

#### **Donor 5815**

#### **Genetic Testing Summary**

Fairfax Cryobank recommends reviewing this genetic testing summary with your healthcare provider to determine suitability.

Last Updated: 05/09/19

Donor Reported Ancestry: Puerto Rican Jewish Ancestry: No

Genetic Test*	Result	Comments/Donor's Residual Risk**
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Chromosome analysis (karyotype)	Normal male karyotype	No evidence of clinically significant chromosome abnormalities
Hemoglobin evaluation	Normal hemoglobin fractionation and MCV/MCH results	Reduced risk to be a carrier for sickle cell anemia, beta thalassemia, alpha thalassemia trait (aa/ and a-/a-) and other hemoglobinopathies
Cystic Fibrosis (CF) carrier screening	Negative by gene sequencing in the CFTR gene	1/212
Spinal Muscular Atrophy (SMA) carrier screening	Negative for deletions of exon 7 in the SMN1 gene	1/1061
Expanded Genetic Disease Carrier Screening Panel attached- 289 diseases by genotyping and gene sequencing	Negative for genes tested	

<sup>\*</sup>No single test can screen for all genetic disorders. A negative screening result significantly reduces, but cannot eliminate, the risk for these conditions in a pregnancy.

<sup>\*\*</sup>Donor residual risk is the chance the donor is still a carrier after testing negative.



Ordering Practice:

Practice Code:

Fairfax Cryobank -

Report Generated: 2018-02-26

Donor 5815

DOB:

Gender: Male

Ethnicity: Latin American Procedure ID: 113081

Kit Barcode

Specimen: Blood, #115344

Specimen Collection: 2018-02-15 Specimen Received: 2018-02-16 Specimen Analyzed: 2018-02-26

**TEST INFORMATION** 

Test: CarrierMap<sup>SEQ</sup> (Genotyping &

Sequencing)

Panel: CarrierMap Expanded v3 -

Sequencing

Diseases Tested: 289 Genes Tested: 278 Genes Sequenced: 273 Partner Not Tested

#### SUMMARY OF RESULTS: NO MUTATIONS IDENTIFIED

Donor 5815 was not identified to carry any pathogenic mutations in the gene(s) tested.

No pathogenic mutations were identified in the genes tested, reducing but not eliminating the chance to be a carrier for the associated genetic diseases. CarrierMap assesses carrier status for genetic disease via molecular methods including targeted mutation analysis and/or next-generation sequencing; other methodologies such as CBC and hemoglobin electrophoresis for hemoglobinopathies and enzyme analysis for Tay-Sachs disease may further refine risks for these conditions. Results should be interpreted in the context of clinical findings, family history, and/or other testing. A list of all the diseases and mutations screened for is included at the end of the report. This test does not screen for every possible genetic disease.

For additional disease information, please visit recombine.com/diseases. To speak with a Genetic Counselor, call 855.OUR.GENES.

Assay performed by Reprogenetics CIIA ID: 31 D1054821

3 Regent Street, Livingston, NJ 07039

Lab Technician: Bo Chu

Recombine CLIA # 31 D2100763 Reviewed by Pere Colls, PhD, HCLD, Lab Director





#### ADDITIONAL RESULTS: NO INCREASED REPRODUCTIVE RISK

The following results are not associated with an increased reproductive risk.

Disease (Gene)	Donor 5815	Partner Not Tested	
Spinal Muscular Atrophy: SMN1	SMN1 Copy Number: 2 or more		·
Linked (SMN1)*	copies		
	Method: dPCR & Genotyping		

#### \*SMA Risk Information for Individuals with No Family History of SMA

	Detection Rate	Pre-Test Carrier Risk	Post-Test Carrier Risk (2 SMN1 copies)	Post-Test Carrier Risk (3 SMN1 copies)
European	95%	1/35	1/632	1/3,500
Ashkenazi Jewish	90%	1/41	1/350	1/4,000
Asian	93%	1/53	1/628	1/5,000
African American	71%	1/66	1/121	1/3,000
Hispanic	91%	1/117	1/1,061	1/11,000

For other unspecified ethnicities, post-test carrier risk is assumed to be <1%. For individuals with multiple ethnicities, it is recommended to use the most conservative risk estimate.





#### Methods and Limitations

**Genotyping:** Genotyping is performed using the Illumina Infinium Custom HD Genotyping assay to identify mutations in the genes tested. The assay is not validated for homozygous mutations, and it is possible that individuals affected with disease may not be accurately genotyped.

Sequencing: Sequencing is performed using a custom next-generation sequencing (NGS) platform. Only the described exons for each gene listed are sequenced. Variants outside of these regions may not be identified. Some splicing mutations may not be identified. Triplet repeat expansions, intronic mutations, and large insertions and deletions may not be detected. All identified variants are curated, and determination of the likelihood of their pathogenicity is made based on examining allele frequency, segregation studies, predicted effect, functional studies, case/control studies, and other analyses. All variants identified via sequencing that are reported to cause disease in the primary scientific literature will be reported. Variants considered to be benign and variants of unknown significance (VUS) are NOT reported. In the sequencing process, interval drop-out may occur, leading to intervals of insufficient coverage. Intervals of insufficient coverage will be reported if they occur.

Spinal Muscular Atrophy: Carrier status for SMA is assessed via copy number analysis by dPCR and via genotyping. Some individuals with a normal number of SMN1 copies (2 copies) may carry both copies of the gene on the same allele/chromosome; this analysis is not able to detect these individuals. Thus, a normal SMN1 result significantly reduces but does not eliminate the risk of being a carrier. Additionally, SMA may be caused by non-deletion mutations in the SMN1 gene; CarrierMap tests for some, but not all, of these mutations. Some SMA cases arise as the result of de novo mutation events which will not be detected by carrier testing.

Limitations: In some cases, genetic variations other than that which is being assayed may interfere with mutation detection, resulting in false-negative or false-positive results. Additional sources of error include, but are not limited to: sample contamination, sample mix-up, bone marrow transplantation, blood transfusions, and technical errors. The test does not test for all forms of genetic disease, birth defects, and intellectual disability. All results should be interpreted in the context of family history; additional evaluation may be indicated based on a history of these conditions. Additional testing may be necessary to determine mutation phase in individuals identified to carry more than one mutation in the same gene. All mutations included within the genes assayed may not be detected, and additional testing may be appropriate for some individuals.

This test was developed and its performance determined by Recombine, Inc., and it has not been cleared or approved by the U.S. Food and Drug Administration (FDA). The FDA has determined that such clearance or approval is not necessary.



#### Diseases & Mutations Assayed

11-Beta-Hydroxylase-Deficient Congenital Adrenal Hyperplasia (CYP11B1): Mutations (1): of Genotyping | c.1343G>A (p.R448H) Sequencing | NM\_000497:1-9

17-Alpha-Hydroxylase Deficiency (CYP17A1): Mutations (20): of Genotyping | c. 157\_159delTTC (p.53delF), c.316T>C (p.S106P), c.715C>T (p.R239X), c.1024C>A (p.P342T), c.286C>T (p.R96W), c.1040G>A (p.R347H), c.1073G>A (p.R358Q), c.51G>A (p.W17X), c.340T>G (p.F114V), c.347A>T (p.D116V), c.1039C>T (p.R347C), c.1084C>T (p.R362C), c.1216T>C (p.W406R), c.985T>G (p.Y329D), c.601T>A (p.Y201N), c.81C>A (p.Y27X), c.287G>A (p.R96Q), c.1226C>G (p.P409R), c.1250T>G (p.F417C), c.278T>G (p.F93C) Sequencing | NM\_000102:1-8

17-Beta-Hydroxysteroid Dehydrogenase Deficiency (HSD17B3): Mutations (8):  $\sigma$ Genotyping | c.695C>T (p.S232L), c.703A>G (p.M235V), c.239G>A (p.R80Q), c.608C>T (p.A203V), c.238C>T (p.R80W), c.166G>A (p.A56T), c.389A>G (p.N130S), c.803G>A (p.C268Y) Sequencing | NM\_000197:1-11

21-Hydroxylase-Deficient Classical Congenital Adrenal Hyperplasia (CYP21A2): Mutations (1): of Genotyping | c.293-13C>G

21-Hydroxylase-Deficient Nonclassical Congenital Adrenal Hyperplasia (CYP21A2): Mutations (1): O' Genotyping | c.1360C>T (p.P454S)

3-Beta-Hydroxysteroid Dehydrogenase Deficiency (HSD3B2): Mutations (6): 07 Genotyping | c.512G>A (p.W171X), c.742\_747delGTCCGAinsAACTA (p.V248NfsR249X), c.745C>T (p.R249X), c.29C>A (p.A10E), c.424G>A (p.E142K), c.664C>A (p.P222T) Sequencing NM\_000198:2-4

3-Methylcrotonyl-CoA Carboxylase Deficiency: MCCA Related (MCCC1): Mutations (2): of Genotyping | c.1155A>C (p.R385S), c.1310T>C (p.L437P) Sequencing | NM\_020166:1-

3-Methylcrotonyl-CoA Carboxylase Deficiency: MCCB Related (MCCC2): Mutations (8): O Genotyping | c.295G>C (p.E99Q), c.499T>C (p.C167R), c.464G>A (p.R155Q), c.569A>G (p.H190R), c.803G>C (p.R268T), c.838G>T (p.D280Y), c.929C>G (p.P310R), c.1309A>G (p.1437V) Sequencing | NM\_022132:1-17

3-Methylglutaconic Aciduria: Type 3 (OPA3): Mutations (3): of Genotyping | c.415C>T (p.Q139X), c.320\_337delAGCAGCGCCACAAGGAGG (p.Q108\_E113del), c.143-1G>C Sequencing | NM\_025136:1-2

3-Phosphoglycerate Dehydrogenase Deficiency (PHGDH): Mutations (7): of Genotyping c.1468G>A (p.V490M), c.403C>T (p.R135W), c.712delG (p.G238fsX), c.1273G>A (p.V425M), c.1117G>A (p.A373T), c.781G>A (p.V261M), c.1129G>A (p.G377S) Sequencing

5-Alpha Reductase Deficiency (SRD5A2): Mutations (10): of Genotyping | c.736C>T (p.R246W), c.164T>A (p.L55Q), c.344G>A (p.G115D), c.547G>A (p.G183S), c.679C>T (p.R227X), c.682G>A (p.A228T), c.586G>A (p.G196S), c.692A>G (p.H231R), c.635C>G (p.P212R), c.591G>T (p.E197D) Sequencing | NM\_000348:1-5

6-Pyruvoyl-Tetrahydropterin Synthase Deficiency (PTS): Mutations (6): of Genotyping | c.46C>T (p.R16C), c.74G>A (p.R25Q), c.155A>G (p.N52S), c.259C>T (p.P87S), c.286G>A (p.D96N), c.347A>G (p.D116G) Sequencing | NM\_000317:1-6

ARSACS (SACS): Mutations (6): of Genotyping | c.12973C>T (p.R4325X), c.7504C>T (p.R2502X), c.9742T>C (p.W3248R), c.8844delT (p.l2949fs), c.5836T>C (p.W1946R), c.3161T>C (p.F1054S) Sequencing | NM\_014363:2-10

Abetalipoproteinemia (MTTP): Mutations (2): & Genotyping | c.2593G>T (p.G865X), c.2211 delT Sequencing | NM\_000253:2-19

Acrodermatitis Enteropathica (SLC39A4): Mutations (7): & Genotyping | c.1223-1227delCCGGG, c.968-971delAGTC, c.318C>A (p.N106K), c.599C>T (p.P200L), c.1120G>A (p.G374R), c.909G>C (p.Q303H), c.989G>A (p.G330D) Sequencing | NM\_130849:1-12

Acute Infantile Liver Failure: TRMU Related (TRMU): Mutations (5): of Genotyping | c.229T>C (p.Y77H), c.815G>A (p.G272D), c.2T>A (p.M1K), c.835G>A (p.V279M), c.1102-3C>G Sequencing | NM\_018006:1-11

Acyl-CoA Oxidase | Deficiency (ACOX1): Mutations (5): of Genotyping | c.372delCATGCCCGCCTGGAACTT, c.832A>G (p.M278V), c.926A>G (p.Q309R), c.442C>T (p.R148X), c.532G>T (p.G178C) Sequencing | NM\_004035:1-14

Adenosine Deaminase Deficiency (ADA): Mutations (22): & Genotyping | c.986C>T (p.A329V), c.872C>T (p.S291L), c.646G>A (p.G216R), c.632G>A (p.R211H), c.631C>T (p.R211C), c.596A>C (p.Q199P), c.536C>A (p.A179D), c.529G>A (p.V177M), c.467G>A (p.R156H), c.466C>T (p.R156C), c.454C>A (p.L152M), c.445C>T (p.R149W), c.419G>A (p.G140E), c.385G>A (p.V129M), c.320T>C (p.L107P), c.302G>A (p.R101Q), c.302G>T (p.R101L), c.301C>T (p.R101W), c.248C>A (p.A83D), c.220G>T (p.G74C), c.58G>A (p.G20R), c.43C>G (p.H15D) Sequencing | NM\_000022:1-12

Alkaptonuria (HGD): Mutations (14): of Genotyping | c.1111\_1112insC, c.16-1G>A (IVS1-1G>A), c.174delA, c.342+1G>A (IVS5+1G>A), c.1102A>G (p.M368V), c.140C>T (p.S47L), c.688C>T (p.P230S), c.481G>A (p.G161R), c.808G>A (p.G270R), c.899T>G (p.V300G), c.990G>T (p.R330S), c.457\_458insG, c.360T>G (p.C 120W), c.1112A>G (p.H371R) Sequencing | NM\_000187:1-14

Alpha Thalassemia (HBA1, HBA2): Mutations (9): & Genotyping | SEA deletion, c.207C>A

(p.N69K), c.223G>C (p.D75H), c.2T>C, c.207C>G (p.N69K), c.340\_351delCTCCCGCCGAG (p.L114\_E117del), c.377T>C (p.L126P), c.427T>C (p.X143Qext32), c.\*+94A>G

Alpha-1-Antitrypsin Deficiency (SERPINA1): Mutations (4): of Genotyping | c.226\_228delTTC (p.76delF), c.1131 A>T (p.L377F), c.187C>T (p.R63C), c.1096G>A (p.E366K) Sequencing | NM\_001127701:1-7

Alpha-Mannosidosis (MAN2B1): Mutations (3): of Genotyping | c.2426T>C (p.L809P), c.2248C>T (p.R750W), c.1830+1G>C (p.V549\_E610del) Sequencing | NM\_000528:1-24

Alport Syndrome: COL4A3 Related (COL4A3): Mutations (3): On Genotyping | c.4420\_4424delCTTTT, c.4441C>T (p.R1481X), c.4571C>G (p.S1524X) Sequencing | NM\_000091:2-52

Alport Syndrome: COL4A4 Related (COL4A4): Mutations (4): of Genotyping | c.3713C>G (p.S1238X), c.4129C>T (p.R1377X), c.4923C>A (p.C1641X), c.3601G>A (p.G1201S) Sequencing | NM\_000092:2-48

Amegakaryocytic Thrombocytopenia (MPL): Mutations (23): of Genotyping | c.79+2T>A (IVS1+2T>A), c.127C>T (p.R43X), c.305G>C (p.R102P), c.823C>A (p.P275T), c.304C>T (p.R102C), c.376delT (F126Lfs), c.268C>T (p.R90X), c.235\_236delCT (p.L79fs), c.367C>T (p.R123X), c.460T>C (p.W154R), c.1305G>C (p.W435C), c.770G>T (p.R257L), c.407C>T (p.P136L), c.407C>A (p.P136H), c.1781T>G (p.L594W), c.311T>C (p.F104S), c.556C>T (p.Q186X), c.1473G>A (p.W491X), c.1499delT (p.L500fs), c.769C>T (p.R257C), c.1904C>T (p.P635L), c.213-1G>A (IVS2-1G>A), c.1566-1G>T (IVS10-1G>T) Sequencing | NM\_005373:1-

Andermann Syndrome (SLC12A6): Mutations (5): of Genotyping | c.2436delG (p.T813fsX813), c.901delA, c.2023C>T (p.R675X), c.3031C>T (p.R1011X), c.619C>T (p.R207C) Sequencing | NM\_133647:1-25

Antley-Bixler Syndrome (POR): Mutations (4): ♂ Genotyping | c.859G>C (p.A287P), c. 1615G>A (p.G539R), c. 1475T>A (p.V492E), c. 1370G>A (p.R457H) Sequencing

Argininemia (ARG1): Mutations (13): of Genotyping | c.365G>A (p.W122X), c.871C>T (p.R291X), c.869C>G (p.T290S), c.703G>C (p.G235R), c.32T>C (p.I11T), c.413G>T (p.G138V), c.57+1G>A, c.61C>T (p.R21X), c.263\_266delAGAA (p.K88fs), c.77delA (p.E26fs), c.844delC (p.L282fs), c.466-2A>G, c.703G>A (p.G235R) Sequencing | NM\_000045:1-8

Argininosuccinate Lyase Deficiency (ASL): Mutations (7): of Genotyping | c.446+1G>A (IVS5+1G>A), c.857A>G (p.Q286R), c.1135C>T (p.R379C), c.1153C>T (p.R385C), c.283C>T (p.R95C), c.532G>A (p.V178M), c.1060C>T (p.Q354X) Sequencing | NM\_000048:2-17

Aromatase Deficiency (CYP19A1): Mutations (10): of Genotyping | c.1222delC (p.K409fs), c.296+1G>A (IVS3+1G>A), c.468delC, c.629-3C>A (IVS4-3C>A), c.743+2T>C (IVS6+2T>C), c. 1123C>T (p.R375C), c. 1303C>T (p.R435C), c. 1094G>A (p.R365Q), c. 1310G>A (p.C437Y), c.628G>A (p.E210K) Sequencing | NM\_000103:2-10

Arthrogryposis, Mental Retardation, & Seizures (SLC35A3): Mutations (2): & Genotyping | c.1012A>G (p.S338G), c.514C>T (p.Q172X) Sequencing | NM\_001271685:1-8 Asparagine Synthetase Deficiency (ASNS): Mutations (1): of Genotyping | c.1084T>G (p.F362V) Sequencing | NM\_001673:3-13

Aspartylglycosaminuria (AGA): Mutations (7): of Genotyping | c.200\_201delAG, c.488G>C (p.C163S), c.214T>C (p.S72P), c.916T>C (p.C306R), c.904G>A (p.G302R), c.302C>T (p.A101V), c.179G>A (p.G60D) Sequencing | NM\_000027:1-9

Ataxia with Vitamin E Deficiency (TTPA): Mutations (14): of Genotyping | c.744delA, c.575G>A (p.R192H), c.400C>T (p.R134X), c.303T>G (p.H101Q), c.358G>A (p.A120T), c.513\_514insTT (p.T172fs), c.219\_220insAT, c.175C>T (p.R59W), c.421G>A (p.E141K), c.661C>T (p.R221W), c.486delT (p.W163Gfs), c.736G>C (p.G246R), c.205-1G>C, c.306A>G (p.G102G) Sequencing | NM\_000370:2-5

Ataxia-Telangiectasia (ATM): Mutations (20): & Genotyping | c.103C>T (p.R35X), c.1564\_1565delGA (p.E522fs), c.3245delATCinsTGAT (p.H1082fs), c.3576G>A (p.K1192K), c.3894insT, c.5712\_5713insA (p.S1905fs), c.5762+1126A>G, c.5908C>T (p.Q1970X), c.5932G>T (p.E1978X), c.7268A>G (p.E2423G), c.7271T>G (p.V2424G), c.7327C>T (p.R2443X), c.7517\_7520delGAGA (p.R2506fs), c.7630-2A>C, c.7638\_7646delTAGAATTTC (p.R2547\_S2549delRIS), c.7876G>C (p.A2626P), c.7967T>C (p.L2656P), c.8030A>G (p.Y2677C), c.8480T>G (p.F2827C), c.7449G>A (p.W2483X) Sequencing | NM\_000051:2-63

Autosomal Recessive Polycystic Kidney Disease (PKHD1): Mutations (40): & Genotyping c.5895insA (p.L1966fsX1969), c.9689delA (p.D3230fs), c.107C>T (p.T36M), c.1486C>T (p.R496X), c.10412T>G (p.V3471G), c.10658T>C (p.I3553T), c.10174C>T (p.Q3392X), c.9530T>C (p.13177T), c.9053C>T (p.S3018F), c.8870T>C (p.12957T), c.8011C>T (p.R2671X), c.6992T>A (p.I2331K), c.5221G>A (p.V1741M), c.4991C>T (p.S1664F), c.3761\_3762delCCinsG (p.A 1254fs), c.2414C>T (p.P805L), c.664A>G (p.1222V), c.10036T>C (p.C3346R), c.383delC, c.4220T>G (p.L1407R), c.11612G>A (p.W3871X), c.5984A>G (p.E1995G), c.10637delT (p.V3546fs), c.3747T>G (p.C1249W), c.5750A>G (p.Q1917R), c.10865G>A (p.C3622Y), c.50C>T (p.A17V), c.8063G>T (p.C2688F), c.10402A>G (p.I3468V), c.1529delG (p.G510fs), c.657C>T (p.G219G), c.5513A>G (p.Y1838C), c.10856delA (p.K3619fs), c.5381-9T>G (IVS33-9T>G), c.3229-2A>C (IVS28-2A>C), c.10505A>T (p.E3502V), c.2269A>C (p.I757L), c.4165C>A (p.P1389T), c.10364delC (p.S3455fs), c.7350+653A>G (IVS46+653A>G) Sequencing | NM\_138694:2-67

Bardet-Biedl Syndrome: BBS1 Related (BBS1): Mutations (3): ♂ Genotyping | c.851delA, c.1645G>T (p.E549X), c.1169T>G (p.M390R) Sequencing | NM\_024649:1-17

Bardet-Biedl Syndrome: BBS10 Related (BBS10): Mutations (3): O' Genotyping |



c.271\_273ins1bp (p.C91fsX95), c.101G>C (p.R34P), c.931T>G (p.S311A) Sequencing | NM\_024685:1-2

Bardet-Biedl Syndrome: BBS11 Related (TRIM32): Mutations (1): of Genotyping | c.388C>T (p.P130S) Sequencing | NM\_001099679:2

Bardet-Biedl Syndrome: BBS12 Related (BBS12): Mutations (5): of Genotyping | c.335\_337delTAG, c.865G>C (p.A289P), c.1063C>T (p.R355X), c.1114\_1115delTT (p.F372X), c.1483\_1484delGA (p.E495fsX498) Sequencing | NM\_152618:1-2

