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The Dual Regulatory Role of Amino Acids Leu⁴⁸⁰ and Gln⁴⁸¹ of Prothrombin*

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Prothrombin (FII) is activated to α -thrombin (IIa) by prothrombinase. Prothrombinase is composed of a catalytic subunit, factor Xa (fXa), and a regulatory subunit, factor Va (fVa), assembled on a membrane surface in the presence of divalent metal ions. We constructed, expressed, and purified several mutated recombinant FII (rFII) molecules within the previously determined fVa-dependent binding site for fXa (amino acid region 473–487 of FII). rFII molecules bearing overlapping deletions within this significant region first established the minimal stretch of amino acids required for the fVa-dependent recognition exosite for fXa in prothrombinase within the amino acid sequence Ser⁴⁷⁸–Val⁴⁷⁹–Leu⁴⁸⁰–Gln⁴⁸¹–Val⁴⁸². Single, double, and triple point mutations within this stretch of rFII allowed for the identification of Leu⁴⁸⁰ and Gln⁴⁸¹ as the two essential amino acids responsible for the enhanced activation of FII by prothrombinase. Unanticipated results demonstrated that although recombinant wild type α -thrombin and rIIa^{S478A} were able to induce clotting and activate factor V and factor VIII with rates similar to the plasma-derived molecule, rIIa^{SLQ→AAA} with mutations S478A/L480A/Q481A was deficient in clotting activity and unable to efficiently activate the pro-cofactors. This molecule was also impaired in protein C activation. Similar results were obtained with rIIa^{ΔSLQ} (where rIIa^{ΔSLQ} is recombinant human α -thrombin with amino acids Ser⁴⁷⁸/Leu⁴⁸⁰/Gln⁴⁸¹ deleted). These data provide new evidence demonstrating that amino acid sequence Leu⁴⁸⁰–Gln⁴⁸¹: 1) is crucial for proper recognition of the fVa-dependent site(s) for fXa within prothrombinase on FII, required for efficient initial cleavage of FII at Arg³²⁰; and 2) is compulsory for appropriate tethering of fV, fVIII, and protein C required for their timely activation by IIa.

In the presence of a procoagulant membrane surface and divalent metal ions, factor Va (fVa)³ binds factor Xa (fXa) to form prothrombinase. Prothrombinase is the two-subunit enzymatic complex where the non-enzymatic regulatory subunit (fVa) controls the rate and directs cleavage of prothrombin (FII) by the catalytic subunit (fXa) at two spatially distinct sites resulting in timely α -thrombin (IIa) formation at the place of vascular injury (1–3). Cleavage at Arg²⁷¹ and Arg³²⁰ of FII is required to form the active serine protease IIa. The essential IIa molecule bears strong homology with other serine protease enzymes, such as activated protein C (APC), chymotrypsin, and fXa. Several different numberings of IIa residues appear in the literature based on either the chymotrypsin numbering (4) or IIa numbering (5, 6) or the entire FII sequence (7). The latter nomenclature is used herein with the appropriate chymotrypsin numbering in parentheses when required for comparison with the existing data in the literature.

Historically, it has been shown that in the absence of fVa, initial cleavage at Arg²⁷¹ of FII results in the generation of the inactive intermediate prethrombin-2 and fragment 1·2. Further cleavage of prethrombin-2 at Arg³²⁰ generates IIa (prethrombin-2 pathway) (8–21). Concurrent with the appearance of excess fVa during clotting and in the presence of a procoagulant surface, the order of cleavages is reversed, and initial cleavage at Arg³²⁰ generates a transient enzymatically active intermediate, meizothrombin, that has much higher catalytic efficiency than IIa toward chromogenic substrates usually employed to assess IIa activity (18, 22–24). Meizothrombin is next cleaved at Arg²⁷¹ resulting in the generation of IIa and fragment 1·2 (meizothrombin pathway). Although efficient cleavage at each site requires the presence of phospholipids, initial cleavage at Arg³²⁰ is entirely fVa-dependent.

In the absence of fVa, the two activation cleavage sites are not readily available, and FII is activated at a slow non-physiological rate by membrane-bound fXa alone. Interactions between fXa and FII are known to exist in the presence and absence of fVa; however, the enhanced activity of fXa within prothrombinase toward both activating cleavage sites is controlled solely by the membrane-bound non-enzymatic cofactor (3, 16, 17, 25, 26).

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³ The abbreviations used are: fVa, factor Va; fV, factor V; FII, prothrombin; fXa, factor Xa; IIa, α -thrombin; r, recombinant; PC, L- α -phosphatidylcholine; PCPS, small unilamellar phospholipids vesicles composed of 75% PC and 25% L- α -phosphatidylserine (w/w); rFII^{WT}, recombinant wild type human prothrombin; rIIa^{WT}, recombinant wild type human α -thrombin; fVIII, factor VIII; DAPA, dansylarginine-N-(3-ethyl-1,5-pentanediy)amide; ABE, anion-binding (pro)exosite; APC, activated protein C; TP, prothrombin time.

Consequently, the innate process of coagulation rests on specific molecular interactions involved in the fVa-dependent activation of FII by prothrombinase. In relation to fXa alone, the relative rate of IIa formation by prothrombinase is increased by 300,000-fold because of the increase in the rates of both FII cleavages. This increase is mainly associated with a large (3,000-fold) increase in the k_{cat} value of fXa within prothrombinase with a 100-fold decrease in the K_m value of the enzyme (16). This substantial increase in enzymatic activity resulting in rapid and physiologically relevant IIa generation at the place of vascular injury is credited through precise and unique interactions of the cofactor with specific amino acids affiliated with both membrane-bound fXa and membrane-bound FII as recently demonstrated (27). Accordingly, the introduction of the non-enzymatic cofactor into prothrombinase equips the organism's coagulation artillery necessary for the explosive arrest of vasculature bleeding.

Factor V (fV) is a large quiescent multidomain (A1–A2–B–A3–C1–C2) protein that circulates in blood at a concentration of 20 nM (28–31). Three sequential cleavages of fV at Arg⁷⁰⁹, Arg¹⁰¹⁸, and Arg¹⁵⁴⁵ (29, 32–34) by IIa and/or fXa (35, 36) release the B domain and promote formation of the active cofactor fVa. FII circulates abundantly in blood at a concentration of 1.4 μM as the zymogen form of the serine protease IIa (7, 37). Mature FII protein is composed of a region containing several post-translationally modified γ -carboxyglutamic acid residues (described as the Gla domain, residues 1–46), followed by two Kringle domains (residues 65–143 and 170–248, respectively) and a serine protease domain (residues 272–579, see Fig. 1). FII contains three linkers as follows: linker 1 (residues 47–64) connects the Gla domain to kringle-1; linker 2 (residues 144–169) connects the two kringles; and linker 3 (residues 249–284) connects kringle-2 to the A-chain portion of IIa (7, 38) (Fig. 1).

The necessary fVa-dependent activation of FII by prothrombinase is a widely studied mechanism of coagulation but is still poorly understood. Numerous fVa-binding sites are acknowledged to exist on FII. Earlier investigations have shown the existence of binding sites on FII for fVa in each of the kringle domains (39–41) and within the Gla domain (42). Furthermore, significant protein-protein interactions between the acidic COOH-terminal region of fVa and a region rich in basic amino acids of FII have been inferred and characterized indirectly by employing molecular techniques involving specific hirudin-like ligands and the anion-binding (pro)exosite I (ABE I) of FII derivatives, as well as directly using a specific acidic peptide derived from the COOH-terminal region of the fVa heavy chain and recombinant fVa molecules (43–49). Site-directed mutagenesis of the basic residues in the proenzyme generated a recombinant FII molecule impaired in its ability to be activated by fully assembled prothrombinase (50). Although a crystal structure and a model of fVa have been available for some time now (51, 52), the crucial interaction of the acidic hirudin-like COOH-terminal portion of the heavy chain of the cofactor with FII required for efficient IIa formation was initially ignored because it was missing from the crystal structure of the cofactor (51). This interaction was further discounted without providing any solid evidence (53) despite initial findings by Guinto and Esmon (54) and more recent original find-

ings from our laboratory (47–49). A very recent model of prothrombinase using as a template the crystal structure of prothrombinase from the snake venom of *Pseudonaja textilis* verified and established the critical role of the acidic COOH-terminal region of fVa heavy chain for optimal rates of FII cleavage at two spatially distinct sites by prothrombinase resulting in timely IIa formation at the place of vascular injury (27, 55).

Additional studies with several recombinant prethrombin-1 molecules, where seven critical basic amino acids within (pro)exosite I were individually changed to glutamic acid, confirmed the interaction of (pro)exosite I with fVa acidic regions (50). Notably, the data revealed that although mutated prethrombin-1 is a poor substrate for prothrombinase, the same molecule was activated by membrane-bound fXa alone with similar rates as wild type prethrombin-1. Supplementary to these studies, Yegneswaran *et al.* (56), utilizing synthetic peptide derived from a highly conserved region of FII, postulated the existence of an fVa-dependent binding exosite for fXa within the sequence 473–487 of FII (chymotrypsin numbering 149D-163 (4)) that is in close spatial arrangement to (pro)exosite I. The same authors have also identified an fVa-independent site for fXa on prothrombin (amino acids 557–571) (57).

This study was initiated to identify and investigate the identity and role of the minimum required amino acid stretch within sequence 473–487 of FII that is conserved in a wide range of mammalian species and regulates peptide bond specificity and FII activation by prothrombinase in an fVa-dependent manner. Our findings identify for the first time two specific amino acids within FII that have a dual role. They are required for efficient fVa-dependent tethering of fXa needed for timely FII cleavage at Arg³²⁰ and IIa formation, while also serving an important role in directing efficient cleavage of IIa's physiological substrates. The latter is a prerequisite for expression of optimal physiological IIa activity.

Experimental Procedures

Materials—Phenylmethylsulfonyl fluoride (PMSF), *o*-phenylenediamine dihydrochloride, Hepes, Trizma (Tris base), and Coomassie Blue R-250 were purchased from Sigma. fV-deficient plasma was purchased from Research Protein Inc. (Essex Junction, VT). Secondary anti-mouse, anti-sheep, and anti-equine IgG coupled to peroxidase were from Southern Biotechnology Associates, Inc. (Birmingham, AL). L- α -Phosphatidylserine (PS) and L- α -phosphatidylcholine (PC) were from Avanti Polar Lipids (Alabaster, AL). Chemiluminescent reagent ECL Plus, heparin-Sepharose, and Mono Q 5/50 columns were from GE Healthcare. Normal reference plasma and chromogenic substrate Spectrozyme-TH were from American Diagnostica Inc. (Greenwich, CT). S-2238 was from AnaSpec (Fremont, CA); recombiPlasTin used in the clotting assays was purchased from Instrumentation Laboratory Co. (Lexington, MA). The reversible fluorescent IIa inhibitor dansylarginine-*N*-(3-ethyl-1,5-pentanedyl)amide (DAPA), human plasma-derived protein C, human plasma-derived IIa, human plasma-derived FII, and FII-deficient plasma were purchased from Hematologic Technologies Inc. (Essex Junction, VT). The purified human plasma-derived protein C preparation used contained both heavy chain isoforms that are activated to APC with similar

rates as described earlier (58, 59). Human fXa was purchased from Enzyme Research Laboratories (South Bend, IN). The plasmid pZEM229R-lite encoding human recombinant prothrombin (rFII) was a generous gift from Dr. Kathleen Berkner (Cleveland Clinic Foundation, Cleveland, OH). QuikChange® II XL site-directed mutagenesis kit was obtained from Agilent Technologies Genomics (Santa Clara, CA). All molecular biology and tissue culture reagents, specific primers, and medium were obtained from Gibco, Invitrogen, or as indicated. Monoclonal antibodies to fV (α HFV_{HC}17 and α HFV_{LC}9), monoclonal antibody α HFV1 coupled to Sepharose used to purify plasma and recombinant fV molecules, and a polyclonal antibody to FII used for immunoblotting experiments during rFII production were provided by Dr. Kenneth G. Mann (Department of Biochemistry, University of Vermont, Burlington). Plasma factor V (fV^{plasma}) and plasma fVa (fVa^{plasma}) were purified as described previously (60–62).

