Development of a Simple Chromogenic Factor VIII Assay for Clinical Use

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Abstract. The aim of this study was the development of a simple chromogenic factor VIII assay for practical clinical use. The criteria that the assay fulfills are: (1) The method is so sensitive that even 1% factor VIII in human plasma is easily detected. (2) The method is linear in the amount of factor VIII from 0 to 200% in plasma. (3) The pipetting scheme is very simple; two reagents are prepared, reagent 1 (factor IXa, thrombin, Ca²⁺ and phospholipids) and reagent 2 (factor X). Then we pipet at t = 0 s, 100 μl diluted plasma + 100 μl reagent 1 in a reaction tube; at t = 30 s, 100 μl reagent 2 in the same tube and at t = 90 s, 200 μl of the reaction mixture in a cuvette with 700 μl EDTA buffer (stop buffer) and the formed factor Xa is measured with a chromogenic substrate. (4) The reaction components are stable during at least a whole working day. Factor VIII was measured in an assay using bovine clotting factors, so one avoids the risk of viral infections, which one might catch by working with clotting factors isolated from human plasma.

Introduction

For practical clinical use we developed an assay for factor VIII in human plasma. Some work concerning the development of this assay [1] and its practical use [2] was already presented at the congress on Thrombosis and Haemostasis in Brussels, 1987. In order to be of practical suitability, the assay must be very sensitive (less than 1% factor VIII must be detectable), the assay must be linear from 0 to 200% factor VIII in plasma, the pipetting steps should be as minimal as possible and the reaction components should be stable during a whole working day.

The activation of the zymogen factor X into factor Xa by the serine protease factor IXa [review in ref. 3] is strongly increased by negatively charged phospholipids, Ca²⁺ ions and activated factor VIII [review on factor VIII in ref. 4]. The complex of factor IXa, factor VIIIa, Ca²⁺ and phospholipids is called the intrinsic factor X activating complex, or intrinsic factor X activator [5–9]. Van Dietjen et al. [10] described the conditions for optimal interaction of bovine factor VIIIa in the assembly of the factor X activating complex. In their article they showed that a complete factor X activating complex will activate about 900 molecules factor X.
per minute. Thus, under conditions that the activated factor VIII is completely bound in
the factor X activating complex, 900 molecules factor Xa are formed per minute per
molecule factor VIIIa. Factor Xa can be
measured with a chromogenic substrate. So,
if one chooses the right conditions a small
amount of factor VIIIa can form a large
amount of factor Xa and in this way a sensi-
tive factor VIII assay is obtained. However,
we deal with human factor VIII which may
act differently from bovine factor VIII, for
which the optimal conditions for a good
assay are described [10]. There are two pos-
sible approaches, one can use either human
clotting factors, or bovine clotting factors in
the factor VIII assay. We strongly favored
the use of bovine clotting factors for the fol-
lowing reasons. Bovine blood is easy to ob-
tain in large quantities in contrast to human
blood, isolation procedures for the purifica-
tion of bovine clotting factors are well de-
scribed and finally one avoids the risk of
viral infections using bovine plasma instead
of human plasma.

We investigated if bovine clotting factors
are suitable for a human factor VIII assay,
which was proven to be the case. To obtain a
system as sensitive as possible we have var-
ied each of the reaction components of the
factor X activating complex to determine
their optimal concentrations.

\[
\text{FX} \cdot \text{FVIIIa} \cdot \text{FIXa} \cdot \text{Ca}^{2+} \cdot \text{PL} \rightarrow \text{FXa}
\]

where FX is the factor X activating complex.

We successively searched for the optimal
conditions for factor VIII activation with
thrombin, the optimal factor IXa concentra-
tions to bind all factor VIIIa, the optimal
phospholipid concentration and composition
to obtain a rapid factor Xa formation and
finally the optimal pH for factor X activation.

To determine the required factor X concen-
tration we measured its \( K_m \) for the complex.

