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Biophysical Investigation of GpIba Binding to Thrombin Anion Binding Exosite II†

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Abstract

Substrates and cofactors of the serine protease thrombin (IIa) employ two anion binding exosites (ABE-I and II) to aid in binding. On the surface of platelets resides the membrane bound receptor complex GpIba/β-GpIX-GpV. IIa's ABE-II is proposed to interact with an anionic portion of GpIba which enhances IIa cleavage of PAR-1 and subsequent activation of platelets. In the current work, 1D and 2D NMR, analytical ultracentrifugation (AUC), and hydrogen deuterium exchange (HDX) coupled with MALDI-TOF MS were performed to further characterize the features of binding to IIa's ABEs. The described work builds upon investigations performed in a prior study with fibrin (ogen)'s γ' peptide and IIa (Sabo, T. M., Farrell, D. H., and Maurer, M. C. (2006) *Biochemistry* 45, 7434–7445). 1D line broadening NMR (¹H and ³¹P) and 2D trNOESY NMR studies indicate that GpIba D²⁷⁴-E²⁸⁵ interact extensively with the IIa surface in an extended conformation. AUC demonstrates that both GpIba (269–286) and γ' (410–427) peptides interact with IIa at 1:1 stoichiometry. Comparing the HDX results to the ABE-I targeting peptide hirudin (54–65), the data implies that GpIba (269–286), GpIba (1–290), and γ' (410–427) are indeed directed to ABE-II. The ABE-II binding fragments reduce HDX for sites distant from the interface suggesting long range conformational effects. These studies illustrate that GpIba and γ' target ABE-II with similar consequences on IIa dynamics, albeit with differing structural features.

At the nexus of procoagulation and anticoagulation resides the serine protease thrombin (IIa) (1–3).¹ The procoagulant activities of IIa are multifaceted and numerous, including the conversion of fibrinogen to fibrin to form the soft clot (4) and the activation of platelets through cleavage of the protease activated receptors (PAR) (5). A vital anticoagulant function of IIa is the activation of protein C which deactivates several procoagulant enzymes (6). The specificity of IIa is a result of the loops and residues that surround the active site (7). IIa contains two electropositive patches that flank the catalytic site termed anion binding exosites I and II (ABE-I and ABE-II) (8). Exosites guide substrate orientation for catalysis that is more efficient and/or provide a separate cofactor interaction site that either enhances or dampens enzyme reactivity.

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Supporting Information Available

Table S1 is Deuterium Incorporation for IIa with the peptides GpIba, γ' and Hirudin (54–65) and the protein fragment GpIba (1–290) 3Y_S and 2Y_S at pH 6.5. Table S2 is Deuterium Incorporation for IIa (pH 7.5 and 5.6) with the peptides GpIba, γ' and Hirudin (54–65) at pH 5.6 This material is available free of charge via the Internet at <http://pubs.acs.org>.

¹Abbreviations: IIa, thrombin; PAR-1, protease-activated receptor 1; ABE-I, anion binding exosite I; ABE-II, anion binding exosite II; GpIba, glycoprotein Iba; LRR, leucine rich repeat; FpA, fibrinopeptide A; FVIII, blood clotting factor VIII; PPACK, D-Phe-Pro-Arg chloromethyl ketone; Y_S, sulfotyrosine; Y_p, phosphotyrosine; NMR, nuclear magnetic resonance; 1D, one dimensional; 2D, two dimensional; TOCSY, total correlation spectroscopy; tr-NOESY, transferred nuclear Overhauser effect spectroscopy; MALDI-TOF MS, matrix-assisted laser desorption-ionization time-of-flight mass spectrometry; HDX, hydrogen-deuterium exchange; AUC, analytical ultracentrifugation; S, Svedberg constant; K_D, equilibrium binding constant.

An important event in coagulation is the initial recruitment of Ila by Glycoprotein Iba α (GpIba α) to the surface of platelets. This glycoprotein is part of the highly glycosylated transmembrane platelet receptor GpIba α / β -GpIX-GpV (9,10). The importance of this receptor is highlighted in Bernard-Soulier syndrome, a bleeding disorder resulting from either dysfunctional or low levels of this complex. GpIba α serves a multitude of functions that include binding coagulation proteins, adhering platelets to the site of damaged tissue, and aiding in leukocyte recruitment to combat infection. Two features of the N-terminus of GpIba α are a leucine-rich-repeat (LRR) domain and an anionic cluster (²⁶⁹DEGDTDL_Y_SDY_SY_SPEEDTEG²⁸⁶) (Table 1 and Figure 1A).

Considerable interest has been directed toward understanding in more detail the nature of the GpIba α -Ila interaction (12,13). Mutagenesis studies have identified the ABE-II residues R⁹³, R⁹⁷, R¹⁰¹, R²³³, K²³⁶ and K²⁴⁰ as playing a role in binding, not the ABE-I residues R⁶⁷, R⁷³, R⁷⁵, R⁷⁶, and R^{77A} (14–16).² Additionally, the ABE-II binding glycosaminoglycan heparin prevents GpIba α interaction with Ila (15). Conversely, others suggest ABE-I as GpIba α 's binding site through competitive inhibition with hirudin (54–65) (17–19). Hirudin is a well-characterized ABE-I targeting Ila inhibitor isolated from the salivary glands of medicinal leeches (20). It should also be noted that GpIba α binding to Ila hinders the release of fibrinopeptide A (FpA) from fibrinogen (14,21,22), enhances PAR-1 hydrolysis (23), and decreases catalysis at R³⁷² in FVIII (24).

Two crystal structures of Ila-GpIba α were published concurrently in 2003 (PDB IDs: 1OOK and 1P8V) (25,26) (Figure 1B). Though both groups report a 1:1 asymmetric unit, symmetry related molecules display unique binding interfaces. Both structures illustrate similar GpIba α interactions with ABE-II. 1OOK shows the C-terminus of GpIba α potentially dimerizing Ila through interactions with both exosites (25). In 1P8V, the second Ila molecule binds to GpIba α at the C-terminal end of the LRR domain (26). Two independent groups have analyzed the available experimental data and have concluded that the stronger interaction likely involves ABE-II and GpIba α residues L²⁷⁵-Y_S²⁷⁹, present in both structures (12,13).

Like GpIba α , fibrin(ogen) has been demonstrated to bind ABE-I (27) and ABE-II (28,29). The large soluble glycoprotein fibrinogen is a dimer comprised of three domains (A α B β γ _A)₂ (4, 30). During coagulation, Ila conversion of fibrinogen to fibrin (α β γ _A)₂ initiates formation of the non-covalently associated soft clot. The N-termini of the six polypeptides begin in the central E-region, supplying a low affinity ABE-I binding site (27). The C-termini of the polypeptides reach laterally from the E-region to form the flanking D regions. The ABE-II binding γ ' chain (28,29) is a variant of the γ _A chain with an insertion of 20 amino acids at the C-terminus (⁴⁰⁸VRPEHPAET_Y_SDSL_Y_SPEDDL⁴²⁷) (31,32) (Table 1).

Between 7–15% of fibrinogen circulates as the γ _A/ γ ' as opposed to γ _A/ γ _A (33,34). The potential physiological consequences associated with elevated levels of γ _A/ γ ' remains unclear. Some research has suggested that an abundance of γ _A/ γ ' is linked to increased incidence of cardiovascular diseases (33,35). Conversely, two independent groups demonstrated that reduced levels of the γ ' chain were indicative of a higher risk for deep vein thrombosis (36, 37).

A previous report utilized nuclear magnetic resonance (NMR) and hydrogen deuterium exchange (HDX) coupled with matrix-assisted laser desorption-ionization time-of-flight mass spectrometry (MALDI-TOF MS) to focus on the details of the γ ' peptide (410–427)-Ila ABE-II interaction (38). These experiments localized γ ' peptide binding within ABE-II and detected perturbations in HDX within ABE-I, the autolysis loop, and the A-chain. Many observations

²Throughout the paper, thrombin residues have been referenced according to the chymotrypsin numbering scheme.

reported from the NMR results correlated extremely well with the subsequently reported X-ray structure of γ' peptide-IIa (39).

In the current investigation, the GpIb α -IIa interaction was examined in further detail. Due to the high degree of similarity between the IIa binding sequences of GpIb α and the γ' chain (Table 1), perhaps these anionic peptides share conformational features. NMR experiments demonstrate that the GpIb α peptide binds to IIa in an extended conformation, unlike the turn structure reported for the γ' peptide (38). In addition, these studies were undertaken to better define the IIa exosite(s) responsible for GpIb α interaction, as well as to examine whether GpIb α binding induces the same long-range allostery that is observed when γ' peptide binds to IIa. The HDX results reveal that binding of both the GpIb α peptide (269–286) and fragment (1–290) to IIa's ABE-II alters the dynamics of IIa HDX analogous to the γ' peptide. Finally, analytical ultracentrifugation (AUC) was implemented to observe the behavior of IIa with GpIb α or the γ' peptide in solution.

Materials and Methods

Synthetic Peptides and GpIb α fragments

Bachem (King of Prussia, PA) synthesized peptides based on residues 269–286 of GpIb α (Table 1). SynPep (Dublin, CA) synthesized a peptide based on residues 410–427 of the human fibrinogen γ' chain variant (Table 1). For both peptides, phosphotyrosines (Y_P) were substituted for the naturally occurring sulfotyrosines (Y_S) without sacrificing binding affinity (28,40). Sigma-Aldrich and Bachem prepared the sulfonated hirudin fragment 54–65 (Table 1). MALDI-TOF MS measurements on an Applied Biosystems Voyager DE-Pro mass spectrometer were used to verify peptide mass. The concentrations of the peptide were determined by quantitative amino acid analysis (AAA Service Laboratory, Boring, OR).

The GpIb α 1–290 $3Y_S$ fragment ($Y_S^{276}Y_S^{278}Y_S^{279}$) and the GpIb α 1–290 $2Y_S$ fragment ($Y_S^{276}Y_S^{279}$) were generous gifts from the Ruggeri Laboratory at Scripps in La Jolla, CA. The procedure for *D. melanogaster* expression and purification of the two GpIb α fragments is detailed in a previous report (25). In order to characterize the GpIb α fragments, MALDI-TOF MS was used to observe the mass of the proteins.

On the Use of Bovine Thrombin

Thrombin was purified from bovine plasma barium sulfate eluate (Sigma-Aldrich) as described by Trumbo and Maurer (41). It should be noted that human and bovine IIa exhibit a high degree of sequence conservation (87% identical) (8). Important regions where there are no differences include the active site, the Na $^+$ binding site, the β -insertion (W^{60D}) loop, and both ABEs.

The interaction of the human derived fibrin(ogen) γ' and GpIb α fragments with bovine IIa should not be influenced by the minor differences between the species. Supporting this statement is an HDX study, where PPACK inhibition of human and bovine IIa affected HDX in a similar manner for analogous peptides from both types of IIa (42). In addition, 2D trNOESY NMR examining the structural features adopted by fibrinopeptide A (43,44) and a PAR1 peptide (45) bound to bovine IIa were in agreement with the corresponding human IIa X-ray structures (46,47). Finally, in the previous study concerning the γ' peptide and IIa, 2D trNOESY NMR results were similar for γ' peptide binding to both bovine α -IIa and human γ -IIa (38).

1H and ^{31}P NMR: General Theory

Sample preparation, $^1H/^{31}P$ NMR analysis, and NMR instrumentation were nearly identical to those described in Sabo et al. (38). The only difference is the exclusive use of the 500 MHz NMR spectrometer during the course of this work. A 1:10 ratio of enzyme (150 μ M) to peptide

(1.5 mM) was maintained for the three complexes studies: IIa-GpIb α (Y_P²⁷⁶Y_P²⁷⁸Y_P²⁷⁹), IIa-GpIb α (Y_P²⁷⁶Y_P²⁷⁹), and IIa-GpIb α (NoP). All complexes were examined at a pH of 5.6.

When free in solution, a peptide possesses very little secondary structural features due to lack of constraining factors, such as a protein in which to interact (48,49). This situation results in $\omega\tau_c \approx 1$ and longer transverse relaxation (T_2) times. The 1D spectrum will display very sharp peaks and the 2D trNOESY will lack the presence of many discernible NOEs. Introduction of the binding partner, in this case IIa, leads to the peptide interacting with the protein. Thus, those protons that contact the protein surface adopt the solution characteristics of the larger macromolecule, i.e. $\omega\tau_c \gg 1$, shorter T_2 times, and large negative NOEs. The peptide now assumes the secondary conformational features associated with binding to the protein. If the k_{off} for the peptide is sufficiently fast, then the peptide will take a “memory” of the bound conformation to the free peptide population. This rejoining with the free peptide population results in 1D line broadening for protons that contact the protein surface and a weighted average of the NOEs of the free and bound ligands in the 2D trNOESY spectrum.

Analytical Ultracentrifugation

IIa was inhibited with a two-fold excess of PPACK for 30 min. An assay involving the chromogenic substrate S2238 was employed to ensure that IIa was inactive. PPACK-IIa was then subjected to overnight dialysis with 150mM NaCl, 25mM phosphate, pH 7.4. The γ' or GpIb α peptide was lyophilized and reconstituted in the same buffer. A series of sedimentation velocity AUC experiments ranging from 1, 2, 5, 10, and 20 times peptide to IIa (4.5–7.0 μ M) in duplicate was performed in a XL-A AUC (Beckman-Coulter). The parameters for each run were 60,000 rpm, 25 °C, and 280 nm. The data was analyzed with the program DC-DT+.

HDX Sample Preparation

IIa was buffer exchanged into the HDX buffer 37.5 mM NaCl, 6.25 mM NaH₂PO₄, pH 5.6, 6.5, or 7.5 (all pH values are within ± 0.1) with an Amicon-Ultra 4 unit (10 kDa MWCO). The concentration of IIa for all preparations was 25 μ M. An aliquot of 24 μ L IIa and 24 μ L HDX buffer were evaporated to dryness using a SpeedVac unit (Savant). The dry aliquots were stored at -70 °C.

For experiments involving peptides, enough GpIb α , γ' , or hirudin (54–65) was added to 24 μ L of buffer exchanged IIa to achieve a 20:1 ratio (40:1 in the case of the γ' peptide at pH 6.5) during HDX. The peptides were pH adjusted prior to addition. The aliquots of buffer-exchanged IIa and peptide were evaporated to dryness using a SpeedVac unit. For experiments involving the proteins GpIb α -3Y_S and GpIb α -2Y_S, the two fragments were exchanged into the pH 6.5 HDX buffer. After the buffer exchange, the GpIb α fragment concentration was 40 μ M. An aliquot of 24 μ L IIa and 24 μ L GpIb α fragment were evaporated to dryness using a SpeedVac unit. The dry aliquots were stored at -70 °C.

The increased γ' (410–427) peptide to IIa ratio raises the possibility of discrepancies between results presented in an earlier report (38) and those described within this paper. Re-analysis of the γ' (410–427)-IIa HDX pH showed that the value was likely below 6.0 in the prior work and then compared to IIa HDX at pH 6.5 (38). The acidity of the γ' peptide stock solution surpassed the buffering capabilities of the 25 mM phosphate solution. Thus, both ABE-II binding and pH dependent HDX rate perturbations were being observed (38). When the pH of the γ' (410–427)-IIa solution was raised to 6.5 in the current project, residence time on the enzyme surface may have decreased. Increasing the peptide to IIa ratio to 40:1 promoted the observation of a single population of ligand-bound IIa. Fortuitously, the carefully monitored pH investigations described in this report validates the conclusions obtained in the previous study for γ' (410–

427)-IIa. Ligand binding to ABE-II decreases HDX at the binding interface, which leads to further decreases in HDX at regions not associated with the interaction site.

