Recommended Uniform Screening Panel Core Conditions (As of January 2023)

X: Condition is in this category --: Condition is not in this category

Core Condition	Metabolic Disorder - Organic acid condition	Metabolic Disorder - Fatty acid oxidation disorder	Metabolic Disorder - Amino acid disorder	Endocrine Disorder	Hemoglobin Disorder	Other Disorder
Propionic Acidemia	Х					
Methylmalonic Acidemia (methylmalonyl-CoA mutase)	х					
Methylmalonic Acidemia (Cobalamin disorders)	х					
Isovaleric Acidemia	Х					
3-Methylcrotonyl-CoA Carboxylase Deficiency	х					
3-Hydroxy-3-Methyglutaric Aciduria	х					
Holocarboxylase Synthase Deficiency	х					
ß-Ketothiolase Deficiency	X					
Glutaric Acidemia Type I	Х					
Carnitine Uptake Defect/Carnitine Transport Defect		x				
Medium-chain Acyl-CoA Dehydrogenase Deficiency		х				
Very Long-chain Acyl-CoA Dehydrogenase Deficiency		х				
Long-chain L-3 Hydroxyacyl-CoA Dehydrogenase Deficiency		х				
Trifunctional Protein Deficiency		Х				
Argininosuccinic Aciduria			Х			
Citrullinemia, Type I			Х			
Maple Syrup Urine Disease			Х			
Homocystinuria			Х			
Classic Phenylketonuria			Х			
Tyrosinemia, Type I			Х			
Guanidinoacetate Methyltransferase Deficiency			х			-
Primary Congenital Hypothyroidism				Х		
Congenital adrenal hyperplasia				Х		
S,S Disease (Sickle Cell Anemia)					Х	
S, βeta-Thalassemia					Х	
S,C Disease					Х	
Biotinidase Deficiency						Х
Critical Congenital Heart Disease						X
Cystic Fibrosis						Х
Classic Galactosemia						Х
Glycogen Storage Disease Type II (Pompe)						X
Hearing Loss						Х

Recommended Uniform Screening Panel Core Conditions (As of January 2023)

Core Condition - continued	Metabolic Disorder - Organic acid condition	Metabolic Disorder - Fatty acid oxidation disorder	Metabolic Disorder - Amino acid disorder	Endocrine Disorder	Hemoglobin Disorder	Other Disorder
Severe Combined Immunodeficiencies						Х
Mucopolysaccharidosis Type I						x
X-linked Adrenoleukodystrophy						х
Spinal Muscular Atrophy due to homozygous deletion of exon 7 in SMN1						х
Mucopolysaccharidosis Type II						Х

Recommended Uniform Screening Panel¹ SECONDARY² CONDITIONS ³ (As of January 2023)

		(A3 OI Janua	, 2020)			
Secondary Condition	Metabolic Disorder - Organic acid condition	Metabolic Disorder - Fatty acid oxidation disorder	Metabolic Disorder - Amino acid disorder	Endocrine Disorder	Hemoglobin Disorder	Other Disorder
Methylmalonic acidemia with homocystinuria	х					
Malonic acidemia	Х					
Isobutyrylglycinuria	Х					
2-Methylbutyrylglycinuria	Х					
3-Methylglutaconic aciduria	Х					
2-Methyl-3-hydroxybutyric aciduria	Х					
Short-chain acyl-CoA dehydrogenase deficiency		х				
Medium/short-chain L-3- hydroxyacyl-CoA dehydrogenase deficiency		x				
Glutaric acidemia type II		Х				
Medium-chain ketoacyl-CoA thiolase deficiency		х				
2,4 Dienoyl-CoA reductase deficiency		х				
Carnitine palmitoyltransferase type I deficiency		х			-	
Carnitine palmitoyltransferase type II deficiency		х			1	-
Carnitine acylcarnitine translocase deficiency		х			-	
Argininemia			Х			
Citrullinemia, type II			Х			
Hypermethioninemia			Х			
Benign hyperphenylalaninemia			Х			
Biopterin defect in cofactor biosynthesis			х			

Secondary Condition – Continued	Metabolic Disorder - Organic acid condition	Metabolic Disorder - Fatty acid oxidation disorder	Metabolic Disorder - Amino acid disorder	Endocrine Disorder	Hemoglobin Disorder	Other Disorder
Biopterin defect in cofactor regeneration			х			
Tyrosinemia, type II			Х			
Tyrosinemia, type III			Х			
Various other hemoglobinopathies					Х	
Galactoepimerase deficiency						х
Galactokinase deficiency						Х
T-cell related lymphocyte deficiencies						Х

^{1.} Selection of conditions based upon "Newborn Screening: Towards a Uniform Screening Panel and System." *Genetic Med.* 2006; 8(5) Suppl: S12-S252" as authored by the American College of Medical Genetics (ACMG) and commissioned by the Health Resources and Services Administration (HRSA).

Disorders that can be detected in the differential diagnosis of a core disorder.
 Nomenclature for Conditions based upon "Naming and Counting Disorders (Conditions) Included in Newborn Screening Panels." *Pediatrics*. 2006; 117 (5) Suppl: S308-S314.