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Fibrinopeptides A and B release in the process of surface fibrin formation

Short title for the running head: fibrinopeptide release from surface fibrin

Tomas Riedel¹, Jiri Suttnar¹, Eduard Brynda², Milan Houska², Leonid Medved³, Jan E.Dyr¹

¹ Institute of Hematology and Blood Transfusion, Prague, Czech Republic
² Institute of Macromolecular Chemistry, Academy of Sciences of the Czech Republic, Prague, Czech Republic
³ Department of Biochemistry and Molecular Biology, University of Maryland School of Medicine, USA

Corresponding author’s full name:
Jan Evangelista Dyr, Institute of Hematology and Blood Transfusion, U nemocnice 1, 128 20 Prague 2, Czech Republic, dyr@uhkt.cz, phone and fax number: +420221977208

Contact information for all co-authors:
Tomas Riedel, Institute of Hematology and Blood Transfusion, U nemocnice 1, 128 20 Prague 2, Czech Republic, riedelt@post.cz, phone and fax number: +420221977208

Jiri Suttnar, Institute of Hematology and Blood Transfusion, U nemocnice 1, 128 20 Prague 2, Czech Republic, stt@uhkt.cz, phone and fax number: +420221977208

Eduard Brynda, Institute of Macromolecular Chemistry, Academy of Sciences of the Czech Republic, Heyrovského nam. 2, 162 06 Prague, Czech Republic, brynda@imc.cas.cz, phone and fax number: +420296809266

Milan Houska, Institute of Macromolecular Chemistry, Academy of Sciences of the Czech Republic, Heyrovského nam. 2, 162 06 Prague, Czech Republic, houska@imc.cas.cz, phone and fax number: +420296809234

Leonid Medved, Department of Biochemistry and Molecular Biology, University of Maryland School of Medicine, 108 N. Greene Street, Baltimore, MD 21201, USA, lmedved@som.umd.edu, phone: 410-706-3220, Fax: 410-706-8297

Fibrinopeptides A and B release in the process of surface fibrin formation

Short title for the running head: fibrinopeptide release from surface fibrin
Abstract

Fibrinogen adsorption on a surface results in the modification of its functional characteristics. Our previous studies revealed that fibrinogen adsorbs onto surfaces essentially in two different orientations depending on its concentration in the solution, “side-on” at low and “end-on” at high concentrations. In the present study, we analyzed the thrombin-mediated release of fibrinopeptides A and B (FpA and FpB) from fibrinogen adsorbed in these orientations; as well as from surface-bound fibrinogen-fibrin complexes, prepared by converting fibrinogen adsorbed in either orientation into fibrin and subsequently adding fibrinogen. The release of fibrinopeptides from surface-adsorbed fibrinogen and from surface-bound fibrinogen-fibrin complexes differed significantly, when compared to that from fibrinogen in solution. The release of FpB occurred without the delay (lag-phase) characteristic for its release from fibrinogen in solution. The amount of FpB released from “end-on” adsorbed fibrinogen and from adsorbed fibrinogen-fibrin complexes was much higher than that of FpA. FpB is known as a potent chemoattractant; therefore, its preferential release suggests the physiological purpose in the attraction of cells to the site of injury. The N-terminal portions of fibrin beta chains including residues Bβ15-42, which are exposed after cleavage of FpB, have been implicated in many processes, including angiogenesis and inflammation.

Keywords

Fibrinopeptide release, adsorbed fibrinogen, thrombin, fibrinopeptide A, fibrinopeptide B, biomaterials, thrombogenicity, inflammation, thrombus formation
Fibrinogen, one of the most abundant proteins in blood, plays a key role in hemostasis, inflammation, wound healing, and additional physiological and pathological processes. Immediately after blood comes in contact with artificial materials or with an injured vessel wall subendothelium, fibrinogen rapidly adsorbs on the surface and interacts with adhered activated platelets and subendothelial proteins. Numerous studies have demonstrated that fibrinogen in solution and fibrinogen adsorbed on various surfaces exhibit different properties [1-4]. For example, surface-adsorbed fibrinogen changes its conformation and thus reveals multiple binding sites that interact with the receptors on platelets and leukocytes [5,6]. These reciprocal interactions participate in the process of blood clot formation and in inflammatory response. Platelet adhesion promoted by the deposition of fibrinogen might contribute to the development of the inflammatory response during ischemia reperfusion. The structural properties of fibrinogen play a key role in its interactions with various biomolecules and cell types.