Bardet-Biedl Syndrome: BBS2 Related (BBS2): Mutations (8): of Genotyping | c.940delA, c.72C>G (p.Y24X), c.224T>G (p.Y75G), c.311A>C (p.D104A), c.1895G>C (p.R632P), c.823C>T (p.R275X), c.814C>T (p.R272X), c.1206\_1207insA (p.R403fs) Sequencing | NM\_031885:1-17

Bare Lymphocyte Syndrome: Type II (CIITA): Mutations (3): of Genotyping | c.1141G>T (p.E381X), c.3317+1G>A (IVS18+1G>A), c.2888+1G>A (IVS13+1G>A) Sequencing | NM\_000246:1-19

Bartter Syndrome: Type 4A (BSND): Mutations (6): 07 Genotyping | c.1A>T, c.22C>T (p.R8W), c.139G>A (p.G47R), c.23G>T (p.R8L), c.28G>A (p.G10S), c.3G>A (p.M1I) Sequencing

Beta Thalassemia (HBB): Mutations (81): 03 Genotyping | c.124\_127delTTCT (p.F42Lfs), c. 17\_18delCT, c.20delA (p.E7Gfs), c.217insA (p.S73Kfs), c.223+702\_444+342del620insAAGTAGA, c.230delC, c.25\_26delAA, c.315+1G>A, c.315+2T>C, c.316-197C>T, c.316-146T>G, c.315+745C>G, c.316-1G>A, c.316-1G>C, c.316-2A>G, c.316-3C>A, c.316-3C>G, c.4delG (p.V2Cfs), c.51delC (p.K 18Rfs), c.93-21G>A, c.92+1G>A, c.92+5G>A, c.92+5G>C, c.92+5G>T, c.92+6T>C, c.93-1G>A, c.93-1G>T, c.-50A>C, c.-78a>g, c.-79A>G, c.-81A>G, c.52A>T (p.K18X), c.-137c>g, c.-138c>t, c.-151C>T, c.118C>T (p.Q40X), c.169G>C (p.G57R), c.295G>A (p.V99M), c.415G>C (p.A139P), c.47G>A (p.W16X), c.48G>A (p.W16X), c.-801>a, c.2T>C, c.75T>A (p.G25G), c.444+111A>G, c.-29G>A, c.68\_74delAAGTTGG, c.92G>C (p.R31T), c.92+1G>T, c.93-15T>G, c.93-1G>C, c.112delT, c.113G>A (p.W38X), c.114G>A (p.W38X), c.126delC, c.444+113A>G, c.250delG, c.225delC, c.383\_385delAGG (p.Q128\_A129delQAinsP), c.321\_322insG (p.N109fs), c.316-1G>T, c.316-2A>C, c.287\_288insA (p.L97fs), c.271G>T (p.E91X), c.203\_204delTG (p.V68Afs), c.154delC (p.P52fs), c.135delC (p.F46fs), c.92+2T>A, c.92+2T>C, c.90C>T (p.G30G), c.84\_85insC (p.L29fs), c.59A>G (p.N20S), c.46delT (p.W16Gfs), c.45\_46insG (p.L16fs), c.36delT (p.T13fs), c.2T>G, c.1A>G (p.M1V), c.-137c>t, c.-136C>G, c.-142C>T, c.-140c>t Sequencing | NM 000518:1-3

Beta-Hexosaminidase Pseudodeficiency (HEXA): Mutations (2): & Genotyping | c.739C>T (p.R247W), c.745C>T (p.R249W) Sequencing | NM\_000520:1-14

Beta-Ketothiolase Deficiency (ACAT1): Mutations (19): & Genotyping | c.1006-1G>C, c.1006-2A>C, c.1083insA, c.826+1G>T, c.278A>G (p.N93S), c.433C>G (p.Q145E), c.814C>T (p.Q272X), c.1136G>T (p.G379V), c.1138G>A (p.A380T), c.547G>A (p.G183R), c.997G>C (p.A333P), c.2T>A (p.M1K), c.935T>C (p.I312T), c.99T>A (p.Y33X), c.149delC (p.T50Nfs), c.253\_255delGAA (p.85delE), c.455G>C (p.G152A), c.380C>T (p.A127V), c.371A>G (p.K124R) Sequencing | NM\_000019:1-12

Biotinidase Deficiency (BTD): Mutations (21): & Genotyping | c.98\_104delGCGGCTGinsTCC (p.C33FfsX68), c.1368A>C (p.Q456H), c.755A>G (p.D252G), c.1612C>T (p.R538C), c.235C>T (p.R79C), c.100G>A (p.G34S), c.1330G>C (p.D444H), c.511 G>A (p.A171T), c.1207T>G (p.F403V), c.470G>A (p.R157H), c.1595C>T (p.T532M), c.1489C>T (p.P497S), c.341G>T (p.G114V), c.1052delC (p.T351fs), c.393delC (p.F131Lfs), c.1049delC (p.A350fs), c.1239delC (p.Y414lfs), c.1240\_1251delTATCTCCACGTC (p.Y414\_V417del), c.278A>G (p.Y93C), c.595G>A (p.V199M), c.933delT (p.S311Rfs) Sequencing | NM\_000060:1-4

Bloom Syndrome (BLM): Mutations (25): O' Genotyping | c.2207\_2212delATCTGAinsTAGATTC (p.Y736Lfs), c.2407insT, c.557\_559delCAA (p.S186X), c.1284G>A (p.W428X), c.1701G>A (p.W567X), c.1933C>T (p.Q645X), c.2528C>T (p.T843I), c.2695C>T (p.R899X), c.3107G>T (p.C1036F), c.2923delC (p.Q975K), c.3558+1G>T, c.3875-2A>G, c.2074+2T>A, c.2343\_2344dupGA (p.781EfsX), c.318\_319insT (p.L107fs), c.380delC (p. 127Tfs), c.3564delC (p. 1188Dfs), c.4008delG (p. 1336Rfs), c.947C>G (p.S316X), c.2193+1\_2193+9del9, c.1642C>T (p.Q548X), c.3143delA (p.1048NfsX), c.356\_357delTA (p.C120Hfs), c.4076+1delG, c.3281C>A (p.S1094X) Sequencing | NM\_000057:2-22

Canavan Disease (ASPA): Mutations (8): of Genotyping | c.433-2A>G, c.854A>C (p.E285A), c.693C>A (p.Y231X), c.914C>A (p.A305E), c.71A>G (p.E24G), c.654C>A (p.C218X), c.2T>C (p.M1T), c.79G>A (p.G27R) Sequencing | NM\_000049:1-6

Carnitine Palmitoyltransferase IA Deficiency (CPT1A): Mutations (10): of Genotyping | c.1079A>G (p.E360G), c.1361A>G (p.D454G), c.1241C>T (p.A414V), c.1436C>T (p.P479L), c.2126G>A (p.G709E), c.2129G>A (p.G710E), c.1493A>G (p.Y498C), c.1339C>T (p.R447X), c.2156G>A (p.G719D), c.96T>G (p.Y32X) Sequencing | NM\_001876:2-19

Carnitine Palmitoyltransferase II Deficiency (CPT2): Mutations (19): of Genotyping | c.109\_110insGC, c.1238\_1239delAG, c.1737delC, c.1923\_1935delGAAGGCCTTAGAA, c.534\_558delGAACCCTGCAAAAAGTGACACTATCinsT, c.1649A>G (p.Q550R), c.1883A>C (p.Y628S), c.359A>G (p.Y120C), c.983A>G (p.D328G), c.149C>A (p.P50H), c.1810C>T (p.P604S), c.1891C>T (p.R631C), c.338C>T (p.S113L), c.370C>T (p.R124X), c.680C>T (p.P227L), c.1646G>A (p.G549D), c.452G>A (p.R151Q), c.520G>A (p.E174K), c.1148T>A (p.F383Y) Sequencing | NM\_000098:1-5

Carnitine-Acylcarnitine Translocase Deficiency (SLC25A20): Mutations (7): & Genotyping | c. 199-10T>G (IVS2-10T>G), c.897\_898insC (p.N300fs), c.496C>T (p.R166X), c.84delT (p.H29Tfs), c.713A>G (p.Q238R), c.576G>A (p.W192X), c.106-2A>T Sequencing |

NM 000387:1-9

Carpenter Syndrome (RAB23): Mutations (2): of Genotyping | c.434T>A (p.L145X), c.408\_409insT (p.136fsX) Sequencing | NM\_016277:2-7

Cartilage-Hair Hypoplasia (RMRP): Mutations (2): & Genotyping | n.71A>G, c.263G>T Sequencing | NR\_003051:1

Cerebrotendinous Xanthomatosis (CYP27A1): Mutations (14): of Genotyping | c.1263+1G>A, c.844+1G>A, c.1016C>T (p.T339M), c.1183C>T (p.R395C), c.1420C>T (p.R474W), c.1435C>T (p.R479C), c.379C>T (p.R127W), c.819delT (p.D273fs), c.1214G>A (p.R405Q), c.1421G>A (p.R474Q), c.434G>A (p.G145E), c.583G>T (p.E195X), c.646G>C (p.A216P), c.1183C>A (p.R395S) Sequencing | NM\_000784:1-9

Chediak-Higashi Syndrome (LYST): Mutations (4): of Genotyping | c.3085C>T (p.Q1029X), c.9590delA (p.Y3197fs), c.1902\_1903insA (p.A635Sfs), c.118\_119insG (p.A40fs) Sequencing NM\_000081:3-53

Cholesteryl Ester Storage Disease (LIPA): Mutations (4): & Genotyping | c.1024G>A (p.G342R), c.894G>A (p.Q298X), c.883C>T (p.H295Y), c.652C>T (p.R218X) Sequencing | NM 001127605:2-10

Choreoacanthocytosis (VPS13A): Mutations (1): ♂ Genotyping | c.6058delC (p.P2020fs) Sequencing | NM\_033305:1-72

Chronic Granulomatous Disease: CYBA Related (CYBA): Mutations (12): of Genotyping | c.354C>A (p.S118R), c.467C>A (p.P156Q), c.281A>G (p.H94R), c.7C>T (p.Q3X), c.70G>A (p.G24R), c.244delC (p.P82fs), c.171\_172insG (p.K58fs), c.373G>A (p.A125T), c.174delG (p.K58fs), c.385\_388delGAGC (p.E129SfsX61), c.369+1G>A (IVS5+1G>A), c.71G>A (p.G24E) Sequencing | NM\_000101:1-5

Citrin Deficiency (SLC25A13): Mutations (8): & Genotyping | c.1180G>A (p.G394S), c.674C>A (p.S225X), c.1766G>A (p.R589Q), c.851\_854delGTAT (p.R284fs), c.1802\_1803insA (p.Y601fs), c.1180+1G>A, c.1663\_1664insGAGATTACAGGTGGCTGCCCGGG (p.A555fs), c.1314+1G>A Sequencing | NM\_001160210:1-18

Citrullinemia: Type I (ASS1): Mutations (11): of Genotyping | c.1194-1G>C, c.970+5G>A, c.928A>C (p.K310Q), c.835C>T (p.R279X), c.1085G>T (p.G362V), c.470G>A (p.R157H), c.539G>A (p.S180N), c.970G>A (p.G324S), c.535T>C (p.W179R), c.1168G>A (p.G390R), c.421-2A>G (IVS6-2A>G) Sequencing | NM\_000050:3-16

Classical Galactosemia (GALT): Mutations (18): ♂ Genotyping | c.253-2A>G, c.563A>G (p.Q188R), c.626A>G (p.Y209C), c.404C>T (p.S135L), c.413C>T (p.T138M), c.505C>A (p.Q169K), c.997C>G (p.R333G), c.607G>A (p.E203K), c.855G>T (p.K285N), c.1138T>C (p.X380R), c.221T>C (p.L74P), c.425T>A (p.M142K), c.512T>C (p.F171S), c.584T>C (p.L195P), c. 134\_138delCAGCT, c.-1039\_753del3162, c.820+51\_\*789del2294ins 12, c.404C>G (p.S135W) Sequencing | NM\_000155:1-11

Cockayne Syndrome: Type A (ERCC8): Mutations (3): of Genotyping | c.966C>A (p.Y322X), c.37G>T (p.E13X), c.479C>T (p.A160V) Sequencing | NM\_000082:1-12

Cockayne Syndrome: Type B (ERCC6): Mutations (7): of Genotyping | c.1550G>A (p.W517X), c.2203C>T (p.R735X), c.1518delG (p.K506Nfs), c.1357C>T (p.R453X), c.972\_973insA (p.E325Rfs), c.1974\_1975insTGTC (p.T659fs), c.1034\_1035insT (p.K345fs) Sequencing | NM\_000124:2-21

Cohen Syndrome (VPS13B): Mutations (9): of Genotyping | c.6578T>G (p.L2193R), c.7051C>T (p.R2351X), c.4471G>T (p.E1491X), c.2911C>T (p.R971X), c.7934G>A (p.G2645D), c.10888C>T (p.Q3630X), c.8459T>C (p.I2820T), c.9259\_9260insT (p.L3087fs), c.3348\_3349delCT (p.C1117fx) Sequencing | NM\_017890:2-51,53-62

Combined Pituitary Hormone Deficiency: PROP1 Related (PROP1): Mutations (11): o' Genotyping | c.218G>A (p.R73H), c.150delA (p.G50fsX), c.358C>T (p.R120C), c. 112\_124delTCGAGTGCTCCAC (p.S38fsX), c.2T>C, c. 157delA (p.R53fsX), c.212G>A (p.R71H), c.217C>T (p.R73C), c.582G>A (p.W194X), c.109+1G>T, c.301delAG (p.S101fsX) Sequencing NM 006261:1-3

Congenital Disorder of Glycosylation: Type 1A: PMM2 Related (PMM2): Mutations (5): of Genotyping | c.357C>A (p.F119L), c.422G>A (p.R141H), c.338C>T (p.P113L), c.691G>A (p.V231 M), c.470T>C (p.F157S) Sequencing | NM\_000303:1-8

Congenital Disorder of Glycosylation: Type 1B: MPI Related (MPI): Mutations (1): of Genotyping | c.884G>A (p.R295H) Sequencing | NM\_002435:1-8

Congenital Disorder of Glycosylation: Type 1C: ALG6 Related (ALG6): Mutations (4): d' Genotyping | c.257+5G>A, c.895\_897delATA, c.998C>T (p.A333V), c.1432T>C (p.S478P) Sequencing | NM\_013339:2-15

Congenital Ichthyosis: ABCA12 Related (ABCA12): Mutations (8): of Genotyping | c.4139A>G (p.N1380S), c.4951G>A (p.G1651S), c.4142G>A (p.G1381E), c.4541G>A (p.R1514H), c.4615G>A (p.E1539K), c.7323delC (p.V2442Sfs), c.6610C>T (p.R2204X), c.3535G>A (p.G1179R) Sequencing | NM\_173076:1-53

Congenital Insensitivity to Pain with Anhidrosis (NTRK1): Mutations (12): of Genotyping | c.1729G>C (p.G577R), c.2339G>C (p.R780P), c.25C>T (p.Q9X), c.1076A>G (p.Y359C), c. 1759A>G (p.M587V), c.207\_208delTG (p.E70Afs), c. 1550G>A (p.G517E), c.717+4A>T, c.429-1G>C, c.1660delC (p.R554fs), c.2046+3A>C, c.2084C>T (p.P695L) Sequencing

Congenital Lipoid Adrenal Hyperplasia (STAR): Mutations (12): of Genotyping | c. 178+1\_178+2insT (IVS2+3insT), c.201\_202delCT, c.466-11T>A (IVS4-11T>A), c.64+1G>T (IVS1+1G>T), c.562C>T (p.R188C), c.772C>T (p.Q258X), c.545G>A (p.R182H), c.545G>T



(p.R182L), c.559G>A (p.V187M), c.650G>C (p.R217T), c.749G>A (p.W250X), c.64+1G>A Sequencing | NM\_000349:1-7

Congenital Myasthenic Syndrome: CHRNE Related (CHRNE): Mutations (12): 0" Genotyping | c.1327delG (p.E443fs), c.865C>T (p.L289F), c.911delT (p.L304fs), c.344+1G>A, c.850A>C (p.T284P), c.422C>T (p.P141L), c.250C>G (p.R84G), c.500G>T (p.R167L), c.991C>T (p.R331W), c.37G>A (p.G13R), c.613\_619delTGGGCCA (p.W205fs), c.1353\_1354insG (p.N452Efs) Sequencing | NM\_000080:1-12

Congenital Myasthenic Syndrome: DOK7 Related (DOK7): Mutations (6): & Genotyping | c.601C>T (p.R201X), c.539G>C (p.G180A), c.548\_551delTCCT (p.F183fs), c.1263\_1264insC (p.S422fs), c.101-1G>T, c.331+1G>T Sequencing | NM\_173660:3-7

Congenital Myasthenic Syndrome: RAPSN Related (RAPSN): Mutations (11): o' Genotyping | c.264C>A (p.N88K), c.41T>C (p.114P), c.807C>A (p.Y269X), c.548\_549insGTTCT (p.183fs), c.46\_47insC (p.16fs), c.133G>A (p.V45M), c.848T>C (p.1283P), c.484G>A (p.E162K), c.490C>T (p.R164C), c.-210A>G, c.193-15C>A (IVS1-15C>A) Sequencing | NM\_005055:1-8

Congenital Neutropenia: Recessive (HAX1): Mutations (6): & Genotyping | c.121\_125insG, c.130\_131insA, c.431insG, c.91delG, c.256C>T (p.R86X), c.568C>T (p.Q190X) Sequencing | NM\_006118:1-7

Corneal Dystrophy and Perceptive Deafness (SLC4A11): Mutations (8): o' Genotyping | c.1459\_1462delTACGinsA (p.487\_488delYAinsT), c.2313\_2314insTATGACAC, c.554\_561delGCTTCGCC (p.R185fs), c.2566A>G (p.M856V), c.1463G>A (p.R488K), c.2528T>C (p.1843P), c.637T>C (p.S213P), c.2321+1G>A Sequencing | NM\_001174090:1-20

Corticosterone Methyloxidase Deficiency (CYP11B2): Mutations (3): of Genotyping | c.1492A>G (p.T498A), c.541C>T (p.R181W), c.1382T>C (p.L461P) Sequencing | NM 000498:1-9

Crigler-Najjar Syndrome (UGT1A1): Mutations (11): Ø Genotyping | c.508\_513delTTC (p.170delF), c.1070A>G (p.Q357R), c.1021C>T (p.R341X), c.1124C>T (p.S375F), c.840C>A (p.C280X), c.991C>T (p.Q331X), c.923G>A (p.G308E), c.1198A>G (p.N400D), c.992A>G (p.Q331R), c.44T>G (p.L15R), c.524T>A (p.L175Q) Sequencing | NM\_000463:1-5

Cystic Fibrosis (CFTR): Mutations (149): of Genotyping | c.1029delC, c.1153\_1154insAT, c.1477delCA, c.1519\_1521delATC (p.507delI), c.1521\_1523delCTT (p.508delF), c.1545\_1546delTA (p.Y515Xfs), c.1585-1G>A, c.164+12T>C, c.1680-886A>G, c.1680-1G>A, c.1766+1G>A, c.1766+1G>T, c.1766+5G>T, c.1818del84, c.1911delG, c. 1923delCTCAAAACTinsA, c. 1973delGAAATTCAATCCTinsAGAAA, c. 2052delA (p. K684fs), c.2052insA (p.Q685fs), c.2051\_2052delAAinsG (p.K684SfsX38), c.2174insA, c.261delTT, c.2657+5G>A, c.273+1G>A, c.273+3A>C, c.274-1G>A, c.2988+1G>A, c.3039delC, c.3140-26A>G, c.325delTATinsG, c.3527delC, c.3535delACCA, c.3691delT, c.3717+12191C>T, c.3744delA, c.3773\_3774insT (p.L1258fs), c.442delA, c.489+1G>T, c.531delT, c.579+1G>T, c.579+5G>A (IVS4+5G>A), c.803delA (p.N268fs), c.805\_806delAT (p.I269fs), c.933\_935delCTT (p.311delF), c.946delT, c.1645A>C (p.S549R), c.2128A>T (p.K710X), c.1000C>T (p.R334W), c.1013C>T (p.T338I), c.1364C>A (p.A455E), c.1477C>T (p.Q493X), c.1572C>A (p.C524X), c.1654C>T (p.Q552X), c.1657C>T (p.R553X), c.1721C>A (p.P574H), c.2125C>T (p.R709X), c.223C>T (p.R75X), c.2668C>T (p.Q890X), c.3196C>T (p.R1066C), c.3276C>G (p.Y1092X), c.3472C>T (p.R1158X), c.3484C>T (p.R1162X), c.349C>T (p.R117C), c.3587C>G (p.S1196X), c.3712C>T (p.Q1238X), c.3764C>A (p.S1255X), c.3909C>G (p.N1303K), c.1040G>A (p.R347H), c.1040G>C (p.R347P), c.1438G>T (p.G480C), c.1558G>T (p.V520F), c.1624G>T (p.G542X), c.1646G>A (p.S549N), c.1646G>T (p.S549I), c.1652G>A (p.G551D), c.1675G>A (p.A559T), c.1679G>C (p.R560T), c.178G>T (p.E60X), c.254G>A (p.G85E), c.271G>A (p.G91R), c.274G>T (p.E92X), c.3209G>A (p.R1070Q), c.3266G>A (p.W1089X), c.3454G>C (p.D1152H), c.350G>A (p.R117H), c.3611G>A (p.W1204X), c.3752G>A (p.S1251N), c.3846G>A (p.W1282X), c.3848G>T (p.R1283M), c.532G>A (p.G178R), c.988G>T (p.G330X), c.1090T>C (p.S364P), c.3302T>A (p.M1101K), c.617T>G (p.L206W), c.14C>T (p.P5L), c.19G>T (p.E7X), c.171G>A (p.W57X), c.313delA (p.1105fs), c.328G>C (p.D110H), c.580-1G>T, c.1055G>A (p.R352Q), c.1075C>A (p.Q359K), c.1079C>A (p.T360K), c.1647T>G (p.S549R), c.1976delA (p.N659fs), c.2290C>T (p.R764X), c.2737\_2738insG (p.Y913X), c.3067\_3072delATAGTG (p.I1023\_V1024delT), c.3536\_3539delCCAA (p.T1179fs), c.3659delC (p.T1220fs), c.54-5940\_273+10250del21080bp (p.S18fs), c.4364C>G (p.S1455X), c.4003C>T (p.L1335F), c.2538G>A (p.W846X), c.200C>T (p.P67L), c.4426C>T (p.Q1476X), c.1116+1G>A, c. 1986\_1989delAACT (p.T663R), c.2089\_2090insA (p.R697Kfs), c.2215delG (p.V739Y), c.263T>G (p.L196X), c.3022delG (p.V1008S), c.3908dupA (p.N1303Kfs), c.658C>T

