

Overview

Useful For

Detection and quantitation of inhibitor to coagulation factor X

This test is not useful for the detection of a lupus-like circulating anticoagulant inhibitor, a nonspecific circulating anticoagulant, or other inhibitors that are not specific for coagulation factors.

Profile Information

Test Id	Reporting Name	Available Separately	Always Performed
10INT	FX Inhib Profile Tech Interp	No	Yes
F_10	Coag Factor X Assay, P	Yes	Yes

Reflex Tests

Test Id	Reporting Name	Available Separately	Always Performed
10AIH	FX Inhib Profile Prof Interp	No	No
10_IS	Factor X Inhib Scrn	No	No
GBETH	General Factor Bethesda Units, P	No	No

Testing Algorithm

Testing begins with coagulation factor X activity assay with dilutions to evaluate assay inhibition; if the factor X activity assay is normal or increased, then a technical interpretation will be provided.

If the factor X activity assay is decreased, then an inhibitor screen will be performed at an additional charge to look for specific factor X inhibition and a professional interpretation will be provided. If specific inhibition is apparent, the titer of the inhibitor will be determined.

Special Instructions

- [Coagulation Guidelines for Specimen Handling and Processing](#)

Method Name

F_10, 10_IS, GBETH: Optical Clot-Based

10INT: Technical Interpretation

10AIH: Medical Interpretation

NY State Available

Yes

Specimen

Specimen Type

Plasma Na Cit

Ordering Guidance

This test is for factor X inhibitors only. If the presence or type of inhibitor is unknown, first order APROL / Prolonged Clot Time Profile, Plasma, except for patients with known hemophilia A or B. When screening studies are needed for patients with known hemophilia A or B, order 8INHE / Factor VIII Inhibitor Evaluation, Plasma; or 9 INHE / Factor IX Inhibitor Evaluation, Plasma; respectively.

Shipping Instructions

Send all vials in the same shipping container.

Specimen Required

Specimen Type: Platelet-poor plasma

Patient Preparation:

1. Fasting: 8 hours, preferred but not required
2. Patient **must not** be receiving Coumadin (warfarin) or heparin therapy

Collection Container/Tube: Light-blue top (3.2% sodium citrate)

Submission Container/Tube: Polypropylene plastic vials

Specimen Volume: 3 mL in 3 plastic vials, each containing 1 mL

Collection Instructions:

1. Specimen must be collected prior to factor replacement therapy.
2. For complete instructions, see [Coagulation Guidelines for Specimen Handling and Processing](#).
3. Centrifuge, transfer all plasma into a plastic vial, and centrifuge plasma again.
4. Aliquot plasma (1-2 mL per aliquot) into 3 separate plastic vials, leaving 0.25 mL in the bottom of centrifuged vial.
5. Freeze plasma immediately (no longer than 4 hours after collection) at -20 degrees C or, ideally, -40 degrees C or below.

Additional Information:

1. A double-centrifuged specimen is critical for accurate results, as platelet contamination may cause spurious results.
2. Each coagulation assay requested should have its own vial.

Forms

If not ordering electronically, complete, print, and send a [Coagulation Test Request](#) (T753) with the specimen.

Specimen Minimum Volume

2 Plastic vials, each containing 1 mL

Reject Due To

Gross hemolysis	Reject
Gross lipemia	Reject
Gross icterus	Reject

Specimen Stability Information

Specimen Type	Temperature	Time	Special Container
Plasma Na Cit	Frozen	14 days	

Clinical & Interpretive**Clinical Information**

Coagulation factor inhibitors arise in patients who are congenitally deficient in a specific factor in response to factor replacement therapy, or they can either occur spontaneously without known cause or in response to a variety of medical conditions, including the postpartum state, immunologic disorders, certain antibiotic therapies, some malignancies, and in the older population.

Inhibitors of factor VIII coagulant activity are the most commonly occurring of the specific factor inhibitors.