Fibrinogen is a 340 kDa plasma glycoprotein with a complex structure. The fibrinogen molecule consists of two identical subunits, each composed of three non-identical polypeptide chains: Aα, Bβ and γ. These chains are linked together by 29 disulfide bonds and form several structural regions; two distal D regions, one central E region, and two αC regions [7]. Each pair of distal nodules is linked with the central nodule by a triple helical coiled-coil connector, composed of the middle portions of all three chains. The COOH-terminal parts of the Aα chains (αC regions) fold back to the central E region to form two interacting αC-regions. The central nodule contains two pairs of polymerization sites (knobs), “A” and “B”; while the complementary polymerization sites (holes) “a” and “b” are located in the distal γ- and β-nodules, respectively. Additionally these nodules, as well as the αC-regions, contain
numerous binding sites that become active after the conversion of fibrinogen into fibrin or the adsorption of fibrinogen on various surfaces [6,10,11].

Conversion of monomeric fibrinogen into polymeric fibrin is mediated by thrombin, which binds to the central region of fibrinogen and catalyzes cleavage of the two short peptides, the 16-residue fibrinopeptide A (FpA) and the 14-residue fibrinopeptide B (FpB); each located at the NH2-termini of the Aα and Bβ chains, respectively [12]. This cleavage exposes knobs “A” and “B”, which interact with the complementary holes “a” and “b” of neighboring molecules to form a fibrin polymer (clot) [13]. According to the current view, the assembly of a fibrin clot in solution occurs in two steps. First, thrombin removes FpA enabling “A”-“a” interactions between individual half-staggered molecules and resulting in the formation of two-stranded protofibrils. Second, protofibrils aggregate laterally to make thicker fibers that coalesce to form a three-dimensional network of fibrin clot. It was shown that the lateral aggregation of protofibrils coincides with the removal of FpB enabling “B”-“b” interactions, and that FpB removal is accelerated by forming polymers [14,15]. Thus during fibrin polymerization in solution, the release of FpA occurs very rapidly, while the release of FpB is delayed and reaches its maximum when fibrin formation is almost complete. Such delay results in the sequential release of fibrinopeptides, and thereby sequential activation of the two sets of polymerization sites [12,16], which is necessary for normal fibrin assembly [15].

It has been suggested that the accelerating effect of polymer formation on FpB cleavage is related mainly to the interaction of the N-terminal parts of the Bβ chains with the dimeric DD regions formed in protofibrils and fibers, bringing FpB into the vicinity of bound thrombin [17-22].

So far, fibrin(ogen) characteristics, such as the release of fibrinopeptides and fibrin-fibrinogen interactions, have been mostly studied in solution [23,24]. However, adsorbed
fibrinogen acquires new properties important for biointeraction, as compared with fibrinogen in solution. The adsorption of fibrinogen at the site of vascular injury and inflammation, or on the artificial surfaces of vascular grafts and other blood-contacting components of medical devices plays a significant role in its interaction with cells and in blood coagulation [25,26]. Using infrared multi-internal reflection spectroscopy and monoclonal antibodies against fibrinogen E and D regions, we have previously found that fibrinogen adsorbs on glass, carbon, polyethylene, and polystyrene surfaces in basically two different orientation: “side-on” (laying on the surface) and “end-on” (standing on the surface) [27-29]. While side-on orientation prevails during adsorption from solutions with low fibrinogen concentrations; the other orientation, in which adsorbed fibrinogen is closely-packed, occurs with high fibrinogen concentrations. The adsorbed fibrinogen molecules in both cases have a reduced accessibility of thrombin to the E region. Therefore, the aim of the present study was to investigate fibrinopeptide release from fibrinogen layers adsorbed at different surface densities or immobilized on adsorbed fibrin.

