Ligation of the Fibrin-binding Domain by β-Strand Addition Is Sufficient for Expansion of Soluble Fibronectin*

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How fibronectin (FN) converts from a compact plasma protein to a fibrillar component of extracellular matrix is not understood. "Functional upstream domain" (FUD), a polypeptide based on F1 adhesin of Streptococcus pyogenes, binds by anti-parallel β-strand addition to discontinuous sets of N-terminal FN type I modules, 1–5FNI of the fibrin-binding domain and 8–9FNI of the gelatin-binding domain. Such binding blocks assembly of FN. To learn whether ligation of 2–5FNI, 8–9FNI, or the two sets in combination is important for inhibition, we tested "high affinity downstream domain" (HADD), which binds by β-strand addition to the continuous set of FNIII modules, 1–5FNIII, comprising the fibrin-binding domain. HADD and FUD were similarly active in blocking fibronectin assembly. Binding of HADD or FUD to soluble plasma FN exposed the epitope to monoclonal antibody mAbIII-10 in the tenth FN type III module (10FNIII) and caused expansion of FN as assessed by dynamic light scattering. Soluble N-terminal constructs truncated after 5FNIII or 2FNIII competed better than soluble FN for binding of FUD or HADD to adsorbed FN, indicating that interactions involving type III modules more C-terminal than 5FNIII limit β-strand addition to 1–5FNIII within intact soluble FN. Preincubation of FN with mAbIII-10 or heparin modestly increased binding to HADD or FUD. Thus, ligation of FNIII modules involved in binding of integrins and glycosaminoglycans, 10FNIII and 12–14FNIII, increases accessibility of 1–5FNIII. AllostERIC loss of constraining interactions among 1–5FNIII, 10FNIII, and 12–14FNIII likely enables assembly of FN into extracellular fibrils.

Fibronectin (FN)2 is a dimeric glycoprotein of blood and extracellular matrix. Each subunit of FN, depicted in Fig. 1A, includes 12 type 1 (FNI), 2 type 2 (FNII), and 15–17 type 3 (FNIII) modules, depending on the differential inclusion of disulfide bonds at the extreme C termini. The N-terminal 70-kDa region (N-5FNIII), which is of particular importance in this paper, includes the N-5FNIII fibrin- and 6FNI-9FNI gelatin-binding domains and is composed of nine FNIII and two FNII modules (Fig. 1A). FNIII modules are found only in chondrates, and tandem FNIII modules are found only in FN (2).

Blood plasma FN is produced mainly by hepatocytes and circulates at near micromolar concentrations (3). Rotary shadowing, sedimentation velocity, and dynamic light scattering (DLS) experiments indicate that plasma FN is in a compact conformation that elongates at increased ionic strength or pH (4, 5). FN conformation can be monitored by monoclonal antibody (mAb) mAbIII-10, which recognizes an epitope in 10FNIII that is cryptic in soluble plasma FN at low ionic strength and exposed when FN is adsorbed to a surface, placed in high ionic strength, or incubated with heparin, gelatin, or gangliosides (6). The compact conformation is likely maintained by interactions among distant modules. Several such interactions have been surmised based on studies of FN fragments, including an intrasubunit interaction between 4FNIII and 3FNIII (7), an inter-subunit interaction between 2–3FNIII and 12–14FNIII (8), and a less defined interaction between N-5FNIII and 12–14FNIII (9, 10). The compact structure has been hypothesized to obscure ligand-binding regions of plasma FN and thereby prevent aberrant interactions in the bloodstream (11, 12).

Diverse cells, including fibroblasts, smooth muscle cells, and adherent platelets, support the assembly of compact, soluble FN into insoluble fibrils that support cell adhesion, growth, and migration (13). FN assembly is a complex process in which FN binds to the cell surface, engages receptors, most notably the α5β1 integrin binding to the RGD motif in 10FNIII, and elongates into fibrils (13, 14). Pieces of FN comprising the fibrin- and gelatin-binding domains or just the fibrin-binding domain, however, bind to cell surface sites of FN assembly and block assembly (15–17). A central question regarding FN assembly is whether FN initially interacts via α5β1 with 10FNIII followed by...
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exposure of N-terminal FNI modules (13) or engages cell surface molecules via N-terminal FNI modules followed by exposure of \(10^6\)FN III (17).

