

Case Definitions for Newborn Screening

Table of Contents

V	letabolic Conditions	5
	Propionic Acidemia (PROP)	6
	Methylmalonic Acidemia (MMA)	7
	Methylmalonic Acidemia (MMA) with Homocystinuria (HCY)	9
	Holocarboxylase Synthase Deficiency (MCD)	.10
	Isovaleric Acidemia (IVA)	.11
	Glutaric Cidemia (GA)	.12
	Primary Carnitine Deficiency/ Carnitine Uptake Defect (CUD)	.13
	Very long-chain acyl-CoA Dehydrogenase Deficiency (VLCAD)	.14
	Trifunctional Protein Deficiency (TFP)-Inclusive of LCHAD	.16
	Argininosuccinic Aciduria (ASA)	.18
	Citrullinemia, type 1-Exclusive of Citrin Deficiency (CIT)	.19
	3-Methylcrotonyl-CoA (3-MCC)	.20
	Tyrosinemia, Type 1 (TYR 1)	.21
	Medium-chain acyl-CoA Dehydrogenase Deficiency (MCAD)	.22
	Maple Syrup Urine Disease (MSUD)	.24
	Cystathionine Beta-Synthase (CBS) Deficiency	.25
	Benign Hyperphenylalaninemia (H-PHE)	.25
	Biotinidase Deficiency (BIOT)	.26
	Classic Galactosemia (GALT)	.27
	Variant Galactosemia	.28
	Arginase Deficiency	.28
Εı	ndocrinology Disorders	.29
	Primary Congenital Hypothyrodism (CH)	.30
	Secondary Congenital Hypothyroidism (CH)	.30
	Thyroxine-binding Globulin (TBG) or other Protein Binding Defect	.31
	21-Hydroxylase Deficiency – Classical Salt Wasting	.32
	21-Hydroxylase Deficiency-Classical Simple Virilizing	.34
Α	lpha thalassemia	.37
	S Alpha Thal	.38
	S Alpha Thal	
	C Alpha Thal	.41
	D Alpha Thal	43

	O _{Arab} Alpha Thal	44
	3 Deletion Alpha Thalassemia (Hgb H disease)	45
	Hgb H Constant Spring (2 alpha gene deletion (cis) plus Constant Spring point mutation (trans))	46
3e	ta Thalassemia	47
	Beta + Thal	48
	E Beta + Thal	50
	D Beta + Thal	51
	C Beta + Thal	53
	OArab Beta + Thal	54
	S Beta (0) Thal	55
	E Beta (0) Thal	56
	D Beta (0) Thal	57
	C Beta (0) Thal	58
	OArab Beta (0) Thal	59
	Beta Thal Major or (Homozygous or Heterozygous for 2 Beta Thal mutations)	60
	HPFH	61
	FSHPFH	62
	FOARABHPFH	63
	FCHPFH	64
	FEHPFH	65
Sic	kle Cell Diseases	66
	SC Disease	67
	SD Disease	68
	SD Disease	69
	SE Disease	70
	SOArab Disease	71
	CD Disease	72
	CD Disease	73
	CE Disease	74
	CO _{Arab} Disease	75
	DE Disease	76
	DO _{Arab} Disease	77
	SS Disease	78
	CC Disease	79

EE Disease	80	
DD Disease	81	
Homozygous OArab Disease	82	
Cystic Fibrosis (CF)	83	
Typical CF	84	
Severe Combined Immunodeficiency (SCID)	87	
Typical SCID	88	
Leaky SCID	90	
Omenn Syndrome	91	
Non-SCID conditions associated with SCID NBS	92	
Pompe Disease	93	
Mucopolysaccharidosis Type I (MPS I)	97	
X-linked Adrenoleukodystrophy (X-ALD)	100	
Spinal Muscular Atrophy (SMA)	109	

Metabolic Conditions Case Definitions Tables September 29, 2013

Definitions created by panel of experts between June 2011 and September 2013. This project was funded in part by Cooperative Agreement # U22MC24078 from the Health Resources and Services Administration (HRSA).

Propionic Acidemia (PROP)

	Classification	Urine organic acids	Plasma	Mutation analysis
Propionic Acidemia			Acylcarnitines	
	Definite	Untested or unknown	Untested or unknown	2 known disease causing variants in the same gene (Allele 1 – variant known to be disease causing and Allele 2 – variant known to be disease causing)
	Definite	Presence ofmethyl citrate and -+/-3OH propionic acid, propionyl glycine, -tiglyglycine and Absence of: -MMA and - methyl crotonyl glycine	Elevated C3	Untested or unknown
	Probable	Presence of -3-OH propionic and Absence of: -MMA and -methyl crotonyl glycine	Elevated C3	2 variants of uncertain significance in the same gene - predicted to be pathogenic [Allele 1 - variant of unknown significance (predicted to be pathogenic) and Allele 2 - variant of unknown significance (predicted to be pathogenic)]
	Possible	Presence of -3-OH Propionic and Absence of -MMA and -methyl crotonyl glycine	Elevated C3	Untested or unknown
	Possible	Presence of -3-OH propionic and Absence of -MMA and -methyl crotonyl glycine	Elevated C3	2 variants of uncertain significance in the same gene (Allele 1 - variant of unknown significance and Allele 2 – variant of unknown significance)
	Possible	Presence of -3-OH propionic and Absence of -MMA and -methyl crotonyl glycine	Elevated C3	1 known disease causing variant (Allele 1 - variant known to be disease causing)
	Possible	Absence of -MMA and -methyl crotonyl glycine	Elevated C3	2 variants of uncertain significance in the same gene (Allele 1 - variant of unknown significance and Allele 2 – variant of unknown significance)
	Possible	Presence of -3-OH propionic and Absence of -MMA and -methyl crotonyl glycine	Elevated C3	No variants found

Methylmalonic Acidemia (MMA)

	Classification	•	Plasma	Maternal	Infant	Mutation analysis	Enzyme analysis
		serum organic acids	Acylcarnitines	Studies	chemistries/studie s		,,
	Definite	Untested or unknown	Untestedor unknown	Untestedor unknown	Untested or unknown	2 known disease causing variants in the same gene (Allele 1 – variant known to be disease causing and Allele 2 – variant known to be disease causing)	Untested or unknown
	Definite	Untested or unknown	Untested or unknown	Untested or unknown	Untestedorunknown	Untested or unknown	complementation studies consistent with corresponding disease
mut-; mut0; CblDv2)	Definite	Elevated MMA for age	Elevated C3	Absence of B12 deficiency	-Absence of B12 deficiency and -Normal homocysteine	2 variants of uncertain significance in the same gene - predicted to be pathogenic [Allele 1 - variant of unknown significance (predicted to be pathogenic) and Allele 2 - variant of unknown significance (predicted to be pathogenic) and the significance (predicted to be pathogenic)]	Untested or unknown
MMA; (CbIA; CbIB, 1	Probable	Elevated MMA for age	Elevated C3	Absence of B12 deficiency	-Absence of B12 deficiency and -Normal homocysteine	2 variants of uncertain significance in the same gene (Allele 1 - variant of unknown significance and Allele 2 – variant of unknown significance)	Untestedorunknown
	Probable	Elevated MMA for age	Elevated C3	Absence of B12 deficiency	-Absence of B12 deficiency and -Normal homocysteine	untested or unknown	Untested or unknown
	Probable	Elevated MMA for age	Elevated C3	Absence of B12 deficiency	-Absence of B12 deficiency and -Normal homocysteine	1 known disease causing variant (Allele 1 - variant known to be disease causing)	Untested or unknown

Possible	Elevated MMA for age	Elevated C3	Absence of B12 deficiency	-Absence of B12 deficiency and -Normal homocysteine	None found	Untested or unknown
Possible	Elevated MMA for age	Elevated C3	Untested or unknown	-Absence of B12 deficiency and -Normal homocysteine	N/A	Untestedorunknown

Methylmalonic Acidemia (MMA) with Homocystinuria (HCY)

	Classificatio		Plasma	Matern	Infant	Mutation analysis	Enzyme analysis
	n	serum organic	Acylcarnitin es	al Studies	chemistries/studie s		
		acids	E 5	Studies			
	Definite	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	2 known diseasecausing variants in the same gene (Allele 1 – variant known to be disease causing and Allele 2 – variant known to be disease causing)	Untested or unknown
	Definite	Untested or unknown	Untested or unknown	Untested or unknown	Untestedorunknown	Untestedorunknown	complementation studies consistent with corresponding disease
cbiF; CbiD)	Definite	Elevated MMA for age	Elevated C3	Absence of B12 deficiency	-Absence of B12 deficiency and - Elevated homocysteine	2 variants of uncertain significance in the same gene - predicted to be pathogenic [Allele 1 - variant of unknown significance (predicted to be pathogenic) and Allele 2 - variant of unknown significance (predicted to be pathogenic)]	Untested or unknown
MIMA with Homocystinuria; (CblC; CblDv1; CblF; CblD)	Probable	Elevated MMA for age	Elevated C3	Absence of B12 deficiency	-Absence of B12 deficiency and -Elevated homocysteine	2 variants of uncertain significance in the same gene (Allele 1 - variant of unknown	Untested or unknown
Homocystinuria	Probable	Elevated MMA for age	Elevated C3	Absence of B12 deficiency	-Absence of B12 deficiency and -Elevated homocysteine	untested or unknown	Untested or unknown
MMAwith	Probable	Elevated MMA for age	Elevated C3	Absence of B12 deficiency	-Absence of B12 deficiency and - Elevated homocysteine	1 known disease causing variant (Allele 1 - variant known to be disease causing)	Untestedorunknown
	Possible	Elevated MMA for age	Elevated C3	Absence of B12 deficiency	-Absence of B12 deficiency and - Elevated homocysteine	None found	Untested or unknown
	Possible	Elevated MMA for age	Elevated C3	Untested or unknown	Absence of B12 deficiency and - Elevated homocysteine	N/A	Untestedorunknown

Holocarboxylase Synthase Deficiency (MCD)

	Classificatio	Urine	Plasma	Infant	Mutation analysis	Enzyme analysis
	n	organic	Acylcarniti	chemistries/studie	iviutation analysis	LIIZYIIIE allalysis
		acids	nes	S		
	Definite	Untested or unknown	Untested or unknown	Untested or unknown	2 known disease causing variants in the same gene (Allele 1 – variant known to be disease causing and Allele 2 – variant known to be disease causing)	Untested or unknown
	Definite	Untested or unknown	Untested or unknown	Untestedorunknown	Untested or unknown	enzyme activity on fibroblasts or WBCs consistentwith disease
disorders	Definite	Elevated -3OH Isovaleric and -3OH Propionic and - 3methylcrotony I glycine	elevated -C3; and -C5-OH	Normal biotinidase studies	Untested or unknown	Untested or unknown
ylase Synthase Deficiency or other biotin disorders	Possible	Elevated -3OH Isovaleric and - 3methylcrotony I glycine	elevated -C3; and -C5-OH	Normal biotinidase studies	Untested or unknown	Untested or unknown
lase Synthase Defici	Possible	Elevated -propionyl glycine and - 3methylcrotony I glycine	elevated -C3; and -C5-OH	Normal biotinidase studies	Untested or unknown	Untested or unknown
Holocarboy	Possible	Normal	elevated -C3; and -C5-OH	Normal biotinidase studies	1 known disease causing variant (Allele 1 - variant known to be disease causing)	Untested or unknown
	Possible	Normal	elevated -C3; and -C5-OH	Normal biotinidase studies	2 variants of uncertain significance in the same gene (Allele 1 - variant of unknown significance and Allele 2 - variant of unknown significance)	Untested or unknown

Isovaleric Acidemia (IVA)

	Classification	Urine organic acids	Plasma Acylcarnitine s	Mutation analysis	Enzyme analysis
	Definite	Unteste d or unknow n	Untested or unknown	2 known disease causing variants in the same gene (Allele 1 – variant known to be disease causing and Allele 2 – variant known to be disease causing)	Untested or unknown
	Definite	Unteste d or unknow n	Untested or unknown	Untested or unknown	enzyme activity on fibroblasts or WBCs consistent with disease
Isovaleric Acidmeia	Definite	Elevated - isoval eryl glycine and - 3-OH isovaleric	elevat ed C5	Untested or unknown	Untested or unknown
Isovaleri	Definite	Eleva ted isoval eryl glycin	elevat ed C5	2 variants of uncertain significance in the same gene - predicted to be pathogenic [Allele 1 - variant of unknown significance (predicted to be pathogenic) and Allele 2 – variant of unknown significance (predicted to be pathogenic)]	Untested or unknown
	Possible	Eleva ted isoval eryl	elevat ed C5	Untested or unknown	Untested or unknown
	Possible	Eleva ted isoval eryl	elevat ed C5	1 known disease causing variant (Allele 1 - variant known to be disease causing)	Untested or unknown
	Possible	Eleva ted isoval eryl	elevat ed C5	2 variants of uncertains ignificance in the same gene (Allele 1 - variant of unknown significance and Allele 2 – variant of unknown significance)	Untested or unknown

Glutaric Cidemia (GA)

	Classification	Urine or serum organic acids	Plasma Acylcarnitine s	Mutation analysis	Enzyme analysis
	Definite	Untested or unknown	Untested or unknown	2 known disease causing variants in the same gene (Allele 1 – variant known to be disease causing and Allele 2 – variant known to be disease causing)	Untested or unknown
	Definite	Untested or unknown	Untested or unknown	Untested or unknown	enzyme activity consistent with disease
l e l	Definite	Elevated - 3-OH Glutaric and - Glutaric	elevat ed C5 -DC	Untested or unknown	Untested or unknown
demiaTyp	Probable	Elevated - 3-OH Glutaric	elevat ed C5 –DC	Untested or unknown	Untested or unknown
Glutaric Acidemia Type I	Probable	Elevated glutaric	elevat ed C5 -DC	2 variants of uncertain significance in the same gene - predicted to be pathogenic [Allele 1 - variant of unknown significance (predicted to be pathogenic) and Allele 2 - variant of unknown significance (predicted to be pathogenic)]	Untested or unknown
	Possible	Elevated glutaric	elevat ed C5 -DC	Untested or unknown	Untested or unknown

Primary Carnitine Deficiency/ Carnitine Uptake Defect (CUD)

	Classification	Uri	Plasma	Special	Mutation analysis	Enzyme analysis
		ne	Carniti	Circumstance		,,
		Car	ne			
	Definite	Untested or unknown	Untested or unknown	Untested or unknown	2 known disease causing variants in the same gene (Allele 1 – variant known to be disease causing and Allele 2 – variant known to be disease causing)	Untested or unknown
	Definite	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	enzyme activity consistent with disease
	Definite	Elevat ed fractio nal	Low free carniti ne	Secondary carnitine loss ruled out	Untested or unknown	Untested or unknown
ne Uptake Defect	Probable	Untested or unknown	Low free carniti ne	Secondary carnitine loss ruled out	2 variants of uncertain significance in the same gene - predicted to be pathogenic [Allele 1 - variant of unknown significance (predicted to be pathogenic) and Allele 2 – variant of unknown significance (predicted to be pathogenic)]	Untested or unknown
y/ Carnitir	Possible	Untested or unknown	Low free carniti	Secondary carnitine loss ruled out	1 known disease causing variant (Allele 1 - variant known to be disease causing)	Untested or unknown
Primary Carnitine Deficiency/ Carnitine Uptake Defect	Possible	Untested or unknown	Low free carniti ne	Secondary carnitine loss ruled out	2 variants of uncertain significance in the same gene (Allele 1 - variant of unknown significance and Allele 2 – variant of unknown significance)	Untested or unknown
Primary C	Possible	Untested or unknown	Low free carniti ne	Secondary carnitine loss ruled out	None found	Untested or unknown
	Possible	Untested or unknown	Low free carniti ne	Secondary carnitine loss ruled out	Untested or unknown	Untested or unknown