HDX Experiments

Dry aliquots were allowed to come to room temperature before beginning the experiment. First, 12 μL of D_2O was added to the dry aliquot yielding the final concentrations: 50 μM IIa, 80 μM GpIb α fragment or 1–2 mM peptide, 150 mM NaCl, 25 mM NaH_2PO_4 , pH 5.6, 6.5, or 7.5. The samples were incubated in a desiccator at room temperature for 1 minute or 10 minutes. After incubation, 114 μL of 0.1% TFA pH 2.5 (on ice) was added to quench the solution. An additional 1 μL of 5% TFA was added to insure that the pH was 2.5 in experiments at pH 7.5. Next, 66 μL of the quenched solution was transferred to a tube of activated pepsin bound to agarose. Digestion occurred on ice for 10 minutes. Centrifugation at 4°C separated the digest from the pepsin and 8.2 μL aliquots were immediately frozen in liquid N_2 . Each frozen HDX aliquot was analyzed as detailed in Sabo et al. with MALDI-TOF MS (38).

Coverage exists for ten peptides in bovine IIa from a non-reducing pepsin digest (42). Figure 2 depicts the location of the ten peptides within IIa relative to the two ABEs. Residues 65–84 represent a portion of ABE-I, while part of ABE-II is covered with three fragments: 85–99, 173–180, and 173–181. The autolysis loop peptide (135–149D) and 46–52 reside near ABE-I residues 65–84. A β strand within the segment 106–116 is located anti-parallel to a β strand near the N-terminus of the ABE-II fragment 85–99. The A-chain residues –13 to –4 of bovine IIa are highly flexible, but when present in an X-ray structure (such as 2A1D (50)), reach toward the C-terminal helix of the B-chain (233–240), which has several ABE-II residues. Lastly, the segment 202–207 fits into the hinge formed by the boomerang shaped A-chain.

The HDX experiments described in the results section cover three topics: 1) the effect of pH on IIa HDX, 2) perturbations in IIa HDX due to peptide binding at pH 5.6 and pH 6.5, and 3) a comparison of GpIb α peptide and GpIb α protein binding to IIa at pH 6.5. The data will be reported in two ways. Raw HDX will be illustrated in a one and ten minute bar graph for the four topics (Figure 6, Figure 8–Figure 10) (38). In addition, the numerical values have been reported in Supplementary Tables 1 and 2. The second method of analysis employs % changes to gauge the significance in fluctuations of HDX (Table 3 and Table 4) (38). This value relates the differences in HDX to the theoretical maximum amount of exchangeable protons present in each observable peptide. These % changes will be classified into three categories: 1) insignificant 0 to $\pm 1.9\%$, 2) modest change $\pm 2.0\%$ to $\pm 4.4\%$, and 3) significant change $\geq \pm 4.5\%$ (38,51–54).

Results

^1H and ^{31}P One-Dimensional Line Broadening NMR

1D NMR experiments were performed to determine which ^1H or ^{31}P nuclei within the GpIb α (269–286) peptide interact with IIa. The first set of 1D NMR spectra examined GpIb α ($\text{Y}_\text{P}^{276}\text{Y}_\text{P}^{278}\text{Y}_\text{P}^{279}$) and displayed the absence of significant line broadening for the peptide in the presence of IIa (data not shown). These findings indicate that the 3 Y_P peptide binds too tightly to IIa, hindering any transfer of information regarding bound structural features to the free peptide population.

Since the GpIb α ($\text{Y}_\text{P}^{276}\text{Y}_\text{P}^{278}\text{Y}_\text{P}^{279}$) peptide spectra did not display line broadening, a 2 Y_P variant was synthesized. The GpIb α ($\text{Y}_\text{P}^{276}\text{Y}_\text{P}^{279}$) peptide still interacts with IIa (demonstrated by the two crystal structures (25,26)), but the absence of one Y_P may sufficiently weaken the interaction for the appearance of 1D line broadening. The spectra in Figure 3A and B focus on the amide region and do illustrate some line broadening in the presence of IIa. Most of the

broadening encompasses residues D²⁷⁴-Y_P²⁷⁹, while a modest amount is observed for residues E²⁸¹-D²⁸³. However, the line broadening is not extensive, with residues D²⁶⁹-D²⁷⁴ and T²⁸⁴-G²⁸⁶ appearing unchanged from the free peptide spectrum. Curiously, the ³¹P spectra do not present evidence of line broadening (data not shown).

A final peptide was synthesized without Y_P. Much more line broadening is evident with GpIbα (No Y_P) (Figure 3C and D). The central portion of the peptide spanning D²⁷⁴-Y²⁷⁹, E²⁸¹, and D²⁸³ display a significant amount of line broadening. The residues E²⁸² and E²⁸⁵ show a modest amount of broadening. Finally, the N-terminal residues D²⁶⁹-T²⁷³ and the C-terminal residues T²⁸⁴ and G²⁸⁶ do not appear to be essential for the interaction of GpIbα (269–286) with IIa.

Two-Dimensional Transferred Nuclear Overhauser Effect NMR

The 2D spectrum of GpIbα (Y_P²⁷⁶Y_P²⁷⁸Y_P²⁷⁹) was indistinguishable from the spectrum with the peptide in the presence of IIa (data not shown). For the Y_P²⁷⁶Y_P²⁷⁹ and No Y_P GpIbα peptides, the 2D trNOESY spectra displayed only nearest neighbor NOEs in the presence of IIa. Figure 4 illustrates some of the key features from the GpIbα (No Y_P)-IIa 2D trNOESY spectra. Detection of only nearest neighbor NOEs suggests that the peptide adopts an extended conformation when binding to IIa. In addition, the Y²⁷⁹-P²⁸⁰ amide bond is in the *trans* conformation. With the GpIbα (Y_P²⁷⁶Y_P²⁷⁹) peptide, slightly fewer inter-residue NOEs are observed in the presence of IIa when compared to GpIbα (No Y_P).

Analytical Ultracentrifugation

Two X-ray studies offer evidence for potential ABE-II binding ligand-induced IIa dimerization (25,39). AUC experiments were performed to monitor the behavior of IIa in the presence of the GpIbα or γ' peptides. In these AUC trials, the concentrations of PPACK inhibited IIa spanned 4.7–7.0 μM. The sedimentation velocity experiments were executed with a range of peptide to IIa molar ratios varying from 0:1 to 20:1. Table 2 presents the data for the average of two runs. Figure 5 displays the distribution of Svedberg constants (S) for the trial examining 2:1 GpIbα peptide to IIa. This figure is representative of all data collected from 1:1 to 20:1 peptide to IIa reported in Table 2.

The Svedberg constant measures the rate of sedimentation, which is related to the size and shape of the molecule (55). When two proteins interact, the surface area is decreased leading to a non-additive increase in the S constant. The calculated Svedberg constants for the peptide binding to the enzyme range from 3.2–3.3, compared to 3.1 for IIa alone. The slight increase in molecular weight and S values with increasing peptide concentration indicate that the peptides bind to IIa but do not instigate protein dimerization under these conditions.

HDX Experiments (I): Observing the Effect of pH on Thrombin's Backbone Amide Solvent Accessibility

The HDX method exploits differences in the rate of chemical exchange for backbone amide protons in proteins (56). Surface-exposed backbone amide protons are more available for HDX than protons buried within the interior of the protein. Through fluctuations in protein structure (temporary protein unfolding and/or local conformational changes), a buried backbone amide proton may become exposed to solvent (57,58). Therefore, protein dynamics influence deuterium exchange to a great degree. The rate of chemical exchange is highly dependent on pH and temperature (59).

For the current project, HDX effects at pH values of 5.6, 6.5, and 7.5 were examined. Work at pH 5.6 allows comparisons to be made with the NMR solution conditions. A pH of 7.5 is physiological, but the backbone amides exchange very rapidly. A more favorable working pH

is 6.5, a condition that slows the backbone exchange 10-fold relative to pH 7.5 (60) and decreases the rate of IIa autolysis. Figure 2 illustrates the coverage obtained from a peptic digest of IIa (see Materials and Methods for description).

Comparing the levels of HDX across this pH spectrum reveals three distinct groups. See Figure 6, Supplementary Tables 1 and 2. The first grouping concerns those fragments that steadily increase in deuterium incorporation as the pH increases, suggesting a significant degree of solvent exposure for some of the amide protons. These peptides include the ABE-II peptides 85–99 and 173–181, as well as 106–116. The second grouping involves –13 to –4, 46–52, and 202–207. These three peptides seem impervious to pH change. Two probable explanations are that most of the amide protons are solvent inaccessible and/or the amide protons exchange too quickly for any apparent pH effect to be noticed. Finally, both the ABE-I peptide 65–84 and the autolysis loop 135–149D show stagnant levels of HDX at ten minutes. At one minute, an increase in pH appears to slow HDX for the autolysis loop peptide, perhaps indicating a more stable configuration for 135–149D at higher pH. As for 65–84, the rate of deuterium incorporation may slightly increase with higher pH at one minute of HDX.

HDX Experiments (II): Observing the Effect of the GpIb α (Y_P²⁷⁶Y_P²⁷⁸Y_P²⁷⁹) Peptide, the γ' Peptide (Y_P⁴¹⁸Y_P⁴²²), and Hirudin (54–65) on Thrombin's Backbone Amide Solvent Accessibility at pH 5.6 and pH 6.5

The first reports describing HDX coupled with MALDI-TOF MS to study solvent accessibility at a ligand-protein interface appeared in 1998 (61,62). With such an HDX experiment, the ligand should be bound to the protein, ideally, for the entire course of the deuterium exchange (61,63). Thus, the K_D describing the strength of the ligand-protein interaction, as well as, the concentration of the binding partners, will determine the lifetime of the complex (63).

The GpIb α , γ' , and hirudin (54–65) peptides were incubated with IIa at room temperature and pH 5.6 or pH 6.5 for one or ten minutes in the presence of 99.99% D₂O. A previous study found that GpIb α residues 265–285 bound to IIa with a K_D of 5.9 nM (21). In addition, the larger GpIb α fragment 1–282 possesses a K_D of 149 nM (64). The weaker K_D may stem from the truncated C-terminus. For comparison purposes, an ABE-I targeting peptide was also examined. Sulfonated hirudin (54–65) has a K_D of 48 nM recently determined through fluorescence studies (65). Thus, 1 mM GpIb α and 1 mM hirudin (54–65) should provide 99.99% occupancy of 50 μ M IIa. As for the γ' peptide, the interaction is weaker (K_D = 680 nM) (28), resulting in 99.93% and 99.97% occupancy at 1 and 2 mM peptide, respectively.

In Figure 7, the isotopic cluster for residues 85–99 of 20:1 γ' peptide to IIa at pH 6.5 has an interesting feature not evident at pH 5.6. The cluster appears to be expanded with a slight shoulder ending near 2115 m/z (not shown is similar shoulder for 173–181). The results suggest the presence of two different IIa populations at pH 6.5 exposed to deuterium. HDX performed at pH 6.5 on a higher ratio (40:1) of the γ' peptide (2 mM) to IIa (50 μ M) led to the reappearance of a single isotopic cluster (see Materials and Methods for explanation). Therefore, pH 6.5 HDX comparisons will be made between 20:1 hirudin (54–65)-IIa, 20:1 GpIb α -IIa, and 40:1 γ' peptide-IIa.

The HDX results for peptide binding to IIa are compiled in Table 3 and Table 4, Supplementary Tables 1 and 2, and Figure 8 and Figure 9. At one and ten minutes of HDX, both the GpIb α and γ' peptides significantly protect the ABE-II fragment 85–99 from deuterium. A unique feature of the data at pH 5.6 is the observed increase in HDX protection at ten minutes for 85–99, either indicating a tighter complex at the lower pH, or representing the slower rate of HDX at this pH. Another segment within ABE-II, 173–181, increases in protection from one to ten minutes of HDX at both pH values in a similar manner. These results suggest that the GpIb α

and γ' peptides are interacting with IIa's ABE-II. Conversely, hirudin (54–65) exclusively targets ABE-I as evident from the dramatic HDX protection at 65–84.

Binding to ABE-I and ABE-II appears to protect regions within IIa unaffiliated with their interactive sites. Hirudin (54–65) interactions with IIa lead to mostly modest decreases in HDX for ABE-II peptides 85–99 and 173–181 over the course of ten minutes. The ABE-I fragment 65–84 is affected by GpIb α and γ' binding to ABE-II to greater degree than hirudin (54–65) ABE-I binding imparts over to ABE-II. From one to ten minutes of HDX at pH 5.6, fragment 65–84 displays slightly less deuterium incorporation with the GpIb α peptide than the γ' peptide. At pH 6.5, peptide interactions at ABE-II show a large degree of protection at one minute for 65–84 that becomes significantly reduced within ten minutes of HDX.

Located parallel to a portion of the ABE-II fragment 85–99 are the residues 106–116 represented by two overlapping fragments. Hirudin (54–65) binding has very little effect on HDX for 106–116 at pH 5.6, but at pH 6.5 noteworthy protection can be localized to residues 106–113. Both the GpIb α and γ' peptides display modest protection at one minute, increasing to a level of greater significance at ten minutes for experiments conducted at pH 5.6. As for pH 6.5, the levels of significant HDX protection remain steady from one to ten minutes for 106–116. These data imply that ABE-II binding is transmitted to this neighboring β -strand.

Near ABE-I is the segment 135–149D, which also encompasses most of the autolysis loop. All three peptides significantly slow HDX for this region of IIa. Hirudin (54–65) imparts a larger effect on the dynamics of this region from ABE-I than the GpIb α and γ' peptides from ABE-II. Another long-range consequence of ABE-I and ABE-II occupation is evident from HDX within the A-chain residues –13 to –4 and the nearby B-chain fragment 202–207. The degree of protection is of a larger magnitude for the γ' and GpIb α , equating to mostly significant deviations from the HDX of IIa alone. Hirudin (54–65) results indicate that pH plays a role in HDX for –13 to –4 and 202–207. A higher pH of 6.5 results in these two peptides remaining largely unaffected by hirudin (54–65) binding to ABE-I, whereas at pH 5.6 significant protection is observed. All three peptides do not change the dynamics of deuterium exchange for the fragment 46–52.

HDX Experiments (III): Observing the Effect of the GpIb α (Y_P²⁷⁶Y_P²⁷⁸Y_P²⁷⁹) Peptide and the GpIb α fragment (1–290) on Thrombin's Backbone Amide Solvent Accessibility at pH 6.5

The Ruggeri laboratory at Scripps generously donated expressed GpIb α (1–290) with 3Y_S (Y_S²⁷⁶Y_S²⁷⁸Y_S²⁷⁹) and 2Y_S (Y_S²⁷⁶Y_S²⁷⁹) with the goal of comparing the HDX of the peptide to a larger protein fragment. The GpIb α (1–290)-IIa HDX data recorded at pH 6.5 is quite similar for the 3Y_S protein and the 3Y_P peptide (Figure 10, Table 4, and Supplementary Table 1). The 2Y_S protein displayed similar trends in HDX, however, the degrees of % change are not as large as with the 3Y_S protein. As with the two ABE-II binding peptides, the GpIb α (1–290) fragment appears to be interacting at ABE-II. Also, long range HDX perturbations are evident toward ABE-I, the A-chain, and the autolysis loop. It must be noted that the HDX results for hirudin (54–65) and GpIb α (1–290) are not similar. These findings further support the notion that GpIb α 's primary destination on IIa is ABE-II.

