Some Effects of Calcium on the Activation of Human Factor VIII/Von Willebrand Factor Protein by Thrombin

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ABSTRACT When Factor VIII/von Willebrand factor (FVIII/vWF) protein is rechromatographed on 4% agarose in 0.25 M CaCl₂, the protein and vWF activity appear in the void volume, but most of the FVIII procoagulant activity elutes later. Recent evidence suggests that the delayed FVIII procoagulant activity is a proteolytically modified form of FVIII/vWF protein that filters anomalously from agarose in 0.25 M CaCl₂. To test whether or not thrombin is the protease involved, the effect of 0.25 M CaCl₂ on FVIII/vWF and its reaction with thrombin was examined. About 30% of the FVIII procoagulant activity was lost immediately when solutions of FVIII/vWF protein were made 0.25 M in CaCl₂. When FVIII in 0.15 M NaCl was activated with 0.04 U thrombin/ml and then made 0.25 M in CaCl₂, the procoagulant activity of a broad range of FVIII/vWF protein concentrations remained activated for at least 6 h. But, in 0.25 M CaCl₂, the increase in FVIII procoagulant activity in response to thrombin was much more gradual and once activated, the procoagulant activity was stabilized by 0.25 M CaCl₂. When thrombin-activated FVIII/vWF protein was filtered on 4% agarose in 0.15 M NaCl, there was considerable inactivation of FVIII procoagulant activity; however, the procoagulant activity that did remain eluted in the void volume. In contrast, when thrombin-activated FVIII/vWF protein was filtered in 0.25 M CaCl₂, the FVIII procoagulant activity eluted well after the void volume and remained activated for 6 h. The procoagulant peak isolated by filtering nonthrombin-activated FVIII/vWF protein on agarose in 0.25 M CaCl₂ was compared to that isolated from thrombin-activated FVIII/vWF protein. Both procoagulant activity peak proteins had about the same specific vWF activity as the corresponding void volume protein. Before reduction, the sodium dodecyl sulfate gel patterns for the two procoagulant activity peak proteins were the same. After reduction, the gel pattern for the nonthrombin-activated procoagulant activity peak protein contained bands of 195,000, 148,000–120,000, 79,000, 61,000, 51,000, and 18,000 daltons whereas the pattern for the reduced thrombin-activated procoagulant activity peak protein always lacked the higher molecular weight bands, but consistently showed the lower four molecular weight bands to be well resolved. Taken together, these results imply that thrombin generates the FVIII procoagulant activity that is stabilized by 0.25 M CaCl₂ and elutes aberrantly from 4% agarose in that solvent.

INTRODUCTION

The initial physicochemical characterization of a purified protein with Factor VIII (FVIII) procoagulant activity appeared to establish that it is a glycoprotein with a mol wt > 1 million daltons and a covalent subunit structure (1–9). Subsequently it was shown that this glycoprotein also had von Willebrand factor (vWF) activity and it was suggested that the two activities (i.e. FVIII and vWF) might be properties of the same molecule and conceivably of each subunit (10–12). More recently it has been shown that when the FVIII/vWF protein(s) is rechromatographed on 4% agarose in certain high ionic strength solvents, particularly 0.25 M CaCl₂, virtually all of the protein and vWF activity elute in the void volume, but most of the FVIII procoagulant activity elutes much later (13–21). These observations have led to the development, explicitly or implicitly, of four different hypotheses for the structure-function relationships of the FVIII/vWF

1Abbreviations used in this paper: FVIII, Factor VIII; SDS, sodium dodecyl sulfate; TAMe, p-toluenesulfonyl-L-arginine methylester; vWF, von Willebrand Factor.

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protein: (a) The FVIII procoagulant activity is associated with a small subunit noncovalently bound to a large "carrier" subunit. This model would imply a definite stoichiometry between the carrier protein and the small, active subunit. (b) The two activities are associated with two different proteins that form a complex. Depending on binding constants, the FVIII procoagulant and vWF proteins might be expected to have several favorable combining ratios, but the stoichiometric requirements of this model would be less rigid than for the first. (c) The FVIII procoagulant and vWF activities are associated with two different proteins that simply co-purify. This model would impose no stoichiometric restrictions on the combining ratios of these two unrelated proteins. (d) The FVIII procoagulant and vWF activities are properties of a single molecule composed of disulfide-bound identical subunit polypeptide chains. In this model, the action of thrombin or thrombin-like enzymes on the molecule gives rise to a derivative which has FVIII procoagulant activity and which elutes later from 4% agarose in 0.25 M CaCl₂.