Very long-chain acyl-CoA Dehydrogenase Deficiency (VLCAD)

	Classification	Plasma Acylcarnitines	Mutation analysis	Functional Studies
	Definite	Untested or unknown	2 known disease causing variants in the same gene (Allele 1 – variant known to be disease causing and Allele 2 – variant known to be disease causing)	Untested or unknown
	Definite	Untested or unknown	Untested or unknown	Functional fibroblast or Enzyme analysis consistent with VLCAD
	Definite	Elevated -C14:1 (on more than one sample) and -C14:2 and -C14	Untested or unknown	Untested or unknown
	Definite	Elevated -C14:1 (on more than one sample)	2 variants of uncertain significance in the same gene - predicted to be pathogenic [Allele 1 - variant of unknown significance (predicted to be pathogenic) and Allele 2 – variant of unknown significance (predicted to be pathogenic)]	Untested or unknown
VLCAD	Probable	Elevated -C14:1 (on more than one sample) and -C14:2	2 variants of uncertain significance in the same gene (Allele 1 - variant of unknown significance and Allele 2 – variant of unknown significance)	Untested or unknown
	Probable	Elevated -C14:1 (on more than one sample)	2 variants of uncertain significance in the same gene (Allele 1 - variant of unknown significance and Allele 2 – variant of unknown significance)	Untested or unknown
	Probable	Elevated -C14:1 (on more than one sample) and -C14:2	1 known disease causing variant (Allele 1 - variant known to be disease causing)	Untested or unknown
	Possible	Elevated -C14:1 (on more than one sample) and -C14:2	Untested or unknown	Untested or unknown

Very long-chain acyl-CoA Dehydrogenase Deficiency (VLCAD)

	Classification	Plasma Acylcarnitines	Mutation analysis	Functional Studies
VLCAD	Possible	Elevated -C14:1 (on more than one sample) and -C14:2	No variants found	Untested or unknown
	Possible	Elevated C14:1 on more than one sample	Untested or unknown	Untested or unknown
	Possible	Elevated C14:1 on more than one sample	No variants found	Untested or unknown
	Possible	Elevated C14:1 on more than one sample	1 known di sease causing variant (Allele 1 - variant known to be di sease causing)	Untested or unknown

Trifunctional Protein Deficiency (TFP)-Inclusive of LCHAD

	Classification	Urine Organics	Plasma Acylcarnitines	Mutation analysis	Functional Studies
	Definite	Untested or unknown	Untested or unknown	2 known disease causing variants in the same gene (Allele 1 variant known to be disease causing and Allele 2 – variant known to be disease causing)	Untested or unknown
	Definite	Untested or unknown	Untested or unknown	Untested or unknown	Functional fibroblast or Enzyme analysis consistent with LCHAD or TFP
Inclusive of LCHAD	Definite	Untested or unknown	Elevated: -C16-OH (on more than one specimen) and -C16:1-OH and -C18-OH and -C18:1-OH	2 variants of uncertain significance in the same gene - predicted to be pathogenic [Allele 1 - variant of unknown significance (predicted to be pathogenic) and Allele 2 - variant of unknown significance (predicted to be pathogenic)]	Untested or unknown
	Probable	Elevated -C12-OH dicarboxylic and -C10-OH dicarboxylic	Elevated: -C16-OH (on more than one specimen)and -C16:1-OH and -C18-OH and -C18:1-OH	Untested or unknown	Untested or unknown
	Probable	Untested or unknown	Elevated: -C16-OH (on more than one specimen) and -C16:1-OH and -C18-OH and -C18:1-OH	1 known disease causing variant (Allele 1 - variant known to be disease causing)	Untested or unknown
ifunctional Protein Deficiency-Inclusive of LCHAD	Probable	Untested or unknown	Elevated: -C16-OH (on more than one specimen) and -C16:1-OH and -C18-OH and -C18:1-OH	2 variants of uncertain significance in the same gene (Allele 1 - variant of unknown significance and Allele 2 – variant of unknown significance)	Untested or unknown
Trifunctional I	Possible	Untested or unknown	Elevated: -C16-OH (on more than one specimen) and -C16:1-OH and -C18-OH and -C18:1-OH	No variants found	Untested or unknown
	Possible	Untested or unknown	Elevated: -C16-OH (on more than one specimen) and -C16:1-OH and -C18-OH and -C18:1-OH	Untested or unknown	Untested or unknown
	Possible	Elevated -C12-OH dicarboxylic and -C10-OH dicarboxylic	Untested or unknown	Untested or unknown	Untested or unknown

Trifunctional Protein Deficiency (TFP)-Inclusive of LCHAD

	Classification	Urine Organics	Plasma Acylcarnitines	Mutation analysis	Functional Studies
al Protein Deficiency- : LCHAD	Possible	Elevated -C12-OH dicarboxylic and -C10-OH dicarboxylic	Untested or unknown	1 known disease causing variant (Allele 1 - variant known to be disease causing)	Untested or unknown
Trifunctional Inclusive of LC	Possible	Untested or unknown	Elevated: -C16-OH (on more than one specimen)	1 known disease causing variant (Allele 1 - variant known to be disease causing)	Untested or unknown

Argininosuccinic Aciduria (ASA)

	Classification	Plasma or urine amino acids	Mutation analysis	Enzyme Studies
Argininosuccinic Aciduria (ASA)	Definite	Untested or unknown	2 known disease causing variants in the same gene (Allele 1 – variant known to be disease causing and Allele 2 – variant known to be disease causing)	Untested or unknown
	Definite	Untested or unknown	Untested or unknown	Enzyme analysis consistent with disease
	Definite	Elevated -ASA and -Citrulline	Untested or unknown	Untested or unknown
	Definite	El eva ted ASA	2 variants of uncertain significance in the same gene - predicted to be pathogenic [Allele 1 - variant of unknown significance (predicted to be pathogenic) and Allele 2 – variant of unknown significance (predicted to be pathogenic)]	Untested or unknown
	Possible	Elevated Citrulline	1 known di sease causing variant (Allele 1 - variant known to be di sease causing)	Untested or unknown
	Possible	El eva ted Citrulline	2 variants of uncertain significance in the same gene (Allele 1 - variant of unknown significance and Allele 2 – variant of unknown significance)	Untested or unknown

Citrullinemia, type 1-Exclusive of Citrin Deficiency (CIT)

	Classification	Plasma amino acids	Blood Ammonia Levels	Mutation analysis	Enzyme Studies
	Definite	Untested or unknown	Untested or unknown	2 known disease causing variants in the same gene (Allele 1 – variant known to be disease causing and	Untested or unknown
sucy	Definite	Untested or unknown	Untested or unknown	Untested or unknown	Enzyme analysis consistent with
Citrullinemia Type I- exclusive of Citrin deficiency	Definite	El evated Citrulline and Absent ASA	Untested or unknown	2 variants of uncertain significance in the same gene - predicted to be pathogenic [Allele 1 - variant of unknown significance (predicted to be pathogenic) and	Untested or unknown
cdusive	Definite	El evated Citrulline and Absent ASA	Elevated	Untested or unknown	Untested or unknown
pe l- e)	Probable	Elevated Citrulline and Absent ASA	Untested or unknown	1 known disease causing variant (Allele 1 - variant known to be	Untested or unknown
inemia Ty _l	Probable	Elevated Citrulline and Absent ASA	Untested or unknown	2 variants of uncertain significance in the same gene (Allele 1 - variant of unknown significance and Allele 2 –	Untested or unknown
Citrul	Possible	Elevated Citrulline and Absent ASA	Untested or unknown	Untested or unknown	Untested or unknown

3-Methylcrotonyl-CoA (3-MCC)

	S internylationally control interp						
	Classificati on	Urine organic acids	Plasma Acylcarnitines	Maternal Studies	Mutation analysis	Enzyme analysis	
	Definite	Untested or unknown	Untested or unknown	Untested or unknown	2 known disease causing variants in the same gene (Allele 1 – variant known to be disease causing and Allele 2 – variant known to be disease causing)	Untested or unknown	
	Definite	Untestedorunknown	Untested or unknown	Untested or unknown	Untested or unknown	enzyme activity consistent with disease	
U	Definite	Elevated 3-OH Isovaleric with or without elevated 3-methylcrotonyl glycine	elevated C5 -OH	Maternal deficiency tested and ruled out	2 variants of uncertain significance in the same gene - predicted to be pathogenic [Allele 1 - variant of unknown significance (predicted to be pathogenic) and Allele 2 – variant of unknown significance (predicted to be pathogenic)]	Untested or unknown	
3-MCC	Probable	Elevated 3-OH Isovaleric with or without elevated	elevated C5 -OH	Maternal deficiency tested and ruled out	2 variants of uncertain significance in the same gene (Allele 1 - variant of unknown significance and Allele 2 - variant of unknown significance)	Untested or unknown	
	Probable	Elevated - 3-OH Isovalericand - 3- methylcroton	elevated C5 -OH	Maternal deficiency tested and ruled out	Untested or unknown	Untested or unknown	
	Possible	Elevated - 3-OH Isovalericand - 3- methylcroton	Untested or unknown	Maternal deficiency tested and ruled out	Untested or unknown	Untested or unknown	
	Possible	Untested or unknown	elevated C5 –OH	Maternal deficiency tested and ruled out	Untested or unknown	Untested or unknown	

Tyrosinemia, Type 1 (TYR 1)

	Classification	Urine or Serum studies	Mutation analysis	Enzyme Studies
	Definite	Untested or unknown	2 known disease causing variants in the same gene (Allele 1 – variant known to be disease causing and Allele 2 – variant known to be disease causing)	Untested or unknown
	Definite	Untested or unknown	Untested or unknown	Enzyme analysis consistent with FAH enzyme deficiency
Tyrosinemia type I	Definite	El eva ted Succinylacetone	Untested or unknown	Untested or unknown
	Possible	El evated tyrosine and Normal Succinylacetone	2 variants of uncertain significance in the same gene - predicted to be pathogenic [Allele 1 - variant of unknown significance (predicted to be pathogenic) and Allele 2 – variant of unknown significance (predicted to be pathogenic)]	Untested or unknown
	Possible	Elevated Tyrosine and Normal Succinylacetone	1 known disease causing variant (Allele 1 - variant known to be disease causing)	Untested or unknown
	Possible	Elevated Tyrosine and Normal Succinylacetone	2 variants of uncertain significance in the same gene (Allele 1 - variant of unknown significance and Allele 2 – variant of unknown significance)	Untested or unknown

Medium-chain acyl-CoA Dehydrogenase Deficiency (MCAD)

	Classification	Urine Organics or	Plasma	Mutation analysis	Functional Studies
		aclyglycines	Acylcarnitines	,	
	Definite	Untested or unknown	Untested or unknown	2 known disease causing variants in the same gene (Allele 1 – variant known to be disease causing and Allele 2 – variant known to be disease causing)	Untested or unknown
	Definite	Untested or unknown	Untested or	Untested or unknown	Functional fibroblast or
	Definite	Elevated hexanoylglycine	Elevated: -C8 and -C8>C10 and -C8>C6 and -C6 and -C10	Untested or unknown	Untested or unknown
	Definite	Untested or unknown	Elevated: -C8 and -C8>C10 and -C8 >C6 and -C6 and -C10	2 variants of uncertain significance in the same gene - predicted to be pathogenic [Allele 1 - variant of unknown significance (predicted to be pathogenic) and Allele 2 – variant of unknown significance (predicted to be pathogenic)]	Untested or unknown
MCAD	Probable	Untested or unknown	Elevated C8 on repeat testing	1 known disease causing variant and 1 variants of uncertain significance in the same gene (Allele 1 - variant known to be disease causing and Allele 2 - variant of unknown significance)	Untested or unknown
	Probable	Elevated hexanoylglycine	Elevated C8 on repeat testing	1 known disease causing variant (Allele 1 - variant known to be disease causing)	Untested or unknown
	Probable	Untested or unknown	Elevated C8 on repeat testing	2 variants of uncertain significance in the same gene (Allele 1 - variant of unknown significance and Allele 2 – variant of unknown significance)	Untested or unknown
	Possible	Elevated hexanoylglycine	Elevated C8 on repeat testing	No variants found	Untested or unknown
	Possible	Elevated hexanoylglycine	Untested or unknown	2 variants of uncertain significance in the same gene (Allele 1 - variant of unknown significance and Allele 2 – variant of unknown significance)	Untested or unknown
	Possible	Elevated Hexanoylglycine	Untested or unknown	No variants found	Untested or unknown
	Possible	Untested or unknown	Elevated C8 on repeat testing	No variants found	Untested or unknown

Possible or Carrier	Untested or unknown	El evated C8	1 known disease causing variant (Allele 1 - variant known to be disease causing)	Untested or unknown
Possible or Carrier	Elevated <i>Hexanoylglycine</i>	Normal	1 known disease causing variant (All ele 1 - variant known to be disease causing)	Untested or unknown

Maple Syrup Urine Disease (MSUD)

		Plasma amino	Urine Organic acids	Mutation analysis	Enzyme Studies
	on	acids			
	Definite	Untested or unknown	Untested or unknown	2 known disease causing variants in the same gene (Allele 1 – variant known to be disease causing and Allele 2 – variant known to be disease causing)	Untested or unknown
	Definite	Untested or unknown	Untested or unknown	Untested or unknown	Enzyme analysis consistent with MSUD
MSUD	Definite	Elevated Alloisoleucine and Leu, and Val, and Ileu	Untested or unknown	Untested or unknown	Untested or unknown
	Definite	Elevated Alloisoleucine	Untested or unknown	2 variants of uncertain significance in the same gene - predicted to be pathogenic [Allele 1 - variant of unknown significance (predicted to be pathogenic) and Allele 2 – variant of unknown significance (predicted to be pathogenic)]	Untested or unknown
	Probable	Elevated Alloisoleucine	Untested or unknown	Untested or unknown	Untested or unknown
	Probable	Elevated Leu and Ile and Val and Leu>Val	Elevated 2- ketoisocaproi c acid and 2-OH Isovaleric and 2-ketomethyl valeric acid	Untested or unknown	Untested or unknown
	Possible	Elevated Leu and Ile and Val and Leu>Val	Untested or unknown	Untested or unknown	Untested or unknown

Cystathionine Beta-Synthase (CBS) Deficiency

	Classificati on	Plasma amino acids	Mutation analysis	Enzyme Studies
	Definite	Untested or unknown	2 known disease causing variants in the same gene (Allele 1 – variant known to be disease causing and Allele 2 – variant known to be disease causing)	Untested or unknown
	Definite	Untested or unknown	Untested or unknown	Enzyme analysis consistent with CBS
CBS deficiency	Definite	Elevated -Methionine and -Homocystine	2 variants of uncertain significance in the same gene - predicted to be pathogenic [Allele 1 - variant of unknown significance (predicted to be pathogenic) and Allele 2 — variant of unknown significance (predicted to be pathogenic)]	Untested or unknown
	Probable	Elevated -Methionine and -Homocystine	2 variants of uncertain significance in the same gene (Allele 1 - variant of unknown significance and Allele 2 – variant of unknown significance)	Untested or unknown
	Probable	Elevated -Methionine and -Homocystine	1 known disease causing variant and 1 variant of uncertain significance in the same gene (Allele 1 - variant known to be disease causing and Allele 2 - variant of unknown significance)	Untested or unknown
	Possible	Elevated -Methionine and -Homocystine	Untested or unknown	Untested or unknown