Cystinosis (CTNS): Mutations (14): Of Genotyping | c.18\_21delGACT, c.198\_218delTATTACTATCCTTGAGCTCCC, c.283G>T (p.G95X), c.414G>A (p.W138X), c.506G>A (p.G169D), c.613G>A (p.D205N), c.473T>C (p.L158P), c.329G>T (p.G110V), c.416C>T (p.S139F), c.589G>A (p.G197R), c.969C>G (p.N323K), c.1015G>A (p.G339R), c.39155\_848del57119, c.199\_219delATTACTATCCTTGAGCTCCCC (p.I67\_P73del) Sequencing | NM 001031681:1.3-13

(p.Q220X), c.868C>T (p.Q290X), c.1526delG (p.G509fs), c.2908+1085\_3367+260del7201,

c.1408\_1417delGTGATTATGG (p.V470fs), c.1585-8G>A, c.2909G>A (p.G970D), c.653T>A

c.3731G>A (p.G1244E), c.535C>A (p.Q179K), c.3368-2A>G, c.455T>G (p.M152R),

(p.L218X), c.1175T>G (p.V392G), c.3139\_3139+1delGG, c.3717+4A>G (IVS22+4A>G)

c.1610\_1611delAC (p.D537fs), c.3254A>G (p.H1085R), c.496A>G (p.K166E)

c.11C>A (p.S4X), c.3878\_3881 delTATT (p.V1293fs), c.3700A>G (p.11234V), c.416A>T (p.H139L), c.366T>A (p.Y122X), c.3767\_3768insC (p.A1256fs), c.613C>T (p.P205S), c.293A>G (p.Q98R),

Cystinuria: Non-Type I (SLC7A9): Mutations [15]: of Genotyping | c.508G>A (p.V170M), c.313G>A (p.G105R), c.583G>A (p.G195R), c.775G>A (p.G259R), c.997C>T (p.R333W), c.131T>C (p.144T), c.782C>T (p.P261L), c.695A>G (p.Y232C), c.544G>A (p.A182T), c.368C>T (p.T123M), c.614\_615insA (p.K205fs), c.604+2T>C, c.605-3C>A (IVS5-3C>A), c.1445C>T (p.P482L), c.368\_369delCG (p.T123fs) Sequencing | NM\_001243036:2-13

Cystinuria: Type I (SLC3A1): Mutations (10): o\* Genotyping | c.1400T>C (p.M467T), c.2033T>C (p.L678P), c.542G>A (p.R181Q), c.1955C>G (p.T652R), c.1843C>A (p.P615T), c.1085G>A (p.R362H), c.1597T>A (p.Y533N), c.647C>T (p.T216M), c.808C>T (p.R270X), c.452A>G (p.Y151C) Sequencing | NM\_000341:1-10

D-Bifunctional Protein Deficiency (HSD17B4): Mutations (6): & Genotyping | c.46G>A (p.G16S), c.63G>T (p.L21F), c.422\_423delAG, c.652G>T (p.V218L), c.1369A>T (p.N457Y), c.1369A>G (p.N457D) Sequencing | NM\_000414:1-24

Diabetes: Recessive Permanent Neonatal (ABCC8): Mutations (2): of Genotyping | c.215A>G (p.N72S), c.1144G>A (p.E382K) Sequencing | NM\_000352:1-39

Du Pan Syndrome (GDF5): Mutations (4): of Genotyping | c.1309delTTG, c.1306C>A (p.P436T), c.1133G>A (p.R378Q), c.1322T>C (p.L441P) Sequencing | NM\_000557:1-2

Dyskeratosis Congenita: RTEL1 Related (RTEL1): Mutations (5): O' Genotyping | c.2869C>T (p.R981W), c.2920C>T (p.R974X), c.1548G>T (p.M5161), c.2216G>T (p.G763V), c.3791G>A (p.R1264H) Sequencing | NM\_001283009:2-35

Dystrophic Epidermolysis Bullosa: Recessive (COL7A1): Mutations (11): 0\* Genotyping | c,2470\_2471 insG, c,5820G>A (p,P1940P), c,933C>A (p,Y311X), c,4039G>C (p,G1347R), c,8393T>A (p,M2798K), c,425A>G (p,K142R), C,8441-

14\_8435delGCTCTTGGCTCCAGGACCCCT, c.4783-1G>A, c.7344G>A (p.V2448X), c.4991G>C (p.G1664A), c.497\_498insA (p.V168GfsX179) Sequencing | NM\_000094:1-118

Ehlers-Danlos Syndrome: Type VIIC (ADAMTS2): Mutations (2): of Genotyping | c.673C>T (p.Q225X), c.2384G>A (p.W795X) Sequencing | NM\_014244:2-22

Ellis-van Creveld Syndrome: EVC Related (EVC): Mutations (10): o' Genotyping | c.919T>C (p.S307P), c.1694delC (p.A565VfsX23), c.734delT (p.L245fs), c.910-911insA (p.R304fs), c.2635C>T (p.Q879X), c.1868T>C (p.L623Q), c.

1858\_1879delTTGGGCCGACTGGGCGGCCTC (p.1620\_1626del), c.1886+5G>T, c.1098+1G>A, c.1018C>T (p.R340X) Sequencing | NM\_153717:2-21

Ellis-van Creveld Syndrome: EVC2 Related (EVC2): Mutations {1}:  $\sigma$  Genotyping | c.3025C>T (p.Q1009X) Sequencing | NM\_147127:1-22

Enhanced S-Cone (NR2E3): Mutations (5): Of Genotyping | c.932G>A (p.R311Q), c.227G>A (p.R76Q), c.119-2A>C, c.226C>T (p.R76W), c.747+1G>C (IVS5+1G>C) Sequencing | NM\_016346:1-8

Ethylmalonic Aciduria (ETHE1): Mutations (4): d' Genotyping | c.505+1G>T, c.487C>T (p.R163W), c.3G>T (p.M1I), c.488G>A (p.R163Q) Sequencing | NM\_014297:1-7

Familial Chloride Diarrhea (SLC26A3): Mutations (6): o' Genotyping | c.344delT (p.11151), c.559G>T (p.G187X), c.951delGGT (p.V318del), c.1386G>A (p.W462X), c.371A>T (p.H124L), c.2023\_2025dupATC (p.I675L) Sequencing | NM\_000111:2-21

Familial Dysautonomia (IKBKAP): Mutations (4): of Genotyping | c.2204+6T>C, c.2741C>T (p.P914L), c.2087G>C (p.R696P), c.2128C>T (p.Q710X) Sequencing | NM\_003640:2-37

Familial Hyperinsulinism: Type 1: ABCC8 Related (ABCC8): Mutations (11): of Genotyping | c.3989-9G>A, c.4159\_4161 delTTC (p.1387delF), c.4258C>T (p.R1420C), c.4477C>T (p.R1493W), c.2147G>T (p.G716V), c.4055G>C (p.R1352P), c.560T>A (p.V187D), c.4516G>A (p.E1506K), c.2506C>T (p.Q836X), c.579+2T>A, c.1333-1013A>G (IVS8-1013A>G) Sequencing | NM\_000352:1-39

Familial Hyperinsulinism: Type 2: KCNJ11 Related (KCNJ11): Mutations (6): o' Genotyping | c.776A>G (p.H259R), c.36C>A (p.Y12X), C.C761T (p.P254L), c.G-134T, c.844G>A (p.E282K), c.440T>C (p.L147P) Sequencing | NM\_000525:1

Familial Mediterranean Fever (MEFV): Mutations (12): c<sup>3</sup> Genotyping | c.2076\_2078delAAT (p.692dell), c.2080A>G (p.M694V), c.2084A>G (p.K695R), c.1437C>G (p.F479L), c.800C>T (p.T267I), c.1958G>A (p.R653H), c.2040G>A (p.M680I), c.2040G>C (p.M680I), c.2082G>A (p.M694I), c.2230G>T (p.A744S), c.2282G>A (p.R761H), c.2177T>C (p.V726A) Sequencing | NM\_000243:1-10

Fanconi Anemia: Type A (FANCA): Mutations (10): & Genotyping | c.295C>T (p.Q99X), c.1115\_1118deITTGG, c.3720\_3724deIAAACA (p.E1240Dfs), c.513G>A (p.W171X), c.1606deIT (p.S536fs), c.3558\_3559insG (p.R1187Efs), c.1615deIG (p.D539fs), c.890\_893deIGCTG (p.C297fs), c.2172\_2173insG (p.T724fs), c.4275deIT (p.R1425fs) Sequencing | NM\_000135:1-43

Fanconi Anemia: Type C (FANCC): Mutations (8): d' Genotyping | c.456+4A>T, c.67delG, c.37C>T (p.Q13X), c.553C>T (p.R185X), c.1661T>C (p.L554P), c.1642C>T (p.R548X), c.66G>A (p.W22X), c.65G>A (p.W22X) Sequencing | NM\_000136:2-15

Fanconi Anemia: Type G (FANCG): Mutations (5): o' Genotyping | c.1480+1G>C, c.307+1G>C, c.1794\_1803delCTGGATCCGT (p.W599Pfs), c.637\_643delTACCGCC (p.Y213K+4X), c.925-2A>G Sequencing | NM\_004629:1-14

Fanconi Anemia: Type J (BRIP1): Mutations (1): of Genotyping | c.2392C>T (p.R798X) Sequencing | NM\_032043:2-20

Fumarase Deficiency (FH): Mutations (1):  $\sigma$  Genotyping | c.1431\_1433insAAA Sequencing | NM\_000143:1-10

 $GM1\text{-}Gangliosidoses \ (GLB1): \ \textit{Mutations (17)}: \ \textit{O}^{\text{T}} \ \textit{Genotyping | c.1480-2A>G},$ 

Sequencing | NM\_000492:1-27



c.75+2\_75+3insT, c.1772A>G (p.Y591C), c.947A>G (p.Y316C), c.1051C>T (p.R351X), c.1369C>T (p.R457X), c.145C>T (p.R49C), c.202C>T (p.R68W), c.245C>T (p.T82M), c.601C>T (p.R201C), c.622C>T (p.R208C), c.1370G>A (p.R457Q), c.176G>A (p.R59H), c.367G>A (p.G123R), c.152T>C (p.151T), c.1771T>A (p.Y591N), c.1577\_1578insG Sequencing | NM\_000404:1-16

GRACILE Syndrome (BCS1L): Mutations (12): d\* Genotyping | c.232A>G (p.S78G), c.103G>C (p.G35R), c.148A>G (p.T50A), c.166C>T (p.R56X), c.133C>T (p.R45C), c.296C>T (p.P99L), c.464G>C (p.R155P), c.547C>T (p.R183C), c.548G>A (p.R183H), c.550C>T (p.R184C), c.830G>A (p.S277N), c.1057G>A (p.V353M) Sequencing | NM\_004328:1-9

Galactokinase Deficiency (GALK1): Mutations (7): of Genotyping | c.1144C>T (p.Q382X), c.1045G>A (p.G349S), c.1031C>T (p.T344M), c.238G>T (p.E80X), c.94G>A (p.V32M), c.82C>A (p.P28T), c.593C>T (p.A198V) Sequencing | NM\_000154:1-8

Gaucher Disease (GBA): Mutations (6): & Genotyping | c.84\_85insG, c.1226A>G (p.N409S), c.1343A>T (p.D448V), c.1504C>T (p.R502C), c.1297G>T (p.V433L), c.1604G>A (n.R535H)

Gitelman Syndrome (SLC12A3): Mutations (11): & Genotyping | c.1926-1G>T, c.2883+1G>T, c.1046C>T (p.P348L), c.1763C>T (p.A588V), c.622C>T (p.R208W), c.1889G>T (p.G629V), c.1961G>A (p.R654H), c.1868T>C (p.L623P), c.1180+1G>T (IVS9+1G>T), c.1670-191C>T, c.2548+253C>T Sequencing | NM\_000339:1-26

Globoid Cell Leukodystrophy (GALC): Mutations [10]: of Genotyping | c.1153G>T (p.E385X), c.857G>A (p.G286D), c.2002A>C (p.T668P), c.1700A>C (p.Y567S), c.1586C>T (p.T529M), c.1472delA (p.K491fs), c.913A>G (p.1305V), c.683\_694delATCTCTGGGAGTinsCTC (p.N228\_S232del5insTP), c.246A>G (p.182M), c.1161+6555\_\*9573del31670bp Sequencing | NM\_000153:2-17

Glutaric Acidemia: Type I (GCDH): Mutations (8): of Genotyping | c.1204C>T (p.R402W), c.1262C>T (p.R421V), c.743C>T (p.P248L), c.1093G>A (p.E365K), c.877G>A (p.A293T), c.1083-2A>C (IVS10-2A>C), c.680G>C (p.R227P), c.1198G>A (p.V400M) Sequencing | NM\_000159:2-12

Glutaric Acidemia: Type IIA (ETFA): Mutations (5): of Genotyping | c.797C>T (p.7266M), c.470T>G (p.V157G), c.346G>A (p.G116R), c.809\_811delTAG (p.V270\_A271delinsA), c.963+1delG Sequencing | NM\_000126:1-12

Glutaric Acidemia: Type IIB (ETFB): Mutations (2): of Genotyping | c.764G>A (p.R255Q), c.655G>A (p.D219N) Sequencing | NM\_001014763:1-5, NM\_001985:1

Glutaric Acidemia: Type IIC (ETFDH): Mutations [8]: O Genotyping | c.1448C>T (p.P483L), c.2T>C (p.M1T), c.250G>A (p.A84T), c.524G>T (p.R175L), c.380T>A (p.L127H), c.524G>A (p.R175H), c.1130T>C (p.L377P), c.36delA (p.A12fs) Sequencing | NM\_004453:1-13

Glycine Encephalopathy: AMT Related (AMT): Mutations (6): & Genotyping | c.959G>A (p.R320H), c.878-1G>A, c.826G>C (p.D276H), c.574C>T (p.Q192X), c.139G>A (p.G47R), c.125A>G (p.H42R) Sequencing | NM\_000481:1-9

Glycine Encephalopathy: GLDC Related (GLDC): Mutations (5): & Genotyping | c.2284G>A (p.G762R), c.2266\_2268delTTC (p.756delF), c.1691G>T (p.S564I), c.1545G>C (p.R515S), c.2T>C (p.M1T) Sequencing | NM\_000170:1-25

Glycogen Storage Disease: Type IA (G6PC): Mutations [13]: a<sup>o</sup> Genotyping | c.376\_377insTA, c.79delC, c.979\_981delTTC (p.327delF), c.1039C>T (p.Q347X), c.247C>T (p.R83C), c.724C>T (p.Q242X), c.248G>A (p.R83H), c.562G>C (p.G188R), c.648G>T, c.809G>T (p.G270V), c.113A>T (p.D38V), c.975delG (p.L326fs), c.724delC Sequencing | NM\_000151:1-5

Glycogen Storage Disease: Type IB (SLC37A4): Mutations (5): & Genotyping | c.1042\_1043delCT, c.796G>T (p.G266C), c.1016G>A (p.G339D), c.1099G>A (p.A367T), c.352T>C (p.W118R) Sequencing | NM\_001164277:3-11

Glycogen Storage Disease: Type II (GAA): Mutations (13): & Genotyping | c.1935C>A (p.D645E), c.2560C>T (p.R854X), c.-32-13T>G (IVS1-13T>G), c.525delT (p.E176Rfs), c.710C>T (p.A237V), c.896T>G (p.L299R), c.953T>C (p.M318T), c.1561G>A (p.E521K), c.1585\_1586delTCinsGT (p.S529V), c.1634C>T (p.P545L), c.1927G>A (p.G643R), c.2173C>T (p.R725W), c.2707\_2709delK (p.903delK) Sequencing | NM\_001079804:2-20

Glycogen Storage Disease: Type III (AGL): Mutations (15): O' Genotyping | c.17\_18delAG, c.4455delT [p.S1486fs], c.1222C>T (p.R408X), c.16C>T (p.Q6X), c.1384delG [p.V462X], c.2039G>A (p.W680X), c.2590C>T (p.R864X), c.2681+1G>A, c.3439A>G [p.R1147G], c.3682C>T (p.R1228X), c.3965delT (p.V1322AfsX27), c.3980G>A (p.W1327X), c.4260-12A>G (IVS32-12A>G), c.4342G>C (p.G1448R), c.2681+1G>T Sequencing | NM\_000642:2-34

Glycogen Storage Disease: Type IV (GBE1): Mutations (3):  $\sigma$  Genotyping | c.986A>C (p.Y329S), c.691+2T>C (IVS5+2T>C), c.986A>G (p.Y329C) Sequencing | NM\_000158:1-16

Glycogen Storage Disease: Type V (PYGM): Mutations (10): & Genotyping | c.2128\_2130delTTC (p.710delF), c.1627A>T (p.K543X), c.1628A>C (p.K543T), c.148C>T (p.K50X), c.255C>A (p.Y85X), c.613G>A (p.G205S), c.2392T>C (p.W798R), c.1827G>A (p.K609K), c.632delG (p.S211fs), c.808C>T (p.R270X) Sequencing | NM\_005609:1-20

Glycogen Storage Disease: Type VII (PFKM): Mutations (4): & Genotyping | c.450+1G>A, c.329G>T (p.R110L), c.283C>T (p.R95X), c.2214delC (p.P739Qfs) Sequencing | NM\_001166686:2-25

Guanidinoacetate Methyltransferase Deficiency (GAMT): Mutations (4): & Genotyping | c.506G>A (p.C169Y), c.327G>A, c.309\_310insCCGGGACTGGGCC (p.199\_A103fs),

c.148A>C (p.M50L) Sequencing | NM\_000156:1-6

HMG-CoA Lyase Deficiency (HMGCL): Mutations (7): d\* Genotyping | c.914\_915delTT, c.122G>A (p.R41Q), c.208G>C (p.V70L), c.835G>A (p.E279K), c.561+1G>A, c.109G>T (p.E37X), c.561+1G>T Sequencing | NM\_000191:1-9

Hemochromatosis: Type 2A: HFE2 Related (HFE2): Mutations (1):  $\sigma^a$  Genotyping | c.959G>T (p.G320V) Sequencing | NM\_213653:2-4

Hemochromatosis: Type 3: TFR2 Related (TFR2): Mutations (4): d' Genotyping | c.2069A>C (p.Q690P), c.750C>G (p.Y250X), c.515T>A (p.M172K), c.88\_89insC (p.E60X) Sequencing | NM\_003227:1-18

Hemoglobinopathy: Hb C (HBB): Mutations (1):  $\sigma$  Genotyping | c.19G>A (p.E7K) Sequencing | NM\_000518:1-3

Hemoglobinopathy: Hb D (HBB): Mutations (1): & Genotyping | c.364G>C (p.E122Q) Sequencing | NM\_000518:1-3

Hemoglobinopathy: Hb E (HBB): Mutations (1): ♂ Genotyping | c.79G>A (p.E27K) Sequencing | NM\_000518:1-3

Hemoglobinopathy: Hb O (HBB): Mutations (1):  $\sigma^a$  Genotyping | c.364G>A (p.E122K) Sequencing | NM\_000518:1-3

Hereditary Fructose Intolerance (ALDOB): Mutations (10): & Genotyping | c.357\_360delAAAC, c.1005C>G (p.N335K), c.524C>A (p.A175D), c.448G>C (p.A150P), c.612T>G (p.Y204X), c.865\_867delCTT (p.289delL), c.720C>A (p.C240X), c.442T>C (p.W148R), c.178C>T (p.R60X), c.10C>T (p.R4X) Sequencing | NM\_000035:2-9

Hereditary Spastic Paraplegia: TECPR2 Related (TECPR2): Mutations (1): of Genotyping | c.3416delT (p.L1139fs) Sequencing | NM\_014844:2-20

Herlitz Junctional Epidermolysis Bullosa: LAMA3 Related (LAMA3); Mutations (1): o\* Genotyping | c.1981C>T (p.R661X) Sequencing | NM\_000227:1-38

Herlitz Junctional Epidermolysis Bullosa: LAMB3 Related (LAMB3): Mutations (6): o\* Genotyping | c.3024delT, c.124C>T (p.R42X), c.1903C>T (p.R635X), c.430C>T (p.R144X), c.727C>T (p.Q243X), c.3247C>T (p.Q1083X) Sequencing | NM\_000228:2-23

Herlitz Junctional Epidermolysis Bullosa: LAMC2 Related (LAMC2): Mutations (1): & Genotyping | c.283C>T (p.R95X) Sequencing | NM\_005562:1-23

Hermansky-Pudlak Syndrome: Type 1 (HPS1): Mutations (1): & Genotyping | c.1470\_1486dup16 (p.H497Qfs) Sequencing | NM\_000195:3-20

Hermansky-Pudlak Syndrome: Type 3 (HPS3): Mutations (4): of Genotyping | c.1189C>T {p.R397W}, c.1691+2T>G, c.2589+1G>C, c.1163+1G>A Sequencing | NM\_032383:1-17

Hermansky-Pudlak Syndrome: Type 4 (HPS4): Mutations (7): of Genotyping | c.1876C>T (p.Q626X), c.526C>T (p.Q176X), c.957\_958insGCTTGTCCAGATGGCAGGAAGGAG (p.E319\_N320ins8), c.634C>T (p.R212X), c.397G>T (p.E133X), c.649G>T (p.E217X), c.2039delC (p.P680fs) Sequencing | NM\_152841:1-12

Holocarboxylase Synthetase Deficiency (HLCS): Mutations (7): of Genotyping | c.1795+5G>A (IVS 10+5G>A), c.780delG, c.710T>C (p.L237P), c.1522C>T (p.R508W), c.1648G>A (p.V550M), c.1513G>C (p.G505R), c.772\_781delACAAGCAAGG (p.T258fs) Sequencing | NM\_001242785:4-12

Homocystinuria Caused by CBS Deficiency (CBS): Mutations (8): of Genotyping | c.919G>A (p.G307S), c.833T>C (p.1278T), c.1006C>T (p.R336C), c.959T>C (p.V320A), c.797G>A (p.R266K), c.572C>T (p.T191 M), c.341C>T (p.A114V), c.969G>A (p.W324X) Sequencing | NM\_001178008:3-17

Hurler Syndrome (IDUA): Mutations (8): of Genotyping | c.1598C>G (p.P533R), c.208C>T (p.Q70X), c.1205G>A (p.W402X), c.979G>C (p.A327P), c.266G>A (p.R89Q), c.1960T>G (p.X654G), c.152G>A (p.G51D), c.1037T>G (p.L346R) Sequencing | NM\_000203:2-8,11-14