Despite its recent appearance in the animal kingdom, FN functions as a modulator of cell adhesion and extracellular matrix formation from early development onward (18), and it is involved in diverse pathophysiologic processes, including vascular disease (3, 19). Remarkably, multiple bacteria target FN and its FNI modules as a vehicle to adhere to tissues and evade host defense (20). The pathophysiologic importance of bacterial FN interactions is highlighted by the recent finding that *Staphylococcus aureus* recovered from patients with infected intravascular devices are enriched in polymorphisms of FN-binding protein A that increase the affinity of the protein for FN (21).

FN-binding protein A of *S. aureus* and the F1 adhesin (allelic variant SfbI) from *Streptococcus pyogenes* are representative of a large number of bacterial FN-binding proteins found in Gram-positive cocci and spirochetes that interact with the fibrin- and gelatin-binding domains (20, 22). Embedded within F1, going from N to C termini, are the following: a nonrepetitive sequence that binds to \(^8\)-FN (23, 24), five FN-binding repeats (FNBR) each of which can bind to \(^2\)-FN (25, 26), and a nonrepetitive sequence that binds to \(^1\)-FN (Fig. 1B) (25, 27, 28), and FN-binding protein A contains multiple FNBRs that bind to \(^2\)-FN (22). “Functional upstream domain” (FUD), a 49-amino acid polypeptide from the F1 adhesin, binds by anti-parallel \(\beta\)-strand addition to discontinuous sets of FNI modules within N-\(^3\)-FN; the N-terminal nonrepetitive upstream region binds to \(^8\)-FN of the gelatin-binding domain (23, 24), and the adjoining first FNBR binds to \(^2\)-FN of the fibrin-binding domain (Fig. 1C) (24, 27). FUD blocks FN assembly (29) and exposes the mAbIII-10 epitope in 10FNIII (30) and thus may mimic cell surface molecules of vertebrate cells that interact with N-terminal modules and expose \(^10\)FN III. It is not known whether the effects of FUD are due to its interaction with \(^2\)-FN, \(^8\)-FN, or both \(^2\)-FN and \(^8\)-FN. To learn the importance of the fibrin and gelatin binding, we compared FUD to “high affinity downstream domain” (HADD), a 49-residue polypeptide similar to SfbI-5 with its adjacent nonrepetitive sequence, *i.e.* HADD was designed to bind tightly to N-\(^3\)-FN by anti-parallel \(\beta\)-strand addition (Fig. 1C).

MATERIALS AND METHODS

Proteins—Plasma FN was purified from a fibrinogen-rich plasma fraction (31). Expression and purification of polyhistidine-tagged monomeric N-\(^3\)-FN, N-\(^5\)-FN, and N-\(^8\)-FNIII and dimeric \(^6\)-FN-C (starts at residue 291 (numbering from the initiating methionine) and contains V89 version of the variable region without extra domains A or B) were accomplished using recombinant baculovirus and affinity chromatography as described previously (24, 32). The recombinant proteins were secreted into medium at concentrations of 5–20 \(\mu\text{g/mL}\), and after purification were pure and migrated as expected when analyzed by SDS-PAGE without and with prior reduction. The molarity of FN and constructs stated throughout the paper were calculated based on the monomer or subunit, assumed to have an average molecular mass of 250 kDa for the subunits of heterodimeric plasma FN. Rat tail type I collagen (Upstate) and gelatin (Sigma) were purchased.

FUD was expressed and purified as described previously (24). For HADD, PCR-based strategies were used to add DNA encoding the 16 C-terminal residues of SfbI-5 to DNA encoding 33 residues of FNBR SfbI-4 (Fig. 1C). HADD was expressed, purified, and characterized as described previously for FUD (24) except for determination of concentration. Because HADD lacks tyrosine or tryptophan, its concentration could not be estimated by absorbance at 280 nm and instead was determined using the bicinchoninic acid (BCA) assay (Pierce) with FUD as the standard (24). Amino acid analysis of HADD performed at University of California-Davis Proteomics Core facility yielded a value that was within 10% of that estimated by the BCA.