Recent evidence from this laboratory was interpreted to support the last model (22, 23). This conclusion was based on our demonstration that the FVIII procoagulant activity appeared larger than 400,000 daltons by Sephadex G-200 chromatography in 0.25 M CaCl₂ and by sodium dodecyl sulfate (SDS) polyacrylamide-gel electrophoresis. Also the quantities of protein associated with the two activities were incompatible with a reasonable stoichiometry, and the FVIII procoagulant activity protein also had vWF activity. The presence of vWF activity in the FVIII procoagulant activity peak and the SDS-gel pattern for the reduced FVIII procoagulant activity peak protein suggested that this protein is a proteolytically modified FVIII/vWF species that interacts with agarose in 0.25 M CaCl₂ and thereby elutes anomalously. This report presents additional studies of the effects of calcium on the activities of the FVIII/vWF protein(s) as well as on its gel filtration and electrophoretic behavior before and after exposure to thrombin.

METHODS

Reagents. Reagent-grade chemicals were used without further purification. Unless otherwise noted, all solutions were buffered to pH 7.35 with 0.05 M Tris hydrochloride (Tris).

Protein concentrations. These were measured by the absorbance at 280 nm, corrected for light scattering, and expressed as absorbance units.

FVIII procoagulant activity assays. FVIII procoagulant activity was measured by the kaolin-activated partial thromboplastin time method using hemophilic plasma with <1% FVIII activity as the substrate (8). The modifications required to measure FVIII procoagulant activity in the presence of 0.25 M CaCl₂ have been described in detail (23); in short, one must both dilute the test sample and change the order of reagent addition. More specifically, the FVIII solution was diluted to 0.043 M CaCl₂ and added to a 37°C mixture of 0.1 ml of 0.05 M Tris-0.15 M NaCl, 0.1 ml of Thrombopax (Ortho Diagnostics, Inc., Baritan N. J.) and 0.1 ml of hemophilic substrate plasma; the time required to form a visible clot was then recorded. The FVIII procoagulant activity in units per milliliter was computed from a standard curve obtained by assaying one of the National Institute of Health (NIH) reference plasmas (0.76 or 1.2 U/ml) in 0.043 M CaCl₂, using the same order of reagent addition. With these modifications, the clotting time for each dilution of reference plasma was longer and the slope of the standard curve of clotting time vs. log percent dilution was steeper than the slope obtained when the assay was performed using the regular order of reagent additions.

Thrombin assays. The esterase activity of thrombin was measured by determining the rate of hydrolysis of p-toluenesulfonyl-L-arginine methyl ester (TAME) by the method of Walsh (24). The change in absorbance at 247 nm for the hydrolysis of 1 μmol of TAME was taken to be 0.409 (1/cm)/(μM) (24).

The protease activity of thrombin was measured by the rate of hydrolysis of the synthetic peptide S-2160, N-benzoyl-L-phenylalanyl-L-valyl-L-arginine-p-nitroanilide hydrochloride (AB Bofors, Molndal, Sweden), by the method of Blomback et al. (25). The substrate was dissolved in deionized water to a concentration of ~0.65 mg/ml. A volume of 0.15 ml or less of the test sample was mixed with 1.0 ml of 0.025 M Tris-0.025 M imidazole-0.15 M NaCl-0.02% sodium azide buffer, pH 8.2, at 37°C; then 0.125 ml of substrate also at 37°C was added. The incubation mixtures were kept at 37°C for 5 min at which time the reaction was stopped by adding 0.15 ml of glacial acetic acid. The absorbance at 405 nm, which is proportional to the concentration of p-nitroaniline released, was then determined and expressed as a function of the thrombin concentration. The assays were performed in test tubes that had been soaked overnight in a 0.1% solution of bovine serum albumin (Sigma Chemical Co., St. Louis, Mo.) in deionized water; this procedure was more effective than silicone coating in preventing the loss of thrombin by adsorption to the test tube wall.