Hypophosphatasia (ALPL): Mutations (5): of Genotyping | c.1559delT, c.1133A>T (p.D378V), c.1001G>A (p.G334D), c.571G>A (p.E191K), c.979T>C (p.F327L) Sequencing | NM\_000478:2-12

Inclusion Body Myopathy: Type 2 (GNE): Mutations (3):  $\sigma$  Genotyping | c.2228T>C (p.M743T), c.1807G>C (p.V603L), c.131G>C (p.C44S) Sequencing | NM\_001128227:1-12 Infantile Cerebral and Cerebellar Atrophy (MED17): Mutations (1):  $\sigma$  Genotyping | c.1112T>C (p.L371P) Sequencing | NM\_004268:1-12

Isolated Microphthalmia: VSX2 Related (VSX2): Mutations (4): 6" Genotyping | c.599G>A (p.R200Q), c.599G>C (p.R200P), c.679C>T (p.R227W), c.371-1G>A Sequencing | NM\_182894:1-5

Isovaleric Acidemia (IVD): Mutations (1): O' Genotyping | c.941C>T (p.A314V) Sequencing |

Joubert Syndrome (TMEM216): Mutations (2): of Genotyping | c.218G>T (p.R73L), c.218G>A (p.R73H) Sequencing | NM\_001173991:1-5

Lamellar Ichthyosis: Type 1 (TGM1): Mutations (1): & Genotyping | c.877-2A>G (IVS5-2A>G) Sequencing | NM\_000359:2-15

Laryngoonychocutaneous Syndrome (LAMA3): Mutations (1): & Genotyping | c.151\_152insG (p.V51GfsX3) Sequencing | NM\_000227:1-38

Leber Congenital Amaurosis: GUCY2D Related (GUCY2D): Mutations (3): of



Genotyping | c.1694T>C (p.F565S), c.2943delG (p.G982V), c.387delC (p.P130Lfx) Sequencing | NM 000180:2-19

Leber Congenital Amaurosis: LCA5 Related (LCA5): Mutations (3): of Genotyping | c.835C>T (p.Q279X), c.1476\_1477insA (p.P493TfsX1), c.1151delC Sequencing | NM 001122769:2-8

Leber Congenital Amaurosis: RDH12 Related (RDH12): Mutations (6): & Genotyping | c.565C>T (p.Q189X), c.184C>T (p.R62X), c.464C>T (p.T155I), c.677A>G (p.Y226C), c.146C>T (p.T49M), c.295C>A (p.L99I) Sequencing | NM\_152443:3-9

Leigh Syndrome: French-Canadian (LRPPRC): Mutations (1):  $\sigma$  Genotyping | c.1061C>T (p.A354V) Sequencing | NM\_133259:1-38

Leukoencephalopathy with Vanishing White Matter: EIF2B5 Related (EIF2B5): Mutations (9): of Genotyping | c.338G>A (p.R113H), c.271A>G (p.T91A), c.1882T>C (p.W628R), c.1157G>T (p.G386V), c.584G>A (p.R195H), c.925G>C (p.V309L), c.944G>A (p.R315H), c.166T>G (p.F56V), c.167T>G (p.F56C) Sequencing | NM\_003907:1-16

Leydig Cell Hypoplasia (Luteinizing Hormone Resistance) (LHCGR): Mutations (13): of Genotyping | c.1822\_1827delCTGGTT (p.608\_609dellV), c.1777G>C (p.A593P), c.1660C>T (p.R554X), c.1060G>A (p.E354K), c.1635C>A (p.C545X), c.391T>C (p.C131R), c.1027T>A (p.C343S), c.1627T>C (p.C543R), c.1505T>C (p.L502P), c.430G>T (p.V144F), c.1847C>A (p.S616Y), c.455T>C (p.1152T), c.537-3C>A Sequencing | NM\_000233:1-11

Limb-Girdle Muscular Dystrophy: Type 2A {CAPN3}: Mutations (6): & Genotyping | c.1715G>A {p.R572Q}, c.1469G>A {p.R490Q}, c.550delA {p.T184fs}, c.2306G>A {p.R769Q}, c.2362\_2363delAGinsTCATCT {p.R788Sfs}, c.1525G>T {p.V509F} Sequencing | NM\_000070:1-24

Limb-Girdle Muscular Dystrophy: Type 2B (DYSF): Mutations (5): Of Genotyping | c.4989\_4993delGCCCGinsCCCC (p.E1663fs), c.2833delG (p.A945fs), c.5830C>T (p.R1944X), c.2271C>A (p.Y758X), c.5174+5G>A Sequencing | NM\_001130987:1-56

Limb-Girdle Muscular Dystrophy: Type 2C (SGCG): Mutations (4): o' Genotyping | c.848G>A (p.C283Y), c.787G>A (p.E263K), c.525delT (p.F175fsX), c.87\_88insT (p.G30fs) Sequencing | NM\_000231:2-8

Limb-Girdle Muscular Dystrophy: Type 2D (SGCA): Mutations (1): O' Genotyping | c.229C>T (p.R77C) Sequencing | NM\_000023:1-9

Limb-Girdle Muscular Dystrophy: Type 2E (SGCB): Mutations (6): o\* Genotyping | c.341C>T (p.S114F), c.452C>G (p.T151R), c.272G>C (p.R91P), c.272G>T (p.R91L), c.299T>A (p.M100K), c.323T>G (p.L108R) Sequencing | NM\_000232:2-6

Limb-Girdle Muscular Dystrophy: Type 2F (SGCD): Mutations (5): d' Genotyping | c.493C>T (p.R165X), c.89G>A (p.W30X), c.784G>A (p.E262K), c.391G>C (p.A131P), c.653delC (p.A218fs) Sequencing | NM\_001128209:2-8

Limb-Girdle Muscular Dystrophy: Type 21 (FKRP): Mutations (1): & Genotyping | c.826C>A (p.1276I) Sequencing | NM\_001039885:1-4

Lipoprotein Lipase Deficiency (LPL): Mutations (1): of Genotyping | c.644G>A (p.G215E) Sequencing | NM\_000237:1-10

Long-Chain 3-Hydroxyacyl-CoA Dehydrogenase Deficiency (HADHA): Mutations (2): o' Genotyping | c.1132C>T (p.G378X), c.1528G>C (p.E510G) Sequencing | NM\_000182:1-20 Lysinuric Protein Intolerance (SLC7A7): Mutations (4): o' Genotyping | c.1228C>T (p.R410X), c.726G>A (p.W242X), c.1384\_1385insATCA (p.R462fs), c.895-2A>T Sequencing |

MTHFR Deficiency: Severe (MTHFR): Mutations (6): & Genotyping | c.1721T>G (p.V574G), c.1408G>T (p.E470X), c.1166G>A (p.W389X), c.652G>T (p.V218L), c.523G>A (p.A175T), c.474A>T (p.G158G) Sequencing | NM\_005957:2-12

Malonyl-CoA Decarboxylase Deficiency (MLYCD): Mutations (5): & Genotyping | c.560C>G (p.S187X), c.8G>A (p.G3D), c.1064\_1065delTT (p.F355fs), c.949-14A>G, c.638\_641delGTGA (p.S213fs) Sequencing | NM\_012213:1-5

Maple Syrup Urine Disease: Type 1A (BCKDHA): Mutations (4): of Genotyping | c.860\_867delGAGGCCCC, c.868G>A (p.G290R), c.1312T>A (p.Y438N), c.288+1G>A Sequencing | NM\_000709:1-9

Maple Syrup Urine Disease: Type 1B (BCKDHB): Mutations (6): of Genotyping | c.1114G>T (p.E372X), c.548G>C (p.R183P), c.832G>A (p.G278S), c.970C>T (p.R324X), c.487G>T (p.E163X), c.853C>T (p.R285X) Sequencing | NM\_183050:1-10

Maple Syrup Urine Disease: Type 2 (DBT): Mulations (15): o\* Genotyping | c.670G>T (p.E224X), c.581C>G (p.S194X), c.1355A>G (p.H452R), c.294C>G (p.198M), c.1448G>T (p.X4831), c.75\_76delAT (p.C26Wfs), c.901C>T (p.R301C), c.363\_364delCT (p.Y122lfs), c.1193T>C (p.L398P), c.1169A>G (p.D390G), c.1209+5G>C (IVS9+5G>C), c.1232C>A (p.P411Q), c.939G>C (p.K313N), c.788T>G (p.M263R), c.1202T>C (p.1401T) Sequencing | NM\_001918:1-11

Maple Syrup Urine Disease: Type 3 (DLD): Mutations (8): of Genotyping | c.104\_105insA, c.685G>T (p.G229C), c.214A>G (p.K72E), c.1081A>G (p.M361V), c.1123G>A (p.E375K), c.1178T>C (p.1393T), c.1463C>T (p.P488L), c.1483A>G (p.R495G) Sequencing | NM\_000108:1-14

Maroteaux-Lamy Syndrome (ARSB): Mulations (6): & Genotyping | c.629A>G (p.Y210C), c.1178A>C (p.H393P), c.284G>A (p.R95Q), c.944G>A (p.R315Q), c.1143-8T>G, c.1143-1G>C Sequencing | NM\_000046:1-8

Meckel Syndrome: Type 1 (MKS1): Mutations (5): of Genotyping | c.1408-35\_1408-

7del29 (p.G470fs), c.80+2T>C (IVS1+2T>C), c.1024+1G>A (IVS11+1G>A), c.417G>A (p.E139X), c.50insCCGGG (p.D19AfsX) Sequencing | NM\_017777:1-18

Medium-Chain Acyl-CoA Dehydrogenase Deficiency (ACADM): Mutations (8): of Genotyping | c.985A>G (p.K329E), c.362C>T (p.T1211), c.583G>A (p.G195R), c.799G>A (p.G267R), c.199T>C (p.Y67H), c.262C>T (p.L88F), c.616C>T (p.R206C), c.617G>A (p.C206H) Sequencing | NM\_001127328:1-12

Megalencephalic Leukoencephalopathy (MLC1): Mutations (6): O' Genotyping | c.176G>A (p.G59E), c.278C>T (p.S93L), c.135\_136insC (p.C46fsX), c.908\_918delTGCTGCTGCTGinsGCA (p.V303GfsX96), c.880C>T (p.P294S), c.178-10T>A Sequencing | NM\_139202:2-12

Metachromatic Leukodystrophy (ARSA): Mutations (18): of Genotyping | c.1210+1G>A, c.465+1G>A (IVS2+1G>A), c.862A>C (p.T288P), c.1136C>T (p.P379L), c.1283C>T (p.P428L), c.827C>T (p.T276M), c.542T>G (p.1181S), c.1232C>T (p.T411I), c.769G>C (p.D257H), c.739G>A (p.G247R), c.641C>T (p.A214V), c.302G>A (p.G101D), c.293C>T (p.S98F), c.257G>A (p.R86Q), c.263G>A (p.G88D), c.1114C>T (p.R372W), c.292\_293delTCinsCT (p.S98L), c.302G>T (p.G101V) Sequencing | NM\_001085425:2-9

Methylmalonic Acidemia: MMAA Related (MMAA): Mutations (14): d' Genotyping | c.64C>T (p.R22X), c.161G>A (p.W54X), c.266T>C (p.L89P), c.283C>T (p.Q95X), c.358C>T (p.Q120X), c.397C>T (p.Q133X), c.433C>T (p.R145X), c.503delC (p.T168MfsX9), c.562G>C (p.G188R), c.650T>A (p.L217X), c.653G>A (p.G218E), c.733+1G>A, c.988C>T (p.R330X), c.1076G>A (p.R359Q) Sequencing | NM\_172250:2-7

Methylmalonic Acidemia: MMAB Related (MMAB): Mutations (11): of Genotyping | c.700C>T (p.Q234X), c.656A>G (p.Y219C), c.572G>A (p.R191Q), c.571C>T (p.R191W), c.569G>A (p.R190H), c.568C>T (p.R190C), c.556C>T (p.R186W), c.403G>A (p.A135T), c.291-1G>A, c.287T>C (p.196T), c.197-1G>T Sequencing | NM\_052845:1-9

Methylmalonic Acidemia: MUT Related (MUT): Mutations (23): of Genotyping | c.2150G>T (p.G717V), c.2099T>A (p.M700K), c.2080C>T (p.R694W), c.2054T>G (p.L685R), c.1867G>A (p.G623R), c.1280G>A (p.G427D), c.1106G>A (p.R369H), c.1105C>T (p.R369C), c.1097A>G (p.N366S), c.935G>T (p.G312V), c.691T>A (p.Y231N), c.655A>T (p.N219Y), c.643G>A (p.G215S), c.607G>A (p.G203R), c.572C>A (p.A191E), c.521T>C (p.F174S), c.322C>T (p.R108C), c.313T>C (p.W105R), c.299A>G (p.Y100C), c.284C>G (p.P95R), c.281G>T (p.G94V), c.278G>A (p.R93H), c.643G>T (p.G215C) Sequencing | NM\_000255:2-13

Methylmalonic Aciduria and Homocystinuria: Type cblC (MMACHC): Mutations (5):  $\sigma$  Genotyping | c.271\_272insA (p.R91KfsX14), c.331C>T (p.R111X), c.394C>T (p.R132X), c.482G>A (p.R161Q), c.609G>A (p.W203X) Sequencing | NM\_015506:1-4

Mitochondrial Complex I Deficiency: NDUFS6 Related (NDUFS6): Mutations (1): o\* Genotyping | c.344G>A (p.C115Y) Sequencing | NM\_004553:1-4

Mitochondrial DNA Depletion Syndrome: MNGIE Type {TYMP}: Mutations (6): o\* Genotyping | c.866A>C (p.E289A), c.433G>A (p.G145R), c.665A>G (p.K222R), c.457G>A (p.G153S), c.516+2T>C (IVS4+2T>C), c.1425\_1426insC (p.S476lfs) Sequencing | NM 001257989:2-8.10

Mitochondrial Myopathy and Sideroblastic Anemia (PUS1): Mutations (2): of Genotyping | c.430C>T (p.R144W), c.658G>T (p.E220X) Sequencing | NM\_025215:1-6

Mitochondrial Trifunctional Protein Deficiency: HADHB Related (HADHB): Mutations (7): & Genotyping | c.182G>A (p.R61H), c.788A>G (p.D263G), c.740G>A (p.R247H), c.1331G>A (p.R444K), c.1364T>G (p.V455G), c.776\_777insT (p.G259fs), c.1175C>T (p.A392V) Sequencing | NM\_000183:2-16

Morquio Syndrome: Type A (GALNS): Mutations (6): OT Genotyping | c.205T>G (p.F69V), c.485C>T (p.S162F), c.1156C>T (p.R386C), c.901G>T (p.G301C), c.337A>T (p.1113F), c.178G>A (p.D60N) Sequencing | NM\_000512:2-14

Morquio Syndrome: Type B (GLB1): Mutations (8): O<sup>3</sup> Genotyping | c.1527G>T (p.W509C), c.1313G>A (p.G438E), c.1445G>A (p.R482H), c.247T>C (p.Y83H), c.1444C>T (p.R482C), c.1498A>G (p.T500A), c.1223A>C (p.Q408P), c.817\_818delTGinsCT (p.W273L) Sequencing | NM\_000404:1-16

Mucolipidosis: Type II/III (GNPTAB): Mutations (3): O' Genotyping | c.3503\_3504delTC (p.L1168QfsX5), c.3565C>T (p.R1189X), c.1120T>C (p.F374L) Sequencing | NM\_024312:1-21

Mucolipidosis: Type IV (MCOLN1): Mutations (5): of Genotyping | c.-1015\_788del6433, c.406-2A>G, c.1084G>T (p.D362Y), c.304C>T (p.R102X), c.244delC (p.L82fsX) Sequencing | NM 020533:1-14

Multiple Pterygium Syndrome (CHRNG): Multions (6): Ø Genotyping | c.715C>T (p.R239C), c.13C>T (p.R239C), c.13C>T (p.R257K), c.320T>G (p.V107G), c.401\_402delCT (p.P134fs), c.1408C>T (p.R470X), c.136C>T (p.R46X) Sequencing | NM\_005199:1-12

Multiple Sulfatase Deficiency (SUMF1): Mutations (1): of Genotyping | c.463T>C (p.S155P) Sequencing | NM\_182760:1-9

 $\label{eq:Muscle-Eye-Brain Disease (POMGNT1): Mutations $$\{3\}$: $\sigma^*$ Genotyping $\mid c.1539+1$ G>A , $c.1324C>T \{p.R442C\}, c.1478C>G \{p.P493R\} Sequencing $\mid NM_001243766:2-23$ }$ 

Navajo Neurohepatopathy (MPV17): Mutations (1): & Genotyping | c.149G>A (p.R50Q) Sequencing | NM\_002437:2-8

Nemaline Myopathy: NEB Related (NEB): Mutations (2): of Genotyping | c.7434\_7536del2502bp, c.8890-2A>G (IVS63-2A>G) Sequencing | NM\_001164508:63-66,86,95-96,103,105,143,168-172, NM\_004543:3-149



Nephrotic Syndrome: Type 1 (NPHS1): Mutations (5): of Genotyping | c.121\_122delCT (p.L41Dfs), c.1481delC, c.3325C>T (p.R1109X), c.3478C>T (p.R1160X), c.2335-1G>A Sequencing | NM\_004646:1-29

Nephrotic Syndrome: Type 2 (NPHS2): Mutations (27): O' Genotyping | c.976\_977insA (p.T326fsX345), c.964C>T (p.R322X), c.948delT (p.A317L), c.871C>T (p.R291W), c.868G>A (p.V290M), c.862G>A (p.A288T), c.855\_856delAA (p.Q285fsX302), c.851C>T (p.A284V), c.779T>A (p.V260E), c.714G>T (p.R238S), c.706\_714del CTAGAGAGG (p.L236\_R238del), c.622G>A (p.A208T), c.555delT (p.F185fsX186), c.538G>A (p.V180M), c.503G>A (p.R168H), c.502C>A (p.R168S), c.502C>T (p.R168C), c.479A>G (p.D160G), c.467delT (p.L156fsX180), c.467\_468insT (p.L156fsX166), c.419delG (p.G140fsX180), c.413G>A (p.R138Q), c.412C>T (p.R138X), c.353C>T (p.P118L), c.274G>T (p.G92C), c.104\_105insG (p.G35fsX69), c.85G>A (p.A29T) Sequencing | NM\_014625:1-8

Neuronal Ceroid-Lipofuscinosis: CLN5 Related (CLN5): Mutations (7): of Genotyping | c.1175\_1176delAT (p.Y392X), c.225G>A (p.W75X), c.835G>A (p.D279N), c.335G>A (p.R112H), c.377G>A (p.C126Y), c.1054G>T (p.E352X), c.1121A>G (p.Y374C) Sequencing | NM 006493:1-4

Neuronal Ceroid-Lipofuscinosis: CLN6 Related (CLN6): Mutations (8): of Genotyping | c.663C>G (p.Y221X), c.460\_462delATC (p.I154del), c.368G>A (p.G123D), c.308G>A (p.R103Q), c.214G>T (p.E72X), c.200T>C (p.L67P), c.139C>T (p.L47F), c.17G>C (p.R6T) Sequencing | NM\_017882:2-7

Neuronal Ceroid-Lipofuscinosis: CLN8 Related (CLN8): Mutations (4): O' Genotyping | c.70C>G (p.R24G), c.789G>C (p.W263C), c.88G>C (p.A30P), c.610C>T (p.R204C) Sequencing

Neuronal Ceroid-Lipofuscinosis: MFSD8 Related (MFSD8): Mutations (2): & Genotyping | c.881 C>A (p.T294K), c.754+2T>A Sequencing | NM\_152778:2-13

Neuronal Ceroid-Lipofuscinosis: PPT1 Related (PPT1): Mutations (8): ♂ Genotyping | c.223A>C (p.T75P), c.364A>T (p.R122W), c.451C>T (p.R151X), c.29T>A (p.L10X), c.656T>A (p.L219Q), c.322G>C (p.G108R), c.236A>G (p.D79G), c.134G>A (p.C45Y) Sequencing

Neuronal Ceroid-Lipofuscinosis: TPP1 Related (TPP1): Mutations (9): Of Genotyping | c.523-1G>A, c.509-1G>C, c.622C>T (p.R208X), c.851G>T (p.G284V), c.1340G>A (p.R477H), c.1094G>A (p.C365Y), c.1093T>C (p.C365R), c.857A>G (p.N286S), c.616C>T (p.R206C) Sequencing | NM\_000391:1-13

Niemann-Pick Disease: Type A (SMPD1): Mutations (6): of Genotyping | c.996delC, c.1493G>T (p.R498L), c.911T>C (p.L304P), c.1267C>T (p.H423Y), c.1734G>C (p.K578N), c.1493G>A (p.R498H) Sequencing | NM\_000543:1-6

Niemann-Pick Disease: Type B (SMPD1): Mutations (3): of Genotyping | c.1828\_1830delCGC (p.610delR), c.880C>A (p.Q294K), c.1280A>G (p.H427R) Sequencing |

Niemann-Pick Disease: Type C1 (NPC1): Mutations (14): of Genotyping | c.2783A>C (p.Q928P), c.3263A>G (p.Y1088C), c.3467A>G (p.N1156S), c.3107C>T (p.T1036M), c.3182T>C (p.11061T), c.2974G>C (p.G992R), c.2932C>T (p.R978C), c.2848G>A (p.V950M), c.2665G>A (p.V889M), c.2324A>C (p.Q775P), c.1133T>C (p.V378A), c.530G>A (p.C177Y), c.337T>C (p.C113R), c.2974G>T (p.G992W) Sequencing | NM\_000271:1-25

Niemann-Pick Disease: Type C2 (NPC2): Mutations (11): & Genotyping | c.58G>T (p.E20X), c.436C>T (p.Q146X), c.358C>T (p.P120S), c.352G>T (p.E118X), c.332delA (p.N1111fs), c.295T>C (p.C99R), c.199T>C (p.S67P), c.190+5G>A, c.141C>A (p.C47X), c.133C>T (p.Q45X), c.115G>A (p.V39M) Sequencing | NM\_006432:1-5

Nijmegen Breakage Syndrome (NBN): Mutations (1): of Genotyping | c.657\_661 delACAAA (p.K219fs) Sequencing | NM\_002485:1-16

Nonsyndromic Hearing Loss and Deafness: GJB2 Related (GJB2): Mutations (29): o Genotyping | c.167delT, c.235delC, c.313\_326delAAGTTCATCAAGGG, c.358delGAG (p. 120delE), c.35delG, c.370C>T (p.Q124X), c.427C>T (p.R143W), c.109G>A (p.V37I), c.231 G>A (p.W77X), c.551 G>C (p.R184P), c.71 G>A (p.W24X), c.229T>C (p.W77R), c.269T>C (p.L90P), c.617A>G (p.N206S), c.299\_300delAT (p.H100Rfs), c.283G>A (p.V95M), c.134G>A (p.G45E), c.139G>T (p.E47X), c.35G>T, c.487A>G (p.M163V), c.250G>C (p.V84L), c.44A>C (p.K15T), c.334\_335delAA (p.K112fs), c.516G>A (p.W172X), c.290\_291insA (p.Y97fs), c.439G>A (p.E147K), c.-23+1G>A, c.550C>T (p.R184W), c.-259C>T Sequencing | NM\_004004:1-2