Mouse anti-human monoclonal antibody mAbIII-10 against \(^10\)FN III was described previously (6), as were mouse anti-human mAbs 4D1, 7D5, 5C3, and 9D2 (24, 30).

Labeling—FN was labeled with fluorescein isothiocyanate FN (FITC-FN) as described previously (33) or with Alexa Fluor 488 (Invitrogen) (A488-FN) as per the manufacturer’s instructions. Biotinylation of FUD or HADD with \(N\)-hydroxysulfosuccinimide-biotin (Pierce) was done as described previously (24); biotinylated probes are designated by the prefix “b-.”

Assembly Assays—Fluorescence microscopic assays of FN assembly were done as described previously (24) except mouse FN\(^{-/-}\) cells were allowed to adhere for 2 h and assembled exogenous FITC-FN for 1 h. For quantitative dose-response assays, human foreskin fibroblasts (strain AH1F) were seeded in microplate wells and incubated with 20 nm A488-FN in the presence or absence of 4 nm to 2.5 \(\mu\text{g/mL}\) FUD or HADD. After 18 h at 37 °C, monolayers were washed, and fluorescence of assembled A488-FN was quantified (excitation, 485 nm; emission, 535 nm) using the Tecan GeniosPro.

Enzyme-linked Immunosorbent Assays—Wells of high binding plates (Costar 3950) were coated overnight with 40 nm FN, N-\(^3\)-FN, \(^6\)-FN-C, or N-\(^8\)-FN in 10 mM Tris, 150 mM NaCl, pH 7.4 (TBS), as indicated in the figure legends. Binding of b-HADD or b-FUD to adsorbed proteins was quantified with the alkaline phosphatase-streptavidin complex as described previously for b-FUD (24). Where indicated, binding of b-HADD was compared with separate wells precoated with 1 \(\mu\text{g/mL}\) b-FUD as a positive control. The experimental setup was altered as described previously to compare the effect of soluble mAb, 1 mM zinc sulfate, FN, or FN constructs on binding of b-HADD or b-FUD to adsorbed FN (24). In some experiments, unfractionated heparin, 6–30 kDa in size (grade 1A from porcine intestine (Sigma)), was incubated with FN or N-\(^8\)-FN prior to the addition of b-HADD or b-FUD. In other experiments, purified mAbIII-10 was preincubated with soluble FN or N-\(^8\)-FN for 1 h prior to the addition of b-HADD or b-FUD. Binding was done in 10 mM Tris, 50–300 mM NaCl, pH 7.4, as indicated in the figure legends. Wells were washed twice with the solution used in the binding step and twice more in 10 mM Tris, 150 mM NaCl, 0.05% Tween, prior to addition of enzyme-linked streptavidin (24). mAb binding to adsorbed FN in the presence of HADD or FUD was quantified as described previously (24).

In experiments looking at the binding of FN to collagen or gelatin, 4 nm FN and 100 nm polypeptide diluted in TBS con-
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Competitive Binding Assays with mAbIII-10—High-binding plates were coated overnight with 50 μl FN, washed with 10 mM Tris, 50 mM NaCl, 0.05% Tween, pH 7.4, and blocked with 1% BSA in washing buffer for 1 h. In some experiments, HADD or FUD was added to FN in 10 mM Tris, 50 mM NaCl, pH 7.4, plus 0.2% BSA for 30 min. To 150 μl of the FN-containing solution, 25 μl of mAbIII-10 ascites (previously diluted to yield a final dilution of 1:50,000) was added in 10 mM Tris, 50 mM NaCl, pH 7.4, plus 0.2% BSA and 0.35% Tween 20 (final concentration 0.05%), and the mixture was incubated for 1 h prior to transfer to FN-coated wells and incubation for 2 h. Plates were washed two times with 10 mM Tris, 50 mM NaCl, 0.05% Tween, pH 7.4, and two times with TBS containing 0.05% Tween. Alkaline phosphatase-conjugated donkey anti-mouse IgG (Jackson ImmunoResearch) at 1:5000 was added to wash buffer and incubated for 1 h. Plates were washed four additional times before the addition of 1 mg/ml 4-nitrophenyl phosphate disodium salt hexahydrate (Sigma) in 10 mM Tris, 150 mM NaCl, pH 9.0. A_{405 nm} was determined using a Tecxan Genios Pro.

