NEWBORN SCREENING REPORT

319/384-5097 or 866/890-5965

Patient		Birth	Date and Gender Cli	ent Reference	Accession #	
LAST10964566, FIRST10964566		4566 202	5-05-01 00:07 10)964566	78461	
		Fen	nale			
218 78TH ST		Colle	ected Re	ceived	Project	
ANYTOWN, IA 52999		202	025-05-02 00:30 2025-05-02 22:30			
					Ordering Health Care Provider	
	ANYTOWN HOSPITAL				WELBY, MARCUS	
6					Primary Care Provider	
รี SEND-OUT LAB				WELBY, MARCUS		
SEND-OUT LAB 1000 MEDICAL ST ANYTOWN, IA 52999					Sample Type	
					Blood spot specimen	
					Sample Note(s)	
Screen	Birth Order	IA Barcode Number	Transfusion Interference	Weight at Collection	Guardian	
Initial		IA2093764	No	2219 grams	MOMLAST10964566, MOMFIRST10964566	

RESULTS OF ANALYSIS - FINAL REPORT

TEST	RESULT	ANALYSIS NOTE(S)
Congenital Adrenal Hyperplasia, Immunoassay		
Congenital adrenal hyperplasia interpretation	Within Normal Limits	
Congenital Hypothyroidism, Immunoassay	MPd Schlamocal Libertia	
Congenital hypothyroidism interpretation	Within Normal Limits	
Biotinidase Deficiency, Immunoassay	MPd. S. Name at Liberta	
Biotinidase deficiency interpretation	Within Normal Limits	
Galactosemia, Enzymatic Assay	Within Normal Limita	
Galactosemia interpretation	Within Normal Limits	4
Hemoglobinopathies, Various Methods	Within Normal Limita	1
Hemoglobin disorders interpretation	Within Normal Limits	0
Cystic Fibrosis, Various Methods	Within Normal Limits	2
Cystic fibrosis interpretation	within Normai Limits	3
Expanded Screening Disorders, Tandem Mass Spectrometry	Within Normal Limits	3
Fatty acid oxidation defects interpretation	Within Normal Limits Within Normal Limits	
Organic acidemias interpretation Amino acidemias interpretation		
•	Within Normal Limits	1
Lysosomal Storage Disorders, Tandem Mass Spectrometry Pompe disease interpretation	Borderline - See second tier test result	4
Mucopolysaccharidosis Type 1 (MPS1)	Within Normal Limits	
disease interpretation	Within Normal Limits	
Mucopolysaccharidosis Type II (MPSII)	Within Normal Limits	
disease interpretation	Within Normal Emiles	
Pompe Disease 2nd Tier NBS, Flow Injection Analysis-Tanden	Mass Spectrometry	5
Pompe Disease 2nd Tier NBS interpretation	See second tier results report attached to the sam	ple in the
·	OpenELIS Web Portal	•
X-linked Adrenoleukodystrophy, Tandem Mass Spectrometry		6
X-linked Adrenoleukodystrophy disease	Within Normal Limits	
interpretation		
Severe Combined Immunodeficiency, Real-Time PCR		7
Severe Combined Immunodeficiency (SCID)	Within Normal Limits	
interpretation		0
Spinal Muscular Atrophy, Real-Time PCR Spinal Muscular Atrophy (SMA) interpretation	Within Normal Limits	8
opinal widecular Atrophy (OWA) interpretation	WILLIAM LIMINS	



NEWBORN SCREENING REPORT

319/384-5097 or 866/890-5965

Patient Name	Birth Date	Client Reference	Accession #
LAST10964566, FIRST10964566	2025-05-01 00:07	10964566	78461

