Binding of Human Factor VII and VIIa to Monocytes

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ABSTRACT Human coagulation Factors VII and VIIa bind with equal affinity to monocytes stimulated with endotoxin. Equilibrium binding studies performed at 0°C using 125I-labeled Factor VII and VIIa showed the dissociation constant (Kd) to be 82 pM with 3,600 binding sites/monocyte. Ca++ was required for Factor VII and VIIa interaction with monocytes (optimal CaCl2 concentration ≥2.5 mM) and binding was reversed by the addition of EDTA. The rate of conversion of Factor X to Xa in mixtures containing Factor VIIa and monocytes was directly related to the quantity of Factor VIIa bound to the monocyte surface. Thus the monocyte binding sites appear to represent tissue factor. Competition experiments showed that Factor VII and VIIa bind to the same monocyte sites and further, that unlabeled Factor VII and VIIa have the same affinity for the binding sites as the 125I-labeled proteins.

INTRODUCTION

Human Factor VII is a single chain, vitamin K-dependent serine protease present in plasma that can be converted to a two-chain form, Factor VIIa, by a number of agents including thrombin, Factor Xa, and Factor XIa (1). This conversion is associated with a 24-40-fold increase in activity when measured in a one-stage coagulation assay (1-3).

Tissue factor is a lipoprotein cofactor that markedly enhances the proteolytic activity of Factors VII and VIIa upon Factors IX and X (4, 5). The activation of Factor X to Factor Xa by tissue factor and Factor VII or VIIa constitutes the classical "extrinsic" pathway of coagulation (6).

Tissue factor has been detected in many tissues and cultured cells including human fibroblasts (7, 8) and smooth muscle cells (9). Blood cells, however, lack significant tissue factor activity unless they are stimulated by a variety of agents—for example, endotoxin (10), immune complexes (11), or complement C5a (12). Although granulocytes or lymphocytes were initially felt to produce this tissue factor activity, more recent studies have established the monocyte as its source (13-15).

In most studies of production of tissue factor by monocytes, total activity is measured following disruption of the cells by freeze-thawing or sonication. However, at least a portion of the tissue factor activity has been found to be available on whole, intact cells following their stimulation. By analogy with the reported binding of Factor Xa to platelets (16) through its cofactor, Factor Va, it appeared possible that Factor VII or VIIa bind to tissue factor at the monocyte surface and there activate Factor X (and Factor IX).

Single chain, zymogen, Factor VII is unique among the vitamin K-dependent proteases in that it appears to contain inherent coagulation activity. Recently, Nemerson et al. (17) have presented a cogent, albeit indirect, argument for this fact using bovine Factor VII and VIIa. Further, using a kinetic system involving the activation of Factor IX, they have shown that Factor VII and Factor VIIa have similar, if not identical, affinities for purified tissue factor.

We now show that human Factor VII and VIIa bind to the surface of stimulated, intact monocytes with essentially the same affinities and furthermore, that the binding of Factor VII or VIIa to monocytes is associated with the proteolytic activation of Factor X.

METHODS

Materials. Sterile, pyrogen-free water was obtained from Travenol Laboratories, Inc. (Deerfield, IL); Ficoll (type 400), bovine serum albumin (crystallized and lyophilized), bovine Factor VII/X-deficient plasma, rabbit brain cephalin, Trizma base, and Hepes from Sigma Chemical Co. (St. Louis, MO); polystyrene culture T-flasks from Corning Medical (Medfield, MA) and sodium Hypaque (50%) from Winthrop Laboratories, New York. Sodium [125I] iodide, carrier free, was purchased from New England Nuclear (Boston, MA). All other chemicals were reagent grade or better products of Sigma Chemical Co., Fisher Scientific Co. (Pittsburgh, PA), or J. T. Baker Chemical Co. (Phillipsburg, NJ).

Solutions. Endotoxin-free water was used in all solutions,
which were sterilized by passage through 0.2 or 0.45-μm filters (Nalgene Co., Nalgene Labware Div., Rochester, NY).

