

Horizon Conditions List



Horizon™
Advanced carrier screening

CONDITION	GENE	AUTOSOMAL RECESSIVE	X-LINKED	SCREENING RECOMMENDATIONS		PANEL AVAILABILITY			
				ACOG†	ACMG	H445	H569‡	H574‡	H613
17-Beta Hydroxysteroid Dehydrogenase 3 Deficiency	<i>HSD17B3</i>	•				•	•	•	•
3-Beta-Hydroxysteroid Dehydrogenase Type II Deficiency	<i>HSD3B2</i>	•				•	•	•	•
3-Hydroxy-3-Methylglutaryl-Coenzyme A Lyase Deficiency	<i>HMGCL</i>	•				•	•	•	•
3-Hydroxyacyl-CoA Dehydrogenase Deficiency	<i>HADH</i>	•				•	•	•	•
3-Methylcrotonyl-CoA Carboxylase 1 Deficiency	<i>MCCC1*</i>	•				•	•	•	•
3-Methylcrotonyl-CoA Carboxylase 2 Deficiency	<i>MCCC2*</i>	•		○		•	•	•	•
3-Phosphoglycerate Dehydrogenase Deficiency	<i>PHGDH</i>	•				•	•	•	•
5-Alpha-Reductase Deficiency	<i>SRD5A2</i>	•					•	•	•
6-Pyruvoyl-Tetrahydropterin Synthase (PTPS) Deficiency	<i>PTS</i>	•				•	•	•	•
ABCA4-Related Conditions	<i>ABCA4</i>	•					•	•	•
Abetalipoproteinemia	<i>MTTP</i>	•				•	•	•	•
Achondrogenesis, Type 1B	<i>SLC26A2</i>	•		○		•	•	•	•
Achromatopsia, CNGB3-Related	<i>CNGB3</i>	•		○		•	•	•	•
Acrodermatitis Enteropathica	<i>SLC39A4</i>	•				•	•	•	•
Action Myoclonus-Renal Failure (AMRF) Syndrome	<i>SCARB2</i>	•				•			•
Acute Infantile Liver Failure, TRMU-Related	<i>TRMU</i>	•				•	•	•	•
Acyl-CoA Oxidase 1 Deficiency	<i>ACOX1</i>	•				•	•	•	•
Adrenal Hypoplasia Congenita, X-Linked	<i>NR0B1</i>		•	○		•	•	•	•
Adrenoleukodystrophy, X-Linked	<i>ABCD1</i>		•	○		•	•	•	•
Agammaglobulinemia, X-Linked	<i>BTK</i>		•			•	•	•	•
Aicardi-Goutières Syndrome	<i>SAMHD1</i>	•				•	•	•	•
Aicardi-Goutières Syndrome, RNASEH2A-Related	<i>RNASEH2A</i>	•				•	•	•	•
Aicardi-Goutières Syndrome, RNASEH2B-Related	<i>RNASEH2B</i>	•		○		•	•	•	•
Aicardi-Goutières Syndrome, RNASEH2C-Related	<i>RNASEH2C</i>	•				•	•	•	•
Aicardi-Goutières Syndrome, TREX1-Related	<i>TREX1</i>	•				•	•	•	•
Alkaptonuria	<i>HGD*</i>	•				•	•	•	•
Alpha-1 Antitrypsin Deficiency	<i>SERPINA1*</i>	•				•	•	•	•
Alpha-Mannosidosis	<i>MAN2B1</i>	•				•	•	•	•
Alpha-Thalassemia	<i>HBA1/HBA2</i>	•		○	○	•	•	•	•
Alpha-Thalassemia Intellectual Disability Syndrome	<i>ATRX</i>		•			•	•	•	•
Alport Syndrome, COL4A3-Related	<i>COL4A3</i>	•				•	•	•	•
Alport Syndrome, COL4A4-Related	<i>COL4A4</i>	•				•	•	•	•
Alport Syndrome, X-Linked	<i>COL4A5</i>		•			•	•	•	•
Alstrom Syndrome	<i>ALMS1</i>	•				•	•	•	•
Amish Infantile Epilepsy Syndrome	<i>ST3GAL5</i>	•				•	•	•	•
Andermann Syndrome	<i>SLC12A6</i>	•				•	•	•	•
Androgen Insensitivity Syndrome	<i>AR</i>		•				•	•	•
Arginine:Glycine Amidinotransferase Deficiency (Agat Deficiency)	<i>GATM</i>	•					•	•	•
Argininemia	<i>ARG1</i>	•				•	•	•	•
Argininosuccinate Lyase Deficiency	<i>ASL</i>	•		○		•	•	•	•
Aromatase Deficiency	<i>CYP19A1</i>	•				•	•	•	•
Arts Syndrome	<i>PRPS1</i>		•			•	•	•	•
Asparagine Synthetase Deficiency	<i>ASNS</i>	•				•	•	•	•
Aspartylglycosaminuria	<i>AGA</i>	•		○		•	•	•	•
Ataxia with Vitamin E Deficiency	<i>TTPA</i>	•				•	•	•	•
Ataxia-Telangiectasia	<i>ATM</i>	•				•	•	•	•
Ataxia-Telangiectasia-Like Disorder 1	<i>MRE11</i>	•				•	•	•	•
Atransferrinemia	<i>TF</i>			○		•	•	•	•
Autism Spectrum, Epilepsy and Arthrogryposis	<i>SLC35A3</i>	•				•	•	•	•
Autoimmune Polyglandular Syndrome, Type 1	<i>AIRE</i>	•		○		•	•	•	•
Autosomal Recessive Congenital Ichthyosis (ARCI), SLC27A4-Related	<i>SLC27A4</i>	•					•	•	•
Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay	<i>SACS</i>	•				•	•	•	•
Bardet-Biedl Syndrome, ARL6-Related	<i>ARL6</i>	•					•	•	•
Bardet-Biedl Syndrome, BBS10-Related	<i>BBS10</i>	•				•	•	•	•
Bardet-Biedl Syndrome, BBS12-Related	<i>BBS12</i>	•				•	•	•	•
Bardet-Biedl Syndrome, BBS1-Related	<i>BBS1</i>	•		○		•	•	•	•
Bardet-Biedl Syndrome, BBS2-Related	<i>BBS2</i>	•		○		•	•	•	•
Bardet-Biedl Syndrome, BBS4-Related	<i>BBS4</i>	•				•	•	•	•
Bardet-Biedl Syndrome, Bbs5-Related	<i>BBS5</i>	•					•	•	•
Bardet-Biedl Syndrome, BBS7-Related	<i>BBS7</i>	•				•	•	•	•
Bardet-Biedl Syndrome, BBS9-Related	<i>BBS9</i>	•				•	•	•	•
Bardet-Biedl Syndrome, TTC8-Related	<i>TTC8</i>	•					•		
Bare Lymphocyte Syndrome, CIITA-Related	<i>CIITA</i>	•				•	•	•	•
Barth Syndrome	<i>TAZ</i>		•			•	•	•	•
Bartter Syndrome, BSND-Related	<i>BSND</i>	•				•	•	•	•
Bartter Syndrome, KCNJ1-Related	<i>KCNJ1</i>	•				•	•	•	•

