

COMPREHENSIVE PANEL

Condition	Gene	Population	Carrier frequency before screening (1 in)	Detection rate	Carrier residual risk after negative result (1 in)
3-Hydroxy-3-Methylglutaryl-Coenzyme A Lyase Deficiency	HMGCL	General population	*500	99%	Reduced
		Portugese	160	99%	15844
		Saudi Arabia	50	99%	4951
3-Methylcrotonyl-CoA Carboxylase Deficiency 1	MCCC1	European	147	99%	14556
3-Methylcrotonyl-CoA Carboxylase Deficiency 2	MCCC2	General Population	120	99%	11883
3-Methylglutaconic Aciduria, Type 3 (Costeff Syndrome)	OPA3	General Population	*500	99%	Reduced
3-Phosphoglycerate Dehydrogenase Deficiency	PHGDH	Ashkenazi Jewish	400	99%	39608
		General Population	*500	99%	Reduced
6-Pyruvoyl-Tetrahydropterin Synthase (PTPS) Deficiency	PTS	East Asian	122	99%	12081
		General Population	*500	99%	Reduced
Abetalipoproteinemia	MTTP	Ashkenazi Jewish	131	99%	12972
Achondrogenesis, Type 1B	SLC26A2	Finnish	75	99%	7427
		General Population	158	99%	15646
Achromatopsia (CNGB3-related)	CNGB3	General Population	146	98%	7301
Acute Infantile Liver Failure (TRMU-related)	TRMU	General Population	*500	99%	Reduced
		Sephardic Jewish - Yemenite	34	99%	3367
Acyl-CoA Oxidase I Deficiency	ACOX1	General Population	*500	99%	Reduced
Adrenoleukodystrophy, X-Linked	ABCD1	General Population	16800	95%	338035
		Sephardic Jewish	*500	99%	Reduced
Aicardi-Goutières Syndrome	SAMHD1	General Population	*500	95%	Reduced
Alpha Thalassemia	HBA1, HBA2	African American	30	90%	276
		Asian	20	90%	184
		Caucasian	*500	90%	Reduced
		General Population	25	90%	230
Alport Syndrome (COL4A3-related)	COL4A3	Ashkenazi Jewish	183	99%	18121
		General Population	354	>98%	17701
Alport Syndrome, X-Linked	COL4A5	General Population	*500	>90%	Reduced
Alstrom Syndrome	ALMS1	General Population	*500	>90%	Reduced
Andermann Syndrome	SLC12A6	French Canadian	23	99%	2278
		General Population	*500	99%	Reduced
Argininosuccinate Lyase Deficiency	ASL	General Population	132	99%	13071
Aromatase Deficiency	CYP19A1	General Population	*500	99%	Reduced
Arthrogryposis Mental Retardation Seizures	SLC35A3	Ashkenazi Jewish	453	99%	44856
Asparagine Synthetase Deficiency	ASNS	General Population	*500	99%	Reduced
		Sephardic Jewish - Iranian	80	99%	7922
Aspartylglycosaminuria	AGA	Finnish	63	99%	6239
		General Population	*500	99%	Reduced
Ataxia with Vitamin E Deficiency	TTPA	General Population	*500	99%	Reduced
Ataxia-Telangiectasia	ATM	General Population	100	>91%	1120
		Romani Population	36	99%	3565
Autoimmune Polyglandular Syndrome, Type 1	AIRE	Finnish	79	>87%	613
		Sardinian	60	>87%	466
		Sephardic Jewish - Iranian	27	>87%	210
Autosomal Recessive Polycystic Kidney Disease	PKHD1	Ashkenazi Jewish	105	99%	10398
		Caucasian	100	99%	9902
		General Population	70	99%	6932

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Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay	SACS	French Canadian - Charlevoix-Saguenay	21	99%	2080
		General Population	*500	99%	Reduced
Bardet-Biedl syndrome (BBS1-related)	BBS1	General Population	330	99%	32677
