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The Integrin Equilibrium: Balancing Protein-Protein and Protein-Lipid interactions

Abstract

On circulating platelets, the integrin fibrinogen receptor &alphaIIb&beta3 favors inactive conformations. Platelets rapidly activate &alphaIIb&beta3 to bind fibrinogen, mediating a platelet clot. Resting &alphaIIb&beta3 is stabilized by interactions between the &alphaIIb and &beta3 transmembrane domains. Binding of talin-1 and kindlin-3 to the integrin cytoplasmic domain stabilizes separation of the TMs and receptor activation. Src family kinases are needed for transmission of extracellular signals into the cell. We have sought to better understand how signals are transmitted across the &alphaIIb&beta3 TM domain. First we characterized the structure and dynamics of the active and inactive integrin cytoplasmic domain to determine how motifs that bind talin-1 and kindlin-3 are affected by the integrin activation state and the membrane environment. The &alphaIIb&beta3 cytoplasmic domain is disorded, while the &beta3 subunit contains two &alpha-helices, which interact with the phospholipid bilayer. The close proximity of &alphaIIb to &beta3 in the inactive state induces a kink that projects the &beta-chain parallel to the membrane surface. This kink is likely to stabilize interactions between helices in &beta3 with the membrane and possibly compete with binding to talin-1 and kindlin-3. The &beta3 cytoplasmic domain becomes increasingly dynamic further from the membrane, suggesting that distal kindlin-3 binding residues should be more accessible to interact with their binding partners. We also studied the interaction between talin-1 and the &beta3 cytoplasmic domain to determine how the membrane environment affects formation of the talin-1/&beta3 complex. The membrane significantly increases the affinity of talin-1 for &beta3, but that the ternary integrin/talin-1/membrane complex is not significantly more stable than the talin-1/membrane complex alone. We also performed preliminary experiments characterizing the interaction between &alphaIIb&beta3 and both kindlin-3 and c-Src. Unlike talin-1, kindlin-3 does not require the membrane to bind the integrin cytoplasmic domain with high affinity. The interaction between c-Src and the integrin was extremely weak suggesting that it might require colocalization through other interactions. Finally, we developed a simplified model showing how thermodynamic coupling between integrin subunits might allow &alphaIIb&beta3 to adopt multiple activation states.

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THE INTEGRIN EQUILIBRIUM: BALANCING PROTEIN-PROTEIN AND

PROTEIN-LIPID INTERACTIONS

David T. Moore

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Dedicated to my family.

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ABSTRACT

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David T. Moore

William F. DeGrado

On circulating platelets, the integrin fibrinogen receptor α Ilb β 3 favors inactive conformations. Platelets rapidly activate α Ilb β 3 to bind fibrinogen, mediating a platelet clot. Resting α Ilb β 3 is stabilized by interactions between the α Ilb and β 3 transmembrane domains. Binding of talin-1 and kindlin-3 to the integrin cytoplasmic domain stabilizes separation of the TMs and receptor activation. Src family kinases are needed for transmission of extracellular signals into the cell. We have sought to better understand how signals are transmitted across the α Ilb β 3 TM domain. First we characterized the structure and dynamics of the active and inactive integrin cytoplasmic domain to determine how motifs that bind talin-1 and kindlin-3 are affected by the integrin activation state and the membrane environment. The α Ilb cytoplasmic domain is disorded, while the β 3 subunit contains two α -helices, which interact with the

phospholipid bilayer. The close proximity of α IIb to β 3 in the inactive state induces a kink that projects the β -chain parallel to the membrane surface. This kink is likely to stabilize interactions between helices in β 3 with the membrane and possibly compete with binding to talin-1 and kindlin-3. The β3 cytoplasmic domain becomes increasingly dynamic further from the membrane, suggesting that distal kindlin-3 binding residues should be more accessible to interact with their binding partners. We also studied the interaction between talin-1 and the β 3 cytoplasmic domain to determine how the membrane environment affects formation of the talin- $1/\beta$ 3 complex. The membrane significantly increases the affinity of talin-1 for β 3, but that the ternary integrin/talin-1/membrane complex is not significantly more stable than the talin-1/membrane complex alone. We also performed preliminary experiments characterizing the interaction between α IIb β 3 and both kindlin-3 and c-Src. Unlike talin-1, kindlin-3 does not require the membrane to bind the integrin cytoplasmic domain with high affinity. The interaction between c-Src and the integrin was extremely weak suggesting that it might require colocalization through other interactions. Finally, we developed a simplified model showing how thermodynamic coupling between integrin subunits might allow α IIb β 3 to adopt multiple activation states.

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Chapter 1 Background

I. Introduction

In multicellular organisms, integral membrane proteins mediate adhesion and signal transduction across the plasma membrane. Such processes are often dynamically regulated and reversible, suggesting that they can be described as linked equilibria. Changes in conformation, oligomeric state. protein-protein in extracellular, or contacts transmembrane (TM), and cytoplasmic domains can shift the equilibrium between different receptor functional states and transmit information across the membrane. In order to develop effective therapies to target signaling receptors, these equilibria must be understood to prevent undesired side effects. The importance of understanding these processes is illustrated by the platelet fibrinogen receptor integrin, α IIb β 3. Small molecules that mimic fibrinogen successfully block ligand binding to α IIb β 3, but stabilize active receptor conformations, resulting in either a paradoxical prothrombotic state or clearance of platelets and severe bleeding. In the following chapters, the regulation of α IIb β 3 is discussed. In particular, we focus on the mechanisms of transmembrane signal transduction and how coupled protein-protein and protein-lipid interactions allow multidomain receptors such as α IIb β 3 to integrate multiple signals to produce surprisingly diverse functions.

II. Platelet integrins.

Platelets are highly specialized cell fragments that are released from bone marrow megakaryocytes and are necessary for normal arterial clotting (4). Functional thrombocytopenia in the presence of normal platelet counts results from loss of the platelet fibrinogen receptor integrin α IIb β 3. The integrin α IIb β 3 mediates formation of the platelet clot and patients whose platelets lack normal allbß3 function present with the bleeding diathesis Glanzmann thrombasthenia (5, 6). Severe bleeding can also result in patients with normal platelet counts and functional integrins via loss of signal transduction molecules that regulate the α IIb β 3 decreased number of circulating (7-10). Finally, а platelets (thrombocytopenia) can result from spurious activation of the receptor by small molecules or autoimmune antibodies (11).

Resting platelets circulate with α Ilb β 3 in an inactive conformation. In the presence of damaged vasculature, numerous signals are integrated by the platelet, culminating in the activation of α Ilb β 3, which binds fibrinogen and von Willebrand's factor to form a platelet plug (12). The integrin family of receptors is ubiquitously expressed in metazoa (13). Functional receptors are heterodimeric, composed of α and β subunits (Figure 1.1). Each subunit has a large ectodomain, a single transmembrane domain, and a relatively short cytoplasmic domain. In

humans, there are eighteen α -chains and eight β -chains that combine to form 24 integrin heterodimers. Although these receptors are differentially expressed, recognize unique ligands, and occupy distinct functional niches, they contain common structural elements. In particular the transmembrane and cytoplasmic domains are highly conserved, consistent with their importance in regulating integrin function through protein-protein interactions (14).



Figure 1.1 Simplified integrin equilibrium.

Integrins are in equilibrium between inactive (left) and active (right) states. This equilibrium is modulated by both intracellular and extracellular events. Integrins possess highly modular domain structures that communicate between each other establishing an elaborate coupled equilibrium. Interactions between the intregrin TM domains stabilize the inactive conformation. Soluble factors can bind to each modular subdomain and shift the equilibrium between states.

On a resting platelet, the transmembrane and membrane-proximal domains of inactive α IIb β 3 form an extended heterodimeric interface (15-17). Activation of α IIb β 3 requires TM domain separation. This separation is dramatic (100 Å) and propagates to the ectodomain, stabilizing significant conformational changes that allow ligand binding (18). The term "inside-out signaling" refers to the classical mode of integrin activation in which intracellular signaling molecules bind to the integrin cytoplasmic domain and stabilize separation of the α and β TM domains, which results in subsequent activation of the extracellular ligand binding site. However, α IIb β 3 can also be activated through extracellular interactions. This suggests that the activation state of α IIb β 3 can be described as a linked equilibrium in which intracellular, TM, and cytoplasmic conformational states are coupled and either stabilized or destabilized by interactions with other molecules.

Inside-out activation requires the evolutionarily related talin and kindlin protein families (19, 20). There are two talin paralogs (talin-1 and 2) and three kindlin paralogs (kindlin-1, 2, and 3). In platelets, α IIb β 3 requires talin-1 and kindlin-3 only (21) (22). Patients who lack kindlin-3 function have Leukocyte Adhesion Deficiency III (LAD-III) in which platelets are unable to activate α IIb β 3 and immune cells are unable to activate β 2 integrins resulting in both severe bleeding and impaired

immune function (7-10). Both talins and kindlins contain phospholipid binding domains, suggesting that their interactions with the inner leaflet may play a role in the regulation of α IIb β 3.

Integrin "outside-in signaling" can refer to one of two processes. The first is stabilization of an active integrin conformation by extracellular ligands. The second refers to signaling events downstream of integrin activation and ligand binding. Generally, these events require integrin oligomerization and involve recruitment of a number of signaling molecules including Src-family kinases (SFKs) (23). However, the dynamics of the signaling complexes remains an area of active research.

Talin and kindlin are members of the membrane-binding FERM (band 4.1/Ezrin/Radixin/Moesin) domain family. Their interactions with the αIlbβ3 cytoplasmic domains are likely to occur simultaneously with binding to the plasma membrane. Therefore, any equilibrium description of integrin protein-protein interactions must include interactions with the inner membrane leaflet. For simplicity, interactions between soluble proteins and the inner leaflet can be divided into two categories. First, proteins can bind using a combination of nonspecific electrostatic interactions with lipid head groups and hydrophobic interactions through insertion of hydrophobic side chains (24, 25). Second, there are a host of domains that differentiate between lipid head groups with a spectrum of specificity (26). However, "non-specific" electrostatics are likely to play a role in even "specific" head group recognition as lipid binding domains are often

positively charged and the inner leaflet of the plasma membrane is greater than 20% negatively-charged phosphatidylserine (PS).

The majority of negatively-charged lipids reside on the inner leaflet of the plasma membrane bilayer. PS is the most prevalent monovalent anionic lipid in mammalian cells and platelets, representing greater than 20% of the inner leaflet (26). The distribution of PS is asymmetric and it is actively transported from outer leaflet to the cytosolic face of the plasma membrane. During apoptosis or platelet activation, PS redistributes such that roughly half is on both the outer and inner leaflets. Exposed PS serves as an inflammatory nidus and a site for the assembly of clotting factor complexes.

The laboratories of Honig, McLaughlin, and Murray have developed models for negatively charged bilayers containing monovalent lipids (27, 28). They found that the molar partition coefficient varies exponentially with the percentage of PS (28, 29), *i.e.* the Δ G of association is linear with percentage charged lipid concentration. These experiments were extended to the interaction of the myristoylated anchor of c-Src a kinase involved in outside-integrin signaling. While the myristoyl tail produced significant thermodynamic driving force for membrane association, electrostatics increased the affinity significantly (29). Phosphoinositides are negatively-charged lipids that exist in much lower concentrations than phosphatidylserine, but are more dynamically regulated to produce local concentrations of specific lipids to regulate signal transduction (26). As

they contain a number of phosphates, they are negatively-charged and contribute a significant amount of electrostatic driving force to protein-lipid interactions (24). Their ability to support highly specific protein-lipid interactions is dependent on the phosphate geometry at different locations on the inositol ring and can be differentiated with surprising precision (26, 30).

Several lines of evidence suggest that PtdIns(4,5) P_2 , plays a significant role in integrin regulation and the turnover of integrin complexes (31) (32). Likewise, it has been suggested that an interaction between talin-1 and PtdIns(4,5) P_2 directly competes with autoinhibitory interactions between the integrin binding FERM domain and its C-terminal rod domain (33). The talin FERM domain binds directly to the C-terminus of phosphatidylinositol-4-phosphate-5-kinase- γ (PIP5KI γ), a lipid kinase that generates PtdIns(4,5) P_2 from PtdIns(4)P, suggesting that talin may recruit PtdIns(4,5) P_2 to integrin signaling complexes. However, PIP5KI γ knockout cell lines continue to support inside-out integrin activation, but exhibit loss of integrity under mechanical stress (34), suggesting that PtdIns(4,5) P_2 may recruit secondary signaling molecules or simply reinforce talin-lipid interactions under mechanical shear.

Chapter 2 NMR analysis of the α IIb β 3 cytoplasmic interaction suggests a mechanism for integrin regulation.

Abstract

The integrin α IIb β 3 is a transmembrane (TM) heterodimeric adhesion receptor that exists in equilibrium between resting and active ligand binding conformations. In resting α IIb β 3, the TM and cytoplasmic domains of α IIb and β 3 form a heterodimer that constrains α IIb β 3 in its resting conformation. To study the structure and dynamics of the cytoplasmic domain heterodimer, we prepared a disulfide-stabilized complex consisting of portions of the TM domains and the full cytoplasmic domains. NMR and hydrogen-deuterium exchange of this complex in micelles showed that the α IIb cytoplasmic domain is largely disordered, but it interacts with and influences the conformation of the β 3 cytoplasmic domain. The β 3 cytoplasmic domain consists of a stable proximal helix contiguous with the TM helix and two distal amphiphilic helices. То confirm the NMR structure in a membrane-like environment, we studied the β3 cytoplasmic domain tethered to phospholipid bilayers. Hydrogendeuterium exchange mass spectrometry and circular dichroism spectroscopy demonstrated that the β 3 cytoplasmic domain becomes more ordered and helical under these conditions, consistent with our NMR

results. Further, these experiments suggest that the two distal helices associate with lipid bilayers, but undergo fluctuations that would allow rapid binding of cytoplasmic proteins regulating integrin activation, such as talin-1 and kindlin-3. Thus, these results provide a framework for understanding the kinetics and thermodynamics of protein interactions involving integrin cytoplasmic domains and suggest that such interactions may act in a concerted fashion to influence integrin stalk separation and exposure of extracellular ligand binding sites.

I. Introduction

Integrins, ubiquitous heterodimeric adhesion receptors, reside on cell surfaces in equilibrium between resting low-affinity and active high-affinity conformations. Interactions involving the transmembrane (TM) and cytoplasmic domains of integrin α and β subunits maintain the low affinity conformation, whereas intracellular signaling events disrupt these interactions, resulting in global conformational changes that activate the integrin (18). Following ligand binding, integrins cluster into complexes that initiate intracellular signaling pathways (35).

On circulating platelets, the integrin α IIb β 3 is maintained in a resting conformation until platelets encounter vascular damage where it is rapidly shifted to its active conformation. This enables α IIb β 3 to mediate platelet aggregation by binding soluble ligands such as fibrinogen and von Willebrand factor (5). Activation occurs when intracellular proteins such as talin-1 and kindlin-3 bind to conserved motifs in the β 3 cytoplasmic

domain, thereby disrupting TM and cytoplasmic domain heterodimers (19). It has been proposed that membrane-proximal α IIb and β 3 cytoplasmic domain sequences form an activation-constraining "clasp", featuring a salt-bridge between α IIb Arg995 and β 3 Asp723 (36). However, studies specifically designed to detect this salt bridge have failed to do so (37).

Although structures for the α IIb β 3 cytoplasmic domain heterodimer have been determined using soluble peptides and NMR, they may not reflect native contacts because they fail to account for constraining TM interactions (38, 39). To investigate the structure and dynamics of the cytoplasmic domain heterodimer as it exists in resting α IIb β 3, we covalently trapped the complex by introducing an inter-subunit disulfide bond near the C-terminal ends of the α IIb and β 3 TM helices. The location of the crosslink was based on a computational model of the α IIb β 3 TM heterodimer (40). While this work was in progress, three structures for the α IIb β 3 TM heterodimer plus short segments of the cytoplasmic domains were reported, confirming the model used to guide the introduction of the disulfide bond (15-17).

We used NMR and hydrogen-deuterium exchange (HDX) mass spectrometry to study the structure and dynamics of the α IIb β 3 cytoplasmic domain heterodimer in micelles and phospholipid bilayers. The α IIb cytoplasmic domain was largely disordered, but the β 3 cytoplasmic domain consisted of a proximal helix contiguous with the TM helix and two distal helices. The conformation of the proximal helix was influenced by the presence of the α IIb cytoplasmic domain, whereas the conformations of the distal helices were similar to those observed previously in a monomeric protein corresponding to the β 3 TM and cytoplasmic regions (37). Further, the distal helices were amphiphilic and likely lie along the inner surface of the platelet membrane. However, their interaction with the membrane is not so stable as to impede rapid binding of cytoplasmic proteins. Thus, our results provide a framework for understanding the kinetic and thermodynamic coupling between the cytoplasmic domains of integrins and intracellular signaling molecules, interactions that act in a concerted fashion to induce cytoplasmic and TM domain separation and the exposure of extracellular binding sites for integrin ligands.

II. Results

Model α IIb β 3 cytoplasmic domain constructs.

Figure 2.1A and 2.1B shows the sequences of the α IIb β 3 TM and cytoplasmic domains. Our laboratory's previous structural studies of the α IIb β 3 cytoplasmic domain focused on the lone β 3 cytoplasmic domain in the absence of α IIb (Figure 2.1C) (37). Given that integrin activation involves separation of the α IIb and β 3 cytoplasmic domains, we reasoned that the structure of this construct would mimic that of the active integrin conformation. In order to understand how α IIb and β 3 might interact with each other in the inactive integrin state in which TM interactions place the

two cytoplasmic domains in close proximity, we designed a disulfide cross-linked NMR construct (Figure 2.1D). Furthermore, to better under stand how the membrane environment might participate in the structure and dynamics of α IIb β 3, we designed two constructs for HDX analysis of the active and inactive α IIb β 3 conformations in the presence of phospholipid bilayers (Figure 2.1E and F).

A β3 Transmembrane and Cytoplasmic Domains

PKGPDILVVLLSVMGAILLIGLAALLI712WKLLI719TIHDRKEFAKFEEERARAKWDTANNPLYKEATSTFTNITYRGT762 Transmembrane Cytoplasmic

B αllb Transmembrane and Cytoplasmic Domains PIWWVLVGVLGGLLLLTILVLAM987WKVGFFK994-RNRPPLEEDDEEGE1008 Cytoplasmic

Transmembrane

C β3 Transmembrane and Cytoplasmic Domain NMR Construct

GSA-PKGPDILVVLLSVMGAILLIGLAALLI712WKLLI719TIHDRKEFAKFEEERARAKWDTANNPLYKEATSTFTNITYRGT762

D αllbβ3 Cytoplasmic Domain NMR Construct

SPE-C987-WKVGFFKRNRPPLEEDDEEGE1008

HisTag-SPE-C⁷¹²-LIWKLLITIHDRKEFAKFEEERARAKWDTANNPLYKEATSTFTNITYRGT⁷⁶²

E β3 Cytoplasmic Domain HDX Construct

GSHMG-C719-TIHDRKEFAKFEEERARAKWDTANNPLYKEATSTFTNITYRGT762 Maleimide-PE (C16)2

F αllbβ3 Cytoplasmic Domain HDX Constructs

GSHMG-C719-TIHDRKEFAKFEEERARAKWDTANNPLYKEATSTFTNITYRGT762 Ac-C994-RNRPPLEEDDEEGE1008

Figure 2.1 The sequences of the α IIb β 3 TM and cytoplasmic domains and experimental constructs used.

(A) The sequence of the β 3 TM and cytoplasmic domains. (B) The sequence of the α IIb TM and cytoplasmic domains. (C) The sequence of the previously reported (37) NMR construct containing the β 3 TM and cytoplasmic domains spanning from residue 691 to 762. The N-terminus contains a Gly-Ser-Ala that remains from the thrombin cleavage site. (D) The sequence of the NMR construct containing the α IIb β 3 cytoplasmic domains spanning from β 3 residue 713 to 762 and α IIb residue 988 to 1008. The β3 sequence contains an N-terminal His-tag as well as a Ser-Pro-Glu helical cap. The αllb sequence contains an N-terminal Ser-Pro-Glu helical cap. (E) The sequence of the lipid-immobilized HDX construct containing the β 3 cytoplasmic domain spanning from residue 720 to 762. It contains an N-termina Gly-Ser-His-Met-Gly from the thrombin cleavage site and the Ndel restriction site. The β 3 cytoplasmic domain was covalently attached to maleimide-functionalized phosphatidylethanolamine via a cysteine substitution at position 719. (F) The sequence of the bilayertethered α IIb β 3 HDX construct spanning β 3 residues 720 to 762 and α IIb residues 995 to 1008. The α IIb β 3 heterodimer contains a disulfide crosslink between β 3 residue 719 and α IIb residue 954. The N-terminus of α IIb is functionalized with a myristoyl group for membrane immobilization. $\{(C)\}$ made by RHL; (D) made by DGM; (E) and (F) made by DTM}

Our NMR construct (Figure 2.1D) contained the complete α IIb and β3 cytoplasmic domains plus a portion of their TM helices. The peptides were crosslinked by a disulfide bond between Cys residues replacing Met987 and Leu712 in the TM domains of α IIb and β 3, respectively. The position for the disulfide bond was based on a model of the resting α IIb β 3 TM heterodimer (41). Recently published NMR structural studies and disulfide crosslinking experiments confirm that a disulfide bond can be accommodated at this position with little change in backbone structure (15-17). For NMR experiments, the α IIb and β 3 peptides were labeled biosynthetically with ¹³C and ¹⁵N in *E. coli* and disulfide-crosslinked after activating the β 3 Cys with 2,2'-dithiobis(5-nitropyridine) (42). The resulting disulfide-crosslinked construct migrated as а single species in polyacrylamide gels with an apparent molecular weight consistent with the mass of the heterodimers. Following disulfide bond reduction, two peptides were observed that migrated with lower apparent molecular weights (Figure 2.2A).



Figure 2.2 The α IIb β 3 cytoplasmic domain heterodimer construct before and after reducing its disulfide bond.