Discussion

On the surface of platelets, the receptor GpIb α forms a complex with IIa, however, the literature presents evidence for interactions at both ABEs. Though mutagenesis data strongly suggest that GpIb α binds exclusively at ABE-II (14–16), binding studies display interactions at both exosites (15,17,19,21,64). Adding to the ambiguity, two crystal structures of GpIb α -IIa illustrate similar interactions at ABE-II for one IIa, while symmetry related IIa binds to GpIb α through ABE-I at different GpIb α interfaces (25,26) (Figure 1B). The studies described

in this report use solution NMR, AUC, and HDX coupled with MALDI-TOF MS to investigate the interaction of GpIb α with IIa.

Analysis of the NMR Data: the GpIb α Peptide (269–286) Binds to Thrombin in an Extended Conformation

To sufficiently weaken the interaction and thus observe the trNOESY effect, the GpIb α peptide (269–286) was dephosphorylated to varying degrees. Conflicting accounts exist regarding the interaction of No Y_P variants of GpIb α and IIa. One report describes GpIb α (No Y_P) as unable to bind to IIa (66), while another group demonstrated competitive inhibition of FVIII hydrolysis by IIa in the presence of GpIb α (No Y_P) (268–282) (24). The present investigation supports the observation that GpIb α (No Y_P) can interact with IIa.

The lack of line broadening observed for GpIb α residues D²⁶⁹-T²⁷³ in the presence of IIa can potentially help resolve the two different conformations depicted for GpIb α 269–277 in the X-ray structures (Figure 11A) (25, 26). Also, a previous study de-emphasizes the importance of GpIb α D²⁶⁹-G²⁷¹ for binding IIa (64). It can be concluded that the GpIb α residues D²⁶⁹-T²⁷³ probably exist as a flexible portion of the glycoprotein, able to adopt multiple conformations, including those observed in the X-ray structures.

Overall, the GpIb α -IIa NMR work is more consistent with the 100K X-ray structure (25), which depicts GpIb α 269–284 in an extended conformation with a *trans* configuration for the Y_P ²⁷⁹-P²⁸⁰ bond (Figure 11A) (25). Electrostatic interactions are observed in 100K for residues D²⁷⁴-T²⁸⁴ with both ABE-I and ABE-II (Figure 11B) (25). Presumably, residues P²⁸⁰-G²⁸⁶ are too disordered in 1P8V to be effectively modeled (26); yet the NMR data displays significant interactions with IIa for several of the residues within this sequence. The current NMR data and the X-ray structure 100K allude to the possibility of GpIb α binding to both exosites of IIa in an extended conformation through dimerization of the protease (Figure 11B).

An analogous situation is apparent when analyzing the NMR and X-ray data for the interaction of the γ' peptide and IIa (38,39) (Figure 11C). In this case, the γ' peptide appears to dimerize IIa in the X-ray structure through utilization of ABE-II of both IIa molecules. The turn structure first described by the NMR work (38) is depicted as being sandwiched between the IIa dimer, perhaps stabilizing this configuration (39). Such interpretations should, however, be viewed with caution since crystal packing effects can influence the structural features being observed.

Analysis of the Analytical Ultracentrifugation Data: GpIb α Peptide (269–286) and γ' Peptide (410–427) binding to Thrombin

The NMR and X-ray work were the motivations to use AUC to further examine how IIa behaves with increasing concentrations of peptide. The S values calculated for PPACK-IIa (3.1) and PPACK-IIa in the presence of the peptides (3.2–3.3) match those previously determined for PPACK-IIa (3.3) (67). The same study shows a fragment of thrombomodulin with a similar molecular weight to IIa binding to the serine protease and increasing S from 3.3 to 4.2 (67). Another group utilized sedimentation equilibrium AUC to demonstrate that human IIa does not dimerize at 4 μ M, matching our results with bovine IIa (68). Taken together, IIa dimerization is not occurring under these conditions.

Analysis of the HDX Data (I): the Influence of pH on HDX dynamics

A series of IIa HDX experiments were performed at pH 5.6, 6.5, and 7.5. The present HDX studies demonstrate that ligand binding to IIa affects HDX in a similar manner at pH 5.6 and 6.5. This finding is important relative to NMR studies performed at pH 5.6 since it suggests that ligand binding effects associated with IIa binding are maintained at lower pH. The highest pH selected was 7.5, a more physiological pH, however, backbone amides exchange at a 10-

fold faster rate than at pH 6.5 (60). The main impetus for focusing on pH 6.5 for the current and previous HDX studies (38) are the slower rate of backbone amide exchange coupled with the decreased rate of IIa autolysis.

It is interesting to compare these results with another HDX study with human IIa at pH 6.6 and 7.9 (63). In agreement with the pH studies of Mandell *et al.*, the ABE-II 85–99 region can be described as completely solvent accessible and the 106–116 segment as fairly solvent accessible. Also, the ABE-I region 65–84 is defined by both groups as undergoing a fast rate of HDX. By contrast, the current HDX results for 65–84 at one minute slows significantly with the lower pH of 5.6, a condition not studied by the Komives group. Some differences in HDX are observed when comparing results for 135–149D from this study and 136–149A from Mandell *et al.* (63). The discrepancies are probably due to the inclusion of the highly solvent accessible autolysis loop (149A–149D) in the current study.

Analysis of the HDX Data (II): GpIb α (269–286), GpIb α (1–290), and the γ' Peptide (410–427) Target ABE-II

Both peptides appear to bind to IIa's ABE-II based on HDX protection for fragments 85–99 and 173–181 over the course of ten minutes. Figure 12 presents the γ' peptide-IIa and GpIb α -IIa X-ray structures viewed from ABE-II. Two residues from the GpIb α mutagenesis studies are present within 85–99 that reduce binding when mutated to A: R⁹³ and R⁹⁷ (14, 15). While R⁹⁷ does not appear to interact with GpIb α in the crystal structures, R⁹³ is depicted as being a prominent residue in binding to Yp²⁷⁹ of both GpIb α crystals (25, 26). Also, the GpIb α structures display the importance of IIa N¹⁷⁹ in binding to GpIb α D²⁷⁷, which is within fragment 173–181 (25, 26). Similarly, the γ' HDX data supports the observed electrostatic interactions between IIa R⁹³ and γ' S⁴²⁰, as well as IIa N¹⁷⁹ and γ' D⁴¹⁹ (39).

Hirudin (54–65) possesses a unique HDX profile when compared to the ABE-II targeting peptides. Numerous X-ray and NMR structures have been reported detailing IIa bound hirudin (69–74). In addition, Myles *et al.* provide an excellent analysis regarding the electrostatic and hydrophobic nature of hirudin binding to IIa (75). Hirudin (54–65) affects IIa HDX in a similar manner as the larger thrombomodulin fragment TMEGF45 (63). The dramatic protection observed for IIa residues 65–84 can be attributed to hirudin (54–65) electrostatic interactions with R⁷³, R⁷⁵, and R^{77a}, hydrophobic contacts with L⁶⁵, T⁷⁴, Y⁷⁶, and I⁸², and the energetically linked residues R⁶⁷ and K⁸¹.

The evidence for a similar destination on IIa for γ' and GpIb α becomes more compelling when considering recent research. In some of these investigations, the γ' peptide competitively inhibits IIa binding to GpIb α , consequently hindering platelet aggregation (76,77). Similarly, Weeterings *et al.* have shown that fibrin bound to IIa's ABE-I can promote platelet aggregation through GpIb α interaction with ABE-II (78). Taken together with the HDX data in this report, GpIb α residues 269–286 and 1–290 appear to preferentially target ABE-II.

Finally, it must be pointed out that others have declared ABE-I as GpIb α 's primary binding site on IIa (17–19). Perhaps the concentration of GpIb α and IIa is a factor to consider when examining GpIb α interactions with IIa. In one of these reports, the investigators used IIa at 0.42 nM and GpIb α at 50 μ M to monitor the effect of the peptide on catalytic activity, 100,000 times the amount of IIa (17). Conversely, a study proclaiming ABE-II as GpIb α 's binding site implemented a much lower 80:1 ratio of IIa (100 nM) to GpIb α (8 μ M) (15). As for the GpIb α protein fragment (1–290), the ratio was much lower at 1.6:1, yielding similar results to the GpIb α peptide (269–286) at 20:1. These observations suggest that protein/peptide concentration plays an important role in determining the destination of GpIb α on IIa.

Analysis of the HDX Data (III): GpIb α (269–286), GpIb α (1–290), and the γ' Peptide (410–427) Affect Thrombin HDX Dynamics in a Similar Manner

The evidence for exosite binding linkage to the active site is well established (15,23,79–81). It has been shown that occupation of ABE-II by GpIb α enhances PAR-1 hydrolysis by Ila (23), yet may non-competitively inhibit FpA release from fibrinogen (15). More recently, research has demonstrated that competitive binding of γ' peptide to ABE-II reduces the amount of PAR-1 cleaved on the platelet surface (76,77). As for energetically linked exosites, conflicting reports exist in the literature. Though work has shown a strong negative thermodynamic linkage between the exosites (29,82), others have found simultaneous exosite occupation without a significant energetic coupling (79,83).

In this context, questions arise about the nature of the reduced HDX observed when these peptides bind to Ila through ABE-II. Three distinct possibilities can be envisioned. Observed decreases in HDX could be the product of a restricted ensemble of conformations, a conformational change, or as previously discussed, a ligand-protein interface. The last scenario implies that these peptides also bind to other sites on the surface of Ila, such as ABE-I. Evidence against this situation has been provided in the earlier discussion.

The second scenario is that ABE binding could trigger a series of conformational changes that emanate from the binding interface. For example, within the autolysis loop peptide (135–149D), residue W¹⁴¹ has been postulated to connect ABE-I to the active site (42,63,84). This line of communication from ABE-I to the active site via W¹⁴¹ has been observed with thrombomodulin binding to ABE-I (63,84) and PPACK inhibition at the active site of Ila (42). The strong HDX perturbations occurring to the loop residues 135–149D when hirudin (54–65), and to a lesser extent γ' and GpIb α , bind may be reflective of linkage between the Ila active site, W¹⁴¹, and the exosites ABE I and/or ABE-II.

An alternative explanation centers on a seminal publication from the Griesinger group which presents strong evidence that proteins (in this case ubiquitin) sample most conformations associated with the observed bound conformation before interacting with a ligand (85). Through evolutionary means, the authors suggest that ligands have exploited specific conformations that are sampled by the target protein (i.e. the conformational ensemble). In the context of ligand binding to IIIa, perhaps binding to ABE-I and ABE-II favors one group of Ila conformers over another. This preference would be reflected in the altered HDX dynamics at sites unaffiliated with ligand binding. Regardless of the explanation, the physiological significance of reduced HDX profiles for ABE-occupied Ila remains an intriguing question. Results collected thus far strongly suggest a role for linkage between the two ABE sites.

Correlating the NMR, AUC, and HDX Results: the Influence of Thrombin Concentration

An important variable exists between the three primary investigations reported in this paper: Ila concentration. The Ila required for the NMR, HDX, and AUC studies amounted to 150 μ M, 50 μ M, and 4 μ M, respectively. Early studies attempting to determine the molecular weight of bovine Ila found that as the concentration of Ila increased, the apparent MW doubled (86, 87). Another group reported Ila dimerization in gel-filtration runs (88), which has been corroborated in the presence of GpIb α by Celikel *et al* (25). Physiologically, research has shown that low levels of Ila activate platelets in a GpIb α dependent manner, whereas higher levels of Ila perform this function independent of GpIb α (18,89). Clearly, Ila concentration has direct consequences on the mechanism of platelet activation.

Remaining to be determined is whether GpIb α induced Ila dimerization has physiological relevance in regard to bridging neighboring platelets (26) or clustering GpIb α together on the surface of a platelet (25). Perhaps Ila dimerization observed in crystallography and implied in

the NMR work is related to the higher amount of material required for these investigations. Recently, serine protease dimerization was reported in IIa-suramin complexes and the effect attributed to crystal packing (90). It is quite intriguing that a potential physiologically relevant dimeric form of prothrombin has been described that retains catalytic activity (91). As several groups have suggested, the K_D and concentrations of all IIa interacting partners must be considered at the site of thrombus formation (76,92,93).

Conclusions

These studies further characterize peptide binding to IIa's anion binding exosites. The NMR results suggest that the GpIb α peptide binds to IIa through residues D²⁷⁴-E²⁸⁵ in an extended conformation. HDX data reveal significant similarities between GpIb α and γ' peptide binding to ABE-II, while hirudin (54–65) demonstrates the HDX profile for an ABE-I targeting peptide. A significant finding from this work concerns the similarity in HDX profiles for the GpIb α peptide and protein fragments, providing strong evidence for the effectiveness of peptide mimics of proteins. By examining γ' peptide (410–427) and GpIb α (269–286) binding to IIa, a clearer understanding emerges concerning the importance of ABE-II in regulating blood coagulation and platelet activation.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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References

1. Di Cera E. Thrombin interactions. *Chest* 2003;124:11S–17S. [PubMed: 12970119]
2. Bode W. The structure of thrombin, a chameleon-like proteinase. *J Thromb Haemost* 2005;3:2379–2388. [PubMed: 15892855]
3. Huntington JA. Molecular recognition mechanisms of thrombin. *J Thromb Haemost* 2005;3:1861–1872. [PubMed: 16102053]
4. Weisel JW. Fibrinogen and fibrin. *Adv Protein Chem* 2005;70:247–299. [PubMed: 15837518]
5. Coughlin SR. Protease-activated receptors in vascular biology. *Thromb Haemost* 2001;86:298–307. [PubMed: 11487018]
6. Esmon CT. The protein C pathway. *Chest* 2003;124:26S–32S. [PubMed: 12970121]
7. Bode W. The structure of thrombin: a janus-headed proteinase. *Semin Thromb Hemost* 2006;32:16–31. [PubMed: 16673263]
8. Bode W, Turk D, Karshikov A. The refined 1.9-Å X-ray crystal structure of D-Phe-Pro-Arg chloromethylketone-inhibited human alpha-thrombin: structure analysis, overall structure, electrostatic properties, detailed active-site geometry, and structure-function relationships. *Protein Sci* 1992;1:426–471. [PubMed: 1304349]
9. Andrews RK, Gardiner EE, Shen Y, Whisstock JC, Berndt MC. Glycoprotein Ib-IX-V. *Int J Biochem Cell Biol* 2003;35:1170–1174. [PubMed: 12757754]
10. Clemetson KJ. A short history of platelet glycoprotein Ib complex. *Thromb Haemost* 2007;98:63–68. [PubMed: 17597992]
11. DeLano, WL. The PyMOL Molecular Graphics System. San Carlos, CA, USA: DeLano Scientific; 2002.