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				ACOG [†]	ACMG	H445	H569 [‡]	H574 [‡]	H613
Barter Syndrome, SLC12A1-Related	<i>SLC12A1</i>	•				•	•	•	•
Batten Disease, CLN3-Related	<i>CLN3</i>	•				•	•	•	•
Bernard-Soulier Syndrome, Type A1/A2	<i>GP1BA*</i>	•				•	•	•	•
Bernard-Soulier Syndrome, Type C	<i>GP9*</i>	•				•	•	•	•
Beta-Hemoglobinopathies	<i>HBB</i>	•	○	○		•	•	•	•
Beta-Mannosidosis	<i>MANBA</i>	•					•	•	•
Beta-Ureidopropionase Deficiency	<i>UPB1</i>	•				•			•
Bilateral Frontal Parietal Polymicrogyria	<i>GPR56</i>	•				•	•	•	•
Biotinidase Deficiency	<i>BTD*</i>	•		○		•	•	•	•
Biotin-Thiamine-Responsive Basal Ganglia Disease (BTBGD)	<i>SLC19A3</i>			○		•	•	•	•
Bloom Syndrome	<i>BLM</i>	•	○	○		•	•	•	•
Brittle Cornea Syndrome 1	<i>ZNF469</i>	•					•	•	•
Brittle Cornea Syndrome 2	<i>PRDM5</i>	•					•	•	•
Canavan Disease	<i>ASPA</i>	•	○	○		•	•	•	•
Carbamoyl Phosphate Synthetase I Deficiency	<i>CPS1</i>	•				•	•	•	•
Carnitine Deficiency	<i>SLC22A5</i>	•				•	•	•	•
Carnitine Palmitoyltransferase IA Deficiency	<i>CPT1A</i>	•				•	•	•	•
Carnitine Palmitoyltransferase II Deficiency	<i>CPT2</i>	•		○		•	•	•	•
Carnitine-Acylcarnitine Translocase Deficiency	<i>SLC25A20</i>	•				•	•	•	•
Carpenter Syndrome	<i>RAB23</i>	•				•	•	•	•
Cartilage-Hair Hypoplasia	<i>RMRP</i>	•				•	•	•	•
Catecholaminergic Polymorphic Ventricular Tachycardia	<i>CASQ2</i>	•				•	•	•	•
CD59-Mediated Hemolytic Anemia	<i>CD59</i>	•					•	•	•
CEP152-Related Microcephaly	<i>CEP152</i>	•					•	•	•
Cerebral Dysgenesis, Neuropathy, Ichthyosis, And Palmoplantar Keratoderma (CEDNIK) Syndrome	<i>SNAP29</i>	•					•	•	•
Cerebrotendinous Xanthomatosis	<i>CYP27A1</i>	•		○		•	•	•	•
Charcot-Marie-Tooth Disease with Deafness, X-Linked	<i>GJB1</i>		•			•	•	•	•
Charcot-Marie-Tooth Disease, Recessive Intermediate C	<i>PLEKHG5</i>	•					•	•	•
Charcot-Marie-Tooth Disease, Type 4D	<i>NDRG1</i>	•				•	•	•	•
Chediak-Higashi Syndrome	<i>LYST</i>	•				•	•	•	•
Choreoacanthocytosis	<i>VPS13A</i>	•				•	•	•	•
Choroideremia	<i>CHM</i>		•			•	•	•	•
Chronic Granulomatous Disease, CYBA-Related	<i>CYBA</i>	•				•	•	•	•
Chronic Granulomatous Disease, NCF2-Related	<i>NCF2</i>	•					•	•	•
Chronic Granulomatous Disease, X-Linked	<i>CYBB</i>		•			•	•	•	•
Ciliopathies, RPGRIP1L-Related	<i>RPGRIP1L</i>	•				•	•	•	•
Citrin Deficiency	<i>SLC25A13</i>	•				•	•	•	•
Citrullinemia, Type 1	<i>ASS1</i>	•				•	•	•	•
CLN10 Disease	<i>CTSD</i>	•				•	•	•	•
Cohen Syndrome	<i>VPS13B</i>	•				•	•	•	•
COL11A2-Related Conditions	<i>COL11A2</i>	•					•	•	•
Combined Malonic and Methylmalonic Aciduria	<i>ACSF3</i>	•				•	•	•	•
Combined Oxidative Phosphorylation Deficiency 1	<i>GFM1</i>	•				•	•	•	•
Combined Oxidative Phosphorylation Deficiency 3	<i>TSFM</i>	•				•	•	•	•
Combined Pituitary Hormone Deficiency 1	<i>POU1F1</i>	•					•	•	•
Combined Pituitary Hormone Deficiency-2	<i>PROP1</i>	•				•	•	•	•
Congenital Adrenal Hyperplasia, 11-Beta-Hydroxylase Deficiency	<i>CYP11B1</i>	•				•	•	•	•
Congenital Adrenal Hyperplasia, 17-Alpha-Hydroxylase Deficiency	<i>CYP17A1</i>	•				•	•	•	•
Congenital Adrenal Hyperplasia, 21-Hydroxylase Deficiency	<i>CYP21A2</i>	•		○		•	•	•	•
Congenital Adrenal Insufficiency, CYP11A1-Related	<i>CYP11A1</i>			○		•	•	•	•
Congenital Amegakaryocytic Thrombocytopenia	<i>MPL</i>	•				•	•	•	•
Congenital Chronic Diarrhea	<i>DGAT1</i>	•					•	•	•
Congenital Disorder Of Glycosylation Type 1, ALG1-Related	<i>ALG1</i>	•					•	•	•
Congenital Disorder of Glycosylation, Type 1A, PMM2-Related	<i>PMM2</i>	•		○		•	•	•	•
Congenital Disorder of Glycosylation, Type 1B	<i>MPI</i>	•				•	•	•	•
Congenital Disorder of Glycosylation, Type 1C	<i>ALG6</i>	•				•	•	•	•
Congenital Dyserythropoietic Anemia Type 2	<i>SEC23B</i>	•					•	•	•
Congenital Finnish Nephrosis	<i>NPHS1</i>	•		○		•	•	•	•
Congenital Hydrocephalus 1	<i>CCDC88C</i>			○		•	•	•	•
Congenital Hyperinsulinism, KCNJ11-Related	<i>KCNJ11</i>	•				•	•	•	•
Congenital Insensitivity to Pain with Anhidrosis (CIPA)	<i>NTRK1</i>	•				•	•	•	•
Congenital Myasthenic Syndrome, CHAT-Related	<i>CHAT</i>	•				•	•	•	•
Congenital Myasthenic Syndrome, CHRNE-Related	<i>CHRNE</i>	•		○		•	•	•	•
Congenital Myasthenic Syndrome, COLQ-Related	<i>COLQ</i>	•				•			•
Congenital Myasthenic Syndrome, DOK7-Related	<i>DOK7</i>	•				•	•	•	•
Congenital Myasthenic Syndrome, RAPSN-Related	<i>RAPSN</i>	•				•	•	•	•
Congenital Nephrotic Syndrome, PLCE1-Related	<i>PLCE1</i>	•				•			•
Congenital Neutropenia, G6PC3-Related	<i>G6PC3</i>	•					•	•	•
Congenital Neutropenia, HAX1-Related	<i>HAX1</i>	•				•	•	•	•
Congenital Neutropenia, VPS45-Related	<i>VPS45</i>	•				•	•	•	•

