

Case Worksheets for Newborn Screening

Version 1.3 | June 2018 | <u>https://www.newsteps.org</u> | <u>newsteps@aphl.org</u>

Case Information Worksheet

Question:	Answer	
INFANTDEMOGRAPHICS		
State Unique ID?(alphanumeric)		
Date of Birth?(mm/dd/yyyy)		
Gestational Age?(in weeks)		
Birth Weight?(in grams)		
Biological Sex? Race? (select all that apply)	 Male Female Unspecified Unknown White Black or African American 	
	 American Indian or Alaskan Native Asian Native Hawaiian or other Pacific Islander Not Reported/Unknown 	
Ethnicity? (select one)	 Hispanic, Latino(a) or Spanish origin Not of Hispanic, Latino(a),or Spanish origin Not Reported/Unknown 	
SCREENINGINFORMATION		
Was prenatal testing done that indicated that this infant was at risk for this disorder?	Yes No Don't Know	
Which newborn screen result indicated this infant was at risk for the disorder?	 Initial Screen 2nd Required Screen Subsequent Screen 	
Was this individual diagnosed later in life (not identified by newborn screening)?	Yes No Unknown	
Date of <u>initial</u> specimen collection (mm/dd/yyyy)?		
Date of receipt by lab of <u>initial specimen (mm/dd/yyyy)?</u>		
Date of release of out of range results of <u>initial</u> specimen (mm/dd/yyyy)?		
Date of <u>subsequent</u> specimen collection (mm/dd/yyyy)?		
Date of receipt by lab of <u>subsequent</u> specimen (mm/dd/yyyy)?		
Date of release of out of range results of <u>subsequent</u> specimen (mm/dd/yyyy)?		
Date of intervention by appropriate medical provider (mm/dd/yyyy)?		
Date of confirmation of diagnosis (mm/dd/yyyy)?		

Newborn Screening Surveillance Case Definitions

Developed by the Health Resources and Services Administration (HRSA) and NewSTEPs in cooperation with the newborn screening medical sub-specialty community, standard surveillance case definitions for newborn screening conditions allow for determination of true prevalence and incidence of disorders, and for comparison of outcomes across states. The case definition forms can be found in the pages to follow, stratified by disorder type. Additionally you can find case definition classification tables linked here that can used as a reference resource.

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Note: standard surveillance case definitions have not been developed for 3-Hydroxy-3-methyglutaric aciduria (HMG) or for ß-Ketothiolase deficiency (ßKT). These are forthcoming.

Metabolic Disorders

Organic Acid Disorders:

Glutaric Acidemia/ Aciduria Type I (GA1) Case Confirmatory Diagnosis Follow-Up

Please answer the following as Yes/No/Don't Know	If Yes
Were urine organic acids tested? Yes No Don't Know	Was 3-OH Glutaric acid level Elevated Normal Unknown Was Glutaric acid level Elevated Normal Unknown
Were serum organic acids tested? ☐ Yes ☐ No ☐ Don't Know	Was 3-OH Glutaric acid level Elevated Normal Unknown Was Glutaric acid level Elevated Normal Unknown
Were plasma acylcarnitines tested? Yes No Don't Know 	Was C5 -DC level Elevated Normal Unknown
Was a mutation analysis performed for Glutaric aciduria type I? Yes No Don't Know	What genes were included in the mutation analysis? GCDH gene Other gene:

MOLECULAR GENETICS REPORT	
Were variants detected in the GCDH gene? Yes No Don't Know	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown
Were variants detected in other genes? Yes No Don't Know 	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown
Was enzyme analysis for Glutaric Acidemia enzyme activity completed? Yes No Don't Know	Was enzyme activity: Consistent with disease Normal activity (not consistent with disease) Unknown

Isovaleric Acidemia/ Aciduria (IVA) Case Confirmatory Diagnosis Follow-Up

Please answer the following as Yes/No/Don't Know	If Yes
Were urine organic acids tested? ☐ Yes ☐ No ☐ Don't Know	Was 3OH Isovaleric acid level Elevated Normal Unknown Was Isovaleryl glycine level Elevated Normal Unknown
Were plasma acylcarnitines tested? Yes No Don't Know 	Was C5 level Elevated Normal Unknown
Was a mutation analysis performed for Isovaleric aciduria? Yes No Don't Know	What genes were included in the mutation analysis? IVD gene Other gene:

MOLECULAR GENETICS REPORT	
Were variants detected in the IVD gene?	Check the types of variants found on:
□ Yes	Allele 1:
□ No	Variant known to be disease causing
Don't Know	Variant of unknown significance
	 Predicted to be pathogenic
	□ Wild Type (Normal)
	Allele 2:
	Variant known to be disease causing
	Variant of unknown significance
	 Predicted to be pathogenic
	□ Wild Type (Normal)
Were variants detected in other genes?	Check the types of variants found on:
Yes	Allele 1:
	□ Variant known to be disease causing
Don't Know	 Variant of unknown significance
	 Predicted to be pathogenic
	□ Wild Type (Normal)
	Unknown
	Allele 2:
	Variant known to be disease causing
	Variant of unknown significance
	 Predicted to be pathogenic
	□ Wild Type (Normal)
Was enzyme analysis for isovaleric acidemia	Was enzyme activity:
enzyme activity completed?	Consistent with disease
Yes	 Normal activity (not consistent with disease)
	Unknown
Don't Know	

3-Methylcrotonyl-CoA Carboxylase Deficiency (3-MCC) Case Confirmatory Diagnosis Follow-Up

Please answer the following as Yes/No/Don't Know	If Yes
Were urine organic acids tested? ☐ Yes ☐ No ☐ Don't Know	Was 3OH Isovaleric acid level Elevated Normal Unknown Was 3-methylcrotonyl glycine level Elevated Normal Unknown
Were plasma acylcarnitines tested? Yes No Don't Know	Was C5-OH level Elevated Normal Unknown
Was maternal 3-MCC level tested and ruled out? Yes No Don't Know 	
Was a mutation analysis performed for 3-MCC? Yes No Don't Know	 What genes were included in the mutation analysis? MCCC1 gene MCCC2 gene Other gene:

MOLECULAR GENETICS REPORT	
Were variants detected in the MCCC1 gene?	Check the types of variants found on:
□ Yes	Allele 1:
□ No	Variant known to be disease causing
Don't Know	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	🗖 Unknown
	Allele 2:
	Variant known to be disease causing
	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
Were variants detected in the MCCC2 gene?	Check the types of variants found on:
□ Yes	Allele 1:
□ No	Variant known to be disease causing
Don't Know	Variant of unknown significance
	 Predicted to be pathogenic
	U Wild Type (Normal)
	Allele 2:
	□ Variant known to be disease causing
	□ Variant of unknown significance
	• Predicted to be pathogenic
	Wild Type (Normal)
Were variants detected in other genes?	Check the types of variants found on:
Yes	Allele 1:
	□ Variant known to be disease causing
Don't Know	Variant of unknown significance
	• Predicted to be pathogenic
	 Wild Type (Normal) Unknown
	Allele 2:
	□ Variant known to be disease causing
	□ Variant of unknown significance
	 Predicted to be pathogenic
	□ Wild Type (Normal)
	Unknown
Was enzyme analysis for 3-MCC enzyme activity	Was enzyme activity:
completed?	Consistent with disease
□ Yes	 Normal activity (not consistent with disease)
Don't Know	

MMA With Homocystinuria; (CblC; CblD; CblF; CblDv1; CblJ) Case Confirmatory Diagnosis Follow-Up

Instructions: Please answer the questions as fully as possible. All testing refers to confirmatory testing following newborn screening. You may need to consult with relevant clinician(s) and/or medical staff to obtain this information.