Bardet Biedl syndrome (BBS12-related)	BBS12	General Population	*500	99%	Reduced
Bare Lymphocyte Syndrome (CIITA-related)	CIITA	General Population	*500	99%	Reduced
Bartter Syndrome (BSND-related)	BSND	General Population	*500	99%	Reduced
Batten Disease (CLN3-related)	CLN3	General Population	230	99%	22775
Beta Thalassemia	HBB	African-American	8	99%	793
		Chinese	29	99%	2872
		Mediterranean	28	99%	2773
		Middle-Eastern	30	99%	2971
		Thai	20	99%	1981
Biotinidase Deficiency	BTM	General Population	120	99%	11883
Bloom Syndrome	BLM	Ashkenazi Jewish	100	99%	9902
Canavan Disease	ASPA	Ashkenazi Jewish	57	99%	5645
		European	3392	99%	Reduced
Carnitine Palmitoyltransferase IA Deficiency	CPT1A	General Population	*500	99%	Reduced
		Hutterite	16	99%	1585
Carnitine Palmitoyltransferase II Deficiency	CPT2	Ashkenazi Jewish	45	99%	4456
		General Population	182	99%	18022
Carpenter Syndrome	RAB23	General Population	*500	99%	Reduced
Cartilage-Hair Hypoplasia	RMRP	Amish	19	99%	1882
		Finnish	76	99%	7526
		General Population	*500	99%	Reduced
Cerebrotendinous Xanthomatosis	CYP27A1	General Population	112	99%	11091
		Sephardic Jewish	76	99%	7526
Choreacanthocytosis	VPS13A	Ashkenazi Jewish	NA	99%	Reduced
Choroideremia, X-Linked	CHM	General Population	*500	>75%	Reduced
Chronic Granulomatous Disease, X-Linked	CYBB	General Population	*500	>90%	Reduced
Citrin Deficiency	SLC25A13	General Population	*500	>94%	8390
Citrullinemia, Type 1	ASS1	General Population	119	>96%	2991
Combined Malonic and Methylmalonic Acidemia	ACSF3	General Population	87	99%	8615
Combined Oxidative Phosphorylation Deficiency 1	GFM1	General Population	*500	99%	Reduced
Combined Oxidative Phosphorylation Deficiency 3	TSFM	Finnish	80	99%	7922
Combined Pituitary Hormone Deficiency 2	PROP1	General Population	45	>98%	2251
Congenital Disorder of Glycosylation, Type 1A (PMM2-related)	PMM2	Ashkenazi Jewish	61	99%	6041
		Caucasian	60	99%	5942
		General Population	190	99%	18814
Congenital Disorder of Glycosylation, Type 1B	MPI	General Population	*500	99%	Reduced
Congenital Disorder of Glycosylation, Type 1C	ALG6	General Population	*500	98%	Reduced
Congenital Finnish Nephrosis	NPHS1	Finnish	46	99%	4555
		General Population	*500	99%	Reduced
		Groffdale Conference Mennonite	12	99%	1189
Congenital Insensitivity to Pain with Anhidrosis	NTRK1	General Population	*500	99%	Reduced
Congenital Myasthenic Syndrome (CHRNE-related)	CHRNE	General Population	200	99%	19804
		Roma - Southeastern European	25	99%	2476
Congenital Myasthenic Syndrome (RAPSN-related)	RAPSN	General Population	283	99%	28023
Congenital Neutropenia (HAX1-related)	HAX1	General Population	*500	99%	Reduced
Congenital Neutropenia (VPS45-related)	VPS45	General Population	*500	99%	Reduced
Corneal Dystrophy and Perceptive Deafness	SLC4A11	General Population	*500	99%	Reduced

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Corticosterone Methyloxidase Deficiency	CYP11B2	General Population	*500	99%	Reduced