Methods

Proteins and Reagents

Fibrinogen and thrombin were purchased from Sigma-Aldrich (Czech Republic). D-phenylalanyl-L-prolyl-L-arginine chloromethyl ketone (PPACK) and hirudin were purchased from Merck (Czech Republic). Solutions were prepared in a 0.05 M Tris buffer, pH 7.4, with 0.1 M NaCl and 2.5 mM CaCl₂ (TBS), and in 0.1 M phosphate buffered saline, pH 7.4 (PBS). Buffers were filtered through a Millipore 0.22 μm filter.

Attenuated Total Reflection Fourier Transform Infrared Spectroscopy (ATR FT-IR)

ATR FT-IR spectra were measured using a Bruker IFS 55 spectrometer equipped with a Wilks Sci ATR attachment. A ZnSe ATR element was spin coated with a polystyrene film,
thickness 60 nm. The kinetics of fibrinogen adsorption on the polystyrene surface from fibrinogen solutions was measured in situ in an ATR liquid flow cell. The amount of adsorbed fibrinogen was estimated by measuring the integral intensity of the fibrinogen Amide II band at 1549 cm\(^{-1}\). The intensity to mass ratio factor was calculated by comparing spectra of dried adsorbed fibrinogen layers with a calibration curve obtained by deposition and drying of known fibrinogen amounts on the polystyrene coated ZnSe.

**Surface Plasmon Resonance (SPR)**

The SPR instrument based on the spectral interrogation of SPR conditions was manufactured at the Institute of Photonics and Electronics, Academy of Sciences of the Czech Republic, Prague. The tested solutions were flowed along the gold surface of SPR chips through the 4-channel flow cell using a peristaltic pump. An increase in the mass of immobilized proteins is expressed by an increase in resonance wavelength \(\lambda_{\text{res}}\) (nm).

**Preparation of adsorbed fibrin(ogen) and fibrinopeptide samples**

Fibrin(ogen) structures were prepared on the bottoms of polystyrene wells in 6-well plates (Polystyrene Non-Tissue Treated Plate, well diameter 3.5 cm, bottom surface area \(\sim 10 \text{ cm}^2\), Falcon Multiwell\textsuperscript{TM}; Becton Dickinson Labware, France). A primary adsorbed fibrinogen layer was prepared by application of a solution of either 20 μg/ml or 500 μg/ml fibrinogen in TBS for 150 or 30 minutes, respectively. The wells were then successively washed with TBS, PBS, and incubated in a thrombin solution (2.5 U/ml in PBS) for 60 minutes. Solutions containing released fibrinopeptides were collected at selected reaction times and consequently the samples obtained for HPLC analysis were frozen and stored. Sample obtained from one well represents one point in the graph. The completeness of release
of all accessible fibrinopeptides after 60 minutes under these conditions was proven earlier [30].

The second fibrinogen layer was prepared on a primary adsorbed fibrinogen layer treated with thrombin for 60 minutes as follows. The well was washed with PBS and TBS; and the surface-bound thrombin was inhibited by incubation with PPACK (10 μM) and hirudin (6 U/ml) for 20 minutes. After washing with TBS, fibrinogen at either 20 μg/ml or 200 μg/ml in TBS was applied for 120 and 60 minutes, respectively. The surface was washed with TBS and PBS, and treated again with thrombin in PBS (2.5 U/ml). The samples containing released fibrinopeptides were obtained and processed in the same manner as in the case of the first fibrinogen layer. The interaction of surface-bound fibrin with fibrinogen was proven previously [31]. Fibrinopeptide release from fibrinogen in solution (3 mg/ml) was performed using 2.5 U/ml thrombin. The reaction was stopped by trichloroacetic acid (final concentration; 5% m/v) at selected times, the precipitates centrifuged; and the supernatants analyzed by HPLC.

