Detection of fibrinogen antigens with two latex techniques applied to urine concentrates

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Synopsis Fibrinogen antigens were measured either with an agglutination inhibition method (using latex particles coated with fibrinogen; Diagen test) or with a direct agglutination technique (using latex particles coated with a mixture of anti-D and anti-E antibodies; Thrombo-Wellcotest). Both methods were compared with the tanned red cell haemagglutination inhibition immunoassay (TRCHII) during progressive degradation of fibrinogen with plasmin and using purified fibrinogen fragments or urine concentrates from chronic glomerulonephritis or transplanted patients. Due to the different sensitivity of the two latex techniques to fibrinogen and its plasmin derivatives, their combined use may be helpful to distinguish the nature of the fibrinogen-like material excreted in urine.

Many studies have recently indicated that urinary excretion of fibrinogen degradation products is a valuable parameter of disease activity in some renal disorders (Clarkson, MacDonald, Petrie, Cash, and Robson, 1971; Dotremont, Vermuyen, Donati, Van Damme, and Michielsen, 1972), may be helpful in evaluating the response of glomerulonephritis patients to treatment (Vermuyen, Dotremont, de Gaetano, Donati, and Michielsen, 1970; Clarkson, MacDonald, Cash, and Robson, 1972), and represents a diagnostic tool during rejection crises following kidney homotransplantation (Braun and Merrill, 1968; Antoine, Neveu, and Ward, 1969; Clarkson, Morton, and Cash, 1970; Bouma, Hedner, and Nilsson, 1971; Haanen, Novakova, Wijdeveld, and van Liebergen, 1971). The development of simple, rapid and sensitive methods for detecting fibrinogen derivatives in urine is therefore of increasing interest.

In this paper two techniques using latex particles will be compared with the tanned red cell haemagglutination inhibition immunoassay (TRCHII) of Merskey, Lalezari, and Johnson (1969) on purified fibrinogen derivatives and on urine concentrates from chronic glomerulonephritis and transplanted patients.

Materials and Methods

Purified human fibrinogen was a gift of Kabi (Stockholm). Solutions of 1% (w/v) clottable protein were prepared in 0.1M phosphate buffer pH 7.4 and incubated at 37°C with human plasmin (Claeys, Molla, and Verstraete, 1973), at a final concentration of 0.5 CTA units/ml. At different time intervals aprotinin (Trasylol, Bayer, Leverkusen) was added at a final concentration of 500 KI units/ml.

Purified D and E fragments from human fibrinogen were obtained by DEAE-cellulose chromatography of an extensively degraded fibrinogen-late (Nussenzweig, Seligmann, Pelmont, and Grabar, 1961; Donati, Molla, and Vermuyen, 1971). Urine concentrates were prepared as previously described (Vermuyen et al, 1970).

The TRCHII, the staphylococcal clumping test (SCT) (Hawiger, Niewiarowski, Gurewich, and Thomas, 1970; Donati, Vermuyen, and Verstraete, 1971), and the measurement of the clottable protein on urine concentrates were performed as described elsewhere (Vermuyen et al, 1970; Donati, Molla, Michielsen, and Vermuyen, 1973). Clottable protein was considered to be present in a sample when the titre in the TRCHII decreased at least eight-fold after incubation with thrombin and proteolysis inhibitor.

The Thrombo-Wellcotest reagent was supplied by Wellcome Reagents Limited (Beckenham and Brussels) as a suspension of latex particles coated with rabbit antibody against D and E fragments.
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Test samples were diluted in a 0·1M glycine/saline buffer pH 8·2 and the test was performed as suggested by the manufacturer, except that neither thrombin nor proteolysis inhibitor were added to the test samples.

The Diagen reagents were a gift of Diagnostic Reagents Limited through the courtesy of Dr K. W. E. Denson.

The principle and the performance of this test have been described by Allington (1971a and b). The agglutination inhibition titre was transformed into concentrations of fibrinogen-like material (μg/ml) by comparison with a simultaneously studied standard normal plasma of known fibrinogen concentration.

Results

Reactivity of purified materials in LATEX TECHNIQUES

The reactivity of purified fibrinogen during plasmin digestion in both latex techniques was compared to that in the TRCHII and the SCT. The figure shows the evolution of the titre in the different tests before and during the proteolysis. With the high plasmin concentration used, the reactivity in the SCT decreased very early (after one hour) whereas the titre in the three other tests remained unchanged during at least eight hours; at the stage of very extensive degradation the reactivity decreased in the TRCHII and the Diagen tests, but increased in the Thrombo-Wellcotest. These results were confirmed by the study of purified D and E products: as shown in Table I, the Diagen test, like the TRCHII, was highly sensitive to fibrinogen, but reacted four to eight times less with extensively proteolysed D and not at all with fragment E. In contrast, the Thrombo-Wellcotest was four and 60 times more sensitive to D and E respectively than to fibrinogen.

Reactivity of urinary fibrinogen-like material in LATEX TECHNIQUES

The reactivity of urinary fibrinogen-related material in the two latex techniques was studied on 67 urine concentrates, 42 from 14 transplanted patients and 25 from 14 chronic glomerulonephritis patients. Table II shows the regression lines obtained when the results of the Diagen test, of the Thrombo-Wellcotest, and of the TRCHII were compared. In view of the different sensitivity of the latex techniques to fibrinogen and its derivatives, shown by the studies on purified systems, the reactivity in these tests of clottable and non-clottable samples was also considered separately. A very close correlation was found between the Diagen test and the TRCHII both when the total group and clottable or non-clottable samples were evaluated (the coefficient being only slightly lower in the correlation calculated for non-clottable samples). In contrast, the correlation coefficients and the regression lines calculated for either the TRCHII or the Diagen and the Wellcotest differed markedly depending on whether the urine concentrates contained clottable fibrinogen derivatives or not. Indeed, the Thrombo-Wellcotest reacted rather poorly when the samples contained clottable fibrin(ogen)-related material.

