

Client: ARUP Example Report Only 500 Chipeta Way Salt Lake City, UT 84108 UNITED STATES

Physician: arup, arup

Patient: Test, MCADPCR 1

DOB

Sex: Male **Patient Identifiers:** 46923 **Visit Number (FIN):** 47261

**Collection Date:** 3/3/2023 11:20

## Medium Chain Acyl-CoA Dehydrogenase (ACADM) 2 Mutations

ARUP test code 0051205

MCADPCR Specimen Whole Blood MCAD Mutation A985G Negative MCAD Mutation T199C Negative

Medium Chain Acyl-CoA Interpretation

See Note

Indication for testing: Carrier screening or diagnostic testing for MCAD deficiency.

Result

A985G: Negative T199C: Negative

This sample is negative for the ACADM variants, c.985A>G and c.199T>C. Persons affected with medium chain acyl-CoA dehydrogenase (MCAD) deficiency may have pathogenic variants not detected by this assay. If the patient has biochemical and/or clinical evidence of MCAD deficiency, plasma acylcarnitine profile testing and/or ACADM gene sequencing is recommended.

This result has been reviewed and approved by

BACKGROUND INFORMATION: Medium Chain Acyl-CoA Dehydrogenase (ACADM) 2 Mutations

CHARACTERISTICS: Limited mitochondrial fatty acid beta-oxidation leading to hypoglycemia, lethargy, seizures hypoketotic aciduria, vomiting, hepatomegaly, hepatic failure, encephalopathy, and sudden death. Manifestations often triggered by prolonged fasting or other metabolic stressors. INCIDENCE: 1 in 15,000 INHERITANCE: Autosomal recessive.

CAUSE: Deleterious ACADM gene mutations. CLINICAL SENSITIVITY: 75 percent for MCAD deficiency.

MUTATIONS TESTED: ACADM mutations c.985A>G (p.K329E, also known as K304E) and c.199T>C (p.Y67H, also known as Y42H).
METHODOLOGY: Polymerase chain reaction (PCR) and fluorescence monitoring.

ANALYTICAL SENSITIVITY AND SPECIFICITY: 99 percent.
LIMITATIONS: Diagnostic errors can occur due to rare sequence
variations. ACADM mutations other than c.985A>G and c.199T>C

will not be detected.

H=High, L=Low, \*=Abnormal, C=Critical

Unless otherwise indicated, testing performed at:



This test was developed and its performance characteristics determined by ARUP Laboratories. It has not been cleared or approved by the US Food and Drug Administration. This test was performed in a CLIA certified laboratory and is intended for clinical purposes.

Counseling and informed consent are recommended for genetic testing. Consent forms are available online

VERIFIED/REPORTED DATES				
Procedure	Accession	Collected	Received	Verified/Reported
MCADPCR Specimen	23-062-105687	3/3/2023 11:20:00 AM	3/3/2023 11:20:17 AM	4/13/2023 1:18:00 PM
MCAD Mutation A985G	23-062-105687	3/3/2023 11:20:00 AM	3/3/2023 11:20:17 AM	4/13/2023 1:18:00 PM
MCAD Mutation T199C	23-062-105687	3/3/2023 11:20:00 AM	3/3/2023 11:20:17 AM	4/13/2023 1:18:00 PM
Medium Chain Acyl-CoA Interpretation	23-062-105687	3/3/2023 11:20:00 AM	3/3/2023 11:20:17 AM	4/13/2023 1:18:00 PM

END OF CHART

H=High, L=Low, \*=Abnormal, C=Critical