

Acknowledgement of Positive Carrier Screening Results: Donor 4768

I, the undersigned recipient, understand that this donor has tested **POSITIVE** as a carrier for the following condition(s). More information regarding these conditions can be found in his Invitae Expanded Carrier Screening results.

- SLC26A4-related conditions (including Pendred Syndrome)
- WNT10A-related conditions

I intend to use sperm samples from this donor for insemination or other assisted conception procedure(s).

I acknowledge that The Sperm Bank of California (TSBC) has made the donor's genetic testing results available to me and my medical providers, and that I have reviewed these results. I understand that TSBC **strongly recommends** that I review these genetic testing results with a Genetic Counselor and my medical providers. I understand that TSBC can refer me to genetic counseling services if desired.

I understand that recipient testing is strongly recommended when a donor has positive carrier screening results and that such testing can reduce but not eliminate risks.

- I acknowledge that I personally assume all risks associated with use of semen samples provided by a donor who has tested **POSITIVE as a carrier for SLC26A4-related conditions (including Pendred Syndrome) and WNT10A-related conditions**
- I further acknowledge that I understand that WNT10A-related Conditions are inherited in both an autosomal recessive and autosomal dominant pattern.

On behalf of myself and my spouse, heirs, representatives, I hereby release and forever hold harmless TSBC and its current and former officers, directors, employees, attorneys, insurers, consultants, agents, and representatives (collectively "Releasees") from any liability or responsibility whatsoever for any and all outcomes, and hereby release and forever discharge Releasees from any and all actions, causes of action, demands, damages, losses, liabilities, suits, expenses, including attorneys' fees and costs, of whatever character, in law or in equity, whether currently known, suspected, unknown or unsuspected, matured or unmatured, arising out of my use of sperm donated by a donor who has tested **POSITIVE as a carrier for WNT10-A Related Conditions that are inherited in both an** <u>autosomal recessive</u> and <u>autosomal dominant</u> pattern and positive as a carrier for SLC26A4-related conditions (including Pendred Syndrome).

This release involves the waiver of all rights and benefits that I may have under California Civil Code section 1542, which states: "A general release does not extend to claims that the creditor or releasing party does not know or suspect to exist in his or her favor at the time of executing the release and that, if known by him or her, would have materially affected his or her settlement with the debtor or released party."

Please select one of the following:

- □ I have been tested for the above named condition(s) and/or I plan to be tested prior to using the samples.
- □ I understand that TSBC **strongly recommends** that I discuss these results with a Genetic Counselor and my medical providers and consider testing for the above named condition(s). At this time I have **declined** testing and/or **do not anticipate being tested**.

I understand that if I transfer my vials (or embryos if applicable) to any other person, including my spouse, that TSBC requires that person (1) register with TSBC and (2) complete an **Acknowledgement of Positive Carrier Screening Results**. I understand that any and all questions as to the legal interpretation, validity or any other aspect of this agreement shall be determined by the laws of the State of California, regardless of the location or residence of any of the parties.

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Recipient's signature
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GENETIC TESTING: POSITIVE CARRIER STATUS

This donor tested **POSITIVE** as a carrier for one or more autosomal recessive conditions as described on the prior page and in the attached genetic testing results.

What does it mean to be a carrier?

<u>All people carry genetic mutations in their DNA</u>. Genetic testing can help to identify some, but not all, of these mutations. While this donor carries a mutation for one or more recessively inherited condition(s), offspring from this donor are not expected to be at risk of developing these condition(s) unless the recipient (or egg provider if different from the recipient) also carries a genetic mutation for the same condition(s). For this reason, we strongly encourage you to discuss carrier screening for yourself (or your egg provider) with your physician and a genetic counselor. Genetic testing can reduce but not eliminate risks.

*NOTE: Some recessive conditions may also have increased risks for people who only have one copy of the gene. Children who inherit one copy the WNT10-A Related Conditions gene variant from 4768 may be affected with a common condition called isolated tooth agenesis. Tooth agenesis is the absence of one or more teeth. This variant has reduced penetrance - people who carry it may not be affected but are more likely than others to have it. It's estimated people who carry the variant are 2-3 times more likely to have isolated tooth agenesis than the general population where it occurs in 2-8% of people. See "Reclassification of a genetic variant in the WNT10A gene" from Invitae attached to the ECS results for more information.

What are my next steps?

1. Download the genetic test results and review with your medical providers

We **strongly recommend** that you discuss this donor's genetic test results with your physician **PRIOR TO SCHEDULING A SHIPMENT OR PICK-UP**, to confirm the donor is suitable for your use. Vials retrieved from the building cannot be exchanged or refunded. The donor's genetic test results are available for free download on the donor's page at <u>https://www.thespermbankofca.org/donor-catalog</u>.

2. We recommend scheduling a genetic counseling session.

A genetic counselor can explain the results in detail including the inheritance pattern, potential risks to your children, and the available testing options that you may want to consider for yourself (or your egg provider). Phone or in person consultations are available for a fee with TSBC's Genetic Counselors at San Francisco Genetic Counseling (<u>https://www.sfgenetics.org/</u>) or you can locate a genetic counselor at <u>www.findageneticcounselor.com</u>.

3. Complete and return the Acknowledgement of Positive Carrier Screening Results

TSBC requires that all recipients selecting this donor complete this acknowledgement form **PRIOR TO SCHEDULING A SHIPMENT OR PICK-UP**. Completing this form documents that you have been informed about this donor's genetic test results and that you are aware of TSBC's recommendation to discuss the genetic test results with your medical providers as noted above.

Reproductive Technologies, Inc. THE SPERM BANK OF CALIFORNIA

EXPANDED CARRIER SCREENING RESULTS DONOR 4768

Expanded carrier screening for 268 autosomal recessive conditions was completed by Invitae and reported on 08/06/2021.

The results were **POSITIVE** for **SLC26A-Related Conditions (including Pendred Syndrome)** and **WNT10-A Related Conditions**. Donor 4768 is a carrier for these conditions.

It is strongly recommended recipients who use this donor's sperm undergo carrier screening for SLC26A-Related Conditions (including Pendred Syndrome) and WNT10A-related conditions.

WNT10A-related conditions are inherited in both an autosomal recessive and autosomal dominant pattern. The variant detected in WNT10A (c.682A>T; p.Phe228Ile) can cause a common condition called isolated tooth agenesis, which affects 2-8% of the general population. Tooth agenesis, also called hypodontia, is the absence of one or more teeth. It's estimated people who carry the variant are 2-3 times more likely to have one or more missing teeth. All offspring of donor 4768 have a 50% chance to inherit the c.682A>T WNT10A variant and be at increased risk for isolated tooth agenesis/hypodontia.

Testing was negative for the remainder of genes screened.

Disease	Result	Residual risk to be a carrier (based on Middle Eastern ethnicity)
Pendred syndrome and other SLC26A4-related conditions	POSITIVE	n/a
WNT10A-related conditions	POSITIVE	n/a
Cystic Fibrosis	Negative	1 in 4,400
Spinal Muscular Atrophy	Negative: 2 copies exon 7 c.*3+80T>G variant not detected	1 in 800
HBB Hemoglobinopathies & Thalassemia	Negative	1 in 2,700
Alpha Thalassemia	Negative	1 in 241

Genetic tests can significantly reduce, but never completely eliminate, the chance that a person is a carrier for a particular disorder.

Please refer to the donor's Invitae expanded carrier test report for more information on WNT10A conditions, the testing completed and the donors test results.

Please also see the Health Problems List for a summary of the information that this donor has provided to us regarding personal and family medical history.





	ype: Saliva	Report date:	08/06/2021
DOB: Sample	collection date: 07/21/2021	Invitae #:	RQ2437678
Sex: Male Sample	accession date: 07/22/2021	Clinical team:	Janine Mash
MRN:			Lorraine Bonner, MD

Reason for testing

Gamete donor

Test performed

Invitae Comprehensive Carrier Screen without X-linked Disorders

- Primary Panel (CF, SMA)
- Add-on Comprehensive Carrier Screen without X-linked Disorders genes



RESULT: POSITIVE

This carrier test evaluated 268 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.

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: SLC26A4-related conditions	SLC26A4	c.578C>T (p.Thr193Ile)	Autosomal recessive	Yes
Carrier: WNT10A-related conditions	WNT10A	c.682T>A (p.Phe228Ile) §	Autosomal recessive	Yes

🖇 This variant is known to have low penetrance. See Clinical summary and/or Variant details on following pages for more information.

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 table below for residual risks, which presumes a negative family history of the conditions listed.
- Genetic counseling is recommended to further explain the implications of this test result and assess family health history, which may point to health information that merits additional consideration.
- 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.



Clinical summary

B RESULT: CARRIER

SLC26A4-related conditions

A single Pathogenic variant, c.578C>T (p.Thr193Ile), was identified in SLC26A4.

What are SLC26A4-related conditions?

SLC26A4-related conditions include Pendred syndrome (PDS) and SLC26A4-related deafness. Pendred syndrome is a condition that causes deafness, and specific bone and endocrine abnormalities. SLC26A4-related deafness affects an individual's ability to hear.

PDS is characterized by severe to profound bilateral deafness caused by damage to structures in the inner ear (sensorineural deafness), which is typically present at birth. Affected individuals also have balance difficulties due to inner ear problems (vestibular dysfunction) and abnormalities of the temporal (skull) bones, specifically an enlarged vestibular aqueduct (EVA) with or without underdevelopment of a part of the inner ear (cochlear hypoplasia). Additionally, individuals with PDS develop enlargement of the thyroid gland (goiter) in late childhood to early adulthood. Symptoms of PDS may vary, even among members of the same family. Intelligence and life span are not typically affected. Digenic inheritance, which occurs when an individual has a genetic change in two different Pendred syndrome-associated genes, has been reported (PMID: 19426954); however, the evidence available at this time is insufficient to confirm this as a mode of inheritance.

SLC26A4-related deafness is characterized by nonsyndromic deafness (DFNB4), enlarged vestibular aqueduct, and vestibular dysfunction; however, thyroid defects are not observed.

Follow-up depends on each affected individual's specific situation, and discussion with a healthcare provider should be considered.

Next steps

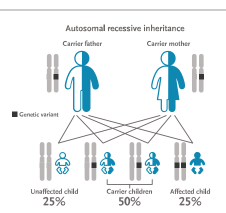
Carrier testing for the reproductive partner is recommended.

+ If your partner tests positive:

In autosomal recessive inheritance, an individual must have disease-causing genetic changes in each copy of the SLC26A4 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 SLC26A4-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 FREQUENCY BEFORE SCREENING	CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
SLC26A4-related conditions (AR)	SLC26A4	Asian	1 in 74	1 in 7300
NM_000441.1	SLC20A4	Pan-ethnic	1 in 80	1 in 7900



RESULT: CARRIER

WNT10A-related conditions

A single Pathogenic (low penetrance) variant, c.682T>A (p.Phe228Ile), was identified in WNT10A. See "What are WNT10A-related conditions?" and Variant details for additional information.

