	None seen

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Date Created and Stored 06/09/23 0916 ET Final Report Page 2 of 3

DOB: Age: Sex: **Male** Patient Report Account Number: 04087470 Ordering Physician: J OLLIFFE



Date Collected: 06/06/2023

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

Bacteria⁰¹

None seen

None seen/Few

Disclaimer

The Previous Result is listed for the most recent test performed by Labcorp in the past 5 years where there is sufficient patient demographic data to match the result to the patient. Results from certain tests are excluded from the Previous Result display.

Icon Legend

🔺 Out of Reference Range 🛛 📕 Critical or Alert

Performing Labs

01: SO - Labcorp San Diego 13112 Evening Creek Dr So Ste 200, San Diego, CA, 92128-4108 Dir: Jenny Galloway, MD 02: BN - Labcorp Burlington 1447 York Court, Burlington, NC, 27215-3361 Dir: Sanjai Nagendra, MD For Inquiries, the physician may contact Branch: 800-859-6046 Lab: 858-668-3700

Patient Details 14527, Donor 8950 VILLA LA JOLLA DR STE B214,STE, LA JOLLA, CA, 920371708

Phone: Date of Birth: Age: Sex: **Male** Patient ID: Alternate Patient ID: Physician Details J OLLIFFE San Diego Sperm Bank 8950 Villa La Jolla Dr Ste B214, La Jolla, CA, 92037

Phone: **858-732-8500** Account Number: **04087470** Physician ID: NPI: **1306838271** Specimen Details Specimen ID: **157-253-3185-0** Control ID: **L2303768710** Alternate Control Number: **L2303768710** Date Collected: **06/06/2023 1131 Local** Date Received: **06/07/2023 0000 ET** Date Entered: **06/07/2023 0336 ET** Date Reported: **06/09/2023 0906 ET**

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