Purification of blood coagulation factors II, VII, IX and X from bovine citrated plasma

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Purification of Blood Coagulation Factors II, VII, IX and X from Bovine Citrated Plasma

P. Riekers and H. C. Hemker
Department of Internal Medicine, Laboratories of Cardiobiochemistry and Blood Coagulation Biochemistry, University Hospital Leiden, Leiden

Abstract. The bovine coagulation factors II, VII, IX and X were separated from other plasma proteins, and each factor was obtained in a largely pure state by various chromatographic procedures in the presence of diisopropylfluorophosphate (DFP). Factor X was separated from factors II, VII, and IX by DEAE Sephadex chromatography, followed by G-100 filtration and hydroxylapatite chromatography. Separation of factors II and IX from factor VII was achieved by chromatography on DEAE cellulose, followed by G-100 gel filtration and hydroxylapatite chromatography. Factors II and IX were separated by preparative polyacrylamide electrophoresis. After DEAE cellulose chromatography, factor VII was purified by gradient elution from hydroxylapatite.

Key Words
- Purification
- Prothrombin
- Factor VII
- Factor IX
- Factor X
- Bovine plasma

Introduction

The problem of obtaining coagulation factors II, VII, IX and X from plasma in a pure state has been a particularly stubborn one, and this difficulty has retarded progress in coagulation biochemistry appreciably. Methods

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giving partial separation and yielding one or two of these factors in a pure state from the same plasma sample have occasionally been published since 1964. In this article, procedures will be described by which each of the four factors can be obtained separately from the same batch of bovine plasma, the resulting products being free of any other coagulation factor activity and very pure with respect to contaminating proteins. The initial steps consist of aluminium hydroxide adsorption and ammonium sulfate precipitation, as described in the literature for human or rabbit citrated plasma [6, 7, 12, 32, 41, 44]. The other methods applied include chromatography, gel filtration and preparative polyacrylamide electrophoresis, partly along the lines indicated by investigations on the purification of single coagulation factors [5, 8, 9, 13, 14, 16, 17, 25, 27, 29, 34–36, 40, 43, 45]. Following a suggestion of Jackson and Hanahan [17], we added DFP, 10⁻⁴ M, to all preparations and buffer solutions after the ammonium sulfate precipitation step to prevent activation of the coagulation factors.

Materials and Methods

Chemicals and Special Reagents

Aluminium hydroxide moist gel batch No.0340600, kaolin light, and DFP were obtained from British Drug House. Trizma base 121, hemoglobin, egg albumin, cytochrome c and tetra-methylene diamine (Temed) were purchased from Sigma. DEAE cellulose (DE-32) was obtained from Whatman Biochemicals. DEAE Sephadex A-50, Sephadex G-100 and dextran blue 2000 were obtained from Pharmacia. Hydroxylapatite Biogel, HTP No. 9112, was obtained from Biorad. Acrylamide and N,N'-methylene-bis-acrylamide were products of Fluka. Liver alcohol dehydrogenase and pancreas trypsin were obtained from Boehringer Mannheim. Rabbit antitotal bovine serum-antiserum was obtained from the Central Laboratory of the Netherlands Red Cross. All other chemicals were of analytical grade and purchased from Merck.

Thromboplastin from bovine brain was prepared according to Ouwren and Aas [33]. Phospholipid was isolated from the cephalin fraction of human brain according to Milstone [30] and suspended in veronal acetate buffer, pH 7.35, to a concentration of 0.9 mg protein per ml immediately before use. Factor-II reagent was prepared analogously to the method of Koller et al. [21], using bovine instead of human materials. Factor-VII reagent was prepared according to the method of Lechner and Deutsch [24], as modified by Swart et al. [44]. Factor IX-deficient plasma was obtained from a patient with severe hemophilia B (factor IX: 1%). Factor-X reagent was prepared either by adding a prepara-

Abbreviations: DFP = diisopropylfluorophosphate; prep. p.a.a. = preparative polyacrylamide electrophoresis; PPSB = preparation containing prothrombin, proconvertin, Stuart-Power factor and antihemophilia factor B.
tion of factors II and VII to BaSO₄-adsorbed oxalated bovine plasma or by neutralizing and precipitating factor X from normal bovine plasma with a specific antiserum against bovine factor X.

Assay of Blood Coagulation Factors

The activities of factors II, VII, and X were determined in one-stage assays as described for factor II by Koller et al. [21]. Factor IX activity was estimated as described by Velten-Kamp et al. [46]. Factor Xa was determined as described by Delange and Hemker [23]. Normal bovine plasma was obtained after centrifugation of blood (15 min, 1,000 g; and 3 min, 1,500 g) drawn into one-tenth by volume of trisodium citrate, 0.14 M. Hematocrit values were about 40%. It is defined that 1 ml of normal bovine plasma with a coagulation factor activity of 100% contains 1 U of activity of each of the factors II, VII, IX or X.