SAMPLE AND ANALYSIS NOTES

- 1. Core conditions screened: Sickle cell disease, Hemoglobin S/C disease, Hemoglobin S beta-thalassemia
- 2. Cystic Fibrosis Transmembrane Conductance Regulator (CFTR) gene variant testing was NOT performed on this sample because the Immunoreactive Trypsinogen (IRT) value was in the normal range.
- 3. Core conditions screened: Fatty acid oxidation defects: Carnitine uptake defect (Carnitine transport defect), Long-chain 3-hydroxy acyl-CoA dehydrogenase deficiency, Medium chain acyl-CoA dehydrogenase deficiency, Trifunctional protein deficiency, Very long-chain acyl-CoA dehydrogenase deficiency; Organic acidemias: Glutaric acidemia type I, 3-Hydroxy 3-methylglutaric aciduria, Isovaleric acidemia, 3-Methylcrotonyl-CoA carboxylase deficiency, Methylmalonic acidemia (methylmalonyl-CoA mutase, cobalamin disorders, vitamin B12 disorders), Beta-Ketothiolase deficiency, Propionic acidemia, Holocarboxylase synthetase deficiency; Amino acidemias: Argininosuccinic aciduria, Citrullinemia type 1, Homocystinuria, Maple Syrup Urine Disease, Classic Phenylketonuria, Tyrosinemia type I. The performance characteristics of this test were determined by the State Hygienic Laboratory. It has not been cleared or approved by the U.S. Food and Drug Administration.
- 4. Core conditions screened: Lysosomal Storage Disorders: Pompe disease, Mucopolysaccharidosis Type 1 (MPS1) disease, Mucopolysaccharidosis Type II (MPSII) disease. A Presumptive Positive interpretation for Pompe, MPS1, or MPSII will automatically reflex to second tier testing if a suitable sample is still available at SHL. A Borderline interpretation for Pompe will automatically reflex to second tier testing if a suitable sample is still available at SHL. If not, program staff will contact with recommendations for second tier testing. A second tier test interpretation will take precedence over the first tier screening interpretation. The performance characteristics of this test were determined by the State Hygienic Laboratory. It has not been cleared or approved by the U.S. Food and Drug Administration.
- 5. The "Pompe Disease 2nd Tier NBS interpretation" will take precedence over the "Pompe disease interpretation". This test was developed and its performance characteristics determined by Mayo Clinic Laboratories, 200 First Street SW, Rochester, MN 55905 (CLIA Certificate: 24D0404292) in a manner consistent with CLIA requirements. This test has not been cleared or approved by the U.S. Food and Drug Administration.
- 6. Core condition screened: X-linked Adrenoleukodystrophy. The performance characteristics of this test were determined by the State Hygienic Laboratory. It has not been cleared or approved by the U.S. Food and Drug Administration.
- 7. The performance characteristics of this test were determined by the State Hygienic Laboratory. It has not been cleared or approved by the U.S. Food and Drug Administration.
- 8. Spinal Muscular Atrophy (SMA) screening uses a deletion in exon 7 of the survival motor neuron 1 (SMN1) gene to assess risk. 95% of SMA cases are caused by deletion of SMN1 exon 7. Thus, the screening assay may miss up to 5% of SMA cases that are not caused by SMN1 exon 7 deletion.

ANALYSIS INFORMATION

<u>TEST</u>	ANALYZED	SITE	RELEASED
Congenital Adrenal Hyperplasia, Immunoassay	2025-05-03 06:00 MBH	10320	2025-05-14 09:41 MBH
2. Congenital Hypothyroidism, Immunoassay	2025-05-03 06:00 MBH	10320	2025-05-14 09:41 MBH
3. Biotinidase Deficiency, Immunoassay	2025-05-03 06:00 MBH	10320	2025-05-14 09:41 MBH
4. Galactosemia, Enzymatic Assay	2025-05-03 06:00 MBH	10320	2025-05-14 09:41 MBH
5. Hemoglobinopathies, Various Methods	2025-05-03 06:00 MBH	10320	2025-05-14 09:41 MBH
6. Cystic Fibrosis, Various Methods	2025-05-03 06:00 MBH	10320	2025-05-14 09:41 MBH
7. Expanded Screening Disorders, Tandem Mass Spectrometry	2025-05-03 06:00 MBH	10320	2025-05-14 09:41 MBH
8. Lysosomal Storage Disorders, Tandem Mass Spectrometry	2025-05-03 06:00 MBH	10320	2025-05-14 09:41 MBH
Pompe Disease 2nd Tier NBS, Flow Injection Analysis-Tandem Mass Spectrometry	2025-05-04 10:30 MBH	10415	2025-05-14 09:41 MBH
10. X-linked Adrenoleukodystrophy, Tandem Mass Spectrometry	2025-05-03 06:00 MBH	10320	2025-05-14 09:41 MBH
11. Severe Combined Immunodeficiency, Real-Time PCR	2025-05-03 06:00 MBH	10320	2025-05-14 09:41 MBH
12. Spinal Muscular Atrophy, Real-Time PCR	2025-05-03 06:00 MBH	10320	2025-05-14 09:41 MBH

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NEWBORN SCREENING REPORT

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Patient Name	Birth Date	Client Reference	Accession #
LAST10964566, FIRST10964566	2025-05-01 00:07	10964566	78461

SITE(S) PERFORMING TESTING

STATE HYGIENIC LABORATORY ANKENY, IOWA LABORATORIES COMPLEX, 2220 S ANKENY BLVD, ANKENY, IA 50023; Phone 515/725-1630; Fax 515/725-1650; Michael A. Pentella, Ph.D., D(ABMM), Director; CLIA ID Number 16D0709302

10415 MAYO CLINIC LABORATORIES, ROCHESTER MAIN CAMPUS, 200 FIRST STREET SW, ROCHESTER, MN 55905; CLIA Certificate: 24D0404292

For questions about resubmission and result interpretation please contact the medical consultants. During regular work hours (8:00 AM to 4:30 PM): 319/384-5097 or toll free in lowa 866/890-5965. 24hr Geneticist on-call number: 319/356-1616. This is a screening test and not indicated for stand-alone purposes; results should be used in conjunction with other available laboratory and clinical information. A false negative or a false positive result must always be considered when screening; therefore, clinical findings and status should be considered whenever interpreting laboratory results. Newborn reference values may not be applicable to older infants, thus screening results should be interpreted with caution in such cases. Information on the conditions screened is available at https://hhs.iowa.gov/programs/programs-and-services/family-health/congenital-inherited-disorders/iowanewborn-screening-program/iowa-newborn-screening-education/blood-spot-screening-information. The result(s) of this report relate only to the items analyzed. Where the laboratory has not been responsible for the sampling stage the results apply only to the sample as received. This report shall not be reproduced except in full without the written approval of the laboratory.

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