12. Kobe B, Guncar G, Buchholz R, Huber T, Maco B, Cowieson N, Martin JL, Marfori M, Forwood JK. Crystallography and protein-protein interactions: biological interfaces and crystal contacts. *Biochem Soc Trans* 2008;36:1438–1441. [PubMed: 19021571]
13. Vanhoorelbeke K, Ulrichs H, Romijn RA, Huizinga EG, Deckmyn H. The GPIIb/IIIa-thrombin interaction: far from crystal clear. *Trends Mol Med* 2004;10:33–39. [PubMed: 14720584]
14. De Cristofaro R, De Candia E, Landolfi R, Rutella S, Hall SW. Structural and functional mapping of the thrombin domain involved in the binding to the platelet glycoprotein Ib. *Biochemistry* 2001;40:13268–13273. [PubMed: 11683636]
15. Li CQ, Vindigni A, Sadler JE, Wardell MR. Platelet glycoprotein Ib alpha binds to thrombin anion-binding exosite II inducing allosteric changes in the activity of thrombin. *J Biol Chem* 2001;276:6161–6168. [PubMed: 11024046]
16. Yun TH, Baglia FA, Myles T, Navaneetham D, Lopez JA, Walsh PN, Leung LL. Thrombin activation of factor XI on activated platelets requires the interaction of factor XI and platelet glycoprotein Ib alpha with thrombin anion-binding exosites I and II, respectively. *J Biol Chem* 2003;278:48112–48119. [PubMed: 12968031]
17. Bouton MC, Thureau C, Guillin MC, Jandrot-Perrus M. Characteristics of the interaction between thrombin exosite I and the sequence 269–287 [correction of 269–297] of platelet glycoprotein Ib alpha. *Thromb Haemost* 1998;80:310–315. [PubMed: 9716158]
18. Jandrot-Perrus M, Bouton MC, Lanza F, Guillin MC. Thrombin interaction with platelet membrane glycoprotein Ib. *Semin Thromb Hemost* 1996;22:151–156. [PubMed: 8807712]
19. Jandrot-Perrus M, Huisse MG, Krstenansky JL, Bezeaud A, Guillin MC. Effect of the hirudin carboxy-terminal peptide 54–65 on the interaction of thrombin with platelets. *Thromb Haemost* 1991;66:300–305. [PubMed: 1746000]
20. Karshikov A, Bode W, Tulinsky A, Stone SR. Electrostatic interactions in the association of proteins: an analysis of the thrombin-hirudin complex. *Protein Sci* 1992;1:727–735. [PubMed: 1363935]
21. De Marco L, Mazzucato M, Masotti A, Ruggeri ZM. Localization and characterization of an alpha-thrombin-binding site on platelet glycoprotein Ib alpha. *J Biol Chem* 1994;269:6478–6484. [PubMed: 8119999]
22. Jandrot-Perrus M, Clemetson KJ, Huisse MG, Guillin MC. Thrombin interaction with platelet glycoprotein Ib: effect of glycofibrin on thrombin specificity. *Blood* 1992;80:2781–2786. [PubMed: 1450405]
23. De Candia E, Hall SW, Rutella S, Landolfi R, Andrews RK, De Cristofaro R. Binding of thrombin to glycoprotein Ib accelerates the hydrolysis of Par-1 on intact platelets. *J Biol Chem* 2001;276:4692–4698. [PubMed: 11084032]
24. De Cristofaro R, De Filippis V. Interaction of the 268–282 region of glycoprotein Ib alpha with the heparin-binding site of thrombin inhibits the enzyme activation of factor VIII. *Biochem J* 2003;373:593–601. [PubMed: 12689334]
25. Celikel R, McClintock RA, Roberts JR, Mendolicchio GL, Ware J, Varughese KI, Ruggeri ZM. Modulation of alpha-thrombin function by distinct interactions with platelet glycoprotein Ib alpha. *Science* 2003;301:218–221. [PubMed: 12855810]
26. Dumas JJ, Kumar R, Sehra J, Somers WS, Mosyak L. Crystal structure of the GPIIb/IIIa-thrombin complex essential for platelet aggregation. *Science* 2003;301:222–226. [PubMed: 12855811]
27. Vali Z, Scheraga HA. Localization of the binding site on fibrin for the secondary binding site of thrombin. *Biochemistry* 1988;27:1956–1963. [PubMed: 3378041]
28. Lovely RS, Moaddel M, Farrell DH. Fibrinogen gamma' chain binds thrombin exosite II. *J Thromb Haemost* 2003;1:124–131. [PubMed: 12871549]
29. Pospisil CH, Stafford AR, Fredenburgh JC, Weitz JI. Evidence that both exosites on thrombin participate in its high affinity interaction with fibrin. *J Biol Chem* 2003;278:21584–21591. [PubMed: 12682049]
30. Mosesson MW, Siebenlist KR, Meh DA. The structure and biological features of fibrinogen and fibrin. *Ann N Y Acad Sci* 2001;936:11–30. [PubMed: 11460466]
31. Chung DW, Davie EW. gamma and gamma' chains of human fibrinogen are produced by alternative mRNA processing. *Biochemistry* 1984;23:4232–4236. [PubMed: 6091742]

32. Fornace AJ Jr, Cummings DE, Comeau CM, Kant JA, Crabtree GR. Structure of the human gamma-fibrinogen gene. Alternate mRNA splicing near the 3' end of the gene produces gamma A and gamma B forms of gamma-fibrinogen. *J Biol Chem* 1984;259:12826–12830. [PubMed: 6092346]
33. Lovely RS, Falls LA, Al-Mondhiry HA, Chambers CE, Sexton GJ, Ni H, Farrell DH. Association of gammaA/gamma' fibrinogen levels and coronary artery disease. *Thromb Haemost* 2002;88:26–31. [PubMed: 12152671]
34. Mosesson MW, Finlayson JS. Biochemical and chromatographic studies of certain activities associated with human fibrinogen preparations. *J Clin Invest* 1963;42:747–755. [PubMed: 13936148]
35. Mannila MN, Lovely RS, Kazmierczak SC, Eriksson P, Samnegard A, Farrell DH, Hamsten A, Silveira A. Elevated plasma fibrinogen gamma' concentration is associated with myocardial infarction: effects of variation in fibrinogen genes and environmental factors. *J Thromb Haemost* 2007;5:766–773. [PubMed: 17263791]
36. Grunbacher G, Weger W, Marx-Neuhold E, Pilger E, Koppel H, Wascher T, Marz W, Renner W. The fibrinogen gamma (FGG) 10034C>T polymorphism is associated with venous thrombosis. *Thromb Res* 2007;121:33–36. [PubMed: 17445871]
37. Uitte de Willige S, de Visser MC, Houwing-Duistermaat JJ, Rosendaal FR, Vos HL, Bertina RM. Genetic variation in the fibrinogen gamma gene increases the risk for deep venous thrombosis by reducing plasma fibrinogen gamma' levels. *Blood* 2005;106:4176–4183. [PubMed: 16144795]
38. Sabo TM, Farrell DH, Maurer MC. Conformational analysis of gamma' peptide (410–427) interactions with thrombin anion binding exosite II. *Biochemistry* 2006;45:7434–7445. [PubMed: 16768439]
39. Pineda AO, Chen ZW, Marino F, Mathews FS, Mosesson MW, Di Cera E. Crystal structure of thrombin in complex with fibrinogen gamma' peptide. *Biophys Chem* 2006;125:556–559. [PubMed: 16962697]
40. Stone SR, Hofsteenge J. Kinetics of the inhibition of thrombin by hirudin. *Biochemistry* 1986;25:4622–4628. [PubMed: 3768302]
41. Trumbo TA, Maurer MC. Examining thrombin hydrolysis of the factor XIII activation peptide segment leads to a proposal for explaining the cardioprotective effects observed with the factor XIII V34L mutation. *J Biol Chem* 2000;275:20627–20631. [PubMed: 10801785]
42. Croy CH, Koeppel JR, Bergqvist S, Komives EA. Allosteric changes in solvent accessibility observed in thrombin upon active site occupation. *Biochemistry* 2004;43:5246–5255. [PubMed: 15122890]
43. Ni F, Zhu Y, Scheraga HA. Thrombin-Bound Structures of Designed Analogs of Human Fibrinopeptide-a Determined by Quantitative Transferred Noe Spectroscopy - a New Structural Basis for Thrombin Specificity. *Journal of Molecular Biology* 1995;252:656–671. [PubMed: 7563081]
44. Ni F, Meinwald YC, Vasquez M, Scheraga HA. High-resolution NMR studies of fibrinogen-like peptides in solution: structure of a thrombin-bound peptide corresponding to residues 7–16 of the A alpha chain of human fibrinogen. *Biochemistry* 1989;28:3094–3105. [PubMed: 2742827]
45. Ni F, Ripoll DR, Martin PD, Edwards BF. Solution structure of a platelet receptor peptide bound to bovine alpha-thrombin. *Biochemistry* 1992;31:11551–11557. [PubMed: 1332764]
46. Martin PD, Robertson W, Turk D, Huber R, Bode W, Edwards BF. The structure of residues 7–16 of the A alpha-chain of human fibrinogen bound to bovine thrombin at 2.3-A resolution. *J Biol Chem* 1992;267:7911–7920. [PubMed: 1560020]
47. Mathews II, Padmanabhan KP, Ganesh V, Tulinsky A, Ishii M, Chen J, Turck CW, Coughlin SR, Fenton JW 2nd. Crystallographic structures of thrombin complexed with thrombin receptor peptides: existence of expected and novel binding modes. *Biochemistry* 1994;33:3266–3279. [PubMed: 8136362]
48. Ni F. 2-Dimensional Transferred Nuclear-Overhauser-Effects with Incomplete Averaging of Free-Ligand and Bound-Ligand Resonances. *Journal of Magnetic Resonance Series B* 1995;106:147–155.
49. Ni F, Scheraga HA. Use of the Transferred Nuclear Overhauser Effect to Determine the Conformations of Ligands Bound to Proteins. *Accounts of Chemical Research* 1994;27:257–264.
50. Friedrich R, Panizzi P, Kawabata S, Bode W, Bock PE, Fuentes-Prior P. Structural basis for reduced staphylocoagulase-mediated bovine prothrombin activation. *J Biol Chem* 2006;281:1188–1195. [PubMed: 16230338]

51. Sabo TM, Brasher PB, Maurer MC. Perturbations in Factor XIII Resulting from Activation and Inhibition Examined by Solution Based Methods and Detected by MALDI-TOF MS. *Biochemistry* 2007;46:10089–10101. [PubMed: 17691819]
52. Chen J, Smith DL. Amide hydrogen exchange shows that malate dehydrogenase is a folded monomer at pH 5. *Protein Sci* 2001;10:1079–1083. [PubMed: 11316888]
53. Wang L, Lane LC, Smith DL. Detecting structural changes in viral capsids by hydrogen exchange and mass spectrometry. *Protein Sci* 2001;10:1234–1243. [PubMed: 11369862]
54. Turner BT Jr, Maurer MC. Evaluating the roles of thrombin and calcium in the activation of coagulation factor XIII using H/D exchange and MALDI-TOF MS. *Biochemistry* 2002;41:7947–7954. [PubMed: 12069584]
55. Lebowitz J, Lewis MS, Schuck P. Modern analytical ultracentrifugation in protein science: a tutorial review. *Protein Sci* 2002;11:2067–2079. [PubMed: 12192063]
56. Resing KA, Hoofnagle AN, Ahn NG. Modeling deuterium exchange behavior of ERK2 using pepsin mapping to probe secondary structure. *J Am Soc Mass Spectrom* 1999;10:685–702. [PubMed: 10439507]
57. Arrington CB, Robertson AD. Correlated motions in native proteins from MS analysis of NH exchange: evidence for a manifold of unfolding reactions in ovomucoid third domain. *J Mol Biol* 2000;300:221–232. [PubMed: 10864511]
58. Clarke J, Itzhaki LS. Hydrogen exchange and protein folding. *Curr Opin Struct Biol* 1998;8:112–118. [PubMed: 9519304]
59. Swint-Kruse L, Robertson AD. Temperature and pH dependences of hydrogen exchange and global stability for ovomucoid third domain. *Biochemistry* 1996;35:171–180. [PubMed: 8555171]
60. Bai Y, Milne JS, Mayne L, Englander SW. Primary structure effects on peptide group hydrogen exchange. *Proteins* 1993;17:75–86. [PubMed: 8234246]
61. Mandell JG, Falick AM, Komives EA. Identification of protein-protein interfaces by decreased amide proton solvent accessibility. *Proc Natl Acad Sci U S A* 1998;95:14705–14710. [PubMed: 9843953]
62. Mandell JG, Falick AM, Komives EA. Measurement of amide hydrogen exchange by MALDI-TOF mass spectrometry. *Anal Chem* 1998;70:3987–3995. [PubMed: 9784743]
63. Mandell JG, Baerga-Ortiz A, Akashi S, Takio K, Komives EA. Solvent accessibility of the thrombin-thrombomodulin interface. *J Mol Biol* 2001;306:575–589. [PubMed: 11178915]
64. De Cristofaro R, De Candia E, Rutella S, Weitz JI. The Asp(272)-Glu(282) region of platelet glycoprotein I α interacts with the heparin-binding site of α -thrombin and protects the enzyme from the heparin-catalyzed inhibition by antithrombin III. *J Biol Chem* 2000;275:3887–3895. [PubMed: 10660541]
65. Kroh HK, Tans G, Nicolaes GA, Rosing J, Bock PE. Expression of allosteric linkage between the sodium ion binding site and exosite I of thrombin during prothrombin activation. *J Biol Chem* 2007;282:16095–16104. [PubMed: 17430903]
66. Marchese P, Murata M, Mazzucato M, Pradella P, De Marco L, Ware J, Ruggeri ZM. Identification of three tyrosine residues of glycoprotein Ib α with distinct roles in von Willebrand factor and α -thrombin binding. *J Biol Chem* 1995;270:9571–9578. [PubMed: 7721887]
67. Olsen PH, Esmo NL, Esmo CT, Laue TM. Ca²⁺ dependence of the interactions between protein C, thrombin, and the elastase fragment of thrombomodulin. Analysis by ultracentrifugation. *Biochemistry* 1992;31:746–754. [PubMed: 1310045]
68. Lonhienne TG, Jackson CM, Winzor DJ. Thermodynamic non-ideality as an alternative source of the effect of sucrose on the thrombin-catalyzed hydrolysis of peptide p-nitroanilide substrates. *Biophys Chem* 2003;103:259–269. [PubMed: 12727288]
69. Grutter MG, Priestle JP, Rahuel J, Grossenbacher H, Bode W, Hofsteenge J, Stone SR. Crystal structure of the thrombin-hirudin complex: a novel mode of serine protease inhibition. *EMBO J* 1990;9:2361–2365. [PubMed: 2369893]
70. Rydel TJ, Ravichandran KG, Tulinsky A, Bode W, Huber R, Roitsch C, Fenton JW 2nd. The structure of a complex of recombinant hirudin and human α -thrombin. *Science* 1990;249:277–280. [PubMed: 2374926]
71. Rydel TJ, Tulinsky A, Bode W, Huber R. Refined structure of the hirudin-thrombin complex. *J Mol Biol* 1991;221:583–601. [PubMed: 1920434]

