Artificial factor VIII deficient plasma: preparation using monoclonal antibodies and its use in one stage coagulation assays

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SUMMARY Monoclonal antibodies to factor VIII antigen (VIII:Ag) and von Willebrand factor (vWF:Ag) were immobilised on Sephacryl S-1000 and tested for their ability to deplete normal human citrated plasma of factor VIII. A combination of two antibodies to VIII:Ag and one antibody to vWF:Ag was required to produce plasma containing less than 0.01 IU/ml. Its performance in the one stage clotting assay of VIII:C was equivalent to that of congenital VIII deficient plasma for the assay of normal and haemophilic plasma and factor VIII concentrates. Storage of freeze dried aliquots of this product at -20°C, +4°C, and 37°C showed that it could be used as a substrate for at least six months when stored at temperatures +4°C and below.

The assay of factor VIII procoagulant activity (VIII:C) is important both in the clinical assessment of patients with haemophilia A and in the production of therapeutic factor VIII concentrates. While VIII:C may be assayed by the two stage clotting,1 or chromogenic assay,2 most laboratories use the one stage clotting assay4 which requires severe haemophilic plasma as a reagent. This has largely been obtained from patients with a congenital deficiency, but the increasing introduction of treatment for use at home has made its supply more uncertain; and concern over viral infectivity of these haemophilic plasmas makes the preparation of artificial haemophilic plasma an attractive proposition.

Artificial haemophilic plasma prepared by mixing partially purified fractions5 has not gained widespread acceptance. Normal plasma has been depleted of VIII:C using polyclonal rabbit antibodies,7 human antibodies,8 and rabbit and human antibodies combined,9 while cryosupernatant has been depleted with rabbit antibodies.10 Unfortunately, these methods are limited by the availability of suitable human or rabbit inhibitors.

In this study murine monoclonal antibodies to factor VIII antigen (VIII:Ag) and von Willebrand factor (vWF:Ag) were used to deplete normal human plasma of VIII:C and the performance of this reagent in the one stage clotting assay was assessed.

Material and methods

Sephacryl S-1000 was obtained from Pharmacia (UK) Ltd. Human plasma was obtained from normal blood donations and was both human immunodeficiency virus (HIV) antibody and hepatitis B antigen negative. Calibrated normal pooled plasma was used as a standard for all assays. Its VIII:C content was 0.9 IU/ml when assayed against the International Plasma Standard for factor VIII (provided by the National Institute for Biological Standards and Control).

Monoclonal antibodies to VIII:Ag, ESH 1-10, were prepared and purified on protein A-Sepharose 4B by Griffin et al.11 Purified monoclonal antibodies to vWF:Ag, EsvWF 8, 9, and 10, were prepared in the same manner as EsvWF 1 to 5.12

Human IgG to VIII:Ag (80 BU/ml plasma) and rabbit IgG to vWF:Ag were isolated from the corresponding antisera (after recalcification of plasma when necessary) by sodium sulphate precipitation.13 The rabbit antiserum was prepared locally.12

IgG was immobilised at 1 to 10 mg/ml settled gel on Sephacryl S-1000 by the low temperature triethylamine-cyanogen bromide method.14 The Sephacryl S-1000 gel was autoclaved at 121°C for 30 minutes, before activation to enhance the subsequent reactivity with CNBr. This improved the activation yield from 5 to 25 μmol cyanate/ml of gel. The gels were washed and heated for 30 minutes at 56°C after coupling of immunoglobulin.

One stage clotting assay of VIII:C, two-site immunoradiometric assay of VIII:Ag, the electroimmunoassay of vWF:Ag, and the radioimmunoassay of vWF:Ag have been described previously.12-15 The microtray chromogenic assay of VIII:C used was that adapted by Prowse et al.3

One stage clotting assays for other factors16 and
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fibrinogen were performed according to established methods.

Endotoxin concentrations were measured by the Limulus assay (Associates of Cape Cod Inc, Massachusetts) and bacterial contamination was measured by adding 1 ml sample to 15 ml tryptone soya agar in a sterile Petri dish at 22°C for 48 hours and counting the colonies.

LYOPHILISATION

Sucrose (2% w/v) was added to the immunodepleted plasma which was freeze dried in 2 ml aliquots and stored under nitrogen at −20°C, +4°C, and 37°C. A donation of congenitally factor VIII deficient plasma was treated similarly for comparison.