Materials and Methods

Materials

FXa-substrate (CH_3OCO-D-CHG-Gly-Arg-pNA-
AcOH) and \( \alpha \)-NAPAP \( \left[ N-\alpha-(2\text{-naphtylsulfo-
nylglucyl})-D,L\text{-amidinophenyl-alanine-piper-
} \right] \text{dide hydroiodide} \) should be added, to inhibit
the large amount of thrombin which is present
in the assay mixture for complete factor VIII
activation.

Stability studies showed lyophilized re-
agents could be stored for several months
without loosing their activity and that recon-
stituted reagents were stable for at least a
whole working day. Moreover, we showed
that the assay was linear from 0 to 200% fac-
tor VIII in plasma and that less than 1% fac-
tor VIII could be well detected.

Materials and Methods

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FXa-substrate (CH_3OCO-D-CHG-Gly-Arg-pNA-
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nylglucyl})-D,L\text{-amidinophenyl-alanine-piper-
} \right] \text{dide hydroiodide} \) were obtained from Pentapharm, Switzerland;
DEAE-Sephadex and Sepharose-4B were from Pharm-
macia, Sweden; Heparin-Agarose was kindly donated
by T. Janssen-Claessen.

All other reagents were of the highest grade avail-
able and were supplied by either Merck (FRG) or
Sigma (USA).

Phospholipids and Phospholipid vesicles

Phosphatidyl choline was extracted from egg yolks
according to Bligh and Dyer [11] and purified as
described by Comfurius and Zwaal [12]. Phosphati-
dyl serine and phosphatidyl ethanolamine were pre-
pared in a similar way, however, extracted from bo-
Vince brains. Vesicles were prepared by sonication of a phospholipid mixture in a buffer containing 30 mM Tris-HCl and 175 mM NaCl (pH 7.9) using an MSE Mark II 150-Watt ultrasonic disintegrator set at 9 µ peak to peak amplitude. The composition of the vesicles was 75 mol% phosphatidyl choline and 25 mol% phosphatidyl serine as indicated.

**Proteins**

Bovine factor X was isolated according to Fuji-kawa et al. [3] and bovine factor IXa as described by van Dieijen et al. [8]. Thrombin was prepared as described by Wagenvoord et al. [4]. Congenital factor VIII deficient plasma was kindly donated by T. Repucci (Centre de Transfusion, Liège, Belgium).

**Plasmas**

Prepared containing more than 100% factor VIII were prepared by mixing reference plasma with a cryoprecipitate (10 U/ml).

**Results**

**Properties of the Chromogenic Substrate CH₃OCO-D-CHG-Gly-Arg-pNA-AcOH**

We have studied the kinetic properties of the factor Xa substrate CH₃OCO-D-CHG-Gly-Arg-pNA-AcOH (FXa-substrate). First we determined the kinetic constants of FXa-substrate hydrolysis by factor Xa. For that reason a Lineweaver-Burk plot was made (fig. 1a). For Kₘ we measured 142 µM and Vₘₐₓ was 386 nM p-nitroanilide liberated/s·nM factor Xa, or 229.5 milliabsorbance units/nM factor Xa·min. Remarkable is the effect of a high NaCl concentration on Kₘ, which becomes 27.3 µM in the presence of 2.5 M NaCl, Vₘₐₓ, however, does not change by the increased NaCl concentration. Thus, by addition of NaCl the affinity of the FXa substrate for factor Xa increases, which leads to a higher rate of hydrolysis.

Since in the factor VIII assay mixture a large amount of thrombin is present, it was necessary to determine the kinetic constants of the FXa substrate hydrolysis by thrombin. Figure 1b shows that FXa substrate is hydrolysed by thrombin, however, much slower than factor Xa. Vₘₐₓ is 12.6 nM p-nitroanilide liberated/s·nM thrombin or 7.52 milliabsorbance units/nM thrombin·min and Kₘ is 59.0 µM. In reagent 1 of the FVIII assay a large amount of thrombin is present (fig. 6), which causes a very rapid hydrolysis of the FXa substrate, so it cannot be used unless an effective thrombin inhibitor is present which does not inhibit factor Xa.