Ficol-Hypaque: Ficol, 57.34 g was dissolved in 180 ml of sodium Hypaque (50%) and 725 ml of water.

Phosphate-buffered saline (PBS): 1 liter contained 0.2 g KCl, 0.2 g KH₂PO₄, 8.0 g NaCl, 2.16 g Na₂HPO₄·7H₂O, and 1.0 g glucose, pH 7.4.

PBS/HSA/EDTA: PBS with 1 mM EDTA, pH 7.4.

DMEM: powered Dulbecco's modified essential medium with l-glutamine (KC Biological, Inc., Lenexa, KS) was reconstituted with water and NaHCO₃ according to manufacturer's instructions. Additional glucose and 1.0 M HEPES were added so that the final concentrations were 4 g/liter glucose and 20 mM Hepes, pH 7.4. Penicillin and streptomycin (Grand Island Biological Co., Grand Island, NY) were added to final concentrations of 50 U/ml and 50 μg/ml, respectively.

HRB: Hepes resuspension buffer contained 0.15 M NaCl, 0.02 M Hepes, pH 7.4, 1 g/liter glucose and 2 g/liter bovine serum albumin (American Red Cross Blood Services).

DMEM: powered Dulbecco's modified essential medium with l-glutamine (KC Biological, Inc., Lenexa, KS) was reconstituted with water and NaHCO₃ according to manufacturer's instructions. Additional glucose and 1.0 M Hepes were added so that the final concentrations were 4 g/liter glucose and 20 mM Hepes, pH 7.4. Penicillin and streptomycin (Grand Island Biological Co., Grand Island, NY) were added to final concentrations of 50 U/ml and 50 μg/ml, respectively.

Procedures

Purification of cells. Human whole blood was obtained from donors following their informed consent. 480 ml of blood were drawn into plastic syringes containing EDTA at a final concentration of 5 mM (1.5 ml of 0.2 M EDTA-NaOH, pH 7.5, per 60 ml syringe). Mononuclear cells and platelets were obtained by centrifugation over Ficol-Hypaque (18), resuspended in 300 ml of cold PBS/EDTA and centrifuged at 500 g in a Beckman J6-B centrifuge (Beckman Instruments, Inc., Spinclo Div., Palo Alto, CA) for 10 min. The cell pellets were resuspended in 50 ml cold PBS/HSA/EDTA and washed again by centrifugation. This pellet was resuspended in 100 ml of PBS/HSA/EDTA and placed on ice for elutriation.

Countercurrent elutriation was performed using a Beckman J21-B high-speed centrifuge equipped with a JE-6 elutriator rotor and strobe unit. A Cole-Parmer model 7013 Masterflex pump (Cole-Parmer Instrument Co., Chicago, IL) was used to control flow through the system which was monitored by a flow meter (model B-5684, Roger Gilmont Instruments, Inc., Great Neck, NY). Before each procedure, the entire system was washed extensively with ethyl alcohol (95%), followed by sterile water, and finally cold PBS/HSA/EDTA. The centrifuge was kept at 4°C and 400 g throughout.

By adjusting flow rates appropriately, lymphocytes and monocytes can be obtained using this elutriation system (19, 20). For the studies reported here, however, the goal was purification of monocytes alone, and an abbreviated elutriation schedule was used. The cell sample was loaded into the elutriation chamber at a flow rate of 9.4 ml/min, followed by 250 ml of PBS/HSA/EDTA at 9.4–10.5 ml/min, and finally 150 ml of buffer at 16 ml/min. The cells elutriated at this final flow rate were collected by centrifugation and resuspended in DMEM. Minor changes in the flow rate during the washing stage were made depending upon the level of the monocytes that could be directly visualized in the elutriation chamber.

