Rationally Designed Integrin β3 Mutants Stabilized in the High Affinity Conformation*

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Integrins are important cell surface receptors that transmit bidirectional signals across the membrane. It has been shown that a conformational change of the integrin β-subunit headpiece (i.e. the β I domain and the hybrid domain) plays a critical role in regulating integrin ligand binding affinity and function. Previous studies have used coarse methods (a glycan wedge, mutations in transmembrane contacts) to force the β-subunit into either the open or closed conformation. Here, we demonstrate a detailed understanding of this conformational change by applying computational design techniques to select five amino acid side chains that play an important role in the energetic balance between the open and closed conformations of αIIbβ3. Eight single-point mutants were designed at these sites, of which five bound ligands much better than wild type. Further, these mutants were found to be in a more extended conformation than wild type, suggesting that the conformational change at the ligand binding headpiece was propagated to the legs of the integrin. This detailed understanding of the conformational change will assist in the development of allosteric drugs that either stabilize or destabilize specific integrin conformations without occluding the ligand-binding site.

Allostery is important in the function of many signaling proteins (1), including cell adhesion molecules, such as integrins (2). Integrins are large, heterodimeric molecules that transmit signals bidirectionally across the plasma membrane and regulate many biological functions, including wound healing, cell differentiation, and cell migration. The conformational changes associated with integrin activation and signaling have been studied structurally and functionally (3–8). Integrins bind ligands at an interface between the α-subunit β-propeller domain and the β-subunit 1 domain in the integrin headpiece (2). An acidic residue in the ligand coordinates with a Mg2+ ion in a metal ion-dependent adhesion site (MIDAS).5 Remodeling of ligand-binding residues in the β I domain is allosterically linked to reorientation at its interface with the hybrid domain.

Crystal structures of integrins have revealed open, liganded (8) and closed, unliganded (3) conformations of the integrin headpiece (Fig. 1). Movement of the β1-α1 and β6-α7 loops, which bind the MIDAS and ADMIDAS (adjacent to MIDAS) metal ions are coupled to movements of the α1 and α7 helices, which are adjacent to one another. Reshaping to the open conformation, which exhibits high affinity for ligand, is allosterically linked to C-terminal piston-like movement of the α7-helix. This linkage is critical for bidirectional propagation of conformational signals between the ligand binding pocket and other integrin domains. The orientation between the β I and hybrid domains appears to represent the critical “translator” for conveying large scale interdomain rearrangements into local conformational changes within the β I domain that regulate affinity for ligand. The piston-like displacement of the α7-helix of the β I domain in the “open” crystal structure results in complete remodeling of the interface with the hybrid domain (8). Relative to the closed conformation, the hybrid domain swings out about 60°, resulting in a 70-Å displacement of the β-subunit knee away from the α-subunit knee.

In the bent integrin conformation, the headpiece is in the closed conformation. After a switchblade-like extension at the integrin knees, the headpiece is found in both closed and open conformations (2). Extensive interfaces between the integrin headpiece and lower legs in the bent conformation are broken both by integrin extension and by hybrid domain swing-out; therefore, headpiece opening is less energetically costly in the extended than in the bent conformation. Conversely, headpiece opening favors integrin extension. This provides a mechanism in integrins for communicating activation signals between the ligand binding site in the headpiece and cytoskeletal protein-binding sites in the α and β-subunit cytoplasmic domains.

Characterization of integrin variants has provided strong evidence linking the “open” state observed in crystal structures, the “extended” integrin morphology observed via negative staining electron microscopy, and the “high affinity for ligand” state observed in cell adhesion assays. One such study introduced an N-glycosylation site at the most acute region of the