Part I: Final Diagnosis as determined by metabolic geneticist or clinician performing the follow-up

Please Select One:

- A. Cobalamin C deficiency (CblC)
- B. Cobalamin D deficiency (CbID)
- C. Cobalamin F deficiency (CbIF)
- D. Cobalamin Dv1 deficiency (CblDv1)
- E. Cobalamin J deficiency (CblJ)
- F. Other cobalamin deficiency not listed above: ______

Please answer the following as Yes/No/Don't Know	If Yes
Was serum MMA level tested?	Was MMA level in serum:
Yes	Elevated
No	Normal
Don't Know	Unknown
Was urine MMA level tested?	Was MMA level in urine:
Yes	Elevated
No	Normal
Don't Know	Unknown
Were plasma acylcarnitines tested? Yes No Don't Know 	Was C3: Elevated Normal Unknown
Was maternal vitamin B12 levels tested?	Was maternal vitamin B12 deficient?
Yes	Yes
No	No
Don't Know	Unknown

Were infant vitamin B12 levels tested? Yes Don't Know Was total plasma homocysteine tested? Yes No Don't Know	Was infant vitamin B12 deficient? Yes NO Unknown Was total plasma homocysteine: Elevated Normal Unknown What genes were included in the mutation analysis?
□ Yes	C2ORF25 gene (cblD)
□ No □ Don't Know	 MMACHC gene LMBRD1 gene (cblF)
	ABCD4 gene (cblJ)
	Other MMA associated gene:
Mole	CULAR GENETICS REPORT
Were variants found in C2ORF25 gene?	Check the types of variants found on:
□ Yes	Allele 1:
□ No	Variant known to be disease causing
Don't Know	Variant of unknown significance
	• Predicted to be pathogenic
	Wild Type (Normal)
	Unknown Allele 2:
	□ Variant known to be disease causing
	 Variant of unknown significance
	 Predicted to be pathogenic
	□ Wild Type (Normal)
	□ Unknown
Were variants found in MMACHC gene?	Check the types of variants found on:
□ Yes	Allele 1:
	Variant known to be disease causing
Don't Know	□ Variant of unknown significance
	• Predicted to be pathogenic
	 Wild Type (Normal) Unknown
	Allele 2:
	□ Variant known to be disease causing
	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	Unknown

Were variants found in LMBRD1 gene?	Check the types of variants found on:
□ Yes	Allele 1:
□ No	Variant known to be disease causing
Don't Know	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	□ Unknown
	Allele 2:
	Variant known to be disease causing
	Variant of unknown significance
	 Predicted to be pathogenic
	□ Wild Type (Normal)
Were variants found in ABCD4 gene?	Check the types of variants found on:
Yes	Allele 1:
	□ Variant known to be disease causing
Don't Know	 Variant of unknown significance
	-
	• Predicted to be pathogenic
	Wild Type (Normal)
	Unknown Allele 2:
	Variant known to be disease causing Variant of unknown significance
	□ Variant of unknown significance
	• Predicted to be pathogenic
	Wild Type (Normal)
Were variants found in other MMA related	Check the types of variants found on:
genes?	Allele 1:
□ Yes	Variant known to be disease causing
□ No	Variant of unknown significance
Don't Know	 Predicted to be pathogenic
	Wild Type (Normal)
	Unknown
	Allele 2:
	Variant known to be disease causing
	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	Unknown
Were enzyme complementation studies completed?	Were complementation studies:
□ Yes	Consistent with disease
	 Normal activity (not consistent with disease)
Don't Know	

MMA Without Homocystinuria; (CblA; CblB; mut-; mut0; CblDv2) Case Confirmatory Diagnosis Follow-Up

Instructions: Please answer the questions as fully as possible. All testing refers to confirmatory testing following newborn screening. You may need to consult with relevant clinician(s) and/or medical staff to obtain this information.

Part I: Final Diagnosis as determined by metabolic geneticist or clinician performing the follow-up

Please Select One:

- G. Cobalamin A deficiency (CbIA)
- H. Cobalamin B deficiency (CblB)
- I. Mutase (-) (mut-0)
- J. Mutase (0) (mut0)
- K. Cobalamin Dv2 (CblDv2)

Please answer the following as Yes/No/Don't Know	If Yes
Was serum MMA level tested? Yes No Don't Know	Was MMA level in serum: Elevated Normal Unknown
Was urine MMA level tested? Yes No Don't Know	Was MMA level in urine: Elevated Normal Unknown
Were plasma acylcarnitines tested? Yes No Don't Know 	Was C3: Elevated Normal Unknown
Was maternal vitamin B12 levels tested? Yes No Don't Know	Was maternal vitamin B12 deficient? Yes No Unknown

Were infant vitamin B12 levels tested? Yes No Don't Know Was total plasma homocysteine tested? Yes No Don't Know	Was infant vitamin B12 deficient? Yes NO Unknown Was total plasma homocysteine: Elevated Normal Unknown What genes were included in the mutation analysis?
□ Yes	METHYLMALONYL-CoA MUTASE
No Don't Know	MMAA gene
	MMAB gene
	Other MMA associated gene:
	CULAR GENETICS REPORT
Were variants found in the METHYLMALONYL-CoA MUTASE gene? Yes No Don't Know	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown
 Were variants found in MMAA gene? Yes No Don't Know 	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance • Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance • Predicted to be pathogenic Wild Type (Normal) Unknown

Were variants found in MMAB gene?	Check the types of variants found on:
□ Yes	Allele 1:
🗆 No	Variant known to be disease causing
Don't Know	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	Unknown
	Allele 2:
	Variant known to be disease causing
	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	□ Unknown
Were variants found in other MMA related	Check the types of variants found on:
genes?	Allele 1:
□ Yes	Variant known to be disease causing
🗆 No	Variant of unknown significance
Don't Know	 Predicted to be pathogenic
	Wild Type (Normal)
	□ Unknown
	Allele 2:
	Variant known to be disease causing
	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	Unknown
Were enzyme complementation studies completed?	Were complementation studies:
□ Yes	Consistent with disease
□ No	Normal activity (not consistent with disease)
Don't Know	□ Unknown

Propionic Acidemia/ Aciduria Case Confirmatory Diagnosis Follow-Up

Please answer the following as Yes/No/Don't Know	If Yes
Were urine organic acids tested?	Please indicate which of the following metabolites were detected:
🗆 Yes	Propionyl glycine:
🗆 No	□ Yes
Don't Know	🗆 No
	🗖 Unknown
	Tiglyglycine:
	□ Yes
	🗆 No
	🗖 Unknown
	Methylcitrate:
	□ Yes
	🗆 No
	🗖 Unknown
	3OH propionic acid:
	□ Yes
	🗆 No
	🗖 Unknown
	MMA:
	□ Yes
	□ No
	Unknown
	Methylcrotonyl glycine:
	□ Yes
	□ No
Were plasma acylcarnitines tested?	Was C3 level
□ Yes	Elevated
□ No	Normal
Don't Know	Unknown

Was a mutation analysis performed for Propionyl- CoA carboxylase (PCC)? Yes No Don't Know	What genes were included in the mutation analysis? PCCA PCCB Other gene:
Molec	CULAR GENETICS REPORT
Were variants detected in the PCCA gene? Yes No Don't Know	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown
 Were variants detected in the PCCB gene? Yes No Don't Know 	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown

Were variants detected in other genes?	Check the types of variants found on:
 Yes No Don't Know 	 Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown

Holocarboxylase Synthetase (Multiple Carboxylase) Deficiency or Other Biotin Disorders (Not Biotinidase Deficiency) (MCD) Case Confirmatory Diagnosis Follow-Up

Instructions: Please answer the question as fully as possible. All testing refers to confirmatory testing following newborn screening. You may need to consult with relevant clinician(s) and/or medical staff to obtain this information.