The α Ilb β 3 cytoplasmic domain heterodimer construct before and after disulfide reduction. (a) Coomassie stained SDS-PAGE of the NMR construct. Lane A, molecular weight markers. Lane B, the disulfide-crosslinked construct. Lane C, the construct following disulfide bond reduction. Differences in band intensity reflect differences in the molecular weights of α Ilb and β 3 peptides, the β 3 peptide is three times larger than α Ilb per mole. (b) Circular dichroism (CD) spectra of the α Ilb β 3 heterodimer before (solid line) and after reduction (dotted line). *{Performed by DGM.}*

The NMR construct was not stable in the absence of detergents and when transferred to phospholipid bilayers resulted in precipitation of the mixture. We reasoned that the NMR construct might be unstable in bilayers due to the incomplete TM domain, which would be unable to span the bilayer. Therefore two new constructs were designed for studies in the presence of phospholipids. The intention of these constructs was to mimic the α Ilb β 3 cytoplasmic domain as it exited the bilayer. The first construct was designed to model the active β 3 cytoplasmic domain in the absence of α IIb. In a recent disulfide crosslinking study (15) that mapped the interface between α IIb and β 3 TM and membrane proximal regions, residue IIe719 was found to be the first ß3 cytoplasmic residue to be accessible to chemical cross-linking, suggesting that it is accessible to the cytosolic compartment and likely at the membrane/cytosol interface. Ile719 was also predicted to be at the membrane/cytosol interface when the β 3 TM and membrane proximal regions were modeled using the E(z) membrane insertion potential (43). Based upon these findings, we reasoned that residue 719 would be appropriate for covalent attachment to the phospholipid headgroups. To do so, we introduced a cysteine residue, which was reacted with the surface of bilayers containing maleimide-functionalized phosphatidylethanolamine (Figure 2.1E, top). The β 3 cytoplasmic domain reacted readily with the bilayers and after two hours, only the immobilized construct was observed (Figure 2.3A).

As for our NMR experiments, a disulfide cross-linked α IIb β 3 construct was designed for HDX studies in the presence of bilayers. Besides placing IIe719 at the membrane surface recent disulfide cross-linking studies (15) also found that position 719 was part of a helical interface between α IIb and β 3. In that study residue 719 readily formed a disulfide bond with position 994 in α IIb. Using similar chemistry as for the NMR construct, the β 3 cytoplasmic domain was attached to α IIb

cytoplasmic domain via a disulfide bond between residues 719 and 994 respectively. As disulfide chemistry was used to form the heterodimer, maleimide chemistry could not be used for immobilization on the membrane surface. Instead, a myristoyl group was covalently attached to the N-terminus of α IIb. The myristoylated α IIb β 3 construct was purified by HPLC (Figure 2.3B).



Figure 2.3 Mass spectra of membrane tethered α IIb β 3 cytoplasmic domains.

(A) The β 3 cytoplasmic domain peptide (MW = 5680) containing an Nterminal cysteine residue was reacted with large unilamellar vesicles (LUVs) for 1.5 hours at pH 6.8 producing the lipid conjugated β 3 cytoplasmic domain (MW = 6592). MALDI-TOF mass spectrometry was performed on the starting material (bottom, magenta), the reaction after 30 minutes (middle, green), and the reaction after 1.5 hours (top, blue). (B) Myristoylated- α IIb β 3 heterodimer was prepared as described in the methods, purified by HPLC, and subjected to MALDI-TOF mass spectrometry. {*Performed by DTM.*}

Association of the α IIb and β 3 cytoplasmic domains influences their conformation.

Due to the inclusion of TM domain residues, the heterodimeric construct was insoluble in aqueous buffers and was studied in dodecylphosphocholine (DPC) micelles, similar to our previous analysis of the monomeric β 3 TM and cytosolic regions (37). CD spectroscopy of the disulfide-linked construct contained bands at 190, 208, and 222 nm consistent with helical secondary structure (Figure 2.2B). Disulfide bond reduction resulted in a decrease in the magnitude of the molar ellipticity at 222 nm, indicating that the disulfide-linked construct was more structured than its component monomers.

The interface between α IIb and β 3 was mapped by comparing ¹³C resonances of the disulfide-linked and monomeric peptides. Like the monomeric β 3 cytoplasmic domain (37), C α resonances of the β 3 cytoplasmic domain in the disulfide-linked construct were consistent with the presence of three helices: a proximal helix contiguous with the β 3 TM domain (TMproxH) and two distal cytoplasmic helices (CytoH1 and CytoH2) (Figure 2.4A). No differences were observed between the distal helices of monomeric and disulfide-linked β 3, but there were significant differences in the residues that define the proximal helix. The largest differences occurred at residues 720 through 724, eight to twelve residues downstream from the disulfide bond (Figure 2.4B), implying that the differences reflect α IIb and β 3 interactions rather than local perturbations caused by the disulfide bond itself. Further, the chemical shifts are

indicative of increased helical content, consistent with the increased helical character in the CD spectrum following disulfide bond formation.



Figure 2.4 Interaction between α IIb and β 3 cytoplasmic domains.

(A) C α chemical shifts define three helical regions of β 3 in the disulfidelinked heterodimer. Differences between the C α resonances in the α IIb β 3 heterodimer and β 3 monomer are shown as bars above the graph. (B) ¹³C HSQC spectra of aliphatic atoms in the α IIb β 3 heterodimer (blue/purple) and the α IIb and β 3 monomers (red and green). {*Performed by DGM.*}

The ¹³C HSQC spectra of the disulfide-linked and reduced states were compared in order to determine the interface between the α and β chains (Figure 2.4B). The chemical shifts of the Lys716 and Ile719 side chains changed when the disulfide bond was reduced, while no differences were observed for neighboring residues, suggesting that Lys716 and IIe719 interact directly with α IIb (further supporting the choice of disulfide cross-link in Figure 2.1F). Thus, Lys716 and Ile719 define a face of TMproxH that includes Asp723, a residue postulated to form a salt bridge with α IIb Arg995 (36). Analysis of the Asp723 side chain neither confirmed nor refuted its interaction with α IIb. We did not identify NOEs between Arg995 guanidine protons and β 3 sidechains. The Arg995 resonances were characteristic of a dynamic side chain and did not change substantially when the disulfide bond was reduced. Further, NMR spectra and HDX results suggest that Asp723 and Arg995 are highly solvent-exposed. Thus, any acid/base interaction would appear to be dynamic and partially screened under physiological salt conditions.

Structure of the $\beta 3$ cytoplasmic domain crosslinked to the αllb peptide

Three-dimensional structures were calculated for the α IIb and β 3 cytoplasmic domains using intra- and interchain NOEs, HN-H α Jcouplings, and statistical analysis of chemical shifts (Figure 2.5A). We were unable to assign more than 5 strong NOEs between α IIb and β 3, precluding calculation of a high-resolution structure. Moreover, the intraamide ¹H-¹⁵N HSQC peak heights for α IIb became progressively greater from its N- to its C-terminus, implying that the α IIb cytoplasmic domain is largely unstructured near its C-terminus (Figure 2.6A). Thus, we focused on defining the structure of β 3 in the presence of α IIb. A structure for β 3 was computed using 16 long-range, 235 medium-range, 259 short-range, and 202 intra NOE restraints, 84 dihedral angle restraints, and 24 hydrogen bonds (Table S2.1).



Figure 2.5 The structure of the β 3 cytoplasmic domain.

(A) Twenty structures in the NMR ensemble, consisting of two regions encompassing residues Leu713-Trp739 and Asp740-Thr762, were aligned separately over well-ordered regions. Domains interacting with α IIb, talin-1, kindlin-3, and c-Src are enclosed by dashed lines and side chains that make up interfaces are depicted as sticks. The two β 3 NPXY motifs are colored magenta and the Src-binding RGT motif is colored cyan. (B) A single residue hinge at Arg724 is stabilized by packing between residues Ile721 and Phe727 (green) and an electrostatic interaction between Arg724 (blue) and Glu726 (red). (C) The first cytosolic helix (residues Glu726-Arg736) is stabilized by a network of complementary charged residues. Positively charged residues are colored blue and negatively charged residues red. {*Structure solved by DGM.*} The C-terminal ends of TMproxH and CytoH1 pack together in a well-structured helix-turn-helix motif. Arg724 serves as a single-residue hinge between the two helices, having ϕ/ψ angles in the β /PPII region of the Ramachandran plot and producing a 100.5 ± 8.8° angle between the two helices. A series of interactions, including strong NOEs between Phe727 and IIe721, as well as electrostatic interaction between Arg724 and Glu726, stabilize the turn (Figure 2.5B). CytoH1 itself is stabilized by a highly favorable arrangement of 12 intra-helical electrostatic interactions between residues *i* and *i*+3/4 (Figure 2.5C).

CytoH1 is followed by a flexible linker spanning residues 738-743. CytoH2 is then initiated by Asn744, the first residue of an NPXY motif consisting of residues Asn744-Tyr747 (44, 45). Although NPXY motifs are often found in β -turns, they are also helix initiation sites (46). Indeed, Asn744-Tyr747 adopts an identical conformation in a crystal structure of the F2F3 domains of talin-1 fused to a portion of the β 3 cytoplasmic domain (pdb code 1MK7, chain A residues 744-747) (44).

Conformational flexibility of the α IIb β 3 cytoplasmic domains in micelles

The dynamics of the cross-linked α IIb and β 3 cytoplasmic domains in DPC micelles were studied by measuring the intensity of intra-amide ¹H-¹⁵N HSQC peaks (Figure 2.6A) (47, 48). α IIb residues Val990-Asn996 and β 3 residues Leu713-Glu726, comprising the TM-proximal helices, displayed similar peak intensities, indicating limited mobility and stable
structures. β 3 CytoH1 was somewhat more dynamic, while CytoH2 showed graded mobility, becoming increasingly less structured near its C-terminus. As noted, ¹H-¹⁵N HSQC peak intensities for α IIb became progressively greater near its C-terminus, implying that the terminal region of the α IIb cytoplasmic domain is largely unstructured. These findings were confirmed by NMR measurements of amide proton exchange with solvent deuterons (HDX, Figure 2.6B) (49). Indeed, H-¹⁵N HSQC spectra of TMproxH were highly protected from HDX, while the distal helices exchanged in a rapid time regime that was not amenable to NMR studies.



Figure 2.6 Backbone dynamics of the α IIb β 3 cytoplasmic domain heterodimer.

(A) Backbone dynamics were inferred from intra-amide ${}^{1}\text{H-}{}^{15}\text{N}$ HSQC peak heights and were compared with the corresponding isolated β 3 construct (37). (B) ${}^{15}\text{N}$ HSQC spectrum of the disulfide-linked complex before and after HDX. The β 3 backbone amides IIe714-IIe721 are protected from HDX, suggesting they form stable hydrogen bonds in a helix. Other helical regions exchange with deuterium on the minute time-scale, suggesting they make excursions from helical conformations. *{Performed by DGM}*

Conformational flexibility of the α IIb β 3 cytoplasmic domain in phospholipid bilayers.

Accordingly, we used mass spectrometry to measure HDX kinetics for the distal helices in phospholipid bilayer vesicles, a condition more closely resembling natural membranes than detergent micelles. This type of experiment has been used to determine membrane protein structure and measure unfolding rates (50, 51). We focused on the β 3 segment spanning residue Thr720 to the C-terminus, because, as discussed above, this represents the portion of β 3 accessible to the cytosolic compartment and the full-length NMR construct was not stable in bilayers. The cytoplasmic domain was studied in aqueous solution as well as tethered to preformed large unilamellar vesicles (LUVs). As with our NMR studies, β 3 was probed in the presence and absence of α IIb in order to determine the effects of α IIb on its structure.

CD spectroscopy revealed that when the β 3 cytoplasmic domain was covalently tethered to bilayers, the peptide's α -helical content increased and the random coil signature decreased (Figure 2.7), indicating that the β 3 cytoplasmic domain interacts with phospholipid bilayers. To measure HDX, free and tethered β 3 peptides were incubated in D₂O buffer for varying periods of time, after which amide deuterium exchange was quenched at pH 2.5. The quenched peptides were injected onto an immobilized pepsin column to generate fragments that were analyzed by tandem LC/MS. Free β 3 peptide exchanged rapidly with D₂O (Figure 2.8, red), indicating that its conformation was largely disordered, consistent with CD results (Figure 2.7). When the β 3 peptide was membrane-bound, the most membrane proximal region, Thr720-Glu732, remained tethered to the lipid after proteolysis and was not observed. However, we observed two fragments that contained portions of CytoH1 and CytoH2. Both fragments had two kinetically distinct classes of amides: one class had unprotected amides that exchanged at a rate indicative of an unstructured backbone; the second class exchanged with a protection factor on the order of 10² to 10³, indicative of a helix making occasional excursions to less well-structured conformations that allow HDX. The fragment containing the C-terminus of CytoH1, the flexible loop, and the N-terminus of CytoH2 (Glu733-Thr753) had approximately 8 protected amides (Figure 2.8A, green). The second fragment contained the C-terminus of CytoH2 (Asn756-Thr762) and had only one protected amide (Figure 2.8B, green). These results confirmed the location of CytoH1, which was identified by NMR, as well as the hierarchy of increasing disorder that progresses from the N- to the C-terminus of the β 3 cytoplasmic domain.



Figure 2.7 Circular dichroism confirms that interactions with phospholipids stabilizes a helical conformation.

CD spectra were taken of free β 3 in the absence of lipid (black dots), membrane- tethered β 3 (solid line), and β 3 mixed with lipids, but not reacted with the maleimide head group (grey dots and lines). CD spectra of β 3 alone were taken of 30 μ M β 3 in 25 mM phosphate 100 mM NaCl, pH 6.8. For lipid conjugated β 3, 15 μ M β 3 was conjugated to 1 mM total lipid containing 20% POPG, 12% maleimide-PE, and 68% POPC in 25 mM phosphate with 100 mM NaCl, pH 6.8. Unconjugated β 3 in the presence of lipid contained 15 μ M β 3 and 1mM total lipid containing 20% POPC. *{Spectra are representative of three independent experiments. Performed by DTM]*

Our NMR results suggest that when α IIb is held in close proximity to β 3 – as would be expected in the inactive α IIb β 3 conformation – a kink is stabilized in β 3. We hypothesized that such a kink would project distal β 3 cytoplasmic elements such as CytoH1 and CytoH2 parallel to the membrane and encourage their interactions with the bilayer beyond the interactions observed with lipid-immobilized β 3. We therefore sought to

determine by HDX whether this kink in the presence of bilayers might stabilize distal structures observed by NMR. As described above, we synthesized and purified a disulfide cross-linked α IIb β 3 construct with an N-terminal myristoyl membrane anchor. This peptide was not water soluble, but was readily solubilized when extruded with phospholipid When HDX experiments were performed on this α IIb β 3 bilayers. construct, similar pepsin cleavage products were observed. Again, the Nterminus was not seen, because of its covalent attachment to allb, but peptides containing residues Glu733-Thr753 and the Asn756-Thr762 were observed. Like β 3 in the absence of α IIb (Figure 2.8A, green), the peptide spanning Glu733-Thr753 was found to have approximately 8 protected amides in the presence of bilayers (Figure 2.8A, blue). The protection pattern for the most distal fragment was quite different, though. The peptide containing Asn756-Thr762 was found to be significantly protected from exchange. In the presence of bilayers, the heterodimeric α IIb β 3 cytoplasmic domain had 3-4 amides with protection factors on the order of 10²-10³ (Figure 2.8B, blue). These results support the NMR-based model that the presence of α IIb stabilizes the interaction between β 3 and the membrane. It is possible that the protection of the distal β 3 segment from HDX could arise from direct interactions with α IIb and as opposed to a helical interaction with the membrane, however, NOEs between α IIb and the β 3 segment were not observed in the NMR.

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Figure 2.8 HDX-MS of the α IIb β 3 cytoplasmic domain in phospholipid bilayers.

Model α IIb β 3 cytoplasmic domain constructs were prepared in pH 6.8 phosphate with 100 mM NaCl. Each sample was diluted 1:10 in pH 6.8 phosphate with 100 mM NaCl containing 90% D₂O on ice. Deuteration was quenched at pH 2.5 on ice and samples were injected onto a pepsin column in tandem with LC-MS to resolve fragments and calculate deuteration. The deuteration reaction for each sample was fit to a double exponential rate equation assuming subpopulations of rapidly and slowly exchanging amides. Peptide fragments containing (A) Glu733-Thr753 and (B) Asn756-Thr762 were observed. For each peptide a theoretical curve based upon the predicted intrinsic exchange rate for a disordered peptide was generated (black). HDX was performed on free β 3 cytoplasmic domain (red), the β 3 cytoplasmic domain covalently immobilized on phospholipid bilayers containing 20% POPG, 12% maleimide-PE, and 68% POPC (green) which is predicted to model the active β 3 state, and disulfide cross-linked α IIb β 3 in phospholipid bilayers containing 20% POPG and 80% POPC (blue) which is predicted to model the inactive state. Residues highlighted in blue are predicted to be protected from exchange based upon the NMR structure. Residues highlighted in orange correspond to regions found to be disorganized in the NMR structure or at the N-termini of helices and not predicted to be protected from exchange. {Experiments performed once by DTM}

The increased helicity of the β 3 cytoplasmic domain (Figure 2.7) in the presence of bilayers and its protection from HDX under these conditions suggest that it interacts with the membrane surface. We used E(z), a depth-dependent potential (43), to assess the binding of the β 3 helices in membranes. As expected, TMproxH was predicted to partition into the bilayer perpendicular to the membrane as an extension of the TM helix, whereas CytoH1 and CytoH2 were predicted to lie along the membrane surface, stabilized by hydrophobic residues Phe727 and Phe730 in CytoH1 and Leu746, Tyr747, Ala750, Phe754, and Ile757 in CytoH2 (Figure 2.9A).

III. Discussion

We have studied the structure and dynamics of the platelet integrin α IIb β 3 cytoplasmic domain. We found the α IIb cytoplasmic domain intrinsically disordered, consistent with its paucity of hydrophobic residues and a recent NMR study using phospholipids bicelles (52). Nonetheless, chemical shift perturbations indicate that α IIb interacts with the TM proximal portion of β 3 over the length of 12 residues, implying that α IIb has a conformational bias toward structures that interact with β 3. This type of intrinsically disordered interaction has been found in proteins that couple multiple signaling events to an overall equilibrium (53). In the case of α IIb β 3, an intrinsically disordered interface would allow the integrin to

rapidly shift between resting and active conformations following a breach in vascular integrity.

The N-terminal portion of the β 3 cytoplasmic domain consists of a highly stable helix contiguous with the C-terminal end of the TM helix and separated from a second surface-associated helix by a single-residue hinge at Arg724. The remainder of the cytoplasmic domain consists of a short unstructured region, a second cytoplasmic helix and a flexible Cterminal tail. CytoH1 and CytoH2 are likely stabilized by interactions with the membrane bilayer (Figure 2.9A), although they are not so stable as to lock the cytoplasmic domain into a single fixed conformation. Instead, CytoH1 and CytoH2 undergo frequent excursions into non-helical conformations within the aqueous environment. The association of CytoH1 and CytoH2 with membranes is consistent with previous NMR experiments observing NOEs between micellar DPC and Trp739/Tyr747 (54). Amphiphilic domains proximal to TM domains have also been detected in proteins such as voltage gated channels (55, 56), phospholamban (57), and the influenza M2 proton channel (58, 59). The association of cytoplasmic helices with membranes can change in response to gating ligands or post-translational modification. For example, the cytoplasmic helix of the pH-activated M2 channel changes its orientation in response to pH. Also, aromatic residues in the cytoplasmic domain of the T-cell receptor $CD3\varepsilon$ subunit were found to insert into membranes, including a tyrosine in its NPXY motif (60). Following T-cell

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receptor activation, the amphiphilic domain is released from the membrane.



Figure 2.9 A linked equilibrium describing the activation of α IIb β 3 by intracellular signaling molecules.

(A) The E(z) membrane insertion potential was used to model the interaction of β 3 with the membrane. It predicts that aromatic residues (green) on one face of CytoH1 and CytoH2 will preferentially bind the membrane. (B) The dynamic nature of the β 3 cytoplasmic domain makes protein-protein interaction motifs accessible. Dissociation of α IIb and extension of the β3 TM-proximal helix to join CytoH1 allows talin-1 to bind and places positively-charged residues on talin-1's upper surface in a favorable position to interact electrostatically with the inner membrane leaflet. B3 is also now available to bind kindlin. Residues that are perturbed by the α IIb cytoplasmic domain are shown in red. (C) An equilibrium model of talin-1 and kindlin-3 binding the integrin cytoplasmic domain. To the left is a resting conformation with helices CytoH1 and CytoH2 interacting with the bilayer. The fully inactive state is not competent to bind talin-1 or kindlin-3 because the membrane obstructs access to β 3. To the right is a fully active conformation with separated TM and cytoplasmic domains. In between exists an ensemble of states in which the integrin is not active, but the β 3 cytoplasmic domain samples a wide range of conformations and is accessible to intracellular molecules. Although talin-1 and kindlin-3 can bind the ensemble of free cytoplasmic domains in the middle column, available data are insufficient to predict the order of binding. Therefore, equilibria depicting talin-1 binding first (top) and kindlin-3 binding first (bottom) are shown. However the fact that both kindlin-3 and talin-1 are necessary for activation suggests that both bind to the integrin cytoplasmic domain.

The marginally stable interactions between the membrane bilayer and the β 3 cytoplasmic helices are likely important for allowing kinetic access to cytoplasmic proteins, such as talin-1 and kindlin-3, responsible for triggering α IIb β 3 activation (19). The structure of β 3 elucidated here presumably represents the resting state which must undergo structural changes to allow binding of these proteins. The talin-1 binding site, extending from the C-terminus of the TM helix through CytoH1 to the NPxY motif near the N-terminus of CytoH2, is masked in the NMR structure by interactions with the α IIb tail and the phospholipid bilayer. Interaction with membranes is facilitated by the bend between TMproxH and CytoH1 at Arg724; if this residue instead continues the TMproxH α -helix (Figure 2.9B), CytoH1 becomes an extension of the TM helix, allowing binding of the talin-1 head domain (61, 62). This orientation not only allows interaction of the talin-1 F3 domain with the β 3 tail, but also electrostatic interactions between acidic lipids on the cytoplasmic leaflet of the bilayer and positively charged residues in talin-1's F2 and F3 domains. Moreover, the interaction of the β 3 tail with talin-1 removes the C-terminal region of β 3 from the membrane and positions its kindlin-binding site for simultaneous interaction with kindlin-3. Such a tandem interaction has recently been demonstrated for dimerization of the PTB domain of the cytoplasmic adaptor protein Dok7 (63). Displacement of this segment from the membrane also destabilizes the C-terminal β 3 helix, allowing it to more easily conform to the requirements for kindlin-3 binding.