In experiments comparing purified FN to FN in plasma, exposure of the mAbIII-10 epitope was monitored by a competitive ELISA similar to that described above. Purified FN was diluted in TBS to the concentration of FN determined to be in the plasma sample by competitive ELISA using mAb 9D2, which is not sensitive to conformation as described previously (30). These samples were then diluted further in 10 mM Tris, 50 mM NaCl, pH 7.4, to 233 mM FN without or with FUD or HADD in 2.5-fold molar excess. After 30 min, the mixtures were diluted further, and mAbIII-10 ascites (final concentration after addition of 1:50,000) was added in a volume that brought the concentration of the FN to 200, 100, 50, 25, or 12.5 nM. After 30 min, these mixtures were added to FN-coated wells, which were incubated for 1.5 h. Washes and incubations to determine binding of mAbIII-10 to adsorbed FN were described above except that the detecting reagent was peroxidase-conjugated goat anti-mouse IgG (Jackson ImmunoResearch), and bound conjugate was quantified by addition of SureBlue TMB microwell peroxidase substrate as above.

Dynamic Light Scattering and Isothermal Titration Calorimetry (ITC)—Measurements were performed using Beckman Coulter (Miami, FL) N5 Submicron Particle Size Analyzer instrument in 20 mM Tris, 100 mM NaCl, pH 7.4, at 25 °C. The buffer was degassed and filtered with 0.22-μm Millex-GP filter (Millipore, Cork, Ireland). FN at a concentration of 1 mg/ml (4 μM FN subunit) was titrated with solutions of 200 μM FUD or 165 μM HADD. After each addition, the mixture was allowed to equilibrate for 10 min, and then scattered intensity of light from the He-Ne laser was measured at 90° for 200 s; each measurement was repeated six times. Data were processed by Photon Correlation Spectroscopy software. The mean size of FN or FN-polypeptide complex was computed assuming a refractive index of 1.33 and viscosity of 0.89 poise. Size distribution processor analysis was performed for each sample, and the mean size of the particle was determined according to the volume (weight) distribution format of size distribution processor analysis. Large particles accounted for <5% of total particles.

ITC was performed as described previously (24) with a VP-ITC microcalorimeter (MicroCal, LLC) at 25 °C. The cell contained 1.4 ml of a solution of FN or N-9FNI, and the syringe contained 600 μl of HADD. The titration was performed in 37 injections (1 of 1 μl, four of 4 μl, and 32 of 8 μl) delivered at 120-s intervals. Data from the initial injection were discarded. HADD, N-9FNI, and FN were dialyzed against PBS, pH 7.4, and adjusted to the concentrations given in Table 1. Data were fit by Lavenberg-Marquardt nonlinear regression with Origin 7.0 using the one-site model.

RESULTS

HADD Binds to N-5FNI—Binding of FUD to 2–5FNI and 8–9FNI of soluble FN blocks FN assembly and induces a conformational change that exposes the epitope to mAbIII-10, which is otherwise partially cryptic at low ionic strength (24, 29, 30). We have speculated that binding of FUD blocks assembly and exposes the mAbIII-10 epitope by breaking the electrostatic interaction between 4FNI and 3FNIII or coupling of binding to 8–9FNI with conformational changes in adjacent 1–3FNIII and linkage of changes in 3FNIII to changes in 4FNIII (24). Learning the contribution of 2–5FNI requires a polypeptide that binds by β-strand addition to 2–5FNI with an affinity that is comparable with the affinity of binding of FUD to N-9FNI. Polypeptides based on individual FNBRs of F1 adhesin (e.g. SfbI-2 or -4) bind to N-9FNI with >10-fold looser affinity than FUD (34). We therefore designed, expressed, and purified HADD, which contains SfbI-4 and the downstream region of SfbI-5 (Fig. 1B).