Final diagnosis as determined by the metabolic geneticist or clinician performing the follow-up:

Please choose one:

- A. Holocarboxylase deficiency
- B. Other biotin disorder (not biotinidase deficiency)_____

Please answer the following as Yes/No/Don't Know	If Yes
Were urine organic acids tested? Yes No Don't Know	Was 3OH Isovaleric acid level Elevated Normal Unknown Was 3OH Propionic acid level Elevated Normal Unknown Was 3-methylcrotonyl glycine level Elevated Normal Unknown
Were plasma acylcarnitines tested? Yes No Don't Know	Was C3 level Elevated Normal Unknown Was C5-OH level Elevated Normal Unknown
Were infant chemistries (biotinidase) studies completed? Yes No Don't Know	What were the Biotinadase results? Normal Abnormal Untested/Unknown

Was a mutation analysis performed for Holocarboxylase Synthetase Deficiency? Ves No Don't Know	What genes were included in the mutation analysis? HLCS gene Other gene:
Molec	ULAR GENETICS REPORT
 Were variants detected in the HLCS gene? Yes No Don't Know 	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance • Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance • Predicted to be pathogenic Wild Type (Normal) Unknown
 Were variants detected in other genes? Yes No Don't Know 	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown
Was enzyme analysis for holocarboxylase synthetase deficiency enzyme activity completed? Yes No Don't Know	Was enzyme activity: Consistent with disease Normal activity (not consistent with disease) Unknown

Fatty Acid Disorders:

Primary Carnitine Deficiency/ Carnitine Uptake Deficiency (CUD) Case Confirmatory Diagnosis Follow-Up

Please answer the following as Yes/No/Don't Know	If Yes
Was urine carnitine tested? Yes No Don't Know	Was fractional excretion of free carnitine level: Elevated Normal Unknown
Were plasma carnitine levels tested? Yes No Don't Know 	Was free carnitine (CO) Low Normal Unknown
 Were other causes for carnitine loss ruled out? Yes No Don't Know 	
Was a mutation analysis performed for carnitine transporter defects? Yes No Don't Know	What genes were included in the mutation analysis? SLC22A5 gene Other gene:
Molec	ULAR GENETICS REPORT
Were variants detected in the SLC22A5 gene? Yes No Don't Know	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown

Were variants detected in other genes?	Check the types of variants found on:
□ Yes	Allele 1:
🗆 No	Variant known to be disease causing
Don't Know	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	Unknown
	Allele 2:
	Variant known to be disease causing
	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	Unknown
Was enzyme analysis for carnitine	Was enzyme activity:
deficiency enzyme activity completed?	Consistent with disease
□ Yes	Normal activity (not consistent with disease)
□ No	□ Unknown
Don't Know	

Medium-chain acyl-CoA Dehydrogenase Deficiency (MCAD) Case Confirmatory Diagnosis Follow-Up

Please answer the following as Yes/No/Don't Know	If Yes
 Were urine organic acids or aclyglycines tested? Yes No Don't Know Were plasma acylcarnitines tested?	Was Hexanoylglycine level Elevated Normal Unknown Was C8 level:
☐ Yes ☐ No ☐ Don't Know	 Elevated Elevated on repeat testing Normal Unknown Was C8>C10 level Elevated Normal Unknown Was C8>C6 level Elevated Normal Unknown Was C6 level Elevated Normal Unknown Was C10 level Elevated Normal Unknown Was C10 level Elevated Normal Unknown
Was mutation analysis performed for MCAD? Yes No Don't Know 	What genes were included in the mutation analysis? ACADM Other:

MOLECULAR GENETICS REPORT		
Were variants detected in the ACADM gene? Yes No Don't Know	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown	
Were variants detected in other genes? Yes No Don't Know	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown	
Was functional analysis of fatty acid oxidation in cultured fibroblasts performed?	Was functional fibroblast analysis: Consistent with disease Normal activity (not consistent with disease) Unknown 	
Was enzyme analysis for MCAD enzyme activity completed? Yes No Don't Know	Was enzyme analysis: Consistent with disease Normal activity (not consistent with disease) Unknown 	

Tri-Functional Protein Deficiency (TFP); Inclusive of LCHAD Case Confirmatory Diagnosis Follow-Up

Instructions: Please answer the questions as fully as possible. All testing refers to confirmatory testing following newborn screening. You may need to consult with relevant clinician(s) and/or medical staff to obtain this information.

Part I: Final Diagnosis as determined by metabolic geneticist or clinician performing the follow-up

Please select one:

- L. Trifunctional Protein Deficiency
- M. Long Chain Acyl CoA dehydrogenase deficiency (LCHAD)

Please answer the following as Yes/No/Don't Know	If Yes
Were urine organic acids tested? Yes No Don't Know	Was C12-OH dicarboxylic acid level Elevated Normal Unknown Was C10-OH dicarboxylic level? Elevated Normal Unknown
Were plasma acylcarnitines tested? Yes No Don't Know	Was C16-OH level Elevated (on more than one sample) Normal Unknown Was C16:1-OH level Elevated Normal Unknown Was C18-OH level Elevated Normal Unknown Was C18:1-OH level I Elevated Normal Unknown

Was mutation analysis performed for Trifunctional Protein deficiency? Yes No Don't Know	What genes were included in the mutation analysis? HADHA HADHB Other:
Molec	ular Genetics Report
Were variants detected in HADHB gene? No Don't Know Were variants detected in HADHA gene? Yes Yes No Don't Know	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Variant of unknown significance Predicted to be pathogenic Variant of unknown significance Variant of unknown significance Unknown
Were variants detected in Other genes?	 Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Check the types of variants found on: Allele 1: Variant known to be disease causing
Don't Know	 Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown

Was enzyme analysis for TFP enzyme activity completed? Yes No Don't Know	Was enzyme analysis: Consistent with disease Normal activity (not consistent with disease) Unknown
Was functional analysis of fatty acid oxidation in cultured fibroblasts performed? Yes No Don't Know	 Was Functional fibroblast analysis: Consistent with disease Normal activity (not consistent with disease) Unknown

Very Long-chain acyl-CoA Dehydrogenase Deficiency Case Confirmatory Diagnosis Follow-Up

Please answer the following as Yes/No/Don't Know	If Yes
Were plasma acylcarnitines tested? Yes No Don't Know	Was C14:1 level Elevated (on more than one sample) Normal Unknown Was C14:2 level Elevated Normal Unknown Was C14 level Elevated Normal Unknown
Was mutation analysis performed for VLCAD? Yes No Don't Know	What genes were included in the mutation analysis? ACADVL Other:

Molecular Genetics Report	
Were variants detected in ACADVL gene? Yes Don't Know Were variants detected in Other genes? Yes No Don't Know	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Vild Type (Normal) Unknown Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Variant of unknown significance
Was enzyme analysis for VLCAD enzyme activity completed? Yes No Don't Know	Was enzyme analysis: Consistent with disease Normal activity (not consistent with disease) Unknown
Was functional analysis of fatty acid oxidation in cultured fibroblasts performed? Yes No Don't Know	 Was functional fibroblast analysis: Consistent with disease Normal activity (not consistent with disease) Unknown

Argininosuccinic Acidemia/ Aciduria (ASA) Case Confirmatory Diagnosis Follow-Up

Please answer the following as Yes/No/Don't Know	If Yes
Were plasma amino acids tested? ☐ Yes ☐ No ☐ Don't Know	Was plasma ASA level Elevated Normal Unknown Was Citrulline level Elevated Normal Unknown
Were plasma urine acids tested? Yes No Don't Know	Was urine ASA level? Elevated Normal Unknown Was urine Citrulline level? Elevated Normal Unknown
Was a mutation analysis performed for ASA? Yes No Don't Know 	What genes were included in the mutation analysis? ASL Other gene:

MOLECULAR GENETICS REPORT	
Were variants detected in the ASL gene?	Check the types of variants found on:
□ Yes	Allele 1:
□ No	Variant known to be disease causing
Don't Know	Variant of unknown significance
	 Predicted to be pathogenic
	□ Wild Type (Normal)
	Allele 2:
	Variant known to be disease causing
	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	□ Unknown
Were variants detected in other genes?	Check the types of variants found on:
□ Yes	Allele 1:
□ No	Variant known to be disease causing
Don't Know	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	□ Unknown
	Allele 2:
	Variant known to be disease causing
	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	Unknown
Was enzyme analysis for ASA enzyme activity	Was enzyme activity:
completed?	Consistent with disease
□ Yes	Normal activity (not consistent with disease)
□ No	Unknown
Don't Know	

Citrullinemia Type I (CIT) Case Confirmatory Diagnosis Follow-Up

Please answer the following as Yes/No/Don't Know	If Yes
Were plasma amino acids tested? □ Yes □ No □ Don't Know	Was plasma ASA: Present Absent Unknown Was Citrulline level Elevated Normal Unknown
Was blood ammonia tested? Yes No Don't Know	Was blood ammonia level Elevated Normal Unknown
Was a mutation analysis performed for Citrullinemia type I? Yes No Don't Know	What genes were included in the mutation analysis? ASS1 gene Other gene:

