Sequence of Fibrinogen Proteolysis and Platelet Release after Intrauterine Infusion of Hypertonic Saline

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Abstract Plasma fibrinopeptide B (Bβ1-14 or FPB) immunoreactivity was studied by radioimmunoassay in patients who received intrauterine infusion of hypertonic saline to terminate pregnancy. FPB immunoreactivity increased with thrombin treatment (TIFPB) suggesting the presence of a larger FPB-containing peptide, since purified FPB is not altered by thrombin, whereas thrombin increases the immunoreactivity of Bβ1-42 (which includes FPB) 10-fold. TIFPB immunoreactivity in plasma, drawn 4 h after hypertonic saline infusion eluted from Sephadex G-50 similarly to isolated Bβ1-42. Streptokinase, incubated with normal plasma progressively generated TIFPB immunoreactivity, which showed a major component which eluted from Sephadex G-50 similarly to Bβ1-42. Streptokinase generated TIFPB much more rapidly in reptilase-treated plasma that contains fibrin I, (which still includes FPB), indicating that fibrin I is prefered over fibrinogen as a substrate for plasmin cleavage of arginine (Bβ42)–alanine (Bβ43). Serial studies were then made in 10 patients receiving intrauterine hypertonic saline. Fibrinopeptide A (FPA) levels rose immediately, reached a peak between 1 and 2 h, were declining at 4 h, and were normal at 24 and 48 h. TIFPB levels rose slightly in the 1st h, reached a peak at 4 h, and had returned to base-line values at 24 h. Serum fibrinogen degradation product levels were unchanged at 1 h, reached their highest level at 4 h, and were still markedly elevated at 24 and 48 h. Fibrinogen levels dropped slightly being lowest at 4 and 24 h. Platelet counts declined in parallel with the fibrinogen levels over the first 4 h, but continued to decrease through 48 h. Beta thromboglobulin (βTG) levels generally paralleled FPA levels whereas platelet factor 4 (PF4) levels showed only slight changes. The data indicate that immediately after intrauterine hypertonic saline infusion thrombin is formed that cleaves FPB from fibrinogen to produce fibrin I and releases βTG and PF4 from platelets. Later plasmin cleaves Bβ1-42 from fibrin I to produce fragment X, which is further degraded to form serum fibrinogen degradation products. This sequence of proteolysis indicates that plasmin action on fibrin I serves as a mechanism that regulates fibrin II formation by removing the Bβ chain cleavage site, which is required for thrombin action in converting fibrin I to fibrin II.

INTRODUCTION

The proteolysis of fibrinogen by thrombin and plasmin and the activation of platelets are important in normal hemostasis, in thrombosis and in disseminated intravascular coagulation. Recently assays have been developed that reflect fibrinogen proteolysis and platelet activation and allow study of these processes in vivo. Proteolysis of fibrinogen by thrombin occurs in two stages (1–3). In the first step thrombin cleaves fibrinogen to produce fibrinopeptide A (FPA)¹ and fibrin I (i.e., fibrinogen lacking FPA but still containing fibrinopeptide B [FPB]) and in the second step it releases FPB (Bβ1-14) from fibrin I to produce fibrin II. Early plasmin action removes two-thirds of the COOH-terminal end of the Aα-chain and also the NH₂-terminal end of the Bβ chain of fibrinogen. Bβ1-42 is the initial specific product of Bβ chain proteolysis by plasmin (4–5). The remaining molecule is termed fragment X (6–8).² Specific cleavage products reflecting platelet

¹Abbreviations used in this paper: βTG, beta thromboglobulin; FDP, fibrinogen degradation products; FPA, fibrinopeptide A; FPB, fibrinopeptide B; PF4, plasma factor 4; TIFPB, thrombin-increasable FPB.

²The term fragment X was originally used to designate a large molecular weight product of early plasmin proteolysis of fibrinogen. The NH₂-terminal portion of the Aα chain is still intact. When fibrin I or fibrin II is the substrate for plasmin action a similar fragment is produced that lacks FPA and may be noncovalently bound in large molecular size polymers.

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activation are not known but two proteins specific to the platelet are secreted from the alpha granules in the energy-dependent platelet release reaction (9-11). These proteins are platelet factor 4 (PF4) and beta thromboglobulin (βTG). Specific assays applicable to clinical blood samples have been developed for FPA (12), PF4 (13-15), and βTG (15, 16). An assay for FPB (3) in vitro has also been developed and this paper describes the use of this assay to identify a plasma-produced peptide, which includes FPB and is derived from the NH2-terminal end of the ββ chain.