Figure 2 shows that the thrombin inhibitor α-NAPAP is able to inhibit thrombin effectively without inhibiting factor Xa very much. At a concentration of 1 µM α-NAPAP, thrombin is inhibited by 99.5%, but factor Xa only by 1%. By using 1 µM α-NAPAP (final concentration in the cuvette), the hydrolysis rate of FXa-substrate by thrombin is brought back to 21.8 nM/min, which is only 1% of the rate to be expected by the factor Xa formed in an assay with normal plasma.

On basis of the results of figures 1 and 2 factor Xa was measured by addition of 100 µl FXa substrate (800 µM) plus 10 µM α-NAPAP to 900 µl of a factor Xa sample.

**Preparation of a Factor VIII Assay Mixture**

Our first goal was the preparation of a mixture containing the purified bovine clotting factors which could detect human plasma factor VIII as sensitive as possible, therefore we varied each of the components of the factor X activating complex to determine their optimal concentration.

In figure 3a we studied the effect of thrombin on factor VIII activation in human plasma. Maximal factor VIII activation oc-
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Fig. 1. Lineweaver-Burk plots of FXa-substrate hydrolysis by factor Xa and thrombin. a Plot of FXa-substrate hydrolysis by factor Xa. The factor Xa concentration was 1 nM. The experiment was done in standard buffer (175 mM NaCl, 50 mM Tris-HCl (pH 7.9), 0.5 mg/ml human serum albumin); b experiment in which a buffer with 2.5 M NaCl is used. b Plot of FXa-substrate hydrolysis by thrombin. The thrombin concentration was 1 nM and standard buffer was used.

curs at thrombin concentrations of 30 nM or more. The presence of more than 30 nM thrombin does not lead to more factor VIIIa formation, however, it is preferable to have a large excess of thrombin in the assay because of the presence of antithrombin III and α₂-macroglobulin in the plasma, which will inactivate the thrombin. An additional advantage of excess of thrombin is that it will saturate all plasma serine protease inhibitors and thus formed factor Xa is not inactivated.

Figure 3b shows the effect of factor IXa concentration on the factor Xa formation by the factor X activating complex. The reaction rate is a reflection of binding of factor VIIIa into the complex. The figure shows that optimal binding occurs when the factor IXa concentration is 50 nM or higher.

In figures 3c and d the effect is shown of phospholipids on the activation of bovine factor X by the factor X activating complex composed of bovine factor IXa, human factor VIIIa and Ca²⁺. Figure 3c shows the effect of varying amounts of phosphatidyl ser-
Fig. 3. Optimization of the human factor VIII assay. a) Activation of human plasma factor VIII with thrombin: to 200 μl factor IXa (200 nM), CaCl₂ (10 mM), phospholipids (40 μM; 75 mol% phosphatidyl choline, 25 mol% phosphatidyl serine) 100 μl thrombin were added. At t = 0 s 50 μl 10 times diluted human plasma were added and at t = 30 s 50 μl factor X (2 μM). Samples of 300 μl were taken at t = 60 s and the factor Xa formation was measured. b) Effect of the factor IXa concentration on the rate of factor Xa formation: to 200 μl thrombin (200 nM), CaCl₂ (10 mM), factor IXa (200 nM) 100 μl phospholipids were added. At t = 0 s 50 μl 10 times diluted human plasma were added and at t = 30 s 50 μl factor X (2 μM). Samples of 300 μl were taken at t = 90 s to measure the formed factor Xa. c) The vesicles concentration was kept at 20 μM, whereas the composition was varied. The phosphatidyl serine content was 0–50 mol% and the phosphatidyl choline content 100–50 mol%. d) Effect of phospholipid concentration on the rate of factor Xa formation. The vesicle composition was kept at 25 mol% phosphatidyl serine and 75 mol% phosphatidyl choline.
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Fig. 4. Lineweaver-Burk plot of factor X activation by the factor X activating complex composed of bovine factor IXa and human factor VIIa. To 200 μl thrombin (200 nM), CaCl₂ (10 mM), factor IXα (200 nM) phospholipids (40 μM; 25 mol% phosphatidyl serine, 75 mol% phosphatidyl choline) at t = 0 s 100 μl 20 times diluted human plasma were added. At t = 30 s 100 μl factor X (67-800 nM) were added. Samples of 300 μl were taken at t = 90 s to measure the formed factor Xa.

