ARUP LABORATORIES | aruplab.com

500 Chipeta Way, Salt Lake City, Utah 84108-1221 phone: 801-583-2787, toll free: 800-522-2787

Tracy I. George, MD, Chief Medical Officer

Patient Report

Patient Age/Gender: Female

Specimen Collected: 30-Aug-21 13:14

Hemophilia A (F8) 2 Inversions, | Received: 30-Aug-21 13:14 Report/Verified: 30-Aug-21 13:18

Fetal

Procedure Result Units Reference Interval

Maternal Contamination Mix<5 percent * f1

Study Fetal Spec

Maternal Contam Study, Whole Blood i1

Maternal Spec

F8 INV FE Specimen Cultured Amnio Hemophilia A (F8) Negative f2 i2

Inversions Interp

Result Footnote

f1: Maternal Contamination Study Fetal Spec

Result: Mixed maternal and fetal genotype Number of informative markers: _

Interpretation: Low level maternal contamination (<5 percent) was observed in the fetal sample.

f2: Hemophilia A (F8) Inversions Interp

This result has been reviewed and approved by

Test Information

il: Maternal Contam Study, Maternal Spec

For quality assurance purposes, ARUP Laboratories will confirm the above result at no charge following delivery. Order Confirmation of Fetal Testing and include a copy of the original fetal report (or the mother's name and date of birth) with the test submission. Please contact an ARUP genetic counselor at (800) 242-2787 extension 2141 prior to specimen submission.

i2: Hemophilia A (F8) Inversions Interp

BACKGROUND INFORMATION: Hemophilia A (F8) 2 Inversions

CHARACTERISTICS: Hemophilia A is characterized by deficiency of factor VIII clotting activity. Less than 1 percent factor VIII activity results in severe deficiency associated with spontaneous joint or deep muscle bleeding. Moderate deficiency (1-5 percent activity) and mild deficiency (6-40 percent activity) are associated with prolonged bleeding after tooth extractions, surgery, or injuries, and recurrent or delayed wound healing. Female carriers of hemophilia A may have increased bleeding tendencies.

EPIDEMIOLOGY: 1 in 5,000 live male births worldwide

CAUSE: Pathogenic F8 germline variants

INHERITANCE: X-linked recessive. In the estimated 30 percent of cases that appear to be de novo, the mother is found to be a carrier at least 80 percent of the time. PENETRANCE: 100 percent in males. Approximately 30 percent of female carriers have factor VIII activity levels of less than 40 percent and are at risk for bleeding symptoms typically consistent with mild hemophilia A.

*=Abnormal, #=Corrected, C=Critical, f=Result Footnote, H-High, i-Test Information, L-Low, t-Interpretive Text, @=Performing lab

Unless otherwise indicated, testing performed at:

ARUP Laboratories

500 Chipeta Way, Salt Lake City, UT 84108 Laboratory Director: Tracy I. George, MD

 ARUP Accession:
 21-242-900109

 Report Request ID:
 15048074

 Printed:
 15-Sep-21 10:47

Page 1 of 2

ARUP LABORATORIES | aruplab.com

500 Chipeta Way, Salt Lake City, Utah 84108-1221

phone: 801-583-2787, toll free: 800-522-2787 Tracy I. George, MD, Chief Medical Officer ·

Patient Report

Patient Age/Gender: Female

Test Information

i2: Hemophilia A (F8) Inversions Interp

CLINICAL SENSITIVITY: 51 percent of variants causing severe hemophilia A are detected by F8 inversion testing. This assay does not detect F8 variants associated with mild or moderate hemophilia A in males.

METHODOLOGY: Intron 22-A and intron 1 inversions detected by inverse PCR and electrophoresis.

ANALYTICAL SENSITIVITY/SPECIFICITY: 99 percent

LIMITATIONS: A negative result does not exclude a diagnosis of or carrier status for hemophilia A. Diagnostic errors can occur due to rare sequence variations. F8 variants, other than the F8 type 1 or type 2 intron 22-A and intron 1 inversions, will not be detected. Rare F8 intron 22-A and intron 1 inversions with different breakpoints may not be detected by this assay.

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.

*=Abnormal, #=Corrected, C=Critical, f=Result Footnote, H-High, i-Test Information, L-Low, t-Interpretive Text, @=Performing lab

Unless otherwise indicated, testing performed at:

ARUP Accession: 21-242-900109 **Report Request ID**: 15048074

Printed: 15-Sep-21 10:47

Page 2 of 2