Nonsyndromic Hearing Loss and Deafness: LOXHD1 Related (LOXHD1): Mutations (2): of Genotyping | c.2008C>T (p.R670X), c.4714C>T (p.R1572X) Sequencing | NM\_144612:1-40

Nonsyndromic Hearing Loss and Deafness: MYO15A Related (MYO15A): Mutations (10): of Genotyping | c.453\_455delCGAinsTGGACGCCTGGTCGGGCAGTGG (p.E152GfsX81), c.7801A>T (p.K2601X), c.6337A>T (p.I2113F), c.3866+1G>T, c.3313G>T (p.E1105X), c.3334delG (p.G1112fs), c.8148G>T (p.Q2716H), c.6331A>T (p.N2111Y), c.3685C>T (p.Q1229X), c.3866+1G>A Sequencing | NM\_016239:2-65

Oculocutaneous Albinism: Type 1 (TYR): Mutations (27): O' Genotyping | c.272G>A (p.C91Y), c.242C>T (p.P81L), c.265T>C (p.C89R), c.1A>G (p.M1V), c.140G>A (p.G47D), c.325G>A (p.G109R), c.568delG (p.G191Dfs), c.707G>A (p.W236X), c.832C>T (p.R278X), c.1118C>A (p.T373K), c.229C>T (p.R77W), c.823G>T (p.V275F), c.32G>A (p.W11X), c.149C>T (p.S50L), c.1467\_1468insT (p.A490Cfs), c.820-2A>G, c.892C>T (p.R298W), c.1064C>T (p.A355V), c.1090A>C (p.N364H), c.1150C>G (p.P384A), c.1184+1G>A, c.1309G>A (p.D437N), c.1469C>A (p.A490D), c.133\_134insC (p.P45fs), c.710delA (p.D237fs), c.978delA (p.Q326fs), c.1138\_1158delTCTGCCAACGATCCTATCTTC (p.S380\_F386del) Sequencing | NM\_000372:1-5

Oculocutaneous Albinism: Type 3 (TYRP1): Mutations (6): & Genotyping | c.1067G>A (p.R356Q), c.497C>G (p.S166X), c.107delT, c.1057\_1060delAACA (p.N353fs), c.1103delA (p.K368fs), c.1120C>T (p.R374X) Sequencing | NM\_000550:2-8

Oculocutaneous Albinism: Type 4 (SLC45A2): Mutations (2): & Genotyping | c.469G>A (p.D157N), c.563G>T (p.G188V) Sequencing | NM\_016180:1-7

Omenn Syndrome: DCLRE1C Related (DCLRE1C): Mutations (1): Of Genotyping | c.597C>A (p.Y199X) Sequencing | NM\_001033855:1-14

Omenn Syndrome: RAG2 Related (RAG2): Mutations (1): of Genotyping | c.685C>T (p.R229W) Sequencing | NM\_000536:1-2

Ornithine Translocase Deficiency (SLC25A15): Mutations (3): & Genotyping | c.562\_564delTTC (p.188delF), c.95C>G (p.T32R), c.535C>T (p.R179X) Sequencing |

Osteopetrosis: TCIRG1 Related (TCIRG1): Mutations (6): & Genotyping | c.1674-1G>A, c.1392C>A (p.C464X), c.117+4A>T, c.1213G>A (p.G405R), c.1331G>T (p.R444L), c.922delC (p.Q308fs) Sequencing | NM\_006019:1-20

POLG Related Disorders: Autosomal Recessive (POLG): Mutations (16): of Genotyping | c.695G>A (p.R232H), c.752C>T (p.T251I), c.1399G>A (p.A467T), c.1760C>T (p.P587L), c.2243G>C (p.W748S), c.2542G>A (p.G848S), c.3488T>G (p.M1163R), c.911T>G (p.L304R), c.8G>C (p.R3P), c.2617G>T (p.E873X), c.2794C>T (p.H932Y), c.3151G>C (p.G1051R), c.2591A>G (p.N864S), c.1491G>C (p.Q497H), c.679C>T (p.R227W), c.3218C>T (p.P1073L) Sequencing | NM\_001126131:2-23

Papillon-Lefevre Syndrome (CTSC): Mutations (11): of Genotyping | c.815G>A (p.R272H), c.96T>G (p.Y32X), c.380A>C (p.H127P), c.1287G>C (p.W429C), c.856C>T (p.Q286X), c.755A>T (p.Q252L), c.628C>T (p.R210X), c.857A>G (p.Q286R), c.890-1G>A, c.1047delA (p.G350Vfs), c.1056delT (p.Y352fs) Sequencing | NM\_001814:1-7

Pendred Syndrome (SLC26A4): Mutations (7): of Genotyping | c.1001+1G>A, c.1151A>G (p.E384G), c.1246A>C (p.T416P), c.2168A>G (p.H723R), c.707T>C (p.L236P), c.716T>A (p.V239D), c.919-2A>G Sequencing | NM\_000441:1-21

Persistent Mullerian Duct Syndrome: Type I (AMH): Mutations (6): & Genotyping | c.1144G>T (p.E382X), c.571C>T (p.R191X), c.1518C>G (p.H506Q), c.1574G>A (p.C525Y), c.17\_18delTC, c.283C>T (p.R95X) Sequencing | NM\_000479:1-4

Persistent Mullerian Duct Syndrome: Type II (AMHR2): Mutations (14): & Genotyping | c.232+1G>A, c.1330\_1356delCTGGGCAATACCCCTACCTCTGATGAG, c.596delA, c.1217G>A (p.R406Q), c.742G>A (p.E248K), c.1277A>G (p.D426G), c.846T>G (p.H282Q), c.1373T>C (p.V458A), c.1471G>C (p.D491H), c.1510C>T (p.R504C), c.118G>T (p.G40X), c.289C>T (p.R97X), c.160C>T (p.R54C), c.425G>T (p.G142V) Sequencing | NM\_020547:1-11

Phenylalanine Hydroxylase Deficiency (PAH): Mutations (61): of Genotyping | c.1066-11 G>A (IVS 10-11 G>A), c. 1315+1 G>A (IVS 12+1 G>A), c. 1241 A>G (p.Y414C), c. 1222C>T (p.R408W), c.754C>T (p.R252W), c.1223G>A (p.R408Q), c.473G>A (p.R158Q), c.782G>A (p.R261Q), c.814G>T (p.G272X), c.143T>C (p.L48S), c.194T>C (p.I65T), c.896T>G (p.F299C), c.842C>T (p.P281L), c.838G>A (p.E280K), c.117C>G (p.F39L), c.3G>A (p.M1I), c.1A>G (p.M1V), c.611A>G (p.Y204C), c.721C>T (p.R241C), c.727C>T (p.R243X), c.1139C>T (p.T380M), c.926C>T (p.A309V), c.898G>T (p.A300S), c.734T>C (p.V245A), c.818C>T (p.S273F), c.997C>T (p.L333F), c.199T>C (p.S67P), c.1042C>G (p.L348V), c.136G>A (p.G46S), c.728G>A (p.R243Q), c.745C>T (p.L249F), c.581T>C (p.L194P), c.722G>T (p.R241L), c.829T>G (p.Y277D), c.899C>T (p.A300V), c.926C>A (p.A309D), c.1045T>C (p.S349P), c.1157A>G (p.Y386C), c.1169A>G (p.E390G), c.331C>T (p.R111X), c.241\_256delACCCATTTGGATAAAC (p.T81fs), c.442-1G>A (IVS4-1G>A), c.463\_464insTGTGTACC (p.R155fs), c.569T>G (p.V190G), c.682G>T (p.E228X), c.755G>A (p.R252Q), c.770G>T (p.G257V), c.781C>T (p.R261X), c.800A>G (p.Q267R), c.842+5G>A (IVS7+5G>A), c.856G>A (p.E286K), c.904delT (p.F302fs), c.913-7A>G (IVS8-7A>G), c.935G>T (p.G312V), c.1068C>G (p.Y356X), c.1238G>C (p.R413P), c. 1301 C>A (p.A434D), c.842+2T>A (IVS7+2T>A), c.764T>C (p.L255S), c.722G>A (p.R241H), c.533A>G (p.E178G) Sequencing | NM\_000277:1-13

Polyglandular Autoimmune Syndrome: Type I (AIRE): Mutations (5): & Genotyping | c.769C>T (p.R257X), c.254A>G (p.Y85C), c.1163\_1164insA (p.M388IfsX36), c.967\_979delCTGTCCCCTCCGC (p.L323SfsX51), c.415C>T (p.R139X) Sequencing

Pontocerebellar Hypoplasia: EXOSC3 Related (EXOSC3): Mutations (4): 07 Genotyping c.395A>C (p.D132A), c.294\_303delTGTTTACTGG (p.V99Wfs), c.92G>C (p.G31A), c.238G>T (p.V80F) Sequencing | NM\_016042:1-4

Pontocerebellar Hypoplasia: RARS2 Related (RARS2): Mutations (3): of Genotyping | c.35A>G (p.Q12R), c.110+5A>G, c.1024A>G (p.M342V) Sequencing | NM\_020320:1-20

Pontocerebellar Hypoplasia: SEPSECS Related (SEPSECS): Mutations (1): & Genotyping c.1001A>G (p.Y334C) Sequencing | NM\_016955:1-11

Pontocerebellar Hypoplasia: TSEN54 Related (TSEN54): Mutations (3): & Genotyping | c.919G>T (p.A307S), c.736C>T (p.Q246X), c.1027C>T (p.Q343X) Sequencing

Pontocerebellar Hypoplasia: VPS53 Related (VPS53): Mutations (2): ♂ Genotyping | c.2084A>G (p.Q695R), c.1556+5G>A Sequencing | NM\_001128159:1-22

Pontocerebellar Hypoplasia: VRK1 Related (VRK1): Mutations (2): & Genotyping



c.1072C>T (p.R358X), c.397C>T (p.R133C) Sequencing | NM\_003384:2-13

Primary Carnitine Deficiency (SLC22A5): Mutations (12): & Genotyping | c.506G>A (p.R169Q), c.396G>A (p.W132X), c.1195C>T (p.R399W), c.1433C>T (p.P478L), c.43G>T (p.G 15W), c.1324\_1325delGCinsAT (p.A4421), c.632A>G (p.Y211C), c.1202\_1203insA (p.Y401fsX), c.844C>T (p.R282X), c.505C>T (p.R169W), c.1196G>A (p.R399Q), c.95A>G (p.N32S) Sequencing | NM\_003060:1-10

Primary Ciliary Dyskinesia: DNAI1 Related (DNAI1): Mutations (5): of Genotyping | c.282\_283insAATA (p.G95Nfs), c.1543G>A (p.G515S), c.48+2\_48+3insT, c.1658\_1669delCCAAGGTCTTCA (p.Thr553\_Phe556del), c.1490G>A (p.G497D) Sequencing |

Primary Ciliary Dyskinesia: DNAI2 Related (DNAI2): Mutations (4): of Genotyping | c.1494+1G>A, c.346-3T>G, c.787C>T (p.R263X), c.1304G>A (p.W435X) Sequencing | NM\_023036:2-13

Primary Congenital Glaucoma (CYP1B1): Mutations (9): ♂ Genotyping | c.1405C>T (p.R469W), c.1093G>T (p.G365W), c.155C>T (p.P52L), c.1064\_1076delGAGTGCAGGCAGA (p.R355Hfs), c.1410\_1422delCATTGGCGAAGAA (p.C470fs), c.862\_863insC, c.1199\_1200insTCATGCCACC, c.182G>A (p.G61E), c.535delG (p.A179fs) Sequencing | NM\_000104:2-3

Primary Hyperoxaluria: Type 1 (AGXT): Mutations (11): of Genotyping | c.508G>A (p.G170R), c.454T>A (p.F152I), c.731T>C (p.I244T), c.121G>A (p.G41R), c.198C>G (p.Y66X), c.245G>A (p.G82E), c.466G>A (p.G156R), c.613T>C (p.S205P), c.697C>T (p.R233C), c.698G>A (p.R233H), c.738G>A (p.W246X) Sequencing | NM\_000030:1-11

Primary Hyperoxaluria: Type 2 (GRHPR): Mutations (3): of Genotyping | c.103delG, c.404+3delAAGT, c.295C>T (p.R99X) Sequencing | NM\_012203:1-9

Primary Hyperoxaluria: Type 3 (HOGA1): Mutations (2): & Genotyping c.944\_946delAGG (p.315delE), c.860G>T (p.G287V) Sequencing | NM\_138413:1-7

Progressive Familial Intrahepatic Cholestasis: Type 2 (ABCB11): Mutations (5): o' Genotyping | c.3767\_3768insC, c.890A>G (p.E297G), c.1723C>T (p.R575X), c.3169C>T (p.R1057X), c.1295G>C (p.R432T) Sequencing | NM\_003742:2-28

Propionic Acidemia: PCCA Related (PCCA): Mutations (13): of Genotyping | c.862A>G (p.R288G), c.937C>T (p.R313X), c.1196G>A (p.R399Q), c.1685C>G (p.S562X), 916\_917insT, c.1192T>C (p.C398R), c.229C>T (p.R77W), c.590G>A (p.G197E), c.1643+1G>A (IVS18+1G>A), c.890A>G (p.Q297R), c.1644-6C>G (IVS18-6C>G), c.1746G>A (p.S582S), c.1268C>T (p.P423L) Sequencing | NM\_000282:1-24

Propionic Acidemia: PCCB Related (PCCB): Mutations (13): & Genotyping | c.280G>T (p.G94X), c.335G>A (p.G112D), c.457G>C (p.A153P), c.502G>A (p.E168K), c. 1218\_1231 delGGGCATCATCCGGCinsTAGAGCACAGGA (p.G407fs), c. 1228C>T (p.R410W), c.1283C>T (p.T4281), c.1304A>G (p.Y435C), c.1495C>T (p.R499X), c.1534C>T (p.R512C), c.1539\_1540insCCC (p.R514PfsX38), c.1556T>C (p.L519P), c.1606A>G (p.N536D) Sequencing NM\_000532:1-15

Pseudocholinesterase Deficiency (BCHE): Mutations (1): & Genotyping | c.293A>G (p.D98G) Sequencing | NM\_000055:2-4

Pycnodysostosis (CTSK): Mutations (2): of Genotyping | c.990A>G (p.X330W), c.926T>C (p.L309P) Sequencing | NM\_000396:2-8

Pyruvate Carboxylase Deficiency (PC): Mutations (15): of Genotyping | c.1892G>A (p.R631Q), c.184C>T (p.R62C), c.2540C>T (p.A847V), c.1351C>T (p.R451C), c.467G>A (p.R156Q), c.1828G>T (p.A610S), c.2229G>T (p.M743I), c.434T>C (p.V145A), c.1748G>T (p.R583L), c.2491\_2492delGT (p.V831fs), c.3409\_3410delCT (p.L1137fs), c.2493\_2494delGT (p.F832Xfs), c.2876\_2877insT (p.F959fs), c.2473+2\_2473+5delTAGG, c.1828G>A (p.A610T) Sequencing | NM\_022172:2-21

Pyruvate Dehydrogenase Deficiency (PDHB): Mutations (2): of Genotyping | c.395A>G (p.Y132C), c.1030C>T (p.P344S) Sequencing | NM\_000925:1-10

Renal Tubular Acidosis and Deafness (ATP6V1B1): Mutations (7): of Genotyping | c.242T>C (p.L81P), c.232G>A (p.G78R), c.1248+1G>C, c.585+1G>A, c.497delC (p.T166fs), c.1037C>G (p.P346R), c.1155\_1156insC (p.I386fs) Sequencing | NM\_001692:1-14

Retinal Dystrophies: RLBP1 Related (RLBP1): Mutations (3): & Genotyping | c.700C>T (p.R234W), c.141G>A (p.K47=), c.141+2T>C Sequencing | NM\_000326:3-9

Retinal Dystrophies: RPE65 Related (RPE65): Mutations (12): & Genotyping | c.1292A>G (p.Y431C), c.1102T>C (p.Y368H), c.11+5G>A, c.700C>T (p.R234X), c.1087C>A (p.P363T), c.1022T>C (p.L341S), c.271C>T (p.R91W), c.1355T>G (p.V452G), c.1543C>T (p.R515W), c.907A>T (p.K303X), c.1067delA (p.N356fs), c.95-2A>T (IVS2-2A>T) Sequencing |

Retinitis Pigmentosa: CERKL Related (CERKL): Mutations (5): O' Genotyping | c.420delT (p.1141Lfs), c.598A>T (p.K200X), c.780delT (p.P261Lfs), c.769C>T (p.R257X), c.238+1G>A (IVS1+1G>A) Sequencing | NM\_201548:1-13

Retinitis Pigmentosa: DHDDS Related (DHDDS): Mutations (1): & Genotyping | c.124A>G (p.K42E) Sequencing | NM\_024887:2-9

Retinitis Pigmentosa: FAM161A Related (FAM161A): Mutations (5): of Genotyping | c.685C>T (p.R229X), c.1309A>T, c.1355\_1356delCA (p.T452fs), c.1567C>T (p.R523X), c.1786C>T (p.R596X) Sequencing | NM\_001201543:1-7

Rhizomelic Chondrodysplasia Punctata: Type I (PEX7): Mutations (8): of Genotyping | c.903+1G>C, c.649G>A (p.G217R), c.875T>A (p.L292X), c.40A>C (p.T14P),

c.45\_52insGGGACGCC (p.H18RfsX35), c.120C>G (p.Y40X), c.345T>G (p.Y115X), c.653C>T (p.A218V) Sequencing | NM\_000288:1-10

Salla Disease (SLC17A5): Mutations (5): & Genotyping | c.802\_816delTCATCATTAAGAAAT (p.L336fsX13), c.406A>G (p.K136E), c.115C>T (p.R39C), c.548A>G (p.H183R), c.1001C>G (p.P334R) Sequencing | NM\_012434:1-11

Sandhoff Disease (HEXB): Mutations (14): of Genotyping | c.76delA, c.445+1G>A, c.850C>T (p.R284X), c.508C>T (p.R170X), c.796T>G (p.Y266D), c.845G>A (p.G282E), c.800\_816delCACCAAATGATGTCCGT (p.T267fs), c.1082+5G>A, c.1250C>T (p.P417L), c.1615C>T (p.R539C), c.1514G>A (p.R505Q), c.1303\_1304delAG (p.R435fs), c.1509-26G>A, c.1597C>T (p.R533C) Sequencing | NM\_000521:1-14

Sanfilippo Syndrome: Type A (SGSH): Mutations (11): & Genotyping | c.734G>A (p.R245H), c.220C>T (p.R74C), c.197C>G (p.S66W), c.449G>A (p.R150Q), c.1339G>A (p.E447K), c.1105G>A (p.E369K), c.1298G>A (p.R433Q), c.383C>T (p.P128L), c.617G>C (p.R206P), c.892T>C (p.S298P), c.1080delC (p.T360fs) Sequencing | NM\_000199:1-8

Sanfilippo Syndrome: Type B (NAGLU): Mutations (10): d' Genotyping | c.2021 G>A (p.R674H), c.889C>T (p.R297X), c.1928G>A (p.R643H), c.1927C>T (p.R643C), c.1562C>T (p.P521L), c.1444C>T (p.R482W), c.1693C>T (p.R565W), c.1694G>C (p.R565P), c.700C>T (p.R234C), c.1876C>T (p.R626X) Sequencing | NM\_000263:2-6

Sanfilippo Syndrome: Type C (HGSNAT): Mutations (13): & Genotyping | c.848C>T (p.P283L,p.P311L), c.962T>G (p.L321X), c.1529T>A (p.M510K), c.1030C>T (p.R344C), c.1553C>T (p.S518F), c.1150C>T (p.R384X), c.493+1G>A (IVS4+1G>A), c.372-2A>G (IVS3-2A>G), c.1622C>T (p.S541L), c.852-1G>A, c.525\_526insT (p.A175fsX), c.1345insG (p.D449fsX), c.234+1G>A (IVS2+1G>A) Sequencing | NM\_152419;2-18

Sanfilippo Syndrome: Type D (GNS): Mutations (5):  $\sigma^2$  Genotyping | c.1063C>T (p.R355X), c. 1168C>T (p.Q390X), c. 1226insG (p.R409fsX), c. 1138insGTCCT (p.D380fsX), c. 1169delA (p.Q390fsX) Sequencing | NM\_002076:1-14

Short-Chain Acyl-CoA Dehydrogenase Deficiency (ACADS): Mutations (5): & Genotyping | c.1058C>T (p.S353L), c.1138C>T (p.R380W), c.1147C>T (p.R383C), c.319C>T (p.R107C), c.575C>T (p.A192V) Sequencing | NM\_000017:1-10

Sickle-Cell Anemia (HBB): Mutations (1): ♂ Genotyping | c.20A>T (p.E7V) Sequencing | NM\_000518:1-3

Sjogren-Larsson Syndrome (ALDH3A2): Mutations (2): of Genotyping | c.943C>T (p.P315S), c.1297\_1298delGA (p.E433fs) Sequencing | NM\_001031806:1-10

Sly Syndrome (GUSB): Mutations (5): of Genotyping | c.526C>T (p.L176F), c.1244C>T (p.P415L), c.1222C>T (p.P408S), c.1856C>T (p.A629V), c.1429C>T (p.R477W) Sequencing NM\_000181:1-12

 $Smith-Lemli-Opitz\ Syndrome\ \{DHCR7\}:\ \textit{Mutations}\ (50):\ \textit{O}'\ Genotyping\ |\ c.964-1G>C,$ c.356A>T (p.H119L), c.1054C>T (p.R352W), c.1210C>T (p.R404C), c.278C>T (p.T93M), c.1055G>A (p.R352Q), c.1139G>A (p.C380Y), c.1337G>A (p.R446Q), c.452G>A (p.W151X), c.453G>A (p.W151X), c.744G>T (p.W248C), c.976G>T (p.V326L), c.326T>C (p.L109P), c.470T>C (p.L157P), c.1342G>A (p.E448K), c.1228G>A (p.G410S), c.906C>G (p.F302L), c.725G>A (p.R242H), c.724C>T (p.R242C), c.506C>T (p.S169L), c.1A>G, c.670G>A (p.E224K), c.818T>G (p.V273G), c.203T>C (p.L68P), c.292C>T (p.Q98X), c.532A>T (p.1178F), c.545G>T (p.W182L), c.682C>T (p.R228W), c.575C>T (p.S192F), c.1295A>G (p.Y432C), c.1039G>A (p.G347S), c.1079T>C (p.L360P), c.1424T>C (p.F475S), c.1190C>T (p.S397L), c.1351T>C (p.C451R), c.853\_855delTTC (p.285delF), c.1327C>T (p.R443C), c.151C>T (p.P51S), c.296T>C (p.L99P), c.443T>G (p.L148R), c.502T>A (p.F168I), c.523G>C (p.D175H), c.536C>T (p.P179L), c.728C>G (p.P243R), c.852C>A (p.F284L), c.861C>A (p.N287K), c.970T>C (p.Y324H), c. 1384T>C (p.Y462H), c. 1406G>C (p.R469P), c. 111G>A (p.W37X) Sequencing