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				ACOG [†]	ACMG	H445	H569 [‡]	H574 [‡]	H613
Congenital Secretory Chloride Diarrhea 1	<i>SLC26A3</i>	•				•	•	•	•
Corneal Dystrophy and Perceptive Deafness	<i>SLC4A11</i>	•				•	•	•	•
Corticosterone Methyloxidase Deficiency	<i>CYP11B2</i>	•				•	•	•	•
Costeff Syndrome (3-Methylglutaconic Aciduria, Type 3)	<i>OPA3</i>	•				•	•	•	•
Cowchock Syndrome	<i>AIFM1</i>		•			•			•
CRB1-Related Retinal Dystrophies	<i>CRB1</i>	•				•	•	•	•
Creatine Transporter Defect (Cerebral Creatine Deficiency Syndrome 1, X-Linked)	<i>SLC6A8</i>		•	○	○	•	•	•	•
Cystic Fibrosis	<i>CFTR</i>	•		○	○	•	•	•	•
Cystinosis	<i>CTNS</i>	•				•	•	•	•
Cytochrome C Oxidase Deficiency, PET100-Related	<i>PET100</i>	•				•	•	•	•
Cytochrome P450 Oxioreductase Deficiency	<i>POR</i>	•					•	•	•
D-Bifunctional Protein Deficiency	<i>HSD17B4</i>	•				•	•	•	•
Deafness, Autosomal Recessive 77	<i>LOXHD1</i>	•				•	•	•	•
Dent Disease, Type 1	<i>CLCN5</i>		•			•			•
Dent Disease, Type 2 / Lowe Syndrome	<i>OCRL</i>		•			•	•	•	•
Developmental And Epileptic Encephalopathy 36	<i>ALG13</i>		•				•	•	•
Dihydropteridine Reductase (DHPR) Deficiency	<i>QDPR</i>	•					•	•	•
Dihydropyrimidine Dehydrogenase Deficiency	<i>DPYD</i>	•				•			•
Donnai-Barrow Syndrome	<i>LRP2</i>			○	•	•	•	•	•
Dubin-Johnson Syndrome	<i>ABCC2</i>	•					•	•	•
Duchenne/Becker Muscular Dystrophy	<i>DMD</i>		•	○	○	•	•	•	•
Dyskeratosis Congenita Spectrum Disorders	<i>TERT</i>	•					•	•	•
Dyskeratosis Congenita, DKC1-Related	<i>DKC1</i>		•			•	•	•	•
Dyskeratosis Congenita, RTEL1-Related	<i>RTEL1</i>	•				•	•	•	•
Dystrophic Epidermolysis Bullosa, COL7A1-Related	<i>COL7A1</i>	•		○	○	•	•	•	•
Early Infantile Epileptic Encephalopathy, CAD-Related	<i>CAD</i>	•					•	•	•
Ehlers-Danlos Syndrome Type VI	<i>PLOD1</i>	•					•	•	•
Ehlers-Danlos Syndrome, Classic-Like, TNXB-Related	<i>TNXB</i>			○	•	•	•	•	•
Ehlers-Danlos Syndrome, Type VIIc	<i>ADAMTS2</i>	•				•	•	•	•
Ellis-van Creveld Syndrome, EVC2-Related	<i>EVC2</i>	•		○	○	•	•	•	•
Ellis-van Creveld Syndrome, EVC-Related	<i>EVC</i>	•				•	•	•	•
Emery-Dreifuss Muscular Dystrophy 1, X-Linked	<i>EMD</i>		•			•	•	•	•
Emery-Dreifuss Muscular Dystrophy 6, X Linked	<i>FHL1</i>		•				•	•	•
Enhanced S-Cone Syndrome	<i>NR2E3</i>	•				•	•	•	•
Epimerase Deficiency (Galactosemia Type III)	<i>GALE</i>	•					•	•	•
Epiphyseal Dysplasia, Multiple, 7 / Desbuquois Dysplasia 1	<i>CANT1</i>	•				•	•	•	•
ERCC6-Related Disorders	<i>ERCC6</i>	•				•	•	•	•
ERCC8-Related Disorders	<i>ERCC8</i>	•				•	•	•	•
Ethylmalonic Encephalopathy	<i>ETHE1</i>	•				•	•	•	•
Fabry Disease	<i>GLA</i>		•	○	○	•	•	•	•
Factor IX Deficiency	<i>F9</i>		•	○	○	•	•	•	•
Factor V Deficiency	<i>F5*</i>	•					•	•	•
Factor XI Deficiency	<i>F11*</i>	•				•	•	•	•
Familial Dysautonomia	<i>IKBKA</i>	•		○	○	•	•	•	•
Familial Hemophagocytic Lymphohistiocytosis, PRF1-Related	<i>PRF1</i>	•		○	○	•	•	•	•
Familial Hemophagocytic Lymphohistiocytosis, STX11-Related	<i>STX11</i>	•				•	•	•	•
Familial Hemophagocytic Lymphohistiocytosis, STXBP2-Related	<i>STXBP2</i>	•				•	•	•	•
Familial Hemophagocytic Lymphohistiocytosis, UNC13D-Related	<i>UNC13D</i>	•					•	•	•
Familial Hypercholesterolemia, LDLRAP1-Related	<i>LDLRAP1</i>	•				•	•	•	•
Familial Hypercholesterolemia, LDLR-Related	<i>LDLR</i>	•				•	•	•	•
Familial Hyperinsulinism, ABCC8-Related	<i>ABCC8</i>	•		○	○	•	•	•	•
Familial Mediterranean Fever	<i>MEFV*</i>	•				•	•	•	•
Familial Nephrogenic Diabetes Insipidus, AQP2-Related	<i>AQP2</i>	•				•	•	•	•
Fanconi Anemia Group J	<i>BRIP1</i>	•					•	•	•
Fanconi Anemia, Group A	<i>FANCA</i>	•		○		•	•	•	•
Fanconi Anemia, Group B	<i>FANCB</i>		•			•	•	•	•
Fanconi Anemia, Group C	<i>FANCC</i>	•		○	○	•	•	•	•
Fanconi Anemia, Group D2	<i>FANCD2</i>	•				•	•	•	•
Fanconi Anemia, Group E	<i>FANCE</i>	•				•	•	•	•
Fanconi Anemia, Group F	<i>FANCF</i>	•				•			•
Fanconi Anemia, Group G	<i>FANCG</i>	•		○		•	•	•	•
Fanconi Anemia, Group I	<i>FANCI</i>	•				•	•	•	•
Fanconi Anemia, Group L	<i>FANCL</i>	•				•	•	•	•
Farber Lipogranulomatosis	<i>ASAHI</i>	•				•			•
Foveal Hypoplasia	<i>SLC38A8</i>	•					•	•	•
Fragile X Syndrome	<i>FMR1</i>		•	○	○	•	•	•	•
Fragile XE Syndrome	<i>AFF2</i>		•		○	•	•	•	•
Fraser Syndrome 3, GRIP1-Related	<i>GRIP1</i>				○	•	•	•	•
Fraser Syndrome, FRAS1-Related	<i>FRAS1</i>	•					•	•	•
Fraser Syndrome, FREM2-Related	<i>FREM2</i>	•					•	•	•