5 The abbreviations used are: MIDAS, metal ion-dependent adhesion site; SASA, solvent-accessible surface area; CHO, Chinese hamster ovary; mAb, monoclonal antibody; FITC, fluorescein isothiocyanate; MFI, mean fluorescence intensity; LIBS, ligand-induced binding site.
interface between the β I domain and the hybrid domain. The resulting glycan “wedge,” designed to shift the conformational balance toward the open state, was indeed found to increase ligand binding affinity (9, 10). This study laid the groundwork for understanding the relationship of the hinge between the open and closed conformation beyond the resolution afforded by the glycan wedge. We explain these differences. Pro85 was originally selected as a site for mutation but has been discarded from the current study (8). In all cases, the range of residues considered was Pro57–Asp432. After the mutational studies were completed, five of these open structures were rerefined, and four further ones were reported: Protein Data Bank codes 2vc2, 2vdk, 2vdl, 2vdm, 2vdo, 2vdp, 2vdq, and 2vdr (13). Moreover, a higher resolution and better refined closed structure of αIIbβ3 was reported.6 The environment of each selected residue was computed retrospectively for these structures as well, by using only the newer or rerefined structures. For cases in which more than one subunit was present in the asymmetric unit, calculations were averaged over all independent subunits. This retrospective analysis shows no qualitative differences to the data collected using the smaller set of structures (Table 1), except for the quality of packing around Leu138 and Glu206 (the new open αIIbβ3 structures exhibit better packing at these positions than the new closed structures). The electron density around these two residue environments is not well resolved in the newer closed structures, however, which may explain these differences. Pro85 was originally selected as a site for mutation but has been discarded from the current study because it was assigned as cis in open αIIbβ3 and trans in closed α5β3 but subsequently cis in closed αIIbβ3; furthermore, the mutation of this cis-Pro to Gly resulted in a lack of expression (data not shown). The side chain solvent-accessible surface area (SASA) was computed using a 1.4-Å probe to search for sites buried in the closed state but exposed in the open state.

The quality of packing of each residue was additionally assessed via “SASAPack.” This measure involves computing the side chain-accessible surface area using a 0.5-Å probe and comparing the value to the average value for side chains with similar SASA (computed using a 1.4-Å probe). This metric has proven effective at detecting packing defects, since holes lead to an increase in the accessible surface area available to a small probe.7

**Plasmid Construction, Expression, and Immunoprecipitation**—Plasmids coding for full-length human αIIb and β3 were subcloned into pEF/V5-HisA and pcDNA3.1/Myc-His (+), respectively (5). Single-residue substitutions of β3 were carried out using site-directed mutagenesis. Constructs were transfected into CHO-K1 cells (American Type Culture Collection) using a Fugene transfection kit (Roche Applied Science) according to

6 Zhu, J., Luo, B. H., Xiao, T., Zhang, C., Nishida, N., and Springer, T. A. (2008) Mol. Cell 32, 849–861.
7 P. Bradley, personal communication.
the manufacturer’s instructions. Stably transfected CHO cells and β3 were detected by flow cytometry staining with the following monoclonal antibodies: 10E5 (anti-αIIb mAb, kindly provided by B. S. Coller, The Rockefeller University, New York, NY) (15), 7E3 (anti-β3 mAb), and AP3 (nonfunctional anti-β3 mAb, American Type Culture Collection), respectively.

**Soluble Ligand Binding**—The activating anti-αIIb mAb PT25-2 was a generous gift from M. Handa (Keio University Hospital, Tokyo, Japan) (16). Soluble binding of FITC-labeled human fibrinogen (Enzyme Research Laboratories, South Bend, IN) and ligand mimetic IgM PAC-1 (BD Biosciences) was determined as described (9). Briefly, transfected cells suspended in 20 mM Hepes-buffered saline (pH 7.4) supplemented with 5.5 mM glucose and 1% bovine serum albumin were incubated with FITC-conjugated human fibrinogen or PAC-1 in the presence of either 5 mM EDTA; 1 mM Ca<sup>2+</sup>, 1 mM Mg<sup>2+</sup>, 1 mM Mn<sup>2+</sup>; 1 mM Ca<sup>2+</sup>, 1 mM Mg<sup>2+</sup> plus 10 μg/ml activating mAb PT25-2; or 1 mM Mn<sup>2+</sup> plus 10 μg/ml PT25-2 at room temperature for 30 min. For PAC-1 binding, cells were washed and stained with FITC-conjugated anti-mouse IgM on ice for another 30 min before being subjected to flow cytometry. Cells were also stained in parallel with anti-β3 mAb AP3 followed by FITC-conjugated anti-mouse IgG. Binding activity is presented as the percentage of MFI of anti-LIBS antibody after subtraction of MFI of the control X63 IgG, relative to MFI of the conformation-independent anti-β3 mAb AP3 after subtraction of the same control.