Benign Hyperphenylalaninemia (H-PHE)

	Classificati on	Plasma amino acids	Special Studies	Mutation analysis	Enzyme Studies
HyperPhe	Definite	Untested or unknown	Untested or unknown	2 known di sease causing variants in the same gene (Allele 1 – variant known to be disease causing and Allele 2 – variant known to be di sease	Untested or unknown
	Definite	Untested or unknown	Untested or unknown	Untested or unknown	Enzyme analysis consistent with PAH
	Definite	Elevated Phe (>120umol/L on unrestricted diet) and Phe/Tyr ratio	Normal biopterin studies	Untested or unknown	Untested or unknown
	Possible	Elevated Phe (>120umol/L on unrestricted diet) and Phe/Tyr ratio	Untested or unknown	Untested or unknown	Untested or unknown

Biotinidase Deficiency (BIOT)

	Disorder	Classification	Enzyme Levels	Mutation analysis
	Profound	Definite	Untested or unknown	2 variants known to be associated with profound enzyme deficiency in the same gene (Allele 1 – variant known to be associated with profound enzyme deficiency and Allele 2 – variant known to be associated with profound enzyme deficiency)
e Deficiency	Partial Definite Untested or unknown			1 variant known to be associated with profound enzyme deficiency and 1 Known to be associated with partial enzyme deficiency ['mild' mutation]
Biotinidase	Partial	Definite	Untested or unknown	2 variants known to be associated with partial enzyme deficiency ['mild' mutation] (Allele 1 and allele – variant known to be associated with partial enzyme deficiency ['mild' mutation (D444H)]
	Profound	Probable	<10% normal activity	Untested or unknown
	Partial	Probable	10-30% normal activity	Untested or unknown

Classic Galactosemia (GALT)

	Classificati on	GALT Levels	Gal-1-P level	Urine Galactitol	Mutation analysis
	Definite				2 known disease causing variants in the same gene (Allele 1 – variant known to be disease causing and Allele 2 – variant known to be disease causing)
ctosemia	Definite		Elevated		2 variants of uncertain significance in the same gene - predicted to be pathogenic [Allele 1 - variant of unknown significance (predicted to be pathogenic) and Allele 2 – variant of unknown significance (predicted to be pathogenic)]
Classic Galactosemia	Definite			Elevated	2 variants of uncertain significance in the same gene - predicted to be pathogenic [Allele 1 - variant of unknown significance (predicted to be pathogenic) and Allele 2 – variant of unknown significance (predicted to be pathogenic)]
	Definite		Elevated		2 variants of uncertain significance in the same gene [Allele 1 - variant of unknown significance and Allele 2 – variant of unknown significance]
	Definite			Elevated	2 variants of uncertain significance in the same gene [Allele 1 - variant of unknown significance and Allele 2 – variant of unknown significance]
	Definite		Elevated		1 known disease causing mutation and 1 mutation of uncertain significance gene (All ele 1 – variant known to be disease causing and Allele 2 – and Allele 2 – variant of unknown significance)
	Definite			Elevated	1 known disease causing mutation and 1 mutation of uncertain significance gene (All ele 1 – variant known to be disease causing and Allele 2 – and Allele 2 – variant of unknown significance)
	Probable		Elevated		
	Probable			Elevated	
	Probable				1 known disease causing mutation (Allele 1 – variant known to be disease causing)
	Probable				2 variants of uncertain significance in the same gene - predicted to be pathogenic [Allele 1 - variant of unknown significance (predicted to be pathogenic) and Allele 2 – variant of unknown significance (predicted to be pathogenic)]
	Probable	<10%			2 variants of uncertain significance in the same gene [Allele 1 - variant of unknown significance and Allele 2 – variant of unknown significance]
	Probable	<10%			

Variant Galactosemia

	Classification	GALT Levels	Gal-1-P level	Urine Galactitol	Mutation analysis	Protein Phenotyping
Biotinidase Deficiency	Definite				1 known classic galactosemia disease causing mutation and 1	
	Definite	10%-30%	Elevated		1 known disease causing mutation and 1 mutation of uncertain significance-	
	Definite 10%-30			Elevated	1 known disease causing mutation and 1 mutation of uncertain significance-	
	Definite	10%-30%	Elevated			phenotype consistent with variant
	Definite	10%-30%		Elevated		phenotype consistent with variant
	Definite	10%-30%	Elevated		1 known disease causing mutation and 1 mutation of uncertain significance	
	Definite	10%-30%		Elevated	1 known disease causing mutation of	
	Probable	10%-30%				phenotype consistent with variant
	Possible	10%-30%				

Arginase Deficiency

	Classification	Plasma amino acids	Mutation analysis	Enzyme Studies
Deficiency	Definite		2 known disease causing variants in the same gene (Allele 1 – variant known to be disease causing and Allele 2 – variant known to be disease causing)	
Arginase I	Definite			Enzyme a nalysis consistent with Arginase deficiency
Arı	Probable	Elevated Arginine	1 known disease causing mutation	
	Possible	Elevated Arginine		

Endocrinology Disorders Case Definitions Tables September 29, 2013

Definitions created by panel of experts between June 2011 and September 2013. This project was funded in part by Cooperative Agreement # U22MC24078 from the Health Resources and Services Administration (HRSA).

Primary Congenital Hypothyrodism (CH)

	Category	Serum TSH mU/L*	Serum Total or Free T4*		
	Definite	TSH > 10	< age established reference range		
roidism	Probable	TSH > 10	normal T4/total T4		
thy	Probable	TSH > 10	Untested or unknown		
Primary Congenital Hypothyroidism	Possible**	TSH 6-10	< age established reference range		
gen	Possible **	TSH 6-10	Normal		
Con	Possible **	TSH 6-10	Untested or unknown		
ary	Incomplete	Untested or unknown	Untested or unknown		
Prim	Incomplete	Untested or unknown	< age established reference		

Secondary Congenital Hypothyroidism (CH)

	coolinary congernarity conference (cry								
	Category	Serum TSH mU/L*	Serum Total or Free T4*	Other studies					
	Definite	TSH < 10	< age established reference	documentation of other pituitary hormone deficiencies or midline defects					
idism	Probable**	TSH < 10	< age established reference range	no other pituitary hormone deficiencies or midline defects					
Secondary Congenital Hypothyroidism	Possible Untested or unknown		< age established reference range	Documentation of other pituitary hormone deficiencies or midline defects					
çenital H _y	Possible	TSH<10	Untested or unknown	Documentation of other pituitary hormone deficiencies or midline defects					
lary Cong	Incomplete	Untested or unknown	Untested or unknown	Documentation of other pituitary hormone deficiencies or midline defects					
Second	Incomplete TSH<10		Untested or unknown	no other pituitary hormone deficiencies or midline defects					
	Incomplete	Untested or unknown	< age established reference range	no other pituitary hormone deficiencies or midline defects					

Thyroxine-binding Globulin (TBG) or other Protein Binding Defect

ct	Category	Serum TSH mU/L	Serum Free T4	Serum Total T4	Other studies
other Protein Binding Defect	Definite	normal	Normal for age	Low for age	LowTBG
TBG or other	Definite	normal	Normal for age	Lowforage	increased T3 or T4 resin uptake

 $^{^{*}}$ The results referenced should be obtained before the initiation of the rapy.

^{**} Since there can be overlap in these 2 categories (possible primary or probable secondary congenital hypothyroidism) based on the laboratory values, the treating clinician should determine which category.

21-Hydroxylase Deficiency – Classical Salt Wasting

	Catagory	Serum 17-	Urinarystaraidnyafilina	Ser	Plas	CYP21A2 Mutation	If
	Category		Urinary steroid profiling	um	ma	Analysis	available
		ОНР		Sod	Reni	Allalysis	-
				iu			Supporti
		baseline			n Activ		ve
		or ACTH		m 			Clinical
		stimulated*		mE ~/!	ity		
				q/L			or
							Laborato
							ry
-	- 0 11	10.000					Evidence
	Definite	> 10,000	Untested or unknown	< 135	Untestedor	Untestedor	Evidence of salt
					unknown	unknown	wasting (present in shock or
							severe failure to
							thrive)
	D - C - : : -	. 10.000	Haladada alama	<135	Untestedor	Untestedor	·
	Definite	> 10,000	Untested or unknown	133	unknown	unknown	ambiguous genitalia in
_							46, XX
	Definite	> 10,000	Untested or unknown	<135	Untestedor	Untestedor	other hormonal
					unknown	unknown	evidence of CAH
-				Untested	Elevated for	Untestedor	Evidence of salt
	Definite	> 10,000	Untested or unknown	or	age	unknown	wasting (present
ഫ				unknown	~Bc		in shock or
ţį							severe failure to
Vas							thrive)
<u>+</u>	Definite	> 10,000	Untested or unknown	Untested	Elevated for	Untestedor	ambiguous genitalia in
Sa	Dellilite	> 10,000	Ontested of unknown	or	age	unknown	
cal				unknown			46, XX
roxylase Deficiency – Classical Salt Wasting	Definite	> 10,000	Untested or unknown	Untested	Elevated for	Untestedor	other hormonal
Ö				or	age	unknown	evidence of CAH
\ \ \				unknown			
enc	Definite	Unteste	Untested or unknown	Untested	Untested or	two classic	Evidence of salt
<u>ij</u>		d or		or	unknown	genemutations or	wasting (present
De		unknow n		unknown		deletions <i>intrans</i>	in shock or severe failure to
Se		"					thrive)
- Ala		Unteste		Untested	Untested or	two classic gone	•
<u> </u>	Definite	d or	Untested or unknown		unknown	two classic gene mutations or deletions <i>in</i>	ambiguous genitalia in
<u> </u>		unknow		or unknown	ulikilowii	trans	46, XX
21-Hyd	_	Unteste		Untested	Untested or		other hormonal
2:	Definite	d or	Untested or unknown	or	unknown	two classic gene mutations or deletions <i>in</i>	evidence of CAH
		unknow		unknown	anknown	trans	CVIDENCE OF CALL
		Unteste	(mass spectrometry)	Untested	Untested or	Untestedor	Evidence of salt
	Definite	d or	indicative of 21-	or	unknown	unknown	wasting (present
		unknow	Hydroxylase Deficiency	unknown	anknown	ananown	in shock or
		n	CAH				severe failure to
							thrive)
	Dofinito	Unteste	(mass spectrometry)	Untested	Untestedor	Untestedor	ambiguous ganitaliaia
	Definite	d or	indicative of 21-	or	unknown	unknown	ambiguous genitalia in
		unknow	Hydroxylase Deficiency	unknown			46, XX
		n	CAH				

21-Hydroxylase Deficiency – Classical Salt Wasting

	,		<u> </u>			avaa.	
	Category	Serum 17-OHP - baseline or ACTH stimulated*	Urinary steroid profiling	Seru m Sodiu m mEq/ L	Plasma Renin Activity	CYP21A2 Mutation Analysis	If available - Supporti ve Clinical or Laborato ry Evidence
	Definite	Untested or unknown	(mass spectrometry) indicative of 21- Hydroxylase Deficiency CAH	Untested or unknown	Untested or unknown	Untested or unknown	other hormonal evidence of CAH
Salt Wasting	Probable	1,000 -10,000	Untested or unknown	<135	Untested or unknown	Untested or unknown	Evidence of salt wasting (presentin shock or severe failure to thrive)
y – Classica	Probable	1,000 -10,000	Untested or unknown	<135	Untested or unknown	Untested or unknown	ambiguous genitalia in 46,XX
e Deficienc	Probable	1,000 -10,000	Untested or unknown	< 135	Untested or unknown	Untested or unknown	other hormonal evidence of CAH
21-Hy21-Hydroxylase Deficiency – Classical Salt Wasting	Probable	1,000 -10,000	Untested or unknown	Untested or unknown	Elevated for age	Untested or unknown	Evidence of salt wasting (presentin shock or severe failure to thrive)
21	Possible	1,000 -10,000	Untested or unknown	Untested or unknown	Elevated for age	Untested or unknown	ambiguous genitalia in 46,XX
	Possible	1,000 -10,000	Untested or unknown	Untested or unknown	Elevated for age	Untested or unknown	other hormonal evidence of CAH

21-Hydroxylase Deficiency-Classical Simple Virilizing

	Category		Urinary Steroid profiling	Serum Sodium mEq/L	Plasma Renin Activity	CYP21 A2 Mutat ion Analy sis	If available - Supportive Clinical or Laboratory Evidence
	Definite	>10,000	Untested or unknown	>135	Untested or unknown	Untested or unknown	Ambiguous genitalia in 46,XX
	Definite	>10,000	Untested or unknown	>135	Untested or unknown	Untested or unknown	no evidence of salt wasting
firilizing	Definite	>10,000	Untested or unknown	>135	Untested or unknown	Untested or unknown	other hormonal evidence of CAH
al Simple Vi	Definite	>10,000	Untested or unknown	Untested or unknown	Normal for age	Untested or unknown	Ambiguous genitalia in 46,XX
- Classica	Definite	>10,000	Untested or unknown	Untested or unknown	Normal for age	Untested or unknown	no evidence of salt wasting
iciency -	Definite	>10,000	Untested or unknown	Untested or unknown	Normal for age	Untested or unknown	other hormonal evidence of CAH
21-Hydroxylase Deficiency – Classical Simple Virilizing	Definite	Untested or unknown	(mass spectrometry) indicative of 21- Hydroxylase Deficiency CAH	>135	Untested or unknown	Untested or unknown	Ambiguous genitalia in 46,XX
21-Hydı	Definite	Untested or unknown	(mass spectrometry) indicative of 21- Hydroxylase Deficiency CAH	>135	Untested or unknown	Untested or unknown	no evidence of salt wasting
	Definite	Untested or unknown	(mass spectrometry) indicative of 21- Hydroxylase Deficiency CAH	>135	Untested or unknown	Untested or unknown	other hormonal evidence of CAH
	Definite	Untested or unknown	(mass spectrometry) indicative of 21- Hydroxylase Deficiency CAH	Untested or unknown	Normal for age	Untested or unknown	Ambiguous genitalia in 46,XX

21-Hydroxylase Deficiency-Classical Simple Virilizing

С	Category	Serum 17-OHP - baseline or ACTH stimulated*	Urinary Steroid profiling	Serum Sodiu m mEq/L	Plasma Renin Activity	CYP21A2 Mutation Analysis	If available - Supportive Clinical or Laboratory Evidence
D	Definite	Untested or unknown	(mass spectrometry) indicative of 21- Hydroxylase Deficiency CAH	Untested or unknown	Normal for age	Untested or unknown	no evidence of salt wasting
D	Definite	Untested or unknown	(mass spectrometry) indicative of 21- Hydroxylase Deficiency CAH	Untested or unknown	Normal for age	Untested or unknown	other hormonal evidence of CAH
D	Definite	Untested or unknown	Untested or unknown	>135	Untested or unknown	two classic gene mutations or deletions in trans	Ambiguous genitaliain 46,XX
D	Definite	Untested or unknown	Untested or unknown	>135	Untested or unknown	two classic gene mutations or deletions in trans	no evidence of salt wasting
D	Definite	Untested or unknown	Untested or unknown	>135	Untested or unknown	two classic gene mutations or deletions in trans	other hormonal evidence of CAH
D	Definite	Untested or unknown	Untested or unknown	Untested or unknown	Normal for age	two classic gene mutations or deletions in trans	Ambiguous genitaliain 46,XX
D	Definite	Untested or unknown	Untested or unknown	Untested or unknown	Normal for age	two classic gene mutations or deletions in trans	no evidence of salt wasting
D	Definite	Untested or unknown	Untested or unknown	Untested or unknown	Normal for age	two classic gene mutations or deletions in trans	other hormonal evidence of CAH

Probable	1,000 -10,000	Untested or unknown	>135	Untested or unknown	Untested or unknown	Ambiguous genitalia in 46,XX or normal genitalia in 46,XY
Probable	1,000 -10,000	Untested or unknown	Untested or unknown	Normal for age	Untested or unknown	Ambiguous genitalia in 46,XX or normal genitalia in 46,XY
Probable	1,000 -10,000	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	no evidence of salt wasting

Alpha thalassemia Case Definitions Tables September 29, 2013

Definitions created by panel of experts between June 2011 and September 2013. This project was funded in part by Cooperative Agreement # U22MC24078 from the Health Resources and Services Administration (HRSA).