On studies of SfbI-5 and its adjacent downstream region or a homologous construct from the Streptococcus dysgalactiae adhesin (28, 35), we anticipated that HADD would bind to N-9FNI with an affinity equivalent to FUD because the favorable energy of binding to 1FNI substitutes for loss of the favorable energy of binding to 8–9FNI (Fig. 1C). Indeed, K_{D} values for binding of HADD to N-9FNI and intact FN were 2.4 and 12.6 μM, respectively, as measured by ITC in 150 mM sodium chloride at 25 °C (Table 1); these affinities are comparable with ITC measurements of binding of FUD to N-9FNI and intact FN (24). Furthermore, the interactions were driven by ΔH values favorable enough to overcome unfavorable ΔS values (Table 1), as in the case of binding to SfbI-5 by β-strand addition to N-9FNI (35).

The specificity of HADD for N-9FNI was assessed by five additional assays. When we examined the ability of b-HADD to bind adsorbed FN, N-9FNI, or 6FNI-C, there was similar binding of b-HADD to FN and N-9FNI and no binding to 6FNI-C (Fig. 2A). As with b-FUD binding (24), mAb 4D1 to 2FNI slightly decreased b-HADD binding to coated FN, whereas 7D5 to 4FNI decreased binding of both polypeptides considerably (Fig. 2B). In contrast, 5C3 to 9FNI decreased b-FUD binding to...
adsorbed FN but increased binding of b-HADD (Fig. 2B). In the reciprocal experiment, FUD or HADD increased binding of 4D1 and decreased binding of 7D5 to coated FN, whereas FUD decreased, and HADD had no effect on binding of 5C3 to coated FN (Fig. 2C). Thus, the cross-competition studies indicate that HADD binds to 2FNI and 4FNI but not 9FNI. Zn²⁺/H₁₁₀₀₁ binds to and grossly alters the global fold of 8FNI (36) and decreases the ability of FUD to bind to adsorbed FN (24). Consistent with the expected binding specificity of HADD versus FUD, 1 mM Zn²⁺/H₁₁₀₀₁ had no effect on binding of b-HADD to adsorbed FN under conditions in which binding of b-FUD was decreased (Fig. 2D). Collagenous sequences bind to an extended site on the gelatin-binding domain that includes 8–9FNI (37). Consistent with this, preincubation of FN with soluble FUD, but not HADD, blocked binding of FN to adsorbed gelatin or collagen (Fig. 2E).

**HADD, Like FUD, Blocks FN Assembly**—Deletion of N⁻⁵FNI from FN prevents the formation of FN fibrils (38), and exogenous N⁻⁵FNI (15), FUD (29), or monoclonal antibody to ⁴FNI or ⁹FNI (24) blocks the assembly of FN. To test whether ligation of ¹⁻⁴FNI by b-strand addition is sufficient to block FN assembly, we asked if HADD blocks FN assembly. FN⁻¹⁻⁴ cells adherent to adsorbed laminin were incubated with 20 nM FITC-FN for 1 h with or without 50 or 500 nM HADD. There was a decrease in FN fibrils when the concentration of HADD was 50 nM and a loss of FN fibrils at 500 nM (Fig. 3A). Dose-response curves of inhibition were performed on deposition of soluble 20 nM A₄₈₈-FN over 18 h by monolayers of human foreskin fibroblasts using fluorescence as a read-out (Fig. 3B). HADD and FUD were similarly active and had near maximum effects at 50 nM.

**Binding of HADD, Like Binding of FUD, Causes Exposure of the mAbIII-10 Epitope and Expansion of Soluble FN**—To determine whether HADD, like FUD, causes conformational change in soluble FN, we monitored exposure of the mAbIII-10 epitope using a competitive ELISA in which binding of mAbIII-10 to substrate-bound FN is inhibited by increasing concentrations of soluble purified FN with or without a 2.5-fold molar excess of the polypeptide. The assay was done in low salt (50 mM NaCl) in which the mAbIII-10 epitope is hidden (6). HADD or FUD caused FN to compete many-fold better than FN alone (Fig. 3A). The competition curves for FN alone were complex, steeper at lower concentrations than at higher concentrations, although not as steep as when HADD or FUD was present. Purified FN has been demonstrated previously to contain multimers that interact preferentially with mAbIII-10; such multimers could account for complexity of the inhibition curve (6). To ascertain if the complexity of the inhibition curves is a result...
of the purification procedure, inhibition was performed with plasma with known FN concentration. The same complexity was found for unfractionated plasma as for purified FN, and enhancement of competition by addition of HADD or FUD to plasma was similar to enhancement of competition in purified FN (Fig. 4A). Thus, FN exists as a variety of conformers in plasma before purification, and some of the conformers compete for mAbIII-10 binding better than others and all or almost all conformers are susceptible to further conformational change upon complex formation with HADD or FUD.