BATCH ADSORPTIONS USING MONOCLONAL ANTIBODIES TO VIII:Ag AND vWF:Ag

To assess whether normal plasma could be depleted of VIII using a single monoclonal antibody, batch immunoabsorption of 1 ml aliquots of normal plasma was performed with 0.1 ml of Sephacryl S-1000 immobilised monoclonal antibody to VIII:Ag (1 mg IgG/ml gel) for four hours at room temperature. Ten monoclonal antibodies were tested and the plasma samples were assayed for residual VIII activities. ESvWF 8, 9, and 10 (monoclonal antibodies to vWF:Ag) were immobilised and tested in a similar way.

COLUMN ADSORPTION

Initial column adsorption experiments involved passing 20 ml of plasma down 1 ml columns of immobilised ESvWF 8 and ESH 8 in separate columns, run in series. ESH 5 was subsequently added to the series and this combination was then tested on a larger scale using 17 mg of each of the three antibodies as follows. Each monoclonal antibody was coupled to Sephacryl S-1000 at 1 mg/ml settled gel and plasma was passed through three serially connected columns (1.5 × 15 cm) containing these gels at a rate of 50 ml/hour—that is 1 column volume per hour. In addition, a precolumn of 10 ml bolland Sephacryl S-1000 was included as a clarifying filter. Columns were pre-equilibrated in citrate-saline (15 mM citrate, 150 mM sodium chloride pH 7.0, 3 mM sodium azide) at room temperature, regeneranted after passage of plasma with 2 M KI, 20 mM imidazole (pH 6.5), and then re-equilibrated and stored in citrate-saline at 4°C.

Fibrinogen concentrations were measured at 10 ml intervals in the initial stages of this work.

TESTS ON DEPLETED AND CONGENITALLY DEFICIENT PLASMA

Normal plasma (200 ml) was processed in one day and the product tested for VIII:C content and for use as a substrate in the one stage assay for VIII:C. Six such products were mixed together and then freeze dried as described earlier. The phials were then divided into three lots and stored at −20°C, +4°C, and +37°C. Each month aliquots were reconstituted and assayed for various coagulation factors and tested for use as a substrate in the one stage assay for VIII:C up to a period of six months.

A donation of congenitally FVIII deficient plasma was freeze dried and treated in the same way for comparison. vWF:Ag was measured on each of the six products that were to be pooled.

Results

BATCH ADSORPTION

Results were compared with the normal pooled plasma standard (100%). Assay of the residual FVIII in the plasma showed that a maximum of 88% VIII:Ag and 68% VIII:C was adsorbed using ESH 8. Only ESH 5 removed a greater percentage of VIII:C than VIII:Ag from plasma. Mixing together of ESH 2, 4, 5 and 8 did not increase these values.

ESvWF 8 and 10 each removed 50% VIII:C and 30% VIII:Ag and more than 90% of the plasma vWF:Ag. ESvWF 9 removed less VIII. Similar experiments using 50 μg of ESvWF 8 combined with 50 μg of monoclonal antibody to VIII:Ag gave increased adsorption with a maximum 96% VIII:Ag and 86% VIII:C being adsorbed when using ESH 8. The main results are given in fig 1.

COLUMN ADSORPTION

Use of column rather than batch adsorption yielded much better results and ESvWF 8 alone removed 98% of VIII:C. ESvWF 8 and ESH 8 together, however, still
gave residual VIII:C values that were too high (> 1%) to allow the product to be used as a substrate in the one stage coagulation assay for VIII:C. This residual activity was shown to be due to true VIII:C as it could be inhibited by human IgG to VIII:C. To show this, 50 µl of the adsorbed plasma was incubated with 50 µl of human anti-VIII:C for two hours at room temperature. At the end of this time, the plasma was assayed for VIII:C activity by the chromogenic assay and this was shown to be less than 1%. Control experiments, in which IgG was replaced by buffer, showed residual VIII activity.

As ESH 5 was the only antibody to VIII:Ag to adsorb a greater percentage of VIII:C than VIII:Ag, it was decided to use this together with ESvWF 8 and ESH 8. Column adsorption was tried with this combination of ESvWF 8, ESH 8, and ESH 5 monoclonal antibody columns and resulted in greater than 99% adsorption of VIII:C (table 1). When plasma was passed down the columns at a rate of 3 column volumes/hour the adsorption was inadequate.