MOLECULAR GENETICS REPORT	
Were variants detected in the ASS1 gene? Yes No Don't Know	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance • Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance • Predicted to be pathogenic Wild Type (Normal) Unknown
Were variants detected in other genes? Yes No Don't Know 	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown
Was enzyme analysis for Cirtullinemia type-I enzyme activity completed? Ves No Don't Know	Was enzyme activity: Consistent with disease Normal activity (not consistent with disease) Unknown

Cystathionine Beta-Synthase (CBS) Deficiency (Classic Homocystinuria) Case Confirmatory Diagnosis Follow-Up

Please answer the following as Yes/No/Don't Know	If Yes
Were plasma amino acids tested? □ Yes □ No □ Don't Know	Was Methionine level Elevated Normal Unknown
Was plasma Homocysteine tested? Yes No Don't Know 	Was plasma Homocysteine level Elevated Normal Unknown
Was a mutation analysis performed for CBS? Yes No Don't Know	What genes were included in the mutation analysis? CBS gene Other gene:

Molec	CULAR GENETICS REPORT
Were variants detected in the CBS gene?	Check the types of variants found on:
□ Yes	Allele 1:
□ No	Variant known to be disease causing
Don't Know	Variant of unknown significance
	 Predicted to be pathogenic
	□ Wild Type (Normal)
	□ Unknown
	Allele 2:
	Variant known to be disease causing
	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	Unknown
Were variants detected in other genes?	Check the types of variants found on:
□ Yes	Allele 1:
□ No	Variant known to be disease causing
Don't Know	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	□ Unknown
	Allele 2:
	Variant known to be disease causing
	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	Unknown
Was enzyme analysis for CBS enzyme activity	Was enzyme activity:
completed?	Consistent with disease
□ Yes	Normal activity (not consistent with disease)
□ No	Unknown
Don't Know	

Maple Syrup Urine Disease (MSUD) Case Confirmatory Diagnosis Follow-Up

Instructions: Please answer the questions as fully as possible. All testing refers to confirmatory testing following newborn screening. You may need to consult with relevant clinician(s) and/or medical staff to obtain this information.

Part I: Final Diagnosis as determined by metabolic geneticist or clinician performing the follow-up

- A. MAPLE SYRUP URINE DISEASE, TYPE IA
- B. MAPLE SYRUP URINE DISEASE, TYPE IB
- C. MAPLE SYRUP URINE DISEASE, TYPE II
- D. MAPLE SYRUP URINE DISEASE, TYPE III

Please answer the following as Yes/No/Don't Know	If Yes
Were plasma amino acids tested? Yes Don't Know	Was Alloisoleucine level Elevated Unknown Was Leucine level Elevated Normal Unknown Was Isoeucine level Elevated Normal Unknown Was Isoeucine level Elevated Normal Unknown Was Valine level Elevated Normal Unknown Was Leu>Val? Yes No Unknown

Were Urine organic acids tested? Yes No Don't Know	Was 2-ketoisocaproic acid level Elevated Normal Unknown Was 2-OH Isovaleric acid level Elevated Normal Unknown Was 2-ketomethyl valeric acid level Elevated Normal Unknown Unknown Was 2-ketomethyl valeric acid level Unknown Unknown Unknown
Was maternal 3-MCC level tested and ruled out? Yes No Don't Know 	
Was a mutation analysis performed for MSUD? Yes No Don't Know 	 What genes were included in the mutation analysis? DBT BCKDHB DLD BCKDHA Other:
Molec	CULAR GENETICS REPORT
Were variants detected in DBT? Yes No Don't Know	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown

Were variants detected in BCKDHD?	Check the types of variants found on:
□ Yes	Allele 1:
🗆 No	Variant known to be disease causing
Don't Know	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	Unknown
	Allele 2:
	Variant known to be disease causing
	Variant of unknown significance
	 Predicted to be pathogenic
	□ Wild Type (Normal)
	Unknown
Were variants detected in <i>DLD</i> ?	Check the types of variants found on:
□ Yes	Allele 1:
□ No	Variant known to be disease causing
Don't Know	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	Unknown
	Allele 2:
	Variant known to be disease causing
	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	Unknown
Were variants detected in BCKDHA?	Check the types of verients found on
	Check the types of variants found on:
Yes	Allele 1:
	Variant known to be disease causing
Don't Know	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	Unknown
	Allele 2:
	Variant known to be disease causing
	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	□ Unknown
	<u> </u>

Were variants detected in Other genes?	Check the types of variants found on:
□ Yes	Allele 1:
□ No	Variant known to be disease causing
Don't Know	Variant of unknown significance
	Predicted to be pathogenic
	Wild Type (Normal)
	Unknown
	Allele 2:
	Variant known to be disease causing
	Variant of unknown significance
	Predicted to be pathogenic
	Wild Type (Normal)
	Unknown
Was enzyme analysis for MSUD enzyme activity	Was enzyme activity:
completed?	Consistent with disease
□ Yes	Normal activity (not consistent with disease)
□ No	Unknown
Don't Know	

Hyperphenylalaninemia (HyperPHE) (Inclusive of Classic PKU) Case Confirmatory Diagnosis Follow-Up

Instructions: Please answer the questions as fully as possible. All testing refers to confirmatory testing following newborn screening. You may need to consult with relevant clinician(s) and/or medical staff to obtain this information.

Final diagnosis as determined by a metabolic geneticist or clinician performing the follow-up:

- A. Classic PKU
- B. Benign HyperPhe
- C. HyperPhe diet controlled

Please answer the following as Yes/No/Don't Know	If Yes
Were plasma amino acids tested? Yes No Don't Know	Was Phe level Elevated (>120umol/L on unrestricted diet) Normal Unknown Was Phe/Tyr ratio Elevated Normal Unknown
Were biopterin studies done? Yes No Don't Know	Were biopterin studies: Normal Abnormal Unknown
Was a mutation analysis performed for Hyperphenylalaninemia? Yes No Don't Know	What genes were included in the mutation analysis? PAH gene Other gene:

MOLECULAR GENETICS REPORT	
Were variants detected in the PAH gene?	Check the types of variants found on:
□ Yes	Allele 1:
□ No	Variant known to be disease causing
Don't Know	Variant of unknown significance
	 Predicted to be pathogenic
	□ Wild Type (Normal)
	Allele 2:
	Variant known to be disease causing
	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	🗖 Unknown
Were variants detected in other genes?	Check the types of variants found on:
□ Yes	Allele 1:
🗆 No	Variant known to be disease causing
Don't Know	Variant of unknown significance
	 Predicted to be pathogenic
	□ Wild Type (Normal)
	□ Unknown
	Allele 2:
	Variant known to be disease causing
	Variant of unknown significance
	 Predicted to be pathogenic
	Wild Type (Normal)
	Unknown
Was enzyme analysis for Hyperphe (inclusive of	Was enzyme activity:
classic PKU) enzyme activity completed?	Consistent with disease
□ Yes	Normal activity (not consistent with disease)
	Unknown
🗖 Don't Know	

Tyrosinemia Type I (TYR-1) Case Confirmatory Diagnosis Follow-Up

Instructions: Please answer the questions as fully as possible. All testing refers to confirmatory testing following newborn screening. You may need to consult with relevant clinician(s) and/or medical staff to obtain this information.

Please answer the following as Yes/No/Don't Know	If Yes
Were plasma organic acids tested? Yes No Don't Know Were urine organic acids tested? Yes No Don't Know	Was plasma Succinylacetone level Elevated Vormal Unknown Was plasma tyrosine level Elevated Normal Elevated Normal Unknown Was urine tyrosine level Elevated Normal Unknown Was urine tyrosine level Elevated Normal Unknown Unknown
Was mutation analysis performed for Tyrosinemia Type I? Yes No Don't Know	What genes were included in the mutation analysis?

Molecular Genetics Report	
Were variants detected in FAH? Yes No Don't Know	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown
Were variants detected in Other genes? Yes No Don't Know 	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown
Was enzyme analysis for Tyrosinemia Type I enzyme activity completed? Yes No Don't Know	Was enzyme analysis: Consistent with disease Normal activity (not consistent with disease) Unknown

Endocrine Disorders

Congenital Hypothyroidism (CH) Case Confirmatory Diagnosis Follow-Up

Instructions: Please answer the questions as fully as possible. All testing refers to confirmatory testing following newborn screening. You may need to consult with relevant clinician(s) and/or medical staff to obtain this information.