When the concentration of soluble FN was 20 nM and exposure of the mAbIII-10 epitope was measured as function of increasing concentrations of HADD, a similarly complex curve was obtained with a near maximal effect when the ratio of HADD/FN subunit was 1:1 (Fig. 4B). To relate epitope exposure to another measure of conformational change, hydrodynamic radius was determined by DLS as 4 μm FN was titrated with HADD or FUD (Fig. 4C). The radius increased linearly as a function of polypeptide concentration, changing from ∼9 to ∼11.5 nm and reaching a maximum at polypeptide/FN subunit ratios of 1:1. These results indicate that HADD or FUD each forms a tight complex with FN in which FN assumes an expanded conformation in which the mAbIII-10 epitope is exposed.

Soluble N-9FNIII and N-2FNIII Compete Better Than Soluble FN for Binding of HADD to Substrate-bound FN—Soluble N-9FNIII competes better than soluble FN for binding of b-FUD to coated FN (24). We hypothesized that this difference is due to intramolecular interactions within FN occluding the FUD-binding site. To determine whether the electrostatic interaction between N-FNI and FNIII (7) is responsible for the difference in ability of FN and N-FNI to compete with b-FUD or b-HADD, we tested competition by soluble N-9FNIII, in which the interaction has been demonstrated (7), for binding of b-FUD or b-HADD to adsorbed FN. Because the FNI-FNIII interaction is sensitive to ionic strength (7), binding assays were done in buffer containing 50 mM NaCl. Increasing concentrations of N-9FNIII competed for binding of b-FUD or b-HADD to adsorbed FN to nearly the same extent as N-9FNII, and both competed ∼10-fold better than soluble FN (Fig. 5, A and B).

mAbIII-10 or Heparin Increases the Ability of Soluble FN to Bind to b-FUD or b-HADD—The observations that HADD or FUD causes exposure of the mAbIII-10 epitope and N-9FNIII and N-2FNIII both compete better than soluble FN for binding of FUD or HADD suggest that the FNI-FNIII interaction is part of a larger network that limits β-strand addition to 1-5FNII. We therefore compared FN and FN-mAbIII-10 1:1 complex as competitors for binding of b-FUD or b-HADD to adsorbed FN. Complex formation with mAbIII-10 increased the ability of FN to compete for binding of b-FUD (Fig. 6A) or b-HADD (Fig. 6B) to N-9FNII. The effect was small, 2-fold for FUD and ∼1.5-fold for HADD, but reproducible.

Heparin binds to 12–14FNIII (1, 39) and is known to expose the mAbIII-10 epitope (6). To investigate the possibility that heparin binding causes a coupled conformational change encompassing both FNIII and the N-terminal FNI modules, we looked at the effect of heparin on the ability of soluble FN to compete with b-FUD or b-HADD for binding to adsorbed FN (Fig. 6, C and D). Using the type of competitive ELISA shown in Fig. 4, heparin, 0.25 mg/ml, ∼10 μM, caused increased exposure of the mAbIII-10 epitope as in published experiments (results not shown) (6). Although it had no effect on competition by soluble N-FNI for binding of b-FUD (Fig. 6C) or b-HADD (Fig. 6D), heparin binding causes a coupled conformational change.
to substrate-bound FN, this concentration of heparin caused ~2-fold enhancement of the ability of soluble FN to compete for binding of b-FUD or b-HADD. The differences between the effects of heparin on the competition by soluble N-9FNII and FN for FUD or HADD binding indicate that the small but reproducible effect of heparin on the mAbIII-10 exposure assay is due to distant binding of heparin to 12–14FNIII rather than direct binding by heparin to the N-9FN region.