**REGENERATION**

After plasma depletion elution of columns with 2 M KI and 20 mM imidazole pH 6.5 yielded 50% of the applied VIII:Ag and vWF:Ag from the gels. Other eluents were tested including 3 M KSCN, 5 M urea, and 0.2 M glycine (pH 3.0), but no better results were obtained. The gels were used and regenerated 10 times and still produced a product containing less than 1% of VIII:C. Between experiments the gels were stored at 4°C in citrate-saline buffer containing azide.

**TESTS ON DEPLETED AND CONGENITALLY DEFICIENT PLASMAS**

The six products, which were pooled and freeze dried for the stability study, were each shown to contain less than 1% VIII:C and VIII:Ag with vWF:Ag, ranging from <1% to 3% of normal pooled plasma. The mean (SD) for other clotting factors was: FII 61 (8); FV 70 (11); FIX 68 (6); FX 93 (22); and FXI 40 (14) (values given as per cent of normal pooled plasma). Fibrinogen concentrations were 2.7 (1) g/l.

Table 2 gives results for the pooled immunodepleted plasma and for the congenital deficient plasma after freeze drying.

<table>
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<tr>
<th>Factor</th>
<th>After freeze drying</th>
<th>6 months at -20°C</th>
<th>6 months at +4°C</th>
<th>2 months at +37°C</th>
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<td></td>
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<td>XI</td>
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Results compared with normal pooled plasma = 100%, except fibrinogen (g/l)
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Fig 3 Assays on haemophilic plasmas, normal plasmas, and VIII concentrates using immunodepleted plasma compared with identical samples assayed using congenital substrate.

(\(\Delta\) = concentrates plotted on 1/5 scale) \(n = 51, r = 0.99\)

\(p < 0.001\) with a regression equation of immunodepleted

\[100 \text{ congenital} - 0.005 (\text{IU/ml})\].

Fig 2 shows standard curves obtained in the one stage assay when using congenitally deficient plasma, deficient plasma produced as described above containing less than 1% VIII:C, plasma adsorbed with ESvWF 8 (containing 2% VIII:C), and plasma adsorbed by using ESH 8 together with ESH 5 (containing 4% VIII:C).

When newly prepared columns were used for the first time \(\approx 50\%\) of fibrinogen was removed from the first 10–20 ml of plasma. The first 30 ml of plasma from such runs was therefore rejected. After regeneration of the column subsequent runs yielded plasma with fibrinogen concentrations that were always within the normal range.

Stability of freeze dried product and congenitally deficient plasma

The freeze dried samples were stored at \(-20^\circ\text{C}, +4^\circ\text{C}\), and \(+37^\circ\text{C}\) and aliquots were reconstituted and tested at monthly intervals for six months. After six months of storage at \(-20^\circ\text{C}\) and \(+4^\circ\text{C}\), the immunodepleted plasma had a concentration of 40% of factor XI compared with a value of 65% for the freeze dried congenitally deficient plasma.

Table 2 shows the residual concentrations of coagulation factors at six months (two months for plasma stored at \(+37^\circ\text{C}\)).

Comparison with congenitally deficient plasma for one stage assays

Fifty one samples, including 29 normal plasmas, eight haemophilia A plasmas (before and after infusion of VIII concentrates), and 14 VIII concentrates, were assayed for VIII:C using both the immunodepleted plasma and congenitally deficient plasma. These showed excellent correlation \(r = 0.99\) (fig 3). Regression analysis of these data showed that the potencies could be fitted to the equation: immunodepleted

\[100 \text{ congenital} - 0.005 (\text{IU/ml})\].

No drift was found in clotting times during the assay period (up to 60 minutes) when using the immunodepleted plasma.

Bacterial testing

Bacteria have deleterious effects on plasma products. Checks were therefore carried out on normal and depleted plasma to check if column passage resulted in contamination. The column had been used and regenerated five times before this test. Passage down the antibody column resulted in a minor increase in endotoxin concentrations from undetectable to 2 ng/ml and in bacterial contamination from 5 to 30 colony forming units/ml.