activity in the vesicles (fixed concentrations) on the factor X activating reaction: maximal activity is found at a phosphatidyl serine concentration of around 25 mol%. In figure 3d one can observe a rapid increase of the enzymatic activity when the phospholipid concentration (with fixed composition) increases from 0 to 10 μM, then becomes maximal and subsequently slows down at still higher phospholipid concentrations. This result is in agreement with an earlier report [8], however, valid for proteins which all were isolated from bovine blood.

We also have investigated the effect of phosphatidyl ethanolamine and cholesterol addition to the vesicles. When a phosphatidyl choline/phosphatidyl serine ratio of 3 is maintained and either phosphatidyl ethanolamine or cholesterol is added, there is no effect at low concentrations of both compounds. However, when phosphatidyl ethanolamine is present in a concentration of more than 15 mol%, the activity of the vesicles drops. For cholesterol the same effect is found at concentrations of more than 20% (w/w). So we can conclude that vesicles of a composition of 25 mol% phosphatidyl serine and 75 mol% phosphatidyl choline have an optimal activity, which cannot be improved by addition of other compounds.

Figure 4 shows the Lineweaver-Burk plot of factor X activation by the complete factor X activating complex. The two kinetic parameters that can be derived from the figure are Kₘ, which is 31.4 nM, and Vₘₐₓ, which is 241 milliabsorbance units/min, when a 20 times diluted reference plasma is used (11.7 nM factor Xa/min).

We also have investigated the effect of the pH on the complete factor X activating complex (containing bovine factors, but human factor VIIIa). The activity showed a broad maximum between pH 7.6 and 8.5, whereas at lower and higher pH values the activity decreased. As a result of these experiments we decided to use buffers with pH 7.9.

In figure 5 we have measured factor Xa formation by time in an assay containing the human factor VIII and the other components of the factor X activating complex, which were present in optimal concentrations as judged from figures 3 and 4. Two reagents were prepared: reagent 1, containing 300 nM factor IXα, 300 nM thrombin, 15 mM CaCl₂ and 60 μM phospholipid vesicles (75 mol% phosphatidyl choline and 25 mol% phosphatidyl serine) and reagent 2 containing 1 μM factor X. By mixing 1 part reagent 1, 1 part diluted plasma and 1 part reagent 2, the final concentrations of each of the clotting factors were optimal for maximal factor Xa forma-
Fig. 5. Linearity of the factor X activating reaction. To 1 ml of 300 nM factor IXa, 300 nM thrombin, 60 μM phospholipids (75 mol% phosphatidyl choline, 25 mol% phosphatidyl serine), 15 mM CaCl₂ at t = 0 s 1 ml 100 times diluted human plasma was added. At t = 30 s 1 ml factor X (1 μM) was added. Samples of 200 μl were taken in time to measure formed factor Xa.