**DISCUSSION**

FN assembly by adherent vertebrate cells and F1 adhesin/SfbI-mediated entrance of *S. pyogenes* into host cells utilize common features of FN, i.e. interactions involving the N-terminal N-9FN region of FN and binding of integrins to FNIII modules, usually α5β1 to 10FNIII (13, 40). To investigate how ligation of the N terminus of FN leads to exposure of 10FNIII for processes including assembly and internalization, we utilized a competitive binding assay to monitor accessibility of the mAbIII-10 epitope in 10FNIII that is cryptic in soluble FN at low ionic strength (6). Previous work showed that the mAbIII-10 epitope becomes available upon incubation of soluble FN with FUD, which binds to 8–9FN and 2–5FN by β-strand addition (24) or denatured collagen (gelatin) (6, 30). In addition, expansion of plasma FN is seen upon binding of cyanogen bromide fragment 7 (CB7) of the α1(I) chain of type I collagen (41). CB7 contains a sequence that binds by β-strand addition to 3FNIII (37). To determine whether ligation of 8–9FN is necessary for exposure of the mAbIII-10 epitope, we designed a polypeptide, HADD, that mimics SfbI-5 in binding to 1–5FN (28, 35). HADD exposed the mAbIII-10 epitope and caused expansion of FN as assessed by DLS, indicating that ligation of 8–9FN is not necessary for FN expansion. Exposure of mAbIII-10 epitope by HADD demonstrates that ligation of the fibrin-binding region alone is sufficient to disrupt intramolecular interactions and cause long range conformational changes that result in the exposure of 10FNIII.

Plasma FN is a heterodimer of subunits that differ in whether the variable region is present (1). The conformations assumed by the 58 modules of plasma FN are presumably controlled by “head-to-tail” interactions between consecutive modules and longer range interactions among nonadjacent modules. Candidate long range interactions have been identified between 4FN and 3FNIII of the same subunit (7), between 2–3FNIII and 12–14FNIII of different subunits (8), and a less characterized interaction between N-5FN and 12–14FNIII (9, 10). Soluble FN as compared with soluble N-9FN has decreased ability to compete for binding of HADD to adsorbed FN. Experiments showing that N-9FN and N-3FNIII compete equally well for HADD suggest that disruption or loss of the interface between 4FN and 3FNIII is not sufficient to explain why soluble FN competes better for mAbIII-10 in the presence of FUD or HADD and indicates the involvement of FN modules C-terminal to 3FNIII.
This finding is compatible with published ITC experiments demonstrating little difference in $K_D$ values or enthalpies for binding of a FUD-like polypeptide to N-9FNI compared with N-3FNIII (34). Fig. 7 describes a “beads on a string” (5) model for compact FN drawn to emphasize the interactions described above. The intra-subunit 4FNI-3FNIII and inter-subunit 2–3FNIII-12–14FNIII interactions constitute a pair of junctions at the core of the molecule. We speculate that disruption of weak 4FNI-3FNIII interactions in both subunits by binding of FUD or HADD may be coupled to more energetically profound disruptions of inter-subunit interactions between 2–3FNIII and 12–14FNIII. These disruptions result in extension of the two subunits, shown at its most extreme in Fig. 7, and exposure of 10FNIII.