Initial Steps of Purification Procedure

After adsorption of factors II, VII, IX and X from bovine citrated plasma onto aluminum hydroxide (1% final concentration) at 4 °C, the adsorbent was washed with Na-EDTA, 0.3 M, pH 8.0; and Na-citrate, 0.1 M, pH 8.0. The factors were eluted from the adsorbent with phosphate buffer, 0.25 M, pH 8.0 [44], and concentrated by ammonium sulfate precipitation between 35 and 65% saturation. For factors II and X, a 200-fold purification was obtained at a yield of approximately 30%. For factors VII and IX, the yield and purification were considerably higher, probably due to activation of these factors. The ammonium sulfate concentrate was dialyzed against 0.02 M K-phosphate + 0.1 M NaCl + DFP, 0.1 mM, pH 6.8.

This preparation was called PPSB-2, and was used for separation of coagulation factors II, VII, IX and X by anion exchange chromatography.

Chromatography

All chromatographic procedures were performed at 4 °C. The buffers contained DFP, 0.1 mM. DEAE Sephadex was equilibrated in potassium phosphate buffer, 0.02 M, pH 6.8 ± NaCl, 0.1 M. DEAE cellulose was equilibrated with potassium phosphate buffer, 0.05 M, pH 6.8. Conditions for elution of coagulation factor activity are described in the legends to the figures.

Gel filtration on Sephadex G-100 was done in Pharmacia K 25/100 columns, using phosphate buffer, 0.2 M, pH 6.8, at a flow rate of 3-6 ml/cm²/h. Assuming a linear relation between the logarithm of the molecular weight and the elution volume of the protein [1], estimation of molecular weights was done on G-100 columns precalibrated with dextran blue 2000, liver alcohol dehydrogenase, egg albumin, pancreas trypsin and cytochrome c.

Hydroxyapatite was equilibrated in potassium phosphate buffer, 0.075 M, pH 6.8. On the column, a linear gradient was established to potassium phosphate, 0.5 M, pH 6.8. Before application to the hydroxyapatite column, the samples were made 0.075 M in phosphate.

Protein contents were estimated after Lowry et al. [26], or by measuring the extinction at 280 nm (Zeiss PMQ II spectrophotometer). Immunoelectrophoresis was performed according to Scheidegger [39] on microscope slides.

Column effluents were analyzed conductometrically with the LKB Conductolyzer. Concentration of protein samples was done in Amicon ultrafiltration cells using UM-10 membranes. The UV absorbancies of LKB Uvicord II monitor.

Polyacrylamide Electrophoresis

Analytical polyacrylamide electrophoresis using a 7.5-percent gel of 2.8% cross-linker. Glycoproteins were stained with PAS. Proteins were stained with Sudan orange III.

Preparation p.a.a. was performed in a Quickfit 'Prep Page' equipment. As described by Gag and Louts [15] was used. Bromophenol blue did not contain DFP.

![Fig. 1. DEAE Sephadex chromatography.](image-url)
Purification of Clotting Factors II, VII, IX and X

membranes. The UV absorbancies of column effluents were measured with the aid of an LKB Uvicord II monitor.

Polyacrylamide Electrophoresis

Analytical polyacrylamide electrophoresis was performed according to Davis [10], using a 7.5-percent gel of 2.8 % cross-linking. Amido black was used for protein staining. Glycoproteins were stained with PAS reagent, according to Keyzer [20]. Lipoproteins were stained with Sudan orange III, according to Pratt and Dangrefield [37].

Prep. p.a.a. was performed in a 7-percent separating gel, 2.8 % cross-linking, using the Quickfit 'Prep Page' equipment. As a buffer system, borate buffer according to Gordon and Louis [15] was used. Bromophenol blue served as a tracking dye. The elution buffer did not contain DFP.

![Graph](image-url)

**Fig. 1.** DEAE Sephadex chromatography of PPSB-2 in 0.02 M K-phosphate + 0.1 M NaCl + 0.1 mM DFP, pH 6.8. 70 ml of protein solution (21 mg/ml) was applied. After 50 ml of effluent had been collected, a gradient of 2 x 350 ml was started, ranging from 0.02 K-phosphate + 0.1 M NaCl to 0.02 M K-phosphate + 1 M NaCl + 0.1 mM DFP, pH 6.8. Fraction volume = 2.8 ml. Bed dimensions of anion exchanger = 35 cm x 2.5 cm.