For monocyte culture, T-flasks (150 cm²) were coated with 10 ml of fetal calf serum for 1–7 days at 4°C. 5 ml of the fetal calf serum was then removed and 45 ml of cells in DMEM and lipopolysaccharide (LPS) added such that the final concentration of cells was 1 × 10⁶/ml and fetal calf serum 10%. The T-flasks were incubated at 37°C under CO₂ (10%) overnight (16–18 h). The cells were then eluted from the flasks by the addition of 0.2 M EDTA-NaOH, pH 7.5 to a concentration of 5 mM and further incubated at 37°C for 60–90 min (without agitation). The medium containing the eluted cells was poured from the T-flasks and the cells collected by centrifugation, resuspended in 50 ml of cold PBS/HSA/EDTA, again collected by centrifugation, and finally resuspended in DMEM at 1 × 10⁶ monocytes/ml and placed on ice.

The final preparations of cells used for these experiments contained 90–96% monocytes, 1–8% lymphocytes, 0–4% basophils, <1% erythrocytes, <1% neutrophils, and <1% platelets. 75–85% of the cultured cells were eluted from the flasks using EDTA. Final yields of monocytes from 480 ml whole blood ranged from 7 to 20 × 10⁵ cells and were >97% viable by trypan blue exclusion.

Identification of cells. Cells were identified by morphology and, initially, nonspecific esterase staining (alpha naphthylbutyrate or Sigma Histozyme kit 90-A1). Subsequently, neutral red was used, as it was faster, more convenient, and gave results very comparable (±2%) to nonspecific esterase staining (21, 22). 200 cells or more were examined to obtain a differential count.

LPS. LPS was a phenol extract of Escherichia coli 026:B6 or 055:B5 (Sigma Chemical Co.). Following resuspension in 0.15 M NaCl at a concentration of 5 mg/ml, the suspension was sterilized by passage through a 0.45-μm filter and stored at 4°C. The concentration of LPS in the stock suspension was assumed to remain 5 mg/ml following filtration. The concentrations of LPS added to the cultures were chosen to stimulate maximal monocyte tissue factor production (10–40 μg/ml).

Proteins. Factor VII was purified using a modification of a previously published method (8). Its molecular weight was assumed to be 48,000 (1) and an extinction coefficient (ε₁cm at 280 nm) of 11.5 was estimated by the method of Babul and Stellwagen (23).

Factor X was purified essentially as previously described (24). Bovine Factor XIIa was a gift from Dr. K. Karachi and Dr. E. Davie (University of Washington, Seattle) and the Factor X coagulant protein from Russell’s viper venom was a gift from K. Levine and Dr. C. Jackson (Washington University, St. Louis).

Crude human brain thromboplastin was prepared as previously described (1).

Identification of Factor VII. Factor VII was labeled with [125I]iodide using a modification of the method of Praker and Speck (25). 10 μg of 1,3,4,6-tetrachloro-3a,6a-diphenylglycoluril (Iodo-gen, Pierce Chemical Co., Rockford, IL) in 100 μl chloroform was dried in the bottom of a 12 × 75-mm polypropylene tube. 100–150 μl of Factor VII, 1 mg/ml, in 0.15 M NaCl, 0.02 M sodium phosphate, 0.005 M benzamidine, 0.001 M EDTA, pH 7.0, was placed in the tube and 2.3–3.0 mCl (4–6 μl) of carrier-free sodium [125I]iodide added. Following incubation for 15 min on ice with occasional agitation, the solution was removed from the tube and dialyzed two times against 1 liter of 0.15 M NaCl, 0.1 M KI, 0.02 M benzamidine, 0.02 M Tris-HCl, pH 7.5 and then two times against 1 liter 0.15 NaCl, 0.02 Tris-HCl, pH 7.5.

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Abbreviations used in this paper: DMEM, Dulbecco’s modified essential medium; HRB, Hepes resuspension buffer; HSA, human serum albumin; LPS, lipopolysaccharide; PBS, phosphate-buffered saline.
A portion of the radiolabeled Factor VII was converted to two-chain Factor Vlla by incubation (100 µg/ml) with 2% (wt/wt) of bovine Factor XIIa in 0.15 M NaCl, 0.02 M Hepes, pH 7.4, with 1 mg/ml bovine serum albumin at 4°C for 48 h. The iodinated single- and two-chain proteins were stored at a concentration of 1 µg/ml in HRB at −70°C.