Serial changes in the plasma concentration of these peptides were studied in patients during the termination of pregnancy by the intravascular infusion of hypertonic saline. This widely used method for interrupting pregnancy has been documented to be associated with transient in vivo coagulation as reflected by reduction in fibrinogen, clotting factor levels, and platelet counts, and by elevation of serum fibrinogen degradation product (FDP) levels (17-20). The application of assays for FPA (reflecting thrombin action on fibrinogen), for NH2-terminal ββ chain peptide (reflecting plasmin action) and for PF4 and βTG (reflecting platelet release) in this syndrome is described in this paper.

METHODS

Reagents. Native FPA was prepared from fibrinogen (grade L, 95% clottable, from A. B. Kabi, Stockholm) as described by Blomback et al. (21). Native FPB and Bβ1-42 were gifts of Dr. Birger Blombäck, Department of Blood Coagulation Research, Karolinska Institutet, Stockholm, Sweden or were isolated in this laboratory. Specific antisera and radiolabeled tyrosinated FPA (22), FPB analogue (3), and PF4 and βTG (15) were prepared as described. Human thrombin was the gift of Dr. John Fenton, New York State Public Health Laboratories, Albany, N. Y. Streptokinase was streptodornase-streptokinase from Lederle Laboratories Div., American Cyanamid Co., Pearl River, N. Y. Heparin for in vitro use was obtained from Hynson, Westcott & Dunning, Inc., Baltimore, Md. Ovalbumin was obtained from ICN Nutritional Biochemicals, Cleveland, Ohio, and charcoal (Norit A) from Amend Drug and Chemical Co., Irvington, N. J. Anhydrous denatured ethanol (Fisher Scientific Co., Pittsburgh, Pa.) was used to precipitate fibrinogen from plasma samples. Bentonite was also obtained from Fisher Scientific Co. and protamine sulfate was obtained from Eli Lilly & Co., Indianapolis, Ind. Sephadex G-50 (superfine) was obtained from Pharmacia Fine Chemicals, Inc., Piscataway, N. J. adenosine from ICN Nutritional Biochemicals, and theophylline from Sigma Chemical Co., St. Louis, Mo. Reptilase was a purified fraction, the gift of Dr. Kurt Stocker, Pentapharm, Zurich, Switzerland.

Patients. The patients involved in this study had all been scheduled for abortion and had provided informed consent to participate in the study. The patients' ages ranged from 19 to 41 yr with a mean of 22 yr and the estimated duration of gestation ranged from 10 to 18 wk with a mean of 16 wk. 200 ml of sterile sodium chloride (20%) was injected into the uterine cavity over a 2- to 3-min period by the patient's obstetrician. All patients aborted within 48 h of injection of the saline. No patient experienced excess local or any generalized bleeding, fever, tachycardia, or hypotension.

Blood collection and processing. Blood samples were collected from the antecubital veins of patients via 19-gauge needles. To insure that blood flow was free the time between the initial entry of the blood into the syringe and its mixture with the anticoagulant solution was recorded with a stop watch. In all instances the time period was 45 s or less. Blood for FPA and FPB assays was mixed with one-tenth volume heparin (1,500 U/ml) and Trasylol (1,000 U/ml; FBA Pharmaceuticals, Inc., New York). For the FPA assays the blood was processed and assayed as described (12). The rate of FPA generation in vitro was measured as described (23). For the FPB assays the plasma was precipitated with ethanol as described for the FPA assay (12). The plasma was precipitated with ethanol within 30 min of blood collection and no tests were made on plasma samples that had been stored frozen. Prompt processing of plasma for the FPB assays is necessary because thrombin-increasable FPB (TIFPB) immunoreactivity is generated in vitro during storage in the frozen state even at -90°C despite the presence of protease inhibitors. After precipitation with ethanol and reconstitution with distilled water the plasma extracts were dialyzed for 48 h and the dialysates were tested for FPB immunoreactivity before and after treatment with thrombin (10 U/ml for 1 h at 37°C) and the results were corrected for incomplete recovery on dialysis to obtain the plasma values. In additional experiments the dialysis step was omitted. The reconstituted samples now contained ≈3 U/ml heparin, which was neutralized by adding protamine sulfate (final concentration 0.15 mg/ml). The extracts were tested for FPB immunoreactivity before and after treatment with thrombin. With this procedure recovery of Bβ1-42 added to blood in vitro averaged 90% (Table I). Blood for FPB and βTG assays was mixed with one-tenth volume 4% trisodium citrate that contained 0.01 M adenosine and 0.02 M theophylline (−0.01 M adenosine −0.02 M theophylline) and centrifuged for 10 min at 1,500 g at room temperature. Platelet-poor plasma was removed and spun at 43,000 g for 10 min at 4°C. The supernatant plasma contained <250 plates/µl and was stored frozen at -60°C till assay. Blood for fibrinogen and serum FDP measurements was mixed with one-tenth volume 3.8% trisodium citrate and centrifuged at 1,500 g for 20 min to deposit the cells. Serum was prepared by clotting the plasma with thrombin as described elsewhere (24).