O—Protein, \(\varepsilon_{280}\) nm; ■—factor-II activity; ▲—factor-VII activity; ●—factor-IX activity; ♦—factor-X activity; ——ionic strength, \(\mu\).
Fig. 2. DEAE cellulose chromatography of semipurified coagulation factor preparation (fractions 125–156 from DEAE Sephadex, fig. 1). 90 ml of protein solution (8.3 mg/ml) in 0.05 M K-phosphate + 0.1 mM DFP, pH 6.8, was applied to DEAE cellulose equilibrated in the same buffer. A gradient was used of 2 x 250 ml, starting at 0.05 M K-phosphate + 0.1 M NaCl + 0.1 mM DFP, pH 8.0, to 0.05 M K-phosphate + 0.5 M NaCl + 0.1 mM DFP, pH 8.0. Bed dimensions of anion exchanger = 35 cm x 2.5 cm. Fraction volume = 4.0 ml. O— Protein, E$_{280}$nm; ■— factor II; ▲— factor VII; ●— factor IX; x— factor X; — ionic strength, $\mu$.

Results

Separation of Factor X by DEAE Sephadex Chromatography

The result of a typical experiment in which PPSB obtained from 8 l of plasma was separated on Sephadex A-50 is shown in figure 1. With the use of a linear NaCl gradient, factors II and IX were eluted at an ionic strength, $\mu = 0.48$, and factor VII at $\mu = 0.50$. Factor-X activity was eluted both at $\mu = 0.50$ and 0.57. By coagulation factor assays, the first peak of factor-X activity was shown to coincide with factor X and the second peak with factor X. This second peak (fractions 157–180) served as a source for preparing pure factor X by procedures to be described later. For further purification of factors II, VII, and IX, fractions 125–156 were pooled and

Fig. 3. Hydroxylapatite chromatography of 25 ml factors II and IX after G-200 gel filtration was diluted twice its volume with 0.075 M, pH 6.8 phosphate buffer, at phosphate concentration of 0.01. Hydroxylapatite bed = 25 cm x 2.5 cm, determined according to SUNNER. O— factor IX; x— factor X; — chromatographed on DEAE buffer, 0.05 M, pH 6.8. As and polyacrylamide-gel electrophoresis determined that there were less than 10 components, including factor VII.

Separation of Factor VII

Figure 2 shows that rec purified coagulation factor elution of factors II and IX activity was found coinciding with factor-X activity and II, IX, and XI, as a source of factors II and IX.
Fig. 3. Hydroxylapatite chromatography of a coagulation factor preparation containing mainly factors II and IX after G-100 gel filtration. Before chromatography, the sample was diluted twice its volume with distilled water. Hydroxylapatite was equilibrated in 0.075 M, pH 6.8 phosphate buffer, + DFP, 0.1 mM. Linear gradient = 2 × 200 ml, starting at phosphate concentration of 0.075–0.5 M, pH 6.8 + DFP, 0.1 mM. Dimensions of hydroxylapatite bed = 25 cm × 2.5 cm. Fraction volume = 4.0 ml. Inorganic phosphate was determined according to SUMNER [42]. ○ protein; ■ factor II; ▲ factor VII; ● factor IX; x factor X; — inorganic phosphate (Pi), mM.

rechromatographed on DEAE cellulose after dialysis against phosphate buffer, 0.05 M, pH 6.8. As observed from immunoelectrophoretic analysis and polyacrylamide-gel electrophoresis, this preparation still contained more than 10 components, including albumin and β-globulins.

Separation of Factor VII by DEAE Cellulose Chromatography

Figure 2 shows that rechromatography on DEAE cellulose of a semi-purified coagulation factor preparation (fractions 125–156, fig. 1) resulted in elution of factors II and IX at ionic strength μ = 0.38. The greater part of factor-VII activity was found to emerge from the column at μ = 0.44, coinciding with factor-X activity. The main protein peak (fractions 40–60) served as a source of factors II and IX; factor VII was further purified from frac-
tions 61–90. This elution profile differed from that observed by Pechet and Smith [35], who found factors II and VII to coincide and to be eluted well before factors IX and X.

Purification of Bovine Factors II and IX

Fractions 40–60 of the eluate from DEAE cellulose which contained factor-II and factor-IX activity (fig. 2) were pooled and further freed of factor VII activity and some contaminating protein by gel filtration on Sephadex G-100. The elution volumes for both factor II and factor IX corresponded to a molecular weight of 82,000 [1].

