

Coagulation Control A - English

INTENDED USE

For use as accuracy control of screening tests (PT/aPTT/TT/Fibrinogen) and determination of various coagulation factors in the interface between normal and abnormal coagulation activity.

Precision control, i.e. the recording of the day to day variation, is an important procedure in coagulation testing; checking the accuracy of the test system is particularly important.

Coagulation Control A can be used for:

Accuracy control of all screening tests, i.e. prothrombin time (PT), partial thromboplastin time (aPTT), thrombin time (TT), fibringen determination (Clauss), and of various coagulation factors in the interface between normal and abnormal coagulation activity.

· Precision control of all parameters listed in the table.

SUMMARY

Coagulation Control A is a lyophilized abnormal human citrated plasma, where the clotting factors are reduced, Coagulation Control A contains stabilizers but no bactericide additives

REAGENTS

The REF 5021055 Coagulation Control A contains:

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	Reagent / Content	Description			
5 x 1 mL	Coagulation Control A	Human plasma, lyophilized			

The REF 5021060 Coagulation Control A contains:

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	Reagent / Content	Description
50 x 1 mL	Coagulation Control A	Human plasma, lyophilized

Material required (not supplied with the kit)

Distilled water

Precision pipette Laboratory timer

Warning and precautions

- RUO for research use only .
- This kit is intended for use by personnel trained in laboratory procedures and universal precautions for the use of chemicals and potentially biohazardous substances. All human blood or plasma products as well as samples must be considered as potentially infectious. They have to be handled with appropriate care and in strict observance of safety regulations. The rules pertaining to disposal
- are the same as applied to disposing hospital waste. Calibrators and control plasmas are made from human blood and any individual plasma involved in the procedure are tested HbsAg, HIV 1/2 Ab and HCV-Ab-negative. However, all human blood products should be handled as potentially infectious material. Get a Material Safety Data Sheet for this product from www.technoclone.com.

Stability and storage

The expiry date printed on the labels is only applicable to storage of the unopened containers at +2...8 $^{\circ}\text{C}.$

Reagent	1825 °C	28 °C	< -20 °C
Coagulation Control A	4 hours	8 hours	1 month

The vials can be only frozen once.

Upon storage, caps should be screwed tightly.

Use the reconstituted plasma in single factor determinations within 2 hours, stored at room temperature.

TEST PROCEDURE

Preparation of Coagulation Control A

Before starting the test, all the required components are to be brought to room temperature.

When reconstituting plasmas, mixing reagents or buffers avoid foaming.

 Coagulation Control A: Reconstitute each control vial with 1.0 mL of distilled water. Allow the reconstituted material
to stand at room temperature for 10 minutes followed by swirling of the vial before use. For standardization a reconstitution time of 30 minutes is recommended

Treat the reconstituted plasma as a citrated sample according to the instructions of the respective test.

LIMITATION OF THE TEST

The figures in the table are only applicable to the indicated lot number of Coagulation Control A and the method has to

adhere strictly as prescribed for the respective reagent. The guidelines for evaluation given in this leaflet have been established assuming that a constant sensitivity of different batches is guaranteed by the manufactures of each reagent.

INTERPRETATION OF RESULTS

The figures in the table are only applicable to the indicated lot number of Coagulation Control A and the method has to adhere strictly as prescribed for the respective reagent.

STANDARDISATION

The reported values were determined over multiple runs using a specific lot of reagent and against an Internal Reference Standard which is traceable to the current International Standards, identified in the acceptance range table.

For the tests where International Standards are not available, these parameters have been assigned against an Internal Reference Standard which is traceable to a frozen normal plasma pool of 100 donors.

LITERATURE

Please contact Technoclone (www.technoclone.com) or your local distributor.

EDITORIAL NOTE

This document is available in several languages. The translations have been done using the master document in English. In the event of doubts or discrepancies, the wording in the master document in English shall take precedence.





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