Final diagnosis as determined by the endocrinologist or clinician performing the follow-up:

- A. Primary Congenital Hypothyroidism
- B. Secondary Congenital Hypothyroidism
- C. TBG Deficiency (Thyroxine Binding Globulin) or other protein binding defect

Please answer the following as Yes/No/Don't Know	If Yes
Was Serum TSH tested? Yes No Don't Know	What was the level: TSH > 10 mU/L TSH 6-10 mU/L TSH <6 mU/L Unknown Was it tested before initiation of treatment? Yes No Unknown
Was Serum Total T4 tested? Yes No Don't Know	 Unknown Was Serum Total T4 below the age-established reference range? Yes No Unknown Was it tested before initiation of treatment? Yes No Unknown
Was Serum <u>Free</u> T4 tested? Yes No Don't Know	 Was Serum Free T4 below the age-established reference range? Yes No Unknown Was it tested before initiation of treatment? Yes No Unknown

Does this baby have other pituitary hormone deficiencies? Yes No Don't Know	
Does this baby have midline defects? Yes No Don't Know	
Was TBG tested? Yes Don't Know Was T3 or T4 resin uptake tested? Yes No Don't Know	 Was TBG below the age established reference range? Yes No Unknown Was T3 or T4 resin uptake above the age established reference range? Yes No Unknown

Congenital Adrenal Hyperplasia (CAH) Case Confirmatory Diagnosis Follow-Up

Instructions: Please answer the questions as fully as possible. All testing refers to confirmatory testing following newborn screening. You may need to consult with relevant clinician(s) and/or medical staff to obtain this information.

Final diagnosis as determined by the endocrinologist or clinician performing the follow-up:

- A. Classic 21-Hydroxylase Deficiency Salt Wasting
- B. Classic 21-Hydroxylase Deficiency Simple Virilizing
- C. Other Adrenal disorder: (Please list) _____

Please answer the following as Yes/No/Don't Know	If Yes
Was a confirmatory serum 17-OHP level obtained? Yes No Don't Know	Was there a value at baseline: 10,000 ng/dl; 1000-10,000 ng/dl; 4 1000 ng/dl; Unknown Was it tested before initiation of treatment? Yes No Was there a result after ACTH stimulation: 10,000 ng/dl; 1000-10,000 ng/dl; 4 1000 ng/dl; Unknown Was it tested before initiation of treatment? Yes No
Was tandem mass spectrometry urinary steroid profile obtained? Ves No Don't Know	 Were the urinary spectrometry steroid profile results: Indicative of 21-Hydroxylase Deficiency CAH Unknown

Was serum sodium level measured before initiation of treatment? Yes No Don't Know Was Plasma renin activity level measured at time of initiation of treatment? Yes No Don't Know	Was the sodium level: <pre> <pre> </pre> < 135 mEq/L > 135 mEq/L Unknown Was it tested before initiation of treatment? </pre> Yes No Was the Plasma renin activity normal for age? Yes No Was it tested before initiation of treatment? Yes No Was it tested before initiation of treatment? Yes No Was it tested before initiation of treatment? Yes No Was it tested before initiation of treatment?
Mouro	ULAR GENETICS REPORT
Wolec Was mutation analysis for 21-Hydroxylase deficiency (CYP21A2) performed? Yes No Don't Know	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown
	ClinicalResults
Is there evidence of salt wasting? (e.g. shock or severe failure to thrive)? Yes No Don't Know	
Is there supportive clinical or laboratory evidence of CAH? Yes No Don't Know	Is the evidence: (check all that apply) Ambiguous genitalia, with 46,XX karyotype Normal genitalia, with 46,XY karyotype Other hormonal evidence of CAH

Hemoglobinopathies

Presence of Hb S Case Confirmatory Diagnosis Follow-Up

Instructions: Please answer the questions as fully as possible. All testing refers to confirmatory testing following newborn screening. You may need to consult with relevant clinician(s) and/or medical staff to obtain this information

Please answer the following:	If Yes
 Final Diagnosis as determined by a clinician performing the follow-up S, Beta + Thalassemia – Hb S/B + Th S, C disease – Hb S/C 	
 Sickle Cell Disease, Hb S only S,S Disease (Sickle Cell Anemia) – Hb SS S, Beta 0-thalassemia – Hb S/B0Th Not Known S, other 	
Was qualitative (IEF or HPLC) testing completed? Yes No Unknown	What were the results? FS FSC FSA FSA ₂ FSAA ₂
	□ Other □ Unknown
Was quantitative (HPLC or electrophoresis) testing completed? Yes No Unknown	What were the results? FS FSC FS with high A ₂ FSA with high A ₂ FSA Other Unknown

Was mutation analysis performed? Yes No Unknown	Check the type of variant found on allele 1: S C Beta + Thal Beta ⁰ Thal Other Unknown Check the type of variant found on allele 2: S C Beta + Thal Beta ⁰ Thal Other Unknown
NBS result Yes No Unknown	What were the results? FS FSC FSA FSA ₂ Other Unknown
Was a CBC performed? Yes No Unknown	What were the results? Normal – high MCV Low MCV Unknown
Were family studies (in parents) done? Yes No Unknown	What were the results? Maternal Status: Carrier S Carrier C Carrier Beta + Thal Carrier Beta ⁰ Thal Other Unknown Paternal Status: Carrier S Carrier C Carrier Beta + Thal Carrier Beta + Thal Carrier Beta ⁰ Thal Other Unknown

Was there a positive family history? Yes No Unknown 	
Were HPLC & IEF tested on the same sample from the infant? Yes No Unknown	What were the results? FS FSC FSA ₂ FSAA ₂ Other Unknown
Were Hgb tests (electrophoresis or HPLC) performed on family members? Yes No Unknown	What were the results? Positive Negative Unknown

Presence of Other Hb Variant Case Confirmatory Diagnosis Follow-Up

Instructions: Please answer the questions as fully as possible. All testing refers to confirmatory testing following newborn screening. You may need to consult with relevant clinician(s) and/or medical staff to obtain this information.

Please answer the following:	If Yes
 Final Diagnosis as determined by a clinician performing the follow-up Hemoglobin C disease Hemoglobin D disease Hemoglobin E disease Hemoglobin O-Arab disease Other hemoglobin disease, please describe 	
Alpha thalassemia present? Yes No Unknown	
Was qualitative (IEF or HPLC) testing completed? ☐ Yes ☐ No ☐ Unknown	What were the results? FC FD FE FO _{ARAB} Other Unknown
Was quantitative (HPLC or electrophoresis) testing completed? Yes No Unknown	What were the results? FC FD FE FO _{ARAB} Other Unknown

Was mutation analysis performed? Yes No Unknown 	Check the type of variant found on allele 1: C D E
	 O-Arab Other Unknown Check the type of variant found on allele 2: C D E O-Arab Beta + Thal Beta⁰ Thal Other Unknown
NBS result Yes No Unknown	What were the results? FC FD FE FO _{ARAB} Other Unknown
Was a CBC performed? Yes No Unknown	What were the results? Normal – high MCV Low MCV Unknown

Were family studies (in parents) done? Yes No Unknown	What were the results? Maternal Status: Carrier C Carrier D Carrier E Carrier O-Arab Carrier Beta + Thal
	 Carrier Beta⁰ Thal Other Unknown Paternal Status:
	Carrier C Carrier D Carrier E Carrier O-Arab Carrier Beta + Thal
	 Carrier Beta⁰ Thal Other Unknown
Was there a positive family history? Yes No Unknown 	
Were Hgb tests (electrophoresis or HPLC) performed on family members? Yes No Unknown	What were the results? Positive Negative Unknown

Lysosomal Storage Disorders

Mucopolysaccharidosis Type I (MPS I) Case Confirmatory Diagnosis Follow-Up

Birth weight (grams):

Gestational age (weeks gestation at time of birth):

State of birth (state reporting the case):

Sex (male/female/unknown):

Instructions: Please answer the questions as fully as possible. All testing refers to confirmatory testing following newborn screening. You may need to consult with relevant clinician(s) and/or medical staff to obtain this information.