72. Vitali J, Martin PD, Malkowski MG, Robertson WD, Lazar JB, Winant RC, Johnson PH, Edwards BF. The structure of a complex of bovine alpha-thrombin and recombinant hirudin at 2.8-Å resolution. *J Biol Chem* 1992;267:17670–17678. [PubMed: 1517214]
73. Ni F, Konishi Y, Scheraga HA. Thrombin-bound conformation of the C-terminal fragments of hirudin determined by transferred nuclear Overhauser effects. *Biochemistry* 1990;29:4479–4489. [PubMed: 2350549]
74. Ni F, Ripoll DR, Purisima EO. Conformational stability of a thrombin-binding peptide derived from the hirudin C-terminus. *Biochemistry* 1992;31:2545–2554. [PubMed: 1547237]
75. Myles T, Le Bonniec BF, Betz A, Stone SR. Electrostatic steering and ionic tethering in the formation of thrombin-hirudin complexes: the role of the thrombin anion-binding exosite-I. *Biochemistry* 2001;40:4972–4979. [PubMed: 11305913]
76. Lovely RS, Rein CM, White TC, Jouihan SA, Boshkov LK, Bakke AC, McCarty OJ, Farrell DH. gammaA/gamma' fibrinogen inhibits thrombin-induced platelet aggregation. *Thromb Haemost* 2008;100:837–846. [PubMed: 18989528]
77. Lancellotti S, Rutella S, De Filippis V, Pozzi N, Rocca B, De Cristofaro R. Fibrinogen-elongated gamma chain inhibits thrombin-induced platelet response, hindering the interaction with different receptors. *J Biol Chem* 2008;283:30193–30204. [PubMed: 18779330]
78. Weeterings C, Adelmeijer J, Myles T, de Groot PG, Lisman T. Glycoprotein Ialpha-mediated platelet adhesion and aggregation to immobilized thrombin under conditions of flow. *Arterioscler Thromb Vasc Biol* 2006;26:670–675. [PubMed: 16357309]
79. Verhamme IM, Olson ST, Tollefsen DM, Bock PE. Binding of exosite ligands to human thrombin. Re-evaluation of allosteric linkage between thrombin exosites I and II. *J Biol Chem* 2002;277:6788–6798. [PubMed: 11724802]
80. Siebenlist KR, Mosesson MW, Hernandez I, Bush LA, Di Cera E, Shainoff JR, Di Orio JP, Stojanovic L. Studies on the basis for the properties of fibrin produced from fibrinogen-containing gamma' chains. *Blood* 2005;106:2730–2736. [PubMed: 16002430]
81. Lane DA, Philippou H, Huntington JA. Directing thrombin. *Blood* 2005;106:2605–2612. [PubMed: 15994286]
82. Fredenburgh JC, Stafford AR, Weitz JI. Evidence for allosteric linkage between exosites 1 and 2 of thrombin. *J Biol Chem* 1997;272:25493–25499. [PubMed: 9325262]
83. Hogg PJ, Jackson CM, Labanowski JK, Bock PE. Binding of fibrin monomer and heparin to thrombin in a ternary complex alters the environment of the thrombin catalytic site, reduces affinity for hirudin, and inhibits cleavage of fibrinogen. *J Biol Chem* 1996;271:26088–26095. [PubMed: 8824251]
84. Koeppe JR, Seitova A, Mather T, Komives EA. Thrombomodulin tightens the thrombin active site loops to promote protein C activation. *Biochemistry* 2005;44:14784–14791. [PubMed: 16274226]
85. Lange OF, Lakomek NA, Fares C, Schroder GF, Walter KF, Becker S, Meiler J, Grubmuller H, Griesinger C, de Groot BL. Recognition dynamics up to microseconds revealed from an RDC-derived ubiquitin ensemble in solution. *Science* 2008;320:1471–1475. [PubMed: 18556554]
86. Cohly MA, Scheraga HA. Aggregation of thrombin. *Arch Biochem Biophys* 1961;95:428–434. [PubMed: 13880342]
87. Winzor DJ, Scheraga HA. Studies of Chemically Reacting Systems on Sephadex. II. Molecular Weights of Monomers in Rapid Association Equilibrium. *The Journal of Physical Chemistry* 1964;68:338–343.
88. Batt CW, Mikulka TW, Mann KG, Guarracino CL, Altieri RJ, Graham RG, Quigley JP, Wolf JW, Zafonte CW. The purification and properties of bovine thrombin. *J Biol Chem* 1970;245:4857–4862. [PubMed: 5528237]
89. De Marco L, Mazzucato M, Masotti A, Fenton JW 2nd, Ruggeri ZM. Function of glycoprotein Ib alpha in platelet activation induced by alpha-thrombin. *J Biol Chem* 1991;266:23776–23783. [PubMed: 1748654]
90. Lima LM, Becker CF, Giesel GM, Marques AF, Cargnelutti MT, de Oliveira Neto M, Queiroz Monteiro R, Verli H, Polikarpov I. Structural and thermodynamic analysis of thrombin:suramin interaction in solution and crystal phases. *Biochim Biophys Acta* 2009;1794:873–881. [PubMed: 19332154]

91. Anderson PJ. A dimeric form of prothrombin on membrane surfaces. *Biochem J* 1998;336(Pt 3):631–638. [PubMed: 9841875]
92. Adam F, Bouton MC, Huisse MG, Jandrot-Perrus M. Thrombin interaction with platelet membrane glycoprotein Ib alpha. *Trends Mol Med* 2003;9:461–464. [PubMed: 14604821]
93. Mann KG, Brummel-Ziedins K, Orfeo T, Butenas S. Models of blood coagulation. *Blood Cells Mol Dis* 2006;36:108–117. [PubMed: 16500122]

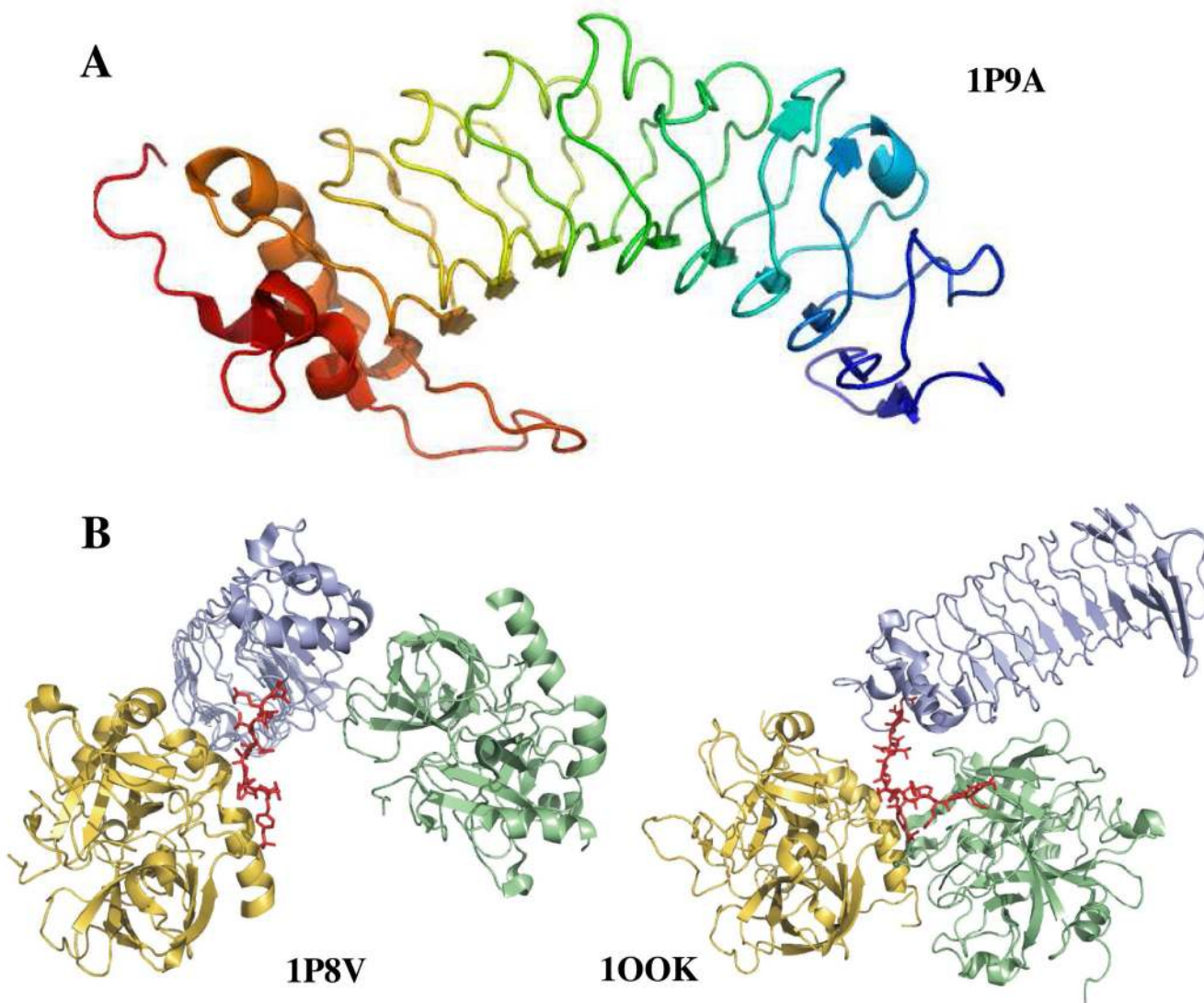


Figure 1.

A. Structure of glycoprotein Iba (1–266) (1P9A) (25). The highly anionic C-terminal tail (269–286) is not modeled in this structure. The most prominent feature of the structure is the leucine rich repeat domain that forms the curvature of the protein. **B.** Two views of GpIba binding to thrombin. The crystal structures depict similar ABE-II interactions but differ in the depiction of thrombin dimerization. In blue is GpIba and the anionic C-terminal tail is in red. Thrombin in gold binds to the C-terminal tail, while thrombin in green binds at different interfaces. The structure on the left is 1P8V (26), the structure on the right 100K (27). These figures were created using PyMol (11).

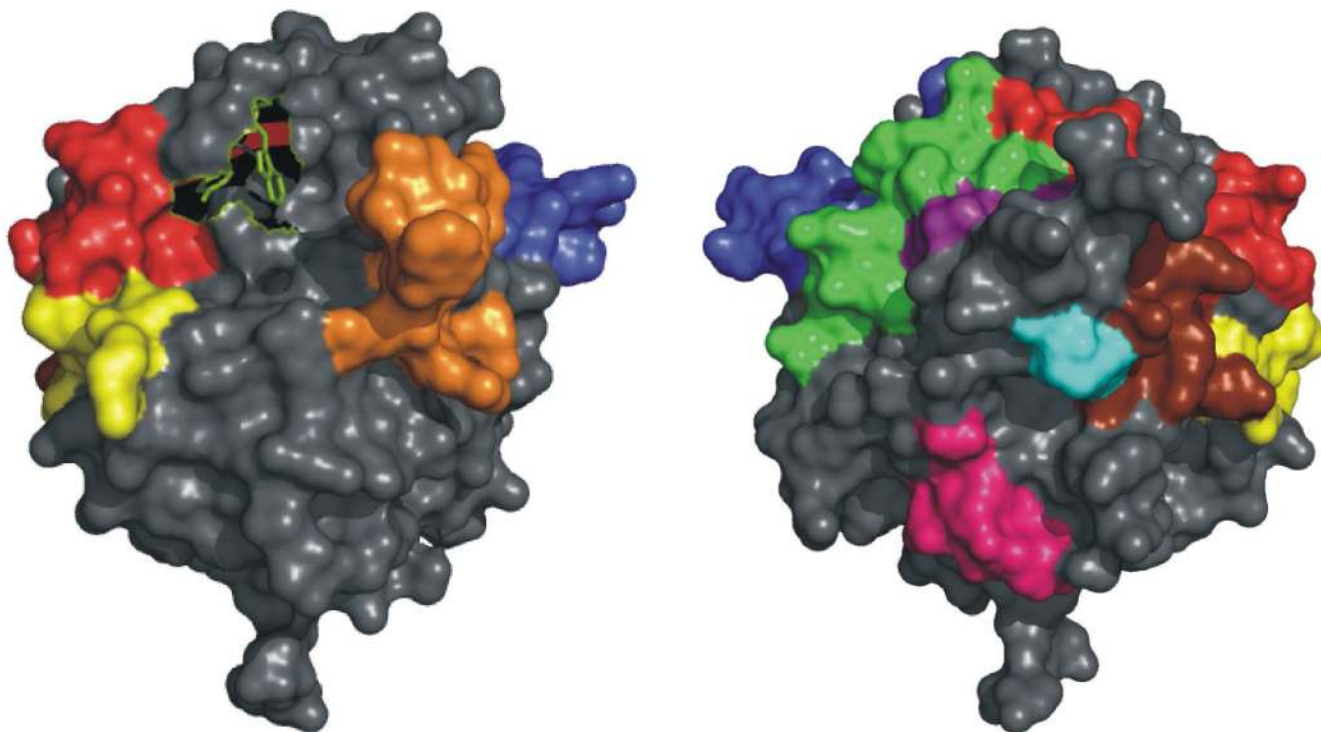
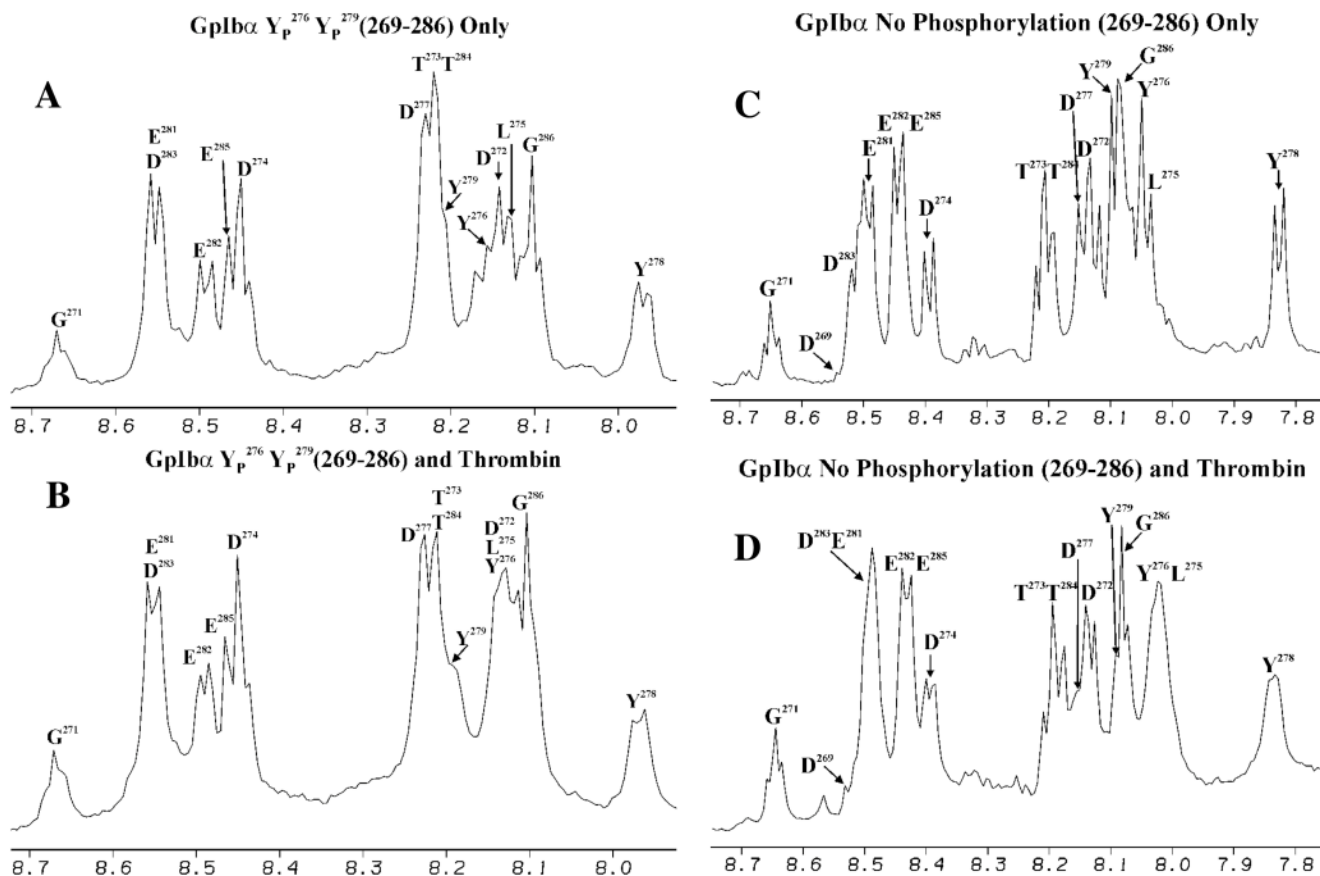
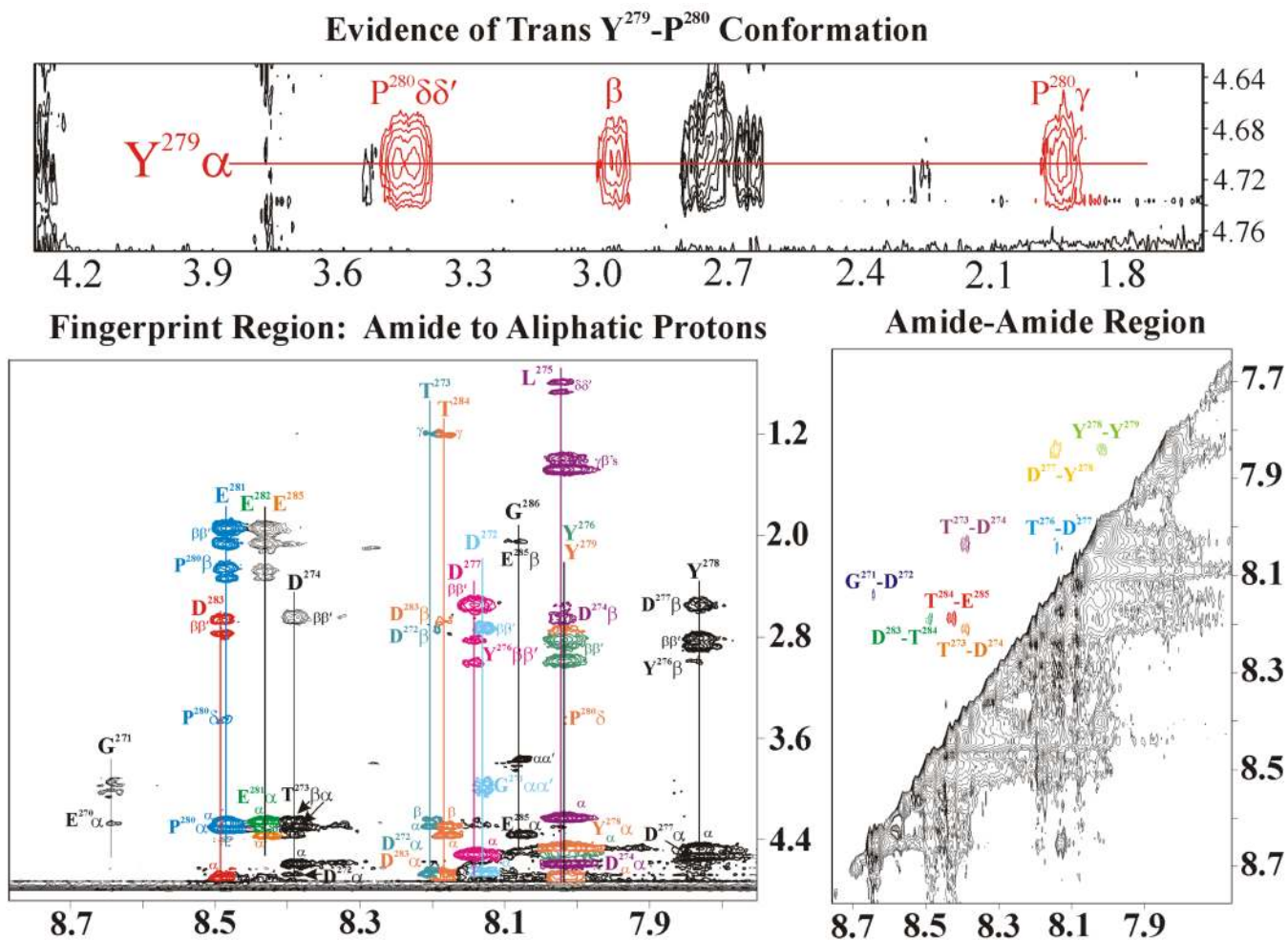