Construction of rFII Molecules—To investigate the importance of amino acid region 473–487 of the serine protease domain of FII, we first constructed a recombinant mutant FII molecule with this region deleted (rFII^{Δ473–487}) using Stratagene's QuikChange® site-directed mutagenesis kit and the pZEM229R-lite plasmid. rFII^{Δ473–487} was constructed using the mutagenic primers 5'-GAG ACG TGG ACA GCC AAC GTT GTG GAG CGG CCG GTC TGC AAG-3' (sense) and 5'-CTT GCA GAC CGG CCG CTC CAC AAC GTT GGC TGT CCA CGT CTC-3' (antisense) (corresponding to the ⁴⁷³GKGQPSVLQV⁴⁸⁷ deletion). The mutation was confirmed by DNA sequencing (DNA Analysis Facility, Department of Molecular Cardiology at The Lerner Research Institute, Cleveland Clinic). To further investigate the minimum sequence of amino acids required for the fVa-dependent fXa binding on FII within the region 473–487 of the serine protease domain, several rFII molecules with the mutations denoted as rFII^{ΔN10} (rFII^{ΔN10} is recombinant human prothrombin missing amino acids GKGQPSVLQV), rFII^{ΔC10}, rFII^{ΔS5V}, rFII^{S478A} (rFII^{S478A} is recombinant human prothrombin with the mutation S478A), rFII^{L480A}, rFII^{SL→AA}, rFII^{SQ→AA}, and rFII^{SLQ→AAA} (where rFII^{SLQ→AA} is recombinant human prothrombin with the mutation S478A/L480A/Q481A) were constructed using Stratagene's QuikChange® site-directed mutagenesis kit and the pZEM229R-lite plasmid. First, overlapping deletions in the region 473–487 were constructed using the mutagenic primers for rFII^{ΔN10} 5'-G ACG TGG ACA GCC AAC GTT GTG AAC CTG CCC ATT GTG GAG-3' (sense) and 5'-CTC CAC AAT GGG CAG GTT CAC AAC GTT GGC TGT CCA CGT C-3' (antisense) (corresponding to the ⁴⁷³GKGQPSVLQV⁴⁸² deletion), whereas mutagenic primers used for rFII^{ΔC10} were 5'-GTT GGT AAG GGG CAG CCC GTG GAG CGG CCG GTC TGC-3' (sense) and 5'-GCA GAC CGG CCG CTC CAC GGG CTG CCC CTT ACC AAC-3' (antisense) (corresponding to the ⁴⁷⁸SVLQVVNLPI⁴⁸⁷ deletion). Similarly, the middle deletion of the overlapping mutations rFII^{ΔS5V} was constructed using the mutagenic primers 5'-GTT GGT AAG GGG CAG CCC GTG AAC CTG CCC ATT GTG-3' (sense) and 5'-CAC AAT GGG CAG GTT CAC GGG CTG CCC CTT ACC AAC-3' (antisense) (corresponding to the ⁴⁷⁸SVLQV⁴⁸² middle deletion). Next, within the sequence

478–482 several point mutations were made based on amino acid solvent exposure and homology within other proteins. The first rFII point alanine mutation rFII^{S478A} was constructed using the mutagenic primers 5'-GGT AAG GGG CAG CCC GCA GTC CTG CAG GTG-3' (sense) and 5'-CAC CTG CAG GAC TGC GGG CTG CCC CTT ACC-3' (antisense) (corresponding to the S478A mutation). The next single point mutation rFII^{L480A} was constructed using the mutagenic primers 5'-GGG CAG CCC AGT GTC GCG CAG GTG GTG AAC CTG CCC-3' (sense) and 5'-GGG CAG GTT CAC CAC CTG CCG GAC ACT GGG CTG CCC-3' (antisense) (corresponding to the L480A mutation). In addition to single point mutations, we constructed two double alanine mutations, rFII^{SL→AA} and rFII^{SQ→AA}, within the stretch 478–482 of FII. For rFII^{SL→AA}, we used the mutagenic primers 5'-GGT AAG GGG CAG CCC GCA GTC GCG CAG GTG GTG AAC CTG-3' (sense) and 5'-CAG GTT CAC CAC CTG CGC CAC TGC GGG CTG CCC CTT ACC-3' (antisense) (corresponding to the S478A/L480A mutation). Also, for rFII^{SQ→AA} (where rFII^{SQ→AA} is recombinant human prothrombin with the mutation S478A/Q481A), we used the mutagenic primers 5'-GGT AAG GGG CAG CCC GCA GTC CTG GCG GTG GTG AAC CTG-3' (sense) and 5'-CAG GTT CAC CAC CGC CAG GAC TGC GGG CTG CCC CTT ACC-3' (antisense) (corresponding to the S478A/Q481A mutation). Finally, we constructed a rFII molecule with a triple mutation, rFII^{SLQ→AAA} with the mutagenic primers 5'-GGT AAG GGG CAG CCC GCA GTC GCG GCG GTG GTG AAC CTG-3' (sense) and 5'-CAG GTT CAC CAC CGC GCG GAC TGC GGG CTG CCC CTT ACC-3' (antisense) (corresponding to the S478A/L480A/Q481A mutation) and an rFII molecule with Ser⁴⁷⁸/Leu⁴⁸⁰/Gln⁴⁸¹ deleted (rFII^{ΔSLQ}) using the mutagenic primers 5'-GGT AAG GGG CAG CCC GTC GTG GTG AAC CTG CCC-3' (sense) and 5'-GGG CAG GTT CAC CAC GAC GGG CTG CCC CTT ACC-3' (antisense). All deletions and point mutations were confirmed by DNA sequencing (DNA Analysis Facility, Department of Molecular Cardiology, The Lerner Research Institute, Cleveland Clinic).

Expression of Wild Type and Mutant rFII Molecules in Mammalian Cells—rFII expression in baby hamster kidney (BHK-21) cells has been described previously in detail (63). Briefly, BHK-21 cells were maintained in Dulbecco's modified Eagle's medium (DMEM) supplemented with fetal bovine serum (10%), and a streptomycin/penicillin (1%) mixture. Isolated plasmids (4–6 μg) for wild type and mutant rFII molecules were transfected into the BHK-21 cells using a lipid-based transfection reagent, Lipofectamine (Invitrogen), according to the manufacturer's instructions. Following an incubation period of 48 h, DMEM was supplemented with fetal bovine serum (10%), streptomycin/penicillin (1%) mixture, and methotrexate (1 μM) and added to the cells. After 3 weeks of treatment with the selection medium, colonies were isolated, grown, and screened for levels of rFII expression by Western blot analysis using a monoclonal antibody and compared with plasma-derived FII as a standard (1 μg/ml). Identification of the highest secreting rFII clone was further used in large scale protein expression with serum-free Opti-MEM supplemented with ZnCl₂ (50 μM), vitamin K₁ (10 μg/ml), and penicillin/strepto-

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mycin/Fungizone (1% v/v) mixture, and the medium were collected every 2 days for 2–3 weeks. Following collection, the media were stored at -80°C until the desired amount (usually 4 liters) was obtained and used for purification.

Purification of rFII Molecules—Purification of rFII was performed through a well established protocol previously described in detail (63). Briefly, collected media were thawed, filtered ($0.45\ \mu\text{m}$), and loaded on a tandem column setup of Amberlite XAD₂ and Q-Sepharose. Following the complete addition of medium to the two columns, the Q-Sepharose column was separated and washed with TBS ($0.02\ \text{M}$ Tris, $150\ \text{mM}$ NaCl, pH 7.4). The bound material containing rFII on the Q-Sepharose was eluted with $0.02\ \text{M}$ Tris, $0.5\ \text{M}$ NaCl, pH 7.4. The material was treated with barium citrate, and the isolated pellet was dissolved in a minimum volume of EDTA ($0.5\ \text{M}$, pH 7.4). The dissolved pellet was dialyzed twice in fresh TBS (two times, 4 liters) and filtered ($0.45\ \mu\text{m}$) prior to being loaded onto a General Electric FPLC instrument, equipped with a strong anionic exchanger Mono Q 5/50 column. The column was equilibrated in TBS, and a stepwise gradient of calcium (0 – $50\ \text{mM}$) in TBS was used to isolate fully γ -carboxylated rFII. Tubes containing the rFII molecules were concentrated using an appropriate Millipore Centricon (Billerica, MA), and aliquots were frozen at -80°C to avoid repeated freeze-thaw cycles. Following purification and before any experiment, all rFII molecules were characterized as extensively described below.

The level of γ -carboxylation of all rFII molecules was determined at the Protein Chemistry Facility, Texas A&M University, by alkaline hydrolysis followed by amino acid analysis as described (64, 65). All purified molecules were found to be properly carboxylated (Table 1). To verify that rFII^{WT} and rFII ^{$\Delta 473-487$} are processed at the appropriate cleavage sites when incubated with prothrombinase or fXa alone and produced the expected fragments, the recombinant proteins were incubated with PCPS vesicles and fXa in the presence and absence of fVa. Following gel electrophoresis, fragments were identified following NH_2 -terminal sequencing from PVDF membranes (see below). All fragments derived from the recombinant FII molecules have the expected NH_2 -terminal sequence following cleavage by either prothrombinase or membrane-bound fXa alone (data not shown).

The fact that the rFII ^{$\Delta 473-487$} molecule contains a 15-amino acid deletion was verified by cDNA sequencing. However, in view of the surprising and unexpected data presented herein, it was important to confirm the existence of the deletion in the purified recombinant protein. This was accomplished by mass spectrometry. Briefly, following activation of rFII ^{$\Delta 473-487$} and FII^{plasma} by prothrombinase, aliquots were analyzed in triplicate under reducing conditions on an SDS-12% PAGE. Following staining/destaining, the B-chain of IIa was excised from the gel, and the proteins were reduced and alkylated with iodoacetamide. Digestion (in gel) was accomplished with porcine trypsin. Analysis of the resulting peptides was performed with an α -cyanohydroxycinnamic acid (matrix) Kratos Axima CFR MALDI-TOF mass spectrometer (reflector mode; 25,000 accelerating voltage) in the Protein Chemistry Laboratory, Texas A & M University, under the direction of Dr. Larry Dangott. The data obtained were compared with the peptide map following

digestion of the B chain of IIa obtained from ExPASy/Swiss-Prot and verified the existence of the deletion in rFII ^{$\Delta 473-487$} . Similar experimental work performed with some other mutant molecules demonstrated that the rFII molecules described herein are fully carboxylated, can be appropriately processed by prothrombinase and fXa alone, and do indeed contain the expected deletion/mutations.

Gel Electrophoresis, Western Blotting, and Amino Acid Sequence from PVDF Membranes—SDS-PAGE was performed according to the method of Laemmli (66), using 9.5% gels following reduction with 2% β -mercaptoethanol. Screening for high levels of rFII-secreting clones was performed by Western blotting using PVDF according to a modified method initially described by Towbin *et al.* (67). Successfully transferred proteins were visualized by chemiluminescence using ECL Plus reagents following incubation with a polyclonal antibody specific to prothrombin-1. In some experiments, proteins were transferred to PVDF membranes and stained with Coomassie Blue, and NH_2 -terminal sequencing analysis was performed at the Biomolecular Resource Facility at the University of Texas Medical Branch (Galveston TX) as described previously in detail by our laboratory (68).

Studies of the Pathway for FII Activation by Gel Electrophoresis—The investigation of the activation rates of plasma-derived and of all rFII molecules, cleavage and activation by fXa alone or prothrombinase was performed according to a protocol previously described by our laboratory using plasma-derived FII or rFII (47, 49, 69). The calculation of the rates of all FII molecules consumption by fXa alone or by prothrombinase were performed as described previously with the software Prizm (GraphPad) (47, 49, 69).

Kinetic Titrations of Prothrombinase—To investigate the kinetic constants (K_m and k_{cat}) of prothrombinase, assays with a set amount of plasma-derived fVa and fXa (as described in the legend to the figures) were executed as described by our laboratory in many instances (47, 49, 69, 70). The initial rate of IIa generation was analyzed with the software Prizm (GraphPad) according to the Michaelis-Menten equation, and all final numbers reported are derived directly from the graphs. Each experiment used to report final numbers was run at least in duplicate, and the goodness of fit (R^2) for every model tested is provided under the “Results.” The change in transition-state stabilization free energy, which measures the effect of the mutations in rFII, was calculated for the double and triple mutants as extensively detailed in the literature and previously reported by our laboratory (71–77).

Recombinant Thrombin Activity—rFII molecules were converted to rIIa by $1\ \text{nM}$ prothrombinase. Full conversion of rFII to rIIa under these conditions was verified by gel electrophoresis. The chromogenic substrate S-2238 was used to assess rIIa activity by employing serial dilutions of the enzyme in Tris-NaCl buffer in the presence of 0.1% PEG 8000. The final concentrations of S-2238 used in the reactions were 0.94, 1.87, 3.75, 7.50, 15, and $60\ \mu\text{M}$. The reaction was started by the addition of $4\ \text{nM}$ rIIa. The data were obtained at 1 min using a SpectraMax M2 plate reader (Molecular Devices). The optical density was automatically adjusted for a 1-cm pathlength, and the V_{max} was calculated from the optical density using the established extinc-

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tion coefficient of S-2238 at room temperature (78) following plotting of the data to the Michaelis-Menten equation using the software Prizm (GraphPad).

Activation of fV and fVIII by rIIa—rFII molecules were converted to rIIa by 1 nM prothrombinase. Full conversion of rFII to rIIa under the conditions described was assessed by gel electrophoresis. The resulting wild type and mutant rIIa were assessed for their ability to cleave and activate the cofactors over time, by SDS-PAGE. Reaction mixtures containing either 500 nM plasma-derived human fV or recombinant human fVIII were diluted in Tris-NaCl buffer in the presence of Ca²⁺. The final concentration of rIIa in the mixtures was 4 nM.

Activation of Protein C by Plasma-derived IIa or rIIa—rFII molecules described herein were converted to IIa by 1 nM prothrombinase. Full conversion of rFII to rIIa under the conditions described was assessed by gel electrophoresis. The resulting IIa molecules were assessed for their ability to cleave and activate protein C in the presence of thrombomodulin and PCPS vesicles according to a procedure previously described (79) in Tris-buffered saline with Ca²⁺. Protein C activation was verified by SDS-PAGE under reducing conditions. The final concentration of IIa in all mixtures was 8 nM. Gels were stained with Coomassie Brilliant Blue.

FII Clotting Assay—To assess the function of all FII molecules in whole plasma, a clotting assay using FII-deficient plasma was employed. The clotting assay was performed as described previously (80), and the time needed for formation of a fibrin clot was monitored at 37 °C using a Diagnostica Stago STart® 4 hemostasis analyzer as described previously (80). The analyzer was set up to automatically measure the time to clot up to 120 s.