Final diagnosis as determined by a metabolic geneticist or clinician performing follow-up:

- 1. Primary targets of NBS
 - a. MPS I -severe
 - b. MPS I severity not determined
 - c. MPS I attenuated

Please answer the following as Yes/No/Unknown	If Yes
Was enzyme activity tested? Yes No Unknown	What was the enzyme level? U Within lab known affected range Unknown
Were urine GAGS tested? Yes No Unknown	What was the urine GAG level? Elevated Normal Unknown

 Clinical symptoms/ lab findings? Symptoms present and documented by specialists. Public health (PH) program continued to collect data through the development of symptoms 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 Unknown 	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, hernia
Molecu	JLAR GENETICS REPORT
Were variants detected in the genes known to be associated with MPS I? Yes No Unknown	Check the types of variant(s) found on: Allele 1: Pathogenic variant and associated with SEVERE disease Pathogenic or likely pathogenic variant Variant known to be associated with ATTENUATED Disease Wild Type (Normal) Untested/Unknown Allele 2: Pathogenic variant and associated with SEVERE disease Pathogenic or likely pathogenic variant Variant of uncertain significance Variant known to be associated with ATTENUATED Disease Wild Type (Normal) Untested/Unknown

Pompe Disease Case Confirmatory Diagnosis Follow-Up

Birth weight (grams):

Gestational age (weeks gestation at time of birth):

State of birth (state reporting the case):

Sex (male/female/unknown):

Instructions: Please answer the questions as fully as possible. All testing refers to confirmatory testing following newborn screening. You may need to consult with relevant clinician(s) and/or medical staff to obtain this information.

Final diagnosis as determined by metabolic geneticist or clinician performing the follow-up:

- 1. Primary targets of NBS
 - a. Infantile Onset (IO) Pompe Disease
 - b. Late Onset (LO) Pompe Disease

Please answer the following as Yes/No/Don't Know	If Yes
Was enzyme activity tested in blood (not DBS sample)? Yes No Unknown	 What was the enzyme level? Within lab known affected range for infantile onset (IO) Low (above affected range for IO, may or may not be in late- onset (LO) range but should not be above LO range)) Within lab known affected range for late onset (LO) Low (above affected range, for LO not normal) Unknown
Was enzyme activity tested in skin/muscle? Yes No Unknown	 Positive skin or muscle biopsy Unknown
Was there Cardiac involvement consistent with Pompe? Yes No Unknown	 Positive findings on Chest X-ray/EKG/ECHO in newborn period Positive findings on Chest X-ray/EKG/ECHO

 Lab findings? Elevated CK/AST/ALT/LDH/Urine Hex4 Not present Unknown Not done Clinical findings? Symptoms present after 1 year of age and documented by specialists. PH program continued to collect data through the development of symptoms Symptoms present before 1 year of age but no cardiac involvement Unknown or not reported to PH by the end of follow-up 	Clinical symptoms consistent with Pompe Disease: progressive muscle weakness, need for respiratory assistance, swaying gait or waddle, Lordosis, kyphosis, or scoliosis
Were variants detected in the genes known to be	Check the types of variant(s) found on:
associated with Pompe Disease?	Allele 1:
NO Unknown	 Pathogenic variant and associated with infantile onset Novel variant that is likely pathogenic Pathogenic variant or likely pathogenic variant, with deletion or duplication consistent with infantile onset Pathogenic and associated with non-classical disease, or variant of uncertain significance Pathogenic or likely pathogenic variant, no other variants found, dup/del testing not done or not known Pathogenic or likely pathogenic variant, no other variants found Wild Type (Normal) Unknown Allele 2: Pathogenic variant and associated with infantile onset Novel variant that is likely pathogenic Pathogenic variant or likely pathogenic Pathogenic or likely pathogenic Pathogenic or likely pathogenic Pathogenic variant and associated with infantile onset Novel variant that is likely pathogenic Pathogenic or likely pathogenic variant, with deletion or duplication consistent with infantile onset Pathogenic or likely pathogenic variant, with infantile onset Pathogenic or likely pathogenic variant, with infantile onset Pathogenic or likely pathogenic variant, no other variants found, dup/del testing not done or not known Pathogenic or likely pathogenic variant, no other variants found, dup/del testing not done or not known Pathogenic or likely pathogenic variant, no other variants found Wild Type (Normal) Unknown

Other Disorders

Biotinidase Deficiency Case Confirmatory Diagnosis Follow-Up

Instructions: Please answer the questions as fully as possible. All testing refers to confirmatory testing following newborn screening. Attach a copy of the laboratory reports supporting these findings. The reports will be stored within the newborn screening program for future reference.

Final Diagnosis as determined by metabolic geneticist or clinician performing the follow-up

- D. Profound Biotinidase deficiency
- E. Partial Biotinidase deficiency

Was enzyme analysis for biotinidase enzyme activity completed? Yes No Don't Know	 What as the enzyme activity? <10% normal activity 10-30% normal activity Normal Unknown
Was mutation analysis performed for Biotinidase deficiency? Pres Don't Know	Check the types of variants found on: Allele 1: Variant known to be disease causing Known to be associated with profound enzyme deficiency Known to be associated with partial enzyme deficiency ['mild' mutation (D444H)] Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Known to be associated with profound enzyme deficiency Known to be associated with partial enzyme deficiency ['mild' mutation (D444H)] Variant of unknown significance o Predicted to be pathogenic Wild Type (Normal) Unknown

Galactosemia Case Confirmatory Diagnosis Follow-Up

Instructions: Please answer the questions as fully as possible. All testing refers to confirmatory testing after newborn screening. Attach a copy of the laboratory reports supporting these findings. The reports will be stored within the newborn screening program for future reference.

Please answer the following as Yes/No/Don't Know	If Yes
Were GALT levels tested? Yes No Don't Know	Was GALT level <10%
Were Gal-1-P tested? Ves No Don't Know	Was Gal-1-P level Elevated Normal Unknown
Was Urine Galactitol tested? Yes No Don't Know	Was Gal-1-P level Elevated Normal Unknown
Was a mutation analysis performed for Galactosemia? Yes No Don't Know	Check the types of variants found on: Allele 1: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Wild Type (Normal) Unknown Allele 2: Variant known to be disease causing Variant of unknown significance Predicted to be pathogenic Vild Type (Normal) Unknown
If Variant Galactosemia, was protein phenotyping completed? Yes No Don't Know N/A	 Did result indicate phenotype consistent with variant phenotype NOT consistent with variant Unknown

If Arginase Deficiency, were enzyme studies completed?	Was enzyme activity:		
□ Yes	 Consistent with disease Normal activity (not consistent with disease) 		
 No Don't Know 	□ Unknown		
□ N/A			

Cystic Fibrosis Case Confirmatory Diagnosis Follow-Up

Instructions: Please answer the questions as fully as possible. All testing refers to confirmatory testing following newborn screening. Attach a copy of the laboratory reports supporting these findings. The reports will be stored within the newborn screening program for future reference.

- D. Typical Cystic Fibrosis (CF)
- E. CFTR-Related Metabolic Syndrome (CRMS)
- F. CFTR-Related Disease

Please answer the following as Yes/No/Don't Know	If Yes
Did the NBS result indicate an elevated IRT? Yes No Don't Know 	
 Were CFTR mutations detected on the <u>newborn</u> <u>screening</u> mutation panel? Yes No Don't Know (* Mutations seen in patients with CF have been classified as disease-causing, neutral, or varying clinical consequences through the CFTR2 project: <u>http://cftr2.org/browse.php</u>. Additional information about the mutation and the association with lower sweat chlorides can also be found at CFTR2.) 	Check the types of variants found on: Allele 1: Variant known to be disease causing in CFTR2 Shown to be associated with lower sweat chlorides Neutral variant Variant of varying clinical consequence in CFTR2 Wild Type (Normal) Unknown (not reported in CFTR2) Allele 2: Shown to be disease causing in CFTR2 Shown to be associated with lower sweat chlorides Neutral variant Variant of varying clinical consequence in CFTR2 Wild Type (Normal) Unknown (not reported in CFTR2)
Did the child have meconium ileus? Yes No Don't Know	