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Friedreich Ataxia	<i>FXN</i>				o	•		•	•
Fructose-1,6-Bisphosphatase Deficiency	<i>FBP1</i>	•					•	•	•
Fucosidosis	<i>FUCA1</i>	•					•	•	•
Fumarase Deficiency	<i>FH</i>	•				•	•	•	•
GABA-Transaminase Deficiency	<i>ABAT</i>	•				•			•
Galactokinase Deficiency (Galactosemia, Type II)	<i>GALK1</i>	•				•	•	•	•
Galactosemia	<i>GALT</i>	•			o	•	•	•	•
Galactosialidosis	<i>CTSA</i>	•				•	•	•	•
Gaucher Disease	<i>GBA</i>	•			o	•	•	•	•
GCH1-Related Conditions	<i>GCH1</i>	•					•	•	•
GDF5-Related Conditions	<i>GDF5</i>	•					•	•	•
Geroderma Osteodysplastica	<i>GORAB</i>	•					•	•	•
Gitelman Syndrome	<i>SLC12A3</i>	•				•	•	•	•
Glanzmann Thrombasthenia	<i>ITGB3</i>	•					•	•	•
Glucose-6-Phosphate Dehydrogenase Deficiency	<i>G6PD*</i>		•			•	•	•	•
Glutaric Acidemia, Type 1	<i>GCDH</i>	•				•	•	•	•
Glutaric Acidemia, Type 2A	<i>ETFA</i>	•				•	•	•	•
Glutaric Acidemia, Type 2B	<i>ETFB</i>	•				•	•	•	•
Glutaric Acidemia, Type 2C	<i>ETFDH</i>	•				•	•	•	•
Glutathione Synthetase Deficiency	<i>GSS</i>	•					•	•	•
Glycine Encephalopathy, AMT-Related	<i>AMT</i>	•				•	•	•	•
Glycine Encephalopathy, GLDC-Related	<i>GLDC</i>	•				•	•	•	•
Glycogen Storage Disease Type IXb	<i>PHKB</i>	•					•	•	•
Glycogen Storage Disease Type IXc	<i>PHKG2</i>	•					•	•	•
Glycogen Storage Disease, Type 1A	<i>G6PC</i>	•		o	o	•	•	•	•
Glycogen Storage Disease, Type 1B	<i>SLC37A4</i>	•		o	o	•	•	•	•
Glycogen Storage Disease, Type 2 (Pompe Disease)	<i>GAA</i>	•			o	•	•	•	•
Glycogen Storage Disease, Type 3	<i>AGL</i>	•				•	•	•	•
Glycogen Storage Disease, Type 4	<i>GBE1</i>	•			o	•	•	•	•
Glycogen Storage Disease, Type 5 (McArdle Disease)	<i>PYGM</i>	•				•	•	•	•
Glycogen Storage Disease, Type 7	<i>PFKM</i>	•				•	•	•	•
GRACILE Syndrome	<i>BCS1L</i>	•				•	•	•	•
Guanidinoacetate Methyltransferase Deficiency	<i>GAMT</i>	•				•	•	•	•
Harlequin Ichthyosis	<i>ABCA12</i>	•				•	•	•	•
Heme Oxygenase 1 Deficiency	<i>HMOX1</i>	•					•	•	•
Hemochromatosis, Type 2A	<i>HFE2</i>	•				•	•	•	•
Hemochromatosis, Type 3, TFR2-Related	<i>TFR2</i>	•				•	•	•	•
Hemophilia A	<i>F8</i>		•		o	•	•	•	•
Hepatocerebral Mitochondrial DNA Depletion Syndrome, MPV17-Related	<i>MPV17</i>	•				•	•	•	•
Hereditary Fructose Intolerance	<i>ALDOB</i>	•			o	•	•	•	•
Hereditary Hemochromatosis Type 1	<i>HFE*</i>	•					•	•	•
Hereditary Hemochromatosis Type 2B	<i>HAMP</i>	•					•	•	•
Hereditary Spastic Paraparesis, Type 49	<i>TECPR2</i>	•				•	•	•	•
Hereditary Spastic Paraplegia, CYP7B1-Related	<i>CYP7B1</i>	•					•	•	•
Hermansky-Pudlak Syndrome, AP3B1-Related	<i>AP3B1</i>	•				•			•
Hermansky-Pudlak Syndrome, BLOC1S3-Related	<i>BLOC1S3</i>	•					•	•	•
Hermansky-Pudlak Syndrome, BLOC1S6-Related	<i>BLOC1S6</i>	•					•	•	•
Hermansky-Pudlak Syndrome, HPS1-Related	<i>HPS1</i>	•			o	•	•	•	•
Hermansky-Pudlak Syndrome, HPS3-Related	<i>HPS3</i>	•			o	•	•	•	•
Hermansky-Pudlak Syndrome, HPS4-Related	<i>HPS4</i>	•				•	•	•	•
Hermansky-Pudlak Syndrome, HPS5-Related	<i>HPS5</i>	•					•	•	•
Hermansky-Pudlak Syndrome, HPS6-Related	<i>HPS6</i>	•					•	•	•
Heterotaxy Syndrome, ZIC3-Related	<i>ZIC3</i>		•				•		•
Holocarboxylase Synthetase Deficiency	<i>HLCS</i>	•				•	•	•	•
Homocystinuria And Megaloblastic Anemia Type Cblg	<i>MTR</i>	•					•	•	•
Homocystinuria due to Deficiency of MTHFR	<i>MTHFR</i>	•				•	•	•	•
Homocystinuria, CBS-Related	<i>CBS</i>	•			o	•	•	•	•
Homocystinuria, Type cbfE	<i>MTRR</i>	•				•	•	•	•
HSD10 Disease	<i>HSD17B10</i>		•				•	•	•
Hydrocephalus Syndrome	<i>HYLS1</i>	•				•	•	•	•
Hyper IgM Syndrome, X-Linked	<i>CD40LG</i>		•			•	•	•	•
Hyper-Igm Immunodeficiency	<i>CD40</i>	•					•	•	•
Hyperornithinemia-Hyperammonemia-Homocitrullinuria (HHH Syndrome)	<i>SLC25A15</i>	•					•	•	•
Hyperphosphatemic Familial Tumoral Calcinosis, GALNT3-Related	<i>GALNT3</i>	•					•	•	•
Hypohidrotic Ectodermal Dysplasia, X-Linked	<i>EDA</i>		•				•	•	•
Hypomyelinating Leukodystrophy 12	<i>VPS11</i>	•					•	•	•
Hypophosphatasia, ALPL-Related	<i>ALPL</i>	•			o	•	•	•	•
Imerslund-Gräsbeck Syndrome 2	<i>AMN</i>	•					•	•	•
Immune Dysregulation, Polyendocrinopathy, Enteropathy, X-Linked (IPEX) Syndrome	<i>FOXP3</i>		•				•		•
Immunodeficiency-Centromeric Instability-Facial Anomalies (ICF) Syndrome, DNMT3B-Related	<i>DNMT3B</i>	•					•	•	•