In four iodination procedures the specific activity of the labeled Factor VII or Vlla ranged from 17,760 to 26,400 dpm/ng. In each instance, >99.5% of the radioactivity was precipitated by 10% TCA. Functional, one-stage coagulation assays of Factor VII, pre- and immediately postiodination showed no change in activity (2 units/µg±10%). Activation of labeled Factor VII to Vlla was associated with a 28-fold increase in activity. When the proteins were stored at a concentration of 1 µg/ml there was a gradual loss of functional activity, such that by 12 wk Factor VII had 50% and Factor Vlla 20% of its original activity when compared with unlabeled standards. If the labeled proteins were stored at higher concentrations, inactivation occurred considerably more rapidly. For the experiments presented here, the iodinated Factor VII and Vlla were used within 4 wk of the labeling procedure. Fig. 1 shows the activation of iodinated Factor VII to Vlla by Factor XIIa.

Binding assays. Although a centrifugation through oil technique, similar to that previously described for platelets (16) was initially planned to assay binding of Factor VII and Vlla to monocytes, it was found to be unsuitable. When these stimulated monocytes were spun through a mixture of N-butyldiphenylamine and ethanol (9:1, density 1.028) it was evident both visually and by following specific radiolabeled Factor VII and Vlla binding that only a portion of the cells pelleted to the bottom of the tube as compared with cells centrifuged in the absence of oil. This problem was most marked at low monocyte concentrations (<2 x 10^6/ml) and could be corrected by the inclusion of 1–2 x 10^5/ml platelets (but not lymphocytes or erythrocytes). Therefore, the two-step centrifugation procedure described below was used in all experiments.

200 µl of monocytes (1 x 10^7/ml) were added to 12 x 75 mm-polypropylene tubes containing 300 µl of HRB, 125I-Factor VII or Vlla and CaCl2. Reaction mixtures routinely contained 4 mM CaCl2 and various concentrations of 125I-Factor VII or VIIa. The mixtures were incubated 60 min at 0°, 20°, or 37°C and, following gentle resuspension, a 400-µl portion was transferred to 1.5-ml polypropylene tubes and centrifuged for 2 min in a microfuge (Brinkmann Instruments, Inc., Westbury, NY) at ambient temperature. There was no detectable loss of cell viability during the incubation procedure itself as determined by trypan blue exclusion. 800 µl of N-butyldiphenylamine was layered over the sample and the tube was centrifuged an additional 2 min. A 200-µl portion of the supernatant (now above the oil layer) was removed for determination of “free” Factor VII or Vlla and the remainder of the supernatant and N-butyldiphenylamine aspirated. The bottom of the tube containing the cell pellet was then cut off and counted for “bound” Factor VII or Vlla.

Nonspecific binding, as determined by the inclusion of 2.4 µg/ml unlabeled Factor VII or Vlla in the reaction mixtures, was a constant fraction of total radioactivity added, and varied from 2.0 to 3.5%. Specific binding was assumed to be the difference between the observed and nonspecifically bound radioactivity. Radioactivity was counted on a Searle model 1190 gamma counter (Searle Radiographics, Inc., Des Plaines, IL) with a counting efficiency of 50%.

Coagulation assays. One-stage coagulation assays for Factor X and Factor VII or Vlla were performed as previously described (1, 16). To determine the rate of Factor X activation in thebinding mixtures, a 200-µl sample was removed, placed in a 12 x 75-mm polypropylene tube and 10 µl of Factor X (final concentration 8 µg/ml) was added. At 30-s intervals thereafter, 10-µl samples were removed, diluted in 0.15 M NaCl, 0.02 M Tris, pH 7.5, 0.001 M EDTA, and 1 mg/ml bovine serum albumin (at least 50-fold) and assayed as previously described for Factor Xa activity (16). A standard curve was constructed using Factor Xa produced from purified Factor X using 1% (wt/wt) Factor X coagulant protein from Russell’s viper venom.