Purification of FVIII/vWF protein(s). The starting material for the purification of FVIII/vWF was either the intermediate or high purity concentrate as prepared by the American National Red Cross Laboratory, Bethesda Md., (courtesy of Dr. M. Wickerhauser and Dr. Y. L. Hao) (26). The intermediate-purity FVIII/vWF was converted to high purity FVIII/vWF by dissolving in 0.2 M e-aminoacaproic acid to about 18 mg protein/ml, adsorbing for 10 min with 10 mg of bentonite (Sigma Chemical Co.,)/ml and then precipitating the protein twice with polyethylene glycol (Carbowax 4,000, Union Carbide Corp., New York) as previously described (23). The final precipitate was washed four times with ice-cold 8% ethanol in 0.05 M Tris-0.15 M NaCl buffer; subsequently the precipitate was dissolved in 0.05 M Tris-0.15 M NaCl at about 20 mg protein/ml and chromatographed as described in a later section. The FVIII/vWF concentrates supplied in the high purity form were not subjected to the bentonite adsorption step, but instead were simply dissolved in 0.2 M e-aminoacaproic acid and chromatographed. The high-purity FVIII/vWF solutions were further purified by gel filtration at room temperature on a 4 x 38-cm column of 4% agarose in 0.05 M Tris-0.15 M NaCl, pH 7.35, (Biogel A15m, Bio-Rad Laboratories, Richmond Calif.) as previously described (23). The volume of the FVIII/vWF sample applied to the column was about 25 ml, the flow rate ~15 ml/h and the fraction vol ~7 ml. The void volume fractions were pooled on the basis of their content of FVIII
procoagulant or vWF activity; the protein was precipitated by adding one-third volume 40% polyethylene glycol at 4°C. The FVIII/vWF precipitate was collected by centrifugation at 6,000 g and 4°C, redissolved in 0.05 M Tris-0.15 M NaCl buffer to a concentration of at least 3 absorbance U/ml and stored at −20°C in 0.1−1 ml aliquots in plastic tubes. Our yield was about 35% relative to the starting concentrate and the product was stable for up to 9 mo at −20°C. The ultra-high purity FVIII/vWF prepared by this method was homogeneous by SDS gel electrophoresis before and after reduction and gave a single immunoprecipitin line when it was diffused or electrophoresed against unabsorbed rabbit antiserum (23). The ultra-high purity FVIII/vWF protein had about 25 FVIII U/absorbance U; however, the level of vWF units was found to vary depending on the methods used to relate unitage to that of whole plasma. Although the construction of a ratio of FVIII activity to vWF activity might seem useful, particularly since it should equal unity, it probably is not a reliable index of the purity of FVIII/vWF protein. There are several reasons why it is difficult to relate these two activities back to normal plasma; (a) the exact concentration of the FVIII/vWF protein(s) in normal plasma is not known; (b) trace proteases quickly destroy FVIII activity but leave most of the vWF activity intact; (c) only about 25% of the FVIII activity in normal plasma is recovered by the majority of purification procedures; (d) other plasma proteins may have differential effects on either of the two activities; and (e) the results of assays measuring the two activities of highly purified FVIII/vWF may not be exactly comparable to those of assays conducted in whole plasma. Hence the vWF activity of our purified FVIII/vWF material is defined essentially the same as previously (23).

Enhancement of FVIII procoagulant activity by thrombin. The purified human thrombin used for these experiments was supplied as a lyophilized powder (568 NIH U/absorbance U at 280 nm; courtesy of Dr. D. L. Aronson, Bureau of Biologics, FDA, Bethesda, Md.). The thrombin was reconstituted in 0.05 M Tris-0.15 M NaCl and stored at a concentration of 200 NIH U/ml at 4° or −20°C. Just before use, the stock solution of thrombin was diluted in polyethylene test tubes which had been previously equilibrated with a thrombin solution as recommended by other investigators (27). Factor VIII was activated by adding 4−10 μl of the desired thrombin dilution per ml of highly purified FVIII/vWF. The reaction mixture was incubated at room temperature in a plastic tube which was gently tilted for the first min to insure adequate mixing. FVIII procoagulant activity in the incubation mixture was then monitored at selected time intervals.

Electrophoresis. Sample preparation and SDS-polyacrylamide gel electrophoresis were done as described by Weber and Osborn (28) with slight modifications as reported from our laboratory (8). Staining, destaining, and gel storage were done as described earlier (8).

Chromatographic studies. Ultra-high purity FVIII/vWF protein was rechromatographed on 4% agarose columns in the solvent systems indicated in Results; a constant flow rate was maintained for each column by the use of a peristaltic pump at the column outlet. Plastic chromatographic columns with agarose bed dimensions of 0.9 × 11 and 0.9 × 26 cm were used in these experiments. In each experiment a sample volume of ~0.5 ml was applied directly to the top of the agarose. Flow rates of 4 and 6 ml/hr and fraction volumes of 0.45 and 0.6 ml were used for the 11- and 26-cm columns, respectively. If the columns were to be run in 0.25 M CaCl2, solid CaCl2 was added to the FVIII/vWF sample just before chromatography. Fractions to be pooled for SDS-gel electrophoresis were dialyzed overnight at 4°C against 2,000 times their vol of distilled water, quick-frozen, lyophilized, and stored at −20°C for future SDS-gel electrophoretic studies. To remove as much residual calcium as possible, samples which contained CaCl2 were routinely dialyzed overnight against 2,000 vol of distilled water containing a fourfold molar excess of EDTA and then dialyzed twice for 2−3 h each time against 2,000 vol of distilled water to remove the EDTA.