CONDITION	GENE	AUTOSOMAL RECESSIVE	X-LINKED	SCREENING RECOMMENDATIONS		PANEL AVAILABILITY			
				ACOG [†]	ACMG	H445	H569 [‡]	H574 [‡]	H613
Immunodeficiency-Centromeric Instability-Facial Anomalies (ICF) Syndrome, ZBTB24-Related	ZBTB24	•				•	•	•	•
Inclusion Body Myopathy 2	GNE	•				•	•	•	•
Infantile Cerebral and Cerebellar Atrophy	MED17	•				•	•	•	•
Infantile Nephronophthisis	INVS	•				•	•	•	•
Infantile Neuroaxonal Dystrophy	PLA2G6	•				•	•	•	•
Infantile Spinal Muscular Atrophy, X-Linked	UBA1		•			•			•
Isolated Ectopia Lentis	ADAMTSL4	•					•	•	•
Isolated Lissencephaly Sequence / Subcortical Band Heterotopia	DCX		•			•			•
Isolated Sulfite Oxidase Deficiency	SUOX	•					•	•	•
Isolated Thyroid-Stimulating Hormone Deficiency	TSHB	•					•	•	•
Isovaleric Acidemia	IVD	•				•	•	•	•
Johanson-Blizzard Syndrome	UBR1	•				•	•	•	•
Joubert Syndrome 2 / Meckel Syndrome 2	TMEM216	•		○	○	•	•	•	•
Joubert Syndrome And Related Disorders (JSRD), TMEM67-Related	TMEM67	•					•	•	•
Joubert Syndrome, AHI1-Related	AHI1	•		○	○	•	•	•	•
Joubert Syndrome, ARL13B-Related	ARL13B	•		○		•			•
Joubert Syndrome, B9D1-Related	B9D1	•		○		•			•
Joubert Syndrome, B9D2-Related	B9D2	•		○		•			•
Joubert Syndrome, C2CD3-Related / Orofaciodigital Syndrome 14	C2CD3	•		○		•			•
Joubert Syndrome, CC2D2A-Related / COACH Syndrome	CC2D2A	•		○	○	•	•	•	•
Joubert Syndrome, CEP104-Related	CEP104	•		○		•			•
Joubert Syndrome, CEP120-Related / Short-Rib Thoracic Dysplasia 13 with or without Polydactyly	CEP120	•		○		•			•
Joubert Syndrome, CEP41-Related	CEP41	•		○		•			•
Joubert Syndrome, CPLANE1-Related / Orofaciodigital Syndrome 6	CPLANE1	•		○		•			•
Joubert Syndrome, CSPP1-Related	CSPP1	•		○		•			•
Joubert Syndrome, INPP5E-Related	INPP5E	•		○		•			•
Junctional Epidermolysis Bullosa, COL17A1-Related	COL17A1	•					•	•	•
Junctional Epidermolysis Bullosa, ITGA6-Related	ITGA6	•					•	•	•
Junctional Epidermolysis Bullosa, ITGB4-Related	ITGB4	•					•	•	•
Junctional Epidermolysis Bullosa, LAMA3-Related	LAMA3	•					•	•	•
Junctional Epidermolysis Bullosa, LAMB3-Related	LAMB3	•					•	•	•
Junctional Epidermolysis Bullosa, LAMC2-Related	LAMC2	•					•	•	•
Juvenile Retinoschisis, X-Linked	RS1		•		○	•	•	•	•
Ketothiolase Deficiency	ACAT1	•			○	•	•	•	•
Krabbe Disease	GALC	•				•	•	•	•
L1 Syndrome	L1CAM		•		○	•	•	•	•
Lamellar Ichthyosis, Type 1	TGM1	•				•	•	•	•
Laron Syndrome	GHR	•					•	•	•
Leber Congenital Amaurosis 2	RPE65	•				•	•	•	•
Leber Congenital Amaurosis Type AIPL1	AIPL1	•					•	•	•
Leber Congenital Amaurosis Type LRAT	LRAT	•					•	•	•
Leber Congenital Amaurosis Type TULP1	TULP1	•					•	•	•
Leber Congenital Amaurosis, IQCB1-Related / Senior-Loken Syndrome 5	IQCB1	•					•		•
Leber Congenital Amaurosis, Type CEP290	CEP290	•			○	•	•	•	•
Leber Congenital Amaurosis, Type LCA5	LCA5	•				•	•	•	•
Leber Congenital Amaurosis, Type RDH12	RDH12	•				•	•	•	•
Leber Congenital Amaurosis Type GUCY2D	GUCY2D	•					•	•	•
Leigh Syndrome, French-Canadian Type	LRPPRC	•					•	•	•
Lesch-Nyhan Syndrome	HPRT1		•			•	•	•	•
Lethal Congenital Contracture Syndrome 1	GLE1	•				•	•	•	•
Leukoencephalopathy with Vanishing White Matter	EIF2B5	•				•	•	•	•
Leukoencephalopathy With Vanishing White Matter, EIF2B1-Related	EIF2B1	•					•	•	•
Leukoencephalopathy With Vanishing White Matter, EIF2B2-Related	EIF2B2	•					•	•	•
Leukoencephalopathy With Vanishing White Matter, EIF2B3-Related	EIF2B3	•					•	•	•
Leukoencephalopathy With Vanishing White Matter, EIF2B4-Related	EIF2B4	•					•	•	•
LIG4 Syndrome	LIG4	•					•	•	•
Limb-Girdle Muscular Dystrophy Type 8	TRIM32	•					•	•	•
Limb-Girdle Muscular Dystrophy, Type 2A	CAPN3	•					•	•	•
Limb-Girdle Muscular Dystrophy, Type 2B	DYSF	•					•	•	•
Limb-Girdle Muscular Dystrophy, Type 2C	SGCG	•					•	•	•
Limb-Girdle Muscular Dystrophy, Type 2D	SGCA	•					•	•	•
Limb-Girdle Muscular Dystrophy, Type 2E	SGCB	•					•	•	•
Limb-Girdle Muscular Dystrophy, Type 2F	SGCD	•					•	•	•
Limb-Girdle Muscular Dystrophy, Type 2I	FKRP	•			○	•	•	•	•
Lipoamide Dehydrogenase Deficiency (Dihydrolipoamide Dehydrogenase Deficiency)	DLD	•			○	•	•	•	•
Lipoid Adrenal Hyperplasia	STAR	•					•	•	•
Lipoprotein Lipase Deficiency	LPL	•					•	•	•
Long Chain 3-Hydroxyacyl-CoA Dehydrogenase Deficiency	HADHA	•					•	•	•
Lung Disease, Immunodeficiency, And Chromosome Breakage Syndrome (LICS)	NSMCE3	•					•	•	•
Lysinuric Protein Intolerance	SLC7A7	•					•	•	•

