Timed fibrin digestion: A simplified technique for the measurement of the fibrinolytic activity of the blood

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SYNOPSIS When the rate of fibrinolysis in vitro was measured by an isotopic technique, it was found that the amount of fibrin digested after incubation for three hours correlated closely with the overall rate of the digestion process. It is suggested that, using an isotopic technique, estimation of fibrin digestion after three hours’ incubation provides a useful method for measuring the fibrinolytic activity of blood. A recommended technique is described and its advantages are discussed briefly.

By employing 125I-labelled fibrinogen in vivo, the fibrinolytic activity of blood can be measured in vitro by observing the rate of release of radioactivity from blood samples diluted in buffer and clotted with thrombin (Hickman, 1971). A description of the technique is given in Figure 1. Studies which are given below show that the amount of fibrin digested after incubation for three hours correlates closely with the overall rate of the reaction. A simplified technique is therefore described for the estimation of the three-hour fibrin digestion, and the experimental justification for its application is given.

Materials and Methods

The isotopic method, the method for estimation of plasma fibrinogen concentration, and the reagents used were as described previously (Hickman, 1971), except that the measurement of fibrin digestion at three hours was performed in triplicate.

Ninety-two studies were included in the analysis. For each study, the rate of the reaction, ie, tan θ or 'slope'(see Fig.1) and the amount of fibrin digested in three hours (hereinafter referred to as the three-hour value) were calculated. These measurements were carried out on some 100 blood samples, obtained largely as part of another study, from patients in the postoperative period. In a few samples total lysis had occurred at three hours: these were excluded from the study leaving 92 samples for analysis.

Results

In the 92 studies both the three-hour value and the reaction velocity (slope) were measured. Figure 2 shows the correlation obtained (r = 0.9116). Samples in which total lysis of fibrin had occurred by three hours were excluded since the three-hour measurement would underestimate the real activity. The mean percentage error from the mean in the 92 triplicate readings was 10.6%.

Discussion

Difficulties inherent in the use of lysis time methods when the plasma fibrinogen concentration undergoes alteration have already been discussed (Hickman, 1971). The isotopic method attempts to overcome these difficulties but as put forward originally is time consuming and demanding technically. Furthermore, unless the individual study of fibrin digestion is characterized sufficiently, involving the preparation of numerous aliquots, there is difficulty in determining accurately the rate of the reaction. Thus the assessment of activity by measuring fibrin digestion per unit time is an attractive alternative with advantages both on grounds of accuracy and of technical simplicity.

This study has shown that the three-hour measurement correlates satisfactorily with the overall rate in 92 studies where total lysis of fibrin occurred in not less than three hours. In those samples in which total lysis of fibrin occurs within the three-hour period, the measurements will reflect the total

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amount of fibrin present and tend to underestimate the real fibrinolytic activity of the sample. However, excluding samples of abnormally low fibrin content, total lysis within this period occurs only in samples possessing considerable activity. Nonetheless it is felt that the widest application of the technique lies in the accurate identification of reduced fibrinolytic activity. In view of current interest in the possible association of defective activity with vascular and thrombotic disease it is suggested that there is considerable scope for the useful application of this method. A recommended technique for the measurement of the three-hour value is given in the Appendix.

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Reference


**Appendix**

**Recommended Method for Estimation of Three-hour Fibrin Digestion**

Ten ml of blood is taken by venepuncture using minimal haemostasis. Immediately 4·0 ml is transferred to a conical flask containing 34 ml ice-cold acetate citrate diluent, pH 7·4, and surrounded by melting ice. The remaining 6 ml is transferred to a plastic test tube (capacity 10 ml) containing 40 mg dry EDTA (Stayne Laboratories), and this sample is used subsequently for measuring the plasma fibrinogen concentration and the specific activity of fibrinogen in this sample.
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the $^{125}$I-labelled fibrinogen preparation. From the conical flask containing the blood diluent mixture, five 4-75 ml aliquots are transferred by pipette into chilled glass test tubes of internal diameter 15 mm, containing 0-25 ml of freshly prepared thrombin solution strength 50 NIHu/ml (Leo Pharmaceuticals Ltd, Denmark). To two of these aliquots (hereafter referred to as the inhibited samples) is added 0·1 ml IM EACA solution (13·12 g EACA dissolved in 100 ml of the acetate citrate diluent). Each sample is mixed carefully and placed in a rack in a constant temperature waterbath at 37°C. After about 15 minutes each tube is rotated between the palms of the hands to detach the clot from the side of the test tube and allow clot retraction to proceed. Thereafter the tubes are untouched until completion of the test. After three hours' incubation the samples are removed from the waterbath and in the three non-inhibited ('test') samples further fibrinolysis is prevented by the addition to each tube of 0·1 ml IM EACA solution. The five test tubes are then placed in a centrifuge at 500 g for 10 minutes. After centrifugation the clear supernatant is removed with a Pasteur pipette and filtered by passage through a Pasteur pipette containing a tightly packed plug of cotton wool. Of this filtered supernatant, 3·0 ml is placed in a plastic counting tube, capacity 10 ml, and the radioactivity counted in a well-type scintillation counter with a background for $^{125}$Iodine between 1·0 and 2·0 counts/s.

Calculations
(1) Radioactivity released by fibrin digestion
(a) Per 3 ml aliquot
(Mean count non-inhibited samples − mean count inhibited samples)
(b) Per 0·5 ml whole blood
(Mean count non-inhibited samples − mean count inhibited samples) × $\frac{5}{3}$

(2) Fibrin digested at three hours (mg/0·5 ml blood)
(Mean count non-inhibited samples − mean count inhibited samples) × $\frac{5}{3}$

Specific activity of fibrinogen
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