

# Universal Panel Disease List

The Myriad Foresight Carrier Screen focuses on serious, clinically-actionable, and prevalent conditions to ensure you are providing meaningful information to your patients.

11-Beta-Hydroxylase-Deficient  
Congenital Adrenal Hyperplasia  
(*CYP11B1*)

6-Pyruvoyl-Tetrahydropterin  
Synthase Deficiency (*PTS*)

ABCC8-Related Familial  
Hyperinsulinism (*ABCC8*)

Adenosine Deaminase  
Deficiency (*ADA*)

Adrenoleukodystrophy:  
X-Linked (*ABCD1*) **X-linked**

Alpha Thalassemia (*HBA1/  
HBA2*)\* **ACOG** **ACMG**

Alpha-Mannosidosis (*MAN2B1*)

Alpha-Sarcoglycanopathy  
(including Limb-Girdle Muscular  
Dystrophy, Type 2D) (*SGCA*)

Alport Syndrome, X-Linked  
(*COL4A5*) **X-linked**

Alstrom Syndrome (*ALMS1*)

AMT-Related Glycine  
Encephalopathy (*AMT*)

Andermann Syndrome  
(*SLC12A6*)

Argininemia (*ARG1*)

Argininosuccinic Aciduria (*ASL*)

Aspartylglycosaminuria (*AGA*)

Ataxia with Vitamin E Deficiency  
(*TTPA*)

Ataxia-Telangiectasia (*ATM*)

ATP7A-Related Disorders  
(*ATP7A*) **X-linked**

Autoimmune Polyglandular  
Syndrome Type 1 (*AIRE*)

Autosomal Recessive  
Osteopetrosis, Type 1 (*TCIRG1*)

Autosomal Recessive Polycystic  
Kidney Disease, PKHD1-  
Related (*PKHD1*)

Autosomal Recessive Spastic  
Ataxia of Charlevoix-Saguenay  
(*SACS*)

Bardet-Biedl Syndrome,  
BBS1-Related (*BBS1*)

Bardet-Biedl Syndrome,  
BBS10-Related (*BBS10*)

Bardet-Biedl Syndrome,  
BBS12-Related (*BBS12*)

Bardet-Biedl Syndrome,  
BBS2-Related (*BBS2*)

BCS1L-Related Disorders  
(*BCS1L*)

Beta-Sarcoglycanopathy  
(including Limb-Girdle Muscular  
Dystrophy, Type 2E) (*SGCB*)

Biotinidase Deficiency (*BTD*)

Bloom Syndrome (*BLM*) **ACMG**

Calpainopathy (*CAPN3*)

Canavan Disease (*ASPA*) **ACOG** **ACMG**

Carbamoylphosphate  
Synthetase I Deficiency (*CPS1*)

Carnitine Palmitoyltransferase  
IA Deficiency (*CPT1A*)

Carnitine Palmitoyltransferase II  
Deficiency (*CPT2*)

Cartilage-Hair Hypoplasia (*RMRP*)

Cerebrotendinous  
Xanthomatosis (*CYP27A1*)

Citrullinemia, Type 1 (*ASS1*)

CLN3-Related Neuronal Ceroid  
Lipofuscinosis (*CLN3*)

CLN5-Related Neuronal Ceroid  
Lipofuscinosis (*CLN5*)

CLN6-Neuronal Ceroid  
Lipofuscinosis, Type 6 (*CLN6*)

CLN8-Related Neuronal Ceroid  
Lipofuscinosis (*CLN8*)

Cohen Syndrome (*VPS13B*)

COL4A3-Related Alport  
Syndrome (*COL4A3*)

COL4A4-Related Alport  
Syndrome (*COL4A4*)

Combined Pituitary Hormone  
Deficiency, PROP1-Related  
(*PROP1*)

Congenital Adrenal  
Hyperplasia, CYP21A2-Related  
(*CYP21A2*)\*

Congenital Disorder of  
Glycosylation, MPI-Related  
(*MPI*)

Congenital Disorder of  
Glycosylation, Type Ia (*PMM2*)

Congenital Disorder of  
Glycosylation, Type Ic (*ALG6*)

Costeff Optic Atrophy  
Syndrome (*OPA3*)

Cystic Fibrosis (*CFTR*) **ACOG** **ACMG**

Cystinosis (*CTNS*)

D-Bifunctional Protein  
Deficiency (*HSD17B4*)

Delta-Sarcoglycanopathy  
(*SGCD*)

Dihydrolipoamide  
Dehydrogenase Deficiency  
(*DLD*)