Throughout this document, the following definitions are used:

- 1. Family studies both parents with HPLC, IEF outside of newborn period and CBC if microcytosis –low MCH, MCV (assuming iron deficiency has been ruled out and A2 is not elevated, then presumptive alpha thal trait) OR DNA
- 2. Family history includes reported history of Hgb variant in the family

S Alpha Thal

Category	Qualitative (IEF or HPLC)	Quantitati ve (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history	HPLC& IEF same sample
Definite	FS+Barts	Untested or unknown	Homozygous S mutation and pathological gene changes found in 1-3 of the alpha genes	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FS+Barts	Homozygous S mutation and pathological gene changes found in 1-3 of the alpha genes	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	Homozygous S mutation and pathological gene changes found in 1-3 of the alpha genes	FS + Barts	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FS+Barts	FS+Barts	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FS+Barts	Untested or unknown	Untested or unknown	FS+Barts	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FS+Barts	Untested or unknown	FS+Barts	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FS+Barts x2	Untested or unknown	Untested or unknown	FS+Barts	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown

Probable	Untested or unknown	Untested or unknown	Untested or unknown	FS+Barts	Low MCV	Both parents with AS & amount of S <35%; low MCH & ruled out iron deficiency	Untested or unknown	Untested or unknown
Probable	FS+Barts	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Both parents with AS & amount of S <35%; low MCH & ruled out iron deficiency	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FS+Barts	Untested or unknown	Untested or unknown	Low MCV	Both parents wh AS & amount of S <35%; low MCH & ruled out iron deficiency	Untested or unknown	Untested or unknown

S Alpha Thal

Category	Qualitative (IEF or HPLC)	Quantitati ve (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history	HPLC& IEF same sample
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FS+Barts	Low MCV	Untested or unknown	positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Untested or unknown	Untested or unknown	FS+Barts

C Alpha Thal

Category	Qualitative (IEF or HPLC)	Quantitativ e (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history	HPLC& IEF same sample
Definite	FC+Barts	Untested or unknown	Known C mutation and Deletion in alpha gene	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FC+Barts	Known C mutation and Deletion in alpha gene	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	Known C mutation and Deletion in alpha gene	FC+Barts	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FC+Barts	FC+Barts	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FC+Barts	Untested or unknown	Untested or unknown	FC+Barts	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FC+Barts	Untested or unknown	FC+Barts	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FC+Barts	Untested or unknown	Untested or unknown	FC+Barts	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	Untested or unknown	FC+Barts	Low MCV	Both carriers	Untested or unknown	Untested or unknown
Probable	FC+Barts	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Both carriers	Untested or unknown	Untested or unknown

Probable	Untested or unknown	FC+Barts	Untested or unknown	Untested or unknown	Low MCV	Both Carriers	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FC+Barts	Low MCV	Untested or unknown	positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Untested or unknown	Untested or unknown	FC+Barts

D Alpha Thal

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history	HPLC& IEF same sample
Definite	FD+Barts	Untested or unknown	Known C mutation and Deletion in alpha gene	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FD+Barts	Known C mutation and Deletion in alpha gene	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	Known C mutation and Deletion in alpha gene	FD + Barts	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FD+Barts	FD+Barts	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FD+Barts	Untested or unknown	Untested or unknown	FD+Barts	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FD+Barts	Untested or unknown	FD+Barts	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	Untested or unknown	FD+Barts	Low MCV	Both carriers	Untested or unknown	Untested or unknown
Probable	FD+Barts	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Both carriers	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FD+Barts	Untested or unknown	Untested or unknown	Low MCV	Both Carriers	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FD+Barts	Low MCV	Untested or unknown	positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Untested or unknown	Untested or unknown	FD+Barts

O_{Arab} Alpha Thal

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history	HPLC& IEF same sample
Definite	FOARAB+Barts	Untested or unknown	Known O _{Arab} mutation and Deletion in alphagene	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FOARAB+Barts	Known OArab mutation and Deletion in alpha gene	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	Known OArab mutation and Deletion in alphagene	FOARAB+Barts	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FOARAB+Barts	Untested or unknown	FOARAB+Barts	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FOARABTraceAA2	FOARAB+Barts	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FOARAB+Barts	Untested or unknown	Untested or unknown	FOARAB+Barts	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	Untested or unknown	FOARAB+Barts	Low MCV	Both carriers	Untested or unknown	Untested or unknown
Probable	FOARAB+Barts	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Both carriers	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FOARAB+Barts	Untested or unknown	Untested or unknown	Low MCV	Both Carriers	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FOARAB+Barts	Low MCV	Untested or unknown	positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Untested or unknown	Untested or unknown	FOARAB+Barts

3 Deletion Alpha Thalassemia (Hgb H disease)

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA-based testing	NBS result	CBC Results	Family DNA Studies	Family history	HPLC& IEF same sample
Definite	Untested or unknown	≥25% Barts by HPLC in newborn period	3 alphagene defects (deletions or mutations)	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	3 alphagene defects (deletions or mutations)	Barts or Hgb H	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	Untested or unknown	Barts or Hgb H	Low MCV	Parents with known carriers of 2 gene deletion and 1 gene deletion or point mutation	History of SAB/miscarriage or early termination of pregnancy	Untested or unknown
Probable	Persistent Barts	Untested or unknown	Untested or unknown	Barts or Hgb H	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Persistent Barts	Untested or unknown	Barts or Hgb H	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Elevated Hgb H	Untested or unknown	Untested or unknown	Barts or Hgb H	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	El evated Hgb H	Untested or unknown	Barts or Hgb H	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Possible	Nml	Untested or unknown	Untested or unknown	Barts or Hgb H	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Nml	Untested or unknown	Barts or Hgb H	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown

Hgb H Constant Spring (2 alpha gene deletion (cis) plus Constant Spring point mutation (trans))

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family DNA Studies	Family history	HPLC& IEF same sample
Definite	Constant Spring band identified	Untested or unknown	3 alpha gene deletions and Constant spring mutation	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Constant Spring band identified	3 alpha gene deletions and Constant spring mutation	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	3 alpha gene deletions and Constant spring mutation	Barts or Hgb H	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Constant Spring band identified	Untested or unknown	Untested or unknown	Barts or Hgb H	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Constant Spring band identified	Untested or unknown	Barts or Hgb H	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	Untested or unknown	Barts or Hgb H	Low MCV	Parents with known carriers of 2 gene and 1 gene deletion and one with Constant Spring mutation	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Constant Spring band identified	Untested or unknown	Untested or unknown	Untested or unknown	Parents with known carriers of 2 gene and 1 gene deletion and one with Constant Spring mutation	Untested or unknown	Untested or unknown
Probable	Constant Spring band identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Parents with known carriers of 2 gene and 1 gene deletion and one with Constant Spring mutation	Untested or unknown	Untested or unknown
Possible	Nml	Untested or unknown	Untested or unknown	Barts or Hgb H	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Nml	Untested or unknown	Barts or Hgb H	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown

Beta Thalassemia Case Definitions Tables September 29, 2013

Definitions created by panel of experts between June 2011 and September 2013. This project was funded in part by Cooperative Agreement # U22MC24078 from the Health Resources and Services Administration (HRSA).

The following notes apply throughout these tables:

- 1. Family studies both parents with HPLC, IEF and CBCOR DNA
- 2. Family history includes reported history of Hgb variant in the family
- 3. Need to exclude iron deficiency if using low MCV as part of criteria

Beta + Thal – (note: need separate samples for column 2 and 3)

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history	HPLC & IEF same sample
Definite	Untested or unknown	Untested or unknown	SBeta + THAL	FSA or FS	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	FSAA2	Untested or unknown	SBeta + THAL	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FSA with high A2	SBeta + THAL	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FSAA2	FSA with high A2	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FSAA2	Untested or unknown	Untested or unknown	FSA or FS	Untested or unknown	Both carriers (1 each of Beta + THAL and S)	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FSA with high A ₂	Untested or unknown	FSA or FS	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FSAA2	Untested or unknown	Untested or unknown	FSA or FS	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FS	FSA	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 each of Beta + THAL and S)	Untested or unknown	Untested or unknown
Probable	FSAA2 x2	Untested or unknown	Untested or unknown	FSA or FS	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FSA	FSA	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 each of Beta + THAL and S)	Untested or unknown	Untested or unknown

Beta + Thal – (note: need separate samples for column 2 and 3)

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history	HPLC & IEF same sample
Probable	Untested or unknown	FSA with high A2	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 each of Beta + THAL and S)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FSA or FS	Low MCV	Both carriers (1 each of Beta + THAL and S)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	FSA	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 each of Beta + THAL and S)	Untested or unknown	Untested or unknown
Possible	FSAA2	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 each of Beta + THAL and S)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FSA or FS	Low MCV	Untested or unknown	Positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Untested or unknown	Untested or unknown	FSAA2

E Beta + Thal

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history	HPLC & IEF same sample
Definite	Untested or unknown	Untested or unknown	E Beta + Thal	FEA or FE	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	FEAA ₂	Untested or unknown	E Beta + Thal	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FEA with high A ₂	E Beta + Thal	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FEAA ₂	FEA with high A ₂	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FEAA ₂	Untested or unknown	Untested or unknown	FEA or FE	Untested or unknown	Both carriers (1 with Beta + thal and one for E)	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FEA with high A ₂	Untested or unknown	FEA or FE	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FEAA ₂	Untested or unknown	Untested or unknown	FEA or FE	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FE	FEA	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 with Beta + thal and one for E)	Untested or unknown	Untested or unknown
Probable	FEAA ₂ x2	Untested or unknown	Untested or unknown	FEA or FE	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FEA	FEA	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 with Beta + thal and one for E)	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FEA with high A ₂	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 with Beta + thal and one for E)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FEA or FE	Low MCV	Both carriers (1 with Beta + thal and one for E)	Untested or unknown	Untested or unknown
Possible	FEAA ₂	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 with Beta + thal and one for E)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FEA or FE	Low MCV	Untested or unknown	Positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Untested or unknown	Untested or unknown	FEAA ₂

D Beta + Thal

Category	Qualitativ e (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history	HPLC & IEF same sample
Definite	Untested or unknown	Untested or unknown	D Beta + Thal	FDA or FD	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	FDAA ₂	Untested or unknown	D Beta + THAL	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FDA with high A ₂	D Beta + THAL	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FDAA ₂	FDA with high A ₂	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FDAA ₂	Untested or unknown	Untested or unknown	FDA or FD	Untested or unknown	Both carriers (1 with Beta + thal and one for D)	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FDA with high A ₂	Untested or unknown	FDA or FD	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FDAA ₂	Untested or unknown	Untested or unknown	FDA or FD	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FD	FDA	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 with Beta + thal and one for D)	Untested or unknown	Untested or unknown
Probable	FDAA ₂ x2	Untested or unknown	Untested or unknown	FDA or FD	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown

Probable	FDA	FDA	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 with Beta + thal and one for D)	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FDA with high A ₂	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 with Beta+ thal and one for D)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	FDA	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 with Beta + thal and one for D)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FDA or FD	Low MCV	Both carriers (1 with Beta+ thal and one for D)	Untested or unknown	Untested or unknown
Possible	FDAA ₂	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 with Beta+ thal and one for D)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FDA or FD	Low MCV	Untested or unknown	Positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Untested or unknown	Untested or unknown	FDAA ₂

C Beta + Thal

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history	HPLC & IEF same sample
Definite	Untested or unknown	Untested or unknown	C Beta + Thal	FCA or FC	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	FCAA ₂	Untested or unknown	C Beta + Thal	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FCA with high A ₂	C Beta + Thal	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FCAA ₂	FCA with high A ₂	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FCAA ₂	Untested or unknown	Untested or unknown	FCA or FC	Untested or unknown	Both carriers (1 with Beta+thal and one for C)	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FCA with high A ₂	Untested or unknown	FCA or FC	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FCAA ₂	Untested or unknown	Untested or unknown	FCA or FC	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FCA	FCA	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 with Beta+thal and one for C)	Untested or unknown	Untested or unknown
Probable	FCAA ₂ x2	Untested or unknown	Untested or unknown	FCA or FC	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FCA	FCA	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 with Beta+thal and one for C)	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FCA with high A ₂	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 with Beta+thal and one for C)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	FCA	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 with Beta+thal and one for C)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FCA or FC	Low MCV	Both carriers (1 with Beta+thal and one for C)	Untested or unknown	Untested or unknown
Possible	FCAA ₂	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 with Beta+thal and one for C)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FCA or FC	Low MCV	Untested or unknown	Positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Untested or unknown	Untested or unknown	FCAA ₂

O_{Arab} Beta + Thal

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history	HPLC & IEF same sample
Definite	Untested or unknown	Untested or unknown	O _{Arab} Beta + THAL	FO _{ARAB} A or FO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	FO _{ARAB} AA ₂	Untested or unknown	O _{Arab} Beta + THAL	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknowr
Definite	Untested or unknown	FO _{ARAB} A with high A ₂	O _{Arab} Beta + THAL	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FO _{ARAB} AA ₂	FO _{ARAB} A with high A ₂	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FO _{ARAB} AA ₂ x2	Untested or unknown	Untested or unknown	FO _{ARAB} A or FO _{ARAB}	Untested or unknown	Both carriers (1 Beta + Thal and O _{ARAB})	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FO _{ARAB} A with high A ₂	Untested or unknown	FO _{ARAB} A or FO _{ARAB}	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FO _{ARAB} AA ₂	Untested or unknown	Untested or unknown	FO _{ARAB} A or FO _{ARAB}	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FO _{ARAB}	FO _{ARAB} A	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 Beta + Thal and O _{ARAB})	Untested or unknown	Untested or unknown
Probable	FO _{ARAB} AA ₂ x2	Untested or unknown	Untested or unknown	FO _{ARAB} A or FO _{ARAB}	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FO _{ARAB} A	FO _{ARAB} A	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 Beta + Thal and O _{ARAB})	Untested or unknown	Untested or unknowr
Probable	Untested or unknown	FO _{ARAB} A with high A ₂	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 Beta + Thal and O _{ARAB})	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FO _{ARAB} A or FO _{ARAB}	Low MCV	Both carriers (1 Beta + Thal and O _{ARAB})	Untested or unknown	Untested or unknowr
Possible	Untested or unknown	FO _{ARAB} A	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 Beta + Thal and O _{ARAB})	Untested or unknown	Untested or unknowr
Possible	FO _{ARAB} AA ₂	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 Beta + Thal and O _{ARAB})	Untested or unknown	Untested or unknowr
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FO _{ARAB} AA ₂	Low MCV	Untested or unknown	Positive	Untested or unknowr
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Untested or unknown	Untested or unknown	FO _{ARAB} AA ₂