Platelet studies. The ristocetin-induced aggregation of washed human platelets was measured with a ChronoLog aggregometer (Chrono-Log Corp., Havertown, Pa.) (23). To compute vWF activity, the maximum slope of percent transmittance vs. time was determined and taken as the initial velocity (v) of platelet aggregation. A standard curve of 1/v vs. 1/FVIII/vWF (as determined from the absorbance at 280 nm) was constructed for dilutions of ultra-high purity FVIII/vWF in 0.15 M NaCl-0.05 M Tris. The reciprocal of the concentration of purified native FVIII/vWF required to produce the 1/v value of the test sample was then determined from the standard curve and taken as an estimate of the vWF activity in the test sample.

RESULTS

Effect of calcium chloride on FVIII procoagulant activity. To examine the effect of 0.25 M CaCl2 on procoagulant activity before thrombin activation, 20 different samples of FVIII/vWF at different concentrations of protein were divided into two sets; one set was left in 0.15 M NaCl-0.05 M Tris and the other set was adjusted to 0.25 M CaCl2-0.05 M Tris with solid CaCl2. Each sample in the two sets was adjusted to 0.043 M CaCl2 (23) just before it was assayed at three dilutions by the modified method for solutions containing calcium. Fig. 1 shows that the samples ex-

**Figure 1** A comparison of the FVIII procoagulant activity of the FVIII/vWF protein in 0.25 M CaCl2 vs. 0.15 M NaCl. The solid and open circles correspond to samples assayed at 15 min and 24 h, respectively, after the addition of 0.25 M CaCl2. The slope of this line indicates that the samples in 0.25 M CaCl2 were ~30% less active than those in 0.15 M NaCl. This loss of FVIII procoagulant activity occurred immediately and no additional losses were observed.
posed to 0.25 M CaCl₂ had only ~70% of the FVIII procoagulant activity of the corresponding samples in 0.15 M NaCl; greater inactivation seemed to occur at low concentrations of the FVIII/vWF protein. It should be noted that the loss of procoagulant activity occurred immediately upon the addition of 0.25 M CaCl₂ and further inactivation could not be detected during the next 24 h.

To test the effect of 0.25 M CaCl₂ on thrombin-activated FVIII, six concentrations of FVIII/vWF protein ranging from 0.2 to 5.0 absorbance U were each incubated with a final thrombin concentration of 0.04 NIH U/ml and assayed for FVIII procoagulant activity with time. Again, all samples were adjusted to 0.043 M CaCl₂ just before they were assayed for procoagulant activity. In Fig. 2, the mean result of six separate experiments confirmed that in 0.15 M NaCl, thrombin rapidly enhanced FVIII procoagulant activity which then gradually fell to an average of ~40% of the control level by 6 h. However, when the FVIII/vWF-thrombin reaction mixture was adjusted to 0.25 M CaCl₂ at ~10 min after activation had occurred, an immediate 25% loss of FVIII procoagulant activity was observed; the procoagulant activity then declined to about 140% of the control value over the next hour and remained essentially at this level for the next 6 h. These results demonstrate that 0.25 M CaCl₂ has a stabilizing effect on the procoagulant activity of the FVIII/vWF protein.

**Effect of calcium on the thrombin-FVIII/vWF interaction.** Solutions of FVIII/vWF protein were prepared in either 0.15 M NaCl or 0.25 M CaCl₂ and adjusted to an absorbance of 1.28 at 280 nm. Thrombin was added to each sample to give a final concentration of 0.04 NIH U/ml. The effect on procoagulant activity was monitored by removing aliquots from each solution, adjusting to 0.043 M CaCl₂ and assaying by the modified method referred to earlier. Fig. 3 shows that in NaCl, FVIII procoagulant activity increased to a maximum of ~300% of the control value in about 10 min. Activity then decreased to control levels by about 2 h and to substantially inactivated levels after 4 h.

Fig. 3 also shows that in 0.25 M CaCl₂, a rapid activation of FVIII procoagulant activity by thrombin did not occur. Instead, there was a gradual increase in activity to ~180% of control levels by about 2 h and then a maintenance of this level for another 4 h. These experiments reproducibly demonstrated that 0.25 M CaCl₂ inhibited the rapid activation of FVIII procoagulant activity by thrombin. Furthermore, as discussed in the section above, 0.25 M CaCl₂ appeared to stabilize FVIII procoagulant activity at a partially activated level.

**Effect of EDTA on FVIII/vWF activities.** Attempts were made to restore the partial loss of FVIII procoagulant activity that occurred immediately when the FVIII/vWF protein was exposed to 0.25 M CaCl₂. Accordingly, samples of FVIII/vWF protein, with and without prior exposure to 0.25 M CaCl₂, were dialyzed twice against 100 vol of 0.15 M NaCl-0.05 M Tris-0.02 M EDTA for 1 h each time, and then three times against 0.15 M NaCl-0.05 M Tris in the same vol ratio and for the same time periods. Similar to earlier studies using whole plasma (29), the EDTA-treated FVIII/vWF solutions retained <1% of the FVIII procoagulant activity of control FVIII/vWF solutions.
which had been dialyzed in an identical manner but never exposed to EDTA. The inactivation of FVIII procoagulant activity was not reversed when the EDTA-exposed FVIII/vWF protein was incubated overnight with calcium or other alkaline earth metals, alkali metals, transition metals, or lanthanides. Dialysis against EDTA had no effect on the vWF activity of FVIII in NaCl.