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				ACOG [†]	ACMG	H445	H569 [‡]	H574 [‡]	H613
Malonyl-CoA Decarboxylase Deficiency	MLYCD	•				•	•	•	•
Maple Syrup Urine Disease, Type 1A	BCKDHA	•		○		•	•	•	•
Maple Syrup Urine Disease, Type 1B	BCKDHB	•		○	○	•	•	•	•
Maple Syrup Urine Disease, Type 2	DBT	•		○		•	•	•	•
McKusick-Kaufman Syndrome	MKKS	•				•	•	•	•
Meckel Syndrome 7 / Nephronophthisis 3	NPHP3	•				•			
Meckel-Gruber Syndrome, Type 1	MKS1	•				•	•	•	•
MECP2-Related Conditions	MECP2		•				•	•	•
MECR-Related Neurologic Disorder	MECR	•					•	•	•
Medium Chain Acyl-CoA Dehydrogenase Deficiency	ACADM	•		○	○	•	•	•	•
MEDNIK Syndrome	AP1S1	•				•	•	•	•
Megalencephalic Leukoencephalopathy with Subcortical Cysts	MLC1	•			○	•	•	•	•
Menkes Syndrome	ATP7A		•			•	•	•	•
Merosin-Deficient Muscular Dystrophy	LAMA2	•				•	•	•	•
Metabolic Encephalopathy and Arrhythmias, TANGO2-Related	TANGO2	•				•	•	•	•
Metachromatic Leukodystrophy, ARSA-Related	ARSA	•			○	•	•	•	•
Metachromatic Leukodystrophy, PSAP-Related	PSAP	•				•	•	•	•
Methylmalonic Acidemia And Homocystinuria Type Cblf	LMBRD1	•					•	•	•
Methylmalonic Acidemia And Homocystinuria Type Cblk	HCFC1		•				•	•	•
Methylmalonic Aciduria and Homocystinuria, Type cblC	MMACHC	•			○	•	•	•	•
Methylmalonic Aciduria and Homocystinuria, Type cbID	MMADHC	•				•	•	•	•
Methylmalonic Aciduria, MCEE-Related	MCEE	•					•	•	•
Methylmalonic Aciduria, MMAA-Related	MMAA	•				•	•	•	•
Methylmalonic Aciduria, MMAB-Related	MMAB	•				•	•	•	•
Methylmalonic Aciduria, Type mut(0)	MUT	•			○	•	•	•	•
Mevalonic Kinase Deficiency	MVK				○	•	•	•	•
Microcephalic Osteodysplastic Primordial Dwarfism Type II	PCNT	•					•	•	•
Microphthalmia/Anophthalmia, VSX2-Related	VSX2	•				•	•	•	•
Mitochondrial Complex 1 Deficiency, ACAD9-Related	ACAD9	•				•	•	•	•
Mitochondrial Complex 1 Deficiency, NDUFAF5-Related	NDUFAF5	•				•	•	•	•
Mitochondrial Complex 1 Deficiency, NDUFS6-Related	NDUFS6	•				•	•	•	•
Mitochondrial complex I Deficiency, Nuclear Type 1	NDUFS4	•				•	•	•	•
Mitochondrial Complex I Deficiency, Nuclear Type 10	NDUFAF2	•					•	•	•
Mitochondrial Complex I Deficiency, Nuclear Type 17	NDUFAF6	•					•		
Mitochondrial Complex I Deficiency, Nuclear Type 19	FOXRED1	•					•	•	•
Mitochondrial Complex I Deficiency, Nuclear Type 3	NDUFS7	•					•	•	•
Mitochondrial Complex I Deficiency, Nuclear Type 4	NDUFV1	•					•	•	•
Mitochondrial Complex IV Deficiency, Nuclear Type 2, SCO2-Related	SCO2				○	•	•	•	•
Mitochondrial Complex IV Deficiency, Nuclear Type 6	COX15	•					•	•	•
Mitochondrial Dna Depletion Syndrome 2	TK2	•					•	•	•
Mitochondrial Dna Depletion Syndrome 3	DGUOK	•					•	•	•
Mitochondrial Myopathy and Sideroblastic Anemia (MLASA1)	PUS1	•					•	•	•
Mitochondrial Trifunctional Protein Deficiency, HADHB-Related	HADHB	•					•	•	•
Molybdenum Cofactor Deficiency Type B	MOCs2	•					•	•	•
Molybdenum Cofactor Deficiency, Type A	MOCs1	•				•	•	•	•
Mucolipidosis II/IIIA	GNPTAB	•			○	•	•	•	•
Mucolipidosis III gamma	GNPTG	•				•	•	•	•
Mucolipidosis, Type IV	MCOLN1	•		○	○	•	•	•	•
Mucopolysaccharidosis, Type I (Hurler Syndrome)	IDUA	•				○	•	•	•
Mucopolysaccharidosis, Type II (Hunter Syndrome)	IDS		•				•	•	•
Mucopolysaccharidosis, Type IIIA (Sanfilippo A)	SGSH	•					•	•	•
Mucopolysaccharidosis, Type IIIB (Sanfilippo B)	NAGLU	•					•	•	•
Mucopolysaccharidosis, Type IIIC (Sanfilippo C)	HGSNAT	•					•	•	•
Mucopolysaccharidosis, Type IID (Sanfilippo D)	GNS	•					•	•	•
Mucopolysaccharidosis, Type IVA (Morquio Syndrome)	GALNS	•					•	•	•
Mucopolysaccharidosis, Type IVB / GM1 Gangliosidosis	GLB1	•					•	•	•
Mucopolysaccharidosis, Type IX	HYAL1	•					•	•	•
Mucopolysaccharidosis, Type VI (Maroteaux-Lamy)	ARSB	•					•	•	•
Mucopolysaccharidosis, Type VII	GUSB	•					•	•	•
Mulibrey Nanism	TRIM37	•					•	•	•
Multiple Pterygium Syndrome, CHRNG-Related / Escobar Syndrome	CHRNG	•					•	•	•
Multiple Sulfatase Deficiency	SUMF1	•					•	•	•
Muscle-Eye-Brain Disease, POMGNT1-Related	POMGNT1	•					•	•	•
Muscular Dystrophy-Dystroglycanopathy	RXYLT1	•					•	•	•
MUSK-Related Congenital Myasthenic Syndrome	MUSK	•					•	•	•
Myoneurogastrointestinal Encephalopathy (MNGIE)	TYMP	•					•	•	•
Myotonia Congenita	CLCN1				○	•	•	•	•
Myotubular Myopathy, X-Linked	MTM1		•			•	•	•	•
N-acetylglutamate Synthase Deficiency	NAGS	•				•	•	•	•
Nemaline Myopathy, NEB-Related	NEB	•			○	•	•	•	•

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				ACOG [†]	ACMG	H445	H569 [‡]	H574 [‡]	H613
Nephrogenic Diabetes Insipidus, AVPR2-Related	AVPR2		•			•	•	•	•
Nephronophthisis 1	NPHP1	•				•	•	•	•
Neuronal Ceroid Lipofuscinosis, CLN5-Related	CLN5	•				•	•	•	•
Neuronal Ceroid Lipofuscinosis, CLN6-Related	CLN6	•				•	•	•	•
Neuronal Ceroid Lipofuscinosis, CLN8-Related	CLN8	•				•	•	•	•
Neuronal Ceroid Lipofuscinosis, MFSD8-Related	MFSD8	•				•	•	•	•
Neuronal Ceroid Lipofuscinosis, PPT1-Related	PPT1	•				•	•	•	•
Neuronal Ceroid Lipofuscinosis, TPP1-Related	TPP1	•				•	•	•	•
Glyc1-Congenital Disorder Of Glycosylation	NGLY1	•					•	•	•
Niemann-Pick Disease, Type C1/D	NPC1	•		o		•	•	•	•
Niemann-Pick Disease, Type C2	NPC2	•		o		•	•	•	•
Niemann-Pick Disease, Types A/B	SMPD1	•		o	o	•	•	•	•
Nijmegen Breakage Syndrome	NBN	•				•	•	•	•
Non-Syndromic Hearing Loss, GJB2-Related	GJB2	•		o		•	•	•	•
Nonsyndromic Hearing Loss, MYO15A-Related	MYO15A	•				•	•	•	•
Nonsyndromic Hearing Loss, OTOA-Related	OTOA	•					•	•	•
Nonsyndromic Hearing Loss, OTOF-Related	OTOF	•					•	•	•
Nonsyndromic Hearing Loss, PJVK-Related	PJVK	•					•	•	•
Nonsyndromic Hearing Loss, SYNE4-Related	SYNE4	•					•	•	•
Nonsyndromic Hearing Loss, TMC1-Related	TMC1	•					•	•	•
Nonsyndromic Hearing Loss, TMPRSS3-Related	TMPRSS3	•					•	•	•
Nonsyndromic Intellectual Disability	CC2D1A	•					•	•	•
Normophosphatemic Tumoral Calcinosis	SAMD9	•					•	•	•
Oculocutaneous Albinism Type III	TYRP1	•					•	•	•
Oculocutaneous Albinism Type IV	SLC45A2	•					•	•	•
Oculocutaneous Albinism, OCA2-Related	OCA2			o		•	•	•	•
Oculocutaneous Albinism, Type 1A and 1B	TYR			o		•	•	•	•
Odonto-Onychodermal Dysplasia / Schopf-Schulz-Passarge Syndrome	WNT10A	•				•	•	•	•
Omenn Syndrome, RAG2-Related	RAG2	•				•	•	•	•
Opitz G/BBB Syndrome, X-Linked	MID1		•		o	•	•	•	•
Ornithine Aminotransferase Deficiency	OAT	•				•	•	•	•
Ornithine Transcarbamylase Deficiency	OTC		•		o	•	•	•	•
Osteogenesis Imperfecta Type VII	CRTAP	•					•	•	•
Osteogenesis Imperfecta Type VIII	P3H1	•					•	•	•
Osteogenesis Imperfecta Type XI	FKBP10	•					•	•	•
Osteogenesis Imperfecta Type XIII	BMP1	•					•	•	•
Osteopetrosis, Infantile Malignant, TCIRG1-Related	TCIRG1	•					•	•	•
Osteopetrosis, OSTM1-Related	OSTM1	•					•	•	•
Pantothenate Kinase-Associated Neurodegeneration	PANK2	•					•	•	•
Papillon Lefèvre Syndrome	CTSC	•					•	•	•
Parkinson Disease 15	FBXO7	•					•	•	•
Pendred Syndrome	SLC26A4	•			o	•	•	•	•
Perlman Syndrome	DIS3L2	•					•		•
Pgm3-Congenital Disorder Of Glycosylation	PGM3	•					•	•	•
Phenylketonuria	PAH	•			o	•	•	•	•
Pign-Congenital Disorder Of Glycosylation	PIGN	•					•	•	•
Pituitary Hormone Deficiency, Combined 3	LHX3	•					•	•	•
PLP1 Disorders	PLP1		•		o	•	•	•	•
POLG-Related Disorders	POLG	•			o	•	•	•	•
Polyzystic Kidney Disease, Autosomal Recessive	PKHD1	•			o	•	•	•	•
Pontocerebellar Hypoplasia, EXOSC3-Related	EXOSC3	•					•	•	•
Pontocerebellar Hypoplasia, RARS2-Related	RARS2	•			o	•	•	•	•
Pontocerebellar Hypoplasia, TSEN2-Related	TSEN2	•					•		
Pontocerebellar Hypoplasia, TSEN54-Related	TSEN54	•					•	•	•
Pontocerebellar Hypoplasia, Type 1A	VRK1	•					•	•	•
Pontocerebellar Hypoplasia, Type 2D	SEPSECS	•					•	•	•
Pontocerebellar Hypoplasia, VPS53-Related	VPS53	•					•	•	•
Primary Ciliary Dyskinesia, CCDC103-Related	CCDC103	•					•	•	•
Primary Ciliary Dyskinesia, CCDC39-Related	CCDC39	•					•	•	•
Primary Ciliary Dyskinesia, DNAH11-Related	DNAH11	•					•	•	•
Primary Ciliary Dyskinesia, DNAH5-Related	DNAH5	•					•	•	•
Primary Ciliary Dyskinesia, DNAI1-Related	DNAI1	•					•	•	•
Primary Ciliary Dyskinesia, DNAI2-Related	DNAI2	•					•	•	•
Primary Congenital Glaucoma / Peters Anomaly	CYP1B1	•					•	•	•
Primary Hyperoxaluria, Type 1	AGXT	•			o	•	•	•	•
Primary Hyperoxaluria, Type 2	GRHPR	•					•	•	•
Primary Hyperoxaluria, Type 3	HOGA1	•					•	•	•
Primary Microcephaly 1, Autosomal Recessive	MCPH1	•			o	•	•	•	•
Progressive Early-Onset Encephalopathy With Brain Atrophy And Thin Corpus Callosum	TBCD	•					•	•	•
Progressive Familial Intrahepatic Cholestasis, ABCB4-Related	ABCB4	•					•	•	•