Further purification and partial separation of factor II and factor IX was achieved by chromatography on hydroxylapatite (fig. 3). Contaminating protein material was eluted at phosphate molarity 0.14 M, together with a small amount of factor-II activity. Factor IX activity was eluted at 0.19 M phosphate. Factor IX activity, whereas factor II had a broad range of phosphate. No activity was eluted at 0.05 M phosphate. 

Judged from the results of figures 4 and 5, gel 2B of hydroxylapatite column had two components. Elution of the coagulation factor II and factor IX was confirmed by polyacrylamide electrophoresis, factor II (fig. 4 and 5, parts 2B).

The pooled fractions 108–140 were diluted with 0.05 M by dilution with 0.5 M of veronal acetate buffer, pH 7.4. Horizontally: clotting times; vertically: excised part of the gel. Gel 3: fractions 95–104 after prep. p.a.a. (fig. 6), containing only factor-II activity. ■—Factor II; ○—factor IX.
small amount of factor-II activity. Activated factor VII was eluted at 0.19 M phosphate. Factor IX emerged in a sharp peak at 0.22 M phosphate, whereas factor II had a broad elution profile ranging from 0.22 to 0.30 M phosphate. No activity was eluted at higher phosphate concentration, as was reported by some authors [19, 27, 44].

Judged from the results of polyacrylamide-gel electrophoresis, the pooled fractions 96–107 (fig. 5, gel 2A) and 108–140 (fig. 4, gel 2A) from the hydroxylapatite column had two components in common. Slicing of the gels and elution of the coagulation factor activity demonstrated that on polyacrylamide electrophoresis, factor IX has a somewhat greater mobility than factor II (fig. 4 and 5, parts 2B).

The pooled fractions 108–140 (fig. 3) were brought to phosphate concentration 0.05 M by dilution with distilled water and concentrated by Amicon
UM-10 membrane filtration. This sample was submitted to preparative polyacrylamide electrophoresis. As is shown in figure 6, factor IX was found to be eluted before factor II. Polyacrylamide electrophoresis gave an extensive loss of factor-II activity. Apart from factors IX and II, at least two other proteins could be detected in the protein pattern. No factor VI or thrombin was found in the eluted protein peaks. Fractions 95–104 were pooled and injected into rabbits to obtain an antiserum against bovine factor II [38]. Fractions 60–94 were pooled and used for the adsorption from the antiserum of those antibodies not directed against bovine prothrombin.

Fractions 95–104 were pooled and injected into rabbits to obtain an antiserum against bovine factor II [38]. Fractions 60–94 were pooled and used for the adsorption from the antiserum of those antibodies not directed against bovine prothrombin.

On analytical polyacrylamide electrophoresis (fig. 4, gel 3), factor-II preparation showed one main band, containing factor-II activity, and a minor, more anodal band, which apparently had no coagulation factor activity. Staining of the gels with PAS reagent demonstrated that the main band contained glycoprotein. The gels could not be stained with Sudan orange III.

Table I shows the overall result of the purification procedure for factor II from bovine plasma in a representative experiment. In a set of five experiments, the results were always qualitatively equal, but large variations were observed in the overall yield of factor II after hydroxylapatite chromatography (between 2 and 14 %) and prep. p. a. a. (between 0.5 and 3 %).
Purification of Clotting Factors II, VII, IX and X

Table I. Purification of factor II from bovine plasma.
Results of a representative experiment starting with 16 l of bovine plasma

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Volume, ml</th>
<th>Concentration, U/ml</th>
<th>Total, U</th>
<th>Protein, mg/ml</th>
<th>Spec.act., U/mg</th>
<th>Yield, %</th>
<th>Puri-</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plasma</td>
<td>16,000</td>
<td>1</td>
<td>16,000</td>
<td>75</td>
<td>0.013</td>
<td>100</td>
<td>1</td>
</tr>
<tr>
<td>PPSB-1</td>
<td>2,140</td>
<td>4.20</td>
<td>8,990</td>
<td>5.2</td>
<td>0.81</td>
<td>56</td>
<td>60</td>
</tr>
<tr>
<td>PPSB-2</td>
<td>80</td>
<td>60</td>
<td>4,800</td>
<td>27</td>
<td>2.23</td>
<td>30</td>
<td>170</td>
</tr>
<tr>
<td>DEAE Sephadex</td>
<td>160</td>
<td>28</td>
<td>3,480</td>
<td>8.4</td>
<td>3.35</td>
<td>22</td>
<td>260</td>
</tr>
<tr>
<td>DEAE cellulose</td>
<td>88</td>
<td>35</td>
<td>3,080</td>
<td>12.1</td>
<td>3.64</td>
<td>19</td>
<td>280</td>
</tr>
<tr>
<td>G-100</td>
<td>92</td>
<td>24</td>
<td>2,200</td>
<td>8.0</td>
<td>3.0</td>
<td>14</td>
<td>230</td>
</tr>
<tr>
<td>Hydroxylapatite</td>
<td>132</td>
<td>12</td>
<td>1,584</td>
<td>0.95</td>
<td>12.8</td>
<td>10</td>
<td>960</td>
</tr>
<tr>
<td>Prep. p.a.a.</td>
<td>504</td>
<td>0.7</td>
<td>358</td>
<td>0.08</td>
<td>8.75</td>
<td>2.5</td>
<td>640</td>
</tr>
</tbody>
</table>