**RESULTS**

Rationale for Mutations to Stabilize β3 Integrin in the Open State—Initially, conventional computational design methodologies (18) were applied to collect a series of mutations predicted to be stabilizing for each of several open β3 state integrin structures. Incorporation of a predicted stabilizing mutation from one open state structure onto a different open state structure, however, seldom proved stabilizing; this was an indication that the design methodology was too sensitive to structural details to be useful for this application. For this reason, mutations at each site were selected as per the rationale described below.

Computational design was based on multiple examples of β3 integrin structures in each of the closed and open conformations (Table 1). Changes in residue environments as measured by SASA or SASApack (see “Materials and Methods”) between closed and open structures were used to identify sites of mutation. Due to the moderate resolution of the crystal structures used (Bragg spacings for these structures range from 2.7 to 3.3 Å), emphasis was placed on disruption of specific stabilizing interactions in the closed conformation rather than design of new stabilizing interactions in the open conformation. Residues near the fibrinogen-binding site were not considered.

**Ligand-induced Binding Site (LIBS) Epitope Expression**—Anti-LIBS mAb AP5, LIBS1, and D3 were kindly provided by M. H. Ginsberg (University of California San Diego, La Jolla, CA) and L. K. Jennings (University of Tennessee Health Science Center, Memphis, TN). LIBS epitope expression was determined as described previously (9). In brief, transfected cells suspended in Hepes-buffered saline supplemented with 5.5 mM glucose and 1% bovine serum albumin were incubated with or without 50 μM Gly-Arg-Gly-Asp-Ser-Pro peptide (GRGDSP) in the presence of 1 mM Ca<sup>2+</sup> plus 1 mM Mg<sup>2+</sup> and 10 μg/ml anti-LIBS antibodies or control X63 IgG. After incubation at room temperature for 30 min, cells were washed and stained with FITC-labeled anti-mouse IgG on ice for 30 min. The stained cells were subjected to flow cytometry, and LIBS epitope expression was expressed as the percentage of MFI of anti-LIBS antibody after subtraction of MFI of the control X63 IgG, relative to MFI of the conformation-independent anti-β3 mAb AP3 after subtraction of the same control.

### Table 1

| Quantifier | Open state structures used in designing mutants (6) | Closed state structures used in designing mutants (4) | Current open state structures (9) | Current closed state structures (5) |
|------------|-----------------------------------------------------|------------------------------------------------------|----------------------------------|-----------------------------------|
| SASA (Å<sup>2</sup>) | Leu<sup>1B8</sup>: 0.0 | Leu<sup>1B8</sup>: 1.3 | Leu<sup>1B8</sup>: 0.0 | Leu<sup>1B8</sup>: 1.7 |
| | Glu<sup>20E</sup>: 47.3 | Glu<sup>20E</sup>: 26.7 | Glu<sup>20E</sup>: 53.1 | Glu<sup>20E</sup>: 25.0 |
| | Ser<sup>24E</sup>: 41.4 | Ser<sup>24E</sup>: 9.1 | Ser<sup>24E</sup>: 39.4 | Ser<sup>24E</sup>: 6.6 |
| | Arg<sup>352</sup>: 99.9 | Arg<sup>352</sup>: 2.7 | Arg<sup>352</sup>: 102.3 | Arg<sup>352</sup>: 3.7 |
| | Lys<sup>417</sup>: 38.0 | Lys<sup>417</sup>: 9.4 | Lys<sup>417</sup>: 33.7 | Lys<sup>417</sup>: 10.7 |

**SASApack (dimensionless)**

| Quantifier | Open state structures used