Spinal Muscular Atrophy: SMN1 Linked (SMN1): Mutations (19): of Genotyping | DEL EXON 7, c.22\_23insA, c.43C>T (p.Q15X), c.91\_92insT, c.305G>A (p.W102X), c.400G>A (p.E134K), c.439\_443delGAAGT, c.558delA, c.585\_586insT, c.683T>A (p.L228X), c.734C>T (p.P245L), c.768\_778dupTGCTGATGCTT, c.815A>G (p.Y272C), c.821C>T (p.T274I), c.823G>A (p.G275S), c.834+2T>G, c.835-18\_835-12delCCTTTAT, c.835G>T, c.836G>T dPCR | DEL

Stargardt Disease (ABCA4): Mutations (16): & Genotyping | c.3083C>T (p.A1028V), c.52C>T (p.R18W), c.5338C>G (p.P1780A), c.1018T>G (p.Y340D), c.2461T>A (p.W821R), c.2565G>A (p.W855X), c.3106G>A (p.E1036K), c.3210\_3211insGT (p.S1071Vfs), c.634C>T (p.R212C), c.3113C>T (p.A1038V), c.1622T>C (p.L541P), c.3364G>A (p.E1122K), c.6079C>T (p.L2027F), c.2588G>C (p.G863A), c.1938-1G>A, c.571-2A>G Sequencing | NM\_000350:1-

Stuve-Wiedemann Syndrome (LIFR): Mutations (9): of Genotyping | c.2472\_2476delTATGT, c.2434C>T (p.R812X), c.2274\_2275insT, c.1789C>T (pR597X), c.1601-2A>G, c.1620\_1621insA, c.756\_757insT (p.K253X), c.653\_654insT, c.170delC Sequencing | NM\_002310:2-20

Sulfate Transporter-Related Osteochondrodysplasia (SLC26A2): Mutations (7): 03 Genotyping | c.1018\_1020delGTT (p.340delV), c.-26+2T>C, c.532C>T (p.R178X), c.835C>T (p.R279W), c.1957T>A (p.C653S), c.398C>T (p.A133V), c.764G>A (p.G255E) Sequencing | NM\_000112:1-3

Tay-Sachs Disease (HEXA): Mutations (78): ♂ Genotyping | c.1073+1G>A, c. 1277\_1278insTATC, c. 1421+1G>C, c.805+1G>A, c.532C>T (p.R178C), c.533G>A (p.R178H), c.805G>A (p.G269S), c.1510C>T (p.R504C), c.1496G>A (p.R499H), c.509G>A (p.R170Q), c.1003A>T (p.I335F), c.910\_912delTTC (p.305delF), c.749G>A (p.G250D), c.632T>C (p.F211S),



c.629C>T (p.S210F), c.613delC, c.611A>G (p.H204R), c.598G>A (p.V200M), c.590A>C (p.K197T), c.571-1G>T, c.540C>G (p.Y180X), c.538T>C (p.Y180H), c.533G>T (p.R178L), c.508C>T (p.R170W), c.409C>T (p.R137X), c.380T>G (p.L127R), c.346+1G>C, c.116T>G (p.L39R), c.78G>A (p.W26X), c.1A>G (p.M1V), c.1495C>T (p.R499C), c.459+5G>A (IVS4+5G>A), c.1422-2A>G, c.535C>T (p.H179Y), c.1141delG (p.V381fs), c.796T>G (p.W266G), c.155C>A (p.S52X), c.426delT (p.F142fs), c.413-2A>G, c.570+3A>G, c.536A>G (p.H179R), c.1146+1G>A, c.736G>A (p.A246T), c.1302C>G (p.F434L), c.778C>T (p.P260S), c.1008G>T (p.Q336H), c.1385A>T (p.E462V), c.964G>A (p.D322N), c.340G>A (p.E114K), c.1432G>A (p.G478R), c.1178G>C (p.R393P), c.805+1G>C, c.1426A>T (p.R476X), c.623A>T (p.D208V), c.1537C>T (p.Q513X), c.1511G>T (p.R504L), c.1307\_1308delTA (p.1436fs), c.571-8A>G, c.624\_627delTCCT (p.D208fs), c.1211\_1212delTG (p.L404fs), c.621T>G (p.D207E), c.1511G>A (p.R504H), c.1177C>T (p.R393X), c.2T>C (p.M1T), c.1292G>A (p.W431X), c.947\_948insA (p.Y316fs), c.607T>G (p.W203G), c.1061\_1063delTCT (p.F354\_Y355delinsX), c.615delG (p.L205fs), c.805+2T>C, c.1123delG (p.E375fs), c.1121A>G (p.Q374R), c.1043\_1046delTCAA (p.F348fs), c.1510delC (p.R504fs), c.1451T>C (p.L484P), c.964G>T (p.D322Y), c.1351C>G (p.L451V), c.571-2A>G (IVS5-2A>G) Sequencing | NM\_000520:1-14

Trichohepatoenteric Syndrome: Type 1 (TTC37): Mutations (9): o\* Genotyping | c.3847G>A (p.D1283N), c.751G>A (p.G251R), c.2251C>T (p.Q751X), c.439C>T (p.Q147X), c.2808G>A (p.W936X), c.2515+1G>C, c.4620+1G>C, c.1632+1delG, c.2578-7delTTTTT Sequencing | NM\_014639:4-43

Tyrosine Hydroxylase Deficiency (TH): Mutations (1): of Genotyping | c.698G>A (p.R233H) Sequencing | NM\_199292:1-14

Tyrosinemia: Type I (FAH): Mutations (10): d\* Genotyping | c.1062+5G>A, c.554-1G>T, c.607-6T>G, c.707-1G>C, c.782C>T (p.P261L), c.1069G>T (p.E357X), c.786G>A (p.W262X), c.698A>T (p.D233V), c.1009G>A (p.G337S), c.192G>T (p.Q64H) Sequencing | NM 000137:1-14

Tyrosinemia: Type II (TAT): Mutations (5): of Genotyping | c.169C>T (p.R57X), c.668C>G (p.S223X), c.1249C>T (p.R417X), c.1085G>T (p.G362V), c.236-5A>G Sequencing | NM\_000353:2-12

Usher Syndrome: Type 1B (MYO7A): Mutations {13}: of Genotyping | c.93C>A (p.C31X), c.448C>T (p.R150X), c.634C>T (p.R212C), c.635G>A (p.R212H), c.700C>T (p.Q234X), c.1797G>A (p.M599I), c.1996C>T (p.R666X), c.2476G>A (p.A826T), c.3719G>A (p.R1240Q), c.5581C>T (p.R1861X), c.6025delG (p.A2009fs), c.640G>A (p.G214R), c.1190C>A (p.A397D) Sequencing | NM\_000260:2-49

Usher Syndrome: Type 1C (USH1C): Mutations (6):  $\sigma$  Genotyping | c.496+1G>A, c.238\_239insC, c.216G>A (p.V72fs), c.91C>T (p.R31X), c.36+1G>T, c.496+1G>T Sequencing | NM\_153676:1-27

Usher Syndrome: Type 1D (CDH23): Mutations (14): O<sup>\*</sup> Genotyping | c.172C>T (p.Q58X), c.3367C>T (p.Q1123X), c.3617C>G (p.P1206R), c.3713\_3714delCT (p.S1238fs), c.3880C>T (p.Q1294X), c.4069C>T (p.Q1357X), c.4488G>C (p.Q1496H), c.4504C>T (p.R1502X), c.5237G>A (p.R1746Q), c.5985C>A (p.Y1995X), c.6307G>T (p.E2103X), c.7549A>G (p.S2517G), c.8230G>A (p.G2744S), c.8497C>G (p.R2833G) Sequencing | NM\_022124:2-68

Usher Syndrome: Type 1F (PCDH15): Mutations [7]: O' Genotyping | c.733C>T (p.R245X), c.2067C>A (p.Y684X), c.7C>T (p.R3X), c.1942C>T (p.R648X), c.1101delT (p.A367fsX), c.2800C>T (p.R934X), c.4272delA (p.L1425fs) Sequencing | NM\_001142763:2-35

Usher Syndrome: Type 2A (USH2A): Mutations (22): d' Genotyping | c.14020A>G (p.R4674G), c.12067-2A>G, c.4338\_4339delCT (p.C1447fs), c.2299delG (p.E767SfsX21), c.2209C>T (p.R737X), c.1256G>T (p.C419F), c.1000C>T (p.R334W), c.923\_924insGCCA (p.H308fs), c.12708T>A (p.C4236X), c.13576C>T (p.R4526X), c.1840+1G>A, c.11328T>G (p.Y3776X), c.5329C>T (p.R1777W), c.9165\_9168delCTAT (p.13055MfsX2), c.9469C>T (p.Q3157X), c.1876C>T (p.R626X), c.7123delG (p.G2375fs), c.9492\_9498delTGATGAG (p.D3165fs), c.6235A>T (p.K2079X), c.1403C>G (p.Y4801X), c.3788G>A (p.W1263X), c.11328T>A (p.Y3776X) Sequencing | NM 206933:2-72

 $\label{thm:continuous} \begin{tabular}{ll} Usher Syndrome: Type 3 (CLRN1): Mutations (5): $d$' Genotyping | c.144T>G (p.N48K), c.131T>A [p.M120K], c.567T>G (p.Y189X), c.634C>T (p.Q212X), c.221T>C (p.L74P) Sequencing | NM_001195794:1-4 | NM_000195794:1-4 | NM_001195794:1-4 | NM$ 

Very Long-Chain Acyl-CoA Dehydrogenase Deficiency (ACADVL): Mutations (29): o' Genotyping | c.779C>T (p.T260M), c.848T>C (p.V283A), c.1144A>C (p.K382Q), c.1226C>T (p.T409M), c.1322G>A (p.G441D), c.1372T>C (p.F458L), c.1405C>T (p.R469W), c.1837C>T (p.R613W), c.553G>A (p.G185S), c.739A>C (p.K247Q), c.37C>T (p.Q13X), c.265C>T (p.P89S), c.272C>A (p.P91Q), c.364A>G (p.N122D), c.388\_391delGAGA (p.E130fs), c.520G>A (p.V174M), c.856A>G (p.R286G), c.1606\_1609delGCAG (p.A536fs), c.1531C>T (p.R511W), c.1512G>T (p.E504D), c.644G>A (p.G222R), c.685C>T (p.R229X), c.577G>C (p.G193R), c.881G>A (p.G294E), c.753-2A>C (IVS8-2A>C), c.1349G>A (p.R450H), c.1358G>A (p.R453Q), c.790A>G (p.K264E), c.1246G>A (p.A416T) Sequencing | NM\_000018:1-20

Walker-Warburg Syndrome {FKTN}: Mutations (5): 0<sup>th</sup> Genotyping | c.1167insA (p.F390fs), c.139C>T (p.R47X), c.748T>G (p.C250G), c.648-1243G>T (IVS5-1243G>T), c.515A>G (p.H172R) Sequencing | NM\_006731:2-10

Werner Syndrome (WRN): Mutations [8]:  $\sigma^{7}$  Genotyping | c.3139-1G>C (IVS25-1G>C), c.3913C>T {p.R1305X}, c.3493C>T {p.Q1165X}, c.1730A>T {p.K577M}, c.1336C>T {p.R368X}, c.3686A>T {p.Q1229L}, c.3915\_3916insA {p.R1306fs}, c.2089-3024A>G Sequencing | NM\_000553:2-35

Wilson Disease (ATP7B): Mutations (17): o' Genotyping | c.1340\_1343delAAAC, c.2304delC (p.M769Cfs), c.2332C>G (p.R778G), c.3207C>A (p.H1069Q), c.2333G>T

(p.R778L), c.2336G>A (p.W779X), c.2337G>A (p.W779X), c.2906G>A (p.R969Q), c.1934T>G (p.M645R), c.2123T>C (p.L708P), c.-370\_-394delTGGCCGAGACCGCGG, c.3191A>C (p.E1064A), c.845delT (p.L282Pfs), c.3817C>T (p.P1273S), c.3683G>C (p.R1228T), c.3809A>G (p.N1270S), c.2293G>A (p.D765N) Sequencing | NM\_000053:1-21

Wolcott-Rallison Syndrome (EIF2AK3): Mutations (5): of Genotyping | c.1409C>G (p.S470X), c.1262delA (p.N421fs), c.1570delGAAA (p.E524fsX), c.478delG (p.A160fs), c.1047\_1060delAGTCATTCCCATCA (p.V350Sfs) Sequencing | NM\_004836:1-17

Wolman Disease (LIPA): Mutations (3):  $\sigma$  Genotyping | c.964C>T (p.Q322X), c.419G>A (p.W140X), c.260G>T (p.G87V) Sequencing | NM\_001127605:2-10

Xeroderma Pigmentosum: Group A (XPA): Mutations (7): of Genotyping | c.172+2T>G, c.323G>T (p.C108F), c.374delC (p.T125fs), c.682C>T (p.R228X), c.619C>T (p.R207X), c.348T>A (p.Y116X), c.390-1G>C Sequencing | NM\_000380:1-6

Xeroderma Pigmentosum: Group C (XPC): Mutations (5): o\* Genotyping | c.1735C>T (p.R579X), c.566\_567delAT (p.Y189fs), c.413-9T>A, c.413-24A>G, c.1643\_1644delTG (p.V548fs) Sequencing | NM\_004628:1-16

Zellweger Spectrum Disorders: PEX1 Related (PEX1): Mutations (3): of Genotyping | c.2528G>A (p.G843D), c.2916delA (p.G973fs), c.2097insT (p.I700fs) Sequencing | NM 000466:1-24

Zellweger Spectrum Disorders: PEX10 Related (PEX10): Mutations (2): of Genotyping | c.764\_765insA, c.874\_875delCT Sequencing | NM\_153818:2-6

Zellweger Spectrum Disorders: PEX2 Related (PEX2): Mutations (1): of Genotyping | c.355C>T (p.R119X) Sequencing | NM\_001172087:1-3

Zellweger Spectrum Disorders: PEX6 Related (PEX6): Mutations (8): 63 Genotyping | c.1130+1G>A (IVS3+1G>A), c.1688+1G>A (IVS7+1G>A), c.1962-1G>A (p.L655fsX3), c.1301delC (p.S434Ffs), c.1601T>C (p.L534P), c.511insT (p.G171Wfs), c.802\_815delGACGGACTGGCGCT (p.D268Cfs), c.1715C>T (p.T572I) Sequencing | NM\_000287:1-17



#### Residual Risk Information

Detection rates are calculated from the primary literature and may not be available for all ethnic populations. The values listed below are for genotyping. Sequencing provides higher detection rates and lower residual risks for each disease. More precise values for sequencing may become available in the future.

Disease	Carrier Rate	Detection Rate	Residua Risk
11-Beta-Hydroxylase-Deficient Congenital Adrenal Hyperplasia	♂ Moroccan Jewish: 1/39	91.67%	1/468
17-Alpha-Hydroxylase Deficiency	♂ Brazilian: Unknown	54.55%	Unknow
	♂ Japanese: Unknown	45.45%	Unknown
17-Beta-Hydroxysteroid Dehydrogenase Deficiency	o⁴ Arab: 1/8	>99%	<1/800
	of Dutch: 1/192	13.89%	1/223
21-Hydroxylase-Deficient Classical Congenital Adrenal Hyperplasia	♂ European: 1/62	27.65%	1/86
	of General: 1/62	29.34%	1/88
21-Hydroxylase-Deficient Nonclassical Congenital Adrenal Hyperplasia	♂ Argentinian: 1/4	<10%	1/4
	o⁴ European: 1/16	<10%	1/16
3-Beta-Hydroxysteroid Dehydrogenase Deficiency	♂ General: Unknown	16.13%	Unknow
3-Methylcrotonyl-CoA Carboxylase Deficiency: MCCA Related	♂ European: 1/146	26.32%	1/198
	♂ General: 1/112	37.50%	1/179
3-Methylcrotonyl-CoA Carboxylase Deficiency: MCCB Related	♂ General: 1/112	35.29%	1/173
	o⁴ Japanese: 1/112	33.33%	1/168
	o⁴ Korean: 1/141	66.67%	1/423
	o⁴ Turkish: 1/112	24.07%	1/148
3-Methylglutaconic Aciduria: Type 3	o⁴ Iraqi Jewish: 1/10	>99%	<1/1,00
3-Phosphoglycerate Dehydrogenase Deficiency	♂ Ashkenazi Jewish: 1/400	>99%	<1/40,0 0
5-Alpha Reductase Deficiency	o⁴ Dominican: Unknown	>99%	Unknow
	♂ Mexican: Unknown	68.75%	Unknow
6-Pyruvayl-Tetrahydropterin Synthase Deficiency	od Chinese: 1/183	78.95%	1/869
	♂ East Asian: 1/180	64.20%	1/503
ARSACS	o⁴ French Canadian: 1/22	95.45%	1/484
Abetalipoproteinemia	♂ Ashkenazi Jewish: 1/131	>99%	<1/13,1 0
Acrodermatitis Enteropathica	o³ Arab: Unknown	40.00%	Unknow
	o³ Egyptian: Unknown	33.33%	Unknow
	♂ French: Unknown	27.78%	Unknow
	♂ Tunisian: Unknown	77.78%	Unknow
Acute Infantile Liver Failure: TRMU Related	o <sup>™</sup> Yemenite Jewish: 1/40	71.43%	1/140
Acyl-CoA Oxidase I Deficiency	♂ General: Unknown	35.00%	Unknow
	o⁴ Japanese: Unknown	42.86%	Unknow
Adenosine Deaminase Deficiency	o' General: 1/388	36.96%	1/615

Disease	Carrier Rate	Detection Rate	Residual Risk
Alkaptonuria	o* Dominican: Unknown	>99%	Unknown
The second secon	o' Finnish: 1/251	60.00%	1/628
	o' Slovak: 1/69	59.38%	1/170
Alpha Thalassemia	o' General: 1/48	50.67%	1/97
Alpha-1-Antitrypsin Deficiency	o' European: 1/35	95.00%	1/700
this Research at photosphe & Photospherical photosp	o' General: Unknown	95.00%	Unknown
Alpha-Mannosidosis	o European: 1/354	30.23%	1/507
	♂ General: 1/354	35.19%	1/546
Alport Syndrome: COL4A3 Related	o Dutch: 1/409	22.73%	1/529
Alport Syndrome: COL4A4 Related	o⁴ General: 1/409	23.33%	1/533
Amegakaryocytic Thrombocytopenia	♂ Ashkenazi Jewish: 1/76	>99%	<1/7,600
	o' General: Unknown	64.81%	Unknown
Andermann Syndrome	o⁴ French Canadian: 1/24	99.38%	1/3,888
Antley-Bixler Syndrome	o⁴ General: Unknown	45.65%	Unknown
	o⁴ Japanese: Unknown	60.47%	Unknown
Argininemia	o' Chinese: Unknown	40.00%	Unknown
	o³ French Canadian: Unknown	75.00%	Unknown
	oʻ Japanese: Unknown	>99%	Unknown
Argininosuccinate Lyase Deficiency	o⁴ European: 1/133	57.41%	1/312
	o⁴ Saudi Arabian: 1/80	51.72%	1/166
Aromatase Deficiency	o' General: Unknown	25.00%	Unknown
Arthrogryposis, Mental Retardation, & Seizures	♂ Ashkenazi Jewish: 1/205	>99%	<1/20,50 0
Asparagine Synthetase Deficiency	o" Iranian Jewish: 1/80	>99%	<1/8,000
Aspartylglycosaminuria	o' Finnish: 1/69	96.12%	1/1,780
Ataxia with Vitamin E Deficiency	o' European: 1/274	80.00%	1/1,370
	o' Italian: 1/224	97.73%	1/9,856
	♂ North African: 1/159	>99%	<1/15,90 0
Ataxia-Telangiectasia	o⁴ Costa Rican: 1/100	68.52%	1/318
	o <sup>™</sup> North African Jewish: 1/81	96.97%	1/2,673
	o⁴ Norwegian: 1/197	50.00%	1/394
	♂ Sardinians: Unknown	85.71%	Unknown
	o⁴ US Amish: Unknown	>99%	Unknown
Autosomal Recessive Polycystic Kidney Disease	♂ Finnish: 1/45	84.21%	1/285
	of French: 1/71	62.50%	1/189
	of General: 1/71	37.11%	1/113
Bardet-Biedl Syndrome: BBS1 Related	o⁴ General: 1/376	70.27%	1/1,265
	o <sup>™</sup> Northern European: 1/376	85.90%	1/2,666
	o <sup>®</sup> Puerto Rican: Unknown	90.00%	Unknown
Bardet-Biedl Syndrome: BBS 10 Related	o' General: 1/404	47.79%	1/774
Bardet-Biedl Syndrome: BBS 11 Related	of Bedouin: 1/59	>99%	<1/5,900
Bardet-Biedl Syndrome: BBS 12 Related	od General: Unknown	50.00%	Unknown