Assays. FPA was assayed as described (12). FPB immunoreactivity was assayed as described (3) using similar assay volumes and preincubation as for the FPA assay. Native FPB was used as a standard preparation and antisem R22 was

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The results shown are the mean data from 3 replicate experiments.

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used throughout these studies. PF4 and βTG were assayed as described (15). A mol wt of 35,400 was used to calculate the molar concentration of βTG and of 30,800 to calculate that of PF4. Fibrinogen levels were measured as described by Ellis and Stransky (25). Serum FDP levels were measured by the tanned erythrocyte hemagglutination inhibition method as described by Merskey et al. (24), the normal level by this technique being up to 1 µg/ml. Platelets were counted by a model S plus Coulter Counter (Coulter Electronics Inc., Hialeah, Fla.).

Sephadex G-50 filtration of FPB, Bβ1-42, and extracts of plasma. 5 ml plasma (with heparin and Trasylol), collected from each patient 4 h after intrauterine injection of hypertonic saline, was dialyzed against 30 ml 0.05 M NH4HCO3 containing 50 µg/ml ovalbumin for 48 h. The dialysate was lyophilized and reconstituted in 1.5 ml 0.15 M Tris-buffered saline (0.1 M NaCl and 0.5 M Tris), pH 5, containing 100 µg ovalbumin. The sample was applied to a 95 × 1.2 cm column packed with Sephadex G-50 (superfine) that had been equilibrated with 0.15 M Tris-buffered saline pH 5.0, with 0.2 M epsilon aminocaproic acid and 0.01% ovalbumin. [14C]glucose was used as a marker and 1.1 ml fractions were collected. Extracts of plasma samples in which Bβ1-42 had been generated in vitro by incubation with streptokinase were also filtered over Sephadex G-50 as described for the patients' plasma. Native FPB and Bβ1-42 were filtered over identical but separately packed columns in the same way. The eluted fractions were tested for FPB immunoreactivity before and after treatment with 1 U thrombin/ml for 60 min at 37°C.

Generation of Bβ1-42 in vitro. Streptokinase (1 U/ml) was incubated at 37°C with 2 ml citrated plasma. At intervals aliquots of the plasma were removed, Trasylol (200 U/ml) was added and the samples were precipitated with 50% ethanol and processed as described (without dialysis) and assayed for FPB immunoreactivity with and without treatment by thrombin. The experiment was then repeated using citrated plasma that had first been incubated with reptilase (1.7 µg/ml) for 10 min at 27°C before incubation with streptokinase. The reptilase had produced a visible clot within 60 s.

RESULTS

Assay and recovery of Bβ1-42. Measurements of FPA, βTG, and PF4 levels in clinical blood samples have been described, but the identification of Bβ1-42 in plasma has not previously been reported. An assay specific for Bβ1-42 is not available, however, the molecule cross-reacts in the FPB assay. FPB and Bβ1-42 react differently in the FPB assay with antiserum R22 (3). The slope of the inhibition curve for Bβ1-42 is different from that of FPB and the immunoreactivity of Bβ1-42 increases 10-fold or more after treatment with thrombin, whereas that of FPB is unaltered (Fig. 1). Hence Bβ1-42 can be distinguished from FPB by an increase in immunoreactivity on treatment with thrombin. Fibrinogen also shows a marked increase in FPB immunoreactivity on treatment with thrombin, so before measuring Bβ1-42 in plasma it is necessary to remove all fibrinogen from the plasma without removing Bβ1-42. This can be done by precipitation of the plasma with ethanol. Precipitation of plasma with 50% ethanol removes detectable fibrinogen from the plasma without significant precipitation of Bβ1-42

