

**Reporting Title:** Factor II Inhib Profile, P

**Performing Location:** Rochester

**Ordering Guidance:**

This test is for factor II 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 IX Inhibitor Evaluation, Plasma; or 9 INHE / Factor IX Inhibitor Evaluation, Plasma; respectively.

**Shipping Instructions:**

Send all vials in the same shipping container.

**Necessary Information:**

If priority specimen, mark request form, give reason, and request a call-back.

**Specimen Requirements:**

**Specimen Type:** Platelet-poor plasma

**Patient Preparation:**

- 1. Patient must not be receiving Coumadin (warfarin) or heparin therapy
- 2. Fasting preferred

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

**Submission Container/Tube:** 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. 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 Type	Temperature	Time	Special Container
Plasma Na Cit	Frozen	14 days	

**Result Codes:**

Result ID	Reporting Name	Type	Unit	LOINC®
F_2	Coag Factor II Assay, P	Numeric	%	3289-6
2INHT	FII Inhib Profile Tech Interp	Alphanumeric		69049-5

LOINC® and CPT codes are provided by the performing laboratory.

Supplemental Report:

No

Components:

Test Id	Reporting Name	CPT Units	CPT Code	Always Performed	Available Separately
2INHT	FII Inhib Profile Tech Interp	1	85390	Yes	No
F_2	Coag Factor II Assay, P	1	85210	Yes	Yes

CPT Code Information:

- 85390
- 85210
- 85335 (if appropriate)
- 85335 (if appropriate)
- 85390 (if appropriate)

Reflex Tests:

Test Id	Reporting Name	CPT Units	CPT Code	Always Performed	Available Separately
2AINH	FII Inhib Profile Prof Interp	1	85390	No	No
F2_IS	Factor II Inhib Scrn	1	85335	No	No
GBETH	General Factor Bethesda Units, P	1	85335	No	No

Result Codes for Reflex Tests:

Test ID	Result ID	Reporting Name	Type	Unit	LOINC®
F2_IS	7806	Factor II Inhib Scrn	Alphanumeric		96454-4
2AINH	607489	Reviewed by	Alphanumeric		18771-6
2AINH	607445	FII Inhib Profile Prof Interp	Alphanumeric		69049-5
GBETH	607434	General Factor Bethesda Units, P	Numeric	BU	13591-3

Reference Values:

FACTOR II ACTIVITY ASSAY

Adults: 75-145%

Normal, full-term newborn infants or healthy premature infants may have decreased levels (> or =25%) that may remain below adult levels for 180 days or more postnatal.\*

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

FACTOR II INHIBITOR SCREEN:

Negative

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GENERAL FACTOR BETHESDA UNITS:  
< or =0.5 Bethesda Units