Dysferlinopathy (*DYSF*)

Dystrophinopathies (including  
Duchenne/Becker Muscular  
Dystrophy)(*DMD*) **X-linked**

ERCC6-Related Disorders  
(*ERCC6*)

ERCC8-Related Disorders  
(*ERCC8*)

EVC-Related Ellis-Van Creveld  
Syndrome (*EVC*)

EVC2-Related Ellis-Van Creveld  
Syndrome (*EVC2*)

Fabry Disease (*GLA*) **X-linked**

Familial Dysautonomia (*IKBKAP*)  
**ACOG** **ACMG**

Familial Mediterranean Fever  
(*MEFV*)

Fanconi Anemia  
Complementation, Group A  
(*FANCA*)

Fanconi Anemia, FANCC-  
Related (*FANCC*) **ACMG**

FKRP-Related Disorders (*FKRP*)

FKTN-Related Disorders  
(including Walker-Warburg  
Syndrome) (*FKTN*)

Fragile X Syndrome (*FMR1*)\*  
**X-linked**

Galactokinase Deficiency  
(*GALK1*)

Galactosemia (*GALT*)

Gamma-Sarcoglycanopathy  
(*SGCG*)

Gaucher Disease (*GBA*)\* **ACMG**

GJB2-Related DFNB1  
Nonsyndromic Hearing Loss  
and Deafness (including two  
GJB6 deletions) (*GJB2*)

GLB1-Related Disorders (*GLB1*)

GLDC-Related Glycine  
Encephalopathy (*GLDC*)

Glutaric Acidemia, GCDH-  
Related (*GCDH*)

Glycogen Storage Disease,  
Type Ia (*G6PC*)

Glycogen Storage Disease,  
Type Ib (*SLC37A4*)

Glycogen Storage Disease,  
Type III (*AGL*)

GNE Myopathy (*GNE*)

GNPTAB-Related Disorders  
(*GNPTAB*)

HADHA-Related Disorders  
(including Long Chain  
3-Hydroxyacyl-CoA  
Dehydrogenase Deficiency)  
(*HADHA*)

Hb Beta Chain-Related  
Hemoglobinopathy (including  
Beta Thalassemia and Sickle  
Cell Disease)(*HBB*) **ACOG**

Hereditary Fructose Intolerance  
(*ALDOB*)

Herlitz Junctional Epidermolysis  
Bullosa, LAMB3-Related  
(*LAMB3*)

Hexosaminidase A Deficiency  
(including Tay-Sachs Disease)  
(*HEXA*) **ACOG** **ACMG**

HMG-CoA Lyase Deficiency  
(*HMGCL*)