FIGURE 1 Immunoreactivity of FPB, and Bβ1-42 with and without thrombin treatment. Inhibition of binding of 125I-labeled tyrosinated FPB analogue by: (A) FPB with or without treatment with thrombin 1 U/ml for 1 h at 37°C and (B) Bβ1-42 with or without treatment with thrombin 1 U/ml for 1 h at 37°C.

In vitro generation of TIFPB. When plasma was incubated with streptokinase, progressive generation of TIFPB immunoreactivity occurred and was much more rapid in reptilase-treated plasma in which fibrin I rather than fibrinogen was the substrate for plasmin (Fig. 2). In these experiments an ≈10-fold
increase in FPB immunoreactivity occurred after thrombin treatment suggesting the presence of FPB-containing material larger than FPB. In addition the slope of the inhibition curve produced by the non-

thrombin-treated samples was different from that of FPB. Gel filtration of streptokinase-treated plasma showed an elution volume for TIFPB immunoreactivity almost identical with that for Bβ1-42 (Fig. 3B and D). However, there was also a later eluting peak of FPB immunoreactivity, the upward slope of which also increased on treatment with thrombin and almost certainly represents Bβ1-21 (26). The downward slope probably represents FPB or desargulln FPB (Fig. 3A and D).

**TIFPB immunoreactivity in patient plasma extracts.** Extracts of plasma samples taken before and at intervals after intrauterine injection of saline showed a modest increase in FPB immunoreactivity which then decreased to baseline levels (Fig. 4). Thrombin treatment of the extracts increased FPB immunoreactivity markedly resembling the response of isolated Bβ1-42 (Fig. 4). The slope of the inhibition curve of non-thrombin-treated samples was different from that of FPB. The molecule responsible for the plasma immunoreactivity was further characterized by filtering an extract of the 4-h plasma sample from one patient over Sephadex G-50 and testing the FPB immunoreactivity of the effluent. Two peaks of immunoreactivity were found (Fig. 3C). The earlier eluting peak showed a major increase in immunoreactivity after treatment with thrombin and exhibited the same elution profile as Bβ1-42. The later peak eluted in the same position as FPB and did not increase in immunoreactivity when treated with thrombin. The plasma extract of a second patient examined in a similar manner gave virtually identical results (data not shown). These findings suggest that most, if not all, the TIFPB immunoreactivity in the patients' plasma represents Bβ1-42, however, the term TIFPB immunoreactivity is used because Bβ1-21 also shows a marked increase in immunoreactivity when treated with thrombin (data not shown).

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**FIGURE 3** Elution patterns from Sephadex G-50 of FPB, Bβ1-42, 4-h sample after intrauterine injection of hypertonic saline and plasma incubated with streptokinase in vitro. (A) Native FPB, (B) Bβ1-42, (C) extract of patient plasma drawn 4 h after intrauterine injection of hypertonic saline, and (D) extract of plasma incubated with streptokinase (SK) (20 U/ml) for 1 h at 37°C. 1.1 ml fractions were collected and tested for FPB immunoreactivity before (-----) and after treatment with thrombin (- - - - -).

**FIGURE 4** Plasma FPB immunoreactivity before and after intrauterine infusion of saline. (-----) before thrombin treatment of plasma extracts. (- - - - -) after thrombin treatment of plasma extracts. The data shown are the mean±SEM in 10 patients.
Serial studies of in vivo coagulation after saline infusion. Fibrinogen and plasma fibrinopeptide levels, as well as platelet counts and platelet protein concentrations, in the 10 subjects are shown in Fig. 5.

![Fibrinogen Proteolysis after Intrauterine Saline Infusion](image)

**FIGURE 5** Changes in plasma concentrations of FPA, TIFPB, fibrinogen, FDP, PF4, βTG, and platelet counts before and 1–48 h after intrauterine injection of hypertonic saline. (A) FPA and TIFPB levels. The TIFPB levels from Fig. 4 have been divided by 0.37 to correct for incomplete recovery on dialysis. (B) Plasma and serum FDP levels. (C) Platelet counts and plasma PF4 and βTG levels. The data are the mean ± SEM in 10 patients.