Table II. Purification of factor IX from bovine citrated plasma.
Results of a representative experiment starting with 16 l of bovine plasma

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Volume, ml</th>
<th>Concentration, U/ml</th>
<th>Total, U</th>
<th>Protein, mg/ml</th>
<th>Spec.act., U/mg</th>
<th>Yield, %</th>
<th>Puri-</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plasma</td>
<td>16,000</td>
<td>1</td>
<td>16,000</td>
<td>75</td>
<td>0.013</td>
<td>100</td>
<td>1</td>
</tr>
<tr>
<td>PPSB-1</td>
<td>2,140</td>
<td>10.2</td>
<td>21,800</td>
<td>5.2</td>
<td>2.0</td>
<td>136</td>
<td>150</td>
</tr>
<tr>
<td>PPSB-2</td>
<td>80</td>
<td>1,600</td>
<td>128,000</td>
<td>27</td>
<td>59</td>
<td>800</td>
<td>4,550</td>
</tr>
<tr>
<td>DEAE Sephadex</td>
<td>160</td>
<td>50</td>
<td>8,000</td>
<td>8.35</td>
<td>6.0</td>
<td>50</td>
<td>460</td>
</tr>
<tr>
<td>DEAE cellulose</td>
<td>88</td>
<td>280</td>
<td>24,600</td>
<td>12.1</td>
<td>23</td>
<td>155</td>
<td>1,780</td>
</tr>
<tr>
<td>G-100</td>
<td>92</td>
<td>240</td>
<td>22,100</td>
<td>8.0</td>
<td>30</td>
<td>138</td>
<td>2,310</td>
</tr>
<tr>
<td>Hydroxylapatite</td>
<td>132</td>
<td>60</td>
<td>7,920</td>
<td>1.8</td>
<td>33</td>
<td>50</td>
<td>2,540</td>
</tr>
<tr>
<td>Prep. p.a.a.</td>
<td>188</td>
<td>40</td>
<td>7,530</td>
<td>0.2</td>
<td>200</td>
<td>47</td>
<td>15,400</td>
</tr>
</tbody>
</table>

Separation of factor IX from contaminating factor-II activity, as found in fractions 96-107 from the hydroxylapatite column (fig. 3), was achieved by prep. p.a.a. Factor IX migrated faster than factor II, as had been observed also for prep. p.a.a. of a factor-II preparation (fig. 6). As is shown in figure 5, gel 3, fractions with maximal factor-IX activity showed one main protein band on analytical polyacrylamide electrophoresis, this band coinciding with factor-IX activity, and being stainable with PAS reagent. These fractions
were used to obtain antibodies against factor IX in rabbits [38]. A minor contamination (about 10%) with possibly factor-II activity was observed.

Table II shows the overall result of the purification procedure for factor IX from bovine plasma in a representative experiment. The high yield and purification are undoubtedly due to activation of factor IX during the purification procedure.

**Purification and Isolation of Bovine Factor VII**

Isolation of factor VII started from fractions 61–90 from DEAE cellulose chromatography (fig.2). Considerable purification of factor VII was achieved by gel filtration on G-100. The elution volume for factor VII corresponded to a molecular weight of about 68,000. No factor-II or factor-IX activity was observed. The fractions having maximal factor-VII activity were pooled, concentrated and used for further purification by hydroxylapatite.

As is demonstrated in figure 7, factor VII was eluted at a phosphate concentration of 0.19 M, whereas the main portion of protein material was not retarded on hydroxylapatite (0.075 M). The elution profile of protein elution profile, indicating other protein material.

On analytical polyacrylamide gel electrophoresis, a protein band had factor-VII activity, another minor, more acidic band, which could not be determined with the band representing factor VII from the DEAE cellulose column.