The interactions of talin-1 and kindlin-3 with β 3 show the hallmarks of a cooperative system; association of either protein with its β 3 binding site occurs at the thermodynamic expense of disrupting favorable β 3membrane binding interactions. Moreover, binding of one protein primes the system for a more favorable interaction with the second. Thus, the sites are both kinetically and thermodynamically linked, as shown in Figure 2.9C. The talin-1 binding event is likely the key step in tail separation, given the proximity of the talin-1 interacting site to the α Ilb β 3 interface. Thus, at sufficiently high talin-1 concentrations, it should be possible to activate α IIb β 3, even in the absence of kindlin-3 (64). However, under physiologic conditions, our model predicts that kindlin-3 also plays a pivotal role for interaction of talin-1 with the β 3 tail. At present, it is unclear whether there is a preferred temporal sequence for binding of kindlin-3 versus talin-1 during physiological α IIb β 3 activation. However, the greater mobility of the kindlin-3 binding site in the β 3 cytoplasmic domain suggests it might have a kinetic advantage over talin-1, assuming both are present in appropriate forms for binding. Figure 2.9C (top and bottom) illustrates both kinetic possibilities for the initial binding step. Once initial binding occurs, subsequent interactions and conformational changes become increasingly favorable, ultimately tipping the equilibrium towards tail separation and α IIb β 3 activation (Figure 2.9C, middle panel). This thermodynamic coupling model could be extended to any number of cytoplasmic events playing negative or positive roles in activation. We predict that direct interactions of the β 3 cytoplasmic domain with α IIb and the phospholipid bilayer help restrain α IIb β 3 in a resting state. Any protein that binds more tightly to the separated cytoplasmic domains than the cytoplasmic domain heterodimer will have an activating effect. Furthermore, proteins that negatively impact talin-1 binding will have an inhibitory effect on α IIb β 3 activation.

IV. Methods

Plasmid Constructs—A cDNA encoding αIIb residues Trp988-Glu1008 was cloned into the vector pET-16b (Novagen) as a C-terminal fusion with the designed protein $\alpha_3 D$ (65). The region between the α IIb sequence and $\alpha_3 D$ included a proline to disrupt secondary structure, two glycines as a flexible spacer, a thrombin cleavage site, an SPE N-terminal helix cap, and a cysteine for crosslinking to the β 3 peptide. The α IIb peptide sequence was: $\alpha_3 D$ -P-GG-LVPR-SPE-C-WKVGFFKRNRPPLEEDDEEGE. A cDNA encoding β 3 residues Leu713-Thr761 was cloned into the vector pET-15b (Novagen) downstream from a polyhistidine tag. The construct included an SPE N-terminal helix cap followed by a cysteine for crosslinking. The β 3 peptide sequence was: *MGSSHHHHHHSSGLVPRGSHM-SPE-C*-

LIWKLLITIHDRKEFAKFEEERARAKWDTANNPLYKEATSTFTNITYRGT. The fidelity of the constructs was confirmed by DNA sequencing.

A shortened β3 cytoplasmic domain (MGSSHHHHHHSSGLVPRGS-|-H

MG-C-TIHDRKEFAKFEEERARAKWDTANNPLYKEATSTFTNITYRGT) was cloned into the pet-28a vector. It contained a cysteine for reaction with phospholipid bilayers or αIIb.

Protein Expression and Purification—The α_3 D-αIIb fusion protein and the His-tagged β3 protein were expressed in *E. coli* strain BL21 and labeled biosynthetically with ¹³C glucose and/or ¹⁵N ammonium chloride (Cambridge Isotope Laboratories). The α 3D- α IIb fusion protein was isolated from the supernatant of bacterial lysates, cleaved with thrombin, and purified by reverse-phase HPLC. The His-tagged β3 protein was isolated on Ni-NTA resin using the QIAExpressionist protocol for denaturing conditions (Qiagen) and purified by reverse-phase HPLC. These protocols generated >10 mg of the α IIb and β 3 proteins per liter culture. The molecular weight of each product was confirmed by mass spectroscopy.

The His- β 3 construct used for HDX was expressed in BL21-DE3 *E. coli* and purified using Ni-NTA chromatography on an Akta FPLC (General Electric Healthcare). Purified His- β 3 was dialyzed into 50mM Tris, 100 mM NaCl, pH 7.5 and cleaved with thrombin overnight at room temperature. The resulting peptide (GSHMG-C-TIHDRKEFAKFEEERARAKWDTANNPLYKEATSTFTNITYRGT) was purified by reverse phase HPLC on a C4 preparative column (Vydac) on a water/acetonitile/ⁱPrOH gradient in 0.1% TFA.

SDS-PAGE of $\alpha IIb\beta 3$ NMR construct-The disulfide cross-linked heterodimer was dissolved in LDS loading buffer and run on a NUPAGE SDS-PAGE gel. Excess TCEP was added to an identical sample and run in an adjacent lane. The gel was washed and stained with Coomassie GelCode stain (Pierce).

Synthesis of the myristoylated α IIb cytoplasmic domain– α IIb-CT was synthesized on Wang resin preloaded with an Asp residue using a Symphony peptide synthesizer at 0.1 mM scale in 25% DMSO/75% NMP

with Fmoc-chemistry. α IIb-CT was myristoylated on resin by reacting the protected peptide with 5-fold excess HBTU activated myristic acid (Sigma) for three hours at room temperature. α IIb-CT was cleaved from the resin TFA/triethylsilane/water (95/2.5/2.5 v/v) for three hours and precipitated into dry-ice cold diethylether. β 3-NITYRGT was purified by reverse phase HPLC (Varian Prostar) using a C4 preparative column (Vydac) on a water/acetonitrile/ⁱPrOH gradient with 0.1% TFA.

Formation of a disulfide-linked $\alpha IIb\beta 3$ cytoplasmic domain heterodimer for NMR—The $\beta 3$ peptide was dissolved at 5 mg/ml in 50 mM MES buffer, pH 6.0, containing 2M guanidine, after which its cysteine thiol was activated with a 5-fold molar excess of 2-2'dithio-bis-(5-nitropyridine) dissolved in DMSO (42). After excess 2-2'dithio-bis-(5-nitropyridine) was removed on a PD-10 column (GE Healthcare), the activated $\beta 3$ was reacted with a slight molar excess of the αIIb protein in guanidine-MES buffer, pH 6.5. Disulfide-linked $\alpha IIb\beta 3$ was purified using reverse phase HPLC in a gradient with 60% iPrOH, 30% ACN, 10% H₂O with 0.1% TFA. Its molecular weight confirmed by mass spectrometry.

Preparation of a disulfide-linked membrane-tethered $\alpha IIb\beta 3$ cytoplasmic domain heterodimer for HDX. The $\beta 3$ cytoplasmic domain was prepared as described earlier. To form the myristoylated $\alpha IIb\beta 3$ heterodimer, the $\beta 3$ cytoplasmic domain was reacted with 2-2'dithio-bis-(5nitropyridine) in 50% acetic acid to produce activated 2-thio-(5nitropyridine) adduct, which was transferred to pH 5 sodium acetate buffer and separated from unreacted 2-2'dithio-bis-(5-nitropyridine) by passage over a PD-10 buffer exchange column (GE Healthcare). The activated β 3 was diluted nine fold with acetonitrile and reacted with a slight excess of Myr- α IIb. It was crucial to perform this reaction in an organic solvent, as Myr- α IIb homodimers were the primary product in aqueous buffers, likely due to formation of micelles or aggregates due to the myristic functional group that produced a high local concentration of Myr- α IIb thiol that displaced β 3 from the α IIb β 3 heterodimer. The reaction was diluted with pH 2 water and immediately frozen and lyophilized for purification by reverse phase HPLC on a gradient with 60% iPrOH, 30% ACN, 10% H₂O with 0.1% TFA.

Preparation of membrane tethered β3. Lyophilized Cys-β3 construct was lyophilized and re-dissolved in bilayer conjugation buffer (25mM sodium phosphate or MOPS, 100 mM NaCl, pH 6.8). Large unilamelar vesicles (LUVs) were generated by evaporating a mixture of 20% 1-palmitoyl-2-oleoyl-sn-glycero-3-phospho-(1'-*rac*-glycerol) (POPG, Avanti Polar Lipids), 4 fold molar excess of 1,2-dipalmitoyl-*sn*-glycero-3-phosphoethanolamine-N-[4-(p-maleimidomethyl)cyclohexane-

carboxamide] (16:0 PE MCC, Avanti Polar Lipids) relative to β 3 peptide, and 1-palmitoyl-2-oleoyl-*sn*-glycero-3-phosphocholine (POPC, Avanti Polar Lipids) to a thin film under nitrogen and re-suspending the dried lipids in the bilayer conjugation buffer. LUVs were prepared by extruding 20 times through a 0.1 μ m filter (Avanti Polar Lipids). β 3 peptide was reacted with 4 fold molar excess 16:0 10 PE MCC. The conjugation reaction was performed at pH 6.8 to minimize lysine amine-maleimide side reaction which is strongly disfavored at below pH 7.

NMR Spectroscopy—Purified, disulfide-linked αIIbβ3 heterodimer was dissolved at 1 mM in 5 mM sodium phosphate buffer, pH 6.0, containing 100 mM perdeuterated DPC (Cambridge Isotope Laboratories), 10% deuterium oxide, and 0.02% sodium azide. Samples were analyzed at 37°C on a 750 MHz NMR spectrophotometer equipped with a Varian probe (66).

CD spectroscopy for the NMR sample—CD spectra of 10 μ M disulfide-linked α IIb β 3 heterodimer in 5 mM sodium phosphate buffer, pH 6.5, containing 10mM dodecylphosphocholine (DPC) were collected on a JASCO J-810 spectropolarimeter with a path length of 1 mm before and after the addition of a 5-fold molar excess of the reducing agent tris(2-carboxyethyl) phosphine hydrochloride (Sigma-Aldrich). Measurements were made stepwise for each 1 nm between 190 nm to 260 nm and averaged over ten scans.

Assignment of chemical shifts—The backbone, aliphatic, and aromatic resonances in the α IIb and β 3 subunits were assigned using a standard battery of experiments.(66) Complete residue-specific assignments for the H α , N, C α , and C β resonances in β 3 were obtained using the HNCACB(67) and CBCA(CO)NH(68) experiments. Carbonyl

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carbons were assigned from HNCO experiments.(69) Aliphatic protons identified using H(CCCO)NH,(70, and carbons were 71) (H)CC(CO)NH,(70, 72, 73) and H(C)CH-TOCSY(74-77) experiments. Pro-chiral leucine and valine methyls were distinguished in the ¹³Ccoupled carbon HSQC of a metabolically labeled sample expressed in media containing 10% ¹³C glucose.(78) Aromatic resonances were assigned using HBCBCGCDHD and HBCBCGCDCEHE experiments,(79) an aromatic carbon HSQC, an aromatic ¹³C-edited NOESY, and other NOESY experiments. Assignments for α IIb residues 988-1000 were completed using the same strategy. Additionally, resonances in the α IIb acidic tail E₁₀₀₁EDDEEGE₁₀₀₈ were identified, but were not useful for structural analysis due to spectral overlap. Proton chemical shifts were referenced to residual water signal, while ¹³C and ¹⁵N chemical shifts were indirectly referenced. Chemical shifts were deposited in the BioMagResBank (80).

Identification of NOEs—Nuclear Overhauser Effect restraints (NOEs) were obtained from an ¹⁵N-editied NOESY, an aliphatic ¹³C-edited NOESY, an aromatic ¹³C-edited NOESY, and a 4D ¹³C-edited NOESY.(81, 82) Mixing times were optimized for different NOESYs and ranged from 80 ms in the ¹³C-edited NOESYs to 100 ms in the ¹⁵N-editied NOESY. The solvent contained 99% D₂O in the 4D NOESY.

 $HN-H\alpha$ J-coupling—Phi dihedral angle restraints were obtained from HN-H α J-couplings measured in the HNHA experiment.(81) Phi angles were calculated using the Karplus equation with coefficients A = 6.51, B = -1.76, and C = 1.60. β 3 residues Lys716-Thr720, Phe727-Arg736, and Leu746-Asn756 were given helical phi restraints based on chemical shift analyses, NOE patterns, and HDX experiments that suggest they reside in helices. Other residues were given J-coupling restraints that restrict the phi dihedral to 1-4 possible angles.

Hydrogen-deuterium exchange (HDX)— HDX was performed by lyophilizing a NMR sample and re-dissolving it in deuterium oxide. Exchange was monitored in the HSQC experiment and the first time-point was taken at 11 minutes.

Residual dipolar couplings (RDCs)— RDCs were obtained for samples aligned in axially stretched polyacrylamide gels. A dried, 6mm diameter, 5.5% polyacrylamide gel was soaked with an NMR sample for 72 hours, then funneled into a 4.2mm diameter NMR tube.(83) C α -C' RDCs were obtained using a C α -coupled HNCO experiment.(69)

Implementation of distance restraints—Distance restraints were derived from NOE experiments and HDX. NOESY cross-peak volumes were converted to upper distance limits by the program CALIBA of the CYANA package (84). NOE peaks were divided into three classes with different calibration functions and constants: a $1/r^6$ dependence was used for backbone-proton-backbone-proton NOE values, and a $1/r^4$ dependence for NOEs involving side-chain protons and methyl groups. The upper and lower cutoffs for NOEs before application of pseudoatom corrections were set as 2.4 and 6.0 Å for strong and weak NOEs, respectively. During the structure calculation and NOE assignment, the calibration curves were refined iteratively based on plots of cross-peak volumes versus average proton-proton distances in the current structures. For each distance constraint, the lower limit was set as 1.8 Å, which corresponds to two hydrogen van der Waals radii. Additionally, 24 hydrogen bonds were used for the regular secondary elements Glu711-Asp723, Glu726-Ala736, and Pro745-Ile758. 5 hydrogen bond restraints, which were also confirmed from HDX, were implemented by setting the N to i+4 O distance to 3.0 Å with 10% error, while the remaining hydrogen bond restraints were held with 20% error.

Implementation of angle restraints—Dihedral angle restraints were derived from TALOS+(85) (86) and J-coupling experiments.(81) The TALOS algorithm predicts backbone phi and psi angles based on protein sequence and assigned resonances from a database of known structures. TALOS-derived phi and psi dihedral restraints were used for β 3 residues Cys712 - Thr758, except for residues Arg724, Lys738 and Trp739, which were predicted to be helical with high confidence, consistent with NOE patterns, HDX, and J-coupling. Additionally, phi dihedral angle restraints were obtained from HN-H α J-couplings. β 3 residues Leu713-Asp723, Glu726-Arg736, and Pro745-Thr758 were given helical phi and psi angle restraints based on chemical shift analysis, NOE patterns, and HDX experiments which suggest they reside in helices. Other residues were given J-coupling restraints that restrict the phi dihedral to 1-4 possible angles.(81)

Implementation of RDC restraints—RDCs were measured and evaluated. Nonetheless, the dynamics of the α IIb β 3 construct prevented fruitful refinement of the structural ensemble using RDCs.

Structure Calculation—Data were processed with FELIX, analyzed with Sparky and Xeasy (87) , and structures were calculated using CYANA (84) and finally refined by XPLOR-NIH.(88) Structure calculations and refinement algorithms were based on the anneal.py and refine.py example scripts that come with the XPLOR-NIH package. Structures were calculated for the β 3 portion of the disulfide linked α IIb/ β 3 construct consisting of the SPE N-terminal helix cap, the cysteine used to form the disulfide crosslink, and β 3 residues 713-762:

SPE-C-

LIWKLLITIHDRKEFAKFEEERARAKWDTANNPLYKEATSTFTNITYRGT

where the italicized residues were engineered and the rest of the construct corresponds to the β 3 cytosolic domain. Each algorithm included an initial, high temperature simulated annealing cycle followed by a longer simulated annealing cycle with both cycles restricted to torsional movements. After the simulated annealing cycles, the output conformation was minimized in both torsional and Cartesian space. The energy function included a repulsive van der Waals term and potentials that enforced bond lengths, bond angles, and improper angles.

Additionally, the refinement algorithm included a potential that enforced backbone geometries that are consistent with Ramachandran space. Other potentials enforced NMR restraints as described below.

No violations >0.5 Å per conformer were observed. The statistics of the structure determination are summarized in Table S2.1 and show that a high-resolution and high-quality structure was obtained. The final structure was deposited in the PDB(89) (accession code: 2KV9).

Calculation of a structural ensemble for the β 3 cytoplasmic domain—NOE distance restraints were derived from a 3D ¹⁵N-edited NOESY, 3D aliphatic ¹³C-edited NOESY, 3D aromatic ¹³C-edited NOESY, and a 4D ¹³C-edited NOESY (75). Hydrogen bonding distances and geometry restraints (90) were implemented for the backbone amides of the β 3 regions Glu711-Asp723, Glu726-Ala736, and Pro745-Ile758, predicted to reside in α -helices based upon NOEs and TALOS+. The ϕ and ψ dihedrals of β 3 residues Cys712-Thr758 (excluding Arg724, Lys738 and Trp739) were restrained based on a statistical analysis of chemical shifts using the TALOS+ algorithm (85). Structures were calculated using CYANA (84) and refined with XPLOR-NIH using a protocol similar to the anneal.py and refine.py example scripts that come with the software package (88, 91). A summary of the structural restraints is reported in Table S2.1.

Circular Dichroism (CD) spectroscopy of lipid-conjugated β 3. CD spectra were taken on a JASCO J-810 spectropolarimeter. 15 μ M β 3 was

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conjugated to 1 mM total lipid and probed in 25 mM phosphate with 100 mM NaCl, pH 6.8. Spectra of free β 3 were obtained of 30 μ M β 3. Measurements were made stepwise for each 1 nm between 200 nm to 240 nm and averaged over three scans.

Pepsin-LC-MS/MS HDX. A stock solution consisted of 15 μM β 3 peptide or β 3 peptide conjugated to lipid in 25 mM phosphate and 100 mM NaCl, pH 6.8. H-to-D exchange was initiated by diluting 5 or 10 μL of peptide stock solution into 15 or 10 μL of deuterated buffer, respectively. The reaction mixtures were then incubated for 10 seconds to 1.4 hours, as were non-deuterated control samples. Peptides fragments were collected on a reverse phase trap column (4 μL bed volume, ExSAR) for 3 min. Peptide fragments were eluted from the trap and separated chromatographically on a reverse phase column (4 μL bed volume, ExSAR) using a linear HPLC gradient of 12% to 45% acetonitrile over 11 minutes (pH 2.5, 0°C) at a 6 μL/min flow rate. Peptides were ionized by ESI for mass spectrometric analysis in a Thermo Fisher OrbiTrap-LTQ.

Resulting mass spectra were analyzed using the DXMS software package (Sierra Analytics). The deuterium values per peptide were corrected for the deuterium content in the reaction conditions, as well as for deuterium loss in liquid handling by assuming an average of 75% deuterium recovery. Intrinsic exchange curves were generated using HXpep (92). Experimental curves were generated from the exchange rates generated by exponential fitting of intrinsic curves. Fitting of the experimental curves gives the slowing factor, which is a measure of the H-

to-D exchange protection when compared to a random coil structure.

Supplemental

Conformationally restricting distance constraints	
Intraresidue $[i = j]$	202
Sequential $[i - j = 1]$	259
Medium Range [1 < <i>i − j</i> < 5]	235
Long Range [<i>i – j</i> => 5]	16
Total	712
Dihedral angle constraints	
φ	42
Ψ	42
Hydrogen bond restraints	24
Average number of constraints per residue	15.2
Average number of distance constraints violations per	
Xplor-NIH conformer	
> 0.5 Å	0
Average number of dihedral-angle constraint violations	
Xplor-NIH conformer	
> 5 [°]	0
Average r.m.s.d. to the mean coordinates [Å]	
Residues 713-723 and 726-736, backbone heavy	0.68 ± 0.25
atoms	
Residues 713-723 and 726-736, all heavy atoms	1.56 ± 0.27
Residues 713-723,726-736 and 745-758 , backbone	4.63 ± 1.37
heavy atoms	
Residues 713-723,726-736 and 745-758, all heavy	5.27 ± 1.46
atoms	
PROCHECK(93) G-factors Z-score for all residues (ϕ and	2.08 / 1.89
ψ / all dihedral angles) ^a	
PROCHECK(93) G-factors Z-score for ordered residues (o	2.68 / 2.60
and ψ / all dihedral angles) ^b	
MOLPROBITY(94) clash Z-score	-1.18/-1.18
(all residues ^a / ordered residues ^b)	
Ramachandran plot summary [%] (all residues ^a / ordered	
residues ^b)	
most favored regions	90.8 / 94.5
Additionally allowed regions	7.8/ 5.2
generously allowed regions	1.1 / 0.3
disallowed regions	0.3 / 0.0

Table S2.2.1 Summary of β3 structure statistics

^a Residues 709 - 762 ^b Residues 711-735, 738-758

Chapter 3 The affinity of talin-1 for the β 3-integrin cytosolic domain is determined by its phospholipid bilayer environment.

Abstract

Binding of the talin-1 FERM domain to the β 3 cytosolic tail causes activation of the integrin α IIb β 3. The FERM domain also binds to acidic phospholipids. Although much is known about the interaction of talin-1 with integrins and lipids, the relative contribution of each interaction to integrin regulation and possible synergy between them remain to be clarified. Here, we examined the thermodynamic interplay between FERM domain binding to phospholipid bilayers and to its binding sites in the β 3 tail. We found that although both the F0F1 and F2F3 subdomains of the talin-1 FERM domain bind acidic bilayers, the full-length FERM domain bound with an affinity similar to F2F3, indicating that F0F1 does not significantly increase the affinity for the membrane. When free in solution, the β 3 tail has weak affinity for the FERM domain. However, appending the tail to acidic phospholipids increased its affinity for the FERM domain by three orders of magnitude. Nonetheless, the affinity of the FERM for the appended tail was similar to its affinity for binding to bilayers alone. Thus, talin-1 binding to the β 3 tail is a ternary interaction dominated by a favorable surface interaction with phospholipid bilayers and set by lipid composition. Nonetheless, interactions between the FERM domain, the β 3 tail, and lipid bilayers were not optimized for a high-affinity synergistic

interaction, even at the membrane surface. Instead, the interactions appeared to be tuned in such a way that the equilibrium between inactive and active integrin conformations can be readily regulated.

I. Introduction

The integrin α IIb β 3 resides on the platelet surface in equilibrium between resting and active conformations (5, 6, 95-97). On circulating platelets, α IIb β 3 is constrained in its inactive conformation to prevent spontaneous platelet aggregation. Similarly, the cytoskeletal protein talin-1, whose binding to the β 3 cytosolic tail (CT) stabilizes the active conformation of α IIb β 3 (98-100), is sequestered away from the β 3 CT (101). Besides interacting with integrins, talin-1 interacts with negativelycharged phospholipids (102-104) and phosphoinositides (31, 33, 105, 106). Stimuli generated at sites of vascular damage recruit talin-1 to the platelet plasma membrane, thereby promoting $\alpha IIb\beta 3$ activation (12, 19). Much is known about the interaction of talin-1 with integrins and lipids, but the relative contribution of each interaction to integrin regulation and possible synergy between them remain to be clarified. Elucidating these interactions is important for understanding events leading to and controlling the formation of integrin complexes and for potential pharmacologic modulation of integrin signaling.