S Beta (0) Thal

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history	HPLC & IEF same sample
Definite	Untested or unknown	Untested or unknown	SBeta 0 THAL	FSA ₂ or FS	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	FSA ₂ or FS	Untested or unknown	SBeta 0 THAL	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FS high A ₂	SBeta 0 THAL	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FSA ₂ or FS	FS high A ₂	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FSA ₂ or FS	Untested or unknown	Untested or unknown	FSA ₂ or FS	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FS high A ₂	Untested or unknown	FSA ₂ or FS	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FS	FS high A ₂	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 each of BetaTHAL and Beta S)	Untested or unknown	Untested or unknown
Probable	FSA ₂ or FS	Untested or unknown	Untested or unknown	FSA ₂ or FS	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	Untested or unknown	FSA ₂ or FS	Low MCV	Both carriers (1 each of BetaTHAL and Beta S)	Untested or unknown	Untested or unknown
Probable	FSA ₂ or FS x2	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 each of BetaTHAL and Beta S)	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FS high A ₂	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 each of BetaTHAL and Beta S)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FSA ₂ or FS	Low MCV	Untested or unknown	positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Untested or unknown	Untested or unknown	FSA ₂

E Beta (0) Thal

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history	HPLC & IEF same sample
Definite	Untested or unknown	Untested or unknown	EBeta 0 THAL	FEA ₂ or FE	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	FEA ₂ or FE	Untested or unknown	EBeta 0 THAL	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FE high A ₂	EBeta 0 THAL	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FEA ₂ or FE	FE high A ₂	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FEA ₂ or FE	Untested or unknown	Untested or unknown	FEA ₂ or FE	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FE high A ₂	Untested or unknown	FEA ₂ or FE	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FE	FE high A ₂	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 each of BetaTHAL and Beta S)	Untested or unknown	Untested or unknown
Probable	FEA ₂	Untested or unknown	Untested or unknown	FEA ₂ or FE	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	Untested or unknown	FEA ₂ or FE	Low MCV	Both carriers (1 each of BetaTHAL and Beta S)	Untested or unknown	Untested or unknown
Probable	FEA ₂ or FE x2	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 each of BetaTHAL and Beta S)	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FE high A ₂	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 each of BetaTHAL and Beta S)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FEA ₂ or FE	Low MCV	Untested or unknown	Positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Untested or unknown	Untested or unknown	FEA ₂

D Beta (0) Thal

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history	HPLC & IEF same sample
Definite	Untested or unknown	Untested or unknown	D Beta 0 Thal	FDA ₂ or FD	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	FDA ₂	Untested or unknown	D Beta 0 THAL	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FD high A ₂	D Beta 0 THAL	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FDA ₂	FD high A ₂	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FDA ₂	Untested or unknown	Untested or unknown	FDA ₂ or FD	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FD high A ₂	Untested or unknown	FDA₂ or FD	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FDA ₂ or FD x	Untested or unknown	Untested or unknown	FDA ₂ or FD	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	Untested or unknown	FDA₂ or FD	Low MCV	Both carriers (1 each of BetaTHAL and Beta D)	Untested or unknown	Untested or unknown
Probable	FDA ₂ x2	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 each of BetaTHAL and Beta D)	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FD high A ₂	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 each of BetaTHAL and Beta D)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FDA₂ or FD	Low MCV	Untested or unknown	Positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Untested or unknown	Untested or unknown	FDA ₂

C Beta (0) Thal

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history	HPLC & IEF same sample
Definite	Untested or unknown	Untested or unknown	C Beta 0 THAL	FCA ₂ or FC	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	FCA ₂	Untested or unknown	C Beta 0 THAL	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FC high A ₂	C Beta 0 THAL	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FCA ₂ or FC	FC high A ₂	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FCA ₂ or FC	Untested or unknown	Untested or unknown	FCA ₂ or FC	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FC high A ₂	Untested or unknown	FCA ₂ or FC	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FCA ₂	Untested or unknown	Untested or unknown	FCA ₂ or FC	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	Untested or unknown	FCA ₂ or FC	Low MCV	Both carriers (1 each of BetaTHAL and Beta C)	Untested or unknown	Untested or unknown
Probable	FCA ₂ or FC x2	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 each of BetaTHAL and Beta C)	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FC high A ₂	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 each of BetaTHAL and Beta C)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FCA ₂ or FC	Low MCV	Untested or unknown	Positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Untested or unknown	Untested or unknown	FCA ₂

O_{Arab} Beta (0) Thal

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history	HPLC & IEF same sample
Definite	Untested or unknown	Untested or unknown	O _{Arab} Beta 0 THAL	FO _{ARAB} A ₂ or FO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	FO _{ARAB} A ₂	Untested or unknown	O _{Arab} Beta 0 THAL	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FO _{ARAB} High A ₂	O _{Arab} Beta 0 THAL	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FO _{ARAB} A ₂	FO _{ARAB} High A ₂	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FO _{ARAB} A ₂	Untested or unknown	Untested or unknown	FO _{ARAB} A ₂ or FO _{ARAB}	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FO _{ARAB} High A ₂	Untested or unknown	FO _{ARAB} A ₂ or FO _{ARAB}	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FO _{ARAB} A ₂ x2	Untested or unknown	Untested or unknown	FO _{ARAB} A ₂ or FO _{ARAB}	Low MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	Untested or unknown	FO _{ARAB} A ₂ or FO _{ARAB}	Low MCV	Both carriers (1 each of BetaTHAL and Beta O Arab)	Untested or unknown	Untested or unknown
Probable	FO _{ARAB} A ₂ x2	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 each of BetaTHAL and Beta O Arab)	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FO _{ARAB} High A ₂	Untested or unknown	Untested or unknown	Low MCV	Both carriers (1 each of BetaTHAL and Beta O Arab)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FO _{ARAB} A ₂ or FO _{ARAB}	Low MCV	Untested or unknown	Positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Untested or unknown	Untested or unknown	FO _{ARAB} A ₂

Beta Thal Major or (Homozygous or Heterozygous for 2 Beta Thal mutations) – clinical definition – transfusion dependency defines this – generally manifests after 6mo

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history
Definite	Untested or unknown	Untested or unknown	Homozygous for Point Mutation	F	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	1 Point Mutation and 1 Partial Deletion	F	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	2 Partial Deletions	F	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	2 heterozygous point mutations	F	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	High A ₂ (higher than normal)	Untested or unknown	F	Low MCV	Both carriers	Untested or unknown
Probable	F or FA (smaller A than expected)	Untested or unknown	Untested or unknown	F	Low MCV	Both carriers	Untested or unknown
Probable	F or FA (smaller A than expected)	High A ₂ (higher than normal)	Untested or unknown	F	Low MCV	Untested or unknown	Untested or unknown
Possible	F	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Both carriers	Untested or unknown
Possible	Untested or unknown	High A ₂ (higher than normal)	Untested or unknown	Untested or unknown	Low MCV-	Both Carriers	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	F	Low MCV-	Untested or unknown	positive

${\sf HPFH-cannot}\ be\ confirmed\ until\ 6\ months\ of\ age\ or\ older\ if\ do\ not\ have\ DNA\ results$

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history
Definite	Untested or unknown	Untested or unknown	Homozygous for Point Mutation	F	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	1 Point Mutation and 1 Deletion	F	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	2 Deletions	F	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	2 heterozygous point mutations	F	Untested or unknown	Untested or unknown	Untested or unknown
Probable	F	F	Untested or unknown	Untested or unknown	Untested or unknown	Both Carriers	Untested or unknown
Probable	F	Untested or unknown	Untested or unknown	F	Untested or unknown	Both Carriers	Untested or unknown
Probable	Untested or unknown	F	Untested or unknown	F	Untested or unknown	Both Carriers	Untested or unknown
Probable	F	Untested or unknown	Untested or unknown	F	Untested or unknown	Both Carriers	Untested or unknown
Possible	F	Untested or unknown	Untested or unknown	F	Normal MCV	Untested or unknown	Untested or unknown
Possible	Untested or unknown	F	Untested or unknown	F	Normal MCV	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	F	Normal MCV	Both carriers	Untested or unknown
Possible	F	Untested or unknown	Untested or unknown	Untested or unknown	Normal MCV	Both carriers	Untested or unknown
Possible	Untested or unknown	F	Untested or unknown	Untested or unknown	Normal MCV	Both carriers	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	F	Normal MCV	Untested or unknown	Positive

${\sf FSHPFH-Cannot}\ be\ confirmed\ until\ older\ than\ 6\ months\ of\ age\ if\ do\ not\ have\ DNA\ results$

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history
Definite	Untested or unknown	Untested or unknown	1 mutation With known S mutation	FS	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	1 deletion and known S mutation	FS	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FS	FS	Untested or unknown	Untested or unknown	Untested or unknown	Documented carriers of HPFH and S	Untested or unknown
Possible	Untested or unknown	FS	Untested or unknown	FS	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FS	Untested or unknown	Untested or unknown	FS	Untested or unknown	Documented carriers of HPFH and S	Untested or unknown
Possible	FS	Untested or unknown	Untested or unknown	FS	Normal MCV	Untested or unknown	Untested or unknown
Possible	Untested or unknown	FS	Untested or unknown	FS	Normal MCV	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FS	Normal MCV	Documented carriers of HPFH and S	Untested or unknown
Possible	FS	Untested or unknown	Untested or unknown	Untested or unknown	Normal MCV	Documented carriers of HPFH and S	Untested or unknown
Possible	Untested or unknown	FS	Untested or unknown	Untested or unknown	Normal MCV	Documented carriers of HPFH and S	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FS	Normal MCV	Untested or unknown	Positive

FO_{A RAB}HPFH – Cannot be confirmed until older than 6 months of age if do not have DNA results

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history
Definite	Untested or unknown	Untested or unknown	1 mutation With known O _{ARAB} mutation	FO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	1 deletion and known O _{ARAB} mutation	FO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FO _{ARAB}	FO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown	Documented carriers of HPFH and O ARAB	Untested or unknown
Probable	FO _{ARAB}	Untested or unknown	Untested or unknown	FO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or Unknown	FO _{ARAB}	Untested or unknown	FO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FO _{ARAB}	Untested or unknown	Untested or unknown	FO _{ARAB}	Untested or unknown	Documented carriers of HPFH and O _{ARAB}	Untested or unknown
Possible	FO _{ARAB}	Untested or unknown	Untested or unknown	FO _{ARAB}	Normal MCV	Untested or unknown	Untested or unknown
Possible	Untested or unknown	FO _{ARAB}	Untested or unknown	FO _{ARAB}	Normal MCV	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FO _{ARAB}	Normal MCV	Documented carriers of HPFH and O _{ARAB}	Untested or unknown
Possible	FO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown	Normal MCV	Documented carriers of HPFH and O _{ARAB}	Untested or unknown
Possible	Untested or unknown	FO _{ARAB}	Untested or unknown	Untested or unknown	Normal MCV	Documented carriers of HPFH and O _{ARAB}	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FO _{ARAB}	Normal MCV	Untested or unknown	Positive

FCHPFH – Cannot be confirmed until older than 6 months of age if do not have DNA results

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history
Definite	Untested or unknown	Untested or unknown	1 mutation With known C mutation	FC	Low MCV	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	1 deletion and known C mutation	FC	Low MCV	Untested or unknown	Untested or unknown
Probable	FC	FC	Untested or unknown	Untested or unknown	Low MCV	Documented carriers of HPFH and C	Untested or unknown
Probable	FC	Untested or unknown	Untested or unknown	FC	Low MCV	Documented carriers of HPFH and C	Untested or unknown
Possible	Untested or unknown	FC	Untested or unknown	FC	Low MCV	Untested or unknown	Untested or unknown
Possible	FC	Untested or unknown	Untested or unknown	FC	Low MCV	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FC	Low MCV	Documented carriers of HPFH and C	Untested or unknown
Possible	FC	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Documented carriers of HPFH and C	Untested or unknown
Possible	Untested or unknown	FC	Untested or unknown	Untested or unknown	Low MCV	Documented carriers of HPFH and C	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FC	Low MCV	Untested or unknown	Positive

FEHPFH – Cannot be confirmed until older than 6 months of age if do not have DNA results

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA sequencing and deletion	NBS result	CBC Results	Family Studies	Family history
Definite	Untested or unknown	Untested or unknown	1 deletion and known E mutation	FE	Low MCV	Untested or unknown	Untested or unknown
Probable	FE	FE	Untested or unknown	Untested or unknown	Low MCV	Documented carriers of HPFH and E	Untested or unknown
Probable	FE	Untested or unknown	Untested or unknown	FE	Low MCV	Documented carriers of HPFH and E	Untested or unknown
Probable	Untested or unknown	FE	Untested or unknown	FE	Low MCV	Documented carriers of HPFH and E	Untested or unknown
Possible	FE	Untested or unknown	Untested or unknown	FE	Low MCV	Untested or unknown	Untested or unknown
Possible	FE	Untested or unknown	Untested or unknown	FE	Low MCV	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FE	Low MCV	Documented carriers of HPFH and E	Untested or unknown
Possible	FE	Untested or unknown	Untested or unknown	Untested or unknown	Low MCV	Documented carriers of HPFH and E	Untested or unknown
Possible	Untested or unknown	FE	Untested or unknown	Untested or unknown	Low MCV	Documented carriers of HPFH and E	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FE	Low MCV	Untested or unknown	Positive

C and E will not have normal MCV with HPFH-do not reference MCV

Sickle Cell Diseases Case Definitions Tables September 29, 2013

Definitions created by panel of experts between June 2011 and September 2013.	This project was
funded in part by Cooperative Agreement # U22MC24078 from the Health Resou	irces and Services
Administration (HRSA).	