The tissue factor assay was performed in a fibrometer (Becton, Dickinson & Co., Rutherford, NJ) at 37°C. 60 µl of Factor VIII-deficient plasma (George King Biomedical, Inc., Overland Park, KS) 60 µl of rabbit brain cephalin (1–9 dilution of stock suspension) and 60 µl of the sample to be assayed diluted appropriately in HRB with 0.005 M sodium citrate, were incubated 30 s at 37°C. The reaction was then started by addition of 60 µl of 25 mM CaCl2 at 37°C. A standard curve was constructed using crude human brain thromboplastin and gave a linear relationship between clotting times and thromboplastin dilution on a log–log plot. The activity of the stock human brain thromboplastin was arbitrarily given a value of 10,000 units/ml (a concentration of 200 units/ml in the assay gave a clotting time of 19.5 s and a concentration of 1 unit/ml, a clotting time of 96 s). The coagulant activity that developed in the monocytes following overnight incubation with LPS required Factor VII for expression.

To measure total tissue factor activity of the monocytes, the cells were lysed by sonication (15 short, 2–3-s bursts of sonication at a setting of 2.5 on a Branson model 350 sonifier, Branson Sonic Power Co., Danbury, CT, with the microtip probe) on ice.

**Polyacrylamide gels.** Sodium dodecyl sulfate (SDS) polyacrylamide gel electrophoresis was performed according to the method of Laemmli (26) with a 10% (ratio acrylamide/
RESULTS

In preliminary experiments, no specific binding of Factor VII or VIIa could be detected to monocytes directly following their isolation from blood, or to cells incubated overnight in the absence of LPS. By 4 h following LPS stimulation, specific binding could be detected, but for convenience, the studies that follow were performed after overnight incubation of the monocytes. With LPS stimulation, the tissue factor activity of whole cells increased from <1 to 35.0 to 62.0 units/1 × 10⁶ monocytes by 16–18 h incubation. Assay of the cells following sonication showed total tissue factor activity to be 50–100 units/1 × 10⁶ monocytes. In each instance, when the monocytes were cultured (Methods), the tissue factor activity available on whole cells was 40–100% of the total activity found following cell disruption. No specific Factor VII or VIIa binding to platelets or lymphocytes could be detected, either before or following their stimulation by LPS.

Incubations of monocytes with radiolabeled Factor VII or VIIa were routinely performed at 0°C. This temperature was chosen to prevent further monocyte tissue factor production, inhibit possible confounding effects of cellular endocytosis, and inhibit adhesion of the monocytes to the plastic test tubes (see below and Discussion).

Characteristics of Factor VII and VIIa binding to monocytes. Binding of Factor VII and VIIa to monocytes is time dependent, with maximal binding being obtained by 60 min, even at concentrations of Factor VII or VIIa as low as 0.4 ng/ml (data not shown). At 0°C the binding of zymogen, single-chain Factor VII to monocytes is very similar to that of activated, two-chain Factor VIIa (Fig. 2). That the binding observed using Factor VII was not due to its conversion to Factor VIIa during the incubation period itself was shown by SDS-polyacrylamide gel electrophoresis (under reducing conditions) and autoradiography of cell pellets during the time course experiment (Fig. 3). Further, when the activation of Factor X to Xa was determined in samples containing Factor VII, a delay in Factor X activation was observed, which presumably represented the feedback activation of Factor VII by Factor Xa (Fig. 4).

When similar experiments were performed at room temperature and 37°C, binding in mixtures containing Factor VII and Factor VIIa was again very similar, but SDS-polyacrylamide gel electrophoresis of the cell pellets showed considerable conversion of Factor VII to Factor VIIa during the 1-h incubation period. Whether this activation was related to a contaminating plasma protease, or one associated with the monocytes themselves, has not yet been determined.

FIGURE 2. Time course of Factors VII and VIIa binding to monocytes. Mixtures containing 4 × 10⁶/ml monocytes, 6 ng/ml ^125I-Factor VII or VIIa and 4 mM CaCl₂ were incubated at 0°C along with duplicate mixtures to which unlabeled Factor VII or VIIa (2.4 μg/ml), respectively, had also been added. Specific binding of Factor VII (○) and Factor VIIa (●) is shown. Nonspecific binding in the Factor VII and VIIa mixtures was 16 and 19%, respectively, of the total maximal binding.