Structural Analysis—To evaluate the structural features of the Ser⁴⁷⁸, Leu⁴⁸⁰, and Gln⁴⁸¹ residues, crystal structures of FII and IIa were superimposed and compared. The three human FII crystal structures that have been reported, show similar conformations for the residues of interest and neighboring regions; the highest resolution of these structures was chosen for detailed analysis (38). From the many human IIa crystal structures that are available, several representative examples in different bound states were compared and found to have similar conformations for the region containing the residues of interest. A high resolution structure of unbound IIa was chosen as the representative structure for detailed analysis (81). The program COOT was used to inspect structural features and determine distances (82). AREAIMOL (83–85) was used to calculate the solvent-accessible surface areas for specific residues, and molecular figures were prepared with the PyMOL Molecular Graphics System, version 1.5.0.4 (Schrödinger, LLC).

Results

rFII Expression—To evaluate the minimal amino acid sequence necessary for the fVa-dependent FII activation by prothrombinase within the 473–487 critical amino acid stretch of FII, we stably transfected BHK-21 cell lines according to a previously well defined protocol (63) with several constructs. Mutant rFII molecules prepared were as follows: rFII^{Δ473–487} (missing residues ⁴⁷³GKGQPSVLQVVNLPI⁴⁸⁷), rFII^{ΔN10} (missing amino acid residues ⁴⁷³GKGQPSVLQV⁴⁸²), and

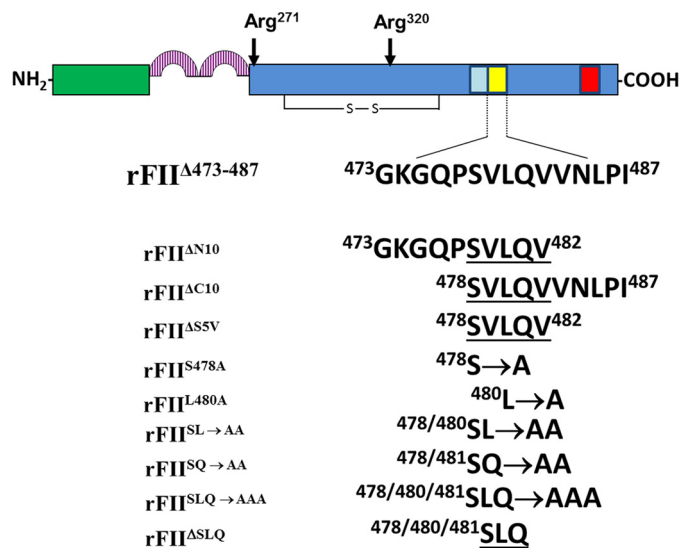


FIGURE 1. Schematic of FII. FII is converted to IIa through two fXa-catalyzed cleavages at Arg²⁷¹ and at Arg³²⁰ resulting in IIa formation. The red rectangle denotes the fVa-independent site for fXa on FII (57), and the yellow rectangle represents the fVa-dependent site for fXa (56, 95) studied herein. The light blue rectangle denotes the amino acids composing (pro)exosite I (50). All mutants created, stably transfected, purified to homogeneity, and used in the study are shown together with their assigned name used throughout this work.

rFII^{ΔC10} (missing amino acid residues ⁴⁷⁸SVLQVVNLPI⁴⁸⁷) (Fig. 1). These three mutant molecules have five amino acids in common (⁴⁷⁸SVLQV⁴⁸²). We thus proceeded to construct, stably express, and purify to homogeneity rFII^{ΔS5V} (a rFII molecule missing amino acids ⁴⁷⁸SVLQV⁴⁸²) (Fig. 1). Preliminary experiments with rFII^{ΔS5V} demonstrated that the mutant molecule had no clotting activity similarly to rFII^{Δ473–487}, rFII^{ΔN10}, and rFII^{ΔC10}. We next proceeded to make single, double, and triple alanine substitutions within this significant region. The following recombinant mutant molecules were made: rFII^{S478A} (S478A), rFII^{L480A} (L480A), rFII^{SL→AA} (S478A/L480A), rFII^{SQ→AA} (S478A/Q481A), and rFII^{SLQ→AAA} (S478A/L480A/Q481A) (Fig. 1). It is important to note that rIIa^{S478A} (where rIIa^{S478A} is recombinant human α-thrombin with the mutation S478A) was previously tested and found to behave as rIIa^{WT} (5, 6). We have thus used this mutation as an internal control for all double and triple alanine substitutions of rFII. Finally, and to verify the results obtained with rFII^{SLQ→AAA}, we have also made an rFII molecule with these three amino acids deleted (rFII^{ASLQ}). In all experiments, results with the mutant molecules were compared with results obtained with rFII^{WT} or FII^{plasma}.

Prothrombin Time—The ability of FII and all rFII molecules to be activated under physiological conditions and to promote fibrin clot formation was first assessed using prothrombin times (PTs) (Fig. 2). The results shown in Table 1 demonstrate that although FII^{plasma}, rFII^{WT}, and rFII^{S478A} had comparable clotting times of 12.7, 12.2, and 11.7 s, respectively, rFII^{L480A} (where rIIa^{L480A} is recombinant human α-thrombin with the mutation L480A) exhibited a minimal but significant prolonged PT of ~30 s (Fig. 2). Surprisingly, although rFII^{SL→AA} (where rFII^{SL→AA} is recombinant human prothrombin with the mutation S478A/L480A) and rFII^{SQ→AA} had slow but compa-

Function of Sequence Leu⁴⁸⁰–Gln⁴⁸¹ of Prothrombin

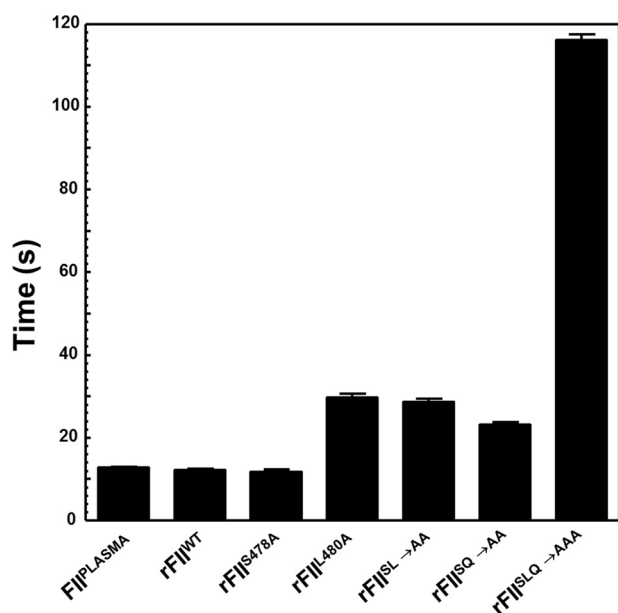


FIGURE 2. **Clotting activity of all forms of FII.** The average clotting time found in four different measurements in FII-deficient plasma is shown for all FII/rFII molecules identified at the bottom of the graph.

rable PTs of ~30 s, the triple mutant rFII^{SLQ->AAA} was severely ineffective in fibrin clot formation (PT ~116 s), whereas rFII^{ΔSLQ} (where rFII^{ΔSLQ} is recombinant human prothrombin with amino acids Ser⁴⁷⁸/Leu⁴⁸⁰/Gln⁴⁸¹ deleted) had a PT around 140 s (data not shown). In contrast, rFII^{Δ473–487}, rFII^{ΔN10}, rFII^{ΔC10}, and rFII^{ΔS5V} were unable to induce clotting under the conditions described. These functional data demonstrate that either rFII^{SLQ->AAA} or rFII^{ΔSLQ} cannot get activated to rIIa in a timely fashion, or that rIIa^{SLQ->AAA} and/or rIIa^{ΔSLQ} formed are catalytically impaired because of the mutations, or both. Because previous data have shown that the S478A transition in IIa is of no consequence for its chromogenic and proteolytic activity (5, 6), overall these results demonstrate for the first time that both Leu⁴⁸⁰ and Gln⁴⁸¹ have a profound effect on IIa generation and/or IIa activity during fibrin clot formation or both.

Activation of rFII Molecules—To ascertain the effect of region 473–487 of FII on its ability to be activated by membrane-bound fXa alone, in the absence of fVa, we assessed the pattern of activation by gel electrophoresis over a 2-h time period (Fig. 3). Fig. 3A shows a control experiment and demonstrates that FII^{PLASMA} activation by membrane-bound fXa alone proceeds following initial cleavage at Arg²⁷¹, through the intermediate prethrombin-2 with very slow gradual appearance of the B-chain of IIa because of inefficient rate of cleavage at Arg³²⁰. Surprisingly, with the removal of amino acids 473–487 from prothrombin (Fig. 3B), there is acceleration of rFII^{Δ473–487} consumption through initial cleavage at Arg²⁷¹ that is evident by the prompt appearance of prethrombin-2. Additional examinations of the intensity of the B-chain of thrombin reveal a substantially delayed cleavage at Arg³²⁰ of rFII^{Δ473–487} compared with rFII^{WT} resulting in insignificant IIa generation. Scanning densitometry of the gels shown in Fig. 3, A and B, showed that the rate of rFII^{Δ473–487} consumption by membrane-bound fXa is ~4-fold increased compared with the rate of cleavage of

rFII^{PLASMA} under similar experimental conditions (Table 1). These data suggest that amino acid sequence 473–487 of prothrombin provides a potential obstruction for efficient initial cleavage of prothrombin at Arg²⁷¹ by membrane-bound fXa alone in the absence of fVa.

To further improve our understanding of the fundamental role of amino acid region 473–487 for FII activation by prothrombinase, we studied the pattern of FII activation by fully assembled prothrombinase with gel electrophoresis over a 2-h time period. A control experiment (Fig. 3C) demonstrates that under the conditions used the activation of FII^{PLASMA} proceeds efficiently following initial cleavage at Arg³²⁰, through the enzymatically active intermediate meizothrombin, as confirmed by the appearance of fragment 1·2-A. Rapid cleavage of this fragment at Arg²⁷¹ leads to the formation of IIa. In contrast, activation of rFII^{Δ473–487} under similar experimental conditions is significantly delayed through the same pathway as verified by the late appearance of the B-chain of IIa (Fig. 3D). Scanning densitometry of the gels shown in Fig. 3, C and D, showed that rFII^{Δ473–487} is consumed with a rate that is ~27-fold slower compared with the rate of FII^{PLASMA} consumption or ~23-fold slower compared with the rate of rFII^{WT} consumption under the experimental conditions used (Table 1). These data suggest that under conditions of saturating amounts of fVa with respect to fXa, amino acid sequence 473–487 of FII plays a preeminent role because it is required for fast and efficient initial cleavage of FII at Arg³²⁰ by prothrombinase.

To further investigate the effect of the deletions and point mutations on rFII cleavage and activation by membrane-bound fXa alone, we studied rFII activation by gel electrophoresis of all mutants detailed in Fig. 1. Fig. 4A shows a control experiment and demonstrates that rFII^{WT} activation by membrane-bound fXa proceeds typically following initial cleavage at Arg²⁷¹, as its plasma counterpart through the intermediate prethrombin-2 with very slow and minimal appearance of the B chain of IIa because of a nonproductive rate of cleavage at Arg³²⁰. With the removal of amino acids 478–482 from rFII^{ΔS5V} (Fig. 4B), there is acceleration of rFII^{Δ478–482} consumption by fXa alone through initial cleavage at Arg²⁷¹ that is evident by the rapid appearance of prethrombin-2. The fact that only trace amounts of B-chain of IIa are apparent under the conditions employed suggests a substantially deferred rate of cleavage at Arg³²⁰ of the deletion mutant compared with cleavage of rFII^{WT} resulting in insignificant amounts of IIa generation. Scanning densitometry of similar gels shown in Fig. 4, A and B, showed that the rate of consumption of all rFII molecules by membrane-bound fXa alone is ~2.3–8-fold increased compared with the rate of cleavage of rFII^{WT} or FII^{PLASMA} (Fig. 4C and Table 1). However, although with rFII^{S478A}, rFII^{L480A}, rFII^{SL->AA}, and rFII^{SQ->AA} minimal amounts of the B-chain of IIa are formed (data not shown), when studying rFII^{ΔN10}, rFII^{ΔC10}, rFII^{ΔS5V}, rFII^{SLQ->AAA}, and rFII^{ΔSLQ} activation, there is accumulation of prethrombin-2 with no significant amounts of B-chain of IIa generated suggesting impaired cleavage of prethrombin-2 at Arg³²⁰ by membrane-bound fXa (Fig. 4, B and D). These data confirm our findings with rFII^{Δ473–487} (Fig. 3) and reveal that the dipeptide Leu⁴⁸⁰–Gln⁴⁸¹ within the 15-amino acid stretch 473–487 of FII appears to be responsible for the sim-

TABLE 1

Physical properties, clotting times, and rate of cleavage of various rFII molecules in the presence of a fixed concentration of membrane-bound fXa or in the presence of prothrombinase