Throughout this document, the following definitions are used:

- 1. Family studies both parents with HPLC, IEF and CBC
- 2. Family history includes reported history of Hgb variant in the family

SC Disease

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA	NBS result	Family Studies	Family history	HPLC& IEF same sample
Definite	FSC	Untested or unknown	Known C and known S mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FSC	Known C and known S mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	Known C and known S mutation identified	FSC	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	Known C and known S mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	FSC
Probable	FSC	FSC	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FSC	Untested or unknown	Untested or unknown	FSC	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FSC	Untested or unknown	FSC	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	Untested or unknown	FSC	Both carriers (1 with C mutation and other with S mutation)	Untested or unknown	Untested or unknown
Probable	FSC	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 with C mutation and other with S mutation)	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FSC	Untested or unknown	Untested or unknown	Both carriers (1 with C mutation and other with S mutation)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FSC	Untested or unknown	positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	FSC

SD Disease

Category	Qualitative (IEF or HPLC)	Quantitativ e (HPLC or electrophoresis)	DNA	NBS result	Family Studies	Family history	HPLC& IEF same sample
Definite	FSD	Untested or unknown	Known D and known S mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FSD	Known D and known S mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	Known D and known S mutation identified	FSD	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	Known D and known S mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	FSD
Probable	FSD	FSD	Untested or unknown	Untested or	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FSD	Untested or	Untested or unknown	FSD	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FSD	Untested or unknown	FSD	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	Untested or unknown	FSD	Both carriers (1 with known S mutation and 1 with known D mutation)	Untested or unknown	Untested or unknown
Probable	FSD	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 with known S mutation and 1 with known D)	Untested or unknown	Untested or unknown

SD Disease

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA	NBS result	Family Studies	Family history	HPLC& IEF same sample
Probabl e	Untested or unknown	FSD	Untested or unknown	Untested or unknown	Both carriers (1 with known S mutation and 1 with known D)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FSD	Untested or unknown	positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	FSD

SE Disease

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA	NBS result	Family Studies	Family history	HPLC& IEF same sample
Definite	FSE	Untested or unknown	Known E and known S mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FSE	Known E and known S mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	Known E and known S mutation identified	FSE	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FSE	FSE	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FSE	Untested or unknown	Untested or unknown	FSE	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FSE	Untested or unknown	FSE	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	Untested or unknown	FSE	Both carriers (1 with known S mutation and 1 with known E)	Untested or unknown	Untested or unknown
Probable	FSE	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 with known S mutation and 1 with known E)	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FSE	Untested or unknown	Untested or unknown	Both carriers (1 with known S mutation and 1 with known E)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FSE	Untested or unknown	positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	FSE

SO_{Arab} Disease

Category	Qualitative	Quantitative	DNA	NBS result	Family Studies	Family history	HPLC& IEF same
catego. y	(IEF or HPLC)	(HPLC or electrophoresis)	5	112313311			sample
Definite	FSO _{ARAB}	Untested or unknown	Known OARAB and known S mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FSOARAB	Known OARAB and known S mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	Known OARAB and known S mutation identified	FSO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FSO _{ARAB}	FSO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FSOARAB	Untested or unknown	Untested or unknown	FSO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FSO _{ARAB}	Untested or unknown	FSOARAB	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	Untested or unknown	FSO _{ARAB}	Both carriers (1 with known S mutation and 1 with known OARAB)	Untested or unknown	Untested or unknown
Probable	FSO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 with known S mutation and 1 with known OARAB)	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FSO _{ARAB}	Untested or unknown	Untested or unknown	Both carriers (1 with known S mutation and 1 with known OARAB)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FSOARAB	Untested or unknown	positive	Untested or unknown
Poss ible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	FSOARAB

CD Disease

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA	NBS result	Family Studies	Family history	HPLC& IEF same sample
Definite	FCD	Untested or unknown	Known C and known D mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FCD	Known C and known D mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	Known C and known D mutation identified	FCD	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FCD	FCD	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FCD	Untested or unknown	Untested or unknown	FCD	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FCD	Untested or unknown	FCD	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	Untested or unknown	FCD	Both carriers (1 with D and 1 with E)	Untested or unknown	Untested or unknown
Probable	FCD	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 with D and 1 with E)	Untested or unknown	Untested or unknown

CD Disease

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA	NBS result	Family Studies	Family history	HPLC& IEF same sample
Probable	Untested	FCD	Untested	Untested	Both carriers (1	Untested or unknown	Untested or unknown
	or		or	or	with D and 1		
	unknown		unknown	unknown	with E)		
Possible	Untested	Untested	Untested	FCD	Untested	positive	Untested or unknown
	or	or	or		or		
	unknown	unknown	unknown		unknown		
Possible	Untested	Untested	Untested	Untested	Untested	Untested or unknown	FCD
	or	or	or	or	or		
	unknown	unknown	unknown	unknown	unknown		

CE Disease

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA	NBS result	Family Studies	Family history	HPLC& IEF same sample
Definite	FCE	Untested or unknown	Known C and known E mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FCE	Known C and known E mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	Known C and known E mutation identified	FCE	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FCE	FCE	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FCE	Untested or unknown	Untested or unknown	FCE	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FCE	Untested or unknown	FCE	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	Untested or unknown	FCE	Both carriers (1 with C and 1 with E)	Untested or unknown	Untested or unknown
Probable	FCE	Untested or unknown	Untested or unknown	Untested or	Both carriers (1 with C and 1 with E)	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FCE	Untested or unknown	Untested or	Both carriers (1 with C and 1 with E)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FCE	Untested or unknown	positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or	Untested or unknown	Untested or unknown	FCE

CO_{Arab} Disease

	COATAD DISCUS						
Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA	NBS result	Family Studies	Family history	HPLC& IEF same sample
Definite	FCO _{ARAB}	Untested or unknown	Known C and known OARAB mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FCO _{ARAB}	Known C and known OARAB mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	Known C and known OARAB mutation identified	FCO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FCO _{ARAB}	FCO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FCOARAB	Untested or unknown	Untested or unknown	FCOARAB	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FCOARAB	Untested or unknown	FCO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	Untested or unknown	FCO _{ARAB}	Both carriers (1 carrier C and 1 carrier OARAB)	Untested or unknown	Untested or unknown
Probable	FCOARAB	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 carrier C and 1 carrier OARAB)	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FCO _{ARAB}	Untested or unknown	Untested or unknown	Both carriers (1 carrier C and 1 carrier OARAB)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FCOARAB	Untested or unknown	positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	FCOARAB

DE Disease

	o E Diocase								
Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA	NBS result	Family Studies	Family history	HPLC& IEF same sample		
Definite	FDE	Untested or unknown	Known D and known E mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown		
Definite	Untested or unknown	FDE	Known D and known E mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown		
Definite	Untested or unknown	Untested or unknown	Known D and known E mutation identified	FDE	Untested or unknown	Untested or unknown	Untested or unknown		
Probable	FDE	FDE	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown		
Probable	FDE	Untested or unknown	Untested or unknown	FDE	Untested or unknown	Untested or unknown	Untested or unknown		
Probable	Untested or unknown	FDE	Untested or unknown	FDE	Untested or unknown	Untested or unknown	Untested or unknown		
Probable	Untested or unknown	Untested or unknown	Untested or unknown	FDE	Both carriers (1 carrier E and 1 carrier D)	Untested or unknown	Untested or unknown		
Probable	FDE	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 carrier E and 1 carrier D)	Untested or unknown	Untested or unknown		
Probable	Untested or unknown	FDE	Untested or unknown	Untested or unknown	Both carriers (1 carrier E and 1 carrier D)	Untested or unknown	Untested or unknown		
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FDE	Untested or unknown	positive	Untested or unknown		
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	FDE		

DO_{Arab} Disease

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA	NBS result	Family Studies	Family history	HPLC& IEF same sample
Definite	FDO _{ARAB}	Untested or unknown	Known OARAB and known S mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FDO _{ARAB}	Known OARAB and known S mutation identified	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	Known OARAB and known S mutation identified	FDO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FDO _{ARAB}	FDO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown
Probable	FDO _{ARAB}	Untested or unknown	Untested or unknown	FDO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FDO _{ARAB}	Untested or unknown	FDO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	Untested or unknown	FDO _{ARAB}	Both carriers (1 carrier C and 1 carrier OARAB)	Untested or unknown	Untested or unknown
Probable	FDO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers (1 carrier C and 1 carrier OARAB)	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FDO _{ARAB}	Untested or unknown	Untested or unknown	Both carriers (1 carrier C and 1 carrier OARAB)	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FDO _{ARAB}	Untested or unknown	positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	Untested or unknown	FDO _{ARAB}

SS Disease

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA-	NBS result	СВС	Family Studies	Family history	Hbg testing (Electrophoresis or HPLC) on family members
Definite	FS	Untested or unknown	SS	Untested or unknown	Untested or unknown	Both carriers S	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FS	SS	Untested or unknown	Untested or unknown	Both carriers S	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	SS	FS	Untested or unknown	Both carriers S	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	SS	Untested or unknown	Untested or unknown	Both carriers S		
Probable	FS	Untested or unknown	Untested or unknown	FS	Nml- high MCV	Untested or unknown	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FS	Untested or unknown	FS	Untested or unknown	Both carriers S	Untested or unknown	Untested or unknown
Probable	FS	Untested or unknown	Untested or unknown	FS	Untested or unknown	Both carriers S	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FS	Nml- high MCV	Untested or unknown	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FS	Untested or unknown	Untested or unknown	positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FS	Untested or unknown	Untested or unknown	Untested or unknown	positive

CC Disease

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA	NBS result	СВС	Family Studies	Family history	Hbg testing (Electrophoresis or HPLC) on family members
Definite	Untested or unknown	FC	СС	Untested or unknown	Nml MCV	Both carriers C	Untested or unknown	Untested or unknown
Definite	FC	Untested or unknown	CC	Untested or unknown	Nml MCV	Both carriers C	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	CC	FC	Nml MCV	Both carriers C	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	СС	Untested or unknown	Untested or unknown	Both Carriers C		
Probable	Untested or unknown	FC	Untested or unknown	FC	Untested or unknown	Both carriers	Untested or unknown	Untested or unknown
Probable	FC	FC	Untested or unknown	Untested or unknown	Untested or unknown	Both Carriers C	Untested or unknown	Untested or unknown
Probable	FC	Untested or unknown	Untested or unknown	FC	Untested or unknown	Both carriers	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FC	Nml MCV	Untested or unknown	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FC	Untested or unknown	Untested or unknown	positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FC	Untested or unknown	Untested or unknown	Untested or unknown	positive

EE Disease

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA- no deletion/duplication analysis	NBS result	CBC	Family Studies	Family history	Hbg testing (Electrophoresis or HPLC) on family members
Definite	FE	Untested or unknown	EE	Untested or unknown	Nml MCV	Both carriers E	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FE	EE	Untested or unknown	Nml MCV	Both carriers E	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	EE	FE	Nml MCV	Both carriers E	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	EE	Untested or unknown	Untested or unknown	Both Carriers E	Untested or unknown	Untested or unknown
Probable	FE	FE	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers E	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FE	Untested or unknown	FE	Untested or unknown	Both carriers	Untested or unknown	Untested or unknown
Probable	FE	Untested or unknown	Untested or unknown	FE	Untested or unknown	Both carriers	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FE	Nml MCV	Untested or unknown	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FE	Untested or unknown	Untested or unknown	positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FE	Untested or unknown	Untested or unknown	Untested or unknown	positive

DD Disease

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA-	NBS result	СВС	Family Studies	Family history	Hbg testing (Electrophoresis or HPLC) on family members
Definite	FD	Untested or unknown	DD	Untested or unknown	Nml MCV	Both carriers D	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FD	DD	Untested or unknown	Nml MCV	Both carriers D	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	DD	FD	Nml MCV	Both carriers D	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	DD	Untested or unknown	Untested or unknown	Both Carriers D	Untested or unknown	Untested or unknown
Probable	FD	FD	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers D	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FD	Untested or unknown	FD	Untested or unknown	Both carriers	Untested or unknown	Untested or unknown
Probable	FD	Untested or unknown	Untested or unknown	FD	Untested or unknown	Both carriers	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FD	Nml MCV	Untested or unknown	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FD	Untested or unknown	Untested or unknown	positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FD	Untested or unknown	Untested or unknown	Untested or unknown	positive

Homozygous O_{Arab} Disease

Category	Qualitative (IEF or HPLC)	Quantitative (HPLC or electrophoresis)	DNA-	NBS result	СВС	Family Studies	Family history	Hbg testing (Electrophoresis or HPLC) on family members
Definite	FO _{ARAB}	Untested or unknown	O _{ARAB} O _{ARAB}	Untested or unknown	Nml MCV	Both carriers O _{ARAB}	Untested or unknown	Untested or unknown
Definite	Untested or unknown	FO _{ARAB}	O _{ARAB} O _{ARAB}	Untested or unknown	Nml MCV	Both carriers O _{ARAB}	Untested or unknown	Untested or unknown
Definite	Untested or unknown	Untested or unknown	$O_{ARAB}O_{ARAB}$	FO _{ARAB}	Nml MCV	Both carriers O _{ARAB}	Untested or unknown	Untested or unknown
Probable	Untested or unknown	Untested or unknown	O _{ARAB} O _{ARAB}	Untested or unknown	Untested or unknown	Both Carriers O _{ARAB}	Untested or unknown	Untested or unknown
Probable	FO _{ARAB}	FO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown	Both carriers O _{ARAB}	Untested or unknown	Untested or unknown
Probable	Untested or unknown	FO _{ARAB}	Untested or unknown	FO _{ARAB}	Untested or unknown	Both carriers	Untested or unknown	Untested or unknown
Probable	FO _{ARAB}	Untested or unknown	Untested or unknown	FO _{ARAB}	Untested or unknown	Both carriers	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FO _{ARAB}	Nml MCV	Untested or unknown	Untested or unknown	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FO _{ARAB}	Untested or unknown	Untested or unknown	positive	Untested or unknown
Possible	Untested or unknown	Untested or unknown	Untested or unknown	FO _{ARAB}	Untested or unknown	Untested or unknown	Untested or unknown	positive

Cystic Fibrosis Case Definitions Tables September 29, 2013

Definitions created by panel of experts between June 2011 and September 2013. This project was funded in part by Cooperative Agreement # U22MC24078 from the Health Resources and Services Administration (HRSA).

Note: CF Disease Causing Mutations should be confirmed on CFTR2 (www.cftr2.org)

CF Disease Causing Mutations should be confirmed on CFTR2 (<u>www.cftr2.org</u>).

Typical CF

Category	Classification	Clinical	Sweat Chloride	Non Newborn Screen Molecular	Newborn Screen Molecular	NBS Result
	Definite		>=60 mmol/L (regardless of age)	Not available or not done	2 CF disease-causing mutations	
	Definite		>=60 mmol/L (regardless of age)	2 CF disease-causing mutations	Not available or not done	
	Definite		No valid sweat chloride result available	2 CF disease-causing mutations	2 CF disease-causing mutations	
	Definite	No known medical condition associated with false positive sweatchloride	TWO results >=60 mmol/L (regardless ofage, two independent results from separatedays)	Not available or not done	Not available or not done	
	Definite		<60 mmol/L	2 CF disease-causing mutations and 1 or both have been shown to have lower chlorides (see CFTR2)	2 CF disease-causing mutations and 1 or both have been previously shown to have lower chlorides, (see CFTR2)	
CF	Probable		No valid sweat chloride result available	Not available or not done	2 CF-causing mutations	
Typical CF	Probable		No valid sweat chloride result available	2 CF-causing mutations	Not available or not done	
F	Probable		>=60 mmol/L (single test, regardlessofage)	Not available or not done	2 Mutations of varying clinical significance	
	Probable		>=60 mmol/L (single test, regardlessofage)	Not available or not done	2 Mutations of unknown clinical significance	
	Probable		>=60 mmol/L (single test, regardlessofage)	2 Mutations of varying clinical consequence	Not available or not done	
	Probable		>=60 mmol/L (single test, regardlessofage)	2 Mutations of unknown clinical significance	Not available or not done	
	Probable		<60 mmol/L	2 CF disease-causing mutations and 1 or both have been previously shown to have lowerchlorides	Not available or not done	
	Probable	obable <60 mmol/L		Not available or not done	CF disease-causing mutations and 1 or both have been previously shown to have lower chlorides,	

CF Disease Causing Mutations should be confirmed on CFTR2 (<u>www.cftr2.org</u>).

Typical CF

Category	Classification	Clinical	Sweat Chloride	Non Newborn Screen Molecular	Newborn Screen Molecular	NBS Result
Typical CF	Possible		No valid sweat chloride result available	2 CF disease-causing mutations CF-causing mutations not yet shown to be <i>in trans</i>	Not available or not done	
Τ	Possible		No valid sweat chloride result available	Not available or not done	2 CF disease-causing mutations CF-causing	
	Definite		<30 if <6mos, <40 if ≥6 mos On 2 occasions	1 CF disease-causing mutation and 1 Mutation of varying clinical consequence <i>in trans</i>		Elevated IRT
	Definite		<30 if <6mos, <40 if >6 mos On 2 occasions	1 CF disease-causing mutation and 1 Mutation of unknown significance		Elevated IRT
CRMS	Definite		30-59 (age<6 mos) On 2 occasions		1 CF disease-causing mutation and 1 Mutation of unknown significance	Elevated IRT
	Definite		40 -59 (age≥6 mos) On 2 occasions		1 CF disease-causing mutation and 1 Mutation of varying clinical consequence	Elevated IRT
	Definite		30-59 (age<6 mos) On 2 occasions			Elevated IRT
	Definite		40 -59 (age>6 mos) On 2 occasions			Elevated IRT

CF Disease Causing Mutations should be confirmed on CFTR2 (<u>www.cftr2.org</u>).