So little material was left after the procedure that pooled fractions 31–33 (fig.7) were used as a factor VII preparation. As is demonstrated in figure 7, factor VII was eluted at a phosphate concentration of 0.19 M, whereas the main portion of protein material was not retarded on hydroxylapatite (0.075 M). The elution profile of protein elution profile, indicating other protein material.

On analytical polyacrylamide gel electrophoresis, a protein band had factor-VII activity, another minor, more acidic band, which could not be determined with the band representing factor VII from the DEAE cellulose column.

So little material was left after the procedure that pooled fractions 31–33 (fig.7) were used as a factor VII preparation. As is demonstrated in figure 7, factor VII was eluted at a phosphate concentration of 0.19 M, whereas the main portion of protein material was not retarded on hydroxylapatite (0.075 M). The elution profile of protein elution profile, indicating other protein material.
Purification of Clotting Factors II, VII, IX and X

not retarded on hydroxylapatite at the starting phosphate concentration (0.075 M). The elution profile of factor-VII activity did not coincide with the protein elution profile, indicating that factor VII was still contaminated with other protein material.

On analytical polyacrylamide electrophoresis (fig. 8, gel 2), the main protein band had factor-VII activity and it could be faintly stained with PAS reagent. Another minor, more anodal protein band was observed, the nature of which could not be determined. The factor-VII band corresponded well with the band representing factor-VII activity in the pooled fractions 61–90 from the DEAE cellulose column (fig. 8, gels 1A and B).

So little material was left after the many stages of the purification procedure that pooled fractions 31–40 after hydroxylapatite chromatography (fig. 7) were used as a factor-VII preparation, although analytical polyacrylamide electrophoresis suggested that further purification was still possible. The preparation was used for immunizing rabbits to obtain an antiserum against bovine factor VII.
Table III. Purification of factor VII from bovine citrated plasma

Results of a representative experiment starting from 16 l of bovine plasma

| Fraction         | Volume, ml | Concentration, U/ml | Total, U | Protein, mg/ml | Spec.act., U/mg | Yield, % | Puri-   |
|------------------|------------|---------------------|----------|----------------|-----------------|----------|fication|
| Plasma           | 16,000     | 1                   | 16,000   | 75             | 0.013           | 100      | 1      |
| PPSB-1           | 2,140      | 5.83                | 12,470   | 5.2            | 1.1             | 77       | 86     |
| PPSB-2           | 80         | 51                  | 4,080    | 27             | 1.9             | 26       | 145    |
| DEAE Sephadex    | 160        | 18                  | 2,880    | 8.4            | 2.2             | 18       | 166    |
| DEAE cellulose   | 128        | 12.5                | 1,600    | 2.5            | 5.0             | 10       | 385    |
| G-100            | 35         | 18.5                | 650      | 0.6            | 31.0            | 4        | 2,380  |
| Hydroxylapatite  | 28         | 18.0                | 500      | 0.1            | 180             | 3        | 14,000 |

Fig. 9. Analytical polyacrylamide-gel electrophoresis. Gel 1: fractions 157–180 (fig. 1) after DEAE Sephadex chromatography, containing factors VII and X; 230 μg protein. Gel 2A: pooled fractions eluted from G-100 between 175–210 ml (fig. 10); 110 μg protein. Gel 2B: factor-VII and factor-X assays in 0.5-cm slices of a gel (ordinate), run parallel to gel 2A with the same sample. Extraction of coagulation factor activity with 0.5 ml veronal acetate buffer, pH 7.4. Horizontally: clotting times; vertically: excised part of the gel. Gel 3: fractions 45–60 after hydroxylapatite chromatography (fig. 11); 125 μg protein. ▲—Factor VII; x—factor X.

Fig. 10. G-100 gel filtration in 0.5-cm slices of a gel, fractions 157–180 after DEAE Sephadex elution (fig. 10). Y—Factor VII; X—factor X.

Table III shows the result of the purification of factor VII from bovine plasma in a representative experiment starting from 16 l of bovine plasma. A 33% yield was obtained with a purification of 14,000.

Purification and Isolation of Factor VII

Fractions 157–180 from DEAE Sephadex chromatography were used as a starting material for the isolation of factor VII. JACkSON et al. [18]. The material was subjected to hydroxylapatite chromatography; the position of the main factor VII peak was determined. Factor X was eluted between 175–210 ml. G-100 gel filtration of this fraction resulted in a further purification of factor X from the contaminating factors. From the elution volume of about 84,000 was calculated, and fractions 157–210 ml were pooled. One should note that the clotting time

Fraction volume = 2.8 ml. O—
Table III shows the result of the purification procedure for factor VII from bovine plasma in a representative experiment. The overall yield was 3% and a purification of 14,000-fold was reached.