		Sephardic Jewish - Iranian	30	99%	2971
CRB1-Related Retinal Dystrophies	CRB1	General Population	112	99%	11091
Creatine Transporter Defect (Cerebral Creatine Deficiency Syndrome 1), X-Linked	SLC6A8	General Population	*500	>98%	Reduced
Crigler Najjar Syndrome, Type I	UGT1A1	Amish	61	99%	Reduced
Cystic Fibrosis	CFTR	General Population	45	97%	1506
Cystinosis	CTNS	French Canadian - Saguenay Lac-St. Jean	39	99%	3862
		General Population	224	99%	22181
		Sephardic Jewish - Moroccan	100	99%	9902
D-Bifunctional Protein Deficiency	HSD17B4	General Population	158	96%	3970
Deafness, Autosomal Recessive 77	LOXHD1	Ashkenazi Jewish	180	99%	17824
		General Population	*500	99%	Reduced
Dystrophic Epidermolysis Bullosa (COL7A1-related)	COL7A1	General Population	370	99%	36638
Dystrophinopathies (Duchenne/Becker Muscular Dystrophy)	DMD	General Population	*500	90%	Reduced
Ehlers-Danlos Syndrome, Type VIIC	ADAMTS2	Ashkenazi Jewish	187	93%	2691
		General Population	*500	93%	Reduced
Emery-Dreifuss Muscular Dystrophy 1, X-Linked	EMD	General Population	200	99%	19804
Enhanced S-Cone Syndrome	NR2E3	General Population	*500	99%	Reduced
Ethylmalonic Encephalopathy	ETHE1	General Population	*500	>84%	Reduced
Fabry Disease	GLA	General Population	*500	99%	Reduced
Factor IX Deficiency	F9	General Population	*500	98%	Reduced
Factor V Leiden Thrombophilia	F5	European	NA	99%	Reduced
		General Population	12-33	99%	1189
		US (African American)	NA	99%	Reduced
		US (Caucasian)	NA	99%	Reduced
		US (Hispanic)	NA	99%	Reduced
Factor XI Deficiency	F11	Ashkenazi Jewish	11	99%	1090
		European	123	99%	12180
Familial Dysautonomia	ELP1	Ashkenazi Jewish	31	99%	3070
Familial Hypercholesterolemia (LDLR-related)	LDLR	Ashkenazi Jewish	69	92%	870
		French Canadian	270	92%	3401
		General Population	250	92%	3149
		South African Afrikaner	72	92%	907
Familial Mediterranean Fever	MEFV	Armenian	17	99%	1684
		Cypriot	25	99%	2476
		Jewish	58	99%	5744
		Turkish	125	99%	12378
Familial Nephrogenic Diabetes Insipidus (AQP2-related)	AQP2	General Population	*500	99%	Reduced
Fanconi Anemia, Type C	FANCC	Ashkenazi Jewish	89	99%	8813
		General Population	417	99%	41292
Fanconi Anemia, Type G	FANCG	French-Canadian	NA	99%	Reduced
		Korean/Japanese	NA	99%	Reduced
		Portuguese-Brazilian	NA	99%	Reduced
Fragile X Syndrome, X-linked	FMR1	African American	267	99%	26439
		Ashkenazi Jewish	102	99%	10101
		Asian	419	99%	41490
		Caucasian	170	99%	16834
		General Population	201	99%	19903
		Hispanic	253	99%	25052

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Galactokinase Deficiency (Galactosemia, Type II)	GALK1	General Population	122	99%	12081
		Roma	47	99%	4654
Galactosemia	GALT	African American	87	99%	8615
		Ashkenazi Jewish	156	99%	15448
		General Population	100	99%	9902
		Irish Travellers	11	99%	1090
Gaucher Disease	GBA	Ashkenazi Jewish	15	99%	1486
		General Population	158	99%	15646
Glutaric Acidemia, Type 1	GCDH	Amish - Pennsylvania	10	99%	991
		General Population	158	99%	15646
		Oji-Cree First Nations - N. Manitoba	10	99%	991