Effect of calcium on thrombin. To determine whether the decreased rate of activation of FVIII procoagulant activity in 0.25 M CaCl₂ was due to an effect on thrombin rather than on FVIII/vWF, the activity of thrombin in 0.25 M CaCl₂ was compared to that in 0.15 M NaCl by three assays: (a) enhancement of FVIII procoagulant activity; (b) TAME hydrolysis; and (c) proteolysis of a synthetic peptide.

Thrombin dissolved in each of the two solvents was used to activate FVIII/vWF protein concentrations of approximately 0.1, 0.5, and 2.0 mg/ml. Since, as discussed above, FVIII procoagulant activity was enhanced only very slowly when the FVIII/vWF was dissolved in 0.25 M CaCl₂, these activation experiments were performed using FVIII/vWF dissolved in 0.15 M NaCl. At a final concentration of 0.04 NIH U/ml, the two thrombin solutions enhanced FVIII procoagulant activity to the same extent and at the same rate. The activation profiles at all three protein concentrations were similar to those shown in Figs. 2 and 3 for activation of FVIII/vWF in 0.15 M NaCl.

The esterase activity of thrombin dissolved in 0.15 M NaCl or 0.25 M CaCl₂ was examined by the TAME assay. The rates of hydrolysis for the two solutions of thrombin were essentially the same: 0.193 and 0.189 TAME U/thrombin NIH U. The two thrombin solutions gave total changes of 0.368 and 0.357 absorbance U at 247 nm/NIH U of thrombin. These results indicate that 0.25 M CaCl₂ has little or no effect on the esterase activity of thrombin.

The proteolytic activity of thrombin dissolved in 0.15 M NaCl or 0.25 M CaCl₂ was assessed spectrophotometrically by monitoring the amount of p-nitroaniline cleaved from the synthetic peptide, N-benzoyl-L-phenylalanyl-L-valyl-L-arginyl-p-nitroanilide. The absorbance at 400 nm, proportional to the concentration of p-nitroaniline released by thrombin proteolysis, was determined after 5–6 min at 37°C. Fig. 4 shows the mean results of four such experiments; the percent loss of activity of thrombin in 0.25 M CaCl₂ is presented as a function of thrombin concentration. In such analyses 0.25 M CaCl₂ had a slight inhibitory effect on the proteolytic activity of thrombin, greatest at low thrombin concentrations, but never more than ~20%. At the thrombin concentration usually used to enhance FVIII procoagulant activity, the proteolytic activity of thrombin in 0.25 M CaCl₂ was ~84% of that for thrombin in 0.15 M NaCl. These three sets of experiments indicate that the inhibitory effect of 0.25 M CaCl₂ on the activation of FVIII/vWF by thrombin is mostly a direct effect on the FVIII/vWF protein.

Gel filtration studies on thrombin-activated FVIII/vWF protein. When thrombin-activated FVIII/vWF was rechromatographed on 4% agarose in 0.15 M NaCl, the elution profile observed was significantly different from that in 0.25 M CaCl₂. These studies were performed by first assaying a sample of FVIII/vWF protein in 0.15 M NaCl for procoagulant activity, activating one-half of the sample with thrombin for ~10 min and then simultaneously gel filtering the two samples on identical-sized 4% agarose columns in 0.15 M NaCl. If the chromatography was to be performed in 0.25 M CaCl₂, the appropriate amount of solid calcium chloride was added to the thrombin-activated sample at the peak of activation and at the same time, to the nonthrombin-activated sample. Then, the two samples were each filtered simultaneously in 0.25 M CaCl₂ on 4% agarose columns of the same size. In all of these chromatography experiments the elution fractions from the paired gel filtration analyses of FVIII/vWF and thrombin-activated FVIII/vWF were assayed immediately for FVIII procoagulant activity. To compare sets of experiments in which different protein concentrations were used, all absorbance or procoagu-

**Calcium and Thrombin Effects on Factor VIII/von Willebrand Protein**

![Figure 4](image-url)
lant values of a given pair were expressed as a fraction of those of the peak absorbance or procoagulant value, respectively, in the control, nonthrombin-activated chromatogram.