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				ACOG [†]	ACMG	H445	H569 [‡]	H574 [‡]	H613
Progressive Familial Intrahepatic Cholestasis, Type 1 (PFIC1)	ATP8B1	•				•	•	•	•
Progressive Familial Intrahepatic Cholestasis, Type 2	ABCB11	•				•	•	•	•
Progressive Familial Intrahepatic Cholestasis, Type 4 (PFIC4)	TJP2	•				•			•
Progressive Pseudorheumatoid Dysplasia	CCN6	•				•	•	•	•
Prolidase Deficiency	PEPD	•				•	•	•	•
Propionic Acidemia, PCCA-Related	PCCA	•				•	•	•	•
Propionic Acidemia, PCCB-Related	PCCB	•				•	•	•	•
Prothrombin-Related Thrombophilia	F2*	•				•	•	•	•
Pseudocholinesterase Deficiency	BCHE	•				•			•
Pseudoxanthoma Elasticum	ABCC6	•				•			•
Pterin-4 Alpha-Carbinolamine Dehydratase (PCD) Deficiency	PCBD1	•					•	•	•
Pycnodysostosis	CTSK	•				•	•	•	•
Pyridoxal 5'-Phosphate-Dependent Epilepsy	PNPO	•				•	•	•	•
Pyridoxine-Dependent Epilepsy	ALDH7A1	•				•	•	•	•
Pyruvate Carboxylase Deficiency	PC	•				•	•	•	•
Pyruvate Dehydrogenase Deficiency, PDHB-Related	PDHB	•				•	•	•	•
Pyruvate Dehydrogenase Deficiency, X-Linked	PDHA1		•			•	•	•	•
Resum Disease, PHYH-Related	PHYH	•				•	•	•	•
Renal Tubular Acidosis and Deafness, ATP6V1B1-Related	ATP6V1B1	•				•	•	•	•
Renal Tubular Acidosis, Proximal, with Ocular Abnormalities and Mental Retardation	SLC4A4	•				•			•
Retinitis Pigmentosa 2	RP2		•				•	•	•
Retinitis Pigmentosa 25	EYS	•				•	•	•	•
Retinitis Pigmentosa 26	CERKL	•				•	•	•	•
Retinitis Pigmentosa 28	FAM161A	•				•	•	•	•
Retinitis Pigmentosa 36	PRCD	•					•	•	•
Retinitis Pigmentosa 59	DHDDS	•			○	•	•	•	•
Retinitis Pigmentosa 62	MAK	•					•	•	•
Retinitis Pigmentosa, X-Linked, RPGR-Related	RPGR		•		○	•	•	•	•
Rhizomelic Chondrodysplasia Punctata, Type 1	PEX7	•				•	•	•	•
Rhizomelic Chondrodysplasia Punctata, Type 2	GNPAT	•				•	•	•	•
Rhizomelic Chondrodysplasia Punctata, Type 3	AGPS	•				•	•	•	•
RLBP1-Related Retinopathy	RLBP1	•				•	•	•	•
Roberts Syndrome	ESCO2	•				•	•	•	•
RYR1-Related Conditions	RYR1	•					•	•	•
Salla Disease	SLC17A5	•				•	•	•	•
Sandhoff Disease	HEXB	•				•	•	•	•
Schimke Immunoosseous Dysplasia	SMARCAL1	•				•	•	•	•
Schindler Disease	NAGA				○	•	•	•	•
Segawa Syndrome, TH-Related	TH	•				•	•	•	•
Senior-Loken Syndrome 4 / Nephronophthisis 4	NPHP4	•				•			•
Sepiapterin Reductase Deficiency	SPR	•					•	•	•
Severe Combined Immunodeficiency (SCID), CD3D-Related	CD3D	•					•	•	•
Severe Combined Immunodeficiency (SCID), CD3E-Related	CD3E	•					•	•	•
Severe Combined Immunodeficiency (SCID), FOXN1-Related	FOXN1	•					•	•	•
Severe Combined Immunodeficiency (SCID), IKBKB-Related	IKBKB	•					•	•	•
Severe Combined Immunodeficiency (SCID), IL7R-Related	IL7R	•					•	•	•
Severe Combined Immunodeficiency (SCID), JAK3-Related	JAK3	•					•	•	•
Severe Combined Immunodeficiency (SCID), PTPRC-Related	PTPRC	•					•	•	•
Severe Combined Immunodeficiency, ADA-Related	ADA	•					•	•	•
Severe Combined Immunodeficiency, RAG1-Related	RAG1	•					•	•	•
Severe Combined Immunodeficiency, Type Athabaskan	DCLRE1C	•					•	•	•
Severe Combined Immunodeficiency, X-Linked	IL2RG		•				•	•	•
Short-Rib Thoracic Dysplasia 3 with or without Polydactyly	DYNC2H1				○	•	•	•	•
Shwachman-Diamond Syndrome, SBDS-Related	SBDS	•					•		
Sialidosis	NEU1	•					•	•	•
Sjögren-Larsson Syndrome	ALDH3A2	•					•	•	•
Smith-Lemli-Optiz Syndrome	DHCR7	•		○	○	•	•	•	•
Spastic Paraparesia, Type 15	ZFYVE26	•				•	•	•	•
Spastic Tetraparesia, Thin Corpus Callosum, and Progressive Microcephaly (SPATCCM)	SLC1A4	•				•	•	•	•
SPG11-Related Conditions	SPG11	•					•	•	•
Spinal Muscular Atrophy	SMN1	•		○	○	•	•	•	•
Spinal Muscular Atrophy With Respiratory Distress Type 1	IGHMBP2	•					•	•	•
Spinocerebellar Ataxia, Autosomal Recessive 10	ANO10				○	•	•	•	•
Spinocerebellar Ataxia, Autosomal Recessive 12	WVOX	•				•			
Spondylocostal Dysostosis 1	DLL3	•					•	•	•
Spondylothoracic Dysostosis, MESP2-Related	MESP2	•					•	•	•
Steel Syndrome	COL27A1	•					•	•	•
Steroid-Resistant Nephrotic Syndrome	NPHS2	•					•	•	•
Stuve-Wiedemann Syndrome	LIFR	•					•	•	•
SURF1-Related Conditions	SURF1	•					•	•	•