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Talin-1 is a 250 kD protein containing a 45 kD N-terminal head domain attached via a flexible linker to a 200 kD C-terminal rod domain (99). Because the head domain is packed against the rod domain in its inactive state (107), talin-1 recruitment to the membrane and to integrin β CTs requires disruption of this interaction (108). At integrin-mediated cellular adhesions, the orientation of talin-1 is highly polarized, with its head domain associated with the plasma membrane and its rod domain making numerous cytoskeletal contacts (108).

The talin-1 head domain is a member of the FERM (4.1/Ezrin/Radixin/Moeisin) domain family (109, 110). Canonical FERM domains, such as that of radixin, consist of F1, F2, and F3 subdomains adopting ubiquitin, acyl-CoA-binding protein, and PTB/PH domain folds, respectively, and are assembled into a trefoil arrangement (Figure 3.1 A and B). The talin-1 FERM domain contains a fourth subdomain, F0, preceding F1 that also adopts a ubiquitin-like fold (111). The four talin-1 subdomains have an extended, nearly linear, configuration (PDB code 3ivf, Figure 3.1C) (104). Whereas radixin binds its targets, such as the CD44 CT, in a groove formed by helix $\alpha 1$ and strand $\beta 5$ of its F3 subdomain in an anti-parallel β - β fashion (Figure 3.1A), the talin- β integrin CT complex contains a second interface with the N-terminal helical portion of the CT (Figure 3.1D) (61, 112). This positions talin to modulate interaction between membrane-proximal cytosolic and transmembrane segments of integrin α and β subunits (1, 61, 112).



Figure 3.1 Comparison of the Talin-1 and Radixin FERM domains.

(A) The radixin FERM domain is composed of three subdomains: F1 (left, orange), F2 (center, yellow), and F3 (right, green). The CD44 cytoplasmic tail (blue) binds to a cleft in F3 formed by helix α 1 and strand β 5. (B) The radixin subdomains adopt a canonical trefoil shape. (C) The talin-1 FERM domain contains F1 (orange), F2 (yellow), and F3 (green) domains, as well as an F0 subdomain (red) that precedes F1. Instead of a trefoil, the talin-1 FERM subdomains are arrayed linearly. (D) Model of talin-1 F3 (green) bound to the integrin β 1D CT (black). Two regions of the β 1D CT interact with talin-1 F3: its C-terminal portion adopts an extended conformation and packs against a ridge formed by the interface of helix α 1 and strand β 5 in F3, while an N-terminal α -helix and a contiguous portion of the β 1 transmembrane (TM) domain packs against the top of F3. Radixin structures were generated from PDB 2zpy. Talin-1 structures were modeled using PDB 3g9w and PDB 3ivf.

Here, we examine the thermodynamic interplay between phospholipid bilayers and β CT binding sites in the talin-1 FERM domain. The impetus for this study arose, in part, from conflicting findings regarding the ability of the talin-1 FERM domain to interact with the β 3 CT. In solution, the affinity of talin-1 for β integrin CTs is nearly 1000-fold weaker than the affinity of radixin for CD44 (113) or of talin-1 for the Cterminus of PIP5KI- γ (61). However, talin-1 binds to the β 3 CT with high affinity when the CT is immobilized on a carboxy-methylated dextran surface (114). This suggests that the interaction of talin-1 with integrins is optimal when the β subunit CT is presented on a hydrophilic negativelycharged surface such as the inner leaflet of plasma membranes.

To assess the interplay between membrane and β CT binding interactions, we tested whether the talin-1 FERM subdomains F0F1 and F2F3 (Figure 3.2B and C), each of which can interact with membranes (111), bind cooperatively or independently to a variety of negatively charged lipids and compared their binding to the complete FERM domain (F0-F3, Figure 3.2A). We then measured the effect of the membrane environment on the affinity of FERM domain binding to the β 3 CT. We found that although the membrane environment increased the overall affinity of the FERM domain for β 3, the interactions between the FERM domain, the β 3 CT, and lipid bilayers were not optimized for a high-affinity synergistic interaction, even at the membrane surface. Instead, the

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interactions appeared to be tuned in such a way that the equilibrium between inactive and active integrin conformations can be readily regulated.



Talin-1 F0F1F2F3: LVPR -|- GSHM-V2ALSLKISI....DIILKKKKS425

Thrombin/MCS



С



Talin-1 F0F1: LVPR -|- GSHM-V²ALSLKISI....LRRKFFYSD²⁰¹ Thrombin/MCS



Talin-1 F2F3: LVPR -|- GSHM-S206RDPVQLNL....DIILKKKKS425

D

BODIPY-β3: LVPR -|- GSHMG-C⁷¹⁹-TIH**DRKEFAKFEEERARAKWDTANNPLY**KEATSTFTNITYRGT⁷⁶² Maleimide-BODIPY-TMR

LIPID-β3: LVPR -|- GSHMG-C⁷¹⁹-TIH**DRKEFAKFEEERARAKWDTANNPLY**KEATSTFTNITYRGT⁷⁶² Maleimide-PE (C16)₂

Ε

FITC-PIP5K1γ: FITC-βAla-S646WVYSPLHY654

Figure 3.2 Talin-1 and α IIb β 3 constructs used.

Talin-1 constructs encompassing the (A) F0, F1, F2, and F3 subdomains, (B) the F0 and F1 subdomains, and (C) the F2 and F3 subdomains. Each construct contains an N-terminal Gly-Ser-His-Met (GSHM) motif enabling cleavage by thrombin. (D) β 3 CT constructs include the entire CT including the talin-1 recognition motif (bold) and preceded by a Cys residue at position 719 as well as an N-terminal GSHMG sequence for cleavage by thrombin (top); a fluorescent β 3 CT construct in which a maleimide-BODIPY-TMR moiety is appended to C719 (middle); and a β 3 CT construct in Cys719 was covalently modified with maleimidefunctionalized phosphatidylethanolamine (bottom). (E) FITC-labeled peptide derived from PIP5Kly. *{All talin and integrin constructs cloned, expressed, and purified by DTM. PIP5K1y was synthesized by HJ.}*

II. Results

The talin-1 FERM domain contains an extended positivelycharged surface.

Projection of an electrostatic surface on the talin-1 FERM domain revealed that it contains an extended positively-charged surface (Figure 3.3) (104). This is especially pronounced for the F2 and F3 subdomains. The F0 and F1 subdomains also contain positively charged surfaces and F1 contains a flexible loop that readily binds to membranes but was removed for crystallization (107). The positively-charged surfaces on F0F1 are not completely aligned with those on F2F3. Although this could reflect a strong bias for maintaining the surfaces in an unaligned state in solution or the selection of a single conformation from a flexible ensemble due to crystal packing, X-ray scattering studies argue against a flexible ensemble of structures (104). Thus, if the solution conformation is maintained, F0F1 should make only modest contributions toward the affinity of the overall complex over that of the F2F3 alone. Alternatively, if F0F1 and F2F3 change their mutual orientation away from the minimum energy solution conformation, this would attenuate the affinity relative to the value expected if they were pre-organized to interact favorably with the surface.

А

 \bigcirc F0 F1 F2 F3 F3 F2 F1 F0 В F0 F1 F2 F3 F3 F2 F1 F0 С F0 F1 F2 F3 F2 F3 F1 FO



As in Figure 3.1, the talin-1 F0-F3 subunit bound to a β -integrin was modeled using the crystal structure of the talin-1 FERM domain (pdb code 3ivf). (A) The interaction of talin-1 with the β integrin CT (black) places its FERM domain in proximity to the inner plasma membrane leaflet. (B)
Projection of an electrostatic surface onto the talin-1 FERM domain. Red denotes negative and blue denotes positive-charge density. (C) Talin-1 FERM domain rotated 90° showing the electrostatic surface as viewed from the membrane (left) and cytosol (right). *{Models were generated by DTM using PyMOL}*

Binding of the talin-1 FERM domain fragment F2F3 to negatively-charged phospholipids.

Before studying the interaction of the talin-1 FERM domain with the β 3 CT tethered to membrane, we investigated whether the F2F3 subdomain, which makes the most direct contact with the β 3 CT, differentially recognizes negatively-charged phospholipids. We used surface plasmon resonance (SPR) to measure the interaction of F2F3 with phospholipid bilayers immobilized on a hydrophobically-modified Biacore L1 SPR chip (115). All measurements were made relative to a surface containing 100% phosphatidylcholine (PC). F2F3 binding to 20% POPS and 80% POPC was measured at steady state (Figure 3.4A) and conformed to a single-site binding model as a function of F2F3 concentration. The dissociation constant (K_d) for F2F3 binding to 20% phosphatidylserine (PS)-containing bilayers was 0.92 μM (Table 3.1). In comparison, the K_d for bilayers containing 10% PtdIns(4,5) P_2 was 0.67 μ M, suggesting that the F2F3 may have a slight preference for PtdIns $(4,5)P_2$ over PS. Nonetheless, F2F3 bound to bilayers containing the diphosphoinositides $PtdIns(3,5)P_2$ and $PtdIns(3,4)P_2$ and the monophosphoinositide PtdIns(5)P with affinities similar to PtdIns(4,5)P₂ and largely within the error of measurements (Figure 3.4B, Table 3.1), indicating that it does not strongly differentiate between the negatively charged head groups.







Figure 3.4 Steady state binding of the talin-1 FERM domain and its subdomains to phospholipid bilayers.

(A) SPR sensorgrams of F2F3 binding to a bilayer composed of 20% POPS and 80% POPC immobilized on a Biacore L1 SPR chip. F2F3 was injected in 2-fold increments up to a concentration of 10 μ M. (B) Steady-state isotherms for F2F3 binding to negatively-charged phospholipids. Diamonds, 10% PtdIns(4,5)*P*₂; squares, 10% PtdIns(3,5)*P*₂; triangles, 10% PtdIns(3,4)*P*₂, Xs, PtdIns(5)*P*, circles, 20% phosphatidylserine. (C) Steady-state isotherms for F0-F3 (squares), F2F3 (triangles), and F0F1 (diamonds) binding to bilayers containing 10% PtdIns(4,5)*P*₂. Data from (B) and (C) were fit to a one-site binding model using BIAevaluation software. {*Three independent experiments were performed. Protein was purified by DTM. SPR was performed by PN. Data analysis was performed by DTM and PN.*}

Table 3.1 Steady-state SPR measurements of talin-1 F2F3 subdomain binding to phospholipids.

Lipid (%)	K _d (μM)	Maximum Binding (RU)
PS (20%)	0.92 ± 0.260	1210 ± 66
PtdIns(4,5)P ₂ (10%)	0.67 ± 0.07	2490 ± 225
PtdIns(3,4)P ₂ (10%)	0.42 ± 0.09	2170 ± 146
PtdIns(3,5)P ₂ (10%)	0.50 ± 0.053	2330 ± 193
PtdIns(5)P (10%)	0.37 ± 0.029	1770 ± 146

Binding to the indicated phospholipids (Figure 3.4B) was fit to a one-site binding model using BIAevaluation. The data represent the mean and standard deviations of three independent experiments.

The total amount of F2F3 bound to bilayers depended on the net negative charge of the membrane, suggesting that electrostatics play a substantial role in the interaction. PtdIns(4,5) P_2 , PtdIns(3,5) P_2 , and PtdIns(3,4) P_2 each have a charge of (-4) per head group, while PtdIns(5)P

has a charge of (-3), and PS has a charge of (-1). If the interaction were dependent of charge, the amount of protein bound to 10% PtdIns(5)*P*, would be expected to be roughly $\frac{3}{4}$ the amount bound to 10% diphosphoinositides. Similarly, $\frac{1}{2}$ as much talin-1 would be expected to bind to 20% PS as to 10% diphosphoinositides. The maximum resonance units (RU) of F2F3 bound to 10% PtdIns(4,5)*P*₂, 10% PtdIns(3,5)*P*₂, and 10% PtdIns(3,4)*P*₂ were similar at 2490, 2330, and 2170 RU, respectively. We found that approximately $\frac{3}{4}$ the amount of F2F3 bound to 10% PtdIns(5)*P* (1770 RU) and $\frac{1}{2}$ the amount bound to 20% PS (1200 RU), confirming that total protein bound depends on the charge of the membrane.

Binding of talin-1 FERM domain and its subdomains to $PtdIns(4,5)P_2$.

Next, we compared the affinity of the complete talin-1 FERM domain (F0-F3) for PtdIns(4,5) P_2 with that of F0F1 and F2F3. PtdIns(4,5) P_2 was chosen as the lipid substrate for these measurements because F2F3 did not differentiate between various phosphoinositides. Phospholipid bilayers containing 10% PtdIns(4,5) P_2 were immobilized on a Biacore L1 chip and all measurements were made relative to a surface containing 100% PC. We found that F0-F3 bound to 10% PtdIns(4,5) P_2 with a K_d of 980 nM, comparable to the K_d for F2F3 of 670 nM, whereas the K_d for F0F1 was 2.6 μ M (Table 3.2). Thus, both F0F1 and F2F3 readily interact with PtdIns(4,5) P_2 , but when combined, the affinity for

bilayers was not enhanced. The relative stoichiometries of F0F1, F2F3, and F0-F3 also suggest that the subdomains do not form a cooperative membrane binding surface. The mass ratios for F0-F3, F2F3, and F0F1 are 2:1.1:1.0. F0-F3 has twice the mass of F2F3 and nearly twice the resonance units of F0-F3 (3960 RU) bound to the sensor chip as F2F3 (2490 RU) (Figure 3.4C, Table 3.2). Thus, F0-F3 and F2F3 have similar stoichiometries for the phospholipid surface. A similar amount of F0F1 bound to the sensor chip (2290 RU) as F2F3, but with lower affinity. Thus, although F0F1 can interact with PtdIns(4,5) P_2 , the affinity and stoichiometry of the full FERM domain for PtdIns(4,5) P_2 is similar to that of F2F3, indicating that F0F1 does not substantially cooperate with F2F3 in FERM domain binding to membranes.

Table 3.2 Steady-state SPR measurements of talin FERM domain binding to phospholipid bilayers containing 10% PtdIns(4,5)P₂.

Binding results were fit to a one-site binding model using BIAevaluation. The data shown represent the mean and standard deviation of three independent experiments.

FERM construct	domain	K _d (μΜ)	Maximum (RU)	Binding
F0-F3		0.98 ± 0.10	3960 ± 258	
F2F3		0.67 ± 0.07	2490 ± 220	
F0F1		2.63 ± 0.84	2290 ± 258	

Enthalpy of talin-1 FERM domain binding to PtdIns(4,5)₂.

It has been reported that the talin-1 FERM domain bind to negatively-charged vesicles in solution (104, 105, 111, 116, 117). To confirm these observations, we used isothermal titration calorimetry (ITC) to measure the enthalpy of F0-F3 binding to large unilamellar vesicles (LUVs) composed of 90% POPC and 10% PtdIns(4,5) P_2 . As a positive control, we measured binding of the pleckstrin homology (PH) domain of PLC- δ 1 to PtdIns(4,5) P_2 (30).

Titration of 100 μ M PLC- δ 1 PH domain into a LUV suspension containing 1%, 3%, and 10% PtdIns(4,5) P_2 generated binding isotherms with molar enthalpies of -11.7 kcal/mol, -10.2 kcal/mol, and -12.8 kcal/mol and dissociation constants of 2.8 μ M, 2.1 μ M, and 4.2 μ M, respectively (Figure 3.5A, Table 3.3). These results are comparable to a previously published binding enthalpy of -11.9 kcal/mol and K_d of 1.1 μ M for PLC- δ 1 PH domain binding to 1% PtdIns(4,5) P_2 bilayers (30).



Figure 3.5 Comparison of PLC- δ 1 PH domain and talin-1 FERM domain binding to PtdIns(4,5) P_2 and Ins(1,4,5) P_2 .

(A) ITC titration of 100 μ M PLC- δ 1 PH domain into LUVs containing 1% (X), 3% (open squares), and 10% (closed squares) PtdIns(4,5) P_2 . The titrations were globally fit to a OneSite binding model using MicroCal/Origin. (B) ITC titration of 190 µM talin-1 FERM domain (F0-F3) into LUVs containing 1% (X), 3% (open squares), and 10% (closed squares) PtdIns(4,5) P_2 No enthalpy was generated for 1% or 3% LUVs. Titration into 10% PtdIns(4,5) P_2 was globally fit to a OneSite binding model using MicroCal/Origin. (C) Steady-state binding isotherms for F0-F3 binding to 1% (triangles), 3% (squares) and 10% (diamonds) PtdIns $(4,5)P_2$ immobilized on a Biacore L1 SPR chip. {PLC- δ 1 ITC titrations were performed once to demonstrate accessibility of the $PtdIns(4,5)P_2$ headgroup. Talin-1 titration into 1% and 3% PtdIns(4,5)P₂ were performed in triplicate independent experiments. Talin-1 titration into 10% PtdIns(4,5) P_2 was performed in triplicate with the same protein sample. SPR data are representative of three independent experiments. ITC experiments were performed by DTM. DTM prepared protein samples for SPR experiments. PN performed SPR experiments. DTM and PN analyzed SPR data.}

Table 3.3 ITC measurements of talin-1 FERM domain and PLC- δ 1 binding to PtdIns(4,5)P₂ and Ins(1,4,5)P₃.

Measurements were performed at 20°C in 25 mM MOPS pH 6.8 with 100 mM NaCl. Background heats of dilution were subtracted. Titrations were fit to OneSite binding isotherms using MicroCal/Origin and errors are reported as fitting errors. There was no net enthalpy for titration of F0-F3 into 1% and 3% PtdIns(4,5)P₂ and 2 mM Ins(1,4,5)P3. {*PLC-*δ1 *ITC experiments were performed once to demonstrate accessibility of PtdIns*(4,5)P₂ *on the bilayer surface – errors represent fitting errors. Talin-1 titration into* 1% *and* 3% *PtdIns*(4,5)P₂ *were performed in triplicate independent experiments. Talin-1 titration into* 10% *PtdIns*(4,5)P₂ *was performed in triplicate with the same protein sample. Talin-1 errors are standard deviations.* }

	K _a (x10 ⁶ M ⁻¹)	K _d (μM)	$\Delta \mathbf{H}$ (kcal/mol)	$\Delta \mathbf{S}$ (cal/mol/K)	n
F0-F3 + 1% PtdIns(4,5) <i>P</i> ₂					
F0-F3 + 3% PtdIns(4,5) <i>P</i> ₂					

F0-F3 + 10% PtdIns(4,5) <i>P</i> ₂	1.3 ± 0.6	0.781	-3.3 ± 0.4	-17.0	0.33 ± 0.03
F0-F3 + 2 mM Ins(1,4,5 <i>)P</i> ₃					
PLC- δ 1 + 1% PtdIns(4,5) P_2	0.36 ± 0.08	2.82	-11.7 ± 1.0	-14.0	1.13 ± 0.07
PLC- δ 1 + 3% PtdIns(4,5) P_2	0.47 ± 0.03	2.11	-10.2 ± 0.2	-8.37	0.97 ± 0.01
PLC- δ 1 +10% PtdIns(4,5) P_2	0.24 ± .02	4.20	-12.8 ± 0.7	-18.3	0.75 ± 0.03
PLC-δ1 + 2 mM Ins(1,4,5) P_3	1.4 ± 0.2	0.72	-17.5 ± 0.7	-30.5	0.80 ± 0.02

Next we titrated 190 μ M F0-F3 into a suspension of LUVs composed of 90% POPC and 10% PtdIns(4,5) P_2 . After subtracting the heat of dilution, F0-F3 binding was slightly exothermic (\approx -3 kcal/mol) with a stoichiometry of \approx 1 F0-F3 to 3 PtdIns(4,5) P_2 phospholipids (Figure 3.5B). Titration of higher F0-F3 concentrations resulted in precipitation of the mixture in the calorimetry cell – confirming the interaction of F0-F3 with membranes, but precluding thermodynamic analysis. Surprisingly, similar experiments using LUVs containing 1% and 3% PtdIns(4,5) P_2 did not generate measurable enthalpy (Figure 3.5B). Therefore, to confirm that F0-F3 actually binds to bilayers containing 1% and 3% PtdIns(4,5) P_2 , we measured the interaction by SPR and found that it bound with high affinity (Figure 3.5C, Table 3.4). Furthermore, the amount of F0-F3 bound to the chip scaled with the amount of PtdIns(4,5) P_2 on the surface. The Rmax for 10% PtdIns(4,5) P_2 was approximately 10 fold that for 1% PtdIns(4,5) P_2 . The Rmax ratio of 10% to 3% PtdIns(4,5) P_2 was 4.4, which was greater than the expected 3.3. All other SPR measurements with F0F1, F2F3, and F0-F3 have Rmax values that scale appropriately with phospholipid charge and protein mass (Tables 3.1 and 3.2), therefore it is unclear why the 3% PtdIns(4,5) P_2 surface would deviate. The experiment was repeated several times with different protein preparations and lipid stocks.

Table 3.4 Steady-state SPR measurements of talin F0-F3 domain binding to phospholipid bilayers containing 10%, 3%, and 1% PtdIns(4,5) P_2 .

Binding results were fit to a one-site binding model using BIAevaluation. The data shown represent the mean and standard deviation of three independent experiments.

Negatively Charged Phospholipid	K _d (μΜ)	Maximum Binding (RU)
10% PtdIns(4,5)P ₂	0.98 ± 0.10	3960 ± 258
3% PtdIns(4,5)P ₂	0.70 ± 0.21	901 ± 57
1% PtdIns(4,5)P ₂	0.65 ± 0.28	427 ± 24

Lastly, we attempted to measure the interaction of F0-F3 with the inositol headgroup of PtdIns(4,5) P_2 , Ins(1,4,5) P_3 , free in solution (Figure 3.6). Two mM Ins(1,4,5) P_3 was titrated into a 150 μ M F0-F3 solution, but

no binding enthalpy was generated. By contrast, titration of 120 μ M Ins(1,4,5) P_3 into a 6 μ M solution of the PLC- δ 1 PH domain generated an exothermic binding curve with thermodynamic parameters comparable to previously published results (Table 3.3) (30).



Figure 3.6 Titration of $lns(1,4,5)P_3$ into talin-1 F0-F3 does not generate significant enthalpy.

ITC titrations of 2 mM Ins $(1,4,5)P_3$ into 150 µM F0-F3 (filled squares) and 120 mM Ins $(1,4,5)P_3$ into 6.0 mM PLC- δ 1 (open squares). The titration into PLC- δ 1 could be fit to a OneSite model (MicroCal/Origin). All samples were prepared in buffer matched 25 mM MOPS 100 mM NaCl, pH 6.8. Background heats of dilution were subtracted. {*Titration of Ins* $(1,4,5)P_3$ *into F0-F3 is representative of two independent experiments. Titration of Ins* $(1,4,5)P_3$ *into PLC-\delta1 was performed once to demonstrate activity of Ins* $(1,4,5)P_3$. Experiments performed by DTM.}

Talin-1 FERM domain binding to the β 3 CT immobilized on negatively charged carboxymethylated dextran.