The protein and FVIII procoagulant activities of control or thrombin-activated FVIII/vWF samples were eluted congruently as sharp peaks in the void volume from 4% agarose in 0.15 M NaCl and no other FVIII procoagulant activity was found in later fractions (Fig. 5). The mean protein elution patterns were identical in the paired chromatograms. Contrary to our expectations, the FVIII procoagulant activity recovered from the chromatography of thrombin-activated FVIII/vWF was always less than or equal to control levels. Considerable variation in recovery of FVIII procoagulant activity was seen in the four experiments illustrated. In each experiment, the loss of FVIII activity during chromatography of thrombin-activated FVIII/vWF was about the same as that of an aliquot of the mixture kept at room temperature for the time required for the chromatography. Such inactivation occurred even when the serine protease inhibitor, p-nitrophenylguanidinobenzoate, was added after 10 min to control and thrombin-activated FVIII/vWF solutions at a 10,000-fold molar excess relative to thrombin.

When similar paired experiments were performed in 0.25 M CaCl₂, very different results were obtained. Fig. 6 compares the mean chromatogram for 10 agarose gel filtrations of nonthrombin-activated FVIII/vWF to the mean chromatogram for 10 identical experiments with thrombin-activated FVIII/vWF solutions. Both agarose columns were eluted simultaneously in 0.25 M CaCl₂ under identical conditions. Shown in Fig. 6, the protein void volume patterns were essentially identical whether or not the FVIII/vWF sample had been exposed to thrombin; however, the void volume FVIII procoagulant activity was much less and the shifted FVIII procoagulant activity peak was substantially larger in the elution pattern for thrombin-activated FVIII/vWF.

When FVIII/vWF or thrombin-activated FVIII/vWF was rechromatographed on 4% agarose in 0.25 M CaCl₂, the elution profiles of the vWF activity were essentially identical. As indicated in Table I, the specific vWF activities were about the same in the void volumes and FVIII procoagulant activity peak regions of the two chromatograms. These data clearly show that despite the enhancement of the procoagulant activity peak by thrombin, the vWF activity is not affected. Hence, these results expand previous reports that thrombin has no inhibitory or stimulatory effect on the vWF activity of the FVIII/vWF protein (18, 22). It should be noted that only about 35% of the vWF activity was observed when FVIII/vWF protein dissolved in 0.25 M CaCl₂ was added to the ristocetin-platelet aggregation assay as opposed to the same concentration of FVIII/vWF protein in 0.15 M NaCl. Preliminary results suggest that the excess calcium ion primarily suppresses platelet reactivity in the ristocetin assay, although some inhibition may also occur on the FVIII/vWF protein as well.

**SDS-gel electrophoretic analyses.** Fig. 7 depicts the SDS-7.5% polyacrylamide gel analyses of fractions from the 0.25 M CaCl₂-4% agarose gel filtration of FVIII/vWF and thrombin-activated FVIII/vWF. Nonreduced, the void volume protein recovered from the agarose-0.25 M CaCl₂ filtration of FVIII/vWF or thrombin-activated FVIII/vWF barely penetrated the top of the gel. The reduced SDS-gel pattern for the two void volume samples were also identical, each showing only one subunit band of 195,000 daltons. The SDS-
Thrombin and filtered simultaneously under the pooled and concentrated fractions of the solutions in CaCl₂ mean absorbance and the filtration patterns of procoagulant FVIII/vWF sample (coltumil pattern with 51,000, 18,000, 61,000-, and 18,000-dalton bands). The major bands, mostly above 100,000 daltons, were clearly present and well resolved. Similar results were obtained when FVIII/vWF was chromatographed on 4% agarose in 0.25 M CaCl₂ and the separated FVIII procoagulant activity peak then activated by thrombin. These experiments suggest a directed correlation between enhancement of FVIII procoagulant activity, the disappearance of the high molecular weight subunit bands, and the presence of the four lower molecular weight bands in the SDS-gel patterns of the reduced FVIII procoagulant activity peak protein. The presence of the same four bands in the reduced SDS-gel pattern of the FVIII procoagulant activity peak isolated from the nonthrombin-activated FVIII/vWF can be explained by some degree of in vivo or in vitro proteolytic modification by trace thrombin.

**DISCUSSION**

The effects of calcium on the FVIII procoagulant activity of FVIII/vWF are complex. After causing an immediate 30% reduction of the FVIII procoagulant activity, calcium stabilized both native and thrombin-activated FVIII/vWF. This stabilization, coupled with the direct observation that FVIII procoagulant activity was rapidly and irreversibly destroyed when FVIII/vWF was incubated with 0.02 M EDTA, suggests that the FVIII/vWF molecule contains bound metal ion(s) that hold it in a specific, relatively stable conformation. Removal of the metal ion(s) apparently results in an irreversible conformational change since the addition of calcium or other cations did not restore procoagulant activity.