FPA levels (5A) were elevated almost 20-fold at 1 h and were at approximately the same level at 2 h. They had decreased significantly at 4 h, were only slightly elevated at 24 h, and were normal at 48 h. TIFPB levels (Fig. 5A) showed a slight increase in the 1st h and then rose more rapidly to reach a peak value at 4 h. At 24 h the levels were similar to the preinfusion levels and then were slightly lower than preinfusion levels at 48 h (Fig. 5A). Fibrinogen levels (Fig. 5B) decreased by a mean of 39 mg/dl (P = 0.007) with the lowest levels at 4 and 24 h. By 48 h preinfusion values had been attained. FDP levels (Fig. 5B) showed no detectable increase at 1 h but a major increase at 2 h with the peak level at 4 h. The FDP levels, although decreasing, were still significantly elevated at 24 and 48 h after saline infusion. Over the first 4 h the decrease in platelet counts (Fig. 5C) paralleled the decline in fibrinogen levels, but thereafter, instead of returning to the preinfusion level, platelet counts inexplicably continued to decline, reaching their lowest point at 48 h. βTG levels (Fig. 5C) rose sharply and had increased about threefold by 1 h (P = 0.0001) with a further increase at 2 h. βTG levels had decreased at 4 h, had reached preinfusion levels at 24 h, and were below initial levels at 48 h. PF4 levels (Fig. 5C) showed a similar pattern of changes, but these were much less marked than with βTG and were distinctly increased above the base-line values only at 1 (P = 0.021) and 2 (P = 0.01) h after infusion. The data shown in Fig. 5 represent mean values but the sequence of changes was the same in all patients studied.

To determine whether the increase in FPA levels preceded that of TIFPB levels an additional four patients were studied at intervals during the 1st h (Fig. 6). FPA levels were elevated immediately (within 3 min of the saline injection) and rose at an almost constant rate (about 0.3 pmol/ml per min) throughout the 1st h to reach a mean of 16 pmol/ml at 1 h. TIFPB levels were not altered immediately after the infusion of saline, but thereafter rose slightly over the 1st h. βTG levels showed some fluctuation in the first 30 min but increased in parallel with the FPA levels between 30 and 60 min.

In vitro FPA generation rates were measured in blood samples taken 1 h after injection of saline. In four patients there was no significant FPA generation (<1 pmol/ml) in vitro and in six patients increased FPA generation (1.1 to 7.0 pmol/ml) occurred (Table II).

**DISCUSSION**

Our results indicate that TIFPB immunoreactivity reflects the presence of material containing the FPB sequence but larger than the FPB (Fig. 1). When the TIFPB immunoreactivity in patient’s blood was further characterized by Sephadex G-50 chromatography it behaved similarly to isolated Bβ1-42 (Fig. 3). However,
the term TIFPB is preferred to Bβ1-42 because there is as yet no absolute way of distinguishing Bβ1-42 from Bβ1-21, which also manifests increased immunoreactivity when treated with thrombin. TIFPB immunoreactivity can theoretically be derived by plasmin cleavage of fibrinogen or fibrin I but not of fibrin II (which contains neither FPA nor FPB) because plasmin is known to cleave the bond between arginine (Bβ42) and alanine (Bβ43) is responsible for the TIFPB immunoreactivity found in patients' plasma although it is conceivable that leukocyte proteases (27) or other enzymes might also cleave FPA containing peptides under appropriate circumstances. TIFPB immunoreactivity was generated much more rapidly when streptokinase was incubated with reptilase-treated plasma than with control plasma (Fig. 2) indicating that fibrin I is preferred over fibrinogen as a substrate for plasmin. This phenomenon is an extension of the well-recognized observation that plasmin proteolyses fibrin more readily than fibrinogen (28). Recent work has shown that plasminogen binds to fibrin via lysine-binding sites. It has been suggested that plasmin binds to fibrin and not to fibrinogen and that the bound plasmin is protected from inhibition by the α2-plasmin inhibitor (29). This could explain why plasmin cleaves Bβ1-42 from fibrin I more readily than from fibrinogen.