Glutaric Acidemia, Type 2A	ETFA	General Population	*500	99%	Reduced
Glycine Encephalopathy (AMT-related)	AMT	General Population	325	99%	32182
Glycine Encephalopathy (GLDC-related)	GLDC	Finnish	NA	>96%	Reduced
		General Population	NA	>96%	Reduced
Glycogen Storage Disease, Type 1A	G6PC	Ashkenazi Jewish	71	99%	7031
		General Population	177	99%	17527
Glycogen Storage Disease, Type 1B	SLC37A4	General Population	354	99%	35053
Glycogen Storage Disease, Type 2 (Pompe Disease)	GAA	General Population	100	>90%	1009
Glycogen Storage Disease, Type 3	AGL	Faroese	30	99%	2971
		Finnish	1580	99%	Reduced
		Sephardic Jewish - Moroccan	35	99%	3466
Glycogen Storage Disease, Type 4	GBE1	Ashkenazi Jewish	68	99%	6734
		General Population	387	>90%	3902
Glycogen Storage Disease, Type 5 (McArdle Disease)	PYGM	General Population	*500	99%	49510
		Sephardic Jewish - Kurdish	84	99%	8318
Glycogen Storage Disease, Type 7	PFKM	General Population	*500	99%	49510
GRACILE Syndrome	BCS1L	Finnish	NA	99%	Reduced
Hemochromatosis, Type 2A	HFE2	General Population	*500	99%	Reduced
Hemochromatosis, Type 3 (TFR2-related)	TFR2	General Population	*500	99%	Reduced
Hereditary Fructose Intolerance	ALDOB	Central European	95	99%	9407
		Northern India	NA	99%	Reduced
		Spanish	NA	99%	Reduced
		US	55-120	99%	5447
Hermansky-Pudlak Syndrome (HPS1-related)	HPS1	General Population	*500	99%	Reduced
		Puerto Rican (Northwestern)	21	99%	2080
Hermansky-Pudlak Syndrome (HPS3-related)	HPS3	Ashkenazi Jewish	235	99%	23270
		General Population	*500	99%	Reduced
Holocarboxylase Synthetase Deficiency	HLCS	Faroese	50	99%	4951
		General Population	224	99%	22181
		Japanese	158	99%	15646
Homocystinuria due to Cystathionine Beta-synthase Deficiency	CBS	General Population	224	98%	11201
		Norwegian	40	98%	2001
		Qatari	21	98%	1051
Homocystinuria-Megaloblastic Anemia, cbIE Type	MTRR	Caucasian	*500	>96%	Reduced
Hydrolethalus Syndrome	HYLS1	Finnish	50	99%	4951
Hypohidrotic Ectodermal Dysplasia, X-Linked	EDA	General Population	112	>90%	1130

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Hypophosphatasia (ALPL-related)	ALPL	General Population	150	98%	7501
		Mennonite	25	99%	2476
Inclusion Body Myopathy, Type 2	GNE	Asian	58	99%	5744
		Iranian Jewish	11-100	99%	1090
Isovaleric Acidemia	IVD	General Population	150	98%	7501
Joubert Syndrome, Type 2	TMEM216	Ashkenazi Jewish	110	99%	10893
Junctional Epidermolysis Bullosa, Herlitz Type	LAMC2	Italian	858	99%	Reduced
Juvenile Retinoschisis, X-Linked	RS1	General Population	*500	>90%	Reduced
Krabbe Disease	GALC	Druze Northern Israel	6	99%	595
		General Population	158	99%	15646
Lamellar Ichthyosis, Type 1	TGM1	General Population	301	99%	29805
		Norwegian	151	99%	14952
Leber Congenital Amaurosis (LCA5-related)	LCA5	Ashkenazi Jewish	100	99%	Reduced
Leber Congenital Amaurosis, Type CEP290	CEP290	General Population	185	97%	6188
Leigh Syndrome, French-Canadian Type	LRPPRC	French Canadian - Saguenay Lac-St. Jean	23	99%	2278
		General Population	*500	99%	Reduced