|   |   |   |   |
|---|---|---|---|
| Holocarboxylase Synthetase Deficiency (HLCS)                      | MKS1-Related Disorders (MKS1)   | Peroxisome Biogenesis Disorder, Type 5 (PEX2)                       | TPP1-Related Neuronal Ceroid Lipofuscinosis (TPP1)                      |
| Homocystinuria, CBS-Related (CBS)                                 | Mucopolysaccharidosis III Gamma (GNPTG)                               | Peroxisome Biogenesis Disorder, Type 6 (PEX10)                      | Tyrosine Hydroxylase Deficiency (TH)                                    |
| Hydrolethalus Syndrome (HYLS1)                                    | Mucopolysaccharidosis IV (MCOLN1) <span>ACMG</span>                   | Phenylalanine Hydroxylase Deficiency (PAH)                          | Tyrosinemia, Type I (FAH)   |
| Hypophosphatasia (ALPL)   | Mucopolysaccharidosis, Type I (including Hurler Syndrome) (IDUA)      | POMGNT-Related Disorders (POMGNT1)                                  | Tyrosinemia, Type II (TAT)  |
| Isovaleric Acidemia (IVD)   | Mucopolysaccharidosis, Type II (IDS) <span>X-linked</span>            | Pompe Disease (GAA)   | USH1C-Related Disorders (USH1C)   |
| Joubert Syndrome 2 (TMEM216)                                      | Mucopolysaccharidosis, Type IIIA (SGSH)                               | PPT1-Related Neuronal Ceroid Lipofuscinosis (PPT1)                  | USH2A-Related Disorders (USH2A)   |
| Junctional Epidermolysis Bullosa, LAMC2-Related (LAMC2)           | Mucopolysaccharidosis, Type IIIB (NAGLU)                              | Primary Carnitine Deficiency (SLC22A5)                              | Usher Syndrome, Type 3 (CLRN1)  |
| Junctional Epidermolysis Bullosa, LAMA3-Related (LAMA3)           | Mucopolysaccharidosis, Type IIIC (HGSNAT)                             | Primary Hyperoxaluria, Type 1 (AGXT)                                | Very Long Chain Acyl-CoA Dehydrogenase Deficiency (ACADVL)              |
| KCNJ11-Related Familial Hyperinsulinism (KCNJ11)                  | MUT-Related Methylmalonic Acidemia (MUT)                              | Primary Hyperoxaluria, Type 2 (GRHPR)                               | Wilson Disease (ATP7B)  |
| Krabbe Disease (GALC)   | MYO7A-Related Disorders (MYO7A)                                       | Primary Hyperoxaluria, Type 3 (HOGA1)                               | X-Linked Congenital Adrenal Hypoplasia (NROB1) <span>X-linked</span>    |
| LAMA2-Related Muscular Dystrophy (LAMA2)                          | NEB-Related Nemaline Myopathy (NEB)                                   | Pycnodysostosis (CTSK)  | X-Linked Juvenile Retinoschisis (RS1) <span>X-linked</span>             |
| Leigh Syndrome, French-Canadian Type (LRPPRC)                     | Nephrotic Syndrome, NPHS1-Related (NPHS1)                             | Pyruvate Carboxylase Deficiency (PC)                                | X-Linked Myotubular Myopathy (MTM1) <span>X-linked</span>               |
| Lipoid Congenital Adrenal Hyperplasia (STAR)                      | Niemann-Pick Disease, SMPD1-Related (SMPD1) <span>ACMG</span>         | Rhizomelic Chondrodysplasia Punctata, Type 1 (PEX7)                 | X-Linked Severe Combined Immunodeficiency (IL2RG) <span>X-linked</span> |
| Lysosomal Acid Lipase Deficiency (LIPA)                           | Niemann-Pick Disease, Type C1 (NPC1)                                  | RTEL1-Related Disorders (RTEL1)                                     | Xeroderma Pigmentosum, Group A (XPA)                                    |
| Maple Syrup Urine Disease, Type Ia (BCKDHA)                       | Niemann-Pick Disease, Type C2 (NPC2)                                  | Salla Disease (SLC17A5)   | Xeroderma Pigmentosum, Group C (XPC)                                    |
| Maple Syrup Urine Disease, Type Ib (BCKDHB)                       | Nijmegen Breakage Syndrome (NBN)                                      | Sandhoff Disease (HEXB)   |   |
| Maple Syrup Urine Disease, Type II (DBT)                          | Ornithine Transcarbamylase Deficiency (OTC) <span>X-linked</span>     | Short Chain Acyl-CoA Dehydrogenase Deficiency (ACADS)               |   |
| Medium Chain Acyl-CoA Dehydrogenase Deficiency (ACADM)            | PCCA-Related Propionic Acidemia (PCCA)                                | Sjogren-Larsson Syndrome (ALDH3A2)                                  |   |
| Megalencephalic Leukoencephalopathy with Subcortical Cysts (MLC1) | PCCB-Related Propionic Acidemia (PCCB)                                | SLC26A2-Related Disorders (SLC26A2)                                 |   |
| Metachromatic Leukodystrophy (ARSA)                               | PCDH15-Related Disorders (including Usher Syndrome, Type 1F) (PCDH15) | Smith-Lemli-Opitz Syndrome (DHCR7)                                  |   |
| Methylmalonic Acidemia, cblA Type (MMAA)                          | Pendred Syndrome (SLC26A4)  | Spastic Paraplegia, Type 15 (ZFYVE26)                               | <span>ACOG</span><br>Indicates disease listed in ACOG guidelines        |
| Methylmalonic Acidemia, cblB Type (MMAB)                          | Peroxisome Biogenesis Disorder, Type 1 (PEX1)                         | Spinal Muscular Atrophy (SMN1)* <span>ACOG</span> <span>ACMG</span> | <span>ACMG</span><br>Indicates disease listed in ACMG guidelines        |
| Methylmalonic Aciduria and Homocystinuria, cblC Type (MMAHC)      | Peroxisome Biogenesis Disorder, Type 3 (PEX12)                        | Spondylothoracic Dysostosis (MESP2)                                 | <span>X-linked</span><br>Indicates X-linked disorders                   |
|   | Peroxisome Biogenesis Disorder, Type 4 (PEX6)                         | Steroid-Resistant Nephrotic Syndrome (NPHS2)                        |   |
|   |   | TGM1-Related Autosomal Recessive Congenital Ichthyosis (TGM1)       |   |

\*Analyzed using custom assay