FIGURE 4. HADD is similar to FUD in exposing the mAbIII-10 epitope of purified FN or FN in plasma and expanding purified FN. A, effect of HADD or FUD on the exposure of the mAbIII-10 epitope in purified FN and FN in plasma as determined by competitive ELISA. Purified FNs (solid lines) or FNs in diluted plasma (dotted lines), 233 nM, were incubated without (●) or with 580 nM FUD (▲) or HADD (□) for 30 min. Prior to the assay, the concentration of FN in neat plasma was found to be 0.65 mg/ml (2600 nM) by competition ELISA with a mAb that is not conformation-sensitive. The six samples were then diluted to the indicated FN concentrations; mAbIII-10 was added, and competition by soluble FN for mAbIII-10 binding to coated FN was determined. Data are expressed as percent of mAbIII-10 binding alone (no FN or polypeptide added) and are representative of two experiments. B, ELISA of competition of binding of mAbIII-10 to coated FN by 20 nM soluble FN alone or preincubated with the indicated concentrations of HADD (□) or with 20 nM FUD (▲). Values are expressed relative to 20 nM soluble FN with mAbIII-10 but no polypeptide and represent mean ± S.D. of three experiments. The inset replots the HADD titration in comparison with the maximum inhibition found with 40 nM HADD. C, HADD (□) or FUD (▲) were titrated into FN solution separately, and the hydrodynamic radius of 4 µM FN or 4 µM FN plus the indicated concentration of polypeptide was calculated. Measurements were performed at 25 °C in 20 mM Tris, 100 mM sodium chloride, pH 7.4. Error bars indicate the standard deviation of six measurements on each sample. The experiment was repeated twice with the same result.
Our experiments and the literature yield additional, albeit limited, insight about the impact of HADD or FUD binding on conformation of a ligated FN subunit versus the overall dimer. Expansion of FN as assessed by DLS was a linear function of the HADD or FUD concentration that reached completion at a polypeptide/FN subunit ratio of 1:1, indicating that ligation of an individual FN subunit results in expansion of that subunit independent of the second subunit. In contrast, the dose response for exposure of the mAbIII-10 epitope by FUD (30) or HADD (Fig. 4B) was curvilinear and ~75% complete at a polypeptide/FN dimer ratio of 1:1 (polypeptide/FN subunit ratio of 0.5:1). This finding suggests that ligation of a single subunit of plasma FN is sufficient to allow binding of mAbIII-10. Published electron microscopic images of FN-mAbIII-10 complexes revealed FN in a V-shaped configuration with mAbIII-10 in the angle of the “V” with Fab arms binding to each subunit of
FN (6). Univalent binding of one Fab arm of mAbIII-10 to one FN subunit therefore may initiate cooperative events to form a complex in which mAbIII-10 and FN are bivalent in relation to each other. Such cooperativity is consistent with the stronger binding of preformed mAbIII-10-FN complexes to HADD or FUD.

HADD, like FUD, blocks FN assembly, suggesting that the fibrin-binding domain of FN mediates the first interaction with the cell surface during FN assembly (15–17, 29). Despite this, there appear to be instances in which ligation of 10FNIII dominates over exposure of the FN N terminus (13). Using competitive ELISAs, we show that binding of mAbIII-10 to FN modestly but significantly increased the ability of soluble FN to interact with FUD or HADD. Heparin, which like FUD or HADD increases exposure of the mAbIII-10 epitope (6), also increased the ability of soluble FN to interact with FUD or HADD. As described above, the modules that bind heparin with high affinity, 12–14FNIII (39), have been deduced to interact with 2–3FNIII in the opposite subunit (8) and with N-5FNII (9, 10). Thus, binding of heparan sulfate to 12–14FNIII may control accessibility of both the RGD cell adhesive sequence in 10FNIII and the N-terminal FNIII modules during assembly. The present findings therefore support the existence of an allosteric network among the fibrin-binding domain 1–5FNII, 10FNIII, and 12–14FNIII in dimeric plasma FN. Such an allosteric network would allow for controlled exposure to binding sites in FN during FN assembly and bacterial host cell invasion.

The allosteric network among the fibrin-binding domain, 10FNIII, and 12–14FNIII is likely different in cellular FN containing differentially spliced EDA and extra domain B FNIII modules (1). For instance, in the 7–14FNIII constructs lacking 2–3FNIII, the presence of EDA between 11FNIII and 12FNIII resulted in dimers because of interaction of EDA with 12–14FNIII, suggesting that the presence of EDA in intact FN may favor an open conformation by competing with 2–3FNIII for binding to 12–14FNIII (8). Such an open conformation may account for the unique ability of EDA-positive FN to be assembled into extensive fibrous networks by CHO cells (42). Interestingly, FN assembly associated with lymphatic valves in mice has been shown to require interaction of α9β1 integrin with the α9β1-recognition sequence in EDA rather than interaction of α5β1 integrin with 10FNIII (43).

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