What are WNT10A-related conditions?

WNT10A-related conditions include autosomal recessive odonto-onycho-dermal dysplasia (OODD) and Schöpf-Schulz-Passarge syndrome (SSPS) and autosomal dominant isolated tooth agenesis. Individuals with a clinically significant variant in this gene are carriers for the autosomal recessive conditions and may be at risk to develop the autosomal dominant condition associated with this gene.

Autosomal recessive WNT10A-related conditions, OODD and SSPS, refer to a spectrum of features associated with ectodermal dysplasia (ED), which causes abnormal development of the skin, hair, nails, teeth, and sweat glands. OODD is characterized by dental abnormalities including either fewer teeth than normal (hypodontia) or in more severe cases, the absence of six or more teeth (oligodontia), as well as a smooth tongue, malformed nails (onychodysplasia), clusters of enlarged blood vessels on the face (facial telangiectasias), and thickened skin (hyperkeratosis) with excessive sweating (hyperhidrosis) of the palms of the hands and soles of the feet. SSPS shares the same features as OODD, and is also associated with an increased risk of skin tumors which may be benign (non-cancerous) or malignant, and multiple eyelid cysts. Symptoms and severity of autosomal recessive WNT10A-related conditions are variable. Intellect and life span are not impacted.

Isolated tooth agenesis is a condition that affects the development of the teeth. It can be caused by changes in different genes. Isolated tooth agenesis is the congenital absence of one or more teeth, most commonly the permanent (secondary) teeth. Additional dental abnormalities may include small and/or irregular shaped teeth. Some individuals with WNT10A-related tooth agenesis have also been reported to have mild symptoms of ectodermal dysplasia (ED), which is a condition associated with abnormal development of the skin, hair, nails, teeth, and sweat glands. The severity of WNT10A-related tooth agenesis is variable, and some affected individuals may not have obvious symptoms (incomplete penetrance). Intellect and life span are not impacted.

Please note, the c.682T>A (p.Phe228Ile) variant identified in this individual is known to have low penetrance for both the associated autosomal recessive and autosomal dominant conditions. This means that not all individuals with this genetic change will show signs or symptoms of the condition.

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.

Due to the potential for personal health risk for this individual associated with this result, follow-up with a medical provider may be warranted.

(+) If your partner tests positive:

The WNT10A gene is associated with conditions that are inherited in both an autosomal recessive and autosomal dominant fashion. In autosomal recessive inheritance, an individual must have disease-causing genetic changes in each copy of the WNT10A gene to be affected. Carriers, who have a disease-causing genetic change in only one copy of the gene, typically do not have symptoms of the autosomal recessive condition. When both reproductive partners are carriers of an autosomal recessive condition, there is a 25% chance for each child to have the condition. In autosomal dominant inheritance, an individual with a disease-causing change in one copy of the WNT10A gene is at risk to be affected with autosomal dominant isolated tooth agenesis. When one parent has a change in the WNT10A gene, there is a 50% chance for each child to inherit the change and be at risk to be affected with the autosomal dominant 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 WNT10A-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 FREQUENCY BEFORE SCREENING	CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
WNT10A-related conditions (AR) NM_025216.2	WNT10A	Pan-ethnic	1 in 305	1 in 30400



Variant details

SLC26A4, Exon 5, c.578C>T (p.Thr193Ile), heterozygous, PATHOGENIC

- This sequence change replaces threonine with isoleucine at codon 193 of the SLC26A4 protein (p.Thr193Ile). The threonine residue is highly conserved and there is a moderate physicochemical difference between threonine and isoleucine.
- This variant is present in population databases (rs111033348, ExAC 0.009%).
- This variant has been observed in individual(s) with SLC26A4-related conditions (PMID: 10878664, 28964290, 26752218, 20597900). It has also been observed to segregate with disease in related individuals. This variant is also known as 801C>T. ClinVar contains an entry for this variant (Variation ID: 4830).
- 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 SLC26A4 protein function.
- Experimental studies have shown that this variant affects SLC26A4 protein function (PMID: 31599023, 26752218).
- For these reasons, this variant has been classified as Pathogenic.

WNT10A, Exon 3, c.682T>A (p.Phe228Ile), heterozygous, Pathogenic (low penetrance)

- This sequence change replaces phenylalanine with isoleucine at codon 228 of the WNT10A protein (p.Phe228IIe). The phenylalanine residue is highly conserved and there is a small physicochemical difference between phenylalanine and isoleucine.
- This variant is present in population databases (rs121908120, ExAC 2.0%), including at least one homozygous and/or hemizygous individual.
- This variant has been observed in many individuals with autosomal recessive forms of ectodermal dysplasia (PMID: 19559398, 28976000, 30974434, Invitae). It has been found in trans (on the opposite chromosome) from many different pathogenic variants. Based on an internal analysis, this variant is associated with reduced penetrance for autosomal recessive disease (15% when in homozygosity and 30-60% when present with another pathogenic variant) compared to other pathogenic or likely pathogenic variants, which have a penetrance of 70-80% (Invitae). In addition, in a large meta-analysis, this variant conferred a 2.25-3.42-fold increased risk (95% CI: 1.39-4.10) for isolated tooth agenesis, the autosomal dominant condition associated with WNT10A (PMID: 29364747).
- ClinVar contains an entry for this variant (Variation ID: 4462).
- Algorithms developed to predict the effect of missense changes on protein structure and function are either unavailable or do not agree on the potential impact of this missense change (SIFT: "Deleterious"; PolyPhen-2: "Probably Damaging"; Align-GVGD: "Class C0").
- In summary, this variant is reported to cause disease. However, because this variant is associated with a lower penetrance form of disease than other pathogenic alleles in the WNT10A gene, and because it is found in homozygosity in healthy individuals, it has been classified as Pathogenic (low penetrance).



Residual risk

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. 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 will vary based on the ethnic background of an individual. For individuals of mixed ethnicity, it is recommended to use the highest residual risk estimate. For any genes marked with an asterisk*, refer to the Limitations section below 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 BEFORE SCREENING	CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
3-hydroxy-3-methylglutaryl-CoA lyase deficiency (AR)	HMGCL	Pan-ethnic	≤1 in 500	Reduced
NM_000191.2	HWIGCL	Portuguese	1 in 160	1 in 15900
ABCB11-related conditions (AR) NM_003742.2	ABCB11	Pan-ethnic	1 in 100	1 in 9900
BCC8-related conditions (AR)		Ashkenazi Jewish	1 in 52	1 in 5100
NM_000352.4 When the mother is a noncarrier, but the father is a		Finnish	1 in 100	1 in 9900
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	1 in 17600
Abetalipoproteinemia (AR)	MTTP	Ashkenazi Jewish	1 in 131	1 in 13000
NM_000253.3	IVITIF	Pan-ethnic	≤1 in 500	Reduced
Achromatopsia (CNGB3-related) (AR) NM_019098.4	CNGB3	Pan-ethnic	1 in 93	1 in 9200
ACOX1-related conditions (AR) NM_004035.6	ACOX1	Pan-ethnic	≤1 in 500	Reduced
Acrodermatitis enteropathica (AR) NM_130849.3	SLC39A4	Pan-ethnic	1 in 354	1 in 35300
Adenosine deaminase deficiency (AR) NM_000022.2	ADA	Pan-ethnic	1 in 224	1 in 2788
Aicardi-Goutieres syndrome 5 (AR) NM_015474.3	SAMHD1	Pan-ethnic	≤1 in 500	Reduced
Aldosterone synthase deficiency (AR)	CYP11B2	Pan-ethnic	≤1 in 500	Reduced
NM_000498.3	CIFIIBZ	Sephardic Jewish (Iranian)	1 in 30	1 in 2900
Alpha-mannosidosis (AR) NM_000528.3	MAN2B1	Pan-ethnic	1 in 354	1 in 35300
		African-American	1 in 30	1 in 291
Alpha-thalassemia (AR)	HBA2/	Asian	1 in 20	1 in 191
NM_000517.4, NM_000558.4	HBA1 *	Caucasian	≤1 in 500	Reduced
		Pan-ethnic	1 in 25	1 in 241
Alport syndrome (COL4A3-related) (AR)		Ashkenazi Jewish	1 in 192	1 in 19100
NM_000091.4	COL4A3	Caucasian	1 in 284	1 in 28300
		Pan-ethnic	1 in 354	1 in 35300
Alport syndrome (COL4A4-related) (AR) NM_000092.4	COL4A4	Pan-ethnic	1 in 353	1 in 35200
Alström syndrome (AR) NM_015120.4	ALMS1	Pan-ethnic	≤1 in 500	Reduced
Arginase deficiency (AR) NM_000045.3	ARG1	Pan-ethnic	1 in 274	1 in 27300
Argininosuccinate lyase deficiency (AR) NM_000048.3	ASL	Pan-ethnic	1 in 133	1 in 1321
Aromatase deficiency (AR) NM_031226.2	CYP19A1	Pan-ethnic	≤1 in 500	Reduced
Asparagine synthetase deficiency (AR) NM_133436.3	ASNS	Pan-ethnic	≤1 in 500	Reduced



DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY BEFORE SCREENING	CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
		Sephardic Jewish (Iranian)	1 in 80	1 in 7900
Aspartylglucosaminuria (AR)	AGA	Finnish	1 in 69	1 in 6800
NM_000027.3	AGA	Pan-ethnic	≤1 in 500	Reduced
Ataxia with vitamin E deficiency (AR)	TTPA	Italian	1 in 274	1 in 2731
NM_000370.3	IIIA	Pan-ethnic	≤1 in 500	Reduced
ATM-related conditions (AR)	ATM	Pan-ethnic	1 in 100	1 in 9900
NM_000051.3	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	Sephardic Jewish	1 in 69	1 in 6800
Autoimmuno polyando svinopothy with condidiosis and		Finnish	1 in 79	1 in 7800
toimmune polyendocrinopathy with candidiasis and codermal dysplasia (AR) 1_000383.3	AIRE	Pan-ethnic	1 in 150	1 in 14900
		Sardinian	1 in 60	1 in 5900
		Sephardic Jewish (Iranian)	1 in 48	1 in 4700
ıtosomal recessive congenital ichthyosis GM1-related) (AR)	TGM1	Norwegian	1 in 151	1 in 3000
NM_000359.2	IGIVIT	Pan-ethnic	1 in 224	1 in 4460
Autosomal recessive spastic ataxia of Charlevoix- Saguenay (AR)	SACS	French Canadian (Saguenay-Lac-St- Jean)	1 in 21	1 in 2000
NM_014363.5		Pan-ethnic	≤1 in 500	Reduced
Bardet-Biedl syndrome (BBS10-related) (AR) NM_024685.3	BBS10	Pan-ethnic	1 in 354	1 in 35300
Bardet-Biedl syndrome (BBS12-related) (AR) NM_152618.2	BBS12	Pan-ethnic	1 in 708	Reduced
BBS1-related conditions (AR)	DDC1	Faroese	1 in 30	1 in 2900
NM_024649.4	BBS1	Pan-ethnic	1 in 330	1 in 32900
BBS2-related conditions (AR)	DDCO	Ashkenazi Jewish	1 in 140	1 in 13900
NM_031885.3	BBS2	Pan-ethnic	1 in 560	Reduced
		Caucasian	1 in 407	1 in 40600
3CS1L-related conditions (AR) NM_004328.4	BCS1L	Finnish	1 in 108	1 in 10700
		Pan-ethnic	≤1 in 500	Reduced
Beta-ketothiolase deficiency (AR)	ACATI	Caucasian	1 in 354	1 in 35300
NM_000019.3	ACAT1	Pan-ethnic	≤1 in 500	Reduced
Biopterin-deficient hyperphenylalaninemia (PTS-related)		Chinese	1 in 122	1 in 12100
(AR) NM_000317.2	PTS	Pan-ethnic	1 in 433	1 in 43200
Bloom syndrome (AR)	BLM	Ashkenazi Jewish	1 in 100	1 in 9900
NM_000057.3	DEIW	Pan-ethnic	≤1 in 500	Reduced
BSND-related conditions (AR) NM_057176.2	BSND	Pan-ethnic	≤1 in 500	Reduced
Canavan disease (AR)		Ashkenazi Jewish	1 in 57	1 in 5600
NM_000049.2	ASPA	Pan-ethnic	1 in 159	1 in 15800
Carbamoyl phosphate synthetase I deficiency (AR) NM_001875.4	CPS1	Pan-ethnic	≤1 in 500	Reduced
Carnitine palmitoyltransferase I deficiency (AR)	60 -	Hutterite	1 in 16	1 in 1500
NM_001876.3	CPT1A	Pan-ethnic	≤1 in 500	Reduced
Carnitine palmitoyltransferase II deficiency (AR)		Ashkenazi Jewish	1 in 45	1 in 4400
NM_000098.2	CPT2	Pan-ethnic	1 in 182	1 in 18100
Carpenter syndrome (RAB23-related) (AR) NM_183227.2	RAB23	Pan-ethnic	≤1 in 500	Reduced
Cartilage-hair hypoplasia-anauxetic dysplasia spectrum		Amish	1 in 10	1 in 900
disorders (AR)	RMRP	Finnish	1 in 76	1 in 7500
NR_003051.3		Pan-ethnic	≤1 in 500	Reduced
CDH23-related conditions (AR) NM_022124.5	CDH23	Pan-ethnic	1 in 202	1 in 4020
CEP290-related conditions (AR) NM_025114.3	CEP290	Pan-ethnic	1 in 185	1 in 18400
Cerebrotendinous xanthomatosis (AR)		Pan-ethnic	1 in 112	1 in 5550
NM_000784.3	CYP27A1	Sephardic Jewish	1 in 76	1 in 3750
CERKL-related conditions (AR)	_	Pan-ethnic	1 in 137	1 in 13600
NM_001030311.2	CERKL	Sephardic Jewish	1 in 24	1 in 2300



DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY BEFORE SCREENING	CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
		African-American - classic CF	1 in 61	1 in 6000
		Ashkenazi Jewish - classic CF	1 in 29	1 in 2800
CFTR-related conditions (AR)		Asian - classic CF	1 in 88	1 in 8700
NM_000492.3	CFTR	Caucasian - classic CF	1 in 28	1 in 2700
		Pan-ethnic - classic CF	1 in 45	1 in 4400
		Pan-ethnic - classic CF and CFTR- related disorders	1 in 9	1 in 800
Charcot-Marie-Tooth disease type 4D (AR) NM_006096.3	NDRG1	Pan-ethnic Roma	≤1 in 500 1 in 22	Reduced 1 in 2100
Chorea-acanthocytosis (AR) NM_033305.2	VPS13A *	Pan-ethnic	≤1 in 500	Reduced
Chronic granulomatous disease (CYBA-related) (AR)		Pan-ethnic	≤1 in 500	Reduced
NM_000101.3	CYBA	Sephardic Jewish (Moroccan)	1 in 13	1 in 1200
		Chinese	1 in 65	1 in 6400
		Japanese	1 in 65	1 in 6400
Citrin deficiency (AR)	SLC25A13	Korean	1 in 112	1 in 11100
NM_014251.2		Pan-ethnic	1 in 313	1 in 31200
		Southern Chinese and Taiwanese	1 in 48	1 in 4700
Citrullinemia type 1 (AR) NM_000050.4	ASS1	Pan-ethnic	1 in 120	1 in 2975
CLN3-related conditions (AR) NM_001042432.1	CLN3	Pan-ethnic	1 in 230	1 in 22900
CLRN1-related conditions (AR)		Ashkenazi Jewish	1 in 120	1 in 11900
NM_174878.2	CLRN1	Pan-ethnic	1 in 533	Reduced
Cobalamin C deficiency (AR) NM 015506.2	ММАСНС	Pan-ethnic	1 in 123	1 in 12200
Cobalamin D deficiency (AR) NM_015702.2	MMADHC *	Pan-ethnic	≤1 in 500	Reduced
Cockayne syndrome A (AR) NM_00082.3	ERCC8	Pan-ethnic	1 in 514	Reduced
Cockayne syndrome B (AR) NM_000124.3	ERCC6	Pan-ethnic	1 in 377	1 in 37600
Cohen syndrome (AR)		Amish (Ohio)	1 in 12	1 in 1100
NM_017890.4	VPS13B	Pan-ethnic	≤1 in 500	Reduced
Combined malonic and methylmalonic aciduria (AR) NM_174917.4	ACSF3	Pan-ethnic	1 in 87	1 in 8600
Combined oxidative phosphorylation deficiency 1 (AR) NM_024996.5	GFM1	Pan-ethnic	≤1 in 500	Reduced
Combined oxidative phosphorylation deficiency 3 (AR)	TOFNAL	Finnish	1 in 80	1 in 1129
NM_001172696.1	TSFM *	Pan-ethnic	≤1 in 500	Reduced
Combined pituitary hormone deficiency (LHX3-related) (AR) NM_014564.4	LHX3	Pan-ethnic	≤1 in 500	Reduced
Combined pituitary hormone deficiency (PROP1-related) (AR) NM_006261.4	PROP1	Pan-ethnic	1 in 45	1 in 2200
Congenital adrenal hyperplasia due to 3-beta- hydroxysteroid dehydrogenase deficiency (AR) NM_000198.3	HSD3B2	Pan-ethnic	≤1 in 500	Reduced
Congenital adrenal hyperplasia due to 21-hydroxylase deficiency (AR) NM_000500.7	CYP21A2 *	Pan-ethnic	1 in 61	1 in 751
Congenital disorder of glycosylation (SLC35A3-related)		Ashkenazi Jewish	1 in 469	1 in 46800
(AR) NM_012243.2	SLC35A3	Pan-ethnic	≤1 in 500	Reduced
		Ashkenazi Jewish	1 in 61	1 in 6000
Congenital disorder of glycosylation type Ia (AR)	PMM2	Caucasian	1 in 60	1 in 5900
NM_000303.2		Pan-ethnic	1 in 190	1 in 18900
Congenital disorder of glycosylation type Ib (AR)				
NM_002435.2	MPI	Pan-ethnic	≤1 in 500	Reduced



DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY BEFORE SCREENING	CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
Congenital disorder of glycosylation type Ic (AR) NM_013339.3	ALG6 *	Pan-ethnic	≤1 in 500	Reduced
Congenital insensitivity to pain with anhidrosis (AR) NM_001012331.1	NTRK1	Pan-ethnic	≤1 in 500	Reduced
Congenital myasthenic syndrome (CHRNE-related)		European Roma	1 in 25	1 in 2400
(AR) NM_000080.3	CHRNE	Pan-ethnic	1 in 200	1 in 19900
		Finnish	1 in 46	1 in 4500
Congenital nephrotic syndrome type 1 (AR) NM_004646.3	NPHS1	Old Order Mennonite	1 in 12	1 in 1100
INIM_004646.3		Pan-ethnic	≤1 in 500	Reduced
Congenital nephrotic syndrome type 2 (AR) NM_014625.3	NPHS2	Pan-ethnic	≤1 in 500	Reduced
Corneal dystrophy and perceptive deafness (AR) NM_032034.3	SLC4A11	Pan-ethnic	≤1 in 500	Reduced
CRB1-related conditions (AR) NM_201253.2	CRB1	Pan-ethnic	1 in 112	1 in 11100
CYP11B1-related conditions (AR)	CYP11B1	Pan-ethnic	1 in 194	1 in 19300
NM_000497.3	СПТЮТ	Sephardic Jewish (Moroccan)	1 in 40	1 in 3900
CYP17A1-related conditions (AR) NM_000102.3	CYP17A1	Pan-ethnic	≤1 in 500	Reduced
Cystinosis (AR)		French Canadian (Saguenay-Lac-St- Jean)	1 in 39	1 in 3800
Cystinosis (AR) NM_004937.2	CTNS	Pan-ethnic	1 in 158	1 in 15700
		Sephardic Jewish (Moroccan)	1 in 100	1 in 9900
DHDDS-related conditions (AR)	DUDDC	Ashkenazi Jewish	1 in 117	1 in 11600
NM_024887.3	DHDDS	Pan-ethnic	≤1 in 500	Reduced
Dihydrolipoamide dehydrogenase deficiency (AR)	DLD	Ashkenazi Jewish	1 in 107	1 in 5300
NM_000108.4	DLD	Pan-ethnic	≤1 in 500	Reduced
Distal renal tubular acidosis with deafness		Pan-ethnic	≤1 in 500	Reduced
(ATP6V1B1-related) (AR) NM_001692.3	ATP6V1B1	Sephardic Jewish	1 in 140	1 in 13900
DYSF-related conditions (AR)	DYSF	Pan-ethnic	1 in 311	1 in 31000
NM_003494.3		Sephardic Jewish (Libyan)	1 in 10	1 in 900
Dyskeratosis congenita spectrum disorders (RTEL1-related) (AR)	RTEL1	Ashkenazi Jewish	1 in 222	1 in 22100
NM_001283009.1	RIELI	Pan-ethnic	≤1 in 500	Reduced
Dystrophic epidermolysis bullosa (AR) NM_000094.3	COL7A1	Pan-ethnic	1 in 370	1 in 12300
Ehlers-Danlos syndrome, dermatosparaxis type (AR)	ADAMTS2	Ashkenazi Jewish	1 in 187	1 in 18600
NM_014244.4	ADAWI 52	Pan-ethnic	≤1 in 500	Reduced
Ellis-van Creveld syndrome (EVC-related) (AR)	EVC	Amish	1 in 8	1 in 700
NM_153717.2		Pan-ethnic	1 in 220	1 in 21900
Ethylmalonic encephalopathy (AR) NM_014297.3	ETHE1	Pan-ethnic	≤1 in 500	Reduced
EVC2-related conditions (AR) NM_147127.4	EVC2	Pan-ethnic	1 in 199	1 in 19800
Familial chylomicronemia syndrome (AR) NM_000237.2	LPL	French Canadian (Saguenay-Lac-St- Jean)	1 in 46	1 in 4500
		Pan-ethnic	≤1 in 500	Reduced
Familial dysautonomia (AR)	ELP1	Ashkenazi Jewish	1 in 36	1 in 3500
NM_003640.3		Pan-ethnic	≤1 in 500	Reduced
		Afrikaner	1 in 72	1 in 7100
Familial hypercholesterolemia (LDLR-related) (AD) NM_000527.4	LDLR	Ashkenazi Jewish	1 in 69	1 in 6800
INIVI_UUU327.4		French Canadian	1 in 270	1 in 26900
		Pan-ethnic	1 in 250	1 in 24900
Familial hypercholesterolemia (LDLRAP1-related) (AR) NM_015627.2	LDLRAP1	Pan-ethnic Sardinian	≤1 in 500	Reduced
		Sardinian Afrikaner	1 in 143	1 in 14200
Fanconi anemia type A (AR) NM_000135.2	FANCA	Pan-ethnic	1 in 83 1 in 345	1 in 8200 1 in 34400



DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY BEFORE SCREENING	CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
		Sephardic Jewish	1 in 133	1 in 13200
		Spanish Roma	1 in 64	1 in 6300
Fanconi anemia type C (AR)	FANCC	Ashkenazi Jewish	1 in 89	1 in 8800
NM_000136.2	TANCC	Pan-ethnic	1 in 417	1 in 41600
Fanconi anemia type G (AR)	FANCG	African-American	1 in 100	1 in 9900
NM_004629.1	FAILCG	Pan-ethnic	≤1 in 500	Reduced
FH-related conditions (AR) NM_000143.3	FH	Pan-ethnic	≤1 in 500	Reduced
Galactokinase deficiency galactosemia (AR)	CALKI	Pan-ethnic	1 in 122	1 in 12100
NM_000154.1	GALK1	Roma	1 in 47	1 in 4600
		African-American	1 in 87	1 in 8600
alactosemia (GALT-related) (AR)	CALT	Ashkenazi Jewish	1 in 156	1 in 15500
NM_000155.3	GALT	Irish Traveller	1 in 11	1 in 1000
		Pan-ethnic	1 in 100	1 in 9900
GBA-related conditions including Gaucher disease (AR)	CDA +	Ashkenazi Jewish	1 in 15	1 in 234
NM_001005741.2	GBA *	Pan-ethnic	1 in 158	1 in 561
GBE1-related conditions (AR)	6051	Ashkenazi Jewish	1 in 68	1 in 6700
NM_000158.3	GBE1	Pan-ethnic	1 in 387	1 in 38600
Gitelman syndrome (AR) NM_000339.2	SLC12A3	Pan-ethnic	1 in 100	1 in 9900
		Ashkenazi Jewish	1 in 13	1 in 1200
GJB2-related conditions (AR) NM_004004.5	GJB2	Pan-ethnic	1 in 50	1 in 4900
		Thai	1 in 9	1 in 800
		Pan-ethnic	1 in 158	1 in 15700
GLB1-related conditions (AR) NM_000404.2	GLB1	Roma	1 in 50	1 in 4900
	GEBT	South Brazilian	1 in 58	1 in 5700
GLE1-related conditions (AR)		Finnish	1 in 100	1 in 9900
NM_001003722.1	GLE1	Pan-ethnic	≤1 in 500	Reduced
		Amish	1 in 9	1 in 800
Glutaric acidemia type I (AR)	GCDH	Oji-Cree First Nations	1 in 9	1 in 800
NM_000159.3	Gebii	Pan-ethnic	1 in 87	1 in 8600
Glutaric acidemia type IIA (AR) NM_000126.3	ETFA	Pan-ethnic	≤1 in 500	Reduced
		Asian	1 in 87	1 in 8600
Glutaric acidemia type IIC (AR) NM_004453.3	ETFDH	Pan-ethnic	1 in 250	1 in 24900
		Finnish	1 in 142	1 in 14100
Glycine encephalopathy (AMT-related) (AR) NM_000481.3	AMT	Pan-ethnic	1 in 325	1 in 32400
Glycine encephalopathy (GLDC-related) (AR)		Caucasian	1 in 141	1 in 14000
NM 000170.2	GLDC	Pan-ethnic	1 in 165	1 in 16400
		Ashkenazi Jewish	1 in 71	1 in 1400
Glycogen storage disease type Ia (AR) NM_000151.3	G6PC	Pan-ethnic	1 in 177	1 in 3520
Glycogen storage disease type Ib (AR) NM_001164277.1	SLC37A4	Pan-ethnic	1 in 354	1 in 7060
11.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1		African-American	1 in 60	1 in 5900
Glycogen storage disease type II (Pompe disease) (AR)		Ashkenazi Jewish	1 in 58	1 in 5700
NM 000152.3	GAA	Asian	1 in 112	1 in 11100
···· <u>-</u> ····		Pan-ethnic	1 in 100	1 in 9900
		Faroese	1 in 28	1 in 540
Glycogen storage disease type III (AR)	AGL	Pan-ethnic	1 in 159	1 in 3160
NM_000642.2	AUL	Sephardic Jewish (Moroccan)	1 in 34	1 in 660
		Caucasian	1 in 158	1 in 15700
Glycogen storage disease type V (AR)	PYGM	Pan-ethnic	1 in 158	
NM_005609.3	PIGN			1 in 17000
		Sephardic Jewish (Kurdish)	1 in 84	1 in 8300
Glycogen storage disease type VII (AR)	PFKM	Ashkenazi Jewish	1 in 250	1 in 24900
NM_000289.5		Pan-ethnic	≤1 in 500	Reduced
GNE-related conditions (AR) NM_001128227.2	GNE	Pan-ethnic	1 in 179	1 in 17800



DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY BEFORE SCREENING	CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
		Sephardic Jewish (Iranian)	1 in 10	1 in 900
GNPTAB-related conditions (AR)	GNPTAB	Irish Traveller	1 in 15	1 in 1400
NM_024312.4	GINPTAD	Pan-ethnic	1 in 200	1 in 19900
Guanidinoacetate methyltransferase deficiency (AR)	GAMT	Pan-ethnic	≤1 in 500	Reduced
NM_000156.5	GAIVIT	Portuguese	1 in 125	1 in 12400
		Finnish	1 in 126	1 in 12500
yrate atrophy of the choroid and retina (AR) M_000274.3	OAT *	Pan-ethnic	≤1 in 500	Reduced
NN_000274.5	-	Sephardic Jewish	1 in 177	1 in 17600
		Caucasian	1 in 250	1 in 24900
DHA-related conditions (AR) 1_000182.4	HADHA	Finnish	1 in 125	1 in 12400
NM_000182.4		Pan-ethnic	1 in 350	1 in 34900
		African-American	1 in 8	1 in 700
	-	Asian	1 in 54	1 in 5300
HBB-related hemoglobinopathies (AR)	-	Caucasian	1 in 373	1 in 37200
NM_000518.4	HBB	Hispanic	1 in 17	1 in 1600
	-	Mediterranean	1 in 28	1 in 2700
		Pan-ethnic	1 in 49	1 in 4800
		African-American	1 in 226	1 in 22500
Hereditary fructose intolerance (AR)	ALDOB	Middle Eastern	1 in 97	1 in 9600
NM_000035.3	, LEOOD	Pan-ethnic	1 in 122	1 in 12100
Hereditary hemochromatosis type 2 (HJV-related) (AR) NM_213653.3	нј∨	Pan-ethnic	≤1 in 500	Reduced
Hereditary hemochromatosis type 3 (AR) NM_003227.3	TFR2	Pan-ethnic	≤1 in 500	Reduced
lermansky-Pudlak syndrome type 1 (AR)	11001	Pan-ethnic	≤1 in 500	Reduced
NM_000195.4	HPS1	Puerto Rican (Northwestern)	1 in 21	1 in 2000
		Ashkenazi Jewish	1 in 235	1 in 23400
Hermansky-Pudlak syndrome type 3 (AR) NM 032383.4	HPS3	Pan-ethnic	≤1 in 500	Reduced
NM_032383.4		Puerto Rican (Central)	1 in 63	1 in 6200
HGSNAT-related conditions (AR) NM_152419.2	HGSNAT	Pan-ethnic	≤1 in 500	Reduced
		Faroese	1 in 20	1 in 1900
Holocarboxylase synthetase deficiency (AR) NM_000411.6	HLCS	Japanese	1 in 158	1 in 15700
NN000411.0		Pan-ethnic	1 in 224	1 in 22300
Homocystinuria due to cobalamin E deficiency (AR) NM_002454.2	MTRR	Pan-ethnic	≤1 in 500	Reduced
Homocystinuria due to cystathionine beta-synthase		Norwegian	1 in 40	1 in 3900
deficiency (AR)	CBS	Pan-ethnic	1 in 224	1 in 22300
NM_000071.2		Qatari	1 in 21	1 in 2000
Homocystinuria due to MTHFR deficiency (AR)	MTHFR *	Pan-ethnic	≤1 in 500	Reduced
NM_005957.4	MITHER *	Sephardic Jewish (Bukharian)	1 in 39	1 in 3800
HSD17B4-related conditions (AR) NM_000414.3	HSD17B4	Pan-ethnic	1 in 158	1 in 15700
Hydrolethalus syndrome type 1 (AR)		Finnish	1 in 40	1 in 3900
NM_145014.2	HYLS1	Pan-ethnic	≤1 in 500	Reduced
Hyperornithinemia-hyperammonemia-homocitrullinuria		Metis (Saskatchewan)	1 in 19	1 in 1800
syndrome (AR) NM_014252.3	SLC25A15	Pan-ethnic	≤1 in 500	Reduced
Hypophosphatasia (AR) NM_000478.5	ALPL	Mennonite Pan-ethnic	1 in 25 1 in 150	1 in 480 1 in 2980
Isovaleric acidemia (AR) NM_002225.3	IVD	Pan-ethnic	1 in 250	1 in 24900
Joubert syndrome and related disorders (MKS1-related)		Finnish	1 in 47	1 in 920
(AR) NM_017777.3	MKS1	Pan-ethnic	1 in 260	1 in 5180
Joubert syndrome and related disorders (RPGRIP1L- related) (AR) NM_015272.2	RPGRIP1L *	Pan-ethnic	1 in 259	1 in 5160



DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY BEFORE SCREENING	CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
Joubert syndrome and related disorders		Ashkenazi Jewish	1 in 92	1 in 9100
(TMEM216-related) (AR) NM_001173990.2	TMEM216	Pan-ethnic	≤1 in 500	Reduced
Junctional epidermolysis bullosa (LAMC2-related) (AR) NM_005562.2	LAMC2	Pan-ethnic	≤1 in 500	Reduced
KCNJ11-related conditions (AR) NM_000525.3	KCNJ11	Pan-ethnic	≤1 in 500	Reduced
Krabbe disease (AR)	GALC *	Druze	1 in 6	1 in 500
NM_000153.3	GALC	Pan-ethnic	1 in 158	1 in 15700
AMA2-related muscular dystrophy (AR) NM_000426.3	LAMA2	Pan-ethnic	1 in 87	1 in 8600
AMA3-related conditions (AR) NM_000227.4	LAMA3	Pan-ethnic	≤1 in 500	Reduced
AMB3-related conditions (AR) NM_000228.2	LAMB3	Pan-ethnic	1 in 317	1 in 31600
Leber congenital amaurosis 5 (AR) NM_181714.3	LCA5	Pan-ethnic	1 in 645	Reduced
Leukoencephalopathy with vanishing white matter (EIF2B5-related) (AR) NM_003907.2	EIF2B5	Pan-ethnic	≤1 in 500	Reduced
.imb-girdle muscular dystrophy (CAPN3-related) (AR) NM_000070.2	CAPN3	Pan-ethnic	1 in 134	1 in 13300
		Caucasian	1 in 571	Reduced
.imb-girdle muscular dystrophy type 2C (AR)		Japanese	1 in 374	1 in 37300
NM_000231.2	SGCG	Moroccan	1 in 250	1 in 24900
		Pan-ethnic	≤1 in 500	Reduced
		Roma	1 in 59	1 in 5800
Limb-girdle muscular dystrophy type 2D (AR) NM_000023.2	SGCA	Caucasian	1 in 286	1 in 28500
		Finnish	1 in 150	1 in 14900
		Pan-ethnic	≤1 in 500	Reduced
.imb-girdle muscular dystrophy type 2E (AR)	SGCB	Caucasian	1 in 404	1 in 5038
VM_000232.4		Pan-ethnic	≤1 in 500	Reduced
.ipoid congenital adrenal hyperplasia (AR) IM_000349.2	STAR	Korean	1 in 170	1 in 16900
NM_000345.2		Pan-ethnic	≤1 in 500	Reduced
ysinuric protein intolerance (AR)	CL C747	Finnish	1 in 120	1 in 2380
íм_001126106.2	SLC7A7	Japanese	1 in 120	1 in 2380
		Pan-ethnic Caucasian	≤1 in 500	Reduced
ysosomal acid lipase deficiency (AR)	LIPA	Pan-ethnic	1 in 112 1 in 359	1 in 1850 1 in 5967
IM_000235.3		Sephardic Jewish (Iranian)	1 in 33	1 in 534
Major histocompatibility complex class II deficiency CIITA-related) (AR) NM_000246.3	CIITA	Pan-ethnic	≤1 in 500	Reduced
Maple syrup urine disease type 1A (AR)	DCKDUA	Mennonite	1 in 10	1 in 900
VM_000709.3	BCKDHA	Pan-ethnic	1 in 373	1 in 37200
Maple syrup urine disease type 1B (AR)	вскрнв	Ashkenazi Jewish	1 in 97	1 in 9600
NM_183050.2	DENDEID	Pan-ethnic	1 in 346	1 in 34500
Ларle syrup urine disease type 2 (AR) NM_001918.3	DBT	Pan-ethnic	≤1 in 500	Reduced
Medium-chain acyl-CoA dehydrogenase deficiency (AR) NM_000016.5	ACADM	Northern European	1 in 40	1 in 3900
		Pan-ethnic	1 in 66	1 in 6500
Aegalencephalic leukoencephalopathy with subcortical systs 1 (AR)	MLC1	Pan-ethnic	≤1 in 500	Reduced
ν́Μ_01̀516́6.3		Sephardic Jewish (Libyan)	1 in 40	1 in 3900
Aetachromatic leukodystrophy (ARSA-related) (AR)		Navajo	1 in 40	1 in 780
VM_000487.5	ARSA	Pan-ethnic	1 in 100	1 in 1980
		Sephardic Jewish	1 in 46	1 in 900
Methylmalonic acidemia (MMAA-related) (AR) NM_172250.2	MMAA	Pan-ethnic	1 in 316	1 in 10500



DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY BEFORE SCREENING	CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
Methylmalonic acidemia (MMAB-related) (AR) NM_052845.3	MMAB	Pan-ethnic	1 in 456	1 in 22750
Methylmalonic acidemia (MUT-related) (AR) NM_000255.3	MUT	Pan-ethnic	1 in 204	1 in 5075
MFSD8-related conditions (AR) NM_152778.2	MFSD8	Pan-ethnic	≤1 in 500	Reduced
Microcephaly, postnatal progressive, with seizures and	MED17	Pan-ethnic	≤1 in 500	Reduced
brain atrophy (AR) NM_004268.4		Sephardic Jewish	1 in 20	1 in 1900
Mitochondrial complex I deficiency 9 (AR)	NDUFS6	Ashkenazi Jewish	1 in 290	1 in 28900
NM_004553.4		Caucasus Jewish	1 in 24	1 in 2300
		Pan-ethnic	≤1 in 500	Reduced
Mitochondrial complex I deficiency 16 (AR)	NDUFAF5	Ashkenazi Jewish	1 in 290	1 in 28900
NM_024120.4		Pan-ethnic	≤1 in 500	Reduced
Mitochondrial complex I deficiency 20/ACAD9 deficiency (AR) NM_014049.4	ACAD9	Pan-ethnic	≤1 in 500	Reduced
Mitochondrial complex IV deficiency / Leigh syndrome, French Canadian type (AR)	LRPPRC	French Canadian (Saguenay-Lac-St- Jean)	1 in 23	1 in 2200
NM_133259.3		Pan-ethnic	≤1 in 500	Reduced
Mitochondrial DNA depletion syndrome-6 (AR)	MPV17	Navajo	1 in 20	1 in 475
NM_002437.4		Pan-ethnic	≤1 in 500	Reduced
Mitochondrial neurogastrointestinal	7.445	Pan-ethnic	≤1 in 500	Reduced
encephalomyopathy (AR) NM_001953.4	ТҮМР	Sephardic Jewish	1 in 158	1 in 15700
MPL-related conditions (AR)	MPL	Ashkenazi Jewish	1 in 57	1 in 5600
NM_005373.2	IVIT E	Pan-ethnic	≤1 in 500	Reduced
Mucolipidosis type III gamma (AR) NM_032520.4	GNPTG	Pan-ethnic	≤1 in 500	Reduced
Mucolipidosis type IV (AR)	MCOLN1	Ashkenazi Jewish	1 in 100	1 in 9900
NM_020533.2		Pan-ethnic	≤1 in 500	Reduced
Mucopolysaccharidosis type I (AR) NM_000203.4	IDUA	Pan-ethnic	1 in 148	1 in 4900
Mucopolysaccharidosis type IIIA (AR)	SGSH	Northern European	1 in 173	1 in 17200
NM_000199.3		Pan-ethnic	1 in 215	1 in 21400
		Taiwanese	≤1 in 500	Reduced
Mucopolysaccharidosis type IIIB (AR) NM_000263.3	NAGLU	Pan-ethnic	1 in 224	1 in 22300
Mucopolysaccharidosis type IIID (AR) NM_002076.3	GNS	Pan-ethnic	≤1 in 500	Reduced
Mucopolysaccharidosis type IX (AR) NM_153281.1	HYAL1	Pan-ethnic	≤1 in 500	Reduced
Mucopolysaccharidosis type VI (AR) NM_000046.3	ARSB	Pan-ethnic	1 in 250	1 in 24900
Multiple sulfatase deficiency (AR) NM_182760.3	SUMF1	Pan-ethnic	≤1 in 500	Reduced
Muscular dystrophy-dystroglycanopathy (FKRP-related)		Norwegian	1 in 116	1 in 11500
(AR) NM_024301.4	FKRP	Pan-ethnic	1 in 158	1 in 15700
Muscular dystrophy-dystroglycanopathy (FKTN-related)	FKTN	Ashkenazi Jewish	1 in 80	1 in 7900
(AR)		Japanese	1 in 188	1 in 18700
NM_001079802.1		Pan-ethnic	≤1 in 500	Reduced
MYO7A-related conditions (AR) NM_000260.3	MYO7A	Pan-ethnic	1 in 200	1 in 3980
Myopathy, lactic acidosis, and sideroblastic anemia 1 (AR) NM_025215.5	PUS1	Pan-ethnic	≤1 in 500	Reduced
N-acetylglutamate synthase deficiency (AR) NM_153006.2	NAGS	Pan-ethnic	≤1 in 500	Reduced
Nemaline myopathy 2 (AR)		Ashkenazi Jewish	1 in 108	1 in 10700
NM_001271208.1	NEB *	Pan-ethnic	1 in 158	1 in 3140



DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY BEFORE SCREENING	CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
Nephrogenic diabetes insipidus (AQP2-related) (AR) NM_000486.5	AQP2	Pan-ethnic	1 in 1118	Reduced
Neuronal ceroid lipofuscinosis type 1 (AR)	PPT1	Finnish	1 in 70	1 in 3450
NM_000310.3		Pan-ethnic	1 in 199	1 in 9900
Neuronal ceroid lipofuscinosis type 2 (AR)	TPP1	Newfoundland	1 in 53	1 in 1734
NM_000391.3		Pan-ethnic	1 in 250	1 in 8300
Neuronal ceroid lipofuscinosis type 5 (AR)	CLN5	Finnish	1 in 115	1 in 11400
NM_006493.2		Pan-ethnic	≤1 in 500	Reduced
Neuronal ceroid lipofuscinosis type 6 (AR) NM_017882.2	CLN6	Pan-ethnic	≤1 in 500	Reduced
Neuronal ceroid lipofuscinosis type 8 (AR)	CLN8	Finnish	1 in 135	1 in 13400
NM_018941.3		Pan-ethnic	≤1 in 500	Reduced
Niemann-Pick disease type C (NPC1-related) (AR) NM_000271.4	NPC1	Pan-ethnic	1 in 183	1 in 18200
Niemann-Pick disease type C (NPC2-related) (AR) NM_006432.3	NPC2	Pan-ethnic	1 in 871	Reduced
Niemann-Pick disease types A and B (AR)	SMPD1	Ashkenazi Jewish	1 in 90	1 in 1780
NM_000543.4	SIVIPUT	Pan-ethnic	1 in 250	1 in 4980
Nijmegen breakage syndrome (AR)	NBN *	Eastern European	1 in 155	1 in 15400
NM_002485.4	INDIN	Pan-ethnic	≤1 in 500	Reduced
Nonsyndromic deafness (LOXHD1-related) (AR)	LOXHD1	Ashkenazi Jewish	1 in 180	1 in 17900
NM_144612.6	LOXIDI	Pan-ethnic	≤1 in 500	Reduced
NR2E3-related conditions (AR) NM_014249.3	NR2E3	Pan-ethnic	≤1 in 500	Reduced
OPA3-related conditions (AR)	OPA3	Pan-ethnic	≤1 in 500	Reduced
NM_025136.3	OPA3	Sephardic Jewish (Iraqi)	1 in 10	1 in 900
		Ashkenazi Jewish	1 in 350	1 in 34900
Osteopetrosis (TCIRG1-related) (AR) NM_006019.3	TCIRG1	Chuvash	1 in 30	1 in 2900
NW_000013.5		Pan-ethnic	1 in 317	1 in 31600
PCDH15-related conditions (AR)	PCDH15	Ashkenazi Jewish	1 in 78	1 in 7700
NM_033056.3	FCDITIS	Pan-ethnic	1 in 400	1 in 39900
PEX7-related conditions (AR) NM_000288.3	PEX7	Pan-ethnic	1 in 157	1 in 15600
		African-American	1 in 111	1 in 11000
		Ashkenazi Jewish	1 in 225	1 in 22400
	-	East Asian	1 in 50	1 in 1225
Phenylalanine hydroxylase deficiency (AR)	PAH	Finnish	1 in 225	1 in 22400
NM_000277.1	РАН	Irish	1 in 33	1 in 3200
		Japanese	1 in 200	1 in 19900
		Pan-ethnic	1 in 58	1 in 5700
		Turkish	1 in 26	1 in 2500
Phosphoglycerate dehydrogenase deficiency (AR)	PHGDH	Ashkenazi Jewish	1 in 400	1 in 39900
NM_006623.3	FIGDI	Pan-ethnic	≤1 in 500	Reduced
Polycystic kidney disease (PKHD1-related) (AR) NM_138694.3	PKHD1	Pan-ethnic	1 in 70	1 in 6900
Polymicrogyria (ADGRG1-related) (AR) NM_005682.6	ADGRG1	Pan-ethnic	≤1 in 500	Reduced
POMGNT1-related conditions (AR) NM_017739.3	POMGNT1	Finnish Don ethnic	1 in 111	1 in 11000
NIVI_V1//37.3		Pan-ethnic	≤1 in 500	Reduced
Pontocerebellar hypoplasia type 2D (AR) NM_016955.3	SEPSECS	Pan-ethnic Sephardic Jewish (Moroccan and	≤1 in 500 1 in 43	Reduced
Pontocerebellar hypoplasia type 6 (AR)	RARS2	Iraqi) Pan-ethnic	≤1 in 500	Reduced
NM_020320.3	MAR JZ			
Primary carnitine deficiency (AR)	CI COOLE	Faroese	1 in 9	1 in 800
NM_003060.3	SLC22A5	Japanese	1 in 100	1 in 9900



DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY BEFORE SCREENING	CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
Primary ciliary dyskinesia (DNAH5-related) (AR) NM_001369.2	DNAH5	Pan-ethnic	1 in 109	1 in 10800
Primary ciliary dyskinesia (DNAI1-related) (AR) NM_012144.3	DNAI1	Pan-ethnic	1 in 250	1 in 24900
Primary ciliary dyskinesia (DNAI2-related) (AR)	DNAI2	Ashkenazi Jewish	1 in 200	1 in 19900
NM_023036.4	DNAI2	Pan-ethnic	1 in 354	1 in 35300
Primary hyperoxaluria type 1 (AR) NM_000030.2	AGXT	Pan-ethnic	1 in 135	1 in 13400
Primary hyperoxaluria type 2 (AR) NM_012203.1	GRHPR	Pan-ethnic	≤1 in 500	Reduced
Primary hyperoxaluria type 3 (AR) NM_138413.3	HOGA1	Pan-ethnic	1 in 354	1 in 35300
Propionic acidemia (PCCA-related) (AR)	РССА	Arab	1 in 100	1 in 2475
NM_000282.3		Pan-ethnic	1 in 224	1 in 5575
Propionic acidemia (PCCB-related) (AR)		Arab	1 in 100	1 in 9900
NM_000532.4	РССВ	Greenlandic Inuit	1 in 20	1 in 1900
		Pan-ethnic	1 in 224	1 in 22300
PSAP-related conditions (AR) NM_002778.3	PSAP	Pan-ethnic	≤1 in 500	Reduced
Pycnodysostosis (AR) NM_000396.3	СТЅК	Pan-ethnic	1 in 438	1 in 43700
Pyruvate carboxylase deficiency (AR)	PC	Algonquian Indian	1 in 10	1 in 180
NM_000920.3		Pan-ethnic	1 in 250	1 in 4980
Pyruvate dehydrogenase complex deficiency (PDHB- related) (AR) NM_000925.3	PDHB	Pan-ethnic	≤1 in 500	Reduced
RAPSN-related conditions (AR) NM_005055.4	RAPSN	Pan-ethnic	1 in 283	1 in 28200
RDH12-related conditions (AR) NM_152443.2	RDH12	Pan-ethnic	1 in 460	1 in 45900
Retinitis pigmentosa 25 (AR) NM_001142800.1	EYS	Pan-ethnic	1 in 129	1 in 12800
		Sephardic Jewish	1 in 42	1 in 4100
Retinitis pigmentosa 28 (AR)	FAM161A	Ashkenazi Jewish	1 in 214	1 in 21300
NM_001201543.1		Pan-ethnic	1 in 289	1 in 28800
		Sephardic Jewish	1 in 41	1 in 4000
Rhizomelic chondrodysplasia punctata type 3 (AR) NM_003659.3	AGPS	Pan-ethnic	≤1 in 500	Reduced
Roberts syndrome (AR) NM_001017420.2	ESCO2	Pan-ethnic	≤1 in 500	Reduced
RPE65-related conditions (AR)	RPE65	Pan-ethnic	1 in 228	1 in 22700
NM_000329.2	KI EOS	Sephardic Jewish	1 in 90	1 in 8900
Sandhoff disease (AR)	НЕХВ	Metis (Saskatchewan)	1 in 15	1 in 1400
NM_000521.3	TIEXE	Pan-ethnic	1 in 180	1 in 17900
Schimke immuno-osseous dysplasia (AR) NM_014140.3	SMARCAL1	Pan-ethnic	≤1 in 500	Reduced
Severe combined immunodeficiency due to DCLRE1C	DOUDERC	Navajo and Apache	1 in 10	1 in 900
(Artemis) deficiency (AR) NM_001033855.2	DCLRE1C	Pan-ethnic	≤1 in 500	Reduced
Severe combined immunodeficiency due to RAG2 deficiency (AR) NM_000536.3	RAG2	Pan-ethnic	≤1 in 500	Reduced
Severe congenital neutropenia due to HAX1 deficiency (AR) NM_006118.3	HAX1	Pan-ethnic	≤1 in 500	Reduced
Severe congenital neutropenia due to VPS45 deficiency (AR) NM_007259.4	VPS45	Pan-ethnic	≤1 in 500	Reduced
Sialic acid storage diseases (AR)	SI C1745	Finnish	1 in 100	1 in 9900
NM_012434.4	SLC17A5	Pan-ethnic	≤1 in 500	Reduced
Sjögren-Larsson syndrome (AR)		Pan-ethnic	≤1 in 500	Reduced
ŃM_000382.2	ALDH3A2	Swedish	1 in 250	1 in 24900



DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY BEFORE SCREENING	CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
SLC12A6-related conditions (AR) NM_133647.1	SLC12A6	French Canadian (Saguenay-Lac-St- Jean)	1 in 23	1 in 2200
		Pan-ethnic	≤1 in 500	Reduced
SLC26A2-related conditions (AR) NM_000112.3	SLC26A2	Finnish	1 in 75	1 in 1480
		Pan-ethnic	1 in 158	1 in 3140
		African-American	1 in 339	1 in 33800
		Ashkenazi Jewish	1 in 41	1 in 4000
Smith Low Chitz and to ma (AD)		Hispanic	1 in 135	1 in 13400
Smith-Lemli-Opitz syndrome (AR) NM_001360.2	DHCR7	Northern European	1 in 50	1 in 4900
		Pan-ethnic	1 in 71	1 in 7000
		Sephardic Jewish	1 in 68	1 in 6700
		Southern European	1 in 83	1 in 8200
Spastic paraplegia type 15 (AR) NM_015346.3	ZFYVE26	Pan-ethnic	≤1 in 500	Reduced
Spastic paraplegia type 49 (AR)	TECODO	Pan-ethnic	≤1 in 500	Reduced
NM_014844.3	TECPR2	Sephardic Jewish - Bukharian	1 in 38	1 in 3700
Spinal muscular atrophy (AR)		African-American	1 in 59	1 in 342
NM_000344.3		Ashkenazi Jewish	1 in 62	1 in 1017
SMN1: 2 copies c.*3+80T>G not detected	SMN1 *	Asian	1 in 50	1 in 701
Carrier residual risks listed are for 2 copy SMN1 results.	SIVINI ^	Caucasian	1 in 45	1 in 880
Carrier residual risk for >2 copies are 5- to 10-fold		Hispanic	1 in 48	1 in 784
lower.		Pan-ethnic	1 in 49	1 in 800
Spondylocostal dysostosis (AR)	MECDO	Pan-ethnic	1 in 224	1 in 22300
NM_001039958.1	MESP2	Puerto Rican	1 in 55	1 in 5400
Steel syndrome (AR)	COL 2741 +	Pan-ethnic	≤1 in 500	Reduced
NM_032888.3	COL27A1 *	Puerto Rican	1 in 51	1 in 5000
Stüve-Wiedemann syndrome (AR) NM_002310.5	LIFR	Pan-ethnic	≤1 in 500	Reduced
		Ashkenazi Jewish	1 in 27	1 in 2600
		Asian	1 in 126	1 in 12500
		Caucasian	1 in 182	1 in 18100
Tay-Sachs disease (AR) NM_000520.4	HEXA	French Canadian	1 in 27	1 in 2600
···· <u>-</u> ····		Irish	1 in 41	1 in 4000
		Pan-ethnic	1 in 250	1 in 24900
		Sephardic Jewish	1 in 125	1 in 12400
Transient infantile liver failure (AR)	TRMU	Pan-ethnic	≤1 in 500	Reduced
NM_018006.4	Thure	Sephardic Jewish (Yemenite)	1 in 34	1 in 3300
Tyrosine hydroxylase deficiency (AR)	тн	Caucasian	1 in 224	1 in 22300
NM_199292.2		Pan-ethnic	≤1 in 500	Reduced
	FAH *	Ashkenazi Jewish	1 in 143	1 in 2840
Tyrosinemia type I (AR)		French Canadian	1 in 66	1 in 1300
NM_000137.2		French Canadian (Saguenay-Lac-St- Jean)	1 in 16	1 in 300
		Pan-ethnic	1 in 125	1 in 2480
Tyrosinemia type II (AR) NM_000353.2	TAT	Pan-ethnic	1 in 250	1 in 24900
	USH1C *	French Canadian/Acadian	1 in 227	1 in 22600
USH1C-related conditions (AR) NM_005709.3		Pan-ethnic	1 in 353	1 in 3521
NNI_003703.3		Sephardic Jewish	1 in 125	1 in 1241
	USH2A	Caucasian	1 in 70	1 in 6900
USH2A-related conditions (AR)		Pan-ethnic	1 in 112	1 in 11100
NM_206933.2		Sephardic Jewish	1 in 36	1 in 3500
Very long-chain acyl-CoA dehydrogenase deficiency (AR) NM_000018.3	ACADVL	Pan-ethnic	1 in 100	1 in 9900
VRK1-related conditions (AR)		Ashkenazi Jewish	1 in 225	1 in 22400
NM_003384.2	VRK1	Pan-ethnic	≤1 in 500	Reduced