Figure 2. Depiction of the coverage obtained from a peptic digest of IIa (2AID) relative to Y^{60A} and W^{60D}, which are green. Left: Looking onto the active site of IIa. Right: The structure rotated 180° about the y-axis. Peptides from the peptic digest are colored as follows: the ABE-II regions 85–99 (red) and 173–181 (yellow), the ABE-I peptide 65–84 (blue), the autolysis loop containing peptide 135–149D (orange), 106–116 (green), the A-chain peptide –13 to –4 (cyan), 202–207 (pink), and 46–52 (purple). The additional ABE-II region not observed in the present HDX study, 233–240, is in brown.

**Figure 3.**

Line broadening spectra for the GpIb α peptides in the presence of thrombin. All NMR samples are in 25 mM H₃PO₄, 150 mM NaCl, 200 μ M EDTA, and 10% D₂O, pH 5.6. **A.** 1D NMR spectrum for 1.5 mM Y_P²⁷⁶Y_P²⁷⁹ peptide in solution. **B.** 1D NMR spectrum for 1.5 mM Y_P²⁷⁶Y_P²⁷⁹ peptide in the presence of 0.148 mM IIa. The amide protons that display the most line broadening are D²⁷⁴-Y_P²⁷⁹. A moderate amount of line broadening is evident for the NHs of E²⁸¹-D²⁸³. Line broadening is not observed for the amide protons of D²⁶⁹-T²⁷³ and T²⁸⁴-G²⁸⁶. **C.** 1D NMR spectrum for 1.5 mM unphosphorylated peptide in solution. **D.** 1D NMR spectrum for 1.5 mM unphosphorylated peptide in the presence of 0.150 mM IIa. The amide protons that display the most line broadening are D²⁷⁴-Y_P²⁷⁹, E²⁸¹, and D²⁸³. A moderate amount of line broadening is evident for the NHs of E²⁸² and E²⁸⁵. Line broadening is not observed for the amide protons of D²⁶⁹-T²⁷³, T²⁸⁴, and G²⁸⁶.

**Figure 4.**

2D trNOESY spectrum of the unphosphorylated GpIb α (1.5 mM) bound to IIa (0.150 mM). The top panel illustrates the evidence for a *trans* Y²⁷⁹-P²⁸⁰ peptide bond. The bottom panels demonstrate that GpIb α exists in an extended structure when bound to IIa due to the presence of only nearest neighbor NOEs. The NOEs are color coded according to the corresponding residue.

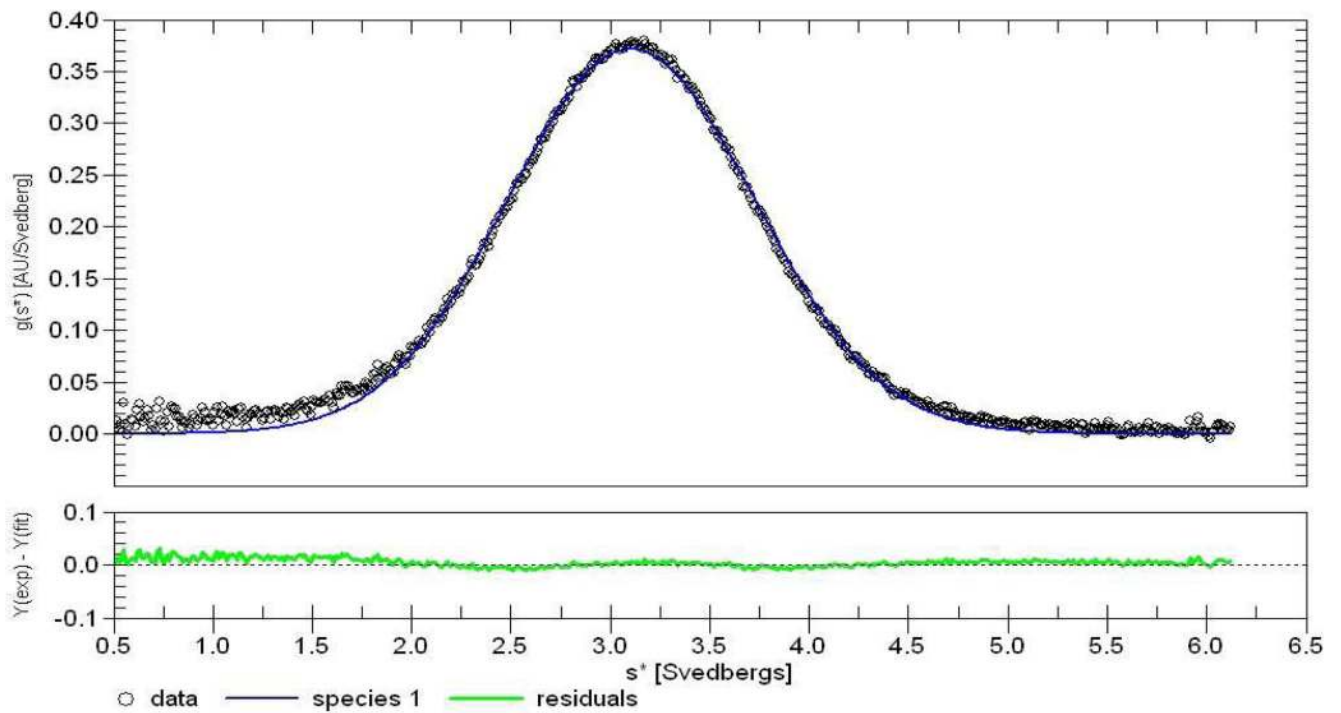


Figure 5. Sedimentation velocity AUC of Gplba peptide (14 μM) and Ila (7 μM). Analysis was performed at 280 nm, 20 $^{\circ}\text{C}$, and 60,000 rpm in 150 mM NaCl, 25 mM phosphate, pH 7.4. The apparent Svedberg constant distribution centered at 3.2 S with the residuals resulting from a global fit of the data using the DCDT+ program.

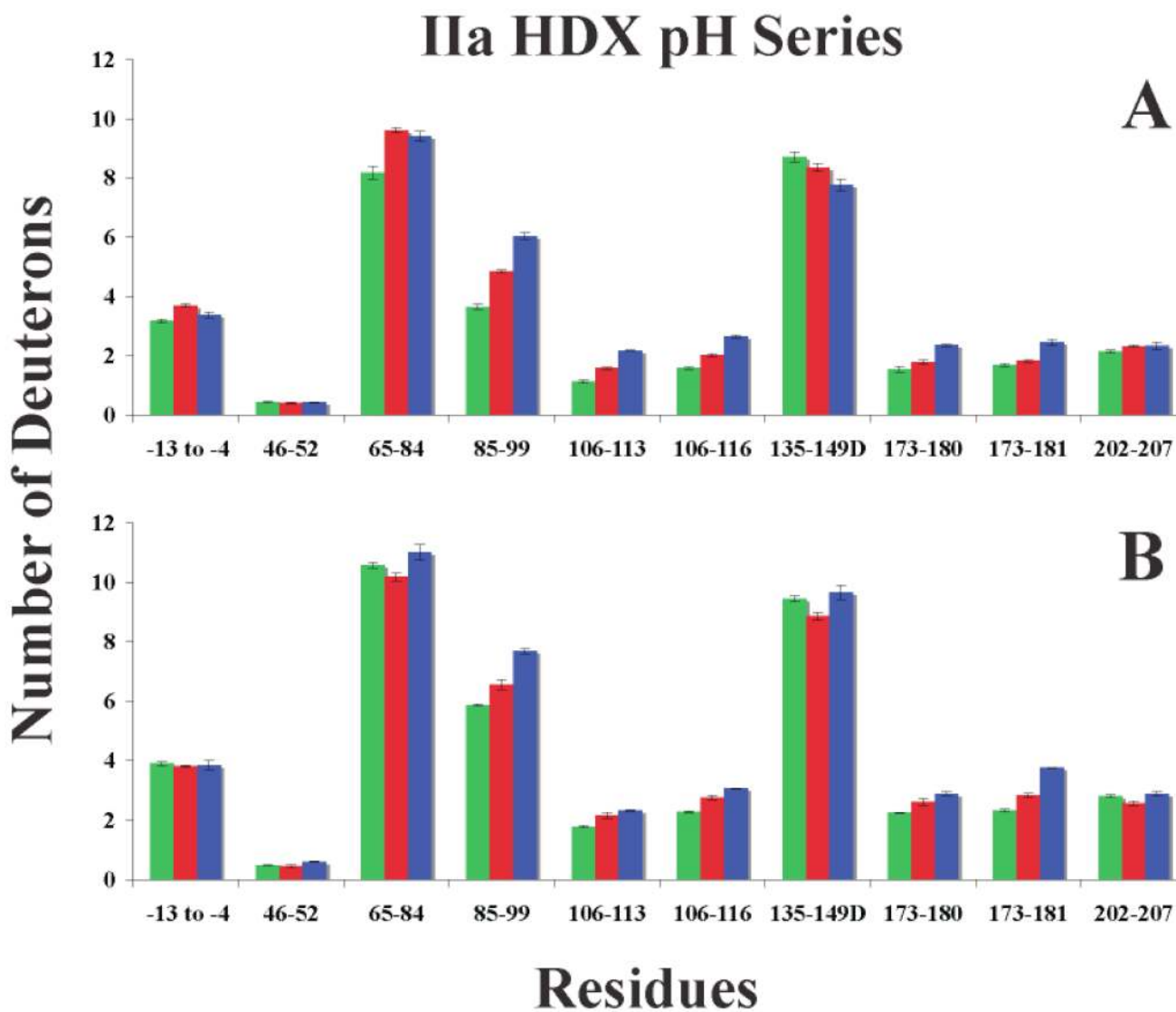
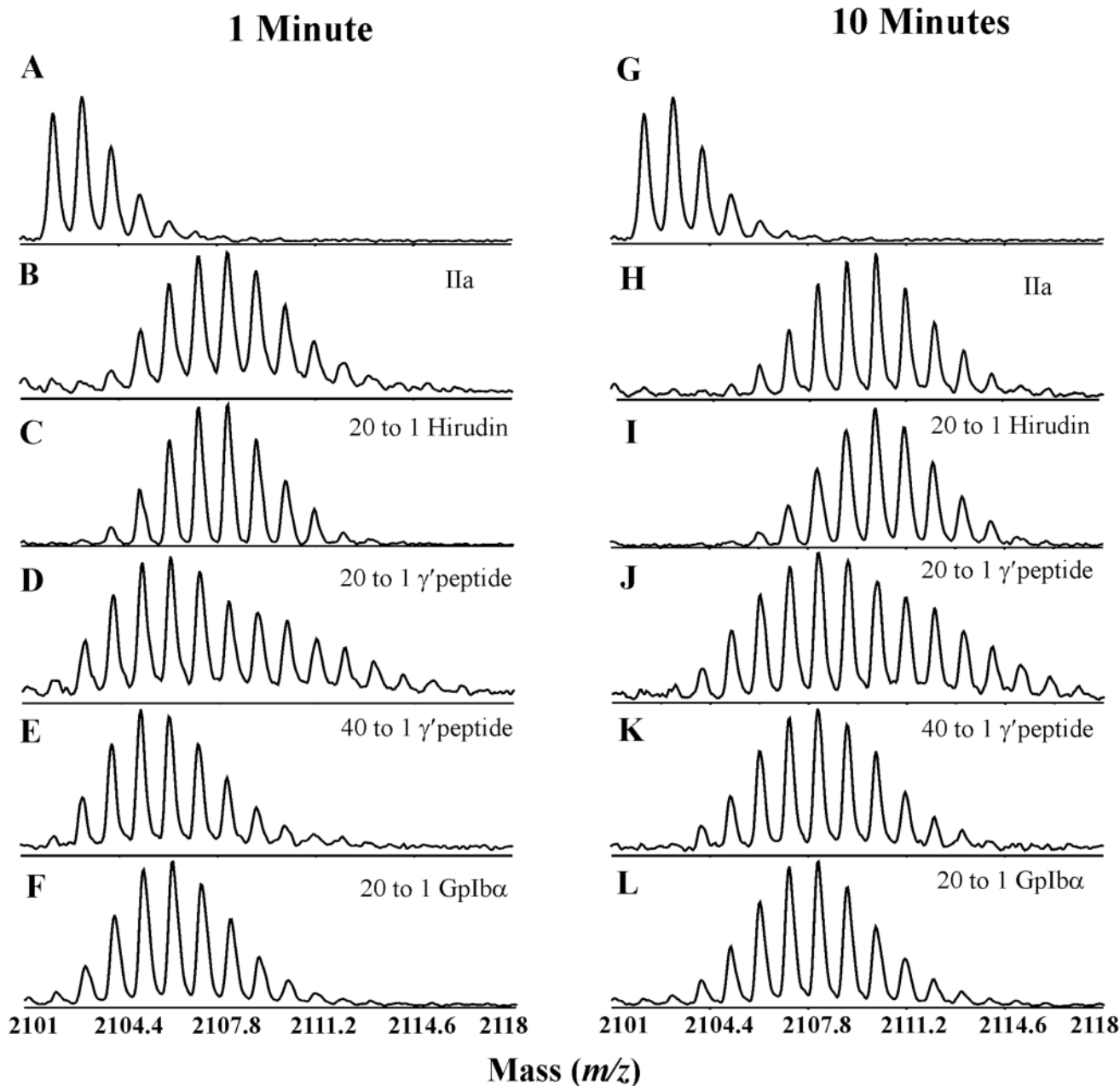