Leukoencephalopathy with Vanishing White Matter	EIF2B5	General Population	*500	99%	Reduced
Leydig Cell Hypoplasia (Luteinizing Hormone Resistance)	LHCGR	Brazilian	NA	99%	Reduced
Limb-Girdle Muscular Dystrophy, Type 2A	CAPN3	General Population	158	>90%	1593
Limb-Girdle Muscular Dystrophy, Type 2B	DYSF	General Population	311	99%	30796
		Sephardic Jewish - Libyan	10	99%	991
Limb-Girdle Muscular Dystrophy, Type 2C	SGCG	General Population	*500	99%	Reduced
		Moroccan	250	99%	24755
		Roma	59	99%	5843
Limb-Girdle Muscular Dystrophy, Type 2D	SGCA	General Population	*500	>95%	Reduced
Limb-Girdle Muscular Dystrophy, Type 2E	SGCB	General Population	*500	>90%	Reduced
Lipoamide Dehydrogenase Deficiency (Maple Syrup Urine Disease, Type 3)	DLD	Ashkenazi Jewish	110	99%	10893
Lipoid Adrenal Hyperplasia	STAR	General Population	*500	99%	Reduced
		Korean	170	99%	16834
Lipoprotein Lipase Deficiency	LPL	African population	308	98%	15401
		French Canadian	46	98%	2301
		General Population	*500	98%	Reduced
Long Chain 3-Hydroxyacyl-CoA Dehydrogenase Deficiency	HADHA	European	210	99%	20795
Lysinuric Protein Intolerance	SLC7A7	Finnish	122	99%	12081
		General Population	*500	99%	Reduced
		Italians	NA	99%	Reduced
		Japanese	119	99%	11784
Maple Syrup Urine Disease, Type 1B	BCKDHB	Ashkenazi Jewish	97	99%	9605
Meckel-Gruber Syndrome, Type 1	MKS1	Finnish	47	>95%	946
		General Population	260	>95%	5232
Medium Chain Acyl-CoA Dehydrogenase Deficiency	ACADM	General Population	66	98%	3301
Megalencephalic Leukoencephalopathy with Subcortical Cysts	MLC1	General Population	*500	99%	Reduced
		Sephardic Jewish - Libyan	40	99%	3961
Metachromatic Leukodystrophy (ARSA-related)	ARSA	General Population	100	99%	9902
		Navajo	40	99%	3961
		Sephardic Jewish - Yemenite	46	99%	4555
Metachromatic Leukodystrophy (PSAP-related)	PSAP	General Population	*500	99%	Reduced
Methylmalonic Acidemia, cbIA Type	MMAA	General Population	316	99%	31291

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Methylmalonic Acidemia, cb1B Type	<i>MMAB</i>	General Population	456	99%	45153
Methylmalonic Acidemia and Homocystinuria, Type cb1C	<i>MMACHC</i>	General Population	123	99%	12180
Methylmalonic Acidemia and Homocystinuria, Type cb1D	<i>MMADHC</i>	General Population	*500	99%	Reduced
Methylmalonic acidemia due to Methylmalonyl-CoA Mutase Deficiency	<i>MMUT</i>	Asian	53	99%	5249
		General Population	383	99%	37925
Microphthalmia/Anophthalmia ( <i>VSX2</i> -related)	<i>VSX2</i>	General Population	*500	99%	Reduced
Mitochondrial Complex 1 Deficiency ( <i>ACAD9</i> -related)	<i>ACAD9</i>	General Population	*500	99%	Reduced
Mitochondrial Complex 1 Deficiency ( <i>NDUFAF5</i> -related)	<i>NDUFAF5</i>	Ashkenazi Jewish	290	99%	28716
		General Population	*500	99%	Reduced
Mitochondrial Complex 1 Deficiency ( <i>NDUFS6</i> -related)	<i>NDUFS6</i>	Caucasus Jewish	24	99%	2377
		General Population	*500	99%	Reduced
Mitochondrial Myopathy and Sideroblastic Anemia (MLASA1)	<i>PUS1</i>	General Population	*500	99%	Reduced