Mutant	mol of Gla/mol of protein	Clotting time ^a	Rate of FII molecule cleavage by membrane-bound fXa alone ^b (nM FII consumed·s ⁻¹ ·fXa ⁻¹)	Rate of FII molecules cleavage by prothrombinase ^b (nM FII consumed·s ⁻¹ ·fXa ⁻¹)
FII ^{plasma}	10 ± 1	12.7 ± 0.12 ^s	0.13 ± 0.03 (0.94) ^c	27.5 ± 4.2 (0.99) ^c
rFII ^{WT}	9.1 ± 0.9	12.2 ± 0.16	0.1 ± 0.012 (0.99)	22.8 ± 3.8 (0.98)
rFII ^{Δ473-487}	9.5 ± 0.1	>120 ^d	0.53 ± 0.013 (0.99)	1.0 ± 0.26 (0.92)
rFII ^{ΔN10}	12.0 ± 1	>120 ^d	0.7 ± 0.03 (0.98)	1.5 ± 0.34 (0.94)
rFII ^{ΔC10}	12.5 ± 1	>120 ^d	0.35 ± 0.04 (0.99)	1.3 ± 0.09 (0.99)
rFII ^{ΔS5V}	11 ± 1	>120 ^d	0.8 ± 0.03 (0.99)	1.8 ± 0.12 (0.99)
rFII ^{S478A}	10.8 ± 1	11.75 ± 0.32	0.25 ± 0.015 (0.99)	24.9 ± 4.6 (0.97)
rFII ^{L480A}	10.1 ± 1	29.8 ± 0.41	0.6 ± 0.02 (0.99)	28.8 ± 2.8 (0.98)
rFII ^{SL→AA}	9.5 ± 0.9	28.7 ± 0.35	0.63 ± 0.03 (0.99)	18.2 ± 3.7 (0.99)
rFII ^{SQ→AA}	10 ± 1	23.2 ± 0.3	0.23 ± 0.025 (0.98)	35.9 ± 3.8 (0.99)
rFII ^{SLQ→AAA}	10.6 ± 1	116.2 ± 0.65	0.6 ± 0.04 (0.99)	1.23 ± 0.12 (0.98)
rFII ^{ΔSLQ}	10.6 ± 1.1	>120 ^d	0.22 ± 0.04 (0.95)	1.84 ± 0.15 (0.99)

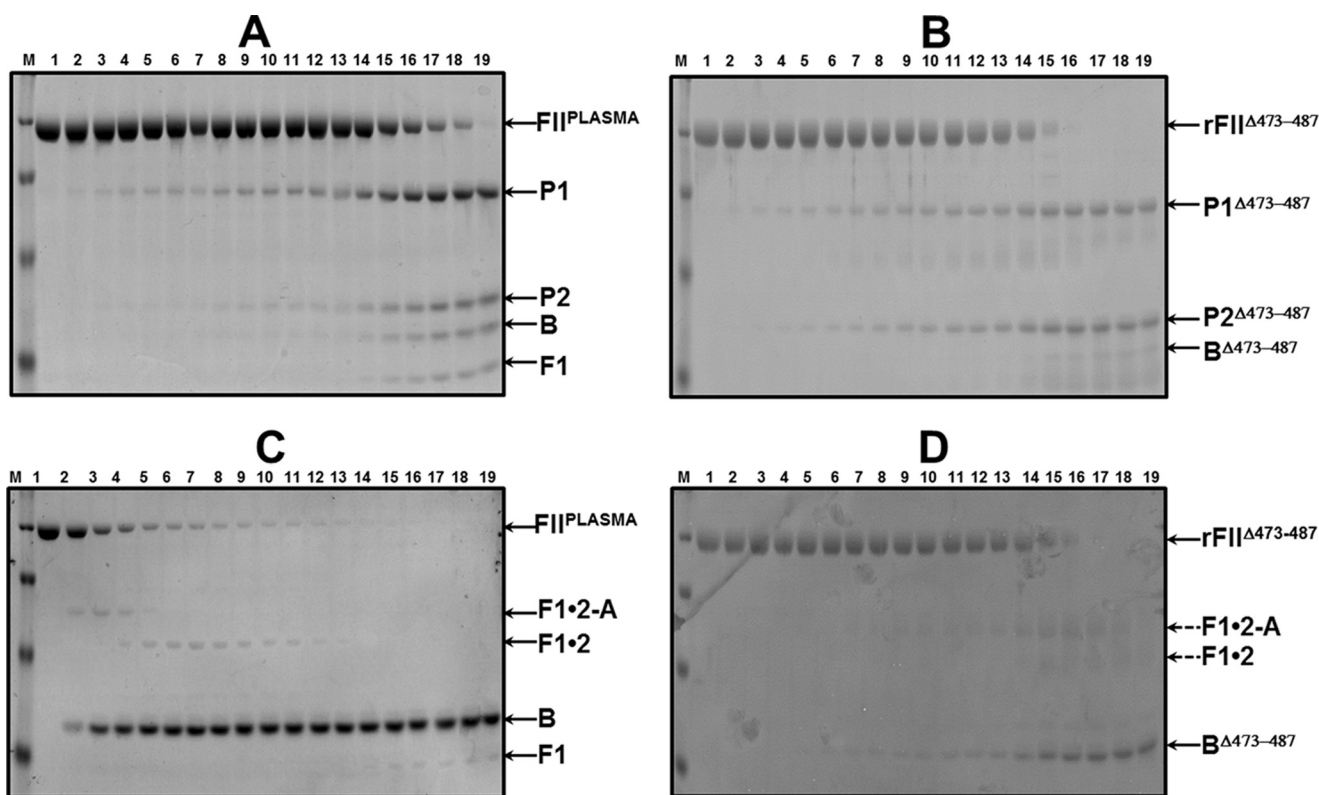
^a Clotting times were determined using FII-deficient plasma as described under "Experimental Procedures" in quadruplicate.^b The rates of rFII consumption were obtained following scanning densitometry of gels studying rFII activation. Some of the gels used are shown in Figs. 3–5. The final rate of rFII consumption in the presence of membrane-bound fXa or prothrombinase was calculated using the apparent first-order rate constant, k (s⁻¹), obtained directly from the graph following plotting of the data as described under "Experimental Procedures."^c The numbers in parentheses represent the goodness of fit (R^2) to the equation representing first-order exponential decay using the software Prism from where the first-order rate constant was obtained.^d No clotting time could be detected following a 120-s incubation time period.

FIGURE 3. SDS-PAGE analyses of FII^{plasma} and rFII^{Δ473-487} activation by membrane-bound fXa alone or prothrombinase. A, rFII^{plasma} (1.4 μM) in the presence of PCPS vesicles, DAPA, and membrane-bound fXa alone (5 nM). B, rFII^{Δ473-487} (1.4 μM) in the presence of PCPS vesicles, DAPA, and membrane-bound fXa alone (5 nM). C, rFII^{plasma} (1.4 μM) in the presence of PCPS vesicles, DAPA, and prothrombinase (1 nM fXa and 20 nM fVa). D, rFII^{Δ473-487} (1.4 μM) in the presence of PCPS vesicles, DAPA, and prothrombinase (1 nM fXa and 20 nM fVa). Aliquots were withdrawn at various time intervals and treated as described (47, 69). M represents the lane with molecular weight markers (from top to bottom): 98,000, 64,000, 50,000, and 36,000, respectively. Lanes 1–19 show samples from the reaction mixture before (0 min) the addition of fXa and 20, 40, 60, 80, 100, 120, 150, 180, 210, and 240 s and 5, 6, 10, 20, 30, 60, 90, and 120 min, respectively, after the addition of fXa. Following scanning densitometry as described under "Experimental Procedures," the data representing FII consumption as a function of time (s) were plotted using non-linear regression analysis according to the equation representing a first-order exponential decay and the rates of FII consumption using the apparent first-order rate constant, k (s⁻¹), obtained directly from the fitted data, were calculated as described (47) and are reported in Table 1. FII-derived fragments are identified to the right of A–D as follows: FII, prothrombin (amino acid residues 1–579); P1, prothrombin-1 (amino acid residues 156–579); F1·2-A, fragment 1·2-A chain (amino acid residues 1–320); F1·2, fragment 1·2 (amino acid residues 1–271); P2, prothrombin-2 (amino acid residues 272–579); B, B-chain of IIa (amino acid residues 321–579); F1, fragment 1 (amino acid residues 1–155).

ilar effects observed with rFII^{ΔN10}, rFII^{ΔC10}, rFII^{ΔS5V}, rFII^{SLQ→AAA}, and rFII^{ΔSLQ} when studying rFII molecular activation by membrane-bound fXa alone in the absence of fVa (Fig. 4D and Table 1).

To improve our understanding of the essential role of amino acids Leu⁴⁸⁰ and Gln⁴⁸¹ for FII activation, we studied the pattern of all rFII molecules activation shown in Fig. 1 by fully assembled prothrombinase (*i.e.* in the presence of an excess of

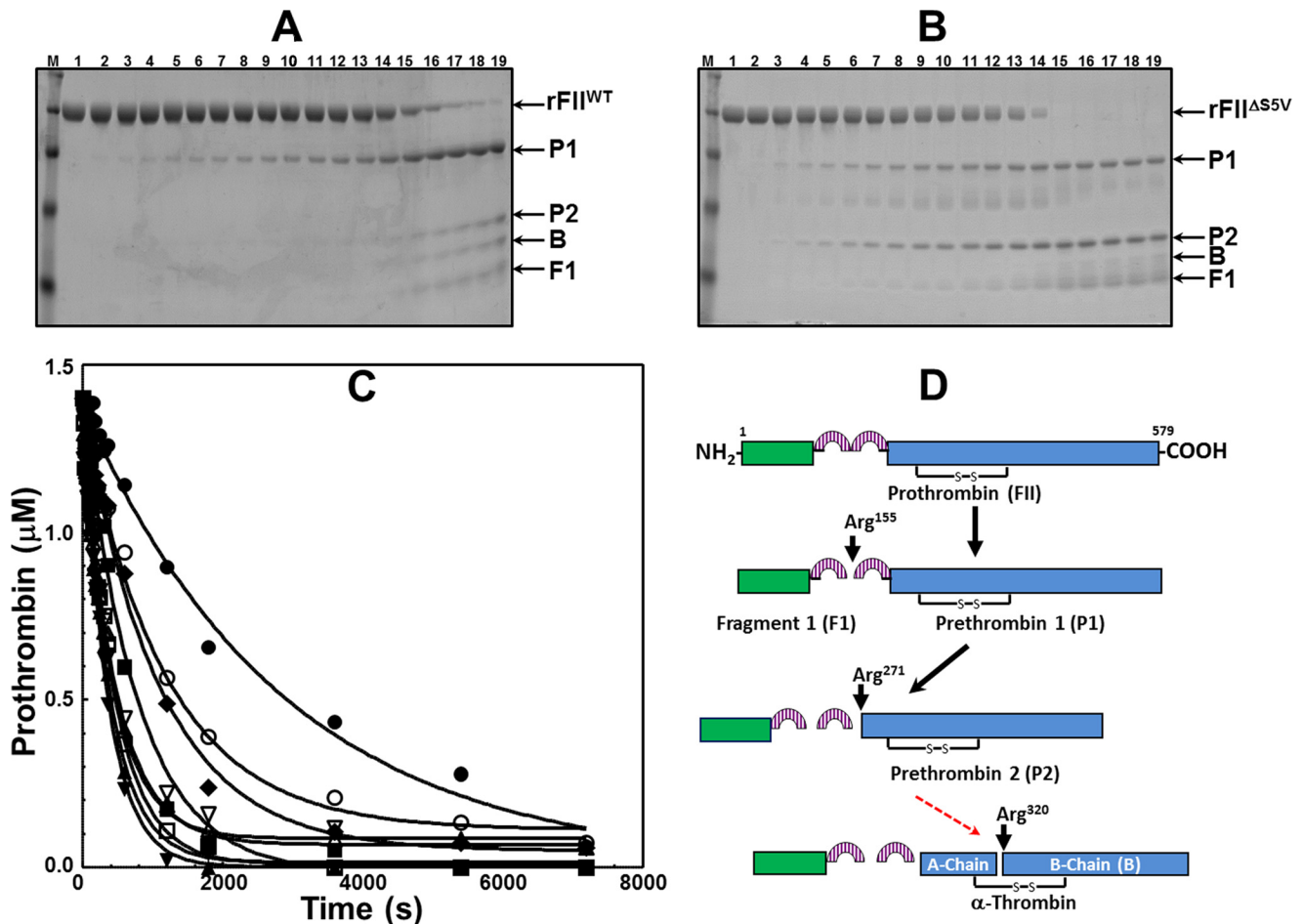


FIGURE 4. SDS-PAGE analyses of rFII molecule activation by membrane-bound fXa alone. *A*, rFII^{WT} (1.4 μM) in the presence of PCPS vesicles, DAPA, and membrane bound fXa alone (5 nM); *B*, rFII^{Δ55V} (1.4 μM) in the presence of PCPS vesicles, DAPA, and membrane-bound fXa alone (5 nM). Aliquots were withdrawn at various time intervals and treated as described (47, 69). *M* represents the lane with molecular weight markers (from top to bottom): 98,000, 64,000, 50,000, and 36,000, respectively. *Lanes 1–19* show samples from the reaction mixture before (0 min) the addition of fXa and 20, 40, 60, 80, 100, 120, 150, 180, 210, and 240 s and 5, 6, 10, 20, 30, and 60, 90, and 120 min respectively, after the addition of fXa. *C*, the two gels shown in *A* and *B* together with similar gels obtained with all rFII studied were scanned, and rFII consumption was recorded as described under “Experimental Procedures.” Following scanning densitometry and normalization to the initial FII concentration, the data representing rFII consumption as a function of time (seconds) were plotted using non-linear regression analysis according to the equation representing a first-order exponential decay using the software Prizm (GraphPad, San Diego). Prothrombinase was assembled with rFII^{WT} (filled circles; *R*² 0.98), rFII^{ΔC10} (filled squares; *R*² 0.98), rFII^{ΔN10} (filled triangles; *R*² 0.99), rFII^{Δ55V} (filled inverse triangles; *R*² 0.99), rFII^{S478A} (filled diamonds; *R*² 0.99), rFII^{L480A} (open squares; *R*² 0.99), rFII^{SQ→AA} (open circles; *R*² 0.98), rFII^{SL→AA} (open triangles; *R*² 0.99), and rFII^{SLQ→AAA} (open inverse triangles; *R*² 0.99). The rates of rFII consumption shown in *C*, using the apparent first-order rate constant, *k* (s⁻¹) obtained directly from the fitted data, were calculated as reported (47), and the data are shown in Table 1. *D*, schematic representation of fragments derived following FII activation by membrane-bound fXa alone. The red arrow indicates impaired cleavage at Arg³²⁰ in rFII^{ΔC10} (filled squares), rFII^{ΔN10} (filled triangles), rFII^{Δ55V} (filled inverse triangles), and rFII^{SLQ→AAA} (open inverse triangles) resulting in prethrombin-2 accumulation. FII-derived fragments are identified to the right of each panel, according to the description provided in the legend of Fig. 3.