FIGURE 3. SDS polyacrylamide gel electrophoresis and autoradiography of bound Factor VII. Samples (400 μl) from the ^125I-Factor VII reaction mixture described in Fig. 2 were removed at 5, 10, 15, 20, 40, and 60 min (left to right in figure) and centrifuged for 2 min in a microfuge. The cells were resuspended in 50 μl of 0.15 M NaCl, 0.010 M EDTA, 0.05 M Tris, pH 8.1, and 0.02 M benzamidine, and centrifuged once again. A sample of each supernatant was reduced with 5% β-mercaptoethanol, and autoradiography performed following SDS polyacrylamide electrophoresis.

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The binding of both $^{125}$I-Factor VII and VIIa could be reversed by the addition of either unlabeled Factor VII or Factor VIIa or by the chelation of calcium with EDTA. Fig. 5 shows the effect of adding a 100-fold excess of unlabeled Factor VIIa or EDTA (final concentration 5 mM) upon the binding of $^{125}$I-Factor VIIa. It is evident from the figure, that a portion of the nonspecific binding (that present in an excess of unlabeled Factor VIIa—see Methods) is calcium dependent. The reversal of binding by EDTA also indicates that endogenous Factor VII or VIIa, which may have bound to the monocytes during their preparation, was removed during the subsequent EDTA washing steps employed before the use of the monocytes in these binding experiments.

Fig. 6 shows the effect of CaCl$_2$ upon Factor VIIa binding. Binding increased as the CaCl$_2$ concentration approached 2.5 mM and then remained constant. Experiments using Factor VII yielded the same results. Additional experiments showed that the amount of specific Factor VII or VIIa binding assayed in the reaction mixtures was directly related to the concentration of monocytes (tested in the range from 1 to 8 x 10$^6$/ml).

**Equilibrium binding of Factor VII and VIIa to monocytes.** The concentration dependence of $^{125}$I-Factor VII and VIIa binding to monocytes at 0°C is shown in Figs. 7 and 8. In seven similar experiments the dissociation constant ($K_d$) and amount bound per 4 x 10$^6$ monocytes for Factor VII was 3.96±1.06 ng/ml (82.4 pM) and 1.14±0.22 ng (3,500 sites/monocyte) respectively (mean±SD). In 10 experiments the $K_d$ for Factor VIIa was 3.89±0.85 ng/ml (81 pM) and the amount bound per 4 x 10$^6$ monocytes was 1.19±0.18 ng (3,700 sites/monocyte). To independently verify the $K_d$ obtained from the equilibrium binding experiments the second order association constant ($K_2$) and first order dissociation constant ($K_1$) were determined for Factor VIIa from the data presented in Fig. 5. The $K_d$ calculated from these kinetic data ($K_d = K_2/ K_1$) was 88 pM, very similar to that obtained by equilibrium binding.

When the equilibrium binding experiments were performed at 37°C, however, double-reciprocal evaluation of the data consistently revealed curvilinear plots (concave upwards) (Figs. 7 and 8). This nonlinearity prevented calculation of dissociation constants, but the apparent saturation binding was always severalfold greater than that noted when the same cells were incubated at 0°C. Intermediate results were obtained when the experiments were performed at 20°C. This effect is being investigated further (Discussion).

To determine if the binding of Factor VIIa to monocytes was associated with its ability to activate Factor X, two experiments were performed. In the first experiment, a direct relationship between the rate of Factor X activation in the reaction mixtures and the quantity of Factor VIIa bound was demonstrated (Fig. 9). In the second experiment, the velocity of Factor
Calcium concentration of trations. After 60 min, monocytes, 6 ng/ml $^{125}$I-Factor VIIa and varying concentrations of CaCl$_2$ were incubated at 0°C. Duplicate mixtures containing additional unlabeled Factor VIIa (final concentration 2.4 µg/ml) were used to determine nonspecific binding. After 60 min, specific binding was determined per 4 x 10$^6$ monocytes.