Typical CF

Category	Classification	Clinical	Sweat Chloride	Non Newborn Screen Molecular	Newborn Screen Molecular	NBS Result
CRD	Definite	CBAVD, recurrent pancreatitis, nasal polyposis, infertility and focal biliary cirrhosis with portal hypertension	30-59 (age<6 mos) On 2 occasions	1 CF disease-causing mutation and 1 Mutation of unknown significance		
כו	Definite	CBAVD, recurrent pancreatitis, nasal polyposis, infertility and focal biliary cirrhosis with portal hypertension	30-59 (age<6 mos) On 2 occasions	1 CF disease-causing mutation and 1 Mutation of varying clinical consequence		

Severe Combined Immunodeficiency Case Definitions Tables May, 2018

Definitions created by panel of experts between June 2011 and September 2013. This project was funded in part by Cooperative Agreement # U22MC24078 from the Health Resources and Services Administration (HRSA).

1. Primary targets of NBS

- a. Typical SCID
- b. Leaky SCID
- c. Omenn Syndrome
- 2. Secondary targets of NBS
 - a. Syndromes with variable immune defects, with some cases having significantly low T-cell numbers
 - b. Secondary T-cell lymphopenia
 - c. Idiopathic T-cell lymphopenia

Typical SCID

Classification: Typical SCID	CD3 T cells/μL	Proliferation to PHA	Maternal engraftment Y/N	Molecular testing	Clinical Presentation
1. Definite	<300 autologous T Cells, undetectable or very few naïve T cells	<10% of normal	Yes	Consistent with SCID [^]	
2. Definite	<300 autologous T Cells, undetectable or very few naïve T cells	<10% of normal	Yes	Unknown or not done	
3. Definite	<300 autologous T Cells, undetectable or very few naïve T cells	Unknown or any	Yes	Consistent with SCID [^]	
4. Definite	Any number	<10% of normal	Yes	Consistent with SCID [^]	
5. Definite	<300 autologous T Cells, undetectable or very few naïve T cells	<10% of normal	No	Consistent with SCID [^]	
6. Probable	<300 autologous T Cells, undetectable or very few naïve T cells	<10% of normal	No	Unknown or not done	

7. Probable	Any number	<10% of normal	No	Consistent with SCID [^]	
8. Probable	Any number	<10% of normal	Yes	Unknown or not done	
9. Probable	Any number	Unknown or any	Yes	None or inconclusive	
10. Probable	Any number	Unknown or any	Yes	Consistent with SCID [^]	
11. Possible	<300 autologous T Cells, undetectable or very few naïve T cells	<10% of normal	Untested or unknown	Untested or unknown	Untested or unknown
12. Possible	<300 autologous T Cells, undetectable or very few naïve T cells	Unknown or any	No	Unknown or not done	
13. Possible	<300 autologous T Cells, undetectable or very few naïve T cells	Unknown or any	No	Consistent with SCID [^]	
14. Probable	<300 autologous T Cells, undetectable or very few naïve T cells	Unknown or any	Yes	Unknown or not done	
15. Possible	Any number	Unknown or any	No	Consistent with SCID [^]	
16. Possible	Any number	<10% of normal	No	None or inconclusive	
17. Uncertain	Any number	Unknown or any	No	Unknown or not done	

[^] Consistent with SCID: Two pathogenic variants in a known SCID gene; pathogenic variant in SCID gene on X chromosome in a male; ruled out 22q11 deletion; ruled out heterozygous TBX1 variants; ruled out homozygous or compound heterozygous FOXN1 mutations

Leaky SCID

Classifi Leaky S	ication: SCID	CD3 T cells/μL	Proliferation	Maternal engraftment Y/N	Molecular testing	Clinical Presentation
1.	Definite	300-1500, few naïve T cells, oligoclonal T cells or poor T cell diversity	10-50% normal PHA	No	Unknown or not done	
2.	Definite	300-1500, few naïve T cells	10-50% normal PHA	No	Consistent with SCID [^]	
3.	Possible	300-1500, few naïve T cells	Unknown or any	No	Unknown or not done	
4.	Definite	300-1500, few naïve T cells	Unknown or any	No	Consistent with SCID [^]	
5.	Possible	Any number	10-30% normal PHA <i>or</i> Absent to Candida/TT	No	Unknown or not done	
6.	Definite	Any number	10-30% normal PHA <i>or</i> Absent to Candida/TT	No	Consistent with SCID [^]	

^{^^} Consistent with Leaky SCID: Two pathogenic variants in a known SCID gene known to be associated with leaky SCID (previously reported or in a gene previously associated with a combined immune deficiency) or one pathogenic variant in SCID gene on X chromosome in a male; ruled out 22q11 deletion; ruled out heterozygous TBX1 variants; ruled out homozygous or compound heterozygous FOXN1 mutations

Omenn Syndrome

Classifi Omenr syndro		CD3 T cells/μL	Proliferation to PHA	Maternal engraftment Y/N	Molecular testing	Clinical Presentation
1.	Definite	>80%CD45RO+	10-50%normal	No	Consistent with OS/SCID^^	Consistent with OS**
2.	Definite	>80%CD45RO+	10-50%normal	No	Untested or unknown	Consistent with OS**
3.	Definite	>80%CD45RO+	10-50%normal	No	No variant reported, ruled out 22q11 and FOXN1	Consistent with OS**
4.	Probable	>80%CD45RO+	Untested or unknown	No	Consistent with OS/SCID^^	Consistent with OS**
5.	Probable	>80%CD45RO+	10-50%normal	Unknown	Untested or unknown	Consistent with OS**
6.	Uncertain	>80%CD45RO+	Untested or Unknown	Untested or Unknown	Untested or Unknown	Consistent with OS**

^{**} Clinical presentation may include Erythroderma with biopsy showing T cell infiltrate; hepatomegaly, splenomegaly or both; adenopathy, eosinophilia, elevated levels of serum IgE antibody

^{^^} Consistent with OS/SCID: Two pathogenic variants in a SCID gene known to be associated with leaky SCID (previously reported or in a gene previously associated with a combined immune deficiency) or one pathogenic variant in SCID gene on X chromosome in a male; ruled out 22q11 deletion; ruled out heterozygous TBX1 variants; ruled out homozygous or compound heterozygous FOXN1 mutations

Non-SCID conditions associated with SCID NBS

Classification	Findings
Syndromes with low T- cell numbers	Recognized genetic syndrome that includes variable immune defects, with some cases having significantly low T-cell numbers (DiGeorge syndrome, FOXN1, CHARGE syndrome, Trisomy 21, Jacobsen syndrome, RAC2 defect, DOCK8 deficiency, Ataxia Telangiectasia, VACTERL association, Barth syndrome, TAR syndrome, Ectrodactyly Ectodermal Dysplasia syndrome, Cartilage Hair Hypoplasia, others)
Secondary T-cell lymphopenia	Congenital malformation or disease process without an intrinsic defect in production of circulating T-cells (e.g. congenital heart disease with vascular leak, hydrops, gastroschisis, chylothorax, intestinal lymphangiectasia, others)
Preterm birth alone	Preterm birth and low birth weight, with low T cell numbers early in life that normalize over time
Idiopathic T-cell lymphopenia (formerly called Variant SCID)	Persistently low T cell numbers for over 3 months without recognized cause

In all of these other conditions there is 1) no maternal engraftment, 2) the T cells are largely naïve, 3) PHA proliferation is usually normal.

Reference List

- 1. Kwan A, Puck JM. History and current status of newborn screening for severe combined immunodeficiency. Semin Perinatol. 2015 Apr;39(3):194-205.
- 2. Dvorak, C.C., et al., The natural history of children with severe combined immunodeficiency: baseline features of the first fifty patients of the primary immune deficiency treatment consortium prospective study 6901. J Clin Immunol, 2013. 33(7): p. 1156-64.
- 3. Shearer W Dynn E, Notarangelo LD, et al. Establishing diagnostic criteria for severe combined immunodeficiency disease (SCID), leaky SCID and Omenn syndrome: The Primary Immune Deficiency Treatment Consortium experience. JACI 2014;133:1092-8.
- 4. Kwan A, Abraham RS, Currier R et al. Newborn screening for severe combined immunodeficiency in 11 screening programs in the United States. *JAMA* 2014;312:729-738.

Pompe Disease Case Definitions Tables May, 2018

Definitions created by panel of experts between June 2011 and September 2013. This project was funded in part by Cooperative Agreement # U22MC24078 from the Health Resources and Services Administration (HRSA).

Pompe Disease

				Enzyme	activity			
	Classification	Disorder	Mutation Status	Blood (not DBS sample)	Skin/Muscle testing	Cardiac Involvement consistent with Pompe	Lab Findings	Clinical Findings
	Definite	Infantile Onset Pompe Disease	Allele 1 –pathogenic and associated with infantile onset and Allele 2 – pathogenic and associated with infantile onset	Within lab known affected range for infantile onset (IO)	Not done or positive skin or muscle bx	Positive findings on Chest X- ray/EKG/ECHO in newborn period	Unknown/N ot Done	
	Definite	Infantile Onset Pompe Disease	Unknown or not done	Within lab known affected range for IO	Not done or positive skin or muscle bx	Positive findings on Chest X- ray/EKG/ECHO in newborn period	Elevated CK/AST/AL T/ LDH/Urine Hex4	
Pompe Disease	Definite	Infantile Onset Pompe Disease	Allele 1 – pathogenic and associated with infantile onset, 1 novel variant that is likely pathogenic	Within lab known affected range for IO	Not done or positive skin or muscle bx	Positive findings on Chest X- ray/EKG/ECHO in newborn period	Elevated CK/AST/AL T/ LDH/Urine Hex4	
Pomp	Definite	Infantile Onset Pompe Disease	Allele 1 – pathogenic and associated with infantile onset and Allele 2 – pathogenic and associated with infantile onset	Within lab known affected range for IO	Not done or positive skin or muscle bx	Positive findings on Chest X- ray/EKG/ECHO in newborn period	Elevated CK/AST/AL T/ LDH/Urine Hex4	
	Definite	Infantile Onset Pompe Disease	1 pathogenic* or likely pathogenic variant, with deletion or duplication consistent with infantile onset	Within lab known affected range for IO	Not done	Positive findings on Chest X- ray/EKG/ECHO	Elevated CK/AST/AL T/ LDH/Urine Hex4	
	Definite	Infantile Onset Pompe Disease	Allele 1 – pathogenic and associated with infantile onset and Allele 2 – pathogenic and associated with non-classical disease, or variant of uncertain significance	Low (above affected range, for IO, may or may not be in late- onset (LO) range but should not be	positive skin or muscle bx	Positive findings on Chest X- ray/EKG/ECHO in newborn period	Elevated CK/AST/AL T/ LDH/Urine Hex4	

			above LO				
			range))				
Probable	Infantile Onset Pompe Disease	Allele 1 – pathogenic and associated with infantile onset and Allele 2 – pathogenic and associated with non-classical disease, or variant of uncertain significance)	Within lab known affected range for IO	Unknown/not done	Positive findings on Chest X- ray/EKG/ECHO	Unknown/no t done	
Probable	Infantile Onset Pompe Disease	1 pathogenic* or likely pathogenic variant, no other variants found, dup/del testing not done or not known	Within lab known affected range for IO	Unknown/not done	Positive findings on Chest X- ray/EKG/ECHO	Elevated CK/AST/AL T/ LDH/Urine Hex4	
Definite	Late Onset Pompe Disease	Allele 1 – pathogenic and Allele 2 – pathogenic and associated with non- classical disease, or variant of uncertain significance)	Within lab known affected range for LO	Unknown/not done	No	Elevated CK/AST/AL T/ LDH/Urine Hex4	Symptoms present after year of age a documented specialists. Fundament of continued to collect data through the development symptoms*
Definite	Late Onset Pompe Disease	Allele 1 – pathogenic and Allele 2 – pathogenic and associated with non- classical disease, or variant of uncertain significance)	Within lab known affected range for LO	Unknown/not done	No	Elevated CK/AST/AL T/ LDH/Urine Hex4	Symptoms present befor year of age b no cardiac involvemen
Probable	Late Onset Pompe Disease	Allele 1 – pathogenic and associated with infantile onset and Allele 2 – pathogenic associated with non-classical disease, or variant of uncertain significance)	Within lab known affected range for LO	Unknown/not done	No	Elevated CK/AST/AL T/ LDH/Urine Hex4	Unknown or r reported to P to program b the end of follow-up
Possible	Late Onset Pompe Disease	Allele 1 – pathogenic* and associated with infantile	Low (above affected	Unknown/not done	No	Not present	

		onset and Allele 2 –	range,for LO				
		pathogenic*)	not normal)				
Possible	Late Onset Pompe Disease	Allele 1 – pathogenic* and associated with infantile onset), no other variants detected; Duplication/deletion testing not completed or unknown	Within lab known affected range for LO	Unknown/not done	No	Not present	
Definite	Late Onset Pompe Disease	Allele 1 – pathogenic* and associated with infantile onset), no other variants detected; Duplication/deletion testing not completed or unknown	Within lab known affected range for LO	Unknown/not done	No	Elevated CK/AST/AL T/ LDH/Urine Hex4	Symptoms present after 1 year of age and documented by specialists. PH program continued to collect data through the development of symptoms**
Possible	Late Onset Pompe Disease	1 pathogenic* or likely pathogenic variant, no other variant found	Within lab known affected range	Unknown/not done	No	Elevated CK/AST/AL T/ LDH/Urine Hex4	
Possible	Late Onset Pompe Disease	1 pathogenic* or likely pathogenic variant, no other variants found	Within lab known affected range	Unknown/not done	No	Not present	

^{*} Pathogenic: classified as pathogenic or likely pathogenic by ACMG Guidelines (2015)

^{**} Clinical symptoms consistent with Pompe Disease: progressive muscle weakness, need for respiratory assistance, swaying gait or waddle, Lordosis, kyphosis, or scoliosis.

MPSI

Case Definitions Tables May, 2018

Definitions created by panel of experts between June 2011 and September 2013. This project was funded in part by Cooperative Agreement # U22MC24078 from the Health Resources and Services Administration (HRSA).