Previous measurements of talin-1 FERM domain binding to the β 3 CT used a β 3 construct beginning at Lys716 and included a linker for attaching the construct to an SPR chip (114). Our β 3 CT construct, designed to mimic the β 3 CT as it exits the membrane, was attached via Cys719 to a CM-5 SPR sensor chip functionalized with N-[Emaleimidocaproic acid]-hydrazide to generate thiol-reactive groups (Figure 3.2). F0-F3 was then flowed over the chip at concentrations up to 1.0 μ M. The resulting sensorgrams (Figure 3.7A) could be fit to two binding events with dissociation constants of 155 nM and 3.5 μ M. The biphasic behavior could have resulted from heterogeneity of \$3 on the chip surface or F0-F3 interaction between heterogeneity of the and the carboxymethylated dextran (118). Regardless, the interaction between the talin-1 FERM domain and the β 3 CT may be sensitive to such heterogeneity since we hypothesize that the FERM domain binds to the β 3 CT and the inner membrane surface simultaneously. We chose not to pursue experiments with carboxy-dextran-immobilized integrin further, as membrane immobilized β 3 is more biologically relevant and immobilization of small peptides on dextran matrices can result in heterogenous interaction surfaces.



Figure 3.7. Talin-1 FERM domain binding to the β 3 CT immobilized on negatively-charged surfaces or free in solution.

(A) The β 3 CT (600 RU) was immobilized on a carboxy-methylated dextran SPR chip and F0-F3 to maximum concentration of 1 μ M was flowed over the chip. (B) FP of F0-F3 titrated into 20 nM BODIPY-TMR β 3. FP was measured at 575 nm after excitation at 530 nm (squares). F0-

F3 was also titrated into a 20 nM FITC-labeled PIP5KI- γ (black circles). FP was measured at 528 nm after excitation at 485 nm. (C) ITC of 140 μ M β 3 CT titrated into buffer-matched 10.5 μ M F0-F3 revealed no significant enthalpy above the heat of dilution. ITC samples were in 25 mM MOPS 100 mM NaCl, pH 6.8. {*SPR data are representative of two independent experiments. Fluorescence polarization experiments were performed in triplicate using the same protein sample. ITC data are representative of three independent experiments – no binding enthalpy was generated in all three experiments. SPR was performed and analyzed by PN using protein prepared by DTM. FP performed by DTM. PIP5K1\gamma peptide was prepared by HJ. Integrin peptide was prepared by DTM. ITC performed by DTM.}*

Binding of the talin-1 FERM domain to the β 3 CT domain tethered to negatively-charged bilayer.

To address whether the talin-1 FERM domain binds the β 3 CT and a negatively-charged surface in a concerted fashion, we first used fluorescence polarization (FP) and ITC to measure the affinity of talin-1 FERM domain for the β 3 CT when both are free in solution.

For FP, the β 3 CT was labeled with maleimide-functionalized BODIPY-TMR at Cys719. When F0-F3, at concentrations up to 100 μ M, was titrated into a 20 nM solution of labeled β 3 CT, the FP increased from 0.1 to 0.16. (Figure 3.7B). These results were fit to a binding isotherm with a K_d of approximately 600 μ M. Since the interaction between talin-1 and β 3 CT was not saturated at the concentrations used, the exact dissociation constant could not be calculated, however other laboratories have reported values on the order of several hundred micromolar (62, 112), which is in line with our results. As a positive control for talin-1 activity, we titrated F0-F3 into a 20 nM solution of a fluorescein isothiocyanate (FITC)-labeled peptide corresponding to the C-terminus of PIP5KI γ (Figure 3.7B). The resulting change in FP was fit to an isotherm with a K_d of 160 nM ± 50 nM, consistent with reported affinities of 170 nM to 1.4 μ M for other PIP5KI γ constructs with slightly different sequences (119, 120).

We also used ITC to confirm that the β 3 CT and F0-F3 does not interact strongly in solution. There was no significant enthalpy above background heats of dilution when F0-F3, up to a final concentration of \approx 30 μ M protein, was titrated into 10 μ M peptide (Figure 3.7C). This contrasts with a recent report in which talin-2 and the β 1D cytoplasmic domain interacted with a K_d of \approx 20 μ M and a molar enthalpy of \approx -17 kcal/mol (112). This difference is associated with several subtle but significant differences between the β 1D CT and β 3 CT sequences. Our result can be interpreted either as a lack of interaction between talin-1 and the β 3 CT at the concentrations used or a lack of significant enthalpy at the concentrations used. Given that the structural similarity of the talin-2 / β 1D interaction (Δ H \approx -17 kcal/mol), we would expect the β 3 interaction to have measurable enthalpy at the concentrations used in this experiment. Furthermore, the lack of observed enthalpy corroborates our FP data and previously published NMR experiments (62, 112).



Figure 3.8. Talin-1 FERM domain binding to the β 3 CT conjugated to phospholipid bilayers containing 1% PtdIns(4,5) P_2 .

The β 3 CT at a final concentration of 5 μ M was immobilized onto the surface of LUVs containing 1% PtdIns(4,5) P_2 (20 μ M)/ 2% maleimide-PE (40 μ M) / 97% POPC (2 mM total lipid). Buffer matched talin-1 FERM domain (F0-F3, 150 μ M) was titrated at 20°C into the suspension of immobilized β 3. Background heats of dilution were subtracted. The titration was globally fit to a OneSite binding model using MicroCal/Origin. All ITC experiments were performed in 25 mM MOPS buffer 100 mM NaCl, pH 6.8. Thermodynamic parameters are summarized in Table 3.5. {Data represent three independent experiments. Experiments performed by DTM.}

Because the interaction between the talin-1 FERM domain and PtdIns(4,5) P_2 occurs with very low enthalpy, we were able to use ITC to

measure talin-1 FERM domain binding to the β 3 CT in the presence of PtdIns(4,5) P_2 . Accordingly, we immobilized the β 3 CT on phospholipid bilayers by reacting a β3 CT construct containing Cys at residue 719 with maleimide-functionalized phosphatidylethanolamine on the surface of LUVs composed of POPC and 1% PtdIns(4,5)P₂, 20% PS, or 20% PS + 1% PtdIns(4,5) P_2 (1). Titration of F0-F3 into a suspension of LUVs containing 1% PtdIns(4,5) P_2 in the absence of immobilized β 3 CT (Figure 3.5B) or into buffer containing the free β 3 CT (Figure 3.4C) generated little or no enthalpy. By contrast, titration of 150 µM F0-F3 into a LUV suspension containing 5 μ M β 3 CT immobilized on 1% PtdIns(4,5) P_2 was exothermic (Figure 3.8). The titration was globally fit to a one-site binding isotherm, revealing a K_d of 860 nM and a molar enthalpy of -25.0 kcal/mol (Table 3.5). When the experiments were repeated with the β 3 CT immobilized on bilayers containing 20% PS or 20% PS with 1% Ptdlns(4,5) P_2 similar results were obtained (Figure 3.9A/B; Table 3.5). Thus, these experiments demonstrate that immobilizing the β 3 CT on phospholipid bilayers increases its affinity for the talin-1 FERM domain by at least three orders of magnitude compared to the β 3 CT free in solution, increasing its affinity from that of a very weak to a high-affinity interaction.



Figure 3.9 FERM domain binding to the β 3 CT conjugated to negatively-charged phospholipid bilayers.

As in Figure 3.5k, the talin-1 FERM domain (F0-F3, 150 μ M) was titrated at 20°C into a buffer-matched suspension of LUVs composed of (A) 20% PS or (B) 20% PS + 1% PtdIns(4,5) P_2 to which 5 μ M β 3 CT had been conjugated via a maleimide-functionalized lipid head group (Methods). The titrations were globally fit to a OneSite binding model using MicroCal/Origin. Thermodynamic parameters are summarized in Table 3.5. {Data are representative of triplicate experiments performed with the same protein sample. Experiments were performed by DTM.}

Table 3.5 ITC measurements of talin-1 FERM domain binding to the β 3 CT attached to large unilamellar phospholipid vesicles (LUVs).

The talin-1 FERM domain was titrated into LUV suspensions containing 5 μ M immobilized β 3 CT (Figures 3.8 and 3.9). All experiments were performed at 20°C in 25 mM MOPS pH 6.8 with 100 mM NaCl and background heats of dilution were subtracted. Titrations were fit to one-site binding isotherms. {*Titration into \beta3 on 1% PtdIns(4,5)P*₂ was performed as three independent experiments and errors are reported as standard deviations. Titration into β 3 on 20% PS and 20% PS with 1% PtdIns(4,5)P₂ was performed once with each sample and the error is reported as the fitting error. All experiments were performed by DTM.}

	К _а (х10 ⁶ М ⁻¹)	K _d (μΜ)	∆H(kcal/mol)	ΔS (cal/mol/K)	n	
β3- 1%PtdIns(4,5) <i>P</i> ₂	1.19 ± 0.21	0.86	-25.0 ± 1.5	-55.8	0.80 0.04	±
β3-20% PS	0.29 ± 0.02	3.4	-32.3 ± 0.9	-83.3	0.8 (held)	
β3-20%PS + 1%PtdIns(4,5) <i>P</i> ₂	0.61 ± 0.11	1.6	-34.0 ± 2.8	-87.5	0.95 .06	±

III. Discussion

Here we found that presentation of the β 3 CT in the context of an acidic bilayer enhances its affinity for the talin-1 FERM domain up to three orders of magnitude, depending on the phospholipid composition of the bilayer. However, FERM domain binding to the β 3 CT contrasts with the stronger interaction of the FERM domain with PIP5K1 γ and with the interaction of the talin-2 FERM domain and the β 1D CT. Small changes in integrin β CT sequences have been shown to significantly increase their affinities for talin family FERM domains (112). This suggests that the interaction between the talin-1 FERM domain and the β 3 CT has been evolutionarily tuned to be intrinsically weak, allowing it to be modulated by changes in phospholipid composition, post-translational modifications, and additional protein-protein interactions.

To understand how these protein-protein and lipid-protein interactions modulate affinity, it is important to consider the various energetic components associated with the interaction of the talin-1 FERM domain and the β 3 CT with lipids, as well as their interaction with each other. Previous studies reported the interaction of the talin-1 FERM domain and the β 3 CT with micelles and phospholipid vesicles (1, 16, 38, 54, 106). Here, we studied this interaction using the β 3 CT tethered to phospholipid bilayers to understand the energetics of the overall process. Previously, we used this construct to study the dynamics of the β 3 CT in the presence of bilayers by hydrogen-deuterium exchange (HDX). Although the construct lacks a transmembrane domain, it was designed computationally to attach via a residue predicted to be at the membrane interface and it contains all of the β 3 residues necessary for interaction with talin-1. While it is possible that the TM domain could affect this interaction, our NMR and HDX studies indicate that the TM helix is broken by a kink as it exits the membrane, implying that the structure of the β 3 CT distal to the helix disruption is unlikely to depend on residues buried within the membrane. Furthermore, our construct contains all of the β 3 residues necessary for formation of the kink (Figure 2.5).

We can consider the overall free energy for the interactions of the bilayer-tethered β 3 CT (RTIn(K_{obs}) = Δ G_{overall}) in terms of several components according to Eq. 1 below. It should be noted that this thermodynamic decomposition is path-independent and does not require kinetic ordering; nevertheless, all interactions here were rapid, on the

second to msec time scale, indicating that they might rapidly interconvert on a physiologically relevant time scale.

Eq. 1: $\Delta G_{\text{overall}} = \Delta G_{\text{talin-bilayer}} + \Delta G_{\text{talin-peptide}} + \Delta G_{\text{reorganization}}$

where:

 $\Delta G_{talin-bilayer}$ is the energy associated with talin-1 head interaction with lipid, as determined directly from the titration of the FERM domains into lipid bilayers;

 $\Delta G_{talin-peptide}$ is the energy associated with talin-1 head interaction with the β 3 CT, as determined directly from the NMR or the fluorescent titrations reported above and which is sequence-specific (112) but only weakly favorable (K_{diss} on the order of 0.2 to 0.6 mM).

 $\Delta G_{reorganization}$ is the free energy required to rearrange the conformations of isolated individual components and binary complexes to allow formation of a ternary complex.

The electrostatic surfaces of F3, F2F3, and the full FERM domain (44, 104) suggest that F2F3 binds negatively-charged bilayers in a manner highly conducive to forming a simultaneous interaction with the β 3 CT; this finding has been supported by detailed coarse-grained molecular dynamics calculations (106). Thus, there is likely little reorganization energy associated with the interaction of these subdomains with bilayers.

By contrast, the β 3 CT, tethered to a bilayer, is in a rapid equilibrium that favors a conformation in which the peptide interacts with the membrane in a way that masks residues important for talin-1 binding and that would otherwise be located 10 – 30 Å into the cytoplasm (1). A second minor conformation is more exposed to the aqueous surroundings, presumably allowing rapid kinetic access to the FERM domain. Hydrogen-deuterium exchange showed that the equilibrium favors the membrane-associated state by 100 to 1000-fold relative to the exposed state (Chapter 2), decreasing the overall affinity for binding cytosolic proteins by an equivalent amount. Thus, we expect that $\Delta G_{rearrange}$ would be dominated by the rearrangement of the β 3 CT within the bilayer.

Consideration of Eq. 1 shows that cytosolic protein binding to the β 3 CT in a proper context depends on an energetic balance, which, in this case, includes a highly favorable interaction between talin-1 and the membrane. A modestly favorable free energy of interaction of the β 3 CT with the talin-1 head is balanced by a modestly unfavorable free energy of rearrangement of the β 3 CT within the bilayer. In net, the observed affinity for formation of the ternary complex remains similar to that for the binary interaction of the talin-1 head with acidic bilayers. Likewise, in the absence of integrin, the binary complex between the FERM domain and the membrane appears to energetically favor the F2F3 subunit, as F0F1 does not affect the overall affinity or stoichiometry of the interaction. This suggests that F0F1 might not interact favorably with the bilayer once it is

tethered in place by the more stable F2F3 interaction. Alternatively, F0F1 might interact extensively when the FERM domain binds, but only after an energetically uphill conformational change that attenuates the overall affinity gain associated with F0F1-bilayer interaction. It is likely not accidental that the binary and ternary complexes are not optimized for cooperativity. Rather they appear to be tuned to be readily regulated by changes in the lipid environment and by post-translational modifications.

In summary, our studies reveal the rich interplay between structure, dynamics, kinetics, and thermodynamics that govern formation of the ternary complex between the β 3 CT, phospholipid bilayers, and talin-1 FERM domain. This energetic dissection provides the basis for understanding how posttranslational modifications and changes in lipid composition might affect such processes. The situation becomes more complex as interactions with additional domains in integrin and talin, as well as other antagonistic or synergistic cytoplasmic proteins, are considered. Nevertheless, the present study illustrates the importance of considering the role of conformational changes, structural rearrangements, and conformational entropy in these processes, and provides a general approach to energetic decomposition that can form the basis for understanding regulation of even more complex systems.

IV. Methods

Design of recombinant talin-1, β 3 CT and PIP5K1 γ constructs. Talin-1, β 3 CT, and PIP5K1 γ constructs were cloned between the Ndel and Xhol restriction sites of the vector pet28a and contained an N-terminal 6xHis tag for purification, followed by a thrombin cleavage site for affinity tag removal. The talin-1 constructs (Figure 3.2, A-C) consisted of the entire talin-1 FERM domain (F0-F3), extending from its N-terminus through residue Ser425, the F0F1 subdomains extending from the N-terminus through residue Asp201, and the F2F3 subdomains spanning residues Ser206 through Ser425.

The β 3 CT construct (Figure 3.2D), encompassing residues Thr720 through the C-terminus at Thr762, was previously described (1). For the current experiments, residue IIe719 was replaced with Cys to enable the construct to be immobilized on the surface of bilayers or to be labeled via maleimide chemistry. A peptide corresponding to residues 646-654 of PIP5K1 γ was also labeled with fluorescein isothiocyanate (FITC) (Figure 3.2E).

Talin-1 and PIP5K1γ were introduced into BL21-DE3 cells (Novagen) and expressed for 20 h at 20°C in the presence of 0.5 mM IPTG. Cells were then lysed by sonication on ice in 20 mM imidazole buffer, pH 7.0, containing 500 mM NaCl, 1 mM EDTA, and a protease inhibitor cocktail (Sigma-Aldrich). After the lysates were cleared by

centrifugation, the supernatants were loaded onto HiTrap Ni-NTA columns (GE Healthcare) and peptides were eluted using an imidazole gradient. Imidazole was dialyzed away overnight into 50 mM Tris buffer, pH 7.5 containing 100 mM NaCl and the 6xHis tag was removed using thrombin. The constructs were then purified by SP ion exchange chromatography (GE Healthcare) and finally size exclusion chromatography with a Superdex 75 preparative column (GE Healthcare).

Preparation of large unilamellar vesicles (LUVs). LUVs were generated by evaporating a mixture of 1-palmitoyl-2-oleoyl-*sn*-glycero-3-phosphocholine (POPC, Avanti Polar Lipids) and 1-palmitoyl-2-oleoyl-*sn*-glycero-3-phospho-L serine (POPS, Avanti Polar Lipids) dissolved were dissolved in chloroform under nitrogen; excess chloroform was removed under high vacuum overnight. Phosphoinositides (Cell Signaling) were dissolved in CHCl₃/MeOH/0.1% HCl, mixed with POPC, and dried under nitrogen. Evaporated lipids were then re-suspended in buffer, vortexed vigorously for 5 minutes, subjected to 3 x freeze- thaw cycles, and extruded 20 to 30 times using 100 nm diameter filters (Avanti Polar Lipids).

Preparation of membrane tethered β **3.** Lyophilized β **3** CT was redissolved in bilayer conjugation buffer (25mM sodium phosphate or MOPS, 100 mM NaCl, pH 6.8) at a concentration of 50 μ M. Large unilamelar vesicles (LUVs) were generated by evaporating a mixture of

1,2-dipalmitoyl-sn-glycero-3-phosphoethanolamine-N-[4-(p-

maleimidomethyl)cyclohexane-carboxamide] (16:0 PE MCC, Avanti Polar Lipids), 1-palmitoyl-2-oleoyl-*sn*-glycero-3-phosphocholine (POPC, Avanti Polar Lipids), and the appropriate negatively charged lipid under nitrogen. The lipids re-suspended in the bilayer conjugation buffer. LUVs were prepared by extruding 20 times through a 0.1 μ m filter (Avanti Polar Lipids). β 3 CT was reacted with 4 fold molar excess 16:0 10 PE MCC. The conjugation reaction was performed at pH 6.8 to minimize lysine amine-maleimide side reactions which are strongly disfavored at below pH 7. After reacting the β 3 CT with the surface of the membrane for three hours, the bilayers were quenched with excess β -mercaptoethanol and dialyzed overnight. The final lipid concentration was calibrated using a POPC standard to calculate the amount of POPC in the β 3-conjugated bilayer.

Isothermal titration calorimetry (ITC). ITC experiments were performed on a MicroCa ITC200 (MicroCal/GE). Samples were prepared in degassed 25 mM 3-(N-morpholino)propanesulfonic acid (MOPS) buffer, pH 6.8, containing 100 mM NaCI. Samples were loaded into cells after vigorous washing with buffer and syringes were loaded with 50 μ L of titrant. Each experiment consisted of 20 x 2.0 μ L injections spaced at intervals of 2-5 minutes. A 0.2 μ L injection preceded all experiments and was not included in fitting. Fitting was performed using the Origin software suite (MicroCal/Origin Systems) as single binding events. Heats of dilution were calculated by titration into the prepared diazylate and subtracted prior to fitting.

Surface plasmon resonance (SPR). All SPR experiments were performed on a Biacore 3000 (GE Healthcare). Each injection was done in triplicate. Kinetic experiments were performed in degassed 10 mM (4-(2-hydroxyethyl)piperazine-1-ethanesulfonic acid (HEPES) buffer, pH 7.2, containing 100 mM NaCl. The CM5 chip was activated using an amine coupling kit (GE Healthcare). N-(2-Aminoethyl)maleimide (pH 7.0) was then injected and non-reacted activated groups were capped with 1 M ethanolamine (pH 7.0). The β 3 CT was immobilized onto the CM5 chip via maleimide crosslinking to \approx 600 RU. Background maleimide was quenched with 2- \Box mercaptoethanol. Talin-1 FERM domain constructs were injected at 50 µL/min with a 2 min injection time and 400 sec dissociation time. Binding data were fit to a double exponential using the BIAvaluation software.

For steady-state experiments, 100 nm LUVs were loaded onto an L1 chip. Talin-1 FERM domain constructs were injected for 10 min with a 120 sec dissociation time and a 10 μ L/min flow rate. Between injections, new lipid surfaces were generated when full dissociation was not observed. Steady state construct binding was compared to a reference channel containing 100% POPC. Binding was fit to a single component isotherm equation: R_{eq} = (KA * [analyte]*RU_{max})/ (KA *[analyte]*n+1)+background. *Fluorescence Polarization (FP).* FP experiments were performed as previously described (121). Briefly, FP was conducted in an ATF105 spectrofluorometer (Aviv Instrument, Inc) using a 0.3 cm path length cuvette. Spectra were measured using 1.0 nm slit widths at 25°C. The β 3 CT was labeled with maleimide-BODIPY-TMR at a cysteine introduced at residue 719. Talin-1 F0-F3 was titrated into 20 nM β 3 BODIPY-TMR and FP was measured at 575 nm after excitation at 530 nm. When F0-F3 was titrated into 20 nM FITC-labeled PIP5KI- γ , a dissociation constant and baseline parameters were derived from the fluorescence anisotropy signal/concentration isotherm using previously described methods (121).

Chapter 4 Transmembrane communication: general principles and lessons integrins and other receptors.

I. Introduction

In this review we explore how the binding of ligands to proteins induces functional conformational changes. Ligand induced conformational changes are fundamental to every aspect of protein function; for example, when an enzyme binds its substrate the act of binding induces changes in the protein's structure and dynamics to improve catalysis (122-124). Thus, it is often difficult to separate binding from catalysis. Similarly, binding of an allosteric ligand can lead to the propagation of conformational changes over long distances. We focus primarily on the generation and transmission of conformational changes in membrane proteins; we first introduce the types and limits of coupled, ligand induced conformational changes. Next we examine in quantitative terms the effects of different coupling mechanisms.

We then explore how ligand-binding affects the structures and functions of various membrane protein including bacterial histidine kinases and integrins, exploring the mechanisms by which multiple proteins, small molecules, metal ions, phospholipids, and post-translational modifications influence their activation and signaling. Finally, we demonstrate how interdomain coupling can generate diverse integrin signals.

