## **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

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)\*

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)

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*)

Homocystinuria, CBS-Related (CBS)

Hydrolethalus Syndrome (HYLS1)

Hypophosphatasia (ALPL)

Isovaleric Acidemia (IVD)

Joubert Syndrome 2 (TMEM216)

Junctional Epidermolysis Bullosa, LAMC2-Related (LAMC2)

Junctional Epidermolysis Bullosa, LAMA3-Related (LAMA3)

KCNJ11-Related Familial Hyperinsulinism (KCNJ11)

Krabbe Disease (GALC)

LAMA2-Related Muscular Dystrophy (*LAMA2*)

Leigh Syndrome, French-Canadian Type (*LRPPRC*)

Lipoid Congenital Adrenal Hyperplasia (*STAR*)

Lysosomal Acid Lipase Deficiency (*LIPA*)

Maple Syrup Urine Disease, Type Ia (*BCKDHA*)

Maple Syrup Urine Disease, Type Ib (BCKDHB)

Maple Syrup Urine Disease, Type II (*DBT*)

Medium Chain Acyl-CoA Dehydrogenase Deficiency (ACADM)

Megalencephalic Leukoencephalopathy with Subcortical Cysts (*MLC1*)

Metachromatic Leukodystrophy (ARSA)

Methylmalonic Acidemia, cblA Type (*MMAA*)

Methylmalonic Acidemia, cblB Type (*MMAB*)

Methylmalonic Aciduria and Homocystinuria, cblC Type (MMACHC)

MKS1-Related Disorders (MKS1)

Mucolipidosis III Gamma (GNPTG)

Mucolipidosis IV (MCOLN1)

Mucopolysaccharidosis, Type I (including Hurler Syndrome) (IDUA)

Mucopolysaccharidosis, Type II (*IDS*) X-linked

Mucopolysaccharidosis, Type IIIA (*SGSH*)

Mucopolysaccharidosis, Type IIIB (*NAGLU*)

Mucopolysaccharidosis, Type IIIC (*HGSNAT*)

MUT-Related Methylmalonic Acidemia (*MUT*)

MYO7A-Related Disorders (MYO7A)

NEB-Related Nemaline Myopathy (*NEB*)

Nephrotic Syndrome, NPHS1-Related (NPHS1)

Niemann-Pick Disease, SMPD1-Related (SMPD1) ACMG

Niemann-Pick Disease, Type C1 (NPC1)

Niemann-Pick Disease, Type C2 (*NPC2*)

Nijmegen Breakage Syndrome (NBN)

Ornithine Transcarbamylase Deficiency (*OTC*) X-linked

PCCA-Related Propionic Acidemia (*PCCA*)

PCCB-Related Propionic Acidemia (*PCCB*)

PCDH15-Related Disorders (including Usher Syndrome, Type 1F) (*PCDH15*)

Pendred Syndrome (SLC26A4)

Peroxisome Biogenesis Disorder, Type 1 (PEX1)

Peroxisome Biogenesis Disorder, Type 3 (*PEX12*)

Peroxisome Biogenesis Disorder, Type 4 (*PEX6*) Peroxisome Biogenesis Disorder, Type 5 (*PEX2*)

Peroxisome Biogenesis Disorder, Type 6 (*PEX10*)

Phenylalanine Hydroxylase Deficiency (*PAH*)

POMGNT-Related Disorders (POMGNT1)

Pompe Disease (GAA)

PPT1-Related Neuronal Ceroid Lipofuscinosis (PPT1)

Primary Carnitine Deficiency (*SLC22A5*)

Primary Hyperoxaluria, Type 1 (AGXT)

Primary Hyperoxaluria, Type 2 (*GRHPR*)

Primary Hyperoxaluria, Type 3 (*HOGA1*)

Pycnodysostosis (CTSK)

Pyruvate Carboxylase Deficiency (*PC*)

Rhizomelic Chondrodysplasia Punctata, Type 1 (*PEX7*)

RTEL1-Related Disorders (RTEL1)

Salla Disease (SLC17A5)

Sandhoff Disease (HEXB)

Short Chain Acyl-CoA Dehydrogenase Deficiency (ACADS)

Sjogren-Larsson Syndrome (*ALDH3A2*)

SLC26A2-Related Disorders (SLC26A2)

Smith-Lemli-Opitz Syndrome (DHCR7)

Spastic Paraplegia, Type 15 (ZFYVE26)

Spinal Muscular Atrophy (SMN1)\* ACOG ACMG

Spondylothoracic Dysostosis (MESP2)

Steroid-Resistant Nephrotic Syndrome (*NPHS2*)

TGM1-Related Autosomal Recessive Congenital Ichthyosis (TGM1) TPP1-Related Neuronal Ceroid Lipofuscinosis (*TPP1*)

Tyrosine Hydroxylase Deficiency (TH)

Tyrosinemia, Type I (FAH)
Tyrosinemia, Type II (TAT)

USH1C-Related Disorders (USH1C)

USH2A-Related Disorders (USH2A)

Usher Syndrome, Type 3 (CLRN1)

Very Long Chain Acyl-CoA Dehydrogenase Deficiency (ACADVL)

Wilson Disease (ATP7B)

X-Linked Congenital Adrenal Hypoplasia (NROB1) X-linked

X-Linked Juvenile Retinoschisis (RS1) X-linked

X-Linked Myotubular Myopathy (MTM1) X-linked

X-Linked Severe Combined Immunodeficiency (*IL2RG*) X-linked

Xeroderma Pigmentosum, Group A (*XPA*)

Xeroderma Pigmentosum, Group C (*XPC*)

## ACOG

Indicates disease listed in ACOG guidelines

## ACMG

Indicates disease listed in ACMG guidelines

## X-linked

Indicates X-linked disorders

\*Analyzed using custom assay