DISORDER (INHERITANCE)	GENE	ETHNICITY	CARRIER FREQUENCY BEFORE SCREENING	CARRIER RESIDUAL RISK AFTER NEGATIVE RESULT
VSX2-related conditions (AR)	VSX2	Pan-ethnic	≤1 in 500	Reduced
NM_182894.2		Sephardic Jewish	1 in 145	1 in 14400
	АТР7В	Ashkenazi Jewish	1 in 67	1 in 3300
		Canary Islander	1 in 25	1 in 1200
Wilson disease (AR) NM 000053.3		Pan-ethnic	1 in 90	1 in 4450
1111_000055.5		Sardinian	1 in 50	1 in 2450
		Sephardic Jewish	1 in 65	1 in 3200
Xeroderma pigmentosum complementation group A	ХРА	Japanese	1 in 100	1 in 9900
(AR) NM_000380.3		Pan-ethnic	1 in 1667	Reduced
Xeroderma pigmentosum complementation group C	XPC	Pan-ethnic	1 in 763	Reduced
(AR) NM_004628.4		Tunisian	1 in 50	1 in 4900
Zellweger spectrum disorder (PEX1-related) (AR) NM_000466.2	PEX1	Pan-ethnic	1 in 144	1 in 14300
Zellweger spectrum disorder (PEX2-related) (AR) NM_000318.2	PEX2	Ashkenazi Jewish	1 in 227	1 in 22600
		Pan-ethnic	≤1 in 500	Reduced
	PEX6	French Canadian	1 in 55	1 in 5400
Zellweger spectrum disorder (PEX6-related) (AR) NM_000287.3		Pan-ethnic	1 in 294	1 in 29300
		Sephardic Jewish	1 in 18	1 in 1700
Zellweger spectrum disorder (PEX10-related) (AR) NM_153818.1	PEX10	Pan-ethnic	1 in 606	Reduced
Zellweger spectrum disorder (PEX12-related) (AR) NM_000286.2	PEX12	Pan-ethnic	1 in 409	1 in 40800

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 below. Enrichment and analysis focus on the coding sequence of the indicated transcripts, 10bp 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.). Pathogenic and Likely Pathogenic variants that do not meet the validated quality thresholds are confirmed. Confirmation technologies may include any of the following: Sanger sequencing, Pacific Biosciences SMRT sequencing, MLPA, MLPA-seq, Array CGH.Array CGH confirmation of NGS CNV calling performed by Invitae Corporation (1400 16th Street, San Francisco, CA 94103, #05D2040778). The following analyses are performed if relevant to the requisition. For GBA and CYP21A2, the reference genome has been modified to mask the sites of polymorphic paralog sequence variants (PSVs) in both the gene and pseudogene. If one or more reportable variants is identified (see Limitations), the gene is amplified by long-range PCR; PacBio sequencing of the long-range amplicons is used to confirm the variant. Gene conversion and fusion events are flagged by our NGS pipeline and reportable pseudogene-derived variants are identified by long-range PCR followed by PacBio sequencing of the long-range amplicons. In some cases, it may not be possible to disambiguate between the gene and pseudogene. 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, triplet repeats are detected by 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. Technical component of confirmatory sequencing is performed by Invitae Corporation (1400 16th Street, San Francisco, CA 94103, #05D2040778).

The following transcripts were used in this analysis. 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: ABCB11 (NM_003742.2), ABCC8 (NM_000352.4), ACAD9 (NM_014049.4), ACADM (NM_000016.5), ACADVL (NM_000018.3), ACAT1 (NM_000019.3), ACOX1 (NM_004035.6), ACSF3 (NM_174917.4), ADA (NM_000022.2), ADAMTS2 (NM_014244.4), ADGRG1 (NM_005682.6), AGA (NM_000027.3), AGL (NM_000642.2), AGPS (NM_003659.3), AGXT (NM_000030.2), AIRE (NM_000383.3), ALDH3A2 (NM_000382.2), ALDOB (NM_000035.3), ALG6 (NM_013339.3), ALMS1 (NM_015120.4), ALPL (NM_000478.5), AMT (NM_000481.3), AQP2 (NM_000486.5), ARG1 (NM_000045.3), ARSA (NM_000487.5), ARSB (NM_000046.3), ASL (NM_000048.3), ASNS (NM_133436.3), ASPA (NM_000049.2), ASS1 (NM_000050.4), ATM (NM_000051.3), ATP6V1B1 (NM_001692.3), ATP7B (NM_000053.3), BBS1 (NM_024649.4), BBS10 (NM_024685.3), BBS12 (NM_152618.2), BBS2 (NM_031885.3), BCKDHA (NM_000709.3), BCKDHB (NM_183050.2), BCS1L (NM_004328.4), BLM (NM_000057.3), BSND (NM_057176.2), CAPN3 (NM_000070.2), CBS (NM_000071.2), CDH23 (NM_022124.5), CEP290 (NM_025114.3), CERKL (NM_001030311.2), CFTR (NM_000492.3), CHRNE (NM_000080.3), CIITA (NM_000246.3), CLN3 (NM_001042432.1), CLN5 (NM_006493.2), CLN6 (NM_017882.2), CLN8 (NM_018941.3), CLRN1 (NM_174878.2), CNGB3 (NM_019098.4), COL27A1 (NM_032888.3), COL4A3 (NM_000091.4), COL4A4 (NM_000092.4), COL7A1 (NM_000094.3), CPS1 (NM_001875.4), CPT1A (NM_001876.3), CPT2 (NM_000098.2), CRB1 (NM_201253.2), CTNS (NM_004937.2), CTSK (NM_000396.3), CYBA (NM_000101.3), CYP11B1 (NM_000497.3), CYP11B2 (NM_000498.3), CYP17A1 (NM_000102.3), CYP19A1 (NM_031226.2), CYP21A2 (NM_000500.7), CYP27A1 (NM_000784.3), DBT (NM_001918.3), DCLRE1C (NM_001033855.2), DHCR7 (NM_001360.2), DHDDS (NM_024887.3), DLD (NM_000108.4), DNAH5 (NM_001369.2), DNAI1 (NM_012144.3), DNAI2 (NM_023036.4), DYSF (NM_003494.3), EIF2B5 (NM_003907.2), ELP1 (NM_003640.3), ERCC6 (NM_000124.3), ERCC8 (NM_000082.3), ESCO2 (NM_001017420.2), ETFA (NM_000126.3), ETFDH (NM_004453.3), ETHE1 (NM_014297.3), EVC (NM_153717.2), EVC2 (NM_147127.4), EVS (NM_001142800.1), FAH (NM_000137.2), FAM161A (NM_001201543.1), FANCA (NM_000135.2), FANCC (NM_000136.2), FANCG (NM_004629.1), FH (NM_000143.3), FKRP (NM_024301.4), FKTN (NM_001079802.1), G6PC (NM_000151.3), GAA (NM_000152.3), GALC (NM_000153.3), GALK1 (NM_000154.1), GALT (NM_000155.3), GAMT (NM_000156.5), GBA (NM_001005741.2), GBE1 (NM_000158.3), GCDH (NM_000159.3), GFM1 (NM_024996.5), GJB2 (NM_004004.5), GLB1 (NM_000404.2), GLDC (NM_000170.2), GLE1 (NM_001003722.1), GNE (NM_001128227.2), GNPTAB (NM_024312.4), GNPTG (NM_032520.4), GNS (NM_002076.3), GRHPR (NM_012203.1), HADHA (NM_000182.4), HAX1 (NM_006118.3), HBA1 (NM_000558.4), HBA2 (NM_000517.4), HBB (NM_000518.4), HEXA (NM_000520.4), HEXB (NM_000521.3), HGSNAT (NM_152419.2), HIV (NM_213653.3), HLCS (NM_000411.6), HMGCL (NM_000191.2), HOGA1 (NM_138413.3), HPS1 (NM_000195.4), HPS3 (NM_032383.4), HSD17B4 (NM_000414.3), HSD3B2 (NM_000198.3), HYAL1 (NM_153281.1), HYLS1 (NM_145014.2), IDUA (NM_000203.4), IVD (NM_002225.3), KCNJ11 (NM_000525.3), LAMA2 (NM_000426.3), LAMA3 (NM_000227.4), LAMB3 (NM_000228.2), LAMC2 (NM_005562.2), LCA5 (NM_181714.3), LDLR (NM_000527.4), LDLRAP1 (NM_015627.2), LHX3 (NM_014564.4), LIFR (NM_002310.5), LIPA (NM_000235.3), LOXHD1 (NM_144612.6), LPL (NM_000237.2), LRPPRC (NM_133259.3), MAN2B1 (NM_000528.3), MCOLN1 (NM_020533.2), MED17 (NM_004268.4), MESP2 (NM_001039958.1), MFSD8 (NM_152778.2), MKS1 (NM_017777.3), MLC1 (NM_015166.3), MMAA (NM_172250.2), MMAB (NM_052845.3), MMACHC (NM_015506.2), MMADHC (NM_015702.2), MPI (NM_002435.2), MPL (NM_005373.2), MPV17 (NM_002437.4), MTHFR (NM_005957.4), MTRR (NM_002454.2), MTTP (NM_000253.3), MUT (NM_000255.3), MYO7A (NM_000260.3), NAGLU (NM_000263.3), NAGS (NM_153006.2), NBN (NM_002485.4), NDRG1 (NM_006096.3), NDUFAF5 (NM_024120.4), NDUFS6 (NM_004553.4), NEB (NM_001271208.1), NPC1 (NM_000271.4), NPC2 (NM_006432.3), NPHS1 (NM_004646.3), NPHS2 (NM_014625.3), NR2E3 (NM_014249.3), NTRK1 (NM_001012331.1), OAT (NM_000274.3), OPA3 (NM_025136.3), PAH (NM_000277.1), PC (NM_000920.3), PCCA (NM_000282.3), PCCB (NM_000532.4), PCDH15 (NM_033056.3), PDHB (NM_000925.3), PEX1 (NM_000466.2), PEX10 (NM_153818.1), PEX12 (NM_000286.2), PEX2 (NM_000318.2), PEX6 (NM_000287.3), PEX7 (NM_000288.3), PFKM (NM_000289.5), PHGDH (NM_006623.3), PKHD1 (NM_138694.3), PMM2 (NM_000303.2), POMGNT1 (NM_017739.3), PPT1 (NM_000310.3), PROP1 (NM_006261.4), PSAP (NM_002778.3), PTS (NM_000317.2), PUS1 (NM_025215.5), PYGM (NM_005609.3), RAB23 (NM_183227.2), RAG2 (NM_000536.3), RAPSN (NM_005055.4), RARS2 (NM_020320.3), RDH12 (NM_152443.2), RMRP (NR_003051.3), RPE65 (NM_000329.2), RPGRIP1L (NM_015272.2), RTEL1 (NM_001283009.1), SACS (NM_014363.5), SAMHD1 (NM_015474.3), SEPSECS (NM_016955.3), SGCA (NM_000023.2), SGCB (NM_000232.4), SGCG (NM_000231.2), SGSH (NM_000199.3), SLC12A3 (NM_000339.2), SLC12A6 (NM_133647.1), SLC17A5 (NM_012434.4), SLC22A5 (NM_003060.3), SLC25A13 (NM_014251.2), SLC25A15 (NM_014252.3), SLC26A2 (NM_000112.3), SLC26A4 (NM_000441.1), SLC35A3 (NM_012243.2), SLC37A4 (NM_001164277.1), SLC39A4 (NM_130849.3), SLC4A11 (NM_032034.3), SLC7A7 (NM_001126106.2), SMARCAL1 (NM_014140.3), SMN1 (NM_000344.3), SMPD1 (NM_000543.4), STAR (NM_000349.2), SUMF1 (NM_182760.3), TAT (NM_000353.2), TCIRG1 (NM_006019.3), TECPR2 (NM_014844.3), TFR2 (NM_003227.3), TGM1 (NM_000359.2), TH (NM_199292.2), TMEM216 (NM_001173990.2), TPP1 (NM_000391.3), TRMU (NM_018006.4), TSFM (NM_001172696.1), TTPA (NM_000370.3), TYMP (NM_001953.4), USH1C