II. Results

The coupling of ligand binding and environmental effects to conformational changes

Ligand binding generally leads to changes in both structure and dynamics, with increases and decreases in dynamics often occurring at different sites of the same protein(125). The effect of ligand binding on protein activity can be viewed in light of classical thermodynamic concepts of allostery, induced fit or conformational selection, and cooperativity(122-127). For simplicity, we consider a two-state transition between an active and resting state. The experimentally observed change in activity depends on several factors including: 1) the free energy difference or "energy gap" between the active and resting states, which determines the fraction of the protein in the activated state in the absence of the ligand, 2) the affinity of the protein for the ligand and their concentrations, and 3) the ability of the ligand to induce a transition when fully bound. Less than complete activation occurs, even when the ligand is fully bound, if it binds to both the activated and the resting conformations; in this case, the extent of activation relates to the relative affinity of the ligand for the activated versus resting state. These concepts of coupled equilibria are well studied and understood. Relating these to such phenomena as partial activation, potentiation, and synergistic ligands, is a simple matter of algebraic bookkeeping, as shown in later sections.

In the next section, we explore in more quantitative terms how these simple binding phenomena lead to a variety of activation curves, when there are multiple ligands capable of activating the protein of interest. In this case, distinct ligand-binding sites at different locations can "couple" to give a variety of outcomes; the binding of one can alter the affinity of the other for the protein, and they can have additive or nonadditive effects on the degree of maximal activation upon ligand saturation.

The coupling of binding and functional transitions refers to the phenomenon by which the binding of a ligand to one site influences the structure, dynamics, and binding affinity at a second site, which relates to the protein's energy gap and the ratio of affinities of the regulatory ligand for the activated versus the resting states. Given these simple requirements, it is not surprising that coupling can be achieved in different ways. One special case is when the binding of a ligand induces a change in association state. When no change in association is observed, it is often implicitly assumed that signaling occurs via "hard connections" in which helices shift, tilt or change register, mechanically driving a conformational change that propagates across a bilayer. A second model involves communication between subunits, which can involve large-scale or subtle changes in interactions between domains or between domains connected by flexible linkers. These different modes of communication can lead to different outcomes, when one measures the effects of mutations at

positions between the binding sites. For example, if a domain shifts as a rigid body, the search for a "pathway" from one location to the other might give rise to different results than if the transition of structural and dynamic information occurs by correlated motions of sidechains that propagate in a two-state manner between the two sites.

Finally, both the magnitude as well as the sign of the energy gap of a transmembrane (TM) protein depends on the membrane environment. Any shift in structure that changes the exposure of specific residues or the width (cross-sectional surface area) of a protein at a given position in the membrane will make the energy gap sensitive to the physical properties of the bilayer. Hence, mechanosensitive channels switch on in response to changes in bilayer tension(128). Potassium channels respond to the transmembrane electrical potential, and the location and coupling of cytoplasmic loops and helices changes in response to the fraction of IP lipids, sphingomyelin, and cholesterol in the bilayer(129-132). Indeed, the hydrophobic length of transmembrane helices is reflected in the width of the bilayers in the organelles to which they localize. Additionally, once located in a given membrane, there is tight coupling between the lipid environment and the conformation of the protein, as it, for example, partitions into cholesterol/sphingomyelin rich phases(133, 134).

Simple models for energetic coupling

<u>Energy gap</u> It is convenient to think of functional proteins as having at least two thermodynamic states – a resting state and an active state. The free-energy gap between these, $\Delta G_{gap} = G_{active} - G_{inactive}$ determines the equilibrium fraction of active protein as $f_{active} = \frac{1}{[1 + \exp(\Delta G_{gap}/RT)]}$. Since the largest possible fraction of the protein in the active state is 1, this also determines the maximal attainable degree of activation due to binding of any ligand or perturbation, via $A = 1 + \exp(\Delta G_{gap}/RT)$. For example, if ΔG_{gap} is 0.41 kcal/mol, then at room temperature the fraction activation is 1/3. Therefore, even the best activating agent can only result in a 3-fold increase in activity. However, if ΔG_{gap} is 6.82 kcal/mol, the maximal degree of activation is 10⁵.

A ligand causes activation when it binds with higher affinity to the active relative to the inactive state (see Figure 4.1A/B). If K_A and K_I are the dissociation constants for the ligand binding to the active and inactive states, respectively, then the ligand is activating if $K_A < K_I$. Moreover, the fraction K_I/K_A determines the activating potential of the ligand. When the system is saturated with ligand, the free energy difference between active and inactive states becomes: $\Delta G_{gap}^{eff} = -RT \ln(K_I/K_A) + \Delta G_{gap}$. For an ideal activating ligand, K_I approaches infinity, such that ligand saturation leads to complete activation, and the degree of activation is only limited by the

intrinsic ΔG_{gap} . If a ligand binds to both states, the maximum degree of activation is $A = (K_I/K_A) \cdot \frac{1 + e^{-\Delta G_{gap}/RT}}{1 + (K_I/K_A) \cdot e^{-\Delta G_{gap}/RT}}$, and is limited by both the intrinsic ΔG_{gap} and the ratio of equilibrium binding constants K_I/K_A (see

Figure 4.1B). For a given protein (and a fixed ΔG_{gap}), this ratio of equilibrium binding constants could differ for different ligands, leading to varying degrees of activation upon ligand saturation. The specifics of a biological process determine how large ΔG_{gap} needs to be and how it is modulated by binding. Large ΔG_{gap} and K_I/K_A lead to a high degree of maximal activation (Figure 4.1B). A large ΔG_{gap} is also useful in situations with multiple effectors that act in a combinatorial manner.

<u>Environment</u> ΔG_{gap} can be modulated by environmental variables, such as the membrane composition. While it may be possible to treat membrane effects as ensembles of binding sites, we treat them as environmental factors modulating ΔG_{gap} in the simplest case.

<u>Coupling</u> Once the receptor ectodomain shifts into its active state, this information is transmitted onto the functional cytoplasmic end of the molecule. This is sometimes referred to as allostery and is equivalent to stating that one end of the molecule is thermodynamically coupled to the

other. This coupling depends on the specifics of the molecule's structure and dynamics, but can be thought of as arising from differences in interdomain interaction free energies when in different states. This interdomain coupling or allostery can therefore be captured with a thermodynamic coupling free energy parameter, $\Delta\Delta G_c$, which determines the extent to which the different domains experience each other's states.

<u>Simple Model 1</u>. A two-state extracellular receptor domain, with an energy gap ΔG_{gap}^{R} , receives input from a signaling ligand, which binds preferentially to the active state of the receptor (Figure 4.1C). The receptor domain is thermodynamically coupled, via the coupling parameter $\Delta\Delta G_{c}^{R}$, to the cytoplasmic domain, which carries out the function of the system in its active state. The intrinsic activity of the cytoplasmic domain is controlled by its own energy gap, ΔG_{gap}^{C} . Figure 4.1D shows how the activity profile of this system (e.g. level of activation as a function of ligand concentration) depends on ΔG_{gap}^{C} and $\Delta\Delta G_{c}^{R}$. Because the function of the values of its gap lead to low levels of background activity, thus more switch-like behavior (compare profiles at low and high ΔG_{gap}^{C} in Figure 4.1C). However, because the two domains are coupled, the energy gap of the receptor domain also influences the level of activation. In fact, the
energy gaps of the two domains can be thought of as jointly modulating the overall effective energy gap of the system, as shown in Figure 4.1C.



Figure 4.1 Simple linked-equilibria models.

(A) A thermodynamic cycle illustrating the coupling between activation of a two-state protein and ligand binding. (B) Activation profiles of a two-state protein as a function of an agonist ligand concentration. Ratio of dissociation constants for the inactive and active states, K_I/K_A , and the intrinsic energy gap, ΔG_{gap} , determine maximal attainable activation. (C) A diagram of a simple signal-transduction molecule. Two domains, a receptor domain and a cytoplasmic functional domain, each with two states, are thermodynamically coupled, with the receptor modulated by an activating ligand. All eight states of the system are explicitly shown and the indicated equilibrium free energy differences correspond to the direction of the arrow with which they are shown. Free energy differences between ligand-bound states (back side of the cube) are omitted for simplicity, but follow simply from thermodynamic cycles. L indicates the concentration of the cytoplasmic ligand. (D) Activation profiles of the system in (C) as a function of the energy gap of the cytoplasmic domain, ΔG_{ean}^{C} , and coupling of the receptor to the cytoplasmic domain, ΔG_{c}^{R} . K_{A} and K_{I} are set to 10 nM and 20 μ M, respectively, and the energy gap of the receptor domain, ΔG_{gap}^{R} , is set to 3.0 kcal/mol. The effective energy for this gap be shown to be $\Delta G_{gap}^{eff} = \Delta G_{gap}^{C} + RT \ln \left(\frac{e^{\left(\Delta G_{gap}^{R} - \Delta \Delta G_{c}^{R} \right) RT} \left(1 + L/K_{I} \right) + \left(1 + L/K_{A} \right)}{e^{\left(\Delta G_{gap}^{R} + \Delta \Delta G_{c}^{R} \right) RT} \left(1 + L/K_{I} \right) + \left(1 + L/K_{A} \right)} \right) - \Delta \Delta G_{c}^{R}.$ Thus, if

 $\Delta\Delta G_c^R$ is zero (i.e. there is no coupling between the domain), the overall gap is just ΔG_{gap}^C , but as long as $\Delta\Delta G_c^R$ is negative (i.e. activation of the receptor promotes the active state of the functional domain), higher values of ΔG_{gap}^R contribute to a higher overall energy gap. Similarly, tighter coupling (lower $\Delta\Delta G_c^R$), leads to a higher overall gap larger degree of maximal activation.

<u>Simple Model 2</u> If the TM domain has active and inactive states, the system has 16 total states (Figure 4.2A), and the activation profile depends on the energy gap of the TM domain and its coupling to the cytoplasmic domain. Whereas larger energy gaps of the cytoplasmic domain always lead to larger overall energy gaps of the system (i.e. larger degree of maximal activation), this is not necessarily the case for the energy gap of the TM domain (see Figure 4.2B), and the effect of

increasing ΔG_{gap}^{TM} can go in either direction, depending on the other parameters. Intuitively, this is because unlike ΔG^{C}_{gap} , ΔG^{TM}_{gap} does not directly modulate the activity of the functional cytoplasmic domain, but rather does so through thermodynamic coupling. Increasing ΔG_{gap}^{TM} leads to lowering of the absolute active fraction of the cytoplasmic domain, by virtue of the two domains being coupled. This serves to silence background activity in the resting state, while in the presence of activating ligand it limits the maximal fraction of active protein. The overall consequence depends on which of these two effects dominates. Because the TM domain lowers activity at resting state, it may provide higher levels of activation compared to a system without a two-state TM domain (Figure 4.2B). This illustrates the advantage of having multiple loosely-coupled domains - it allows for strong switching behavior without resorting to large energy barriers between active and inactive states. Intermediate domains also provide additional opportunities for modulation of the signal. Additional environmental conditions, such as the state of the membrane or the presence of specific ligands, can be integrated into the final output of the signaling system.

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(A) In addition to the receptor and functional cytoplasmic domains as in Figure 4.1, here the TM domain is explicitly modeled as having two states. The intrinsic fraction of TM domain in the active state is defined by ΔG_{gap}^{TM} , and it is coupled to the cytoplasmic domain via ΔG_{c}^{TM} . The sixteen states

of the system along with the corresponding free energies are shown in the table on the right. (B) The dependence of activation profiles on ΔG_{gap}^{TM} and ΔG_c^{TM} . K_A and K_I are set to 10 nM and 20 μ M, respectively, and values of other parameters are shown. The line with circles demonstrates the case where no two-state TM domain is present, and the receptor domain couples directly to the cytoplasmic domain with the same ΔG_{gap}^{R} as in other models.

Simple Model 3 Extracellular and intracellular signals can be integrated by the same receptor. The VirA/VirG two-component system from Agrobacterium tumefaciens integrates multiple inputs with both its extracellular and cytoplasmic domains, including plant phenolic signaling molecules, monosaccharides, and pH(135). This can be modeled with the addition of a two-state cytoplasmic linker domain (Figure 4.3A). An intracellular ligand, such as a phenolic molecule, binds stronger to the activated state than the inactive one (ligand C in Figure 4.3A), whereas a different ligand, such as a sugar, modulates the states of the receptor domain (ligand L in Figure 4.3A). Both the maximal activation as well as the concentration of half-maximal activity (IC50) change for one ligand in response to different concentrations of the other (Figure 4.3B). Such cumulative modulation of activity by both ligands effectively amounts to signal integration. It can be used by the cell to perform certain actions only in cases, for example, where both ligands are present at certain concentrations. In this case, both ligands were chosen to bind more tightly to the active states of their respective domains, but repressing signals can also modulate receptor function.

A)



Figure 4.3 A realistic model of signal transduction, inspired by VirA protein of the VirA/VirG two-component system.

(A) In addition to the domains and states in the model presented in Figure 4.2, there is an additional cytoplasmic linker domain, with its own active and inactive states, which is modulated by the signaling ligand C, different from ligand *L* which modulates the receptor. The total number of states in this system is 64, and for clarity only the fully inactive and the fully active states are diagramed. (B) Activation, relative to resting state, is plotted against increasing concentration of ligand L, in the presence of different concentrations of C. This illustrates the ability to integrate signals from multiple ligands. Though this model does not explicitly treat the receptor as being dimeric (as is the case for the VirA protein), it can be easily extended to do so. In this case, binding of ligand may or may not be coupled of receptor dimerization and there may exist cooperativity (negative or positive) between the two ligand-binding sites. Potential scenarios that can be explored include: 1) binding of a single ligand may push the system towards the active state, whereas binding two ligands symmetrically may favor the inactive state 2) alternatively, the receptor may signal only when two ligands are bound, or 3) binding of the second ligand may proceed independently of the first one.

Inside-to-Outside Coupling in Integrins Receptors

Integrins are α - β heterodimeric adhesion receptors that mediate interactions between the extracellular environment and the intracellular cytoskeleton (6, 95, 136-138) and demonstrate how bidirectional transmembrane signal transduction is fine-tuned by multiple coupled equilibria in the extracellular, transmembrane, and cytosolic domains (Figure 4.4A). Integrin ectodomains contain a large globular N-terminal ligand binding "head" and distinct C-terminal α and β "legs" that cross the plasma membrane as single TM domains and terminate in short cytoplasmic domains that bind intracellular molecules. The inactive integrin adopts a bent conformation placing the head in proximity to the membrane and the distal legs. Upon activation, an extended conformation is favored and the α and β legs, TM, and cytoplasmic domains separate.



Figure 4.4 Thermodynamic model of integrin signaling.

(A) At the coarsest level, extracellular, TM, and intracellular domains of integrins can be thought as being either active or inactive, giving rise to eight coupled states as illustrated. The various effector molecules modulate integrin state by preferentially binding to select domain conformations. "Inside-out" signaling refers to modulation of the extracellular domain conformation via interactions between an integrin's cytosolic tails and intracellular signaling molecules. "Outside-in" events sometimes refer to the activation of oligomerization dependent signaling pathways subsequent to binding polyvalent ligands. However, keeping with our model, here it will refer to the ability of extracellular ligands to affect the conformation and function of intracellular domains. (B) Using reasonable values for the energy gaps and coupling constants associated with these states as well as equilibrium binding constants for the active and inactive states of different effector molecules, one can interrogate the thermodynamics of integrin signaling in a coarse, semi-approximate manner. Shown here is the level of integrin activation (separation of TM/tail regions) in response to increasing amounts of an activating antibody, as a function of Mn²⁺ concentration. This illustrates outside-in signaling. (C) AND (D) Inside-out signaling as illustrated by the fraction of extracellular domain bound to native ligand as a function of different amounts of active intracellular talin and kindlin. The binding affinity of the ligand as well as the amounts of talin/kindlin all couple to determining the activation state of the integrin.

Interactions between the α and β chain legs, TM, and membrane proximal cytosolic domains, stabilize the inactive integrin. Single point mutations result in receptor activation (15, 18, 139, 140) and membrane proximal ab disulfide cross-linking completely abrogates TM signal transduction in either direction (141, 142). While structural studies suggest that there is a distinct TM interface, the membrane proximal regions are dynamic (15, 16, 40, 142, 143), befitting the need for rapid signal transduction.

The "loose" dynamic interactions between the α and β chains near the membrane give way to an extensive interface between globular portions of the extracellular domain, which can be perturbed via mutagenesis (144). Thermodynamic coupling occurs throughout the globular ectodomain. Crystallization with and without ligand resulted in active-extended (145) and inactive-bent (146, 147) conformations respectively. Soaking the inactive crystals with a ligand resulted in only small-scale changes in the binding site without leg extension due to crystal contact constraints (148). Without such constraint, anti-integrin drugs can generate adverse side effects by activating the receptor and generating immune reactive epitopes or Ligand Induced Binding Sites (LIBS) in the

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integrin legs (138). Small molecule drugs can also uncouple subdomains. Allosteric inhibitors that compete for inter-domain contacts generate functionally split integrins in which the head is locked into an inactive conformation, but distal regions generate active LIBS(149). LIBS antibodies also shift the integrin equilibrium to an active conformation. Constraining the integrin C-termini stabilizes the bent conformation and the inactive ligand binding site, but the addition of LIBS antibodies and Mn²⁺ (an second allosteric activator) shifts the head domain equilibrium to an active state despite the distal constraints(150). Demonstrating bidirectionallity, LIBS antibodies induce TM separation when not constrained(18). The combination of LIBS antibody and a second activator, Mn²⁺, further increased activation, while Mn²⁺ alone did not significantly affect TM separation. Therefore, as in the case of VirA/VirG (Figure 4.3), multiple stimuli are integrated to produce different intracellular responses (modeled Figure 4.4B).



Figure 4.5 Coupled equilibria linking integrin TM interactions to cytosolic events.

Integrin $\alpha\beta$ TM heterodimers (left) stabilize inactive ectodomain conformations and TM separation (right) stabilize active ectodomain conformations. The $\alpha\beta$ heterodimer stabilizes interactions between the β -cytoplasmic domain and the inner leaflet of the bilayers (top). Cytosolic proteins talin-1 and kindlin-3 shift the equilibrium towards an active conformation by the stabilizing extended cytoplasmic domain and separated TM domains (bottom right).

While the integrin cytoplasmic domain is much shorter (~5-6 kD for β -integrins) than the ectodomain, it also integrates multiple signals to modulate extracellular function. There is convincing evidence of a defined α IIb β 3 TM interface, however, the membrane proximal and cytoplasmic domains have not provided consistent structures (15, 16, 38, 39, 54, 151). In Chapter 2, we describe the NMR structure and the dynamics of the α IIb and β 3 membrane proximal domains (1). The α IIb cytoplasmic domain

was disordered, but caused a kink in β 3 that was predicted to stabilize the interaction of two β 3 cytoplasmic helices with the bilayer. Significantly, these dynamic membrane interacting helices contain the primary motifs that bind the evolutionarily related talin and kindlin families of proteins (reviewed in (20) and (19)). Figure 4.5 outlines a simple linked equilibrium model, which is quantitatively described in Figure 4.4C/D, for how coupling between talin and kindlin binding motifs allows differentiation between ligands with different affinities for the receptor, effectively recapitulating an intermediate affinity state without the need for distinct extracellular conformations (149, 150). This model is further elaborated upon below.

IV. Coupling between integrin and kindlin subdomains produces distinct integrin activation states.

Our model of α IIb β 3 describes the integrin as series of thermodynamically coupled subdomains (Figure 4.6A). Each has an intrinsic ΔG_{gap} and a coupling parameter $\Delta \Delta G_c$ that describes communication between subunits. The goal of this model is to demonstrate that the presence of multiple intracellular regulatory domains can result in a range of apparent extracellular affinity states that do not require distinct ectodomain conformations. The studies reviewed above

demonstrate that there is interdomain communication in the ectodomain that is coupled to the TM state. Our HDX results with model integrin cytoplasmic domains demonstrate that despite its short sequence, there is also communication between the TM and both the talin binding site and the distal kindlin site. This establishes a path of communication from the kindlin and talin sites to the ligand binding site.



В

Domain	Conformation x	Occupancy =	# States
1) Ligand binding site	Active/Inactive	+/- Ligand	4
2) Metal binding site	Active/Inactive	+/- Mn ²⁺	4
3) Leg, LIBS antibody site	Active/Inactive	+/- LIBS antibody	4
4) Transmembrane domain	Active/Inactive		2
5) Cyto Helix 1	Active/Inactive	+/- Talin	4
6) Cyto Helix 2	Active/Inactive	+/- Kindlin	4

Figure 4.6 Simplified domain structure of α IIb β 3.

 α IIb β 3 contains a series of thermodynamically coupled subdomains that bind allosteric regulators. (A) The β 3 chain was modeled containing six subunits from the N-terminal ligand binding site (1) to the distal C-terminal

kindlin binding site (6). Each subunit an intrinsic equilibrium (ΔG_{gap}). Adjacent subunits are thermodynamically coupled ($\Delta \Delta G_C$). (B) With the exception of the TM, each subdomain recognizes a specific ligand that modulates the equilibrium between active and inactive conformations. In the case of sudomain (1), the natural ligand would be fibrinogen. Subdomain (2) includes the Mn²⁺ binding site (Mn²⁺ increases ligand fibrinogen to α IIb β 3). Subdomain (3) can bind LIBS antibodies. The combined active/inactive and ligand bound/ligand free states result in four states per domain. The TM does not have a ligand and therefore has only 2 states. Talin-1 and kindlin-3 ind regions (5) and (6) respectively.

As actual thermodynamic parameters for each subdomain are unknown, this model is mearly illustrative and intended to demonstrate the functional effects of communication between integrin subunits and not to make specific arguments as to their actual values. Thermodynamic parameters were chosen to conform to published data when possible and to fit with our experimental results. Without TM constraints, the integrin ectodomain equilibrium lies only slightly towards the inactive state (150) (152). Therefore a ΔG_{eav} of only 1 kcal/mol for each extracellular subdomain was used. We assumed that the coupling energy between subdomains was relatively high, lying between 3 kcal/mol and 6 kcal/mol. The ΔG_{gap} for the TM domain was chosen to be 3 kcal/mol as this site is the most sensitive to mutation and removal of the TM results in partial activation. Each subdomain can bind a ligand with different affinities for the active and inactive states. This ratio, K_I/K_A , which was held to 1000 i.e. activating ligands prefer the active conformation 1000 fold more than the inactive. Each subdomain can populate four states: 1) active without ligand; 2) active with ligand; 3) inactive without ligand; and 4) inactive with ligand. The TM domain, does not have a ligand and populates only two states, active (separated) and inactive (heterodimeric) states. The affinity for talin-1 and kindlin-3 was assumed to be 100 nM, which is of the order of magnitude observed in our ITC experiments in Chapter 3.

Thermodynamic coupling between extracellular, TM, and cytoplasmic domains allows dynamic regulation of ligand binding.