Figure 6. Graphs of deuterium incorporation for IIa at pH 5.6 (green), pH 6.5 (red), and pH 7.5 (blue). **A.** One minute of HDX and **B.** Ten minutes of HDX. Errors correspond to the standard deviation of the mean for three independent experiments.

Residues 85-99

**Figure 7.**

Mass spectra of residues 85–99 after one and ten minutes of HDX. All of the depicted HDX experiments are with 50 μM IIa in 150 mM NaCl, 25 mM NaH_2PO_3 , pH 6.5. **A and G:** the undeuterated peak cluster, **B and H:** IIa spectra in the absence of ligands, **C and I:** 20:1 hirudin (54–65) bound to IIa, **D and J:** 20:1 γ' peptide bound to IIa, **E and K:** 40:1 γ' peptide bound to IIa, and **F and L:** 20:1 GpIb α peptide bound to IIa. At the ratio of 20:1, both the γ' and GpIb α peptides exhibit a similar degree of protection for these ABE-II residues; however in spectra D and J a small shoulder is present suggesting the presence of two IIa populations. Increasing the amount of γ' peptide completely abrogates the shoulder suggesting fully occupied ABE-II.

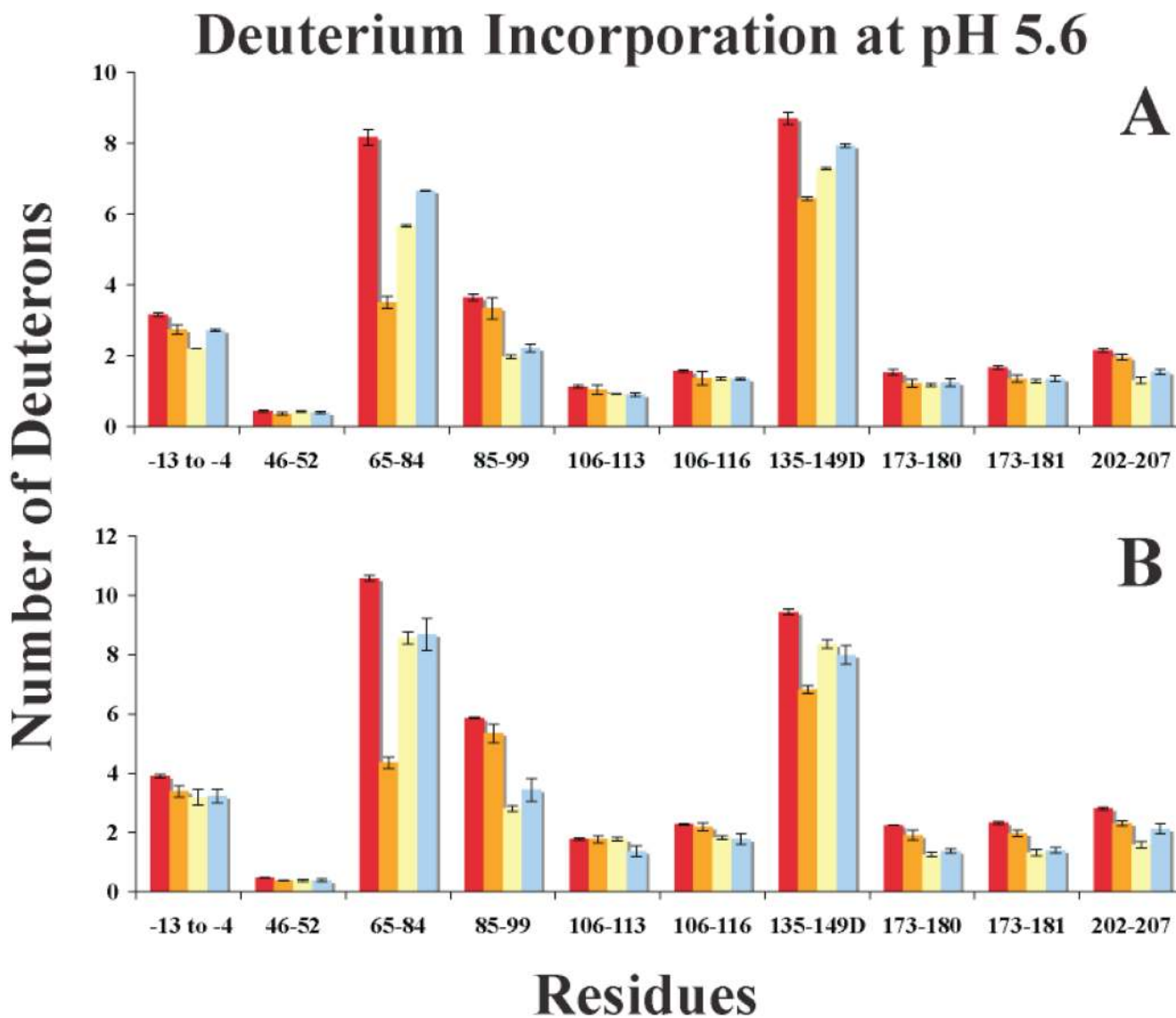
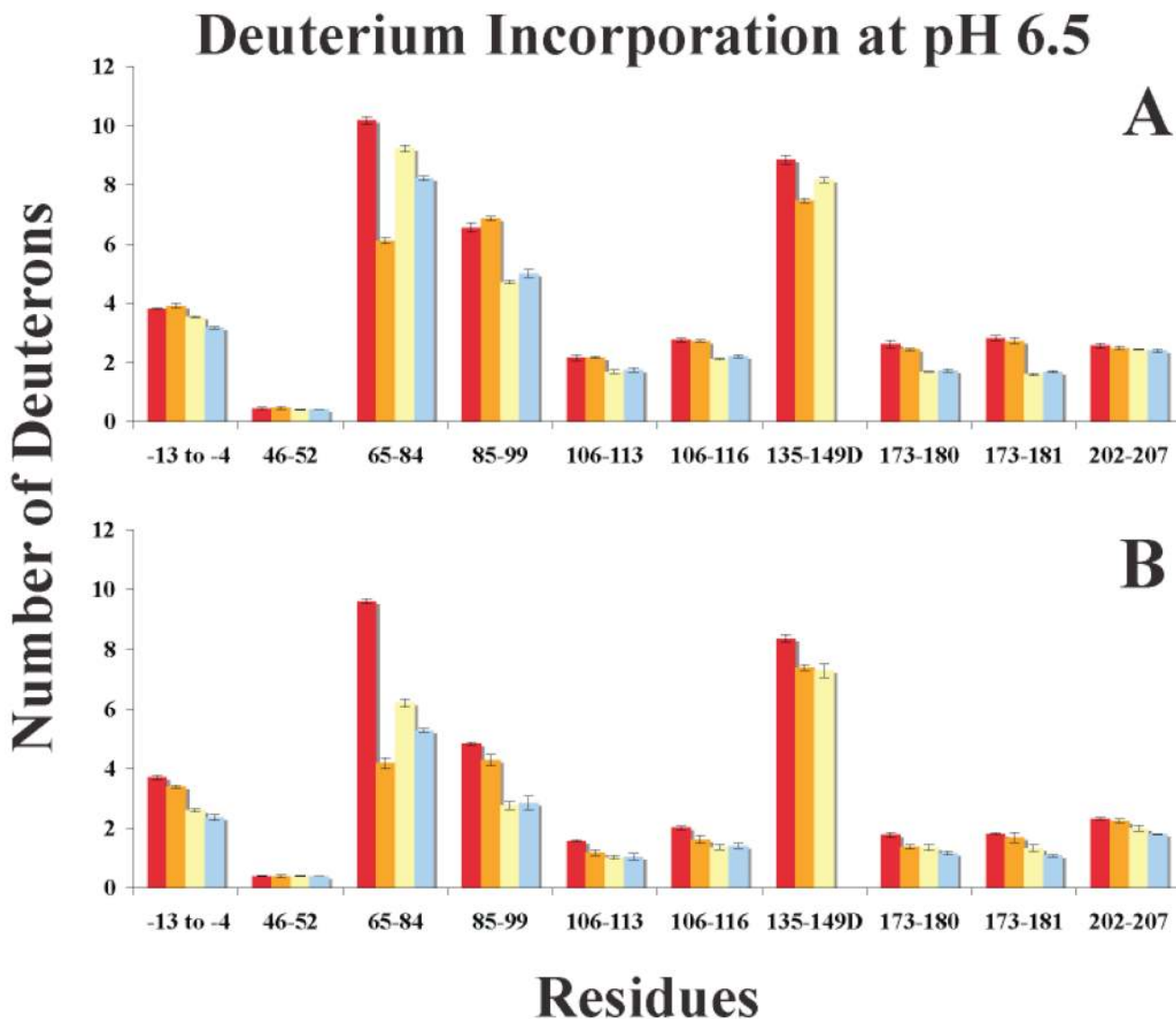
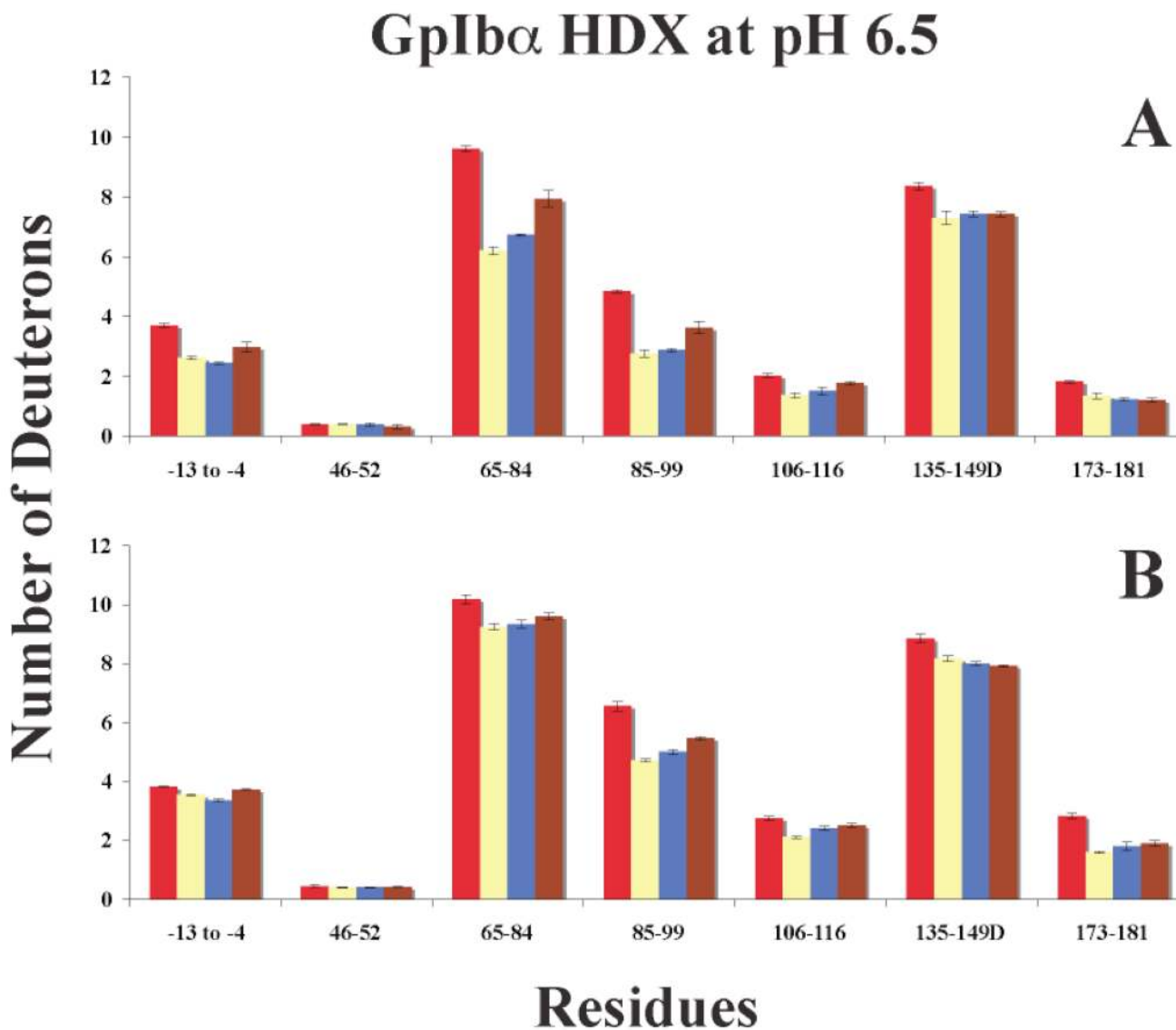


Figure 8.