X activation in the whole reaction mixture (4 x 10$^6$/ml monocytes, 6 ng/ml $^{125}$I-Factor VIIa, 4 mM CaCl$_2$ at 20°C) was compared with the velocities obtained in supernatants of the mixture separated from the cells by centrifugation at 12,000 g for 2 min in a microfuge (as in the binding studies), or at high speed (105,000 g for 30 min). The rate of Factor X activation in the supernatant following microfuge centrifugation was 4% of that of the whole mixture. Essentially no Factor X activation could be detected in the 105,000 g supernatant of the same reaction mixture. The converting activity present in the 12,000 g x 2 min supernatant was apparently “particle” bound as it could be removed by a centrifugation speed commonly used to sediment membranes. Although this activity could represent active monocyte secretion or membrane shedding (27), it appears as likely that it is related to mechanical manipulation of the monocytes during the in vitro incubation itself. The percentage of the total Factor Xa-converting activity, which remained in the microfuge supernatants, could be increased by raising the incubation temperature to 37°C, or by repeated, vigorous agitation of the mixtures.

### DISCUSSION

Before beginning the Factor VII and VIIa binding studies reported here, a number of experiments were performed to determine the optimal conditions for tissue factor production in monocyte cultures (data not shown). In agreement with previous reports, our preliminary experiments showed that monocytes, but not lymphocytes, neutrophils, or platelets produced tissue factor activity following stimulation with LPS. In contrast to other studies (14, 15, 28, 29), however, no dose-dependent enhancement of monocyte tissue factor production was found when elutriator-purified lymphocytes were added to the monocyte cultures. As our monocyte preparations contained small numbers of lymphocytes (1–8%), no comment can be made upon the claim that lymphocytes are absolutely required for stimulation of monocyte tissue factor production (15, 30). The addition of platelets to the monocyte cultures did enhance the amount of tissue factor activity produced (31). This effect appeared to be substantial when...
a one-stage coagulation assay was used to determine tissue factor activity, but only modest (twofold in cultures containing $1 \times 10^8$ platelets and $5 \times 10^5$ monocytes per ml) when a more specific, two stage, chromogenic assay, using purified Factor VIIa and Factor X was used.\(^3\) Because the presence of platelets in the monocytic cultures made subsequent elution of the monocytes considerably more difficult, cultures of monocytes alone were used for the binding studies.

With the use of indirect functional assays, the apparent calcium-dependent binding of Factor VII to crude tissue factor preparations (5, 32) and a human monocytic tumor cell line (33) has been reported previously. In fact, radioactively labeled bovine Factor VII has been used as a probe to localize tissue factor in tissue slices (34). Since the binding of Factor VIIa to the monocyte surface reported here was directly related to the velocity of subsequent Factor X activation, in all probability (essentially by definition), the Factor VII and VIIa binding site represents tissue factor.

Equilibrium binding studies consistently yielded linear double-reciprocal plots when they were performed at 0°C. Curvilinear plots, however, were obtained when the same experiments were performed at 37°C.

\(^3\) The two stage, chromogenic assay for tissue factor activity was performed by incubating the sample to be tested with 100 ng/ml purified Factor VIIa, 6 μg/ml purified Factor X, and 5 mM CaCl₂ at 37°C. After 1 min, the reaction was stopped by the addition of EDTA and a sample was diluted and assayed for Factor Xa activity with the chromogenic substrate S-2222 (Ortho Pharmaceutical, Raritan, NJ).

This phenomenon is being investigated further and preliminary experiments suggest it is not related to cellular endocytosis. The monocytes are metabolically active at 37°C and although "positive cooperativity" or possible "up regulation" of binding sites during the 60-min monocyte incubation period might be invoked to explain this phenomenon, a number of experiments are necessary to insure truly equilibrium conditions, to exclude ligand or receptor site degradation, and to rule out an artifactual effect of the in vitro binding assay itself. These studies are now under way.