**Purification and Isolation of Bovine Factor X**

Fractions 157–180 from DEAE Sephadex chromatography (fig. 1) served as a starting material for the isolation of factor X, as previously reported by Jackson et al. [18]. The material contained some factor-VII activity as well. Polyacrylamide electrophoresis revealed the presence of at least four protein bands; the position of the main protein band correlated well with the position where maximal factor-X activity was found in sliced gels (fig. 9, gels 1 and 2b).

G-100 gel filtration of this preparation resulted in poor separation of factor X from the contaminating factor-VII activity (fig.10). Considerable separation of factor X from other proteins emerging near the void volume was obtained.

From the elution volume of factor-X activity, a molecular weight of about 84,000 was calculated. The fractions between the elution volumes 175–210 ml were pooled. On polyacrylamide-gel electrophoresis, this pre-
Fig. 11. Hydroxylapatite chromatography of fractions 175–210 (fig 10) in phosphate buffer, 0.075 M, pH 6.8. Linear gradient: 2 × 150 ml phosphate, 0.075–0.5 M, pH 6.8 + DFP, 0.1 mM. Phosphate concentration was measured after SUMNER [42]. Dimensions of hydroxylapatite bed—22 cm × 1.5 cm. Fraction volume = 2.8 ml. — Protein; ▲— factor VII; x— factor X; — phosphate concentration, M.

Preparation showed two poorly separated bands and a weak third band. The two bands coincided with maximal factor-VII and factor-X activity in sliced gels (fig.9, gels 2A and B). No factor-II or factor-IX activity was found in this preparation.

As is shown in figure 11, factor-X activity, present in the G-100 effluent, was eluted from hydroxylapatite at a phosphate concentration of 0.25 M; other protein material was not retarded by hydroxylapatite or was eluted at a phosphate concentration of 0.19 M.

Simultaneously with factor X, some factor-VII activity was eluted at 0.25 M phosphate. However, in a factor-Xa assay, factor X appeared to be partly activated; consequently, shortening of clotting time in a factor-VII assay may be expected. The complete coincidence and constant fractional ratio of factors VII and X argued in favor of this factor-VII activity to be an artefact.
On polyacrylamide gels, the factor-X preparation from the hydroxylapatite column revealed one main protein band having factor-X activity. Glycoprotein staining with PAS reagent gave a positive reaction at the same position as the main band observed after protein staining with amido black. Three minor, more cathodal contaminants were present. No further purification of this preparation was attempted, because monospecific antiserum against bovine factor X was obtained by injecting into rabbits those portions of sliced gels containing maximal factor-X activity.

Table IV shows the overall result of the purification procedure for factor X from bovine plasma in a representative experiment. Although considerable purification was obtained by DEAE Sephadex chromatography, this step was accompanied by a considerable loss of factor X because only the second part of the eluted bovine factor-X activity was used for further purification. The final yield of bovine factor X was in the order of 4%, and an approximately 5,000-fold purification was reached.

Discussion

Our aim was to find schemes by which factors II, VII, IX and X can be isolated from the same batch of bovine plasma as separate entities (fig. 12). Under our conditions, DEAE Sephadex chromatography resulted in separation of coagulation factor activity from many other plasma proteins, in-
including albumin, γ-globulin and fibrinogen. In accordance with other authors [13, 18, 34, 45], a partial separation was effected between the factors II and X, the latter having a much higher affinity for the anion exchanger.

The presence of DFP appeared to be necessary for prevention of activation of coagulation factors due to chromatographic procedures, as has been established by JACKSON and HANAHAN [17]. Even in the presence of DFP, factor X eluted in a double peak as was observed previously by JACKSON et al. [18] and by MILSTONE et al. [31]. We can confirm the finding of JACKSON and HANAHAN [17] that both kinds of factor X have the same mobility on polyacrylamide electrophoresis. We were able to demonstrate that the protein peak eluted at the lower ionic strength contained factor X₁, the one at higher ionic strength only factor X [4]. Thus, DFP did not completely abolish this activated factor X. In some experiments, the peak with factor X₁ activity gave rise to double-peak formation on G-150 gel filtration, the two species having molecular weights of 115,000 and 88,000. Chromatography or electrophoresis in the presence of sodium dodecyl sulfate (SDS) did not give any indication of the occurrence of aggregation of factor X with other plasma material. By gel filtration, the molecular weight for factor X of about 90,000 was also made for our factor-X preparation.