With no allosteric modulator, fractional occupancy of α IIb β 3 is dependent on the intrinsic equilibrium of the receptor, ΔG_{gap} and $\Delta \Delta G_{c}$, affinity for the ligand, and the ligand concentration (Figure 4.7A). Any regulatory protein, such as talin-1, that binds preferentially to the active conformation of the cytosolic domain will shift the extracellular ligand binding curve to the left, increasing the apparent affinity of the receptor (comparing Figure 4.7A and 4.7B). Likewise, a negative regulator can stabilize the inactive conformation and shift the equilibrium to the right (comparing Figure 4.7A and 4.7C).

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Domain	∆G _{gap} (kcal/mol)	∆∆G _{C, N+1} (kcal/mol)	[Ligand] (µM)
1) Ligand binding site	1	6	variable
2) Metal binding site	1	6	0
3) Leg, LIBS antibody site	1	6	0
4) Transmembrane domain	3	6	0
5) Cyto Helix 1	2	2	0
6) Cyto Helix 2	1	NA	0

1	6	
		variable
1	6	0
1	6	0
3	6	0
2	2	10
1	NA	0
	1 1 3 2 1	1 6 1 6 3 6 2 2 1 NA



Domain	∆G _{gap} (kcal/mol)	∆∆G _{C, N+1} (kcal/mol)	[Ligand] (µM)
1) Ligand binding site	1	6	variable
2) Metal binding site	1	6	0
3) Leg, LIBS antibody site	1	6	0
4) Transmembrane domain	3	6	0
5) Cyto Helix 1	2	2	10
6) Cyto Helix 2	1	NA	0

Figure 4.7 Binding of regulatory proteins to the integrin cytoplasmic tail affects receptor occupancy.

The fractional occupancy of a model integrin receptor was plotted as as a function of extracellular ligand concentration. Four different ligands were modeled ($K_d = 10$ nM, solid; 100 nM, dashed; 1 μ M, solid ball; 10 μ M, dashed ball). Thermodynamic parameters for integrin subdomains are displayed to the right. (A) Receptor occupancy without a regulatory protein. (B) Receptor occupancy in the presence of a distal activator such as talin ($K_A = 100$ nM, $K_I = 100$ μ M). (C) Receptor occupancy in the presence of a distal inhibitor such as talin ($K_A = 100$ μ M).

On the surface of platelets, $\alpha IIb\beta 3$ is surrounded a high affinity ligand, fibrinogen, which has a serum concentration between 5 and 10 μ M, well above its approximate K_d of 3-10 nM (153). Therefore, the intrinsic equilibrium of the receptor must favor the inactive state strongly to prevent spurious clotting and activation requires allosteric regulation via talin-1 and kindlin-3. In the absence of communication between the intracellular and extracellular domains ($\Delta\Delta G_c = 0$), receptor occupancy is independent of the talin-1 concentration and depends only on the overall ΔG_{eap} of the ectodomain, the ligand concentration, and the ligand's affinity for the integrin (Figure 4.8A). Coupling through the TM results in ligand binding that is dependent on the concentration of active talin-1 (Figure 4.8B). There is minimal binding to even strong ligands ($K_d = 10$ nM) in the absence of talin-1, while receptor occupancy approaches 100% upon talin-1 activation. This produces switch-like behavior, in which the receptor up-regulates affinity for a given ligand in the presence of minimal background activity. Even weaker ligands, with affinities of 100 nM and 1 μ M, have occupancies approaching 80% and 30%, respectively. If the

coupling energy is reduced from 6 kcal/mol to 3 kcal/mol, then dynamic range of the receptor decreases significantly. There would be 50% occupancy for a strong ligand ($K_d = 10$ nM) in the absence of talin and 90% occupancy at saturating concentrations of talin-1 (Figure 4.8C). A slightly weaker ligand with an affinity of 100 nM would have approximately 10% occupancy in the absence of talin-1 and 40% at saturating talin-1 concentrations. The 1 μ M ligand would have neglible binding even at saturating talin-1 concentrations.



Domain	∆G _{gap} (kcal/mol)	∆∆G _{C, N+1} (kcal/mol)	[Ligand] (µM)
1) Ligand binding site	1	6	10
2) Metal binding site	1	6	0
3) Leg, LIBS antibody site	1	6	0
4) Transmembrane domain	3	0	0
5) Cyto Helix 1, Talin	2	2	variable
6) Cyto Helix 2, Kindlin	1	NA	0



Domain	∆G _{gap} (kcal/mol)	$\Delta\Delta G_{C, N+1}$ (kcal/mol)	[Ligand] (µM)
1) Ligand binding site	1	6	10
2) Metal binding site	1	6	0
3) Leg, LIBS antibody site	1	6	0
4) Transmembrane domain	3	6	0
5) Cyto Helix 1, Talin	2	2	variable
6) Cyto Helix 2, Kindlin	1	NA	0

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Domain	∆G _{gap} (kcal/mol)	∆∆G _{C, №1} (kcal/mol)	[Ligand] (µM)
1) Ligand binding site	1	3	10
2) Metal binding site	1	3	0
3) Leg, LIBS antibody site	1	3	0
4) Transmembrane domain	3	3	0
5) Cyto Helix 1, Talin	2	2	variable
6) Cyto Helix 2, Kindlin	1	NA	0

Figure 4.8 Thermodynamic coupling between subdomains increases ligand binding dynamic range.

The fractional occupancy of a model integrin receptor as a function of talin-1 concentration in the presence of 10 μ M extracellular ligand. Four different ligands were modeled (K_d = 10 nM, solid; 100 nM, dashed; 1 μ M, solid ball; 10 μ M, dashed ball). Thermodynamic parameters for integrin subdomains are displayed to the right. (A) Receptor occupancy without coupling through the TM. (B) Binding of ligands with affinities between 10 nM and 10 μ M to the integrin with 6 kcal/mol coupling between subdomains as a function of talin-1 concentration. (C) Receptor occupancy with 3 kcal/mol coupling between subdomains as a function.

Coupling between talin-1 and kindlin-3 produces distinct functional states.

The ability of cells to discriminate between different integrin ligands has been attributed to the existence of distinct conformations in the ectodomain (150, 154). However, this effect can also be produced by coupling between multiple regulatory domains. To demonstrate this, weak thermodynamic coupling was introduced between the talin-1 and kindlin-3 binding sites ($\Delta\Delta G_c = 2$ kcal/mol). Binding of talin-1 (Figure 4.9A) or kindlin-3 (Figure 4.9B) to the integrin cytoplasmic tail results in distinct extracellular functional states. Kindlin-3 produces much lower ligand occupancy than talin-1, since its communication to the TM and the rest of the receptor is dampened by the ΔG_{gap} of the talin-binding site. Already, this produces two distinct extracellular states. The talin-only state has higher occupancy than the kindlin-only state. For example, in the presence of 100 μ M talin, the integrin would bind the 10 nM and 100 nM ligands with 90% and 50% occupancy, respectively (Figure 4.9A, black line). In contrast, the same concentration of kindlin-3 would produce only 10% and ~1% occupancy, respectively (Figure 4.9B, black line). Simultaneous binding of talin-1 and kindlin-3 to the integrin would produce further activation states. For each talin-1 concentration (white/grey/black bars) an increase in occupancy can be produced by the addition of kindlin-3 (Figure 4.9A and 4.9C). For example, if a cell were to activate both talin-1 and kindlin-3, there would be 50% occupancy of a ligand with a 1 μ M dissociation constant (Figure 4.9C, black line). By down regulating kindlin-3, but maintaining the same level of talin-1, a cell could release the 1 μ M ligand, but still bind higher affinity ligands (Figure 4.9A, black line).



Figure 4.9 Multiple integrin regulators can produce diverse activation states.

The fractional occupancy of a model integrin receptor as a function of talin-1 or kindlin-3 concentration in the presence of 10 μ M ligand. Four different ligands were modeled (K_d = 10 nM, solid; 100 nM, dashed; 1 μ M,

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solid ball; 10 μ M, dashed ball). Thermodynamic parameters for integrin subdomains are displayed to the right. (A) Receptor occupancy in the presence of talin-1, without kindlin-3 binding. White, grey, and black vertical lines mark 1 μ M, 10 μ M, and 100 μ M of talin-1, respectively. (B) Receptor occupancy in the presence of kindlin-3, without talin-1. White, grey, and black vertical lines mark 1 μ M, 10 μ M, and 100 μ M of talin-3, without talin-1. White, grey, and black vertical lines mark 1 μ M, 10 μ M, and 100 μ M of kindlin-3, respectively. (C) Receptor occupancy as a function of talin-1 concentration, with a constant kindlin concentration (5 μ M). White, grey, and black vertical lines mark 1 μ M, 10 μ M, and 100 μ M of talin-1.

Chapter 5. Conclusions

We hypothesized that cytosolic regulation of the integrin α Ilb β 3 is highly dependent on the membrane environment and uses coupled equilibria to transmit signals to the extracellular compartment. To test this, we studied the dynamics and structure of model α Ilb β 3 cytoplasmic domains in the presence of phospholipids and detergents (Chapter 2). We showed that interactions in the integrin TM domain are coupled to structural changes in the α Ilb β 3 cytoplasmic domain. The interaction between α Ilb and β 3 stabilizes a kink in the β 3 cytoplasmic domain, which is likely to encourage interactions with the inner leaflet of the plasma membrane. These interactions occur in two helical regions that include the talin-1 and kindlin-3 binding sites. This demonstrates that both the talin-1 and kindlin-3 sites are coupled to the interactions between the α Ilb and β 3 TM domains, suggesting a pathway whereby both talin-1 and kindlin-3 can affect the TM state and therefore the affinity of the ectodomain for fibrinogen.

We also investigated the interactions between α Ilb β 3 and its two regulators, talin-1 (Chapter 3) and kindlin-3 (Appendix A1). Preliminary studies showed that kindlin-3 binds the distal segment strongly in the absence of membrane. The distal kindlin-binding sequence is very dynamic and likely samples the cytoplasmic compartment frequently (Chapter 2). This suggests that kindlin-3 may bind the β 3 cytoplasmic domain without requiring a concerted interaction with the membrane, though membrane colocalization could serve to increase its affinity. This contrasts with talin-1. We showed that talin-1 binding to the integrin cytoplasmic domain is dependent on concerted interactions with the bilayer as its affinity for β 3 increases nearly 1000 fold when the integrin is immobilized onto negatively charged membranes. Though the membrane significantly increases the affinity of talin-1 for the β 3 cytoplasmic domain, its affinity for the integrin in the presence of membranes is not significantly higher than its affinity for negatively charged phospholipids alone. Similarly, we found that though talin-1 contains multiple lipid binding subdomains and that the interactions between these subunits and the bilayer are not cooperative. These results suggest that activation of the integrin by talin-1 is not opitimized for extremely high affinity as would be expected for perfectly cooperative interactions. Rather the energetics of the system suggest that the interactions are evolved for reversibility and sensitivity to changes in structure, post-translational modification, and lipid environment.

Fine-tuning of the integrin cytoplasmic domain for regulation and reversibility is befitting a receptor that must make dynamic interactions with the extracellular environment. Using a simple linked equilibrium model, we showed that coupling between the TM and the talin-1 and kindlin-3 binding sites can produce a range of extracellular functional states. Different affinity states for extracellular ligands are due to a change

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in the distribution of active and inactive states rather than adoption of specific intermediate extracellular conformations. Integrins can access a large range of affinity states by simply shifting the relative amounts of talin-1 and kindlin-3 interacting with the cytoplasmic tail.

Appendix 1. Preliminary investigations of the interaction between kindlin-3 and α IIb β 3

I. Introduction.

Among FERM domain proteins, the kindlins (kindlin-1, kindlin-2, kindlin-3) are evolutionarily the closest to the talin family. Like talin-1, kindlin-3 binds to the β 3 integrin cytoplasmic domain and is necessary for platelet integrin activation (7-10, 22). As the role of kindlin-3 in integrin function has only recently been appreciated, its mechanism of integrin regulation remains largely unclear. Talin-1 and kindlin-3 F3 subunits bind adjacent regions within the ß3 cytoplasmic domain though it is unknown whether they bind simultaneously (22, 155). Talin-1 recognizes a segment that spans from the end of the TM to Tyr747 and includes TMproxH and CytoH1. Kindlin-3 recognizes a distal segment that immediately follows the talin-1 binding sequence, spanning from Ser752-Tyr759, and including CytoH2 (Figure 2.5). Both proteins require intact NPxY or NxxY motifs to bind (22, 155). The spacing between the talin-1 and kindlin-3 NPxY motifs is absolutely conserved in nearly all integrins, with the exception of the β 5 integrins.

Like talin-1, the kindlin-3 FERM domain contains an additional F0 subunit. Whether kindlin-3 adopts a linear arrangement of like talin-1 (Figure 3.1) is unknown, since the structure of kindlin-3 has not been

solved. The kindlin-3 FERM domain also contains an inserted pleckstrin homology (PH) domain in its F2 subunit, which talin-1 lacks. The function of the PH domain has yet to be demonstrated, though PH domains – and FERM domains in general – are often involved in phospholipid interactions (26).

Our HDX data suggest that the β 3 cytoplasmic domain makes significant interactions with the bilayer. In particular, the portion that binds talin-1 is significantly protected from HDX in the presence of phospholipids. Furthermore, our NMR experiments show that α IIb induces a kink in the β 3 cytoplasmic domain, which is likely to orient the β 3 CT parallel to the membrane and further stabilize the interaction. We also have shown that the interaction between talin-1 and β 3 is guite weak and requires a simultaneous interaction between talin-1 and the membrane to be significantly populated. In contrast, our HDX and NMR experiments have shown that the kindlin-3 binding sequence in β 3, which is connected to the talin-1 sequence by a flexible linker, is highly dynamic and less helical and therefore less likely to bind the competing membrane surface. Based upon these results we hypothesized that the kindlin-3 sequence is more accessible than the talin-1 sequence, whether or not the protein/peptide interaction is more thermodynamically stable. A more dynamic kindlin-3 binding sequence would also not adopt a particular orientation relative the inner surface of the bilayer. This would suggest that unlike talin-1, kindlin-3 binding to the integin might be less dependent

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on simultaneous interactions with the membrane. If this were the case, then kindlin-3 might bind the β 3 CT with reasonably high affinity in the absence of membranes. Below are initial experiments attempting to characterize the interaction between kindlin-3 and β 3 to test the above model. We describe method for the purification full-length kindlin-3 from platelets and demonstrate that kindlin-3 binds to the β 3 CT in the absence of bilayers. These are initial experiments that are intended to be the starting points for future lab members.

II. Results

Purification of kindlin-3 from human platelets.

Kindlin-3 constructs were cloned into the pet28a vector and expressed in BL21-DE3 *E. coli.* However, expression of constructs containing full-length kindlin-3, the F2PHF3 subunit, and the F3 subunit resulted in insoluble inclusion bodies. We varied our expression conditions with temperatures ranging from 8°C to 37°C and IPTG concentrations ranging from 0.05 mM to 2 mM. However, only the isolated PH domain could be stably expressed in *E. coli.* We therefore purified full-length kindlin-3 from outdated platelets. Given its close homology to talin-1 and its localization to membranes, it was likely that kindlin-3 also contain positively charged surfaces. Therefore, the purification scheme (Methods) was designed around separation based on charge. After several rounds of SP-Sepharose and size exclusion chromatography, we produced nearly 10 mg of 95% pure kindlin-3 from six units of outdated platelets (Figure A1.1A/B). The predicted molecular weight for kindlin-3 is 75.4 kD and the purified protein had a calculated molecular weight of 74 kD by equilibrium analytical ultracentrifugation (Figure A1.1C).



Figure A1.1 Purification of Kindlin-3 from platelets.

(A) Kindlin-3 was purified by several rounds of cation exchange chromatography (Methods) and refined by size exclusion on an analytical Superdex-200 column. (B) Analytical size exclusion chromatography fractions were run on NuPAGE SDS 10% Bis-Tris gels and stained with Coomassie Blue stain. Higher order aggregates eluted in the void volume (Fraction 1). Kindlin-3 eluted between 13 and 15 minutes (Fractions 2-8) and was found between the 66 kD and 97 kD bands of a Mark12 molecular weight standard. (C) Equilibrium analytical ultracentrifugation of the purified protein. Kindlin-3 was prepared in 25 mM MOPS 100 mM NaCl, pH 7 at 3 μ M (left) and 1.5 μ M (right). The samples were sedimented at 15000 rpm, red/green, 20000 rpm, dark blue/black, and 25000 rpm, light blue/brown. Data were globally fit using the Igor software

program to a MW of 74 kD. {AUC experiments were performed once. Performed by DTM with the guidance of PB.}

Preliminary experiments suggest kindlin-3 binds β 3 with high affinity in the absence of bilayers.

Our HDX and NMR measurements of the β 3 cytoplasmic domain showed that the kindlin-3 binding sequence is very dynamic and likely adopts many conformations relative to the membrane surface. We therefore hypothesized that, unlike talin-1 interaction with β 3, kindlin-3 might not depend on a coordinated interaction with the bilayer surface. If this were the case, then the complex between kindlin-3 and β 3 should be stable at much lower concentrations than the interaction between talin-1 and β 3. We used analytical ultracentrifugation to measure the interactions of labeled β 3 with both talin-1 and kindlin-3 in the absence of membranes. In each case 1 μ M of BODIPY TMR labeled β 3 CT (MW 6242 kD, prepared as in Chapter 2 for FP) was sedimented in the presence of excess kindlin-3 (3.5 μ M or 1.75 μ M) or excess talin-1 (10 μ M). In the presence of 10 μ M talin-1, labeled β 3 cytoplasmic domain sedimented with an apparent molecular weight of 7.7 kD (Figure A1.2A), which is close to its predicted monomeric 6.2 kD mass. In the presence of kindlin-3, there was a clear shift towards higher molecular weight species. Global fitting of both kindlin-3 concentrations resulted in estimation of a dissociation constant between 100 nM and 3 µM (Figure A1.2B). Single species fits produced molecular weights of 43 kD and 12 kD for the 3.5:1 and 1.75:1 experiments, respectively, suggesting a Kd on the order of several micromolar. However, without higher concentrations of kindlin-3 and β 3, it is impossible to calculate the affinity for β 3 with higher accuracy. This would require more kindlin-3 than could be prepared for the experiment.

Figure A1.2 Equilibrium AUC of Kindlin-3 and β 3.

(A) Equilibrium AUC of 10 μ M talin-1 and 1 μ M BODIPY TMR- β 3. Sedimentation was monitored at 600 nm. The sample was sedimented at four velocities (10000 rpm black, 15000 rpm blue, 20000 rpm green, 25000 rpm red). Data were globally fit to a single species of 7.7 kD using Igor. (B)Kindlin-3 and BODIPY- β 3 were combined in two ratios: 3.5 μ M kindlin-3 and 1 μ M BODIPY- β 3 (left) and 1.75 μ M kindlin-3 and 1 μ M BODIPY- β 3 (right). Sedimentation AUC was performed while monitoring 600 nm absorbance at three velocities (15000 rpm red/brown, 20000 rpm dark blue/light blue, and 25000 rpm green/black). Sedimentation was globally fit to a two species 1:1 binding model with the ϵ 600 for Kindlin held to 700 cm⁻¹M⁻¹ and ϵ 600 for BODIPY- β 3 held to 70000 cm⁻¹M⁻¹. Due to the low protein concentration and absorbance relative to background, sedimentation fit equally well to dissociation constants between 100 nM and 3 μ M. {*AUC performed once by DTM under the guidance of PB.*}

III. Methods

Kindlin-3 purification. Six to 10 units of outdated platelets (Hospital of the University of Pennsylvania Blood Bank) were lysed by N₂ bomb on ice in the presence of mammalian protease inhibitors and 10 mM EDTA, 100 mM NaCl. EDTA was necessary to prevent significant proteolysis of kindlin-3. Lysates were centrifuged at either 30k x g or 100k x g to clear insoluble debris. After clearance, lysates were mixed with Q-Sepharose batch-wise for 1 hour at 4 degrees Celsius to clear serum and negatively charged proteins. The cleared lysate was loaded onto SP-sepharose (25 mL) in 50 mM NaCl batchwise and poured into a high-pressure FPLC column for elution on gradient between 50 mM NaCl and 400 mM Arginine, pH 7 on a Akta FPLC (GE Healthcare). The kindlin-3 containing fractions were then concentrated and loaded onto a high resolution SP-

sepharose column (16/10, 25 mL bed volume) (GE Healthcare) and eluted over a 400 mL gradient. At this point, the primary impurities were full length talin and talin head domain, which were largely removed by size exclusion on Superdex200 or Superdex75 analytical column (200 uL injection volume) (GE Healthcare). A final high resolution SP sepharose step was performed on a 400 mL gradient to 400 mM Arginine.

Equilibrium Analytical Ultracentrifugation. Purified peptide and protein samples were prepared in 25 mM MOPS 100 mM NaCl, pH 7.0. Experiments were performed at 25°C. Samples were loaded into a 6-channel carbon-epoxy centerpiece and centrifuged at 10000 rpm, 15000 rpm, 20000 rpm, and 25000 rpm in a Beckman XL-I analytical ultracentrifuge. Samples were monitored at 280 nm and 600 nm in a 1.2 cm path length cell. The molecular weight and partial specific volumes of kindlin-3 were calculated to be 75.4 kD and 0.73 ml/g, respectively using SEDINTERP. The buffer density was calculated to be 1.003 g/ml. Data were globally fit using Igor Pro (Wavemetrics).
Appendix 2. NMR studies of the c-Src SH3 domain and the β 3 cytoplasmic domain.

I. Introduction

Src family kinases (SFKs) are a family of non-receptor tyrosine kinases implicated in a number of biological processes (156, 157). There are nine human SFKs expressed—often redundantly—in nearly all human tissues (158). Of the SFKs, c-Src is the best studied, being discovered as the first proto-oncogene in the 1970s (159). Redundancy in SFK paralog expression has resulted in single-gene knockout phenotypes that are unremarkable, especially in the case of platelets (158). In order to observe a significant platelet phenotype, it was necessary to generate platelets deficient in six kinases (c-Src, Fyn, Fgr, Hck, Lyn, and Yes), which resulted in reduced platelet spreading on the α IIb β 3 ligand fibrinogen (160). These gene-deleted platelets and wild-type platelets treated with SFK inhibitors do not have a defect in integrin activation or ligand binding. Rather they have a downstream defect in tyrosine phosphorylation and cytoskeletal mobilization after initial adhesion to fibrinogen, suggesting that SFKs participate in platelet integrin outside-in signaling, which is supported by the fact that SFKs are recruited to the integrin associated cytoskeleton after platelet aggregation (161, 162) and that c-Src activation requires binding of α IIb β 3 to fibrinogen (160).