Mucopolipidosis II/III	<i>GNPTAB</i>	General Population	200	99%	19804
		Irish Traveller	15	99%	1486
Mucopolipidosis III Gamma	<i>GNPTG</i>	General Population	*500	97%	Reduced
Mucopolipidosis, Type IV	<i>MCOLN1</i>	Ashkenazi Jewish	100	97%	3345
		General Population	*500	97%	Reduced
Mucopolysaccharidosis, Type II (Hunter Syndrome), X-Linked	<i>IDS</i>	General Population	*500	>82%	Reduced
Mucopolysaccharidosis, Type IIIB (Sanfilippo B)	<i>NAGLU</i>	General Population	224	99%	22181
Mucopolysaccharidosis, Type IIIC (Sanfilippo C)	<i>HGSNAT</i>	Caucasians	259	97%	8663
Mucopolysaccharidosis IIID (Sanfilippo Syndrome, Type D)	<i>GNS</i>	General Population	*500	92%	Reduced
Mucopolysaccharidosis, Type IX	<i>HYAL1</i>	General Population	*500	99%	Reduced
Multiple Sulfatase Deficiency	<i>SUMF1</i>	Ashkenazi Jewish	279	99%	27627
		General Population	*500	99%	Reduced
Myoneurogastrointestinal Encephalopathy (MNGIE)	<i>TYMP</i>	Sephardic Jewish - Iranian	158	99%	15646
		General Population	*500	99%	Reduced
Myotubular Myopathy, X-Linked	<i>MTM1</i>	General Population	*500	95%	Reduced
N-acetylglutamate Synthase Deficiency	<i>NAGS</i>	General Population	*500	99%	Reduced
Navajo Neurohepatopathy (MPV17-related Hepatocerebral Mitochondrial DNA Depletion Syndrome)	<i>MPV17</i>	Navajo	20	99%	1981
Neurological Ceroid Lipofuscinosis ( <i>TPP1</i> -related)	<i>TPP1</i>	General Population	250	99%	24755
		Newfoundland	53	99%	5249
Neuronal Ceroid Lipofuscinosis ( <i>CLN5</i> -related)	<i>CLN5</i>	Finnish	115	>98%	5751
		General Population	*500	>98%	Reduced
Neuronal Ceroid Lipofuscinosis ( <i>CLN6</i> -related)	<i>CLN6</i>	General Population	*500	99%	Reduced
Neuronal Ceroid Lipofuscinosis ( <i>CLN8</i> -related)	<i>CLN8</i>	Finnish	135	99%	13368
		General Population	*500	99%	Reduced
Neuronal Ceroid Lipofuscinosis ( <i>MFSD8</i> -related)	<i>MFSD8</i>	General Population	*500	99%	Reduced
		Turkish	NA	99%	Reduced
Neuronal Ceroid Lipofuscinosis ( <i>PPT1</i> -related)	<i>PPT1</i>	Finnish	70	99%	6932
		General Population	199	99%	19705
Niemann-Pick Disease, Types A/B	<i>SMPD1</i>	Ashkenazi Jewish	115	99%	11388
		General Population	250	99%	24755
Niemann-Pick Disease, Type C1/D	<i>NPC1</i>	General Population	183	98%	9151
Niemann-Pick Disease, Type C2	<i>NPC2</i>	General Population	*500	99%	Reduced
Nijmegen Breakage Syndrome	<i>NBN</i>	Eastern European	115	99%	15349
		North American	NA	99%	Reduced
Non-Syndromic Hearing Loss ( <i>GJB2</i> -related)	<i>GJB2</i>	Ashkenazi Jewish	13	99%	1288
		General Population	50	99%	4951
Non-Syndromic Hearing Loss ( <i>GJB6</i> -related)	<i>GJB6</i>	General Population	423	99%	41886



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Odonto-Onycho-Dermal Dysplasia / Schopf-Schulz-Passarge Syndrome	<i>WNT10A</i>	General Population	305	99%	30202
Omenn Syndrome (RAG2-related)	<i>RAG2</i>	General Population	*500	99%	Reduced
Ornithine Aminotransferase Deficiency	<i>OAT</i>	General Population	*500	>96%	12564