When thrombin-activated FVIII/vWF was rechromatographed on 4% agarose in 0.15 M NaCl or 0.25 M CaCl₂, the FVIII procoagulant activity was again stabilized in CaCl₂, but not in NaCl. Furthermore, the peak of FVIII procoagulant activity eluted in the void volume with the protein peak in 0.15 M NaCl, but considerably after the void volume in 0.25 M CaCl₂. As shown in Fig. 6, comparison of the 4% agarose-0.25 M CaCl₂ chromatograms of FVIII/vWF and thrombin-

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**TABLE I**

Relative Specific vWF Activities of Native FVIII/vWF, Thrombin-Activated FVIII/vWF and Pools from 4% Agarose-0.25 M CaCl₂ Columns

<table>
<thead>
<tr>
<th></th>
<th>Native</th>
<th>Thrombin activated</th>
</tr>
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<tbody>
<tr>
<td>FVIII/vWF in CaCl₂</td>
<td>1.00</td>
<td>1.00</td>
</tr>
<tr>
<td>CaCl₂ void volume protein</td>
<td>0.89</td>
<td>0.75</td>
</tr>
<tr>
<td>CaCl₂ activity peak protein</td>
<td>0.75</td>
<td>0.78</td>
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* Mean of three experiments.

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**FIGURE 6** Comparison of the 4% agarose-0.25 M CaCl₂ filtration patterns of FVIII/vWF and thrombin-activated FVIII/vWF (column size: 0.9 × 11 cm). Each panel shows the mean absorbance and the mean FVIII procoagulant activity for 10 sets of experiments. For any given set, one-half of a FVIII/vWF sample was filtered on 4% agarose in 0.25 M CaCl₂ while the other half of the sample was activated by thrombin and filtered simultaneously under the same conditions on an identical-sized column. Absorbance at 280 nm and FVIII procoagulant activity were expressed as fractions of the peak values, respectively, for the control chromatogram. The fractions in the cross-hatched portion of the descending limb of the FVIII procoagulant peak in each chromatogram were pooled and concentrated for analyses by SDS-gel electrophoresis.
activated FVIII/vWF indicates that prior thrombin treatment of the FVIII/vWF protein lessens the amount of FVIII procoagulant activity in the void volume and increases the amount that elutes later. These chromatographic experiments imply that the FVIII procoagulant activity separated by 4% agarose-0.25 M CaCl₂ chromatography of purified, presumably “native,” FVIII/vWF is in fact a very small amount of thrombin-activated FVIII/vWF protein; however, if filtered on 4% agarose in 0.15 M NaCl, this procoagulant-active species still elutes in the void volume with the major protein peak. This latter finding cannot be reconciled with the report from another group that thrombin promoted the formation of FVIII-procoagulant-active species which eluted well into the inner volume of 4% agarose in 0.15 M NaCl (30).

Our SDS-gel analyses of the reduced FVIII procoagulant activity peak protein provide evidence of a change in FVIII/vWF structure resulting from thrombin cleavage. Previous reports which showed that the gel pattern of thrombin-activated FVIII/vWF was identical to that of nonthrombin-activated FVIII/vWF (7,8) could indicate that: (a) no proteolysis had occurred; (b) very minor proteolysis had occurred; or (c) only a small proportion of FVIII/vWF molecules were actually cleaved by thrombin. The results presented in Fig. 7 support the last interpretation since the SDS-gel analyses of the very small amount of protein in the shifted FVIII procoagulant activity peak separated by 4% agarose-0.25 M CaCl₂ chromatography clearly demonstrated structural differences between the nonthrombin- and thrombin-activated species. In both cases, the nonreduced FVIII procoagulant activity peak protein was too large to penetrate the SDS gels. After reduction of disulfide bonds, several subunit bands were seen in both samples; however, compared to the gel of the reduced FVIII procoagulant activity protein from FVIII/vWF not exposed to thrombin, the gel of the protein from thrombin-activated FVIII/vWF showed a lack of higher molecular weight bands and well-resolved lower molecular weight bands. This pattern suggests that additional thrombin treatment caused more cleavages in a molecule already partially modified by thrombin. A similar reduced SDS-gel pattern was obtained when FVIII procoagulant activity protein was thrombin-activated after separation by 4% agarose-0.25 M CaCl₂ chromatography of nonthrombin-treated FVIII/vWF.

vWF activity data provide further evidence that the procoagulant activity peak protein is a thrombin-modified form of the void volume protein. The vWF specific activities of the void volume and FVIII procoagu-
lant activity peak proteins from normal and thrombin-activated FVIII/vWF are essentially identical; thus the observed vWF activity is directly proportional to protein concentration. The presence of vWF activity in the FVIII procoagulant protein from thrombin-activated FVIII/vWF demonstrates that an intact 195,000-dalton subunit is not necessary for vWF activity; this result agrees with our report that FVIII/vWF, which was degraded by plasmin and contained no intact 195,000-dalton subunits, retained ~70% of its original vWF activity (31). The presence of vWF activity despite the absence of the 195,000-dalton subunit in the FVIII procoagulant protein from thrombin-activated FVIII/vWF also demonstrates that vWF activity is a property of the FVIII procoagulant activity peak protein rather than of protein trailing from the void volume. When considered in toto, these results support and expand our earlier interpretation that the FVIII procoagulant protein is a proteolytically modified FVIII/vWF molecular species (23).