SFKs contain well-conserved domain structures (Figure A2.1). Each SFK contains an N-terminal membrane anchor followed by a flexible

linker (black), a Src Homology 3 (SH3, purple) domain, a flexible linker (blue), a Src Homology 2 (SH2, green) domain, a flexible linker (yellow), the kinase domain (orange), and then finally a C-terminal regulatory tail Inactive SFKs adopt compact structures that are stabilized by (red). multiple intramolecular interactions (163). In the inactive state, the phosphotyrosine recognizing SH2 domain (green) binds to the phosphorylated C-terminal tail (red). The inactive conformation is also stabilized by an interaction between the SH3 (purple) domain and a second internal ligand in the linker (yellow) between the SH2 and kinase domains. C-terminal Src Kinase (CSK) phosphorylates the C-terminal tail (164), while protein-tyrosine phosphatase-1B (PTP-1B) activates the kinase by dephosphorylating the C-terminus (165). Activation also requires phosphorylation of a regulatory loop in the kinase domain.

In the active state, the SH2 and SH3 domains are free to interact with other proteins. Such interactions stabilize the active kinase state and determine the localization of SFKs (156). Localization also depends on the N-terminal membrane anchor. All SFKs are myristoylated and a subset are palmitoylated (166, 167). C-Src is not palmitoylated and contains a conserved patch of positively charged Arg and Lys residues (Figure A2.1), which has been demonstrated necessary for binding of c-Src bilayers containing negatively charged phospholipids such as phosphatidylserine and phosphatidylglycerol (25) (29).



+ + + +++ N-terminus: Myristoyl-GSNKSKPKDASQRRR

Figure A2.1 Domain structure of c-Src in its inactive conformation.

From N- to C-terminus: 1) N-terminal membrane anchor that is posttranslationally myrstoylated (Black). C-Src contains a stretch of basic amino acids (bottom). 2) SH3 domain (purple) 3) Interdomain linker (blue). 4) SH2 domain (green). 5) SH2-kinase linker (yellow) 6) Kinase domain (orange). 7) C-terminal regulatory tail (red).

SFK substrate specificity appears to be less dependent on the substrate sequence than physical proximity (168). Therefore, the specificity of SFK protein-protein and protein-lipid interactions is extremely important in determining function. Recruitment to integrin signaling complexes could potentially involve the N-terminal membrane anchor, the SH3 domain, and the SH2 domain. Both the SH3 and SH2 subunits of c-Src have been shown to interact with Focal Adhesion Kinase (FAK) (169) and p130Cas(170, 171), two proteins found in integrin signaling complexes. A number of studies have also suggested that c-Src binds directly to the last three residues (Arg760-Gly761-Thr762) of the β 3 integrin cytoplasmic domain (23). Transgenic platelets containing mutations in this region exhibited defects in spreading and down-stream integrin events (137, 172). The Arg-Gly-Thr motif implicated in the c-Src / β3 interaction is quite different from the canonical polyproline sequence recognized by the c-Src SH3 domain (Figure A2.2B) (173). The polyproline backbone packs against a groove composed of aromatic residues (Tyr93, Tyr95, Tyr139) (174, 175). A specificity pocket defined by Trp121 and Tyr134 is at the end of the aromatic groove and is flanked by the n-Src loop and the RT-loop, which affect substrate specificity (173).

We hypothesized that the β 3 cytoplasmic domain may bind to the c-Src SH3 domain either along the Tyr-groove or in the aromatic specificity pocket. We characterized integrin/SH3 interaction using biophysical techniques sensitive to perturbation at these residues. We were surprised to find that the interaction is much weaker than predicted by published biological and biochemical studies(23, 137, 172), suggesting that the interaction may depend on other proteins or the lipid environment.



Figure A2.2. The interaction between the c-Src SH3 domain and a polyproline peptide.

Interaction between the c-Src SH3 domain and a polyproline peptide (yellow) is shown down the axis of the peptide (A) and rotated 90° (B). The interaction requires packing of the polyproline backbone against an aromatic groove and "specificity pocket" composed tyrosine and tryptophan residues (purple). Residues from the RT-loop and the n-Src loop interact with the backbone and side chains of the peptide and help determine the sequence specificity of the interaction (PDB code 1nlp)

II. Results

C-Src, β 3 integrin, and polyproline constructs.

The c-Src SH3 domain was cloned between residues Ala83 and Ser143 into the pet28a vector with an N-terminal 6xHis tag followed by a thrombin cleavage site for removal of the affinity tag. The β 3 cytoplasmic domain (β 3-NITYRGT) between residues Asn756 and Thr762 was

synthesized on Wang resin generating a short seven amino acid peptide containing the Arg-Gly-Thr binding motif c-Src binding motif. Wang resin was chosen because the resulting peptide contained a C-terminal carboxylate like the natural sequence. The polyproline peptide Arg-Pro-Leu-Pro-Pro-Leu-Pro with a C-terminal amide was synthesized on Rinkamide resin.

Circular dichroism and Trp-fluorescence of c-Src SH3 domain and the β 3 C-terminus.

It has been reported that c-Src co-immunoprecipitates α IIb β 3, that the SH3 domain interacts with the β 3 C-terminus by ELISA with a micromolar binding binding constant, and that mutations to the β 3 Cterminal Arg-Gly-Thr sequence results in outside-in integrin signaling defects (23, 137, 172). Though the integrin sequence is quite different from the canonical polyproline ligand, we hypothesized that the interaction might also occur along the Tyr-groove or the Trp-containing specificity pocket. Changes in the local environment surrounding aromatic residues can often be measured by changes in CD signal or Trp fluorescence (176). Binding of polyproline ligands to this region was previously observed using these techniques (177) (178). We therefore attempted similar experiments with our β 3-NITYRGT peptide, reasoning that such studies would support binding to the aromatic region of the SH3 domain.

Solutions containing 40 μ M SH3 domain and 650 μ M β 3-NITYRGT were prepared in pH 7.0 25 mM phosphate 100 mM NaCl. CD spectra

were taken of each sample (Figure A2.3). The SH3 domain was then mixed 1-to-1 with the β 3-NITYRGT peptide and the CD spectrum of the mixture was taken (Figure A2.3A, green). The expected spectrum for the mixture in the absence of interaction was also calculated by averaging the SH3 and β 3-NITYRGT spectra (Figure A2.3A, green dash). The difference between the expected and experimental CD spectra was then calculated (Figure A2.3A, red). There was no significant difference between the two, demonstrating that mixing the β 3-NITYRGT peptide and SH3 domain resulted in no change in CD signal at the concentrations used. However, when similar experiments were performed with 650 μ M polyproline RPLPPLP (Figure A2.3B), a clear shift in the CD was observed (Figure A2.3B, red). The difference spectrum for RPLPPLP has a maximum at 225 nm, which replicates previously published results (177).



Figure A2.3 Comparison of CD and Trp fluorescence changes in the presence of a polyproline peptide and β 3-NITYRGT.

Protein and peptide samples were prepared in 25mM phosphate 100 mM NaCl, pH 7.0. (A) CD spectra were taken of 40 µM c-Src SH3 domain (dark blue), 650 μM β3-NITYRGT peptide (light blue), and a 1:1 mixture of the two samples with 20 μ M SH3 and 325 μ M β 3-NITYRGT (green). The expected spectrum in the absence of interaction was calculated by averaging the SH3 and β 3 spectra (dashed green). The difference between the actual and expected spectra was calculated (red). (B) CD spectra were taken of 40 µM c-Src SH3 domain (dark blue), 650 µM polyproline (light blue), and a 1:1 mixture of the two samples with 20 μ M SH3 and 325 µM polyproline (green). The expected spectrum in the absence of interaction was calculated by averaging the SH3 and polyproline spectra (dashed green). The difference between the actual and expected spectra was calculated (red). (C) Tryptophan emission spectra of the c-Src SH3 domain. SH3 solutions with or without peptide were excited at 295 nm and emission spectra were recorded from 300-400 nm. Spectra were recorded for 50 μM SH3 domain alone (light blue), 50 μM SH3 domain with 200 μM polyproline (dark blue), 50 μM SH3 domain with 200 μ M β 3-NITYRGT (green), and 50 μ M SH3 domain with 200 μ M polyproline and 200 μM β3-NITYRGT (magenta). {CD and fluorescence experiments are representative of three independent experiments. Protein and peptide samples were made by DTM. Experiments were performed by DTM.}

Binding of a polyproline ligand to the c-Src SH3 domain results in a blue shift in tryptophan fluorescence (178). We reasoned that though mixing of the c-Src SH3 domain and β 3-NITYRGT did not result in a significant change in circular dichroism, if the β 3 peptide bound to the tryptophan containing specificity pocket in the SH3 domain, it might induce in a similar blue shift in fluorescence. A 50 μ M solution of SH3 was excited at 295 nm and an emission spectrum was recorded between 300 and 400 nm, with an emission maximum at 355 nm (Figure A2.3C, light blue). The same measurement was performed on a second solution

containing 50 μ M SH3 and 200 μ M β 3-NITYRGT, with an emission maximum at 356 nm (Figure A2.3C, green). In comparison, the addition of 200 μ M RPLPPLP shifted the SH3 emission maximum to 347 nm (Figure A2.3C, dark blue). Finally, the emission maximum for a sample containing 50 μ M SH3, 200 μ M β 3-NITYRGT, and 200 μ M RPLPPLP was at 346 nm. These results show that unlike the polyproline peptide, the integrin cytoplasmic domain peptide does not cause a blue shift in the SH3 domain Trp-emission spectrum. Likewise, it does not compete for binding of the SH3 domain to the polyproline sequence.

NMR titration of the β 3 cytoplasmic domain the c-Src SH3 domain.

Our CD and Trp fluorescence experiments did not demonstrate an interaction between the c-Src SH3 domain and β 3-NITYRGT at concentrations as high as 350 μ M β 3-NITYRGT. We reasoned that the interaction was either extremely weak and not observed at the concentrations used or it might occur outside of the aromatic groove or specifity pocket and therefore CD and Trp fluorescence would not be well suited for the measurement. We chose to use NMR, which has successfully been used to study weak protein-protein interactions (179). The c-Src SH3 domain was expressed in ¹⁵N M9 minimal media and purified by nickel chromatography. The 6xHis tag was removed using thrombin protease and the purified protein was transferred into pH 6.0

phosphate buffer containing 100 mM NaCl and concetrated. Two NMR samples were prepared for ¹H-¹⁵N HSQC NMR. The first contained 1 mM ¹⁵N SH3 (Figure A2.4, red) and the second contained 1 mM ¹⁵N-SH3 and 15.7 mM β 3-NITYRGT peptide (Figure A2.4, magenta). In the presence of 15.7 mM β 3-NITYRGT, a number of SH3 residues exhibited changes in chemical shift. The interaction appears to be in fast exchange, as only shifts in peak location were observed. Slow exchange binding would be expected to generate two sets of peaks corresponding to the bound and unbound forms, while intermediate exchange would produce broadened signals of both species with potentially a loss of intensity. In this titration, peaks remained sharp and intense.



Figure A2.4 Changes in SH3 chemical shift in the presence of $\beta3\text{-}$ NITYRGT

(A) ¹H-¹⁵N HSQC spectra of samples containing 1 mM c-Src SH3 domain (red) and 1 mM ¹⁵N-labeled c-Src SH3 domain with 15.7 mM β 3-NITYRGT (magenta). Both samples were prepared in pH 6.0 20 mM phosphate containing 100 mM NaCl. {*Samples prepared by DTM. Experiment performed by DTM with help of HC.*}

The difference in the HSQC spectrum for each residue was calculated by the relationship $\Delta ppm = (\Delta^1 H^2 + (\Delta^{15} N/5)^2)^{1/2}$ (Figure A2.5A). The standard deviation for the per residue chemical shift change was 0.04. Sixteen residues had chemical shifts greater than one standard deviation: Tyr95, Glu96, Arg98, Thr99, Glu100, Thr101, Asp102, Leu103, Phe105, Asn115, Asn116, Glu118, Trp121, Ser126, Gly130, and Tyr134. SH3 residues were color coded according to their chemical shift change and mapped onto a previously reported structure of the c-Src SH3 domain (Figure A2.5B/C). The majority of residues perturbed by β 3-NITYRGT were in the n-Src and RT-loops and centered around the specificity pocket (Trp121 and Tyr134). The largest chemical shift changes occurred in residues in or adjacent to the RT-loop, especially residues Arg98 (0.25 ppm), Glu100 (0.15 ppm), and Asp102 (0.13 ppm). Tyr93, Tyr95, and Tyr139 form the aromatic polyproline binding groove (Figure A2.2, magenta) that binds polyproline ligands. Of these, only Tyr95, which is adjacent to the specificity pocket, was perturbed by β 3-NITYRGT. Tyr93 and Tyr139 were unchanged (Figure A2.5B/C, magenta).



Figure A2.5 β 3-NITYRGT causes chemical shift changes centered around the SH3 specificity pocket.

(A) The change in chemical shift in the presence of 15.7 mM β 3-NITYRGT (Figure A2.4) was calculated by the equation $\Delta ppm = (\Delta^1 H^2 + (\Delta^{15}N/5)^2)^{1/2}$. (B) Chemical shift changes were color coded and mapped onto a previously solved NMR structure of the c-Src SH3 domain (pdb

code 1nlp). The maximum chemical shift change was 0.25 ppm (red, Arg98). The minimum cutoff was 0.05 (dark blue, Tyr95). Tyr95 is part of the aromatic groove that binds polyproline ligands. The remaining residues in the aromatic groove (Tyr93 and Tyr139) did not have significant changes in chemical shift and are colored magenta.

The SH3 NMR sample containing 15.7 mM β3-NITYRGT was serially titrated against the sample containing only SH3 domain. This produced NMR samples with constant 1 mM SH3 concentrations and β3-NITYRGT concentrations ranging between 5.3 mM and 15.7 mM. HSQC spectra were recorded for each sample. Figure A2.6A shows the change in Arg98 chemical shift with the addition of β 3-NITYRGT. The relationship was linear (Figure A2.6B), indicating that the titration had not reached saturation and that the majority of SH3 domain was unbound, even in the presence of 15.7 mM β 3-NITYRGT. Since the majority of the protein was not bound, $[SH3]_{total}$ approximates $[SH3]_{free}$ and $[\beta 3]_{total}$ approximates Therefore, the relationship Δ (chem. shift) = Δ ppm_{max} * $[\beta 3]_{\text{free}}$. $([SH3]_{free}*[\beta 3]_{free})/(K_D*[SH3]_{total})$ simplifies to Δ (chem. shift) =[Ligand]* $(\Delta ppm_{max}/Kd)$. The term Δppm_{max} represents the change in chemical shift when the SH3 domain is fully bound by ligand. As the slope is linear, it can be assumed that the maximum chemical shift change is on the order of 10 fold the observed change, or 2.5 ppm. Since the maximum reasonable change in chemical shift is 5 ppm for any protein-protein interaction, this places the Kd between 175 mM and 350 mM.



Figure A2.6 ${}^{1}N{}^{15}N$ HSQC titration of β 3-cytoplasmic domain and c-Src SH3.

(A) ¹H-¹⁵N HSQC spectrum c-Src SH3 residue Arg98 backbone amide without (red) and with 5.3 mM (yellow), 8.4 mM (orange), and 15.7 mM (purple) β 3-NITYRGT. (B) Relationship between Arg98 chemical shift change and total concentration of β 3-NITYRGT. The change in chemical shift was calculated according to the equation Δ ppm = $(\Delta^{1}H^{2} + (\Delta^{15}N/5)^{2})^{1/2}$. The data was fit to a linear curve with a slope 0.0153 ppm/(mM titrant). {*Samples prepared by DTM. Experiments performed by DTM.*}

III. Discussion.

Though SFKs are necessary for integrin outside-in signaling, how they are recruited to these signaling complexes and the degree of specificity built into the nine different kinases remains unclear. SFKs contain multiple subunits capable of making protein-protein and proteinlipid interactions, therefore targeting to a particular signaling complex could potentially require the coincidence of multiple relatively weak interactions. There is clearly a degree of redundancy built into SFK signaling in platelets as six kinases are required to be knocked out to generate platelets that are unable to spread on fibrinogen. As a starting point for understanding the regulation of these processes, we have investigated the interactions between c-Src and the β 3 integrin cytoplasmic domain.

We sought to characterize the interaction between the c-Src SH3 domain and the β 3 integrin cytoplasmic domain C-terminus. Several lines of evidence have suggested that the c-Src SH3 domain specifically binds to the β 3 integrin cytoplasmic domain. It has been shown that mutations to the terminal Arg-Gly-Thr sequence result in decreased spreading of both transfected cells (23, 180, 181) and platelets on a surface coated with the α IIb β 3 ligand fibrinogen(137); that mutations to this sequence disrupt co-immunoprecipitation of c-Src and α IIb β 3(160); that mutations to this sequence block solid-phase binding of GST-SH3 domain to the β 3 cytoplasmic domain (23, 181); and that a synthetic membrane permeable peptide containing the Arg-Gly-Thr sequence disrupts platelet c-Src signaling (182). The interaction was studied in solution using monomeric c-Src and β 3 constructs in contrast to the previously reported solid-phase experiments, which used a dimeric GST-SH3 fusion protein(23). There was no evidence of the interaction by circular dichroism or tryptophan fluorescence and β 3 did not compete with binding by a polyproline ligand. It is possible that an interaction between β 3 and the SH3 occurs away from the canonical polyproline binding pocket, therefore we chose to study the interaction by NMR. We found that addition of the β 3 C-terminus does in fact cause chemical shift changes in the SH3 domain in the region of the specificity pocket. However this requires extremely high concentrations of β 3 and even 15.7 mM β 3 does not saturate the SH3 domain and the calculated dissociation constant may be as high as several hundred millimolar.

Typical polyproline ligands contain Lys or Arg residues that pack against the Trp and Tyr in the specificity pocket and make electrostatic interactions negatively charged residues provided by the n-Src or RT-loop. The β 3 sequence provides a conserved Arg residue at position 759 in the Arg-Gly-Thr sequence and replacement of this sequence with a Glu-Gly-Lys is reported to disrupt c-Src signaling and interactions with β 3 (181). Further NMR studies are necessary with similar mutant peptides to determine whether such a sequence substitution will also disrupt the interaction seen by NMR.

Our measured interaction is much weaker than that reported for binding of solid-supported β 3. It is possible that the differences in the β 3 cytoplasmic domain constructs used can account for the significant variation between our results and those reported previously. We chose to use the C-terminal seven residues for our NMR experiments as these residues have previously been shown necessary for binding. The discrepancy in apparent β 3 binding affinity may partially be due to avidity effects of surface binding by dimeric GST-SH3 fusion proteins (23, 181). It is also possible a third species (or more) may be necessary for the apparent interaction to occur under biological conditions. The adjacent

sequence in the β 3 cytoplasmic domain also recognizes members of the kindlin and filamin families of protein and therefore could conceivably cooperate in binding. C-Src also binds other integrin signaling molecules including FAK (169) and p130Cas(170, 171). Further studies are needed to characterize the recruitment of SFKs to integrin adhesions and to determine whether the membrane environment plays a role.

IV. Methods

Protein and peptide constructs used. The c-Src SH3 domain was cloned between Ala83 and Ser143 into a pet28a vector between the Ndel and Xhol restriction sites. Expression was performed in BL21-DE3 cells (Novagen). They were grown in LB-media to an OD600 of 0.5 at 37°C and 1 mM IPTG was added. After 3 hours of expression, cells were lysed by tip sonication on ice for 20 minutes in pH 7, 500mM NaCl, 20mM imidazole with a protease inhibitor cocktail (Sigma-Aldrich) containing 1 mM EDTA. The lysates were cleared by centrifugation at 30,000 g for 30 minutes. The supernatant was loaded onto a HiTrap Ni-NTA column (GE Healthcare) and eluted over a 50 mL gradient to 500 mM imidazole. Imidazole was dialyzed away overnight into pH 7.5 Tris, 100 mM NaCl, in the presence of thrombin protease (GE Healthcare) to remove the 6xHis tag. The SH3 domain then purified away from the 6xHis tag by DEAE ion exchange chromatography. β3-NITYRGT was synthesized on Wang resin preloaded with a Thr residue using a Symphony peptide synthesizer at 0.1 mM scale in 25% DMSO/75% NMP with Fmoc-chemistry. β3-NITYRGT was N-terminally acetylated and cleaved from the resin TFA/triethylsilane/water (95/2.5/2.5 v/v) for three hours and precipitated into dry-ice cold diethylether. β3-NITYRGT was purified by reverse phase HPLC (Varian Prostar) using a C4 preparative column (Vydac) on a water/acetonitrile gradient with 0.1% TFA.

RPLPPLP was synthesized on Rink-amide resin using a Symphony peptide synthesizer at 0.1 mM scale in 25% DMSO/75% NMP with Fmocchemistry. RPLPPLP was N-terminally acetylated and cleaved from the resin TFA/triethylsilane/water (95/2.5/2.5 v/v) for three hours and precipitated into dry-ice cold diethylether. β 3-NITYRGT was purified by reverse phase HPLC (Varian Prostar) using a C4 preparative column (Vydac) on a water/acetonitrile gradient with 0.1% TFA.

Circular dichroism. Purified c-Src SH3 domain was transferred to CD buffer (mM phosphate 100 mM NaCl, pH 7.0) by dialysis and a stock containing 40 μ M protein was made (calculated by absorption at 280 nm). Stocks of β 3-NITYRGT and RPLPPLP were made in the CD buffer by weighing out the respective peptides dissolving them in the appropriate volume of buffer. CD spectra were taken on a JASCO J-810

spectropolarimeter. Measurements were made stepwise for each 1 nm between 215 nm to 240 nm and averaged over three scans at 25°C.

Tryptophan fluorescence. Stocks of purified c-Src SH3, β 3-NITYRGT, and RPLPPLP were prepared in CD buffer. Three samples were prepared: 50 μ M SH3, 50 μ M SH3 + 200 μ M RPLPPLP, and 50 μ M SH3 + 200 μ M β 3-NITYRGT. All fluorescence measurements were performed at 25°C in an Aviv Associates fluorometer. Samples were excited at 295 nm using a 2 nm band-pass filter and the emission spectrum was measured between 300 and 400 nm with 1 nm stepping and a 2 nm band-pass filter.

15N-1H HSQC NMR. SH3 grown in M9 minimal media supplemented with 15N NH₄Cl was transferred into pH 6.0 phosphate, 100 mM NaCl and concentrated to 1 mM. NMR data were collected on a Bruker DMX-600 spectrometer equipped with a 5-mm x,y,z-shielded pulse-field gradient triple-resonance probe (Bruker). Experiments were performed at 20 degrees Celsius. Assignments were based upon published c-Src SH3 chemical shifts (175). 1H-15N HSQC were acquired and analyzed in the Sparky software suite. Changes in chemical shift were calculated according to the relationship dCS = sqrt(dH²+(dN/5)²).

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