Assays on plasma from patients with von Willebrand’s disease

Four patients with von Willebrand’s disease had their VIII:C concentrations assayed by the one stage method using both congenitally deficient and immunodepleted plasma (table 3).

Discussion

Monoclonal antibodies to vWF:Ag and VIII:Ag provided a means of producing an FVIII deficient substrate from fresh plasma which was both HIV antibody and hepatitis B antigen negative and which was also readily available in quantity.

Takase et al recently reported a similar method for the preparation of VIII-immunodepleted plasma using a monoclonal antibody to vWF:Ag and two to VIII:Ag. Their process, however, depletes cryosupernatant plasma. While the procedure described in this study was also capable of removing all detectable VIII from cryosupernatant, the product obtained had
reduced concentrations of fibrinogen which could possibly result in long clotting times when used as a substrate in the one stage VIII assay. The antibodies used in this study are also available commercially (Bioscot Ltd) and the method is thus potentially applicable in any reasonably equipped laboratory.

The two anti-VIII:Ag monoclonal antibodies used by Takase et al are directed to opposite ends of the VIII:Ag molecule. Epitope maps were not available for the antibodies used in this study and the antibodies were chosen on the basis of the results summarised in fig 1, although recent preliminary results suggest that they also are directed to the heavy (ESH 5) and light (ESH 8) chains of factor VIII. The anti-vWF:Ag antibody will adsorb the forms of VIII:Ag which are complexed with vWF:Ag. As ESH 8 adsorbed the greatest amount of VIII:Ag it was the natural choice for this work and ESH 5 was chosen as being the only antibody to adsorb a greater proportion of VIII:C than VIII:Ag. This antibody possibly recognises a form of VIII:Ag which is also biologically highly active. These three antibodies together were successful in removing more than 99% of VIII:C from plasma.

As the results show the same antibodies give greater adsorption in column experiments than in the batch ones. A column of ESHV 8 alone was capable of removing 98% VIII:C from plasma. As ESHV 8 has been shown by methods described in Hornsey et al to bind only to vWF and not to VIII:Ag, then either this 98% VIII:C must have been complexed with the vWF or, more likely, some of the adsorption observed in this small scale trial was due to non-specific binding to the column matrix.

The performance of the columns did not seem to be reduced after 10 runs, even though regeneration experiments only indicated 50% recovery of the VIII:Ag and vWF:Ag from the columns. This could indicate a possible underutilisation of the gel capacity and it may be possible to increase the load of plasma processed before regeneration becomes necessary. A proportion of the two antigens, however, may be destroyed or changed during regeneration, resulting in an underestimate of antigen yield.

The substrate produced is also very low in vWF:Ag, with values of between 1% and 3% of normal plasma. It has been suggested that these low values of vWF:Ag will affect the results obtained when assaying VIII:C in plasma from patients with von Willebrand's disease who have low concentrations of both vWF:Ag and VIII:C. Muntean et al found an increase in VIII:C values when assaying plasma low in vWF by using a substrate which has been depleted of both VIII:C and vWF. Mertens et al and McPherson et al have shown that vWF does not affect the VIII:C promoted activation of factor X by IXa in a purified system. The small number of results obtained here (table 3) indicate that this substrate may be used in assaying plasma concentrations of VIII:C in patients with von Willebrand's disease.

The freeze dried product was tested in seven laboratories in the United Kingdom. Three laboratories reported that it was perfectly acceptable for all their purposes; two said it could be used for assay down to FVIII values of 0-01 IU/ml; the remaining two found the clotting times too long to be acceptable, possibly because they used manual one stage assay techniques with fairly short activation times. These long clotting times could result from the somewhat reduced concentrations of factor XI, and possibly the other contact factors in the product and, therefore, care should be taken during production to avoid such losses by, for example, siliconising all columns. Even with the long clotting times, standard curves were reported which were of comparable slope to those obtained using congenitally deficient plasma or other immunodepleted commercially available substrates.

This substrate can be stored for at least six months at temperatures +4°C and below. At +37°C a fairly rapid loss of performance as a substrate was seen (table 2), possibly due to inactivation of factor V and fibrinogen. This was seen in both the immunodepleted product and the congenitally deficient plasma.

The factor VIII depleted plasma described should be of great value to laboratories performing routine automated one stage coagulation assays for VIII:C in normal plasma, VIII concentrations, and haemophilic plasmas.

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References
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