Based on the above considerations, the pattern of changes shown in Figs. 5 and 6 is interpreted as follows. Within minutes of infusion of saline, thrombin is formed which cleaves fibrinogen to produce FPA and fibrin I and releases βTG and PF4 from platelets. Thrombin activity reaches its highest level between 1 and 2 h, has decreased at 4 h, and is normal at 24 and 48 h. Plasmin action starts after thrombin action with the production of TIFPB implying formation of fragment X (4–8). Because the data shown in Fig. 2 indicate that fibrin I is preferred to fibrinogen as a substrate for plasmin it is likely that the major source of TIFPB immunoreactivity is fibrin I. NH2-terminal Bβ chain cleavage (reflecting fragment X formation) precedes the formation of fragment Y, which in turn precedes the generation of fragments D and E (4–8) and it is the latter fragments (Y, D, and E), which are measured in the FDP assay. One would, therefore, anticipate that Bβ1-42 levels would rise earlier than FDP levels and the data shown in Fig. 5 suggest but do not prove this. More detailed studies over the 2nd h and more sensitive assays for FDP will be required. FDP levels declined more slowly than the TIFPB levels presumably both because of slower plasma clearance of the larger molecules and of continued lysis of fibrin.

The slow clearance of FDP results in the 24-h pattern in which maximum reduction in fibrinogen levels is present while FDP levels are still high. Low fibrinogen and high FDP levels form the principal laboratory criteria for disseminated intravascular coagulation (30, 31), yet at 24 h thrombin action as reflected by FPA levels was virtually normal. This distinction in the time sequence between thrombin action in vivo and the principal laboratory criteria for diagnosing disseminated intravascular coagulation may be part of the explanation for the contradictory results of heparin therapy. If a similar sequence of proteolyces occurs in clinical disseminated intravascular coagulation, one

### Table II

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<th>Patient</th>
<th>In vitro FPA generation rate (pmol/ml)</th>
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would anticipate that heparin would be useful only when thrombin action is significantly increased and not when it is normal.

The rapid in vitro FPA generation in the peripheral blood of 6 of 10 patients 1 h after injection of saline indicates the presence of thrombin activity in the blood sample (23) implying systemic dilution of thrombin or of activated factors that were initially formed in the uterus. Similarly increased thrombin activity was found in the peripheral blood of a patient with a localized hemangioma in whom prothrombin activation almost certainly starts in the hemangioma (23).

In vitro studies showed that addition of thrombin to citrated platelet-rich plasma or to whole blood resulted in FPA cleavage at thrombin concentrations 100-fold less than were required to release \( \beta TG \) and PF4 (15) but in spontaneously clotting blood FPA cleavage and \( \beta TG \) and PF4 release occurred in parallel (11). The data in the present study more closely resemble the pattern occurring in spontaneously clotting whole blood. The difference in the magnitude of the changes in the plasma concentrations of \( \beta TG \) and PF4 may be caused by differences in the plasma clearance rate of the two proteins. It is not clear why the platelet count continued to decline between 24 and 48 h despite normal \( \beta TG \) and PF4 levels and a rising fibrinogen level.

Aside from the practical implications concerning disseminated intravascular coagulation the data also suggest a theory relating the proteolysis of fibrinogen by thrombin and plasmin in vivo. The theory is illustrated in Fig. 7. Thrombin initially cleaves fibrinogen to produce free FPA and fibrin I, which polymerizes to form a visible clot. Thrombin then cleaves fibrin I to yield free FPB and fibrin II. If plasmin rather than thrombin cleaves the B8 chain on fibrin I, B81-42 and fragment X result. The data in Fig. 2 indicate that plasmin removes B81-42 much more rapidly from fibrin I than from fibrinogen. There are no data as yet concerning the influence of polymerization on the rate of plasmin cleavage of the arginine (B842)–alanine (B843) bond. However, there is evidence that polymerization is important for thrombin to release FPB from fibrin I (1). Fibrin II forms thicker strands than fibrin I (32) and may be more likely to result in occlusive thrombosis. The formation by plasmin of fragment X from fibrin I serves as a mechanism to regulate fibrin II formation. It is suggested that the changes in fibrinogen after intrauterine infusion of saline and in the clinical syndrome of disseminated intravascular coagulation represent the effects of a defense mechanism for coping with thrombin generated in the blood as a response to injury. Thrombin produces fibrin I, which is principally converted to fragment X, thus limiting fibrin II formation and consequent intravascular occlusion by thrombus.

**ACKNOWLEDGMENTS**

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