Transmission electron microscopy (TEM)

The samples of fibrinogen layers for transmission electron microscopy were prepared on a carbon-coated mica using the same procedure as described above. Side-on and end-on attachment of fibrinogen at low and at high concentration on carbon surface was proven earlier [27,29]. Saturated primary monolayers of fibrinogen molecules were prepared. Fibrinogen layers were washed with buffer, and contrasted with 5% uranyl acetate in water for 5 min [32]. They were then washed with water, dehydrated with a series of water-ethanol solutions with an increasing ethanol concentration (0, 25, 50, 75, 100%), and dried in a flow box [33]. The carbon film was floated from the mica at water level and deposited on a
microscope grid. The samples were observed using JEOL JEM-1010 and JEOL 200CX transmission electron microscopes at 100 kV.

**HPLC analysis**

Collected supernatants containing released fibrinopeptides were dried with a centrifugal vacuum concentrator, reconstituted in mobile phase A, filtered using Millipore 0.2 μm centrifugal filters, and analyzed by the RP-HPLC method, essentially according to Suttnar et al. [34]. Peptide concentrations were obtained using calibration curves of pure substances.

**Results**

**Preparation and characterization of fibrinogen surfaces.**

To prepare fibrinogen surfaces for the study of the thrombin-mediated cleavage of fibrinopeptides from adsorbed fibrinogen, we first characterized the adsorption of fibrinogen to the polystyrene surface (formation of a primary fibrinogen layer) and the immobilization of a second fibrinogen layer onto the primary layer.

To coat a surface with the primary layer, fibrinogen solutions at high (500 μg/ml) or low (20 μg/ml) concentrations were incubated with the polystyrene surface. The amount of fibrinogen adsorbed on the surface was determined at different time points using ATR FT-IR Spectroscopy (Fig. 1). Based on previous studies [27-29,32,33], one can assume that when fibrinogen adsorption was performed from the high or low concentration solutions, the polystyrene surface was coated with a saturated monolayer of fibrinogen molecules oriented in “end-on” or “side-on” manner, respectively.

To monitor the formation of an additional (secondary) layer of fibrinogen on the primary layer, we used surface plasmon resonance (SPR). The SPR signal, which reflects the accumulation of fibrinogen on the surface, revealed that fibrinogen adsorption from the high
or low concentration almost reached saturation after 30 or 150 min, respectively (Fig. 2). The SPR signal revealed the additional immobilization of fibrinogen that presumably occurred through the specific fibrinogen-fibrin interaction. This interaction involves always reactive holes “a” and “b” located in the D regions of the added fibrinogen, and the complementary knobs “A” and “B” of the E regions of the adsorbed primary layer, left exposed by the thrombin treatment. Only a very small drop in the SPR signal was observed when the fibrinogen solution was replaced with TBS (Fig. 2, curve 1, right arrow labeled “TBS”). There was no change in the amount of immobilized fibrinogen when the surface was further contacted with fibrinogen at 500 μg/ml, or with a less concentrated fibrinogen solution of 200 μg/ml (not shown). This suggests that the binding of the secondary fibrinogen layer to the thrombin-treated primary layer, adsorbed from the high concentration of fibrinogen, was specific and reached saturation.