Discussion

This study offers an evaluation of two simple and rapid techniques, using latex particles, for the measurement of fibrin(ogen)-related material. Their

<table>
<thead>
<tr>
<th>Material Studied</th>
<th>TRCHII</th>
<th>SCT</th>
<th>Wellcotest</th>
<th>Diagen</th>
</tr>
</thead>
<tbody>
<tr>
<td>Human purified fibrinogen (Kabi) (100 μg/ml)</td>
<td>32</td>
<td>80</td>
<td>10</td>
<td>16</td>
</tr>
<tr>
<td>Fragment D (100 μg/ml)</td>
<td>2</td>
<td>&lt;1</td>
<td>40</td>
<td>4</td>
</tr>
<tr>
<td>Fragment E (100 μg/ml)</td>
<td>&lt;1</td>
<td>&lt;1</td>
<td>600</td>
<td>&lt;2</td>
</tr>
</tbody>
</table>

Table I Reactivity of purified fibrinogen derivatives in different tests

The sensitivity of the different tests was expressed as the reciprocal of the highest dilution of test material still reacting in the test. Protein concentration was determined by ultraviolet absorption at 280 μν using values of 1 cm

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Fig Evolution of the titre in different tests during proteolysis of human fibrinogen by plasmin.
reactivity has been compared with that of the TRCHII, and of the SCT, already two widely used techniques.

Experiments with purified materials have shown that the two latex tests differ markedly in sensitivity to fibrinogen and to its derivatives at different stages of degradation. The Diagen test, like the TRCHII, would measure fibrinogen antigen $d^3$ and therefore, as the TRCHII, be fairly insensitive to stage four digest (Molla, Donati, and Vermilyen, 1973); indeed, like the TRCHII, it did not react with E and poorly with extensively degraded D. In contrast, the Wellcotest had a very high sensitivity to later digests, a finding also reported by Hoq and Cash (1973). The different reactivity of the two latex tests can be explained by differences of the antisera involved: in the Diagen test and in the TRCHII an antiserum directed against whole fibrinogen, in the Thrombo-Wellcotest a mixture of anti-D and anti-E antibodies is used. The increased reactivity of the extensively degraded fibrinogen fragments, compared to unaltered fibrinogen, in the Thrombo-Wellcotest suggests that part of the anti-D and anti-E antibodies is directed against neo-antigens exposed during cleavage of fibrinogen. The existence of such neo-antigens has recently been shown (Plow and Edgington, 1973).

The different characteristics of the two latex techniques appeared also from the study of urine concentrates: the Diagen was highly correlated with the TRCHII whatever the nature of the fibrinogen derivatives present in urine, whereas the Thrombo-Wellcotest was much better correlated with the TRCHII or the Diagen, when only non-clottable fibrinogen-related material was measured; this observation reemphasizes the relatively poor reactivity of unaltered fibrinogen in the Thrombo-Wellcotest. A very satisfactory correlation between the Thrombo-Wellcotest and the TRCHII has been found by other workers in serum (Arocha-Pinango, 1972; Hoq and Cash, 1973) and in urine concentrates (Pitcher, 1972; Hoq and Cash, 1973). The latter authors have performed the test on urine previously treated with thrombin, therefore on samples containing only non-clottable fibrinogen derivatives. Our results thus are in agreement with their findings.

As urine both from glomerulonephritis and from transplanted patients can contain clottable protein (Yatzidis, 1964; Clarkson et al., 1972; Donati et al., 1973), the combined use of the two latex techniques on non-thrombin treated urine concentrates may prove helpful in distinguishing easily the type of fibrinogen-related material excreted in urine.

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References


Table II Regression lines obtained by comparing the results with the Diagen test, Thrombo-Wellcotest, and TRCHII on 67 urine concentrates containing clottable or non-clottable protein

<table>
<thead>
<tr>
<th>Material Studied</th>
<th>No.</th>
<th>x</th>
<th>y</th>
<th>Regression Line</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total group</td>
<td>67</td>
<td>x</td>
<td>y</td>
<td>$\log y = -0.222 + 1.065 \log x$</td>
</tr>
<tr>
<td></td>
<td>67</td>
<td>y</td>
<td>$\log y = 0.1060 + 0.8254 \log x$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>67</td>
<td>$\log y = 0.304 + 0.762 \log x$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-clottable</td>
<td>42</td>
<td>x</td>
<td>y</td>
<td>$\log y = -0.230 + 0.053 \log x$</td>
</tr>
<tr>
<td></td>
<td>42</td>
<td>y</td>
<td>$\log y = -0.158 + 0.986 \log x$</td>
<td></td>
</tr>
<tr>
<td>Clottable</td>
<td>25</td>
<td>x</td>
<td>y</td>
<td>$\log y = 0.202 + 0.938 \log x$</td>
</tr>
<tr>
<td></td>
<td>25</td>
<td>y</td>
<td>$\log y = 0.9504 + 0.539 \log x$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>25</td>
<td>$\log y = 0.967 + 0.523 \log x$</td>
<td></td>
<td></td>
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</tbody>
</table>
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