

NEW TEST

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Prolonged Clot Time Reflexive Profile

3006383, CLOT RFLX

Specimen Requirements:	

Patient Preparation: N/A

Collect: At least five light blue (sodium citrate) tubes. Refer to

Specimen Handling at aruplab.com for hemostasis/thrombosis

Effective Date: May 15, 2023

specimen handling guidelines.

Specimen Preparation: Transfer five 1 mL aliquots of platelet-poor plasma to five

ARUP standard transport tubes and label as sodium citrate.

(Min: 1 mL/aliquot and 5 mL total)

Transport Temperature: CRITICAL FROZEN. Separate specimens must be submitted

when additional tests codes are ordered.

Unacceptable Conditions: Anything other than sodium citrated plasma. Specimens

containing anticoagulant medications. Clotted or hemolyzed

specimens.

Remarks: Submit the Patient History form for the Prolonged Clot Time

Reflexive Profile.

Stability: Ambient: Unacceptable; Refrigerated: Unacceptable; Frozen at

-202 weeks; Frozen at -706 months

Methodology: Electromagnetic Mechanical Clot

Detection/Immunoturbidimetry/Microlatex Particle-Mediated

Immunoassay/Platelet Agglutination/Chromogenic Assay

Performed: Sun-Sat

Reported: 2-10 days

Note: Submission of a completed Patient History form with test order

will allow for optimal panel interpretation. The Patient History form for the Prolonged Clot Time Reflexive Profile is available on the ARUP web site or by contacting ARUP Client Services at 800-522-2787. Initial testing will include D-Dimer (0030057), Fibrinogen (0030130), and Lupus Anticoagulant Reflexive Panel (0030181). Depending on these initial findings, a pathologist will order one or more reflexive tests to provide a comprehensive interpretation. Additional testing may include



Factor II, Activity (Prothrombin) (0030007); Factor V, Activity (0030075); Factor VII Activity (0030080), Factor VIII Activity (0030095), Chromogenic Factor VIII, Activity (3002343); Factor VIII Activity with Reflex to Bethesda Quantitative, Factor IX Activity with Reflex to Bethesda Quantitative, Factor IX Activity with Reflex to Bethesda Quantitative, Factor IX (0030032); Factor X, Activity (0030105); Factor XI, Activity (0030110); Factor XII, Activity (0030115); von Willebrand Factor Activity (Ristocetin Cofactor) (0030250); von Willebrand Factor Antigen (0030285); Fibrinogen Antigen (0030135); Inhibitor Assay, PT with Reflex to PT 1:1 Mix (2003260); and Inhibitor Assay, PTT with Reflex to PTT 1:1 Mix, with Reflex to 1-Hour Incubation (2003266). Additional charges apply.

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CPT Codes:

85390-26; additional CPT codes may apply: 85210; 85220; 85230; 85240; 85245; 85246; 85250; 85260; 85270; 85280; 85335; 85379; 85384; 85385; 85525; 85597; 85598; 85610; 85611; 85613; 85635; 85670; 85730; 85732

New York DOH Approval Status:

This test is New York DOH approved.

Interpretive Data:

Refer to report.

Reference Interval:

Refer to individual components.

HOTLINE NOTE: Refer to the Hotline Test Mix for interface build information.