fVa) by gel electrophoresis over a 2-h time period (Fig. 5). A control experiment (Fig. 5A) demonstrates that under the conditions used, rFII^{WT} proceeds as its plasma counterpart following initial cleavage at Arg³²⁰, through the enzymatically active intermediate meizothrombin, as confirmed by the appearance of fragment 1·2-A. Rapid cleavage of this fragment at Arg²⁷¹ leads to the formation of rIIa. Similar results were found when using rFII^{S478A} (Fig. 5B) demonstrating that the S478A transition alone is of no consequence for timely FII activation by prothrombinase. In contrast, activation of rFII^{Δ55V} and rFII^{SLQ→AAA} under similar experimental conditions was significantly delayed through the same pathway as verified by the lingering of fragment 1·2-A at the late time points and the late appearance of the B-chain of rIIa (Fig. 5, C and D). Similar results were obtained with rFII^{ΔSLQ} (Table 1). A systematic analysis of the activation of all rFII mutant molecules by pro-

thrombinase using similar experimental procedures, followed by scanning densitometry of the gels and calculation of the rate of rFII consumption, revealed the existence of two groups as follows: a group of molecules represented by FII^{plasma}, rFII^{WT}, and rFII^{S478A} (also containing rFII^{L480A}, rFII^{SL→AA}, and rFII^{SQ→AA}) that are efficiently activated by prothrombinase; and a second group of proteins represented by rFII^{Δ55V} and rFII^{SLQ→AAA} (including rFII^{ΔN10}, rFII^{ΔC10}, and rFII^{ΔSLQ}) that are activated by fully assembled prothrombinase with a rate that is ~13–18-fold slower than the rate observed with rFII^{WT} (Fig. 6A, inset, and Table 1). The data suggest that under conditions of saturating amounts of fVa with respect to fXa, the dipeptide Leu⁴⁸⁰-Gln⁴⁸¹ of prothrombin plays a leading role during FII activation because it is required for fast and efficient initial cleavage at Arg³²⁰ by prothrombinase (Fig. 6B, the deficient step is represented by the red arrow).

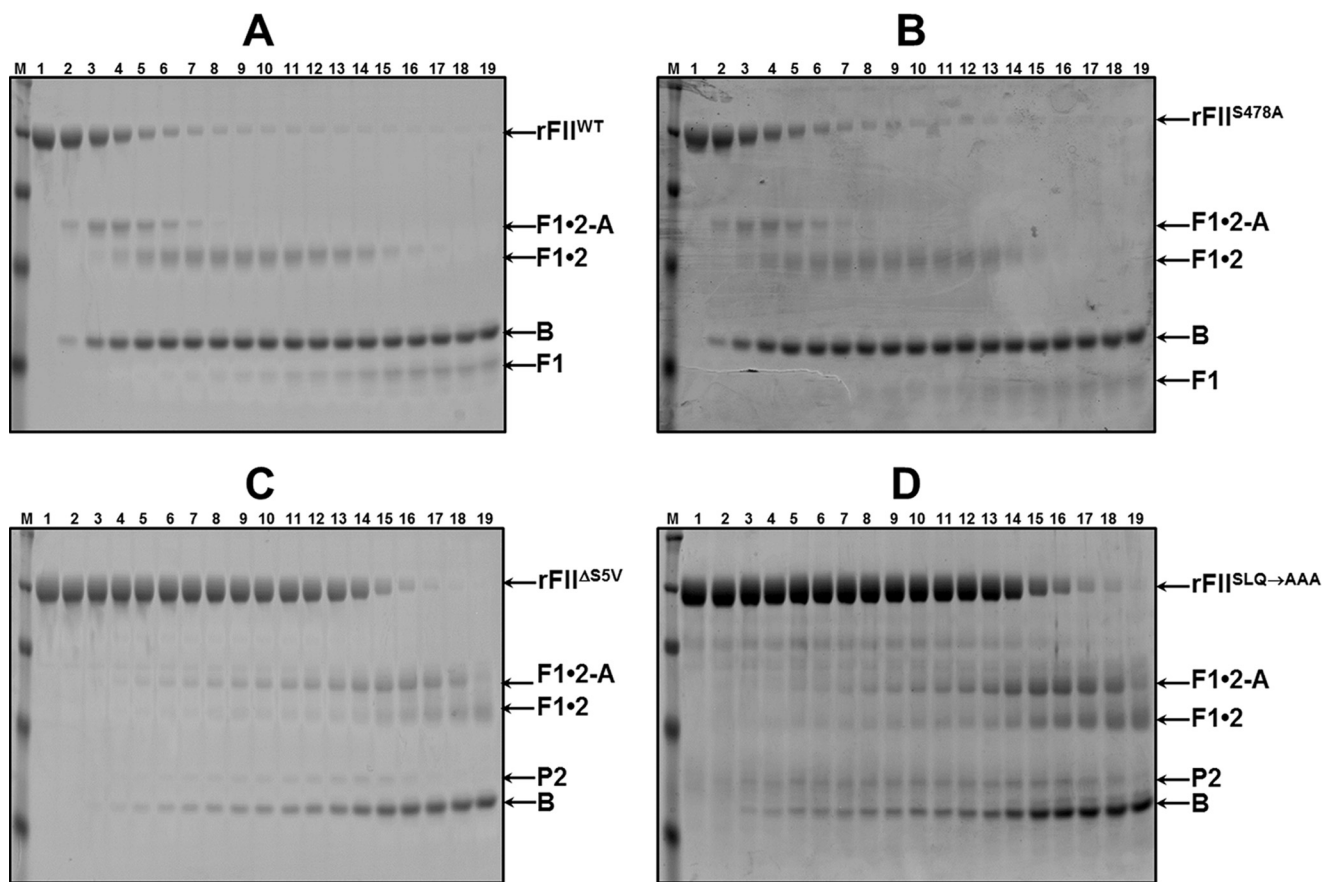


FIGURE 5. SDS-PAGE analyses of rFII molecule activation by prothrombinase. A, rFII^{WT} (1.4 μ M) in the presence of PCPS vesicles, DAPA, and prothrombinase (1 nM fXa and 20 nM fVa). B, rFII^{S478A} same conditions as in A. C, rFII^{ΔS5V} same conditions as in A. D, rFII^{SLQ→AAA} same conditions as in A. Aliquots were withdrawn at various time intervals and treated as described (47, 69). Same time points as described in the legend to Fig. 4. FII-derived fragments are identified to the right of each panel, according to the description provided in the legend of Fig. 3.

Kinetic Analyses of the Activation of rFII Molecules—To understand the effect of the S478A/L480A/Q481A substitutions on the activity of prothrombinase in activating the rFII molecules, we first examined the rates of rIIa formation from all rFII molecules under similar experimental conditions. Historically, this method was designed to identify any deficiency in fVa or fXa as part of prothrombinase in cleaving and activating FII and is measured indirectly by using IIa generation as a reporting probe with a chromogenic substrate. The comprehensive kinetic data for several mutants are shown in Fig. 7 with the kinetic constants derived directly from the fitted data reported in Table 2. The combined findings demonstrate that whereas the single and double alanine substitutions rFII mutants are activated by prothrombinase similarly, providing kinetic constants comparable with the wild type or plasma FII molecules, kinetic analyses of prothrombinase activation of rFII^{SLQ→AAA} demonstrate a modest 2.7-fold decrease in the k_{cat} and a large 21-fold increase in the K_m value of the reaction. Similar experiments studying rFII^{ΔSLQ} activation by fully assembled prothrombinase revealed a 29-fold increase in K_m with a concomitant 16-fold decrease in the k_{cat} values of the reaction. A direct comparison between the data obtained with rFII^{SLQ→AAA} with the data obtained with rFII^{ΔSLQ} strongly suggest an important contribution of the backbone structure of the peptide bond between these three amino acids for efficient rFII activation by prothrombinase.

To quantify the interaction between the two sets of double mutations (S478A/L480A and S478A/Q481A) and to confirm their apparent synergistic detrimental effect on prothrombinase function for activation of rFII^{SLQ→AAA}, we have further calculated the difference in free energy of the transition state analog ($\Delta\Delta G_{int}$) for the triple mutant as described previously by our laboratory (75, 76). The large positive value of $\Delta\Delta G_{int}$ (+2.4 kcal/mol) for the combination of the mutations at Leu⁴⁸⁰ and Gln⁴⁸¹ together with the sizable 55-fold decrease in the second-order rate constant of prothrombinase for rFII^{SLQ→AAA} activation signify that there is a deficiency in recognition between prothrombinase and rFII^{SLQ→AAA}. These findings solidify our previous conclusion that these substitutions are detrimental to the activation of rFII bearing the triple amino acid substitution by fully assembled prothrombinase. However, it is important to note that it is also possible that rIIa^{SLQ→AAA} may also be deficient in its own catalytic activity as observed with rIIa^{ΔSLQ}, and the effect observed with rFII^{SLQ→AAA} activation may be likewise due to the deficiency of rIIa in cleaving the chromogenic substrate. Thus, although we cannot yet assign the poor performance of prothrombinase in cleaving rFII^{SLQ→AAA} solely to a deficiency in recognizing the mutated substrate, and because the S478A transition is of no consequence for either rFII^{S478A} activation or rIIa^{S478A} activity, the overall data presented thus far suggest that amino acid sequence Leu⁴⁸⁰–Gln⁴⁸¹ may have a dual effect in properly directing prothrombinase rec-

Function of Sequence Leu⁴⁸⁰–Gln⁴⁸¹ of Prothrombin

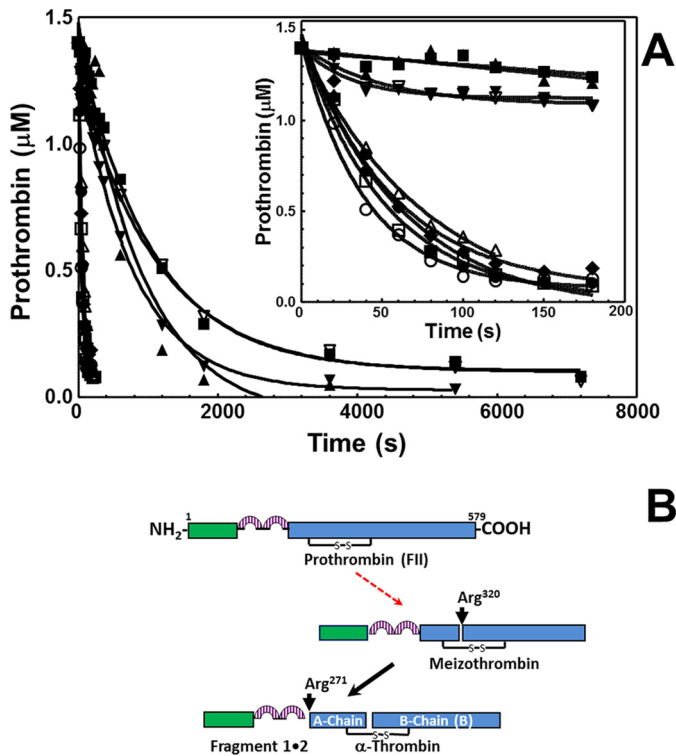


FIGURE 6. Analyses of the rates of activation of rFII by prothrombinase. *A*, gels shown in Fig. 5, together with similar gels obtained with all rFII studied, were scanned, and rFII consumption was recorded as described under “Experimental Procedures.” Following scanning densitometry, the numbers were normalized to the initial concentration of rFII studied, and the data representing rFII consumption as a function of time (seconds) were plotted using nonlinear regression analysis according to the equation representing a first-order exponential decay using the software Prizm (GraphPad, San Diego). rFII^{WT} (filled circles; R^2 0.98), rFII^{ΔC10} (filled squares; R^2 0.99), rFII^{ΔN10} (filled triangles; R^2 0.94), rFII^{ΔS5V} (filled inverse triangles; R^2 0.99), rFII^{S478A} (filled diamonds; R^2 0.97), rFII^{L480A} (open squares; R^2 0.98), rFII^{SQ→AA} (open circles; R^2 0.99), rFII^{SL→AA} (open triangles; R^2 0.99), and rFII^{SLQ→AAA} (open inverse triangles; R^2 0.99) are shown. The inset shows the progress of the reaction during the first 180 s. The rates of rFII consumption using the apparent first-order rate constant k (s^{-1}), obtained directly from the fitted data, were calculated as reported (47) and shown in Table 1. *B*, schematic representation of fragments derived following rFII activation by prothrombinase in the presence of excess fVa with respect to fXa. The red arrow indicates impaired cleavage (at Arg³²⁰) in rFII^{ΔC10} (filled squares), rFII^{ΔN10} (filled triangles), rFII^{ΔS5V} (filled inverse triangles), and rFII^{SLQ→AAA} (open inverse triangles).

ognition of FII, as well as providing the resulting enzyme with the appropriate surface required for proper substrate tethering and cleavage. However, it is also possible that these two amino acids are allosterically involved in both prothrombinase interactions with FII as well as the expression of the enzymatic activity of IIa.