Reference Values

FACTOR X ACTIVITY ASSAY

Adults: 70-150%

Normal, full-term newborn infants or healthy premature infants may have decreased levels (> or =15-20%) that may not reach adult levels for 180 days or more postnatal.*

*See Pediatric Hemostasis References section in [Coagulation Guidelines for Specimen Handling and Processing](#).

FACTOR X INHIBITOR SCREEN:

Negative

GENERAL FACTOR BETHESDA UNITS:

< or =0.5 Bethesda Units

Interpretation

Normally, there is no inhibitor, ie, negative result.

If the screening assays indicate the presence of an inhibitor, it will be quantitated and reported in Bethesda (or equivalent) units.

Cautions

Occasionally, a potent lupus-like anticoagulant may cause false-positive testing for a specific factor inhibitor (eg, factor VIII or IX).

Clinical Reference

1. Hoffman R, Benz Jr EJ, Silberstein LE, et al, eds: Hematology: Basic Principles and Practice. 7th ed. Elsevier; 2018
2. Kasper CK. Treatment of factor VIII inhibitors. Prog Hemost Thromb. 1989;9:57-86

Performance

Method Description

Screening for inhibitors of specific coagulation factors is represented by the inhibitor assay for factor X. This assay consists of measuring the difference in factor X activity (prothrombin time based assay) before and after incubation of a mixture of normal plasma and patient's plasma for 1 hour at 37 degrees C. For optimal sensitivity, the factor X value of the normal plasma is adjusted to approximately 20%, because the factor X assay is more sensitive in this area of the curve. In addition, an excess of patient's plasma will make the test more sensitive to small amounts of inhibitors.(Owen CA Jr, Bowie EJW, Thompson JH Jr. *The Diagnosis of Bleeding Disorders*. 2nd ed. Little, Brown, and Company; 1975:143-145; Cielsa B. Defects of plasma clotting factors. In: *Hematology in Practice*. 3rd ed. FA Davis; 2019:chap 17)

If the inhibitor screen is positive for an inhibitor of factor X, the inhibitor will be quantitated by the "Bethesda assay." In the Bethesda procedure, inhibitors are quantified by mixing equal volumes of serially diluted plasma with normal plasma. This mixture is incubated 2 hours at 37 degrees C, and its factor X activity is measured and compared to a control run at the same time. The difference between the factor X activity of the patient's incubation mixture and that of the control is used to calculate titer. The residual factor X activity is converted to "Bethesda units": 50% residual factor X is equal to 1 Bethesda unit.(Kasper CK, Aldedort LM, Counts RB, et al. A more uniform measurement of factor VIII inhibitors. *Thromb Diath Haemorrh*. 1975;34:869-872; Cielsa B. Defects of plasma clotting factors. In: *Hematology in Practice*. 3rd ed. FA Davis; 2019:chap 17)

PDF Report

No

Day(s) Performed

Monday through Friday

Report Available

1 to 3 days

Specimen Retention Time

7 days

Performing Laboratory Location

Mayo Clinic Laboratories - Rochester Main Campus

Fees & Codes

Fees

- Authorized users can sign in to [Test Prices](#) for detailed fee information.
- Clients without access to Test Prices can contact [Customer Service](#) 24 hours a day, seven days a week.
- Prospective clients should contact their account representative. For assistance, contact [Customer Service](#).

Test Classification

This test has been modified from the manufacturer's instructions. Its performance characteristics were determined by Mayo Clinic in a manner consistent with CLIA requirements. This test has not been cleared or approved by the US Food and Drug Administration.

CPT Code Information

85390
85260
85335 (if appropriate)
85335 (if appropriate)
85390 (if appropriate)

LOINC® Information

Test ID	Test Order Name	Order LOINC® Value
10INE	Factor X Inhib Profile, P	96457-7

Result ID	Test Result Name	Result LOINC® Value
F_10	Coag Factor X Assay, P	3218-5
10INT	FX Inhib Profile Tech Interp	69049-5