Was a valid sweat chloride result available? Yes No Don't Know	What were the sweat test results (please report the highest sweat chloride value from one sweat test)? >=60 mmol/L (regardless of age) If <60 mmol/L If age< 6 months is a start of the selfage.com/lemondle is a start of the selfage.com/lemondle is a start of the selfage.com/lemondle If <60 mmol/L If age is a start of the selfage.com/lemondle If age is a start of the selfage.com/lemondle If age is a start of the selfage.com/lemondle If age If age is a start of the selfage.com/lemondle If age If age <
If a valid sweat test was not available, were there attempts to obtain a sweat chloride that were quantity not sufficient (QNS)? Yes No Don't Know	
Was a sweat chloride repeated on a separate day (results from different arm on the same day should not be reported here)? Yes No Don't Know	What were the repeat sweat test results (please report the highest sweat chloride value from one sweat test)? >=60 mmol/L (regardless of age) If <60 mmol/L If age< 6 months <pre></pre>

Was a CFTR mutation panel completed after the	Check the types of variants found on:
newborn screening mutation panel?	Allele 1:
 Yes No Don't Know (* Mutations seen in patients with CF have been classified as disease-causing, neutral, or varying clinical consequences through the CFTR2 project: <u>http://cftr2.org/browse.php</u>. Additional information about the mutation and the association with lower sweat chlorides can also be found at CFTR2.) 	 Variant known to be disease causing in CFTR2 Shown to be associated with lower sweat chlorides Neutral variant Variant of varying clinical consequence in CFTR2 Wild Type (Normal) Unknown (not reported in CFTR2) Allele 2: Variant known to be disease causing in CFTR2 Shown to be associated with lower sweat chlorides Neutral variant Variant of varying clinical consequence in CFTR2 Wild Type (Normal) Unknown (not reported in CFTR2)
Question Below to be answered on	ly if child was diagnosed after the newborn period
If child was diagnosed after the newborn period, were clinical symptoms associated with CFTR Related Disease present? Yes No Don't Know	Symptoms include: CBAVD Recurrent pancreatitis Nasal polyposis Infertility Focal biliary cirrhosis with portal hypertension

Summary of common variants as reported on CFTR2 (this is not an exhaustive list; please visit <u>www.CFTR2.org</u> for the latest updated list).

Variant name - HGVS nomenclature	Protein name	Variant legacy name	On ACMG Screening Panel	CFTR2 final call	Associated with lower sweat chloride
c.3717+12191C>T	p.Phe316LeufsX12	1078delT	No	CF-causing	NO
c.579+3A>G	p.Phe342HisfsX28	1154insTC	No	CF-causing	NO
c.3454G>C	No protein name	1717-1G->A	Yes	CF-causing	NO
c.3208C>T	No protein name	1811+1.6kbA->G	No	CF-causing	NO
c.3154T>G	No protein name	1898+1G->A	Yes	CF-causing	NO
c.1585-1G>A	p.Leu671X	2143delT	No	CF-causing	NO
c.1680-1G>A	p.Lys684SerfsX38	2183AA->G	No	CF-causing	NO
c.1766+1G>A	p.Lys684AsnfsX38	2184delA	Yes	CF-causing	NO
c.2490+1G>A	p.Gln685ThrfsX4	2184insA	No	CF-causing	NO
c.2988+1G>A	p.Glu726ArgfsX4	2307insA	No	CF-causing	NO
c.1736A>G	No protein name	2789+5G->A	Yes	CF-causing	NO
c.1408A>G	No protein name	3120+1G->A	Yes	CF-causing	NO
c.1841A>G	No protein name	3120G->A	No	CF-causing	NO
c.2991G>C	No protein name	3272-26A->G	No	CF-causing	NO
c.489+1G>T	p.Lys1177SerfsX15	3659delC	Yes	CF-causing	NO
c.350G>A	No protein name	3849+10kbC->T	Yes	CF-causing	NO
c.4242+1G>T	p.Leu1258PhefsX7	3905insT	No	CF-causing	NO
c.3718-1G>A	p.Leu88llefsX22	394delTT	No	CF-causing	NO
c.1240C>T	No protein name	5T	No	Indeterminat e	YES
c.2260G>A	No protein name	621+1G->T	Yes	CF-causing	NO
c.1727G>C	No protein name	711+1G->T	Yes	CF-causing	NO
c.220C>T	No protein name	711+5G->A	No	CF-causing	NO
c.2834C>T	p.Ala455Glu	A455E	Yes	CF-causing	NO
c.1675G>A	p.Ala559Thr	A559T	No	CF-causing	NO
c.1127 1128insA	p.Ser18ArgfsX16	CFTRdele2,3	No	CF-causing	NO
c.1202G>A or c.1203G>A	p.Asp1152His	D1152H	No	Indeterminat e	YES
c.1923 1931del9insA	p.Glu60X	E60X	No	CF-causing	NO
c.1679G>C	p.Phe508del	F508del	Yes	CF-causing	NO
c.3160C>G	p.Gly1244Glu	G1244E	No	CF-causing	NO
c.4046G>A	p.Gly178Glu	G178R	No	CF-causing	NO
c.4196_4197delTC	p.Gly542X	G542X	Yes	CF-causing	NO
c.3731G>A	p.Gly551Asp	G551D	Yes	CF-causing	NO
c.3197G>A	p.Gly85Glu	G85E	Yes	CF-causing	NO
c.2657+2_2657+3insA	p.lle1027Thr	I1027T	No	Not CF- causing	NO
c.1673T>C	p.lle148Thr	I148T	No	Not CF- causing	NO

c.3763T>C	p.Ile336Lys	I336K	No	CF-causing	NO
c.1558G>T	p.lle507del	1507del	Yes	CF-causing	NO
c.3230T>C	p.Leu1077Pro	L1077P	No	CF-causing	NO
c.1040G>A	p.Leu206Trp	L206W	No	CF-causing	NO
c.3302T>A	p.Met1101Lys	M1101K	No	CF-causing	NO
c.274G>A	p.Asn1303Lys	N1303K	Yes	CF-causing	NO
c.617T>G	p.Pro67Leu	P67L	No	CF-causing	NO
c.2764_2765insAG	p.Gln220X	Q220X	No	CF-causing	NO
c.1973_1985del13insAGAA A	p.Gln493X	Q493X	No	CF-causing	NO
c.3196C>T	p.Arg1066Cys	R1066C	No	CF-causing	NO
c.4296_4297insGA	p.Arg1158X	R1158X	No	CF-causing	NO
c.1692delA	p.Arg1162X	R1162X	Yes	CF-causing	NO
c.1055G>A	p.Arg117Cys	R117C	No	CF-causing	NO
c.1466C>A	p.Arg117His	R117H	Yes	Indeterminat e	YES
c.1013C>T	p.Arg334Trp	R334W	Yes	CF-causing	NO
c.532G>A	p.Arg347His	R347H	Yes	CF-causing	NO
c.1040G>C	p.Arg347Pro	R347P	No	CF-causing	NO
c.2908G>C	p.Arg352Gln	R352Q	No	CF-causing	NO
c.2424_2425insAT	p.Arg553X	R553X	Yes	CF-causing	NO
c.2780T>C	p.Arg560Thr	R560T	Yes	CF-causing	NO
c.349C>T	p.Ser1251Asn	S1251N	No	CF-causing	NO
c.1000C>T	p.Ser549Asn	S549N	No	CF-causing	NO
c.3752G>A	p.Ser945Leu	S945L	No	CF-causing	NO
c.1645A>C or c.1647T>G	p.Val520Phe	V520F	No	CF-causing	NO
c.274G>T	p.Trp1282X	W1282X	Yes	CF-causing	NO
c.2128A>T	p.Tyr1092X	Y1092X	No	CF-causing	NO
c.2195T>G	p.Tyr122X	Y122X	No	CF-causing	NO

Severe Combined Immunodeficiencies Case Confirmatory Diagnosis Follow-Up

Birth weight (grams):

Gestational age (weeks gestation at time of birth):

State of birth (state reporting the case):

Sex (male/female/unknown):

Instructions: Please answer the questions as fully as possible. All testing refers to confirmatory testing following newborn screening. You may need to consult with relevant clinician(s) and/or medical staff to obtain this information.