Analyses of the Activity of rIIa Molecules—Although many investigations have identified the specific amino acid residues from FII/IIa participating in either prothrombinase recognition or IIa activity toward its physiological substrates, respectively, few studies have shown that identical residues are involved in both FII recognition by prothrombinase and IIa activity. To understand the effect of the deletions/mutations on IIa activity, we further assessed the amidolytic and biological activity of selected rIIa molecules generated herein toward the chromogenic substrate S-2238 and toward thrombin’s natural substrates, fV, fVIII, and protein C.

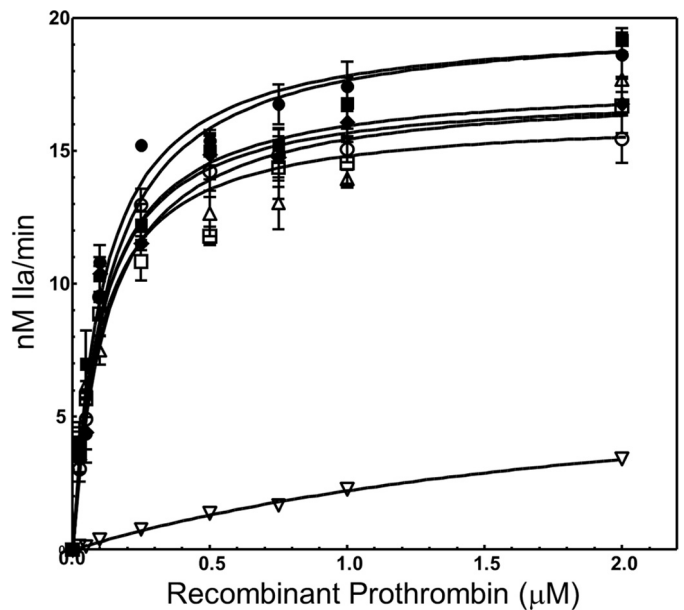


FIGURE 7. Determination of kinetic parameters of prothrombinase catalyzing cleavage and activation of various FII molecules. IIa generation experiments were conducted as described under “Experimental Procedures” by varying the substrate concentration and using a chromogenic substrate. The solid lines represent the nonlinear regression fit of the data using Prizm GraphPad software according to the Henri Michaelis-Menten equation ($V_o = V_{max} \cdot [FII] / (K_m + [FII])$) to yield the K_m and k_{cat} ($k_{cat} = V_{max} / E_{tot}$, where E_{tot} is the concentration of fully assembled prothrombinase, in this case is 10 pM, Table 2). Prothrombinase activity with various rFII molecules is shown as follows: rFII^{WT} filled circles; rFII^{PLASMA} filled squares; rFII^{S478A} filled diamonds; rFII^{L480A} open squares; rFII^{SL→AA} open triangles; rFII^{SQ→AA} open circles; rFII^{SLQ→AAA} open inverse triangles. Kinetic constants reported in the text and in Table 2 were extracted directly from the fitted data shown herein.

To understand the effect of the mutations on the amidolytic activity of IIa, we determined the kinetic constants for the hydrolysis of S-2238 by the rIIa molecules under steady state conditions. The data shown in Table 3 reveal the following: 1) rIIa^{WT} produced under the conditions described by our laboratory has similar activity as previously found with other rIIa^{WT} preparations, and 2) rIIa^{S478A} has similar catalytic efficiency (k_{cat}/K_m) as rIIa^{WT}, as demonstrated previously (6, 86). In addition, we also found that whereas rIIa^{SLQ→AAA} was devoid of activity toward S-2238, rIIa^{SQ→AA} (where rIIa^{SQ→AA} is recombinant human α-thrombin with the mutation S478A/Q481A) has similar amidolytic activity as rIIa^{WT}, whereas rIIa^{L480A} and rIIa^{SL→AA} (where rIIa^{SL→AA} is recombinant human α-thrombin with the mutation S478A/L480A) are the most deficient in S-2238 hydrolysis among the single and double alanine mutants when compared with rIIa^{WT} or rIIa^{S478A} (Table 3). The combined data clearly demonstrate that amino acid Leu⁴⁸⁰ plays an important role during the expression of IIa amidolytic activity and that the integrity of amino acid sequence Leu⁴⁸⁰–Gln⁴⁸¹ is required for optimal expression of this activity.

The data shown in Figs. 8 and 9 demonstrate that whereas rIIa^{WT} and rIIa^{S478A} cleave and activate fV and fVIII with similar rates (Figs. 8, A and C, and 9, A and C), rIIa^{ΔC10} (rIIa^{ΔC10} is recombinant human prothrombin missing amino acids SVLQVVNLPI), and rIIa^{ΔS5V} are totally deficient in cleaving both cofactor molecules over a 3-h incubation period (Figs. 8, B and H, and 9, B and H). These data are in complete agreement

TABLE 2

Kinetic constants of plasma FII and various rFII mutant molecules activation by prothrombinase

FII species	K_m^a	$k_{cat}^{a,b}$	R^2 /points/titrations ^c	k_{cat}/K_m	Decrease ^d
	μM	min^{-1}		$(M^{-1}\cdot s^{-1})\cdot 10^8$	-fold
FII ^{plasma}	0.13 ± 0.02	1992 ± 68	0.93/30/3	2.5	
rFII ^{WT}	0.11 ± 0.015	1976 ± 57	0.97/20/2	2.9	
rFII ^{WTe}	0.11 ± 0.02	2054 ± 84	0.97/10/1	3.1	
rFII ^{Δ473-487}	NP ^f				
rFII ^{ΔN10}	NP				
rFII ^{ΔC10}	NP				
rFII ^{ΔS5V}	NP				
rFII ^{S478A}	0.10 ± 0.013	1764 ± 45	0.97/20/2	2.9	1.0
rFII ^{L480A}	0.10 ± 0.015	1630 ± 49	0.96/20/2	2.7	1.1
rFII ^{SL→AA}	0.12 ± 0.02	1734 ± 65	0.92/30/3	2.4	1.2
rFII ^{SQ→AA}	0.10 ± 0.015	1727 ± 52	0.94/30/3	2.9	1.0
rFII ^{SLQ→AAA}	2.3 ± 0.5	730 ± 101	0.96/36/4	0.053	55
rFII ^{ΔSLQe}	3.2 ± 1.5	128 ± 34	0.96/9/1	0.0067	463

^a The K_m and k_{cat} values of prothrombinase assembled with saturating concentrations of recombinant fVa molecules were determined as described under "Experimental Procedures" according to the Michaelis-Menten equation using the software Prizm with several different preparations of rFII molecules (representative experiments are shown in Fig. 7). Kinetic constants were derived directly from the fitted data.

^b $k_{cat} = V_{max}/[enzyme]$; the enzyme concentration of prothrombinase (fXa-fVa complex on the membrane surface in the presence of Ca²⁺) under the conditions employed herein was 10 pM.

^c R^2 is the goodness of fit of the data points to the Michaelis-Menten equation using the software Prizm. Points and titrations studied represent 10 measurements/graph for all experiments (up to 4 μM plasma-derived FII or rFII molecules) except experiments with rFII^{SLQ→AAA} (nine measurements/graph, up to 2 μM prothrombin) and with rFII^{ΔSLQ} (nine measurements/graph, up to 4 μM prothrombin).

^d The -fold decrease is the ratio of the second-order rate constant (k_{cat}/K_m) of prothrombinase catalyzing rFII^{WT} activation compared with the second-order rate constant of prothrombinase catalyzing activation of all other rFII molecules.

^e Experiments with these two preparations of recombinant molecules were performed in parallel with same reagents (fXa and PCPS vesicles). The results shown are representative of four separate titrations with three different preparations of rFII^{ΔSLQ} compared with either rFII^{WT} or FII^{plasma}.

^f NP is no plot; data could not be plotted to the Michaelis-Menten equation using the software Prizm.

TABLE 3

Kinetic constants of wild-type and selected rIIa mutant molecules toward S-2238

α -Thrombin species	K_m^a	k_{cat}^b	R^2^c	k_{cat}/K_m
	μM	s^{-1}		$(M^{-1}\cdot s^{-1})\cdot 10^6$
rIIa ^{WT}	4.7 ± 1.8	22.7 ± 2.7	0.93	4.8
rIIa ^{S478A}	8.5 ± 1.5	36.5 ± 2.4	0.98	4.3
rIIa ^{L480A}	8.9 ± 1.7	15.5 ± 1.5	0.98	1.7
rIIa ^{SL→AA}	7.7 ± 3.0	14.7 ± 2.0	0.92	1.9
rIIa ^{SQ→AA}	10.8 ± 4.2	33.1 ± 4.9	0.93	3.0
rIIa ^{SLQ→AAA}	NP ^d	NP	NP	

^a The K_m value for S-2238 is listed for wild-type rIIa and selected rIIa mutants and determined as described under "Experimental Procedures" according to the Michaelis-Menten equation using the software Prizm. Kinetic constants shown were derived directly from the fitted data.

^b $k_{cat} = V_{max}/[enzyme]$; the V_{max} was calculated as described under "Experimental Procedures," and the enzyme concentrations of rIIa was 4 nM for all experiments shown.

^c R^2 is the goodness of fit of the data points to the Michaelis-Menten equation using the software Prizm.

^d NP is no plot; no data could be plotted to the Michaelis-Menten equation using the software Prizm.

with our findings shown in Table 1, explaining the fact that rFII^{ΔC10} and rFII^{ΔS5V} are devoid of clotting activity, and further attest to the crucial dual role of the dipeptide Leu⁴⁸⁰-Gln⁴⁸¹ during coagulation. Further analyses of the single or double mutants reveal a slight differentiation in cleavage and activation of the two cofactors by the various rIIa molecules. Although rIIa^{L480A} and rIIa^{SL→AA} appear devoid of activity toward fV (Fig. 8, D and E), both molecules slowly cleave fVIII at the Arg³⁷²- and Arg¹⁶⁸⁹-activating cleavage sites (Fig. 9, D and E) (87–90). Similarly, although rIIa^{SLQ→AAA} has no apparent activity toward fV (Fig. 8G) over a 3-h time period, the mutant enzyme cleaves fVIII slowly at the non-activating Arg⁷⁴⁰ cleavage site (Fig. 9G). Finally, although rIIa^{SQ→AA} cleaves fV efficiently at Arg⁷⁰⁹ to produce the heavy chain of fV and an M_r 220,000 intermediate (Fig. 8F), the enzyme is also efficient in cleaving fVIII at the Arg³⁷²- and Arg¹⁶⁸⁹-activating cleavage sites (Fig. 9F). These two cofactors have strategic functions within the amplified coagulation response to vascular damage and must be activated to perform accordingly within their respective enzymatic complexes. The combined data explain the impaired procoagulant activity of rFII^{SLQ→AAA} (Fig. 2), which is deficient in producing large amounts of rIIa in a timely fashion (Fig. 5D). However,

even when rIIa^{SLQ→AAA} is generated, the recombinant enzyme is deficient in activating the pro-cofactors.

We next assessed the capability of the rIIa molecules in the presence of thrombomodulin to activate protein C and produce APC. Fig. 10 shows the results of such an experiment and demonstrates that although rIIa^{ΔS5V} (rIIa^{ΔS5V} is recombinant human α -thrombin with region SVLQV deleted) cannot cleave and activate protein C, rIIa^{SLQ→AAA} has small but significant activity generating minute amounts of APC (Fig. 10, lanes 8 and 9), which in turn can cleave fV at Arg⁵⁰⁶/Arg³⁰⁶ and produce the characteristic M_r 30,000 fragment (data not shown) (68, 91). All other rIIa mutant molecules tested, for APC generation, have similar activities as rIIa^{WT} or plasma-derived IIa under the condition described (Fig. 10).

These data demonstrate a differential requirement of IIa for cleavage and activation of both the pro-cofactor molecules and protein C and attest to the sensitive requirements of fV for cleavage and activation by IIa. Overall these results demonstrate that amino acids Leu⁴⁸⁰ and Gln⁴⁸¹ within the serine protease domain of FII serve a dual purpose, and thus both are required for efficient cleavage at Arg³²⁰ by prothrombinase and may be involved in the presentation of an obligatory exosite for timely fV, fVIII, and protein C activation by IIa.