The immobilization of a secondary layer of fibrinogen on the primary layer adsorbed from the low (20 μg/ml) fibrinogen concentration (presumably “side-on” molecule orientation in the primary layer) was performed similarly, except that the secondary layer was formed by the application of fibrinogen at two differing concentrations, high (200 μg/ml) and low (20 μg/ml) (Fig. 2, curves 2 and 4). The immobilization of fibrinogen from the high concentration in this case also reached saturation. The binding was determined to be tight, as practically no drop in SPR signal was observed when the fibrinogen solution was replaced with TBS (Fig. 2, curve 2, right arrow labeled “TBS”). In a control experiment, when the primary “side-on” layer was not treated with thrombin, and fibrinogen at high concentration (200 μg/ml) was applied, some additional binding of fibrinogen was detected (Fig. 2, curve 3). However, this binding was about two times lower than the binding of fibrinogen applied at the same concentration to the thrombin-treated primary layer (Fig. 2, curve 2), suggesting that at least half of the latter occurred through the specific fibrinogen-fibrin interaction.
Imaging of the surface fibrinogen-fibrin layers by transmission electron microscopy revealed that their appearance depended on the arrangement of the primary layer adsorbed on the surface. Small protofibrils were observed much more clearly when the primary fibrinogen layer was adsorbed from the low solution concentration of 20 μg/ml, treated with thrombin and thrombin inhibitors, PPACK and hirudin, and subsequently fibrinogen was attached by incubating it in 200 μg/ml (Fig. 3A); in contrast with the mostly homogeneous layer when the primary layer was adsorbed from the high fibrinogen concentration of 500 μg/ml, treated with thrombin, PPACK and hirudin, and incubated in a 200 μg/ml fibrinogen solution (Fig. 3B). TEM images of primary layers prepared by adsorption from low or high fibrinogen concentration revealed completely coated surfaces (data not shown).

**Thrombin-mediated release of fibrinopeptides from fibrinogen surfaces.**

When the primary layer was formed by the adsorption of fibrinogen from the higher concentration (presumably “end-on” adsorption), the release of both fibrinopeptides occurred very rapidly and reached their maximum after 60 min (Fig. 4A). The initial rates of release of both fibrinopeptides were similar; however, the amount of released FpA was more than 40% lower than that of FpB (17 versus 30 pmol). In contrast, when the primary layer was formed by the adsorption of fibrinogen from the lower concentration (presumably “side-on” adsorption), both the initial rate and the amount of FpA released were higher than those of FpB (Fig. 4B). Nevertheless, in 60 min the amount of released FpB approached that of FpA (about 10 pmol). Such kinetics of FpA and FpB release is reminiscent of that of their release from fibrinogen in solution (shown in Fig. 4C for comparison); however, the delay in FpB release characteristic for the latter was not observed with fibrinogen adsorbed from both low and high concentrations. Altogether, these results suggest that the mechanism of thrombin-
mediated fibrinopeptide release from adsorbed fibrinogen differs from that of fibrinogen in solution. It also indicates that such a mechanism depends on the orientation of the adsorbed fibrinogen.

The results presented in Fig. 4A suggest that not all FpA are accessible for thrombin in the fibrinogen adsorbed from the high concentration. To evaluate the accessibility of FpA, as well as that of FpB, we calculated the amount of fibrinogen adsorbed on the surface of polystyrene wells, using the value of the coating density determined by ATR FT-IR Spectroscopy (Fig. 1). Thrombin cleaved only approximately 37% of FpA and 64% of FpB from the fibrinogen adsorbed from the high concentration and approximately 36% of FpA and 35% of FpB from the fibrinogen adsorbed from the low concentration (Fig.4). Thus, in both adsorbed fibrinogens (adsorbed from both the high and low concentrations) only a portion of the molecules had fibrinopeptides accessible for cleavage by thrombin. Presumably these molecules were converted into fibrin upon treatment with thrombin, and bound the added fibrinogen to form the secondary fibrinogen layer (or surface-bound fibrinogen-fibrin complexes), as described in the previous section (SPR measurement, Fig 2, curves 1 and 2).

The release of fibrinopeptides from the secondary fibrinogen layers immobilized on fibrin primary layers markedly differed from that of the original primary layers (Fig.5).