Final diagnosis as determined by a metabolic geneticist or clinician performing the follow-up:

- 1. Primary targets of NBS
 - a. Classic SCID
 - b. Leaky SCID
 - c. Omenn Syndrome

Please answer the following as Yes/No/Don't Know	If Yes
Was the CD3 T cell level tested?	What was the CD3 T cell level?
 Yes No Unknown/untested 	 <300 autologous T cells, undetectable or very few naïve T cells 300-1500, few naïve T cells, oligoclonal T cells or poor T cell diversity >80% CD45RO+ Any number (not zero) Untested/Unknown
Was proliferation to PHA test done?	Proliferation to PHA:
 Yes No Unknown/untested 	 <10% of normal 10-50% of normal PHA 10-30% normal PHA or Absent to Candida/TT <30% of normal Any/Unknown

Was Maternal engraftment documented?	
 □ Yes □ No □ Unknown/Untested 	
Was a mutation analysis performed in the genes known to be associated with SCID?	
☐ Yes☐ No☐ Unknown/untested	
Molec	ULAR GENETICS REPORT
Were variants detected in the genes known to be associated with SCID? Yes No Unknown/untested	 Check the types of variant(s) found on: Allele 1: Pathogenic variant in a known SCID gene Pathogenic variant in a known SCID gene on X chromosome in a male Pathogenic variant in a known SCID gene known to be associated with leaky SCID (previously reported or in a gene previously associated with combined immunodeficiency) Wild Type (Normal)
	Allele 2:
	 Pathogenic variant in a known SCID gene known to be associated with leaky SCID (previously reported or in a gene previously associated with immunodeficiency) Wild Type (Normal) Untested/Unknown
	Was 22q11 deletion ruled out?
	□ Yes □ No □ Don't Know

Were homozygous or compound heterozygous FOXN1 mutations ruled out? Yes No Don't Know
Were heterozygous TBX1 variants ruled out? Yes No Don't Know

Critical Congenital Heart Disease Case Confirmatory Diagnosis Follow-Up

Instructions: Please answer the questions as fully as possible. All testing refers to confirmatory testing following newborn screening. Attach a copy of the laboratory reports supporting these findings. The reports will be stored within the newborn screening program for future reference.

What was the Final Diagnosis? (Check all that apply)

- □ CCHD (Specify)
 - □ Truncus Arteriosus
 - Total Anomalous Pulmonary Venous Connection
 - □ Tetralogy of Fallot
 - Pulmonary Atresia
 - Ebstein's Anomaly
 - □ Hypoplastic Left Heart Syndrome
 - □ Single Ventricle
 - Tricuspid Atresia
 - □ Transposition of the Great Arteries
 - Double Outlet Right Ventricle
 - Coarctation of Aorta
 - □ Interrupted Arch
 - □ Aortic Valve Disease
- □ Non critical CHD (Please specify)
- □ Other (Please specify)

Please answer the following:	If Yes, what were the re	esults of the echocardiogram?	
Was a Postnatal Echocardiogram Completed? Yes No Unknowm/untested	Truncus Arteriosus Truncus arteriosus Truncus arteriosus + Interrupted aortic arch Total Anomalous Pulmonary Venous Connection (TAPVC)		
	 Type1 (supracardiac) Type 2 (cardiac) Type 3 (infracardiac) Type 4 (mixed) 		
	Tetralogy of Fallot (TOF)		
	 TOF TOF, Pulmonary stenosis TOF, AVCanal (AVSD) TOF, Absent pulmonary valve Interrupted aortic arch + AP windov 	v (aortopulmonary window)	
	 Pulmonary Artesia Pulmonary atresia Pulmonary atresia, IVS Pulmonary atresia, VSD (Including T Pulmonary atresia, VSD-MAPCA 	OF, PA)	
	Ebstein's Anomaly Ebstein's anomaly		
	Hypoplastic Left Heart Syndrome (HLHS) Hypoplastic left heart syndrome)	
	 Single Ventricle Single ventricle, DILV Single ventricle, DIRV Single ventricle, Mitral atresia Single ventricle, Unbalanced AV canal 	 Single ventricle, Heterotaxia syndrome Single ventricle, Other Single ventricle + Total anomalous pulmonary venous connection (TAPVC) 	
	Tricuspid Artesia Single ventricle, Tricuspid atresia		
	Transposition of the Great Arteries (TGA) d-TGA, IVS d-TGA, IVS-LVOTO d-TGA, VSD d-TGA, VSD	A)	

	Double Outlet Right Ventricle (DORV) DORV, VSD type	 DORV, Remote VSD (uncommitted VSD) DORV + AVSD (AV Canal) DORV + MC
	 DORV, TOF type DORV, TGA type 	DORV, IVS
	Coarctation of Aorta Coarctation of aorta Aortic arch hypoplasia VSD + Aortic arch hypoplasia VSD + Coarctation of aorta	
	Interrupted Arch ☐ Interrupted aortic arch ☐ Interrupted aortic arch + VSD ☐ Interrupted aortic arch + AP window	(aortopulmonary window)
	Aortic Valve Disease ☐ Aortic Stenosis receiving interventio ☐ Pulmonary Stenosis receiving interve	
Please answer the following:		lf Yes
Was a Prenatal Echocardiogram Completed? Yes No Don't Know	Did the Prenatal Echo findings suggest CC □ Yes □ No	HD?

X-Linked Adrenoleukodystrophy (X-ALD) Case Confirmatory Diagnosis Follow-Up

Birth weight (grams):

Gestational age (weeks gestation at time of birth):

State of birth (state reporting the case):

Sex (male/female/unknown):

Instructions: Please answer the questions as fully as possible. All testing refers to confirmatory testing following newborn screening. You may need to consult with relevant clinician(s) and/or medical staff to obtain this information.

Final diagnosis as determined by a metabolic geneticist or clinician performing the follow-up:

- 1. Primary targets of NBS
 - a. X-Linked Adrenoleukodystrophy (in Males)
 - b. Contiguous ABCD1 DXS1357E deletion syndrome (CADDS)
 - c. X-Linked Adrenoleukodystrophy (in Females)
 - d. Zellweger Spectrum Disorder
 - e. Peroxisomal Disorder
 - f. Acyl-CoA Oxidase Deficiency
 - g. D-Bifunctional Protein Deficiency
 - h. Dyamin-like protein 1 (DLP1)
 - i. ABDC5
 - j. Non-peroxisomal Disorder

Please answer the following as Yes/No/Don't Know	If Yes	
Was plasma VLCFA tested? Yes No Unknown	What was the VLCFA level? Elevated Slightly elevated Normal Low Unknown	
Clinical symptoms? Present Not present Not present at birth Unknown/ Not available 	Symptoms may include: neonatal hypotonia, neonatal seizures, liver disease, neonatal cholestasis, sensorineural deafness, failure to thrive, craniofacial abnormalities	
Was plasmalogen testing done? Yes No Unknown	Plasmalogen level? Normal Low Unknown/ Not available	
Family History done? Yes No Unknown	 Family history results: Family history present or family VLCFA studies suggestive of X-linked ALD Unknown 	
Were fibroblast studies done? Yes No Unknown	 Fibroblast study results: Consistent with Zellweger Spectrum Disorder Consistent with Acyl-CoA Oxidase Deficiency Consistent with D-Bifunctional Protein Consistent with DLP1 Consistent with ABCD5 Unknown 	
	JLAR GENETICS REPORT	
Were variants detected in the ABCD1 gene? Yes No Unknown/untested 	 Check the types of variant(s) found: Pathogenic variant Deletion/ duplication identified No mutation on sequencing, deletion/duplication not done No mutation on sequencing, deletion/duplication not done; rule out other disorders of peroxisomal beta oxidation Variant of uncertain significance Deletion identified in ABCD1 and DXS1357E Unknown 	

 Were variants detected in the PEX1 gene? Yes No Unknown/untested 	Check the types of variant(s) found: Two pathogenic variants in the PEX1 gene Unknown
 Were variants detected in the ACOX1 gene? Yes No Unknown/untested 	Check the types of variant(s) found: Two pathogenic mutations in the ACOX1 gene Unknown
 Were variants detected in the HSD17B4 gene? Yes No Unknown/untested 	Check the types of variant(s) found: Two pathogenic mutations in the HSD17B4 gene Unknown
Mutational analysis done on other genes? Yes No Unknown/untested 	 Check the types of variant(s) found: No mutation on sequencing, deletion/duplication not found Heterozygous, dominant-negative Two disease causing mutations Mutation in one of the 7 known genes for Aicardi-Goutiéres Syndrome Unknown