Function of Sequence Leu⁴⁸⁰-Gln⁴⁸¹ of Prothrombin

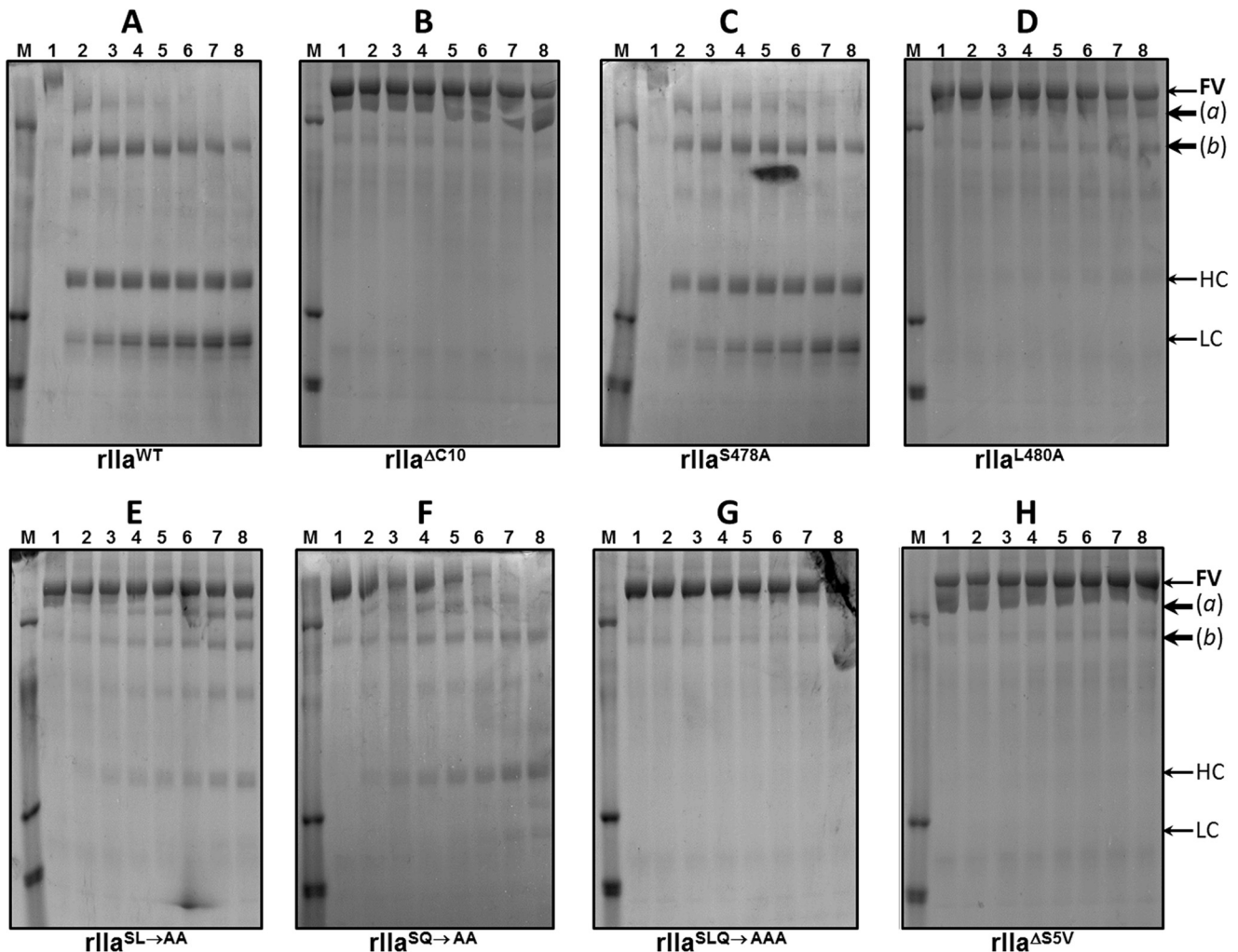


FIGURE 8. **Activation of plasma-derived fV by rIIa.** Plasma-derived fV (500 nM) was incubated with rIIa (4 nM) as described under "Experimental Procedures." At selected time intervals, aliquots of the mixtures were removed, mixed with 2% SDS, heated for 5 min at 90 °C, and analyzed on a 4–12% SDS-PAGE followed by staining with Coomassie Blue. Lane 1 in all panels depicts aliquots of the mixture withdrawn from the reaction before the addition of rIIa. Lanes 2–8 represent aliquots of the reaction mixture withdrawn at 10, 20, 30, 45, 60, 120, and 180 min. The positions of all fV fragments and of the heavy (HC) and light chains (LC) of the active cofactor are indicated on the right. Fragments *a* and *b* of fV are identified as previously described (80). The rIIa molecule used each time is indicated under each panel.

Discussion

Our data demonstrate that amino acid region 473–487 of FII is required for timely activation of FII through the meizothrombin pathway. Although prior work using synthetic peptides suggested that this region of the cofactor may contain an fVa-dependent fXa-binding site for FII, the data presented herein with recombinant FII molecules provide for the first time a mechanistic interpretation of these findings and identify the crucial amino acids from this sequence responsible for the effect observed (56).

To elucidate the number and identity of the required amino acids within amino acid sequence 473–487 of FII, we constructed, expressed, purified to homogeneity, and studied several rFII molecules with deletions and point mutations within this important regulatory region. We first investigated the effects of the 15-amino acid deletion with rFII^{Δ473–487}, followed by experiments with rFII molecules containing overlapping deletions within this segment (rFII^{ΔN10} and rFII^{ΔC10} and rFII^{ΔS5V} and rFII^{ΔSLQ}). Several rFII molecules bearing single mutations (rFII^{S478A} and rFII^{L480A}), double mutations (rFII^{SL→AA} and rFII^{SQ→AA}), and a triple mutation

(rFII^{SLQ→AAA}) were subsequently made. Membrane-bound fXa cleaves FII sequentially at Arg²⁷¹ followed by Arg³²⁰, forming small amounts of IIa. Under these conditions, the activation of the deletion mutants rFII^{Δ473–487}, rFII^{ΔN10}, rFII^{ΔC10}, rFII^{ΔS5V}, the triple alanine mutant, and the deletion mutant (rFII^{SLQ→AAA} and rFII^{ΔSLQ}) resulted in a modest increase of the rate of activation. In addition, activation of these six rFII molecules by fXa alone resulted in accumulation of prothrombin-2, with very little IIa formed. In contrast, activation of all these rFII mutants by fully assembled prothrombinase is significantly delayed. The combined data suggest that amino acids Leu⁴⁸⁰ and Gln⁴⁸¹ within region 473–487 of FII either represent or are responsible for the presentation of an fVa-dependent site for fXa on FII, which is essential for optimal rate of cleavage at Arg³²⁰, which in turn is required for timely IIa formation at the place of vascular injury.

The autolysis loop of APC bears strong homology with the FII sequence 473–487 (chymotrypsin numbering 149D–163) (92). Replacement of several basic amino acids from this homologous region in APC by site-directed mutagenesis to alanine demonstrated the ability of this exosite to interact with its sub-

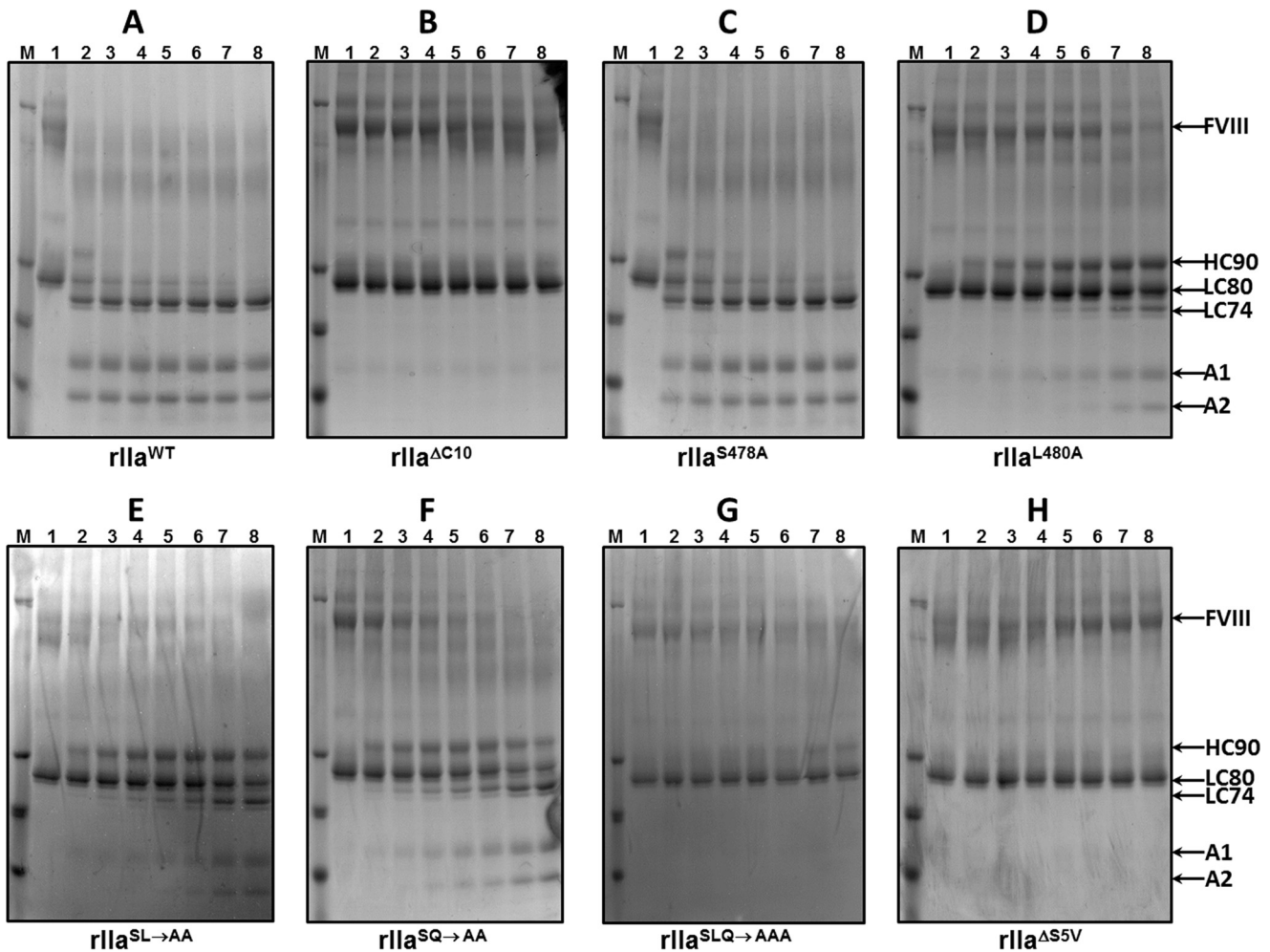


FIGURE 9. **Activation of recombinant FVIII by rIIa.** rFVIII (500 nM) was incubated with rIIa (4 nM) as described under "Experimental Procedures." At selected time intervals, aliquots of the mixtures were removed, mixed with 2% SDS, heated for 5 min at 90 °C, and analyzed on a 4–12% SDS-PAGE followed by staining with Coomassie Blue. Lane 1 in all panels depicts aliquots of the mixture withdrawn from the reaction before the addition of rIIa. Lanes 2–8 represent aliquots of the reaction mixture withdrawn at 10, 20, 30, 45, 60, 120, and 180 min. The positions of all rFVIII fragments are indicated on the right. Fragments from rFVIII are identified as previously demonstrated (80). The rIIa molecule used each time is indicated under each panel.

strate fVa and to differentiate between the Arg⁵⁰⁶ and Arg³⁰⁶ cleavage sites (93). Yegneswaran *et al.* (56) using synthetic peptides provided initial evidence that sequence 473–487 of FII is able to disrupt prothrombinase assembly only in an fVa-dependent manner. Chen *et al.* (50) identified a sequence within pro-exosite I of prothrombin containing basic residues Arg³⁵, Lys³⁶, Arg⁶⁷, Lys⁷⁰, Arg⁷³, Arg⁷⁵, and Arg⁷⁷ (chymotrypsin numbering), which is in close spatial proximity to region 473–487 of FII. These investigations revealed that following replacement of all basic residues from pro-exosite I with Glu, there was a significant effect on fXa within prothrombinase when compared with fXa alone in cleaving and activating FII, suggesting that these amino acids are specific fVa-dependent recognition sites for fXa on FII. Further kinetic studies by Chen *et al.* (50) using the hirudin COOH-terminal peptide (hirugen) showed that the peptide inhibited wild type prethrombin-1 activation by prothrombinase, whereas hirugen had no inhibitory effect on the activation of the mutated zymogen lacking the basic residues in pro-exosite I by fXa alone. The combined studies of Chen *et al.* (50) and Yegneswaran *et al.* (56) suggest the requirement of

both sites for optimum productive interaction of prothrombinase with FII and timely Ila formation.