Disease	Carrier Rate	Detection Rate	Residual Risk	Disease	Carrier Rate	Detection Rate	Residual Risk
Bardet-Biedl Syndrome: BBS2 Related	oʻ Ashkenazi Jewish: Unknown	>99%	Unknown		o <sup>a</sup> Moroccan Jewish: 1/234	>99%	<1/23,40 0
	♂ General: 1/638	38.46%	1/1,037	Citrin Deficiency	♂ Japanese: 1/70	>99%	<1/7,000
	♂ Middle Eastern: Unknown	>99%	Unknown	Citrullinemia: Type I	♂ European: 1/120	18.18%	1/147
Bare Lymphocyte Syndrome: Type II	o⁴ General: Unknown	66.67%	Unknown		♂ General: 1/120	52.27%	1/251
Bartter Syndrome: Type 4A	♂ General: 1/457	81.82%	1/2,514		♂ Japanese: Unknown	64.71%	Unknown
Beta Thalassemia	♂ African American: 1/75	84.21%	1/475		o⁴ Mediterranean: 1/120	50.00%	1/240
	♂ Indian: 1/24	74.12%	1/93	Classical Galactosemia	♂ African American: 1/78	73.13%	1/290
	o' Sardinians: 1/23	97.14%	1/804	8	o⁴ Ashkenazi Jewish: 1/127	>99%	<1/12,70
	o⁴ Spaniard: 1/51	93.10%	1/739		30.1.10	75 470/	0
Beta-Hexosaminidase Pseudodeficiency	o³ Ashkenazi Jewish: Unknown	>99%	Unknown		o <sup>a</sup> Dutch: 1/91 o <sup>a</sup> European: 1/112	75.47% 88.33%	1/371 1/960
	♂ General: Unknown	>99%	Unknown		o⁴ General: 1/125	80.00%	1/625
Beta-Ketothiolase Deficiency	oʻ Japanese: Unknown	58.33%	Unknown		o <sup>7</sup> Irish: 1/76	91.30%	1/874
	♂ Spaniard: Unknown	90.00%	Unknown		o' Irish Travellers: 1/14	>99%	<1/1,400
Biotinidase Deficiency	o' General: 1/123	78.32%	1/567	Cockayne Syndrome: Type A	o³ Christian Arab: Unknown	50.00%	Unknown
Bloom Syndrome	♂ Ashkenazi Jewish: 1/134	96.67%	1/4,020	Cockayne Syndrome: Type B	o' General: 1/378	19.30%	1/468
	oʻ European: Unknown	66.22%	Unknown	Cohen Syndrome	♂ European: Unknown	19.05%	Unknown
	oʻ Japanese: Unknown	50.00%	Unknown		o⁴ Finnish: 1/140	67.24%	1/427
Canavan Disease	♂ Ashkenazi Jewish: 1/55	98.86%	1/4,840		o⁴ US Amish: 1/12	>99%	<1/1,200
	oʻ European: Unknown	53,23%	Unknown	Combined Pituitary Hormone	o' European: 1/45	93.29%	1/671
Carnitine Palmitoyltransferase IA Deficiency	♂ General: Unknown	38.89%	Unknown	Deficiency: PROP1 Related	o' General: 1/45	82.35%	1/255
	o Hutterite: 1/16	>99%	<1/1,600	Congenital Disorder of Glycosylation:	o' Danish: 1/71	90.00%	1/710
	♂ Japanese: 1/101	66.67%	1/303	Type 1 A: PMM2 Related		Name of Section 2017	10074.000
Carnitine Palmitoyltransferase II	o' Ashkenazi Jewish:	>99%	Unknown		o' Dutch: 1/68	39.29%	1/112
Deficiency	Unknown	71 1001	OF LOSS		of European: 1/71	55.33%	1/159
Carnitine-Acylcarnitine Translocase	o' General: Unknown o' Asian: Unknown	71.43% 95.45%	Unknown Unknown	Congenital Disorder of Glycosylation: Type 1B: MPI Related	♂ French: Unknown	54.17%	Unknown
Deficiency	o' General: Unknown	18,75%	Unknown	Congenital Disorder of Glycosylation: Type 1C: ALGó Related	o⁴ French: Unknown	59.09%	Unknown
Carpenter Syndrome	o' Brazilian: Unknown	40.00%	Unknown		o' General: Unknown	86.21%	Unknown
carpenier syndrome	of Northern European:	85.00%	Unknown	Congenital Ichthyosis: ABCA 12 Related	o <sup>*</sup> North African; Unknown	>99%	Unknown
	Unknown	83.00%	Ulkilowii		o" South Asian: Unknown	66.67%	Unknown
Cartilage-Hair Hypoplasia	of Finnish: 1/76	93.33%	1/1,140	Congenital Insensitivity to Pain with Anhidrosis	♂ Japanese: Unknown	56.52%	Unknown
Cerebrotendinous Xanthomatosis	of US Amish: 1/19 of Dutch: Unknown	>99% 78.57%	<1/1,900 Unknown		o⁴ Moroccan Jewish: Unknown	>99%	Unknown
	♂ Italian: Unknown	45.95%	Unknown	Congenital Lipoid Adrenal Hyperplasia	o' Japanese: 1/201	51.11%	1/411
	♂ Japanese: Unknown	92.86%	Unknown		o' Korean: 1/251	63.64%	1/690
	o' Moroccan Jewish: 1/6	87.50%	1/48	Congenital Myasthenic Syndrome:	o' European Gypsy: 1/26	>99%	<1/2,600
Chediak-Higashi Syndrome	o' General: Unknown	19.64%	Unknown	CHRNE Related			
Cholesteryl Ester Storage Disease	o' General: 1/101	68.97%	1/325		of North African: Unknown	60.87%	Unknown
Choreoacanthocytosis	♂ Ashkenazi Jewish: Unknown	66.67%	Unknown	Congenital Myasthenic Syndrome: DOK7 Related	o⁴ European: 1/472	19.05%	1/583
Chronic Granulomatous Disease:	♂ Iranian: Unknown	71.43%	Unknown		o' General: 1/472	18.75%	1/581
CYBA Related	O' Japanese: 1/274	>99%	<1/27,40	Congenital Myasthenic Syndrome: RAPSN Related	o General: 1/437	88.57%	1/3,824
	o⁴ Korean: 1/105	>99%	0 <1/10,50 0		oʻ Non-Ashkenazi Jewish: Unknown	>99%	Unknown



Disease	Carrier Rate	Detection Rate	Residual Risk	Disease	Carrier Rate	Detection Rate	Residua Risk
Congenital Neutropenia: Recessive	♂ English: Unknown	11.76%	Unknown		♂ Saudi Arabian: 1/38	>99%	<1/3,80
	o⁴ Japanese: Unknown	22.22%	Unknown	Familial Dysautonomia	♂ Ashkenazi Jewish: 1/31	>99%	<1/3,10
	o⁴ Turkish: Unknown	89.47%	Unknown	Familial Hyperinsulinism: Type 1:	♂ Ashkenazi Jewish: 1/52	98.75%	1/4,160
Corneal Dystrophy and Perceptive Deafness	o <sup>*</sup> General: Unknown	71.43%	Unknown	ABCC8 Related	♂ Finnish: 1/101	45.16%	1/184
Corticosterone Methyloxidase Deficiency	♂ Iranian Jewish: 1/32	>99%	<1/3,200	Familial Hyperinsulinism: Type 2: KCNJ 11 Related	o⁴ Arab: Unknown	40.00%	Unknown
Crigler-Najjar Syndrome	o⁴ Sardinians; Unknown	80.00%	Unknown	Familial Mediterranean Fever	♂ Arab: 1/4	51.27%	1/8
	of Tunisian: Unknown	>99%	Unknown		of Armenian: 1/5	94.51%	1/91
Cystic Fibrosis	♂ African American: 1/62	69.99%	1/207		♂ Ashkenazi Jewish: 1/81	40.95%	1/137
	oʻ Ashkenazi Jewish: 1/23	96.81%	1/721		♂ Iraqi Jewish: 1/4	76.92%	1/17
	o' Asian: 1/94	65.42%	1/272		♂ Israeli Jewish: 1/5	62.67%	1/13
	o⁴ European: 1/25	94.96%	1/496		♂ Lebanese: 1/6	91.67%	1/72
	♂ Hispanic American: 1/48	77.32%	1/212		o' North African Jewish: 1/5	95.69%	1/116
	o' Native American: 1/53	84.34%	1/338		♂ Syrian: 1/6	85.14%	1/40
Cystinosis	o' Dutch: 1/194	73.08%	1/721		of Turkish: 1/5	74.43%	1/20
	of French Canadian: 1/40	75.00%	1/160	Fanconi Anemia: Type A	on Moroccan Jewish: 1/100	>99%	<1/10,00
	o' General: 1/194	54.51%	1/426		o⁴ Spanish Gypsy: 1/67	>99%	<1/6,700
Cystinuria: Non-Type I	o' European: 1/42	61.11%	1/108	Fanconi Anemia: Type C	o Ashkenazi Jewish: 1/101	>99%	<1/10,10
	o' General: 1/42	37.50%	1/67	Tulicom Alleima. Type C	o / talkenezi seriali. 17 101	,,,,	0
	o⁴ Libyan Jewish: 1/26	93.48%	1/399		♂ General: Unknown	30.00%	Unknown
	o⁴ United States: 1/42	56.25%	1/96	Fanconi Anemia: Type G	o⁴ Black South African:	81.82%	1/556
Cystinuria: Type I	o' European: 1/42	46.67%	1/79		1/101		
	o' Swedish: 1/159	55.88%	1/360		♂ French Canadian: Unknown	87.50%	Unknown
D-Bifunctional Protein Deficiency	o' General: 1/159	38.64%	1/259		o⁴ Japanese: Unknown	75.00%	Unknown
Diabetes: Recessive Permanent Neonatal	♂ General: Unknown	25.00%	Unknown		o⁴ Korean: Unknown	66.67%	Unknown
Du Pan Syndrome	o⁴ Pakistani: Unknown	>99%	Unknown	Fanconi Anemia: Type J	of General: Unknown	86.36%	Unknown
Dyskeratosis Congenita: RTEL1 Related	o' Ashkenazi Jewish: 1/203	>99%	<1/20,30	Fumarase Deficiency	o' General: Unknown	30.00%	Unknown
-	♂ General: 1/501	50.00%	0	GM1-Gangliosidoses	♂ Eurodescent Brazilian: 1/66	62.15%	1/174
Dystrophic Epidermolysis Bullosa:	o' Italian: Unknown	45.00%	Unknown		o' European: 1/194	50.00%	1/388
Recessive					♂ General: 1/194	20.00%	1/243
	o Mexican American: 1/345	56.25%	1/789		o⁴ Hispanic American: 1/194	58.33%	1/466
Ehlers-Danlos Syndrome: Type VIIC	o⁴ Ashkenazi Jewish:	>99%	Unknown		o <sup>a</sup> Japanese: Unknown	62.82%	Unknown
	Unknown			GRACILE Syndrome	o* Finnish: 1/109	97.22%	1/3,924
Ellis-van Creveld Syndrome: EVC Related	o' General: 1/123	32.14%	1/181	Galactokinase Deficiency	o Japanese: 1/501	50.00%	1/1,002
Ellis-van Creveld Syndrome: EVC2 Related	♂ General: Unknown	<10%	Unknown	•	o⁴ Roma: 1/51	>99%	<1/5,100
Enhanced S-Cone	♂ Ashkenazi Jewish:	90.48%	Unknown	Gaucher Disease	♂ Ashkenazi Jewish: 1/15	87.16%	1/117
Elilidiced 3-Colle	Unknown	70.4070	Olikilowii		of General: 1/112	31.60%	1/164
	o" General: Unknown	52.50%	Unknown		♂ Spaniard: Unknown	44.29%	Unknown
Ethylmalonic Aciduria	of Arab/Mediterranean:	29.17%	Unknown		♂ Turkish: 1/236	59.38%	1/581
	Unknown			Gitelman Syndrome	♂ European: 1/100	35.00%	1/154
r	of General: Unknown	38.24%	Unknown		o³ European Gypsy: Unknown	>99%	Unknown
Familial Chloride Diarrhea	of Finnish: 1/51	>99%	<1/5,100		♂ General: 1/101	30.00%	1/144
	o' Kuwaiti: 1/38	90.00%	1/380				



## $CarrierMap^{TM}$

Disease	Carrier Rate	Detection Rate	Residual Risk	Disease	Carrier Rate	Detection Rate	Residual Risk
Globoid Cell Leukodystrophy	o⁴ Dutch: 1/137	60.98%	1/351	Hemochromatosis: Type 2A: HFE2	o³ European: Unknown	69.23%	Unknown
	o⁴ European: 1/150	26.47%	1/204	Related	# 1	70 700/	
53	♂ Japanese: 1/150	36.00%	1/234		of Mediterranean: Unknown	72.73%	Unknown
Glutaric Acidemia: Type I	o³ European: 1/164	57.78%	1/388	Hemochromatosis: Type 3: TFR2 Related	o³ Italian: Unknown	73.21%	Unknown
	♂ General: 1/164	25.51%	1/220	Hemoglobinopathy: Hb C	♂ African American: 1/51	>99%	<1/5,100
	o⁴ US Amish: 1/12	>99%	<1/1,200	Hemoglobinopathy: Hb D	♂ Canadian: 1/64	>99%	<1/6,400
Glutaric Acidemia: Type IIA	of General: Unknown	71.43%	Unknown		♂ Indian: 1/16	>99%	<1/1,600
Glutaric Acidemia: Type IIB	o <sup>™</sup> General: Unknown	33.33%	Unknown		♂ Iranian: 1/11	>99%	<1/1,100
Glutaric Acidemia: Type IIC	♂ Taiwanese: Unknown	>99%	Unknown	Hemoglobinopathy: Hb E	o⁴ Cambodia: 1/4	>99%	<1/400
	o⁴ Turkish: Unknown	80.00%	Unknown		♂ Chinese: 1/13	>99%	<1/1,300
Glycine Encephalopathy: AMT Related	♂ General: Unknown	40.91%	Unknown		o' Indian: 1/10	>99%	<1/1,000
Glycine Encephalopathy: GLDC Related	o⁴ Finnish: 1/118	78.00%	1/536		♂ Thai: 1/9	>99%	<1/900
Keldled	o' General: 1/280	12.50%	1/320	Hemoglobinopathy: Hb O	♂ African American: 1/87	>99%	<1/8,700
Glycogen Storage Disease: Type IA	o <sup>™</sup> Ashkenazi Jewish: 1/71	>99%	<1/7,100		o' Middle Eastern: Unknown	>99%	Unknown
Olycogen Slorage Disease: Type IA	of Chinese: 1/159	80.00%	1/795	Hereditary Fructose Intolerance	o⁴ European: 1/81	72.73%	1/297
	of European: 1/177	76.88%	1/765		♂ Italian: 1/81	90.91%	1/891
	o' Hispanic American:	27.78%	1/245		of Slavic: 1/81	>99%	<1/8,100
	1/177	27.7078	1/ 243	Hereditary Spastic Paraplegia: TECPR2	♂ Bukharan Jewish: 1/75	>99%	<1/7,500
	♂ Japanese: 1/177	89.22%	1/1,641	Related			
	♂ Australian: 1/354	50.00%	1/708	Herlitz Junctional Epidermolysis Bullosa: LAMA3 Related	o⁴ Pakistani: Unknown	>99%	Unknown
	♂ European: 1/354	45.74%	1/652	Herlitz Junctional Epidermolysis	o⁴ European: Unknown	70.00%	Unknown
	o Japanese: 1/354	39.13%	1/582	Bullosa: LAMB3 Related			
Glycogen Storage Disease: Type II	♂ African American: 1/60	45.83%	1/111		d' General: 1/781	52.27%	1/1,636
	o <sup>™</sup> Chinese: 1/112	72.00%	1/400	Herlitz Junctional Epidermolysis Bullosa: LAMC2 Related	♂ Italian: Unknown	28.57%	Unknown
	o⁴ European: 1/97	51.76%	1/201	Hermansky-Pudlak Syndrome: Type 1	o Puerto Rican: 1/22	94.95%	1/436
	o⁴ North African: Unknown	60.00%	Unknown	Hermansky-Pudlak Syndrome: Type 3	o' Ashkenazi Jewish: 1/235	>99%	<1/23,50
Glycogen Storage Disease: Type III	o' Faroese: 1/30	>99%	<1/3,000	Trefinansky-rodiak cyndronic. Type o	O ASIRONOZIJEWISH. 17 200	- 7770	0
	♂ General: 1/159	39.81%	1/264		♂ European: 1/434	12.50%	1/496
	o' North African Jewish:	>99%	<1/3,500	Hermansky-Pudlak Syndrome: Type 4	o' European: Unknown	54.17%	Unknown
Glycogen Storage Disease: Type IV	1/35 O <sup>*</sup> Ashkenazi Jewish: 1/35	>99%	<1/3,500	Holocarboxylase Synthetase Deficiency	o' European: 1/148	83.33%	1/888
	o General: 1/461	18.60%	1/566		♂ Japanese: 1/159	76.92%	1/689
Glycogen Storage Disease: Type V	o⁵ Caucasus Jewish: Unknown	>99%	Unknown	Homocystinuria Caused by CBS Deficiency	o⁴ European: 1/224	64.29%	1/627
	♂ European: 1/159	60.71%	1/405		of Irish: 1/128	70.59%	1/435
	♂ General: Unknown	74.10%	Unknown		o⁴ Italian: 1/224	35.71%	1/348
	♂ Spaniard: 1/159	67.11%	1/483		o⁴ Norwegian: 1/41	84.38%	1/262
	♂ Yemenite Jewish: Unknown	75.00%	Unknown	11 ×	♂ Qatari: 1/22	>99%	<1/2,200
Glycogen Storage Disease: Type VII	♂ Ashkenazi Jewish: 1/250	>99%	<1/25,00		♂ Saudi Arabian: Unknown	92.31%	Unknown
Guanidinoacetate Methyltransferase	o' General: Unknown	29.41%	Unknown	Hurler Syndrome	o⁴ Czech: 1/190	52.50%	1/400
Deficiency	O Celleral, Clikilowii	27.4170	Olkhowii		♂ European: 1/194	81.71%	1/1,061
HMG-CoA Lyase Deficiency	♂ General: 1/159	40.00%	1/265		o⁴ General: 1/194	62.50%	1/517
	♂ Japanese: Unknown	30.00%	Unknown		o' Italian: 1/194	61.11%	1/499
	o³ Portuguese: Unknown	86.36%	Unknown		o⁴ Japanese: 1/194	23.68%	1/254
	o³ Saudi Arabian: Unknown	93.33%	Unknown		♂ Moroccan Jewish: 1/194	92.31%	1/2,522
					o⁴ Scandinavian: 1/194	79.41%	1/942



## $Carrier Map^{tM}$

Disease	Carrier Rate	Detection Rate	Residual Risk	Disease	Carrier Rate	Detection Rate	Residual Risk
	♂ Spaniard: 1/194	52.50%	1/408	Limb-Girdle Muscular Dystrophy: Type	♂ Brazilian: Unknown	57.14%	Unknown
Hypophosphatasia	♂ Canadian Amish: 1/26	>99%	<1/2,600	2E			2.0
	♂ European: 1/159	19.23%	1/197		o⁴ European: 1/539	25.00%	1/719
	o⁴ Japanese: Unknown	54.55%	Unknown		♂ General: Unknown	12.50%	Unknown
Inclusion Body Myopathy: Type 2	♂ General: Unknown	85.83%	Unknown		of US Amish: Unknown	>99%	Unknown
	♂ Iranian Jewish: 1/16	>99%	<1/1,600	Limb-Girdle Muscular Dystrophy: Type 2F	o⁴ Brazilian: Unknown	>99%	Unknown
	♂ Japanese: Unknown	71.88%	Unknown		o⁴ General: Unknown	83.33%	Unknown
	♂ Korean: Unknown	72.50%	Unknown	Limb-Girdle Muscular Dystrophy: Type	♂ Brazilian: Unknown	34.62%	Unknown
nfantile Cerebral and Cerebellar Atrophy	♂ Caucasus Jewish: 1/20	>99%	<1/2,000	21	o <sup>a</sup> Danish: 1/100	85.53%	1/691
solated Microphthalmia: VSX2 Related	♂ Middle Eastern: Unknown	71.43%	Unknown		♂ General: Unknown	43.18%	Unknown
sovaleric Acidemia	o⁴ General: 1/251	47.37%	1/477		♂ German: 1/300	82.50%	1/1,714
loubert Syndrome	o⁴ Ashkenazi Jewish: 1/92	>99%	<1/9,200	Lipoprotein Lipase Deficiency	♂ French Canadian: 1/44	28.95%	1/62
amellar Ichthyosis: Type 1	o⁴ Norwegian: 1/151	81.40%	1/812	St. I and I decided a Co. I allowed a special	of General: Unknown	20.00%	Unknown
aryngoonychocutaneous Syndrome	oʻ Pakistani: Unknown	>99%	Unknown	Long-Chain 3-Hydroxyacyl-CoA	♂ European: 1/126	88.98%	1/1,144
eber Congenital Amaurosis: CEP290	o⁴ European: 1/251	47.32%	1/476	Dehydrogenase Deficiency			
Related		772-2727			♂ General: 1/126	56.25%	1/288
eber Congenital Amaurosis: GUCY2D Related		>99%	Unknown	Lysinuric Protein Intolerance	o⁴ Finnish: 1/123	>99%	<1/12,30 0
eber Congenital Amaurosis: LCA5 telated	o⁴ Pakistani: Unknown	83.33%	Unknown		o⁴ Italian: 1/120	45.45%	1/220
eber Congenital Amaurosis: RDH 12	o' General: 1/560	38.37%	1/909		♂ Japanese: 1/115	37.93%	1/185
elated	,				o' North African: Unknown	>99%	Unknown
eigh Syndrome: French-Canadian	o⁴ French Canadian: 1/23	95.45%	1/506	MTHFR Deficiency: Severe	♂ Bukharan Jewish: 1/39	>99%	<1/3,900
eukoencephalopathy with Vanishing Vhite Matter: EIF2B5 Related	o³ Cree: Unknown	>99%	Unknown	Malonyl-CoA Decarboxylase Deficiency	o⁴ General: Unknown	33.33%	Unknown
	o⁴ European: Unknown	65.22%	Unknown	Maple Syrup Urine Disease: Type 1A	♂ US Amish: 1/10	97.73%	1/440
eydig Cell Hypoplasia (Luteinizing	♂ Brazilian: Unknown	>99%	Unknown	Maple Syrup Urine Disease: Type 1B	♂ Ashkenazi Jewish: 1/97	>99%	<1/9,700
dormone Resistance)				Maple Syrup Urine Disease: Type 2	♂ General: 1/481	42.31%	1/834
imb-Girdle Muscular Dystrophy: Type !A	of Basque: 1/61	61.46%	1/158		♂ Norwegian: 1/481	50.00%	1/962
	of Croatian: 1/133	76.00%	1/554		♂ Turkish: 1/112	58.33%	1/269
	o* European: 1/103	17.23%	1/124	Maple Syrup Urine Disease: Type 3	♂ Ashkenazi Jewish: 1/94	>99%	<1/9,400
	o' General: 1/103	26.47%	1/140		of General: Unknown	68.75%	Unknown
	o' Italian: 1/162	35.71%	1/252	Maroteaux-Lamy Syndrome	♂ Argentinian: 1/274	75.00%	1/1,096
	o* Russian: 1/103	53.33%	1/221		♂ General: 1/388	61.54%	1/1,009
	o' US Amish: Unknown	>99%	Unknown		♂ Spaniard: 1/274	29.17%	1/387
imb-Girdle Muscular Dystrophy: Type	o' Caucasus Jewish: 1/25	>99%	<1/2,500	Meckel Syndrome: Type 1	♂ European: 1/212	72.22%	1/763
28					♂ Finnish: 1/48	>99%	<1/4,800
	♂ Libyan Jewish: 1/19	>99%	<1/1,900	Medium-Chain Acyl-CoA	♂ European: 1/50	90.91%	1/550
imb-Girdle Muscular Dystrophy: Type C	♂ European Gypsy: 1/50	>99%	<1/5,000	Dehydrogenase Deficiency	o³ Saudi Arabian: 1/68	95.00%	1/1,360
	♂ General: Unknown	60.00%	Unknown		♂ United Kingdom: 1/51	90.00%	1/510
	♂ Tunisian: Unknown	>99%	Unknown	Megalencephalic	♂ Japanese: Unknown	50.00%	Unknown
imb-Girdle Muscular Dystrophy: Type 2D	o <sup>*</sup> Brazilian: Unknown	64.29%	Unknown	Leukoencephalopathy	o" Libyan Jewish: 1/40	>99%	<1/4,000
	o⁴ European: 1/288	22.22%	1/370		of Turkish: Unknown	20.00%	Unknown
	o' Finnish: 1/150	95.45%	1/3,300	Metachromatic Leukodystrophy	of European: 1/150	43.88%	1/267
	o' General: Unknown	26.09%	Unknown		of Habbanite Jewish: 1/5	50.00%	1/10
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## $Carrier Map^{TM}$