The apparent equal affinity ($K_d \approx 82$ pM) of both

![Equilibrium binding of \(^{125}\)I Factor VII and VIIa to monocytes. Double reciprocal plot of the data presented in Fig. 7.](http://www.jci.org)
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Figure 10 Inhibition of 125I-Factor VII and VIIa binding by unlabeled Factor VII or VIIa. Mixtures containing 4 × 10^6/ml monocytes, 4 mM CaCl_2, 6 ng/ml 125I-Factor VII and increasing concentrations of unlabeled Factor VII (●) or Factor VIIa (○), and similar mixtures containing 6 ng/ml 125I-Factor VIIa plus increasing concentrations of unlabeled Factor VII (▲) or Factor VIIa (△) were incubated at 0°C. At 60 min, specific binding was determined and plotted as a percentage of that bound in the absence of added, unlabeled Factor VII and VIIa. (100% binding of 125I-Factor VII and VIIa was 0.52 and 0.56 ng, respectively, per 4 × 10^6 monocytes). Binding in mixtures containing 2.4 μg/ml unlabeled Factor VII or VIIa was assumed to represent non-specific binding.

Factor VII and VIIa for their monocyte binding sites (presumably tissue factor) is unique among the known coagulation factors. Whether human Factor VII possesses intrinsic coagulation activity is unclear, as purified preparations undoubtedly contain traces of Factor VIIa, and activation of Factor VII may occur during the assay procedure itself. This problem has been studied extensively in the bovine system by Nemerson and his colleagues (35–37), where it is apparent that bovine zymogen Factor VII possesses at most <1% of the catalytic activity of activated Factor VII (17). Since zymogen Factor VII has only a fraction of the catalytic activity of Factor VIIa, but competes equally with Factor VIIa for monocyte binding (tissue factor), it would be predicted that the zymogen would function as an inhibitor of Factor X (or Factor IX) activation. Nemerson et al. (17) have recently described such zymogen inhibition in an in vitro system using bovine Factors VII and VIIa, purified tissue factor, and measurement of the velocity of Factor IX activation. Furthermore, these investigators found that diisopropyl fluorophosphate inactivated bovine Factor VII and VIIa, inhibited the activation of Factor IX by the Factor VIIa-tissue factor complex with very similar K_{1/2} suggesting that they have essentially equal affinities for tissue factor (17). Thus, the direct binding experiments of human Factors VII and VIIa presented here are in agreement with kinetic experiments presented for the bovine factors.

If these in vitro studies can be applied to in vivo hemostasis, it would be anticipated that in areas where relatively low numbers of binding sites (i.e., less than the amount of tissue factor required to saturate the total Factor VII and VIIa) became available, single-chain Factor VII would dampen subsequent coagulation as its plasma level is >400 ng/ml (1–3), 100-fold above its K_I for inhibition Factor VIIa binding to tissue factor. At larger wounds, however, where it is conceivable that sufficient tissue factor binding sites might be exposed to saturate the available Factor VII and VIIa, coagulation would proceed rapidly.

A number of proteases have been shown to activate Factor VII in purified, and in vitro plasma systems (38–42). The kinetically most efficient of these appears to be Factor Xa, at least when purified reagents are used (38). Whether this is also the case in vivo, and further, the extent of Factor VII activation that occurs in vivo, is not clear. It is of interest, however, that when plasma is induced to clot in vitro by small amounts of tissue factor, Factors IX and VIII are required for optimal Factor X activation and thrombin generation (43–45). Along these same lines, the possible role of activated Factor VII as the Factor VIII bypassing activity in prothrombin complex concentrates has been suggested previously and is now being investigated further (46, 47).

When the level of zymogen Factor VII is low, or the amount of Factor VII to Factor VIIa decreased, one would expect the dampening effect of Factor VII to be diminished. It is tempting to speculate that the loss of this inhibitory effect of zymogen Factor VII might contribute to the thrombotic complications associated with congenital Factor VII deficiency (48–50), loading dose coumarin therapy (51, 52), and infusion of prothrombin complex concentrates, which may contain high levels of activated Factor VII (46, 53–62).

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References


Aronson, D. Personal communication.