Ornithine Transcarbamylase Deficiency	<i>OTC</i>	General Population	*500	99%	Reduced
Ornithine Translocase Deficiency [Hyperornithinemia-Hyperammonemia-Homocitrullinuria (HHH) Syndrome]	<i>SLC25A15</i>	French-Canadian	20	99%	1981
		Japanese	NA	99%	Reduced
Pendred Syndrome	<i>SLC26A4</i>	Asian	74	99%	7328
		General Population	80	99%	7922
Peroxisome Biogenesis Disorders Zellweger Syndrome Spectrum (PEX1-related)	<i>PEX1</i>	General Population	134	>98%	6701
Peroxisome Biogenesis Disorders Zellweger Syndrome Spectrum (PEX2-related)	<i>PEX2</i>	Ashkenazi Jewish	227	99%	22478
Phenylalanine Hydroxylase Deficiency	<i>PAH</i>	Ashkenazi Jewish	250	99%	24755
		Caucasian	50	99%	4951
		East Asian	50	99%	4951
		European	50	99%	4951
		Finland	200	99%	19804
		General Population	140	99%	13863
		Ireland	33	99%	3268
		Japanese	200	99%	19804
Turkish	26	99%	2575		
Pituitary Hormone Deficiency, Combined 3	<i>LHX3</i>	General Population	*500	99%	Reduced
Pontocerebellar Hypoplasia (RARS2-related)	<i>RARS2</i>	General Population	*500	99%	Reduced
Pontocerebellar Hypoplasia, Type 1A	<i>VRK1</i>	Ashkenazi Jewish	225	99%	22280
		General Population	*500	99%	Reduced
Pontocerebellar Hypoplasia, Type 2D	<i>SEPSECS</i>	General Population	*500	99%	Reduced
		Sephardic Jewish - Moroccan, Iraqi	43	99%	4258
Pontocerebellar Hypoplasia, Type 2E	<i>VPS53</i>	Moroccan Jewish	37	99%	3664
Primary Ciliary Dyskinesia (DNAH5-related)	<i>DNAH5</i>	Amish or Mennonite	NA	99%	Reduced
		General Population	106-164	99%	10497
Primary Ciliary Dyskinesia (DNAI1-related)	<i>DNAI1</i>	Ashkenazi Jewish	357	99%	35351
		General Population	150-232	99%	14853
Primary Ciliary Dyskinesia (DNAI2-related)	<i>DNAI2</i>	Ashkenazi Jewish	200	99%	19804
		General Population	*500	99%	Reduced
Primary Hyperoxaluria, Type 1	<i>AGXT</i>	European	173	97%	5787
Primary Hyperoxaluria, Type 2	<i>GRHPR</i>	General Population	*500	99%	Reduced
Primary Hyperoxaluria, Type 3	<i>HOGA1</i>	General Population	309	99%	30598
Pycnodysostosis	<i>CTSK</i>	Danish	NA	99%	Reduced
Pyruvate Dehydrogenase Deficiency (PDHB-related)	<i>PDHB</i>	General Population	*500	99%	Reduced
Pyruvate Dehydrogenase Deficiency, X-Linked	<i>PDHA1</i>	General Population	*500	99%	Reduced
Renal Tubular Acidosis and Deafness (ATP6V1B1-related)	<i>ATP6V1B1</i>	General Population	*500	99%	Reduced
		Sephardic Jewish - Syrian	140	99%	13863
Retinal Dystrophy (RLBP1-related) (Bothnia Retinal Dystrophy)	<i>RLBP1</i>	Swedish	84	99%	8318
Retinitis Pigmentosa 25 (EYS-related)	<i>EYS</i>	Ashkenazi Jewish	189	99%	Reduced
Retinitis Pigmentosa 26	<i>CERKL</i>	General Population	137	99%	13566
		Sephardic Jewish - Yemenite	24	99%	2377
Retinitis Pigmentosa 28	<i>FAM161A</i>	Ashkenazi Jewish	214	99%	21191
		General Population	289	99%	28617
		Sephardic Jewish - Libyan, Moroccan, Tunisian, Bulgarian	41	99%	4060

Condition	Gene	Population	Carrier frequency before screening (1 in)	Detection rate	Carrier residual risk after negative result (1 in)