Discussion

The interaction of fibrinogen with various proteins and cell types, which is of great importance in many vital processes, often occurs after the immobilization of fibrinogen on a surface. Numerous previous studies have characterized fibrinogen properties, including the release of fibrinopeptides and subsequent fibrin formation, mainly in solution. In the present study, we examined the properties of fibrinogen molecules adsorbed on a surface or immobilized on previously adsorbed fibrin. Our results clearly show that the thrombin-
mediated release of fibrinopeptides from adsorbed/immobilized fibrinogen differs significantly from their release from fibrinogen in solution. Namely, (i) the delay of FpB release, characteristic of fibrinogen in solution, was not observed with adsorbed/immobilized fibrinogen; and (ii) the amount of FpB released from “end-on” adsorbed fibrinogen or from the secondary fibrinogen layer immobilized on adsorbed fibrin, was significantly higher than that of FpA. Two main questions arise from our results: (1) why is the release of fibrinopeptides from surface-bound fibrinogen so different from that of fibrinogen in solution; and (2) is this of any significant (patho)physiological importance or relevance?

To address the first question, one should consider the possible arrangement of fibrinogen molecules on the surface. When fibrinogen was adsorbed from the lower concentrations in a “side-on” manner, only about one-thirds of fibrinopeptides were released by thrombin. It seems that fibrinogen adsorption was random; and only this fraction of adsorbed fibrinogen had its fibrinopeptides accessible for thrombin. As in the case with fibrinogen in solution (Fig.4c), the initial rate of FpA release from “side-on” adsorbed fibrinogen was higher than that of FpB, while the final amounts of released fibrinopeptides were similar (Fig.3b). This may be due to the non-substrate interaction of thrombin with adsorbed fibrinogen, which was implicated in the sequential cleavage of fibrinopeptides from fibrinogen in solution [22]. At the same time, no delay of FpB release (lag-phase) from “side-on” adsorbed fibrinogen was observed. Since such a delay was proposed to be connected with conformational changes accompanying FpA release [21] and protofibril formation [22], such changes seem to already occur in adsorbed fibrinogen. This may also be a reason for the absence of a lag-phase in FpB release from the “end-on” adsorbed fibrinogen or fibrinogen attached to the primary fibrin layer (Fig. 4a and 5).

The situation with “end-on” adsorbed fibrinogen was quite different, as the initial rates of FpA and FpB release were similar, while the final amount of released FpA was
significantly (by 40%) lower than that of FpB. This may be explained by the different accessibility of FpA and FpB for thrombin. Indeed, fibrinopeptides A are located in the central part of the molecule [20,22], which should be difficult to access for thrombin in the “end-on” configured adsorbed fibrinogen. The N-terminal segments of the Bβ chains containing FpB were proposed to be flexible [9,35], and according to the results of molecular modeling, may stretch as far as to the outer nodules [22]. Thus, the accessibility of FpB in “end-on” adsorbed fibrinogen should be superior to that of FpA. Similarly, when the fibrinogen from solution of the high concentration (200 μg/ml) is attached to the primary layer of “end-on” adsorbed thrombin-treated fibrinogen, the accessibility of FpB for thrombin should also be greater. This was observed in our experiments (Fig. 5a).

Our results show that FpB is preferentially released from the secondary fibrinogen layers on the surface of growing fibrins. The tendency for reduced FpA accessibility increased with the increase of the coating concentration (Fig. 5b, c). The explanation for such difference in the accessibility of fibrinopeptides is not entirely clear. One can only speculate that this may be a result of the conformational changes in fibrinogen upon its immobilization, or steric hindrance arising from the fibrinogen-fibrin complex formation. The role of the αC region, if any, in the described process is not known and remains to be evaluated. The interaction between fibrin monomers and fibrin(ogen), or vice versa, is characterized by a rather high affinity (K_D ~10^{-8}M) and a change of standard Gibbs energy of about -10 kcal/mol [36]. These data demonstrate that the equilibrium is thermodynamically favored for the formation of a stable fibrin-fibrinogen complex. We have shown previously that the overall structure of fibrin arising with the preferential release of FpB by a specific snake venom enzyme is essentially the same as the structure of fibrin initiated by thrombin [37].

It should be noted that both our results and those of others have previously indicated no lag-phase (delay) in the thrombin-mediated release of FpB from the fibrinogen adsorbed
on glass surfaces, and on negatively and positively charged or hydrophobic and hydrophilic surfaces [27,28,31,38]. Also no delay in the release of FpB was observed by Blombäck and Bark [24] in their experimentation with whole blood and platelet-rich plasma, in which FpB was released almost as quickly as FpA. To explain these observations, they suggested that the binding of fibrinogen to the platelet receptor GPIIb/IIIa may induce conformational changes in bound fibrinogen, resulting in the exposure of a thrombin-susceptible cleavage site and thus facilitating FpB release.

Our data, in accordance with many other authors [2,5] suggest that the adsorption of fibrinogen causes changes in its conformation that may have a further impact on its behavior toward other proteins and cell surface receptors. In addition, the TEM experiments revealed that thrombin-treated fibrinogen molecules, adsorbed “side-on” to the surface from the low concentration fibrinogen solution, promoted the formation of a surface-attached fibrin(ogen) network better than the layer of closely packed thrombin-treated fibrinogen molecules, adsorbed “end-on” from the higher concentration (Fig 3).

As to the (patho)physiological relevance of the observed properties of adsorbed/immobilized fibrinogen, one should take into account the following: First, the N-terminal portions of fibrin β chains (amino acid residues β15-42), which are exposed after cleavage of FpB, have been implicated in a number of important processes including angiogenesis and inflammation [39-48]. Second, the released FpB is a potent chemoattractant [42]; and therefore, its preferential release may indicate the physiological purpose in the attraction of cells to the site of injury.

It has been shown that the association of soluble fibrinogen with the fibrin clot, results in the reduced adhesiveness of such fibrinogen-fibrin matrices towards leukocytes and platelets. [49,50]. Our finding that fibrinopeptide A is less accessible for thrombin in surface-bound fibrinogen-fibrin complexes may be interpreted as a novel supplementary mechanism...
preventing the rapid conversion of bound fibrinogen into fibrin, and thereby extending its anti-adhesive properties, providing an additional level of protection of thrombi from premature dissolution.

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Authorship and Conflict of Interest Statements

Authorship contribution:

J.E.Dyr designed the project, performed SPR experiments, analyzed the data, and wrote the paper; T.Riedel prepared adsorbed fibrin(ogen) and fibrinopeptie samples, designed and performed SPR, TEM and ATR FT-IR experiments, analyzed the data and wrote the paper, J.Suttnar performed HPLC analysis and analyzed the data, E.Brynda designed TEM and ATR FT-IR experiments and analyzed the data, M.Houska designed SPR experiments and analyzed the data, L.Medved analyzed the data and wrote the paper.

Conflict-of-interest disclosure:

The authors declare no competing financial interests.

References


Figure Legends

**Fig. 1. Adsorption of a primary fibrinogen layer observed by ATR FT-IR Spectroscopy.**
Fibrinogen (Fbg) was adsorbed onto polystyrene from high (500 μg/ml) and low (20 μg/ml) concentration fibrinogen solutions (open and solid squares, respectively). Inserts show an illustrative arrangement of fibrinogen molecules adsorbed in the “end-on’ (upper insert) and “side-on” orientations.

**Fig. 2. SPR-observed adsorption of primary and secondary fibrinogen layers.**
A primary layer was formed by the adsorption of fibrinogen onto the gold surface, followed by washing with TBS and PBS; a secondary fibrinogen layer was formed by the immobilization of fibrinogen onto the primary layer, which was treated with thrombin to convert the adsorbed fibrinogen into fibrin (see text). Arrows indicate the replacement of solutions: Curve 1 - fibrinogen (Fbg) at 500 μg/ml, TBS, PBS; thrombin at 2.5 U/ml (THR), PBS, TBS; mixture of PPACK at 10 μM and hirudin at 6 U/ml, TBS; fibrinogen at 200 μg/ml, TBS. Curve 2 - fibrinogen at 20 μg/ml, TBS, PBS; thrombin at 2.5 U/ml, PBS, TBS; PPACK at 10 μM and hirudin at 6 U/ml, TBS; fibrinogen at 200 μg/ml, TBS. Curve 3 - fibrinogen at 20 μg/ml, TBS, PBS; fibrinogen at 200 μg/ml, TBS. Curve 4 - fibrinogen 20 μg/ml, TBS, PBS; thrombin 2.5 U/ml, PBS, TBS; PPACK 10 μM and hirudin 6 U/ml, TBS; fibrinogen 20 μg/ml, TBS.

**Fig. 3. Morphology of fibrinogen-fibrin layers on carbon, observed by TEM**
Fibrinogen was adsorbed on carbon-coated mica at a concentration of 20μg/ml for 150min (a) and 500μg/ml for 30min (b), treated with thrombin at 2.5 U/ml for 60 min, mixture of PPACK at 10μM and hirudin at 6U/ml for 20min, and with fibrinogen at a concentration of 200μg/ml for 60 min. The layers were contrasted with uranyl acetate, dehydrated with a series of water–ethanol solutions, and dried.

**Fig. 4. Kinetics of thrombin-mediated release of fibrinopeptides from the primary fibrinogen layers adsorbed on polystyrene.**

The amounts of FpA (open squares) and FpB (solid squares) released from the primary fibrinogen layers adsorbed from the high (500 μg/ml, 30 min) and low (20 μg/ml, 150 min) fibrinogen concentrations (panels a and b, respectively), and from fibrinogen in solution (panel c) was determined by HPLC, as described in Materials and Methods. The concentration of thrombin in all experiments was 2.5 U/ml. Each point represents a mean value obtained from three independent experiments in which the total amount released in one well at the indicated time interval was measured. Inserts in panels a and b show an illustrative arrangement of fibrinogen molecules adsorbed in “end-on’ and “side-on” orientations, respectively; and in panel c fibrinogen in solution.

**Fig. 5. Kinetics of thrombin-mediated release of fibrinopeptides from the secondary layer of surface-bound fibrinogen-fibrin complexes.**

All fibrinogen-fibrin complexes were formed by the immobilization of the secondary fibrinogen layer on thrombin-treated (2.5 U/ml thrombin for 60 min, followed by a mixture of PPACK and hirudin for 20 min) primary fibrinogen layers adsorbed onto polystyrene. (a) The primary layer was adsorbed from 500 μg/ml fibrinogen for 30 min; the secondary layer was immobilized at 200 μg/ml fibrinogen for 60 min. (b) The primary layer was adsorbed from
20 μg/ml fibrinogen for 150 min; the secondary layer was immobilized at 20 μg/ml fibrinogen for 120 min. (c) The primary layer was adsorbed from 20 μg/ml fibrinogen for 150 min; the secondary layer was immobilized at 200 μg/ml fibrinogen for 60 min. The amounts of FpA (open squares) and FpB (solid squares) released from the secondary layers upon incubation with 2.5 U/ml thrombin were determined by HPLC, as described in Materials and Methods. Each point represents a mean value obtained from three independent experiments.
Figure 1

The graph illustrates the Fbg surface coverage (μg/cm²) over time (min). The data points are shown as squares, with the black line representing the trend. The inset on the right side of the graph provides a visual representation of the surface coverage over time.
Figure 2

Resonant wavelength shift (nm)

Time (min)

Fbg, TBS, PBS, THR, PPACK/Hirudin, Fbg, TBS, TBS

1, 2, 3, 4
Figure 4

(a) 

Fibrinopeptides (pmol)

Time (min)

(b) 

Fibrinopeptides (pmol)

Time (min)

(c) 

Fibrinopeptides (pmol)

Time (min)
Figure 5

a

b

c

Fibrinopeptides (pmol)

0 1 2 3 4 5 6 7

Time (min)

0 10 20 30 40 50 60

Fibrinopeptides (pmol)

0 2 4 6 8 10 12 14

Time (min)

0 10 20 30 40 50 60

Fibrinopeptides (pmol)

0 2 4 6 8 10 12 14

Time (min)

0 10 20 30 40 50 60

Legend:

- FpA
- FpB
Fibrinopeptides A and B release in the process of surface fibrin formation

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