There have been two recent attempts to examine this hypothesis by partially purifying FVIII/vWF in the presence of protease inhibitors and then examining the gel filtration properties of the product (32, 33). In the absence of protease inhibitors, Beck et al. (32) found that FVIII procoagulant activity could be separated from vWF activity by 4% agarose gel filtration in 1.0 M NaCl. But FVIII/vWF prepared in the presence of kallikrein inhibitor (Trasylol, FBA Pharmaceuticals, Inc., New York) could no longer be separated into its component activities when filtered in 1.0 M NaCl (32). In a similar study, Poon and Ratnoff (33) found that FVIII/vWF prepared in the presence of benzamidine, heparin, soybean trypsin inhibitor, e-amino-caproic acid, and hirudin could still be separated into its component activities on 4% agarose in 0.25 M CaCl₂. The difference between these two studies may reflect differences in the FVIII/vWF preparations, the effectiveness of the inhibitors or the effects of 1.0 M NaCl and 0.25 M CaCl₂. In both of these studies a small amount of activated FVIII/vWF could have been generated before the inhibitors were added, either in vivo or during the venipuncture.

Ultracentrifugation of plasma at high ionic strength has also been reported to produce slow-sedimenting FVIII procoagulant activity (15). It is difficult to draw conclusions about molecular size from such experiments, since FVIII/vWF in whole plasma might exist as a lipoprotein complex. As such, the lipid content might enhance the procoagulant activity of the FVIII/vWF protein as well as induce flotation effects during centrifugation. This objection was examined by sucrose-gradient experiments on isolated FVIII procoagulant activity peak protein that suggested a low molecular weight for this species (34). Unfortunately the low levels of FVIII activity recovered and the broad peak observed in those experiments make it difficult to estimate molecular weights. But the most serious problem with those studies is the method by which the FVIII procoagulant activity peak was processed before centrifugation. The albumin-stabilized FVIII procoagulant activity peak was “dissociated” from highly purified FVIII/vWF by CaCl₂-agarose chromatography and then collected into a solution of only partially purified albumin. Since the authors indicate that the albumin contained proteins not precipitated by 50% saturated ammonium sulfate, there is a good possibility that other clotting factors such as Factor Xa or lipids might be present among the contaminants and hence might shorten the Factor VIII assay. Furthermore contaminant lipoproteins might bind the FVIII procoagulant protein and give rise to flotation effects to account for an anomalously low molecular weight for the procoagulant activity. Finally, the FVIII procoagulant activity peak protein might possibly interact with sucrose in the same way that it interacts with agarose and thereby sediment aberrantly. The anomalies observed in the 4% agarose-0.25 M CaCl₂ gel filtration of FVIII/vWF should provide a similar caution for the sucrose-gradient experiments: the proof of molecular weight for the slowly sedimenting activity in sucrose-gradient experiments must await isolation of this species and its characterization by such methods as SDS-gel electrophoresis and or analytical ultracentrifugation in denaturing solvents.

Our results reported here, in conjunction with those of others can be explained by the following model: First, the FVIII/vWF protein present in circulating blood is proposed to be procoagulant-inactive until activated by thrombin. Second, when blood is withdrawn from the circulation, a small amount of thrombin or thrombin-like protease forms as a consequence of the wound or venipuncture and activates some of the FVIII/vWF protein. Third, because FVIII procoagulant activity, but not vWF activity, is very susceptible to destruction by trypsin-like enzymes such as plasmin, the FVIII/vWF protein must be continuously degraded in vivo such that its procoagulant potential is totally destroyed while virtually all of its vWF activity remains intact. FVIII/vWF would then be isolated as a mixture of nonactivated precursor FVIII/vWF, proteolytically inactivated FVIII/vWF and proteolytically activated FVIII/vWF. When the FVIII/vWF protein is filtered on 4% agarose in 0.25 M CaCl₂, the large protein peak that elutes in the void volume contains the precursor FVIII/vWF and proteolytically inactivated FVIII/vWF species. The FVIII procoagulant activity protein, resulting from thrombin cleavage of precursor FVIII/vWF, elutes later from 4% agarose-0.25 M CaCl₂ as a result of interaction with the agarose gel matrix and perhaps self-aggregation. Finally, our data clearly demonstrate that calcium stabilizes the procoagulant activity of the FVIII/vWF molecule. This effect has been essential for the isola-
tion of FVIII procoagulant protein with an extremely high specific activity and likely will be very important in the recovery of sufficient material for continuing structure function studies on the FVIII procoagulant protein.

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REFERENCES


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