Retinitis Pigmentosa 59 (DHDDS-related)	<i>DHDDS</i>	Ashkenazi Jewish	117	99%	11586
Retinitis Pigmentosa, X-linked	<i>RPGR</i>	General Population	259	99%	25647
Rhizomelic Chondrodysplasia Punctata, Type 1	<i>PEX7</i>	General Population	158	99%	15646
Rhizomelic Chondrodysplasia Punctata, Type 3	<i>AGPS</i>	General Population	*500	99%	Reduced
Roberts Syndrome	<i>ESCO2</i>	General Population	*500	99%	Reduced
Salla Disease	<i>SLC17A5</i>	Finnish	100	>95%	2013
		General Population	*500	93%	Reduced
Sandhoff Disease	<i>HEXB</i>	General Population	180	>90%	1815
Schimke Immunoosseous Dysplasia	<i>SMARCAL1</i>	General Population	*500	99%	Reduced
Segawa Syndrome (TH-related)	<i>TH</i>	Caucasian	224	99%	22181
		General Population	*500	99%	Reduced
Severe Combined Immunodeficiency, Type Athabaskan	<i>DCLRE1C</i>	Navajo and Apache Native Americans	48	99%	4753
Severe Combined Immunodeficiency, X-Linked	<i>IL2RG</i>	General Population	*500	99%	Reduced
Sickle-Cell Disease	<i>HBB</i>	General Population	NA	99%	Reduced
Sjögren-Larsson Syndrome	<i>ALDH3A2</i>	Swedish	100	99%	9902
		Dutch	NA	99%	Reduced
Smith-Lemli-Opitz Syndrome	<i>DHCR7</i>	African American	183	99%	18121
		Ashkenazi Jewish	43	99%	4258
		General Population	71	99%	7031
		Hispanic	167	99%	16537
		Northern European	54	99%	5348
		Southern European	66	99%	6536
Spinal Muscular Atrophy	<i>SMN1, SMN2</i>	Caucasian	35	95%	705
		Ashkenazi Jewish	41	90%	414
		Hispanic	117	90%	1180
		Asian	53	90%	535
		African American	66	71%	230
Steroid-Resistant Nephrotic Syndrome	<i>NPHS2</i>	General Population	*500	99%	Reduced
Stuve-Wiedemann Syndrome	<i>LIFR</i>	General Population	*500	99%	Reduced
Tay-Sachs Disease	<i>HEXA</i>	Ashkenazi Jewish	25	99%	2476
		General Population	250	99%	24755
Tyrosinemia, Type 1	<i>FAH</i>	Finnish	122	99%	12081
		French Canadian	66	99%	6536
		French Canadian - Saguenay Lac-St. Jean	16	99%	1585
		General Population	125	99%	12378
Usher Syndrome, Type 1C	<i>USH1C</i>	French Canadian / Acadian	227	99%	22478
		General Population	353	99%	34954
Usher Syndrome, Type 1F	<i>PCDH15</i>	Ashkenazi Jewish	40 - 126	99%	3961
Usher Syndrome, Type 2A	<i>USH2A</i>	Bukharan Jewish	NA	99%	Reduced
		General Population	158	95%	3180
		Sephardic Jewish - Iraqi, Iranian	36	99%	3565
Usher Syndrome, Type 3	<i>CLRN1</i>	Ashkenazi Jewish	120	99%	11883
		Finnish	70	99%	6932
Wilson Disease	<i>ATP7B</i>	Ashkenazi Jewish	67	99%	6635
		Canary Islands	25	99%	2476
		General Population	90	99%	8912
		Sardinian	42	99%	4159
		Sephardic Jewish - North African, Iraqi, Yemenite, Iranian, Bukharian	65	99%	6437



Condition	Gene	Population	Carrier frequency before screening (1 in)	Detection rate	Carrier residual risk after negative result (1 in)
Wolman Disease	LIPA	General Population	*500	99%	Reduced
		Sephardic Jewish - Iranian	32	99%	3169
Zellweger Spectrum Disorders (PEX6-Related)	PEX6	French Canadian	55	99%	5447
		General Population	280	99%	27726
Zellweger Spectrum Disorders (PEX10-related)	PEX10	General Population	*500	97%	Reduced

\* <1 in 500