The best purification of factor X was from the first DEAE Sephadex peak, with the factor-X peak from DEAE cellulose and the factor-X peak from DEAE Sephadex A-50. The presence of DFP appeared to be necessary for prevention of activation of coagulation factors due to chromatographic procedures, as has been established by JACKSON and HANAHAN [17]. Even in the presence of DFP, factor X eluted in a double peak as was observed previously by JACKSON et al. [18] and by MILSTONE et al. [31]. We can confirm the finding of JACKSON and HANAHAN [17] that both kinds of factor X have the same mobility on polyacrylamide electrophoresis. We were able to demonstrate that the protein peak eluted at the lower ionic strength contained factor X₁, the one at higher ionic strength only factor X [4]. Thus, DFP did not completely abolish this activated factor X. In some experiments, the peak with factor X₁ activity gave rise to double-peak formation on G-150 gel filtration, the two species having molecular weights of 115,000 and 88,000. Chromatography or electrophoresis in the presence of sodium dodecyl sulfate (SDS) did not give any indication of the occurrence of aggregation of factor X with other plasma material. By gel filtration, the molecular weight for factor X of about 90,000 was also made for our factor-X preparation.

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other plasma material. By gel filtration, several authors determined a molecular weight for factor X of about 85,000 [13, 18, 34] which observation was also made for our factor-X preparation.

The best purification of factor X was achieved by taking the second part of the factor-X peak from DEAE Sephadex. The resulting factor-X preparation showed one main band on analytical polyacrylamide electrophoresis, and one precipitation line on Ouchterlony plates against rabbit antitotal bovine serum-antisemur or rabbit antitbovine factor-X antiserum and on immunoelectrophoresis [38].

The yield was poor, because a part of factor-X activity was left in the first DEAE Sephadex peak, which was used for the purification of factors II, VII, and IX. In the factor-X preparation, the relative amounts of factor X and factor VII have repeatedly been found in a ratio of 10:1. It is known that pure factor X also shortens the recalcification time of factor VII-deficient plasma containing 50% factor X. We cannot conclude from clotting assays that factor VII was absent in the factor-X preparation, but the factor-VII activity measured can be explained by the influence of factor X on the factor-VII assay.

The factor-VII preparation obtained after hydroxylapatite chromatography does not significantly shorten the recalcification time of factor X-deficient bovine plasma. So, we concluded factor X to be absent in the factor-VII preparation. The molecular weight of bovine factor VII ranged from 72,000 to 62,000, as judged from G-100 gel filtration; this was in agreement with the observations of other investigators [11]. The yield of factor VII was poor, mainly because chromatographic procedures gave rise to activation phenomena in the factor-VII molecules, making the factor more labile. When DFP was added to the eluting phosphate buffer after aluminum hydroxide adsorption of factor VII from bovine plasma, a lower factor-VII activity was found in the eluate than when DFP was omitted.

Factor II and factor IX could not be separated by DEAE cellulose chromatography [9, 35, 45]. Partial separation of these factors was achieved by hydroxylapatite chromatography [27], as has been shown also for the human coagulation factors [5, 43]. In our experiments, factor II eluted at a lower phosphate concentration than was found for human prothrombin.

On both analytical and preparative electrophoresis, factor IX showed a higher electrophoretic mobility than factor II; this result was not obtained by COX and HANAHAN [9].

Our results are in agreement with the observation of PECHET and SMITH [35], who showed factor IX to have a lower isoelectric point than factor II.
The molecular weights of bovine factors II and IX were determined by G-100 gel filtration to be approximately 82,000 for both factors [9, 45]. Controversially, no indication of multiple factor-II elution patterns or factor-II polymorphism was obtained in the present study [2, 3, 22, 28].

The factor-II preparation still contained contaminating proteins. On immunoelectrophoresis of this preparation against antitotal bovine serum antiserum, two precipitation lines were observed, as was also the case with rabbit antihuman prothrombin antiserum, obtained by injecting this prothrombin preparation into rabbits [38]. On polyacrylamide electrophoresis of the factor-II preparation, one main protein band, coinciding with factor-II activity, was observed, together with a thin, slightly more anodal protein band.

Factor-IX preparations still contained about 5% factor-II activity. On polyacrylamide electrophoresis, one protein band coinciding with factor-IX activity was observed. Immunodiffusion experiments of the preparation against rabbit antihuman factor-IX antiserum showed only one precipitation line, as was also the case for immunoelectrophoresis experiments with the same antiserum [38].

References

Purification of Clotting Factors II, VII, IX and X


Request reprints from: Dr. P. Reekers, Blood Transfusion Service, St. Radboud Ziekenhuis, Nijmegen (The Netherlands)