MPSI

	Classification	Disorder	Mutation Status	Enzyme Activity	Urine GAGS	Clinical Symptoms/Lab Findings
	Definite	MPS I – severe	Allele 1 – pathogenic* and associated with severe disease and Allele 2 – pathogenic and associated with severe disease #	Within lab known affected range	Elevated	
	Definite	MPS I - severity not determined	Allele 1 – pathogenic* or likely pathogenic and Allele 2 – variant with uncertain significance	Within lab known affected range	Elevated	
	Definite	MPS I – severity not determined	Not Done/unknown	Within lab known affected range	Elevated	
MPSI	Probable	MPS I – severe	Allele 1 – pathogenic* and associated with severe disease and Allele 2 – pathogenic* and associated with severe disease #	Within lab known affected range	Not done/UNKNOWN	
	Probable	MPS I – severe	Allele 1 – pathogenic* and associated with severe disease and Allele 2 – pathogenic* and associated with severe disease #	Unknown	Not done/UNKNOWN	
	Definite	MPS I Attenuated	Allele 1 – pathogenic* and associated with severe disease and Allele 2 – variant known to be associated with ATTENUATED Disease #	Within lab known affected range	Elevated	Symptoms present and documented by specialists. PH program continued to collect data through the development of symptoms**
	Definite	MPS I Attenuated	Allele 1 – variant known to be associated with ATTENUATED Disease and Allele 2 – variant	Within lab known affected range	Elevated	

		known to be associated with ATTENUATED Disease #			
Probable	MPS I Attenuated	Allele 1 – pathogenic* and associated with severe disease and Allele 2 – variant known to be associated with ATTENUATED Disease #	Unknown/Not Done	Unknown/Not Done	Symptoms present and documented by specialists. PH program continued to collect data through the development of symptoms**
Possible	MPS I Attenuated	Allele 1 – pathogenic* and associated with severe disease and Allele 2 – variant known to be associated with ATTENUATED Disease #	Within lab known affected range	Not done/UNKNOWN	UNKNOWN
Possible MPS Attenuar		Allele 1 – pathogenic* and associated with severe disease and Allele 2 – variant known to be associated with ATTENUATED Disease #	Unknown	Not done/UNKNOWN	
Possible MPS I - Seve not determin		Allele 1 – pathogenic* and associated with severe disease and Allele 2 – variant of unknown significance	Unknown	Not done/UNKNOWN	Symptoms present and documented by specialists. PH program continued to collect data through the development of symptoms**
Possible	MPS I Attenuated	Allele 1 – pathogenic* and associated with severe disease and Allele 2 – variant of unknown significance	Unknown	Not done/UNKNOWN	No symptoms by the time the PH Program closes follow-up (either due to child being lost to follow- up OR program policy on follow-up time

^{*} Pathogenic: Reported in cases known to have severe cases previously.

[#] All reports of two variants determined to be disease causing are assumed to bin in trans, and appropriate testing was completed as necessary

^{**} Clinical symptoms consistent with MPS-I include: Hepatosplenomegaly, Coarse facial features, Hydrocephalus, Skeletal deformities (dysostosis multiplex), Corneal clouding, Large tongue, Prominent forehead, Joint stiffness, Short stature, frequent ear infections and hearing loss, hemia

X-ALD Newborn Screening Case Definitions Tables September 2018

Definitions created by a panel of experts is supported by the Health Resources and Services Administration (HRSA) under Cooperative Agreement # UG9MC30369 New Disorders Implementation Project.

X-ALD

In Males:

	Category	Plasma VLCFA	Clinical	Plasmalogen	Mutation analysis	Family History
			Symptoms			
	Definite	Elevated^	Not present	Untested or	Pathogenic Variant in ABCD1	
				unknown	gene	
	Definite	Elevated^	Not present	Normal	Deletion/duplication	
					identified in ABCD1 gene	
trophy	Definite	Elevated^	Not present	Normal	No mutation on sequencing, deletion/duplication not done	Family history or family VLCFA studies suggestive of X-linked ALD
leukodys	Definite	Elevated^	Not present	Normal	Variant of uncertain significance in <i>ACBD1</i> gene	Family history or family VLCFA studies suggestive of X-linked ALD
dreno	Possible	Elevated^	Not present	Normal	Variant of uncertain significance in <i>ACBD1</i> gene	
X-linked Adrenoleukodystrophy	Possible	Elevated^	Not present	Normal	No mutation on sequencing, deletion/duplication not done; Rule out other disorders of peroxisomal beta oxidation	
	Possible	Elevated^	Not present	Normal	Untested or unknown	
	Probable	Not available	Not available	Not available	Pathogenic Variant in ABCD1 gene	
	Possible	Not available	Not present	Not available	Variant of uncertain significance in <i>ACBD1</i> gene	Family history or family VLCFA studies suggestive of X-linked ALD#

[^] In the pathogenic range

[#] Family history may include multiple relatives consistent with X-linked transmission as determined by clinical specialist: Maternal grandfather, maternal aunts, mother

[^] In the pathogenic range

^{**} Symptoms may include: neonatal hypotonia, neonatal seizures, liver disease, neonatal cholestasis, sensorineural deafness

In Females:

	Category	Diagnosis	Plasma VLCFA	Plasmalogen	Mutation analysis	Fibroblast studies	Additional Comments
	Definite	Carrier Female	Normal	Normal	Pathogenic variant ABCD1 gene	Untested or unknown	
yhdc	Definite	Carrier Female	Elevated^	Normal	Pathogenic variant ABCD1 gene	Untested or unknown	
dystro	Possible	Carrier Female	Elevated^	Not done/unknown	Not done/unknown		
X-linked Adrenoleukodystrophy	Possible	Carrier Female	Not done/unknown	Not done/unknown	Not done/unknown	Not done/unknown	Family history or family VLCFA studies suggestive of X-linked ALD
X-lin	Definite	Carrier Female	Elevated^	Normal	Variant of uncertain significance ABCD1 gene		
	Possible	Carrier Female	Normal	Normal	Variant of uncertain significance <i>ABCD1</i> gene		

[^] In the pathogenic range

	Category	Plasma VLCFA	Plasmalogen	Clinical symptoms	Mutation analysis	Fibroblast studies
-a	Definite	Elevated^	Low	Present*	Two pathogenic variants in the same PEX gene	Untested or unknown
Disorder	Definite	Elevated^	Low	Present*	Untested or unknown	Consistent with ZSD
Zellweger Spectrum L	Definite	Elevated^	Low	Not present	Two pathogenic variants in the same PEX gene	Untested or unknown
eger Sp	Definite	Elevated^	Low	Not present	Untested or Unknown	Consistent with ZSD
Zellw	Definite	Elevated^	Low	Present*	Untested or Unknown	Untested or unknown
	Possible	Elevated^	Low	Not present	Untested or Unknown	Untested or unknown
	Possible	Elevated^	Normal	Not present	Untested or Unknown	Untested or unknown

[^] In the pathogenic range

^{*} Clinical symptoms may include: Hypotonia in newborn period, failure to thrive, craniofacial abnormalities, abnormal liver function tests.

ıal -	Category	Plasma VLCFA	Clinical Symptoms	Plasmalogen	Mutation analysis	Family History
Peroxisom Disorder	PROBABLE	Elevated^	Not present	Normal	No mutation on sequencing, deletion/duplication not found	

[^] In the pathogenic range

e	Category	Plasma	Plasmalogen	Mutation	Fibroblast studies	Clinical Symptoms
dase V		VLCFA		analysis		
-CoA Oxid Deficiency	Definite	Elevated^	Normal	Two pathogenic mutations in the ACOX1 gene	Untested or unknown	Not present at birth
Acyl	Possible	Elevated^	Normal	Untested or unknown	Consistent with Acyl- CoA Oxidase Deficiency	Not present at birth

[^] In the pathogenic range

Protein 3y	Category	Plasma VLCFA	Plasmalogen	Mutation analysis	Fibroblast studies	Clinical Symptoms
Bifunctional P Deficiency	Definite	Elevated^	Normal	Two pathogenic mutations in the HSD17B4 gene	Untested or unknown	Present*
D-Bifur	Possible	Elevated^	Normal	Untested or unknown	Consistent with D- Bifunctional Protein	Present*

[^] In the pathogenic range

^{*} Clinical symptoms may include: Hypotonia in newborn period, failure to thrive, craniofacial abnormalities, abnormal liver function tests.

1 0	Category	Plasma VLCFA	Plasmalogen	Mutation analysis	Fibroblast studies	Clinical Symptoms
ABDC5	Definite	Elevated^	Normal	Two disease causing mutations	Untested or unknown	Not present at birth
	Definite	Elevated^	Normal	Untested or unknown	Consistent with ABCD5	Not present at birth

[^] In the pathogenic range

oxisomal ırder	Category	Plasma VLCFA	Plasmalogen	Mutation analysis	Fibroblast studies	Clinical Symptoms
Non-pero Diso	Definite	Normal	Normal	Mutation in one of the 7 known genes for Aicardi-Goutiéres Syndrome	Untested or unknown	Present**

^{**} Clinical symptoms may include: Hypotonia in newborn period, failure to thrive, on CT scan, intracranial calcifications.

- References:
- www.x-ald.nl (https://adrenoleukodystrophy.info)
- Berger, J, et al. (2014). "Mini-review; Pathophysiology of X-linked adrenoleukodystrophy." Biochemie 98: 135-142.
- Engelen, Met al. (2014). "X-linked adrenoleukodystrophy in women: a cross-sectional cohort study." Brain 137: 693-706.
- Wang, Y et al. (2011) "X-linked adrenoleukodystrophy: ABCD1 de novo mutations and mosaicism." Molecular Genetics and Metabolism 104: 160-166.
- Engelen, M. et al. (2012). "X-linked adrenoleukodystrophy (X-ALD): clinical presentation and guidelines for diagnosis, follow-up and management." Orphanet J. of Rare Diseases **7**:51
- Ferdinandusse S, et al. (2017) "ACBD5 deficiency causes a defect in peroxisomal very long-chain fatty acid metabolism." J Med Genet; **54**:330-337.
- Armangue T et al. (2017) "Neonatal detection of Aicardi-Goutiéres Syndrome by increased C26:0 lysophosphatidylcholine and interferon signature on newborn screening blood spots." Molecular Genetics and Metabolism
- Kemper AR et al. (2017) "Newborn screening for X-linked adrenoleukodystrophy: evidence summary and advisory committee recommendation." Genetics in Medicine 19:1
- Vogel BH et al. (2015) "Newborn screening for X-linked adrenoleukodystrophy in New York State: Diagnostic protocol, surveillance protocol, and treatment guidelines." Molecular Genetics and Metabolism
- Moser AB et al. (2016) "Newborn screening for X-linked adrenoleukodystrophy." Intl. J. of Neonatal Screening: 2;15
- Moser AB et al. (1999). "Plasma very long chain fatty acids in 3,000 peroxisome disease patients and 29,000 controls. "Ann. Neurol; **45**; 100-110.
- Waterham HR et al. (2007). "A lethal defect of mitochondrial and peroxisomal fission." N Engl. J Med: 356; 17.
- Corzo, D., et al. (2002). "Contiguous deletion of the X-linked adrenoleukodystrophy gene (ABCD1) and DXS1357E: a novel neonatal phenotype similar to peroxisomal biogenesis disorders." Am J Hum Genet **70**(6): 1520-1531.
- Waterham HR, Ebberink MS. Genetics and molecular basis of human peroxisome biogenesis disorders. *Biochimica et Biophysica Acta*. 2012;1822:1430-1441.
- Braverman NE, Raymond GV, Rizzo WB, Moser AB, Wilkinson ME, Stone EM, Steinberg SJ, Wangler MF, Rush ET, Hacia JG, Bose M. Peroxisome biogenesis disorders in the Zellweger spectrum: An overview of current diagnosis, clinical manifestations, and treatment guidelines. *Mol Genet Metab.* 2016;117:313.321.

SMA Newborn Screening Case Definitions Tables November 2019

Definitions created by a panel of experts is supported by the Health Resources and Services Administration (HRSA) under Cooperative Agreement # UG9MC30369 New Disorders Implementation Project.

Classification	Newborn Screen Molecular*		Post-Newborn Screen Molecular		Parental Molecular Testing Family History/Parental Genetic Testing	Clinical Symptoms at the time of Presentation**
	SMN1	SMN2 Copy Number	SMN1	SMN2 Copy Number		
Definite	Zero copies of SMN1 (presumed homozygous deletion/conversion) ^	Any	Zero copies of SMN1 (presumed homozygous deletion)	Any		
Definite	2 pathogenic variants	Any	2 pathogenic variants	Any	Phasing is complete and confirms that variants are in trans or both parents are known to be carriers of the pathogenic variants identified	
Definite	2 pathogenic variants observed on two independently collected NBS specimens	Any	Unknown/ Not Done	Any	Phasing is complete and confirms that variants are in trans or both parents are known to be carriers of the	

					pathogenic variants identified
Definite	Unknown/ Not Done/Screen Negative	Any	2 pathogenic variants observed on 2 independently collected specimens	Any	Phasing is complete and confirms that variants are in trans or both parents are known to be carriers of the pathogenic variants identified
Definite	Zero copies of SMN1 (presumed homozygous deletion/conversion) ^		Unknown/ Not Done	Unknown/ Not Done	Both parents are known carriers of SMN1 deletion
Definite	Unknown/ Not Done /Screen Negative	Any	Zero copies of SMN1 (presumed homozygous deletion/conversion)	Any	Both parents are known carriers of SMN1 deletion
Definite	Unknown/ Not Done /Screen Negative	Any	Zero copies of SMN1 (presumed homozygous deletion/conversion) ^ observed on two independently collected specimens	Any	
Definite	Zero copies of SMN1 (presumed homozygous deletion/conversion) ^ observed on two independently collected NBS specimens	Any	Unknown/ Not Done		

Definite	Unknown/ Not Done /Screen Negative	Any	Zero copies of SMN1 (presumed	Any		Clinical Symptoms
			homozygous deletion/conversion) ^			present (see list)
Probable	Zero copies of SMN1 (presumed homozygous deletion/conversion) ^	Any	Unknown/ Not Done	Any		Clinical Symptoms Present (see list)
Probable	2 pathogenic variants	Any	2 pathogenic variants		Phasing not done or not known	Clinical symptoms present (see list)
Probable	2 pathogenic variants observed on two independently collected NBS specimens	Any	Unknown/ Not Done		Phasing not done or not known	Clinical symptoms present (see list)
Probable	Unknown/ Not Done/Screen Negative	Any	2 pathogenic variants observed on 2 independent collected specimens		Phasing not done or not known	Clinical symptoms present (see list)
Possible	Zero copies of SMN1 (presumed homozygous deletion/conversion)	Any	Unknown/ Not Done		Unknown/ Not Done	Unknown/ Not Done
Possible	Unknown/ Not Done /Screen Negative	Any	Zero copies of SMN1 (presumed homozygous deletion/conversion) ^	Any	Unknown/ Not Done	Unknown
Possible	2 pathogenic variants observed on two independently collected NBS specimens	Any	Unknown/ Not Done			

Possible	Unknown/ Not	Any	2 pathogenic variants		
	Done/Screen Negative		observed on 2		
			independently		
			collected specimens		
Possible	2 pathogenic variants	Any	2 pathogenic variants		
Possible	2 pathogenic variants	Any	Unknown/ Not Done	Phasing not done or not known	Clinical symptoms present (see list)
Possible	Unknown/ Not Done	Any	2 pathogenic variants	Phasing not done or not known	Clinical symptoms present (see list)
Possible	1 pathogenic variant and 1 variant of unknown significance	Any	1 pathogenic variant and 1 variant of unknown significance		
Possible	1 pathogenic variant and 1 variant of unknown significance	Any	1 pathogenic variant and 1 variant of unknown significance		Clinical symptoms present (see list)
Possible	1 pathogenic variant and 1 variant of unknown significance	Any	1 pathogenic variant and 1 variant of unknown significance	Phasing is complete and confirms that variants are in trans or both parents are known to be carriers of the variants identified	with or without clinical symptoms
Possible	2 variants of unknown significance	Any	2 variants of unknown significance		
Possible	2 variants of unknown significance	Any	2 variants of unknown significance	Phasing is complete and confirms that variants are in trans or both parents are known to be carriers of the variants identified	with or without clinical symptoms
Possible	Unknown/ Not Done /Screen Negative	Any	2 variants of unknown significance		Clinical symptoms present (see list)

KEY:

- ^ presumed homozygous deletion/conversion: true deletion of exon 7 (or larger) or for which there has been a gene conversion of exon 7 (or more)
- * Programs need to ensure specimens are valid, taking into account NICU status and inhibitor use
- ** Clinical symptoms include Electromyography evidence of motor neuron disease, Absent reflexes, Fasciculations, Feeding difficulty, Hypotonia, Respiratory Difficulty, Weakness