Stability of the Factor VIII Assay Reagents

A point of practical importance is the stability of the reagents after preparation, lyophilization, storage and reconstitution. We prepared reagents 1 and 2. Reagent 1 containing 300 nM thrombin, 300 nM factor IXa, 15 mM CaCl₂ and 60 μM vesicles (75 mol% phosphatidyl choline and 25 mol% phosphatidyl serine), was lyophilized in portions of 2 ml. Also reagent 2 containing 1 μM factor X was lyophilized in this way. Both reagents were stored at 4 °C. After reconstitution they were tested. We determined the protein concentrations in the reagents and the activity of the reagents in the factor VIII assay.

We determined the thrombin and factor IXa concentration in reagent 1 after reconstitution. The thrombin concentration varied from 258 to 290 nM and the factor IXa concentration.
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Fig. 7. Linearity of the FVIII assay. Plasmas were prepared containing 0–200% factor VIII (normal pooled plasma = 100%) which were diluted 10 times. The reagents described in figure 6 were used and the same pipetting sequence was done, however, 200-µl samples were taken at 90 s.

concentration from 281 to 295 nM. After reconstitution we measured the factor X concentration in reagent 2, which was 0.8–1.0 µM. Also in this case there was not much loss of activity. Because the concentration of the proteins was chosen so that small changes in their concentration would not effect the factor VIII assay, we expected no change in the activity. In figure 6 this is shown. Before lyophilization and after reconstitution the activity was the same and, moreover, the activity remained at the starting level even after half year of storage of the lyophilized reagents.

We also tested the stability of the reagent after reconstitution during the working day and after storage at 4 ºC during the night. The activities are expressed in percentages: before lyophilization, 100%; on day 1, after reconstitution of the reagents, the activities were 101% at 10.00 h and 101% at 17.00 h. At day 3 at 11.00 h, the activity was 93%. In other experiments similar results were found.

Sensitivity of the Factor VIII Assay

Finally we have tested the sensitivity of the chromogenic factor VIII assay by measuring plasmas containing 0–200% factor VIII. In figure 7 the results of these tests are shown. One can observe a complete linearity between the factor VIII content of the plasma and the factor Xa formation in the reaction mixture. Plasma containing 1% factor VIII gives rise to a 2–3 times higher factor Xa formation than fully factor VIII deficient plasma, so one can easily measure 1% factor VIII in plasma. Using the same dilutions it is possible to measure 200% without underestimating the reaction rate. When we use normal pooled plasma (100% factor VIII), about 12 nM factor Xa is formed in the assay mixture, so in the cuvette 2.4 nM factor Xa will be present, which gives an increase in absorbance of about 200 milliabsorbance units/min under the described reaction condition.

Discussion

In this article we described the development of a chromogenic factor VIII assay, which can be done in only a few pipetting steps. The test is linear from 0 to 200% factor VIII in human plasma. The reagents are stable when they are lyophilized for at least half a year and after reconstitution they do not lose activity during a whole working day. After reconstitution they can be stored at 4 ºC and used during at least 3 days. Because of its simplicity the test can be auto-
mated. A suggestion to minimize the pipetting steps is to mix the diluted plasma with factor X (reagent 2) and add the reagent 1 to this mixture. Then stop the reaction after 1–1.5 min with the EDTA buffer. When this sequence of pipetting is used, only two steps should be done at fixed times. In this case factor VIII activation precedes the formation of the complete factor X activating complex. However, this is not a real problem, because the time necessary for complete factor VIII activation is only a few seconds, since thrombin is present in high concentration and when the lag time is the same for all factor VIII concentrations, the linearity of the measurement is fully maintained.

If we compare human factor VIII with bovine factor VIII in respect to their action in the factor X activating complex, their resemblance is very close. The turnover rate of the factor X activating complex composed of only bovine factors is about 900 min⁻¹ [10]. We can estimate this number from our results. In human plasma about 0.25 nM factor VIII is present [10], so in the assay of figure 6, 4.2 pM factor VIIIa is present which gives a rate of 6 nM FXa/min, so the turnover number is about 1,400 min⁻¹, a value which is in the same order as was found for the bovine factor X activating complex.

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References


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