CONDITION	GENE	AUTOSOMAL RECESSIVE	X-LINKED	SCREENING RECOMMENDATIONS		PANEL AVAILABILITY			
				ACOG [†]	ACMG	H445	H569 [‡]	H574 [‡]	H613
Surfactant Dysfunction, ABCA3-Related	<i>ABCA3</i>				o	•	•	•	•
Tay-Sachs Disease	<i>HEXA</i>	•		o	o	•	•	•	•
TBCE-Related Conditions	<i>TBCE</i>	•					•	•	•
Thiamine-Responsive Megaloblastic Anemia Syndrome	<i>SLC19A2</i>	•					•	•	•
Thyroid Dyshormonogenesis 1	<i>SLC5A5</i>	•					•	•	•
Thyroid Dyshormonogenesis 2A	<i>TPO</i>	•					•	•	•
Thyroid Dyshormonogenesis 3	<i>TG</i>	•					•	•	•
Thyroid Dyshormonogenesis 6	<i>DUOX2</i>	•					•	•	•
Transcobalamin II Deficiency	<i>TCN2</i>	•					•	•	•
Trichohepatoenteric Syndrome, SKIV2L-Related	<i>SKIIC2</i>	•					•	•	•
Trichohepatoenteric Syndrome, TTC37-Related	<i>TTC37</i>	•				•	•	•	•
Trichothiodystrophy 1 / Xeroderma Pigmentosum, Group D	<i>ERCC2</i>	•			o	•	•	•	•
Trimethylaminuria	<i>FMO3</i>				o	•	•	•	•
Triple A Syndrome	<i>AAAS</i>	•				•	•	•	•
TSHR-Related Conditions	<i>TSHR</i>	•					•	•	•
Tyrosinemia Type III	<i>HPD</i>	•					•	•	•
Tyrosinemia, Type 1	<i>FAH</i>	•			o	•	•	•	•
Tyrosinemia, Type 2	<i>TAT</i>	•				•	•	•	•
Usher Syndrome, Type 1B	<i>MYO7A</i>	•				•	•	•	•
Usher Syndrome, Type 1C	<i>USH1C</i>	•				•	•	•	•
Usher Syndrome, Type 1D	<i>CDH23</i>	•				•	•	•	•
Usher Syndrome, Type 1F	<i>PCDH15</i>	•		o	o	•	•	•	•
Usher Syndrome, Type 1J / Deafness, Autosomal Recessive, 48	<i>CIB2</i>	•				•			
Usher Syndrome, Type 2A	<i>USH2A</i>	•			o	•	•	•	•
Usher Syndrome, Type 2C	<i>ADGRV1</i>	•				•	•	•	•
Usher Syndrome, Type 3	<i>CLRN1</i>	•		o	o	•	•	•	•
Very Long-Chain Acyl-CoA Dehydrogenase Deficiency	<i>ACADVL</i>	•			o	•	•	•	•
Vitamin D-Dependent Rickets Type 2A	<i>VDR</i>	•					•	•	•
Vici Syndrome	<i>EPG5</i>	•					•	•	•
Vitamin D Dependent Rickets, Type 1A	<i>CYP27B1</i>	•			o	•	•	•	•
Vldlr-Associated Cerebellar Hypoplasia	<i>VLDLR</i>	•					•	•	•
Walker-Warburg Syndrome, FKTN-Related	<i>FKTN</i>	•			o	•	•	•	•
Walker-Warburg Syndrome, ISPD-Related	<i>ISPD</i>	•				•			
Walker-Warburg Syndrome, LARGE1-Related	<i>LARGE1</i>	•				•	•	•	•
Walker-Warburg Syndrome, POMT1-Related	<i>POMT1</i>	•				•	•	•	•
Walker-Warburg Syndrome, POMT2-Related	<i>POMT2</i>	•				•	•	•	•
Warsaw Breakage Syndrome	<i>DDX11</i>	•					•	•	•
Werner Syndrome	<i>WRN</i>	•				•	•	•	•
Wilson Disease	<i>ATP7B</i>	•			o	•	•	•	•
Wiskott-Aldrich Syndrome	<i>WAS</i>		•			•	•	•	•
Wolcott-Rallison Syndrome	<i>EIF2AK3</i>	•				•	•	•	•
Wolman Disease	<i>LIPA</i>	•				•	•	•	•
Woodhouse-Sakati Syndrome	<i>DCAF17</i>	•					•	•	•
Xeroderma Pigmentosum Variant Type	<i>POLH</i>	•					•	•	•
Xeroderma Pigmentosum, Group A	<i>XPA</i>	•				•	•	•	•
Xeroderma Pigmentosum, Group C	<i>XPC</i>	•			o	•	•	•	•
X-Linked Chondrodysplasia Punctata 1	<i>ARSL</i>		•			•	•	•	•
X-Linked Lissencephaly with Abnormal Genitalia	<i>ARX</i>		•		o	•	•	•	•
Zellweger Spectrum Disorder, PEX13-Related	<i>PEX13</i>	•					•	•	•
Zellweger Spectrum Disorder, PEX16-Related	<i>PEX16</i>	•					•	•	•
Zellweger Spectrum Disorder, PEX5-Related	<i>PEX5</i>	•					•	•	•
Zellweger Spectrum Disorders, PEX10-Related	<i>PEX10</i>	•					•	•	•
Zellweger Spectrum Disorders, PEX12-Related	<i>PEX12</i>	•					•	•	•
Zellweger Spectrum Disorders, PEX1-Related	<i>PEX1</i>	•					•	•	•
Zellweger Spectrum Disorders, PEX26-Related	<i>PEX26</i>	•					•	•	•
Zellweger Spectrum Disorders, PEX2-Related	<i>PEX2</i>	•					•	•	•
Zellweger Spectrum Disorders, PEX6-Related	<i>PEX6</i>	•					•	•	•

*Low penetrance genes with variable presentation. Horizon 556 is available to clinicians who would like to exclude screening for these 13 genes.

[†]Note that ACOG screening recommendations listed here include diseases in ACOG Committee Opinion 690 example expanded carrier screening panel, as well as the diseases listed in ACOG Committee Opinion 691

[‡]Panel is named based on the total number of genes, counting *HBA1* and *HBA2* separately.

1. American College of Obstetricians and Gynecologists, Committee Opinion # 690, March 2017.

2. American College of Obstetricians and Gynecologists, Committee Opinion # 691, March 2017.

3. Gregg et al. Screening for autosomal recessive and X-linked conditions during pregnancy and preconception: a practice resource of the American College of Medical Genetics and Genomics (ACMG), July 2021.