Graphs of deuterium incorporation for IiA and 20:1 peptide-IiA at pH 5.6. **A.** One Minute of HDX and **B.** Ten Minutes of HDX. The bars in the graph correspond to IiA alone (red), Hir-IiA (orange), GpIb α -IiA (yellow), and γ' peptide-IiA (blue). Errors correspond to the standard deviation of the mean for three independent experiments.

**Figure 9.**

Graphs of deuterium incorporation for IiA and peptide-IiA at pH 6.5. **A.** One Minute of HDX and **B.** Ten Minutes of HDX. The bars in the graph correspond to IiA alone (red), 20:1 Hir-IiA (orange), 20:1 GpIb α -IiA (yellow) and 40:1 γ' peptide-IiA (blue). Errors correspond to the standard deviation of the mean for three independent experiments.

**Figure 10.**

Graphs of deuterium incorporation for Ia, GpIb α peptide-Ia, and GpIb α protein-Ia at pH 6.5. **A.** One Minute of HDX and **B.** Ten Minutes of HDX. The bars in the graph correspond to Ia alone (red), 20:1 GpIb α peptide-Ia (yellow), 1.6:1 GpIb α 3S protein-Ia (blue), and 1.6:1 GpIb α 2S protein-Ia (brown). Errors correspond to the standard deviation of the mean for three independent experiments.

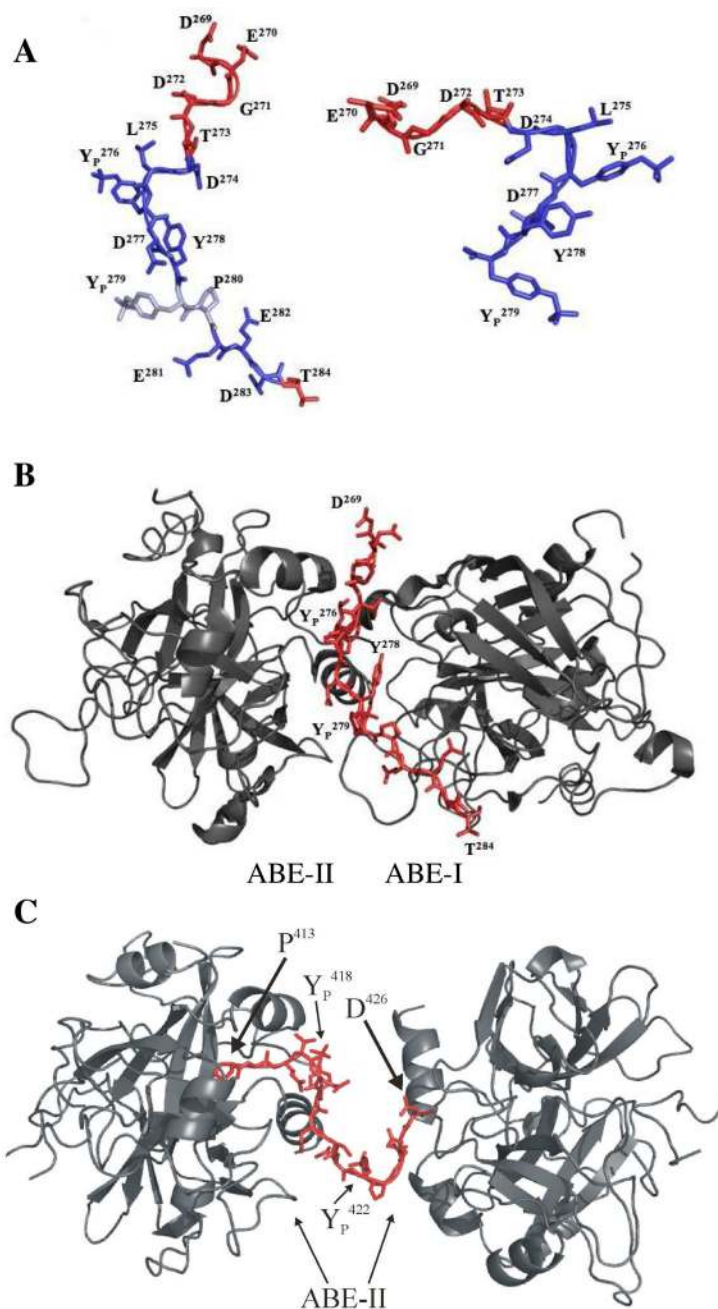


Figure 11.

A. Representation of the GpIb α peptide isolated from thrombin. The peptide on the left is from 100K and the structure on the right is from 1P8V. The individual amino acids are labeled. The *trans* bond between Y_P²⁷⁹-P²⁸⁰ in 100K is in light blue. Residues experiencing line-broadening in the 1D NMR spectra are in blue. **B.** Illustration of GpIb α peptide induced dimerization of thrombin in 100K. The data suggests this structure more accurately represents the NMR results. The C-terminus of GpIb α is in an extended conformation and residues D²⁷⁴-T²⁸⁴ interact with both ABE-II and ABE-I of opposing thrombin monomers. **C.** Illustration of γ' peptide induced dimerization of IIa depicted in the structure 2HWL. The turn

conformation is situated between the two symmetry related IIa monomers, both at ABE-II. These figures were created using PyMol (11).

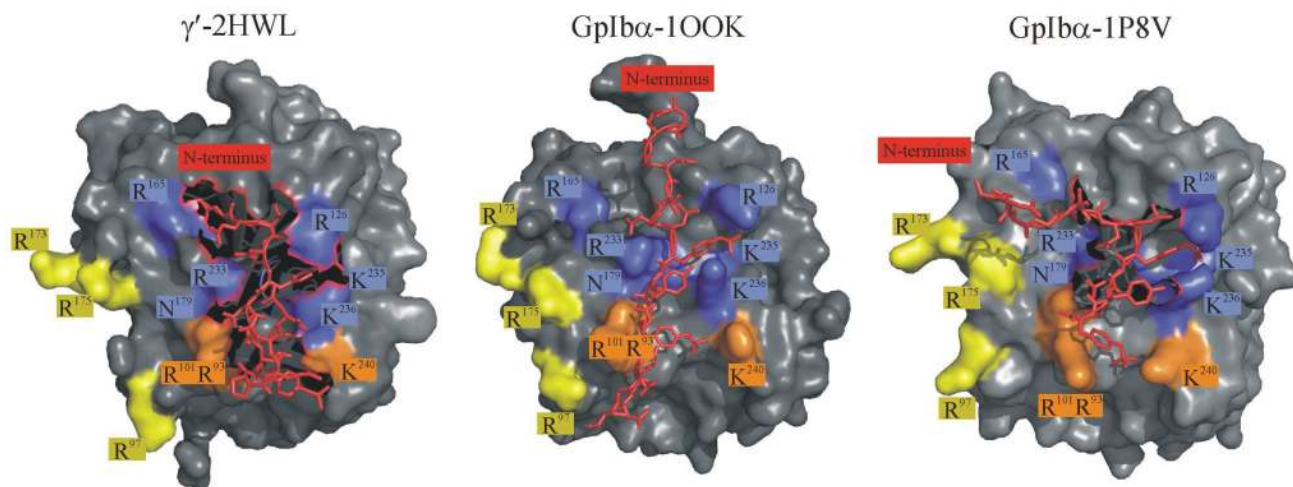


Figure 12.

Crystallographic views of the γ' peptide (2HWL) and Gplb α (100K and 1P8V) bound to thrombin's ABE-II. A prominent feature involves Gplb α Y_P²⁷⁶ and γ' Y_P⁴¹⁸ engulfed by an electropositive pocket formed by IIa R¹²⁶, K²³⁵, and K²³⁶. In addition, Gplb α D²⁷⁷ and γ' D⁴¹⁹ reside near IIa N¹⁷⁹. Finally, Y_P²⁷⁹ and γ' Y_P⁴²² point toward IIa K²³⁶. These structures illustrate similar bound characteristics for the central regions of the bound peptide, however differ in regard to the termini. These figures were created using PyMol (11).

Table 1

Peptides Utilized in IIa Binding Studies

| | |
|---|---|
| GpIb α Y _P ²⁷⁶ Y _P ²⁷⁸ Y _P ²⁷⁹ | ²⁶⁹ DEGDTDLY _P DY _P Y _P PEEDTEG ²⁸⁶ |
| GpIb α Y _P ²⁷⁶ Y ²⁷⁸ Y _P ²⁷⁹ | ²⁶⁹ DEGDTDLY _P DY _P PEEDTEG ²⁸⁶ |
| GpIb α Y ²⁷⁶ Y ²⁷⁸ Y ²⁷⁹ | ²⁶⁹ DEGDTDLYDYYPEEDTEG ²⁸⁶ |
| γ' Y _P ⁴¹⁸ Y _P ⁴²² | ⁴¹⁰ PEHPAETELY _P DSL _P Y _P PEDDL ⁴²⁷ |
| Hirudin Y _S ⁶³ | ⁵⁴ GDFEEIPEEY _S LQ ⁶⁵ |

Table 2

Analytical Ultracentrifugation Data

| | Sedimentation Coefficient (S) | Molecular Weight (kDa) |
|---------------------------|-------------------------------|------------------------|
| I α alone | 3.11 \pm 0.00 | 36.2 \pm 0.3 |
| Gplb α :I α | | |
| 1:1 | 3.19 \pm 0.02 | 37.3 \pm 0.0 |
| 2:1 | 3.18 \pm 0.01 | 37.5 \pm 0.9 |
| 5:1 | 3.27 \pm 0.05 | 37.9 \pm 0.7 |
| 10:1 | 3.25 \pm 0.02 | 37.2 \pm 1.1 |
| 20:1 | 3.22 \pm 0.01 | 38.1 \pm 1.4 |
| γ :I α | | |
| 1:1 | 3.19 \pm 0.02 | 39.3 \pm 1.0 |
| 2:1 | 3.22 \pm 0.03 | 38.6 \pm 0.4 |
| 5:1 | 3.23 \pm 0.03 | 39.9 \pm 0.2 |
| 10:1 | 3.27 \pm 0.01 | 39.9 \pm 0.6 |
| 20:1 | 3.28 \pm 0.01 | 40.9 \pm 0.7 |

Table 3
Changes in % Deuteration for GpIb α -IIa and γ' peptide-IIa at 1 and 10 Minutes Relative to IIa pH 5.6^a

| Residues | GpIb α | | | γ' | | | Hirudin | | |
|-----------|---------------------------|--------------|--------------|-------------|--------------|--------------|--------------|--------------|--------------|
| | 1 m | 10 m | 10 m | 1 m | 10 m | 10 m | 1 min | 10 m | 10 m |
| -13 to -4 | -11.4 ^b | -8.5 | -8.0 | -5.2 | -8.0 | -8.0 | -4.9 | -6.1 | -6.1 |
| 46-52 | -0.2 | -1.5 | -1.4 | -0.5 | -1.4 | -1.4 | -1.1 | -1.4 | -1.4 |
| 65-84 | -11.5 | -9.3 | -8.7 | -6.9 | -8.7 | -8.7 | -21.4 | -28.5 | -28.5 |
| 85-99 | -11.1 | -20.1 | -16.0 | -9.4 | -16.0 | -16.0 | -2.0 | -3.4 | -3.4 |
| 106-113 | -3.1 | -6.0 | -6.1 | -3.3 | -6.1 | -6.1 | -1.3 | -0.1 | -0.1 |
| 106-116 | -2.2 | -4.6 | -5.1 | -2.4 | -5.1 | -5.1 | -2.1 | -0.9 | -0.9 |
| 135-149D | -7.3 | -5.6 | -7.5 | -4.0 | -7.5 | -7.5 | -11.7 | -13.5 | -13.5 |
| 173-180 | -4.7 | -12.7 | -11.2 | -3.7 | -11.2 | -11.2 | -4.0 | -4.4 | -4.4 |
| 173-181 | -4.4 | -11.5 | -10.6 | -3.8 | -10.6 | -10.6 | -3.8 | -4.1 | -4.1 |
| 202-207 | -12.5 | -17.9 | -10.0 | -8.8 | -10.0 | -10.0 | -2.8 | -7.4 | -7.4 |

^aThe % change for a particular peptide is calculated as described in (38).

^bThe values in bold represent significant changes in deuteration of greater than -4.5%.

Table 4
Changes in % Deuteration for Peptide and Protein Binding to Ila at 1 and 10 Minutes Relative to Free Ila pH 6.5^a

| Residues | 20:1 | | | 40:1 | | | 1.6:1 | | | | | | | | |
|-----------|--------------------------|--------------|--------------|--------------|--------------|--------------|--------------|--------------|-------------|--------------|-----|------|----------|------|-----|
| | Hirudin | | | Gplba | | | γ | | | Gplba 3S | | | Gplba 2S | | |
| | 1 m | 10 m | 1 m | 10 m | 1 m | 10 m | 1 m | 10 m | 1 m | 10 m | 1 m | 10 m | 1 m | 10 m | 1 m |
| -13 to -4 | -3.5 | 1.2 | -12.8 | -3.2 | -15.8 | -7.6 | -14.8 | -5.4 | -8.5 | -1.2 | | | | | |
| 46-52 | -0.2 | 0.3 | 0.0 | -0.6 | -0.3 | -0.6 | -0.2 | -0.6 | -1.4 | -0.3 | | | | | |
| 65-84 | -24.9^b | -18.6 | -15.6 | -4.3 | -19.9 | -8.9 | -13.2 | -3.9 | -7.7 | -2.7 | | | | | |
| 85-99 | -3.6 | 2.0 | -13.7 | -12.1 | -13.1 | -10.2 | -13.0 | -10.3 | -8.0 | -7.2 | | | | | |
| 106-113 | -5.8 | 0.1 | -8.4 | -7.0 | -7.9 | -6.1 | n/a | n/a | n/a | n/a | | | | | |
| 106-116 | -4.1 | -0.3 | -6.7 | -6.6 | -6.2 | -5.6 | -2.6 | -3.6 | -2.6 | -2.6 | | | | | |
| 135-149D | -5.0 | -7.1 | -5.5 | -3.6 | n/a | n/a | -4.8 | -4.4 | -4.8 | -4.8 | | | | | |
| 173-180 | -5.1 | -2.2 | -5.4 | -11.9 | -7.8 | -11.5 | n/a | n/a | n/a | n/a | | | | | |
| 173-181 | -1.6 | -1.1 | -5.5 | -14.0 | -8.4 | -13.0 | -7.0 | -11.6 | -7.0 | -10.6 | | | | | |
| 202-207 | -1.0 | -1.2 | -4.7 | -1.9 | -7.6 | -2.5 | n/a | n/a | n/a | n/a | | | | | |

^aThe % change for a particular peptide is calculated as described in (38)

^bThe values in bold represent significant changes in deuteration of greater than -4.5%.