(NM_005709.3), USH2A (NM_206933.2), VPS13A (NM_033305.2), VPS13B (NM_017890.4), VPS45 (NM_007259.4), VRK1 (NM_003384.2), VSX2 (NM_182894.2), WNT10A (NM_025216.2), XPA (NM_000380.3), XPC (NM_004628.4), ZFYVE26 (NM_015346.3).

- Variants of uncertain significance are not included in this report; however, if additional evidence becomes available to indicate that a previously uncertain variant is clinically significant, Invitae will 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) 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 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. In very rare cases (such as circulating hematolymphoid neoplasm, bone marrow transplant, recent blood transfusion, or maternal cell contamination), the analyzed DNA may not represent the patient's constitutional genome.</p>
- COL27A1: Deletion/duplication analysis is not offered for exons 46-47. NBN: Deletion/duplication analysis is not offered for exons 15-16. GALC: Deletion/duplication analysis is not offered for exon 6. MMADHC: Deletion/duplication analysis is not offered for exons 5-6. 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. 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. HBA2: Sequencing analysis is not offered for exons 1-2. USH1C: Deletion/duplication analysis is not offered for exons 5-6. 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.lle237Asn), 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. ALG6: Deletion/duplication analysis is not offered for exons 11-12. 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.Gly241Arg), 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. Sensitivity to detect these variants if they result from complex gene conversion events may be reduced. 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. OAT: Deletion/ duplication analysis is not offered for exon 2. TSFM: Sequencing analysis is not offered for exon 5. FAH: Deletion/duplication analysis is not offered for exon 14. RPGRIP1L: Sequencing analysis is not offered for exon 23. SMN1 or SMN2: NM_000344.3:c.*3+80T>G variant only. 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. VPS13A: Deletion/duplication analysis is not offered for exons 2-3, 27-28.

This report has been reviewed and approved by:

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Reclassification of a genetic variant in the WNT10A gene:

What it means for patients and their families

Why am I receiving this amended report?

Millions of tiny differences exist between the DNA sequences of any two individuals. Although some of these changes, known as variants, can cause disease, most of them do not. Genetic test results, including your carrier screening results, do not mention these "benign" variants because they are not expected to cause health issues.

Yet the scientific community's understanding of which variants do and do not cause disease can change with time.

You are receiving this amended report because the understanding of a variant that we identified in your DNA-the c.682T>A (p.Phe228Ile) variant in the WNT10A gene-has changed and is now classified as "pathogenic (low penetrance)."

Why did the understanding of this variant change?

As you may know, as more people undergo genetic testing, we learn more about how genes vary from person to person. Sometimes this new information will change our understanding of a particular DNA variant. In rare cases, variants thought likely to cause disease may eventually be shown not to cause disease and, conversely, variants thought unlikely to cause disease may be shown to cause disease. When our understanding of a variant changes, we call this a variant reclassification. When this happens at Invitae, all reports that include that variant are automatically updated, and the updated reports are released to the healthcare provider who ordered the test.

Why are you sharing this information with me?

At Invitae, we feel that providing variant reclassifications to healthcare providers and patients is important for a variety of reasons. For carrier screening, the new information may change the reproductive risks for current or future pregnancies or may impact your or your family members' personal risk for disease. Providing amended reports also reduces the need for you to repeat the testing at a later time. Invitae's philosophy is that if we become aware of new medically important information that is relevant to you, that information should not be withheld.

What condition is the c.682T>A (p.Phe228Ile) variant in the WNT10A gene associated with?

The WNT10A gene, where your variant is located, is known to be associated with two conditions.

The first condition is autosomal recessive, which means that a child has to inherit two disease-causing variants—one from each biological parent—to be affected. In this scenario, the child would have a type of **ectodermal dysplasia**, which means abnormal growth of tissues arising from a tissue layer called the ectoderm. Forms of ectodermal dysplasia include odonto-onycho-dermal dysplasia and Schöpf-Schulz-Passarge syndrome, both of which can involve dental anomalies (extra, missing, or malformed teeth), weak hair and nails, and excessive sweating due to abnormal development of sweat glands.^{1,2} Treatment is focused on managing symptoms. With early diagnosis and adequate management, many children with these conditions can have a normal life.

The second condition is autosomal dominant, meaning that a child needs to inherit only one disease-causing genetic change to be affected. Some patients with a single variant in WNT10A may have a few missing teeth, though others show no observable signs or symptoms of disease.³⁻⁵



What does pathogenic (low penetrance) mean?

Pathogenic means that a variant is capable of causing disease. Low penetrance means that only some people with the pathogenic variant will show signs and symptoms of the disease. The c.682T>A (p.Phe228Ile) variant is a pathogenic (low penetrance) variant in the WNT10A gene. This means that not all individuals with this specific genetic change will show the signs or symptoms of a WNT10A-related condition.

Individuals who carry one copy of the c.682T>A (p.Phe228Ile) variant and have no other disease-causing genetic changes typically do not have symptoms, although they may have been missing one or more teeth since birth.⁶

Individuals who carry the c.682T>A (p.Phe228Ile) variant along with another disease-causing variant in the WNT10A gene show signs and symptoms of ectodermal dysplasia somewhere between 15% and 60% of the time.

What does this reclassification mean for me?

If you have a single copy of this variant and no other changes in this gene, it is possible that you may have some mild dental differences, such as missing or underdeveloped teeth. However, not all people with a single variant have noticeable signs or symptoms. If you have a single copy of the variant, we encourage you to inform your doctor. Although it is unlikely to affect your medical care, it's important for your doctor to be aware.

What does the reclassification mean if I am pregnant?

The risk that your child will be affected with a WNT10A-related condition depends on multiple factors, including the number of variants you and your partner carry and the number of variants the child inherits. If you are pregnant, carrier testing for your reproductive partner is recommended, if it has not yet been completed.

If your partner does not have a change in the WNT10A gene, there is a 50% chance that your child will inherit the change. As described above, this typically results in mild dental differences or no signs or symptoms at all.

If your partner does have a change in the WNT10A gene, there is a 25% chance that the child will inherit two diseasecausing variants. In this case, they may have a form of ectodermal dysplasia but, as described above, they may or may not show signs or symptoms of the disease, given the low penetrance of your WNT10A variant. However, there are other scenarios to consider. If your partner carries a WNT10A variant, you should consult a genetic counselor or another qualified healthcare provider to discuss these implications.

What does it mean if I am not pregnant?

If you or any of your children have signs or symptoms of a WNT10A-related condition, you may benefit from a clinical evaluation by a healthcare provider. If you are considering a pregnancy in the future, carrier testing for your reproductive partner is recommended.

Should my family members be tested?

Your biological relatives may wish to consider genetic testing, especially if they have any signs or symptoms of ectodermal dysplasia or are planning a pregnancy themselves. If they are planning a pregnancy and genetic testing shows that they carry the WNT10A variant, carrier testing for their reproductive partners should be considered.

What if I have more questions?

Please send us an email at <u>clientservices@invitae.com</u> or contact Client Services at (800) 436-3037. We are available to discuss any further questions you may have.

References

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