Disease	Carrier Rate	Detection Rate	Residual Risk	Disease	Carrier Rate	Detection Rate	Residual Risk
Methylmalonic Acidemia: MMAA Related	o'' General: 1/274	63.51%	1/751	Neuronal Ceroid-Lipofuscinosis: CLN5 Related	o⁴ Finnish: 1/101	>99%	<1/10,10 0
Methylmalonic Acidemia: MMAB Related	o'' General: 1/396	71.25%	1/1,377	Neuronal Ceroid-Lipofuscinosis: CLN6 Related	o⁴ European: 1/159	36.36%	1/250
Methylmalonic Acidemia: MUT Related	♂ General: 1/177	43.62%	1/314	8	d' General: 1/159	59.52%	1/393
	o³ Chinese: Unknown	61.39%	Unknown		of Portuguese: 1/128	81.00%	1/674
Homocystinuria: Type cblC				Neuronal Ceroid-Lipofuscinosis: CLN8	o' Finnish: 1/135	>99%	<1/13,50
	♂ General: 1/159	65.74%	1/464	Related			0
	o⁴ Italian: Unknown	75.00%	Unknown		of Italian: 1/212	33.33%	1/318
	o <sup>™</sup> Portuguese: Unknown	91.18%	Unknown		O' Turkish: Unknown	77.78%	Unknown
Mitochondrial Complex I Deficiency: NDUFS6 Related	o Caucasus Jewish: 1/24	>99%	<1/2,400	Neuronal Ceroid-Lipofuscinosis: MFSD8 Related	♂ General: 1/159	56.25%	1/363
	♂ Ashkenazi Jewish: Unknown	>99%	Unknown	Neuronal Ceroid-Lipofuscinosis: PPT 1 Related	o⁴ Finnish: 1/58	97.62%	1/2,436
	o⁴ General: Unknown	47.37%	Unknown		d' General: 1/159	72.50%	1/578
	♂ Iranian Jewish: Unknown	>99%	Unknown	Neuronal Ceroid-Lipofuscinosis: TPP1	o' Canadian: 1/159	67.50%	1/489
	♂ Iranian Jewish: Unknown	>99%	Unknown	Related			
Sideroblastic Anemia	5.2 province state of the between				o' European: 1/159	75.00%	1/636
Mitochondrial Trifunctional Protein Deficiency: HADHB Related	o <sup>r</sup> Japanese: Unknown	60.00%	Unknown		of General: 1/159	50.00%	1/318
Morquio Syndrome: Type A	o' Colombian: 1/257	85.00%	1/1,713	NI DI T	of Newfoundlander: 1/43	85.29%	1/292
	o⁴ European: 1/257	20.97%	1/325	Niemann-Pick Disease: Type A	of Ashkenazi Jewish: 1/101	95.00%	1/2,020
	o' Finnish: 1/257	50.00%	1/514	Niemann-Pick Disease: Type B	of Czech: 1/276	83.33%	1/1,656
	o' Latin American: 1/257	36.11%	1/402		of General: Unknown	19.82%	Unknown
Morquio Syndrome: Type B	o³ European: Unknown	83.33%	Unknown		of North African: Unknown	86.67%	Unknown
Mucolipidosis: Type II/III	o⁴ General: 1/158	24.60%	1/210	NI BILDI T CI	of Spaniard: Unknown	38.10%	Unknown
	o Japanese: 1/252	51.25%	1/517	Niemann-Pick Disease: Type C1	of Acadian: Unknown	>99%	Unknown
	o³ Korean: Unknown	30.00%	Unknown		of General: 1/194	15.60%	1/230
	of Portuguese: 1/176	50.00%	1/352		of Japanese: Unknown	18.18%	Unknown
Mucolipidosis: Type IV	o <sup>a</sup> Ashkenazi Jewish: 1/97	96.15%	1/2,522	NI BILDI T CO	of Portuguese: 1/194	25.00%	1/259
Multiple Pterygium Syndrome	o' European: Unknown	41.67%	Unknown	Niemann-Pick Disease: Type C2	o' General: 1/194	75.00%	1/776
9	o' Middle Eastern: Unknown	60.00%	Unknown	Nijmegen Breakage Syndrome	of Eastern European: 1/155	>99%	<1/15,50 0
1.9	o⁴ Pakistani: Unknown	50.00%	Unknown	Nonsyndromic Hearing Loss and	o⁴ Ashkenazi Jewish: 1/20	95.83%	1/480
Multiple Sulfatase Deficiency	♂ Ashkenazi Jewish: 1/320	95.00%	1/6,400	Deafness: GJB2 Related			
5	o' General: 1/501	18.18%	1/612		of Chinese: 1/100	82.26%	1/564
Muscle-Eye-Brain Disease	♂ European: Unknown	54.17%	Unknown		o' European: 1/53	82.47%	1/302
9	o' Finnish: 1/112	97.37%	1/4,256		o⁴ Ghanaian: Unknown	90.91%	Unknown
	o⁴ General: Unknown	23.53%	Unknown		o⁴ Indian: Unknown	66.98%	Unknown
	of United States: Unknown	25.00%	Unknown		o" Israeli: 1/16	93.10%	1/232
Navajo Neurohepatopathy	o' Navajo: 1/39	>99%	<1/3,900		of Japanese: 1/75	75.00%	1/300
Nemaline Myopathy: NEB Related	o <sup>a</sup> Ashkenazi Jewish: 1/108	>99%	<1/10,80	8	o' Roma: Unknown	>99%	Unknown
			0		of United States: 1/34	45.22%	1/62
	of Finnish: 1/45 of US Amish: 1/12	76.84% 50.00%	1/194	Nonsyndromic Hearing Loss and Deafness: LOXHD1 Related	♂ Ashkenazi Jewish: 1/180	>99%	<1/18,00 0
	o' Israeli-Arab: Unknown	55.56%	Unknown	Nonsyndromic Hearing Loss and Deafness: MYO15A Related	o' Balinese: 1/6	>99%	<1/600
Nephrotic Syndrome: Type 2		20.00%	Unknown	Deditiess; MITO IOA Reidled	o* Pakistani: 1/77	24.00%	1 /101
	o⁴ Pakistani: Unknown				O rakisiani: 1///	14 1111/0	1/101
	of Polish: Unknown	16.18%	Unknown	Oculocutaneous Albinism Tuno 1			1 /127
			Unknown Unknown	Oculocutaneous Albinism: Type 1	of European: 1/101 of Hutterite: 1/7	26.32%	1/137



## $CarrierMap^{tM}$

Disease	Carrier Rate	Detection Rate	Residual Risk	Disease	Carrier Rate	Detection Rate	Residual Risk
	♂ Puerto Rican: Unknown	91.67%	Unknown		♂ Italian: Unknown	27.78%	Unknown
Oculocutaneous Albinism: Type 3	o Black South African: 1∕47	94.74%	1/893		o' Norwegian: 1/142	47.92%	1/273
Oculocutaneous Albinism: Type 4	o Japanese: 1/146	58.33%	1/350		♂ Sardinians: 1/61	81.82%	1/336
Omenn Syndrome: DCLRE1C Related	o* Apache: 1/29	>99%	<1/2,900		o⁴ United Kingdom: Unknown	70.00%	Unknown
	o⁴ Navajo: 1/29	97.22%	1/1,044		♂ United States: Unknown	65.62%	Unknown
Omenn Syndrome: RAG2 Related	o³ Arab: Unknown o³ Non-Ashkenazi Jewish:	40.00% 70.00%	Unknown	Pontocerebellar Hypoplasia: EXOSC3	♂ General: Unknown	83.33%	Unknown
	Unknown	70.00%	Unknown	Related Pontocerebellar Hypoplasia: RARS2	o' Sephardic Jewish:	>99%	Unknown
Ornithine Translocase Deficiency	♂ French Canadian: 1/20	95.00%	1/400	Related	Unknown	- 7770	Olikilowii
	♂ Italian: Unknown	18.75%	Unknown	Pontocerebellar Hypoplasia: SEPSECS	♂ Iraqi Jewish: 1/42	>99%	<1/4,200
	♂ Japanese: Unknown	60.00%	Unknown	Related			
Osteopetrosis: TCIRG1 Related	o⁴ Ashkenazi Jewish: 1/350	>99%	<1/35,00 0	Pontocerebellar Hypoplasia: TSEN54 Related	o⁴ European: 1/250	95.65%	1/5,750
	♂ Costa Rican: Unknown	>99%	Unknown	Pontocerebellar Hypoplasia: VPS53 Related	♂ Moroccan Jewish: 1/37	>99%	<1/3,700
	♂ General: 1/251	25.00%	1/335	Pontocerebellar Hypoplasia: VRK 1	♂ Ashkenazi Jewish: 1/225	>99%	<1/22,50
POLG Related Disorders: Autosomal Recessive	♂ Belgian: Unknown	85.00%	Unknown	Related			0
	o⁴ Finnish: 1/140	>99%	<1/14,00	Primary Carnitine Deficiency	♂ European: 1/101	58.33%	1/242
			0		of Faroese: 1/9	53.95%	1/20
	♂ General: Unknown	93.10%	Unknown	Manual Control Manual Control Control Control	o⁴ General: Unknown	20.22%	Unknown
	o⁴ Norwegian: Unknown	>99%	Unknown	Primary Ciliary Dyskinesia: DNAI 1 Related	♂ European: 1/211	52.38%	1/443
Papillon-Lefevre Syndrome	♂ General: Unknown	35.29%	Unknown	Primary Ciliary Dyskinesia: DNAI2	♂ Ashkenazi Jewish: 1/200	>99%	<1/20,00
	♂ Indian Jewish: Unknown	>99%	Unknown	Related	, ,, , ,		0
	♂ Turkish: Unknown	50.00%	Unknown	Primary Congenital Glaucoma	♂ Moroccan: Unknown	>99%	Unknown
Pendred Syndrome	♂ European: 1/58	42.11%	1/100		♂ Saudi Arabian: 1/23	91.67%	1/276
	♂ Japanese: Unknown	45.83%	Unknown	*	o⁴ Turkish: 1/51	70.59%	1/173
	oʻ Pakistani: Unknown	29.82%	Unknown	Primary Hyperoxaluria: Type 1	of Dutch: 1/174	62.12%	1/459
Persistent Mullerian Duct Syndrome: Type I	♂ General: Unknown	28.12%	Unknown		♂ General: 1/189	52.68%	1/399
Persistent Mullerian Duct Syndrome:	♂ General: Unknown	78.12%	Unknown	Primary Hyperoxaluria: Type 2	♂ General: Unknown	70.31%	Unknown
Туре II				Primary Hyperoxaluria: Type 3	o⁴ Ashkenazi Jewish: Unknown	>99%	Unknown
Phenylalanine Hydroxylase Deficiency		46.08%	Unknown		o⁴ European: Unknown	25.00%	Unknown
	♂ Ashkenazi Jewish: 1/224	44.44%	1/403	Progressive Familial Intrahepatic	o⁴ European; Unknown	33.33%	Unknown
	o⁴ Brazilian: 1/71	56.41%	1/163	Cholestasis: Type 2			
	of Chinese: 1/51	76.57%	1/218	Propionic Acidemia: PCCA Related	♂ Japanese: 1/102	86.67%	1/765
	o⁴ Cuban: 1/71	69.64%	1/234	Propionic Acidemia: PCCB Related	♂ General: 1/182	42.86%	1/319
	♂ European: 1/51	73.00%	1/189		♂ Greenlandic Inuit: 1/16	58.33%	1/38
	of French Canadian: 1/80	76.27%	1/337		♂ Japanese: 1/102	78.00%	1/464
	♂ Iranian: 1/31	66.94%	1/94		♂ Korean: Unknown	56.25%	Unknown
	o' Korean: 1/51	51.52%	1/105		o⁴ Latin American: 1/182	75.00%	1/728
	♂ Non-Ashkenazi Jewish: Unknown	63.64%	Unknown	Pseudocholinesterase Deficiency	o" Spaniard: 1/182 o" General: 1/33	52.38% 65.00%	1/382 1/94
	♂ Slovakian Gypsy: 1/39	>99%	<1/3,900	r seddocholinesierose Deficiency	o' Iranian Jewish: 1/9	>99%	<1/900
	♂ Spanish Gypsy: 1/4	93.75%	1/64	Pugnodurostaria	o' Danish: Unknown	87.50%	Unknown
	♂ Taiwanese: Unknown	83.10%	Unknown	Pycnodysostosis			
	♂ US Amish: 1/16	86.84%	1/122	Pyruvate Carboxylase Deficiency	of General: 1/251	62.50%	1/669
Polyglandular Autoimmune Syndrome: Type I	o⁴ Finnish: 1/80	90.48%	1/840	Pyruvate Dehydrogenase Deficiency	d' Native American: 1/10 d' General: Unknown	>99% 50.00%	<1/1,000 Unknown
9	o⁴ Iranian Jewish: 1/48	>99%	<1/4,800				
				t and the second			



## $Carrier Map^{TM}$

Disease	Carrier Rate	Detection Rate	Residual Risk	Disease	Carrier Rate	Detection Rate	Residual Risk
Renal Tubular Acidosis and Deafness	o' Colombian (Antioquia):	92.86%	Unknown		♂ General: Unknown	75.00%	Unknown
Retinal Dystrophies: RLBP1 Related	Unknown  O' Newfoundlander: 1/106	>99%	<1/10,60	Sulfate Transporter-Related Osteochondrodysplasia	of Finnish: 1/51	95.83%	1/1,224
			0		♂ General: 1/100	70.00%	1/333
	of Swedish: 1/84	>99%	<1/8,400	Tay-Sachs Disease	o⁴ Argentinian: 1/280	82.35%	1/1,587
Retinal Dystrophies: RPE65 Related	of Dutch: 1/32	>99%	<1/3,200	100	♂ Ashkenazi Jewish: 1/29	99.53%	1/6,177
	o <sup>a</sup> North African Jewish: Unknown	>99%	Unknown		o' Cajun: 1/30	>99%	<1/3,000
Retinitis Pigmentosa: CERKL Related	of Yemenite Jewish: Unknown	>99%	Unknown	li A	o' European: 1/280	25.35%	1/375
Retinitis Pigmentosa: DHDDS Related	♂ Ashkenazi Jewish: 1/91	>99%	<1/9,100		d' General: 1/280	32.09%	1/412
Retinitis Pigmentosa: FAM 161 A Related	o⁴ Ashkenazi Jewish: Unknown	>99%	Unknown		o' Indian: Unknown	85.71%	Unknown
	o" Non-Ashkenazi Jewish:	>99%	<1/3,200		o' Iraqi Jewish: 1/140	56.25%	1/320
	1/32	-7776	1/3,200		of Japanese: 1/127	82.81%	1/739
Rhizomelic Chondrodysplasia	of General: 1/159	72.68%	1/582		of Moroccan Jewish: 1/110	22.22%	1/141
unctata: Type I	200			1 1	of Portuguese: 1/280	92.31%	1/3,640
alla Disease	o' European: Unknown	33.33%	Unknown		of Spaniard: 1/280	67.65%	1/865
	of Scandinavian: 1/200	94.27%	1/3,491		of United Kingdom: 1/161	71.43%	1/564
Sandhoff Disease	of Argentinian: Unknown	95.45%	Unknown	Trichohepatoenteric Syndrome: Type 1	of European: 1/434	42.86%	1/760
	of Cypriot: 1/7	80.00%	1/35	T	of South Asian: 1/434	66.67%	1/1,302
	o <sup>*</sup> Italian: Unknown	29.17%	Unknown	Tyrosine Hydroxylase Deficiency	of General: Unknown	36.11%	Unknown
	♂ Spaniard: Unknown	64.29%	Unknown	Tyrosinemia: Type I	o³ Ashkenazi Jewish: 1/158	>99%	<1/15,80 0
Sanfilippo Syndrome: Type A	o⁴ Australasian: 1/119	44.12%	1/213		♂ European: 1/166	57.14%	1/387
	o* Dutch: 1/78	63.10%	1/211		♂ Finnish: 1/123	97.22%	1/4,428
	of European: 1/159	35.16%	1/245		♂ French Canadian: 1/64	96.30%	1/1,728
	of United States: 1/159	32.14%	1/234		o⁴ Pakistani: Unknown	92.86%	Unknown
Sanfilippo Syndrome: Type B	of Australasian: 1/230	28.00%	1/319	Tyrosinemia: Type II	♂ General: 1/251	40.00%	1/418
	of Dutch: Unknown	42.31%	Unknown	Usher Syndrome: Type 1B	♂ European: 1/166	39.29%	1/273
	o' European: Unknown	52.38%	Unknown		♂ General: 1/143	12.89%	1/164
	of Japanese: 1/200	81.82%	1/1,100		o' North African: Unknown	66.67%	Unknown
Sanfilippo Syndrome: Type C	of Dutch: 1/346	75.00%	1/1,384		o⁴ Spaniard: 1/152	12.16%	1/173
	o' Greek: 1/415	25.00%	1/553	Usher Syndrome: Type 1C	of Acadian: 1/82	98.86%	1/7,216
	of Moroccan: Unknown	80.00%	Unknown		♂ French Canadian: 1/227	83.33%	1/1,362
	o' Spaniard: Unknown	64.29%	Unknown	Usher Syndrome: Type 1D	♂ General: 1/296	23.17%	1/385
anfilippo Syndrome: Type D	o' General: 1/501	83.33%	1/3,006	Usher Syndrome: Type 1F	♂ Ashkenazi Jewish: 1/126	93.75%	1/2,016
hort-Chain Acyl-CoA Dehydrogenase Deficiency	♂ Ashkenazi Jewish: 1/15	65.00%	1/43	Usher Syndrome: Type 2A	o⁴ Chinese: Unknown	83.33%	Unknown
iickle-Cell Anemia	o' African American: 1/10	>99%	<1/1,000		♂ European: 1/136	40.00%	1/227
	of Hispanic American: 1/95	>99%	<1/9,500		o'' French Canadian: Unknown	66.67%	Unknown
Sjogren-Larsson Syndrome	o⁴ Dutch: Unknown	25.86%	Unknown		of General: 1/136	46.92%	1/256
	o⁴ Swedish: 1/205	>99%	<1/20,50		o General: 17 130 o Japanese: Unknown	55.56%	Unknown
Sly Syndrome	of General: 1/251	35.71%	0 1/390		o Japanese, Onknown o Japanese, Onknown o Japanese, Onknown	61.11%	Unknown
Smith-Lemli-Opitz Syndrome	o' Brazilian: 1/94	79.17%	1/451		Unknown		0.00000
Olimin Colin Copin Copin Colin	o' European: 1/71	84.72%	1/465		♂ Scandinavian: 1/125	39.22%	1/206
	o' Japanese: Unknown	71.43%	Unknown		♂ Spaniard: 1/133	39.02%	1/218
	o <sup>™</sup> United States: 1/70	95.00%	1/1,400	Usher Syndrome: Type 3	♂ Ashkenazi Jewish: 1/120	>99%	<1/12,00 0
Stargardt Disease	♂ General: 1/51	17.51%	1/62		♂ Finnish: 1/134	>99%	<1/13,40
Stuve-Wiedemann Syndrome	o' Emirati: 1/70	>99%	<1/7,000			W 5555	0





Disease	Carrier Rate	Detection Rate	Residual Risk
Very Long-Chain Acyl-CoA Dehydrogenase Deficiency	o' General: 1/87	65.28%	1/251
Walker-Warburg Syndrome	♂ Ashkenazi Jewish: 1/150	>99%	<1/15,00 0
Werner Syndrome	of General: 1/224	31.25%	1/326
	♂ Japanese: 1/87	65.62%	1/253
Wilson Disease	o⁴ Ashkenazi Jewish: 1/100	>99%	<1/10,00 0
	o⁴ Canarian: 1/26	68.75%	1/83
	of Chinese: 1/51	55.97%	1/116
	of Cuban: Unknown	22.22%	Unknown
	o' European: 1/93	41.64%	1/159
	o' Greek: 1/90	44.94%	1/163
	o⁴ Korean: 1/88	51.53%	1/182
	♂ Spaniard: 1/93	38.18%	1/150
Wolcott-Rallison Syndrome	og Saudi Arabian: Unknown	66.67%	Unknown
Wolman Disease	♂ Iranian Jewish: 1/33	>99%	<1/3,300
Xeroderma Pigmentosum: Group A	♂ Japanese: 1/75	97.62%	1/3,150
	o <sup>®</sup> North African: Unknown	87.50%	Unknown
	♂ Tunisian: 1/112	90.91%	1/1,232
Xeroderma Pigmentosum: Group C	♂ Moroccan: 1/71	76.19%	1/298
	♂ Tunisian: 1/51	>99%	<1/5,100
Zellweger Spectrum Disorders: PEX1 Related	of European: 1/139	70.27%	1/468
	o⁴ General: 1/139	67.84%	1/432
Zellweger Spectrum Disorders: PEX10 Related	o³ Japanese: Unknown	40.74%	Unknown
Zellweger Spectrum Disorders: PEX2 Related	o <sup>a</sup> Ashkenazi Jewish: 1/123	>99%	<1/12,30 0
Zellweger Spectrum Disorders: PEX6 Related	o' General: 1/288	30.00%	1/411