Research with discontinuous assays using a chromogenic substrate for Ila revealed that when fVa is incorporated into the prothrombinase complex, the resulting K_m value of the reaction was decreased by 100-fold (corresponding to a 100-fold increase in affinity of prothrombinase for FII as compared with the affinity of fXa alone for the substrate), whereas the catalytic efficiency (k_{cat}) of fXa was increased by 3,000-fold resulting in a 300,000-fold overall increase in the activity of prothrombinase (second-order rate constant) for FII compared with the activity of fXa alone toward FII (16). The significant increase in affinity of prothrombinase for its substrate is attributed to tighter binding of the enzymatic complex to FII because of its localization on the membrane surface by fVa. The longstanding hypothesis that fVa "localizes and positions" FII in an optimum position for efficient catalysis by fXa consistent with the classical role of a cofactor for catalysis was recently confirmed by computational studies with prothrombinase by Shim *et al.* (27). These studies demonstrated that the acidic COOH-terminal portion of the

Function of Sequence Leu⁴⁸⁰–Gln⁴⁸¹ of Prothrombin

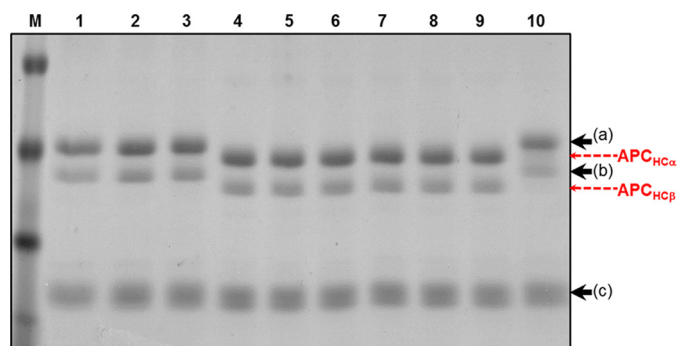


FIGURE 10. Activation of protein C by rIIa. Plasma-derived protein C (80 nM) was incubated with rIIa (8 nM), thrombomodulin, and PCPS as described under "Experimental Procedures." Following a 3-h incubation period, each individual solution was dried with a vacuum concentrator, resuspended in Tris buffer, mixed with 2% SDS and 2% β -mercaptoethanol, heated for 5 min at 90 °C, and analyzed on a 5–15% SDS-PAGE followed by staining with Coomassie Blue. *Lane 1*, protein C alone no IIa; *lane 2*, protein C alone, no IIa incubated with buffer for 3 h; *lane 3*, protein C and rIIa ^{Δ 55V}; *lane 4*, protein C and plasma-derived IIa; *lane 5*, protein C and rIIa^{WT}; *lane 6*, protein C and rIIa^{S478A}; *lane 7*, protein C and rIIa^{L480A}; *lane 8*, protein C and rIIa^{SL→AA}; *lane 9*, protein C and rIIa^{SQ→AA}; and *lane 10*, protein C and rIIa^{SLQ→AAA}. Positions of protein C and APC heavy and light chain fragments are indicated at the right (*a/b* heavy chains, and *c* light chain). The two heavy chains of protein C in plasma (*a* and *b*) have been identified earlier, differ by one glycosylation site, and have been extensively studied (58, 59).

heavy chain of fVa that is contiguous to the A2 domain of fVa is essential in its ability to interact and snare the serine protease domain of FII. In that manner, this acidic amino acid sequence reposition the Arg³²⁰ cleavage site at an optimum position for timely cleavage by fXa and FII activation at the site of vascular injury as earlier suggested (54) and more recently experimentally demonstrated by our laboratory with synthetic peptides and recombinant fVa molecules mutated at these specific sites (45, 47, 48).

We show that following removal of the amino acid sequence 473–487 from FII, prothrombinase loses the ability to efficiently form IIa because of impaired fVa-dependent cleavage of FII by fXa at Arg³²⁰. One easy explanation of these results was that elimination of such a huge portion of the molecule results in significant structural changes of the molecule that in turn have deleterious effects on FII molecular conformation resulting in deficient prothrombinase activity. Despite the fact that rFII ^{Δ 473–487} was activated following the same pathways as rFII^{WT} in the presence or absence of fVa, albeit with different rates, and in the absence of a crystal structure of rFII ^{Δ 473–487}, there was still doubt about the structural integrity and function of a molecule bearing such a large deletion. Experiments using more modest overlapping deletions (with rFII ^{Δ N10}, rFII ^{Δ C10}, and rFII ^{Δ S55V}) as well as with a triple alanine mutant (rFII^{SLQ→AAA}) and a triple deletion mutant (rFII^{SLQ}) demonstrated that these molecules are also hindered in their fVa-dependent cleavage at Arg³²⁰ to a similar level as rFII ^{Δ 473–487} (Table 1 and Fig. 6). These data provide original evidence demonstrating that the minimal sequence of FII required for the 3,000-fold increase in the catalytic efficiency of prothrombinase, as defined ~36 years ago (16) for efficient cleavage of FII by prothrombinase at Arg³²⁰, is carried at least partially by amino acid sequence Leu⁴⁸⁰–Gln⁴⁸¹ of FII. The findings presented herein silence the notion that the effect seen with rFII ^{Δ 473–487} may be due to a structural change of the mutant

molecule rather than to specific amino acid(s) missing from rFII ^{Δ 473–487}.

The kinetic findings presented herein revealed comparable K_m and k_{cat} constants for prothrombinase when rFII molecules bearing the single and double alanine mutations were used as substrate. However, when rFII^{SLQ→AAA} was the substrate for prothrombinase in the same discontinuous assay, there was a significant 21-fold increase in the K_m value and a modest 2.7-fold decrease in the k_{cat} of the enzyme. Similar results were obtained with rFII ^{Δ SLQ}. Furthermore, rFII^{SLQ→AAA} and rFII ^{Δ SLQ} were also found to be substantially deficient in clot formation in an assay using FII-deficient plasma, whereas rIIa^{SLQ→AAA} was also deficient in S-2238 hydrolysis. rIIa^{SLQ→AAA} was also impaired in cleaving fV, fVIII, and to a lesser extent protein C. These data dovetail nicely with results obtained with rIIa ^{Δ S55V} and rIIa ^{Δ 473–487}. We can thus hypothesize that the substantial increase in the K_m of prothrombinase toward rFII^{SLQ→AAA} is due to a deficiency in prothrombinase in recognizing the mutant molecule because of the lack of Leu⁴⁸⁰–Gln⁴⁸¹, whereas the decrease in enzymatic activity of the resulting rIIa^{SLQ→AAA} molecule is also the result of the absence of these two important amino acid side chains. Additional data with rFII ^{Δ SLQ} provide further evidence of the crucial role of amino acids Leu⁴⁸⁰–Gln⁴⁸¹ and the peptide bond backbone between these two amino acids because, when these residues are completely eliminated, the K_m value of the reaction increases by 32-fold, and the k_{cat} value decreases by 16-fold (Table 2). Keeping in mind that the S478A substitution is of no consequence on both rFII activation and rIIa function, these results provide strong evidence in favor of the dual role of amino acids Leu⁴⁸⁰ and Gln⁴⁸¹. Moreover, these amino acids are required by prothrombinase to efficiently promote cleavage of FII at Arg³²⁰ and are also required by IIa for optimum amidolytic activity as well as to proficiently cleave and activate fV, fVIII, and protein C. Finally, the possibility that elimination of these two residues from rFII results in an allosteric transition of the amino acids around/within the active site of rIIa, thus modifying the critical distances between the specific residues of the catalytic triad resulting in impaired catalysis, cannot be eliminated.

A comparison of crystal structures of FII, meizothrombin, IIa, prethrombin-1, and prethrombin-2 was carried out to identify structural differences in/near the Gly⁴⁷³–Ile⁴⁸⁷ segment comprising the fVa-dependent fXa-binding site. These residues adopt similar conformations in all of the crystal structures, with the NH₂-terminal residues Gly⁴⁷³–Gln⁴⁷⁶ being quite solvent-accessible or flexible, and residues Pro⁴⁷⁷–Ile⁴⁸⁷ being variable in their degree of solvent exposure. Residue Ile⁴⁸⁷ is significantly more exposed in prothrombin (accessible surface area of >30 Å² compared with ≤10 Å² in meizothrombin and thrombin), as well as the adjacent Pro⁴⁸⁶ (accessible surface area of ~15–30 Å² reducing to <10 Å² in meizothrombin and thrombin). The amount of solvent exposure of Ile⁴⁸⁷ and Pro⁴⁸⁶ appears to be heavily influenced by the flanking loops encompassing residues Ala⁴⁴⁶–Tyr⁴⁵⁴ and Lys⁵¹¹–Ser⁵²⁵, which adopt different conformations upon FII activation (Fig. 11). Recently, Pozzi *et al.* (94) used the crystal structure of Gla-domainless FII with active site S525A to demonstrate that fVa has recognition sites in close proximity to Arg³²⁰ (Arg¹⁵ chymotrypsin number-

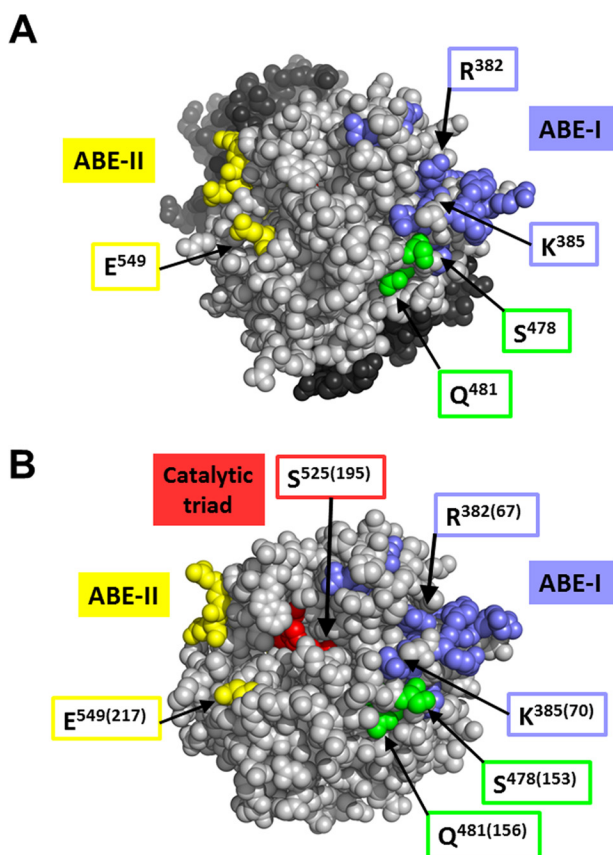


FIGURE 11. Location of exosites in FII and thrombin structures. Representation of the high resolution crystal structures of FII^{WT} (A) and IIa (B). A, space-filling representation of human FII (38). Residues Ser⁴⁷⁸, Leu⁴⁸⁰, and Gln⁴⁸¹ are colored green; ABE-I and ABE-II residues (6, 50) are purple-blue and yellow, respectively; the amino acids composing the catalytic triad residues are not solvent-accessible and thus not visible. Other catalytic domain residues are in light gray, and those in fragment-1 and fragment-2 are in dark gray. B, space-filling representation of IIa (81). Residues are colored as for FII with the addition of catalytic triad residues His³⁶³, Asp⁴¹⁹, and Ser⁵²⁵ (red), which are partially solvent-accessible. In parentheses are the corresponding numbers according to the chymotrypsin numbering of IIa (4). Distances between the active Ser⁵²⁵ side chain hydroxyl and several other amino acids of interest are as follows: 17 Å to Gln⁴⁸¹ OE1/NE2; 15 Å to Glu⁵⁴⁹ OE1; 17 Å to Arg³⁸² NH₂; 18 Å to Lys³⁸⁵ NH₂. The polar atoms at the end of the side chains were used as a reference because these would presumably be involved in intermolecular interactions.

ing). These sites create a strong electrostatic potential due to a number of basic residues described by Chen *et al.* (50). Through analysis of this published crystal structure, we have located this basic region to be in the vicinity of the Leu⁴⁸⁰–Gln⁴⁸¹ amino acid sequence of FII that we found to be required for efficient initial cleavage at Arg³²⁰ by prothrombinase. It is noteworthy that a very recent study by Pozzi *et al.* (38) demonstrated a crucial role for linker 2 for the rate of activation of FII by prothrombinase and suggested that this region may be involved in the interaction of FII with the cofactor. These data are in complete accord with data showing that fragment 1, more precisely the kringle 2 region, is involved in the interaction of fVa as part of prothrombinase with FII (39, 41). Finally, a close comparison of crystal structures of FII and IIa revealed that residues Ser⁴⁷⁸, Leu⁴⁸⁰, and Gln⁴⁸¹ adopt similar conformations in both structures. The Ser⁴⁷⁸ side chain is exposed on the surface of both molecules, whereas the Leu⁴⁸⁰ side chain is surrounded by other residues and is not accessible to solvent. Gln⁴⁸¹ is partially

solvent-exposed in both FII and IIa. The Ser⁴⁷⁸/Leu⁴⁸⁰/Gln⁴⁸¹ residues are near ABE-I (Fig. 11), but >15 Å from the catalytic Ser⁵²⁵ residue, and even more distant from ABE-II.

In conclusion, in this study we provide evidence for the dual effect of amino acids Leu⁴⁸⁰ and Gln⁴⁸¹ of FII. Future mutagenesis studies within the amino acids uncovered herein, paired with selected mutations within pro-exosite-I and/or pro-exosite-II of FII, should be able to elucidate the intermolecular communications within FII, required for both optimal fVa-dependent activation of FII and subsequent IIa catalytic activity toward its numerous physiological substrates. Finally, our results provide evidence for the production of large quantities of rFII^{ΔS5V}, rFII^{SLQ→AAA}, or rFII^{ΔSLQ} that could be used as therapeutic agents because these molecules would compete with the natural substrate *in vivo*, when infused in individuals with prothrombotic tendencies.

Author Contributions—J. R. W. designed, performed, and analyzed most experiments and participated in the writing of the paper; J. H. designed, performed, and analyzed some of the experiments; V. C. Y. designed and produced the structural pictures of prothrombin and thrombin shown in Fig. 11; M. K. conceived and coordinated the study and wrote the paper. All authors reviewed the results and approved the final version of the manuscript.

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The Dual Regulatory Role of Amino Acids Leu⁴⁸⁰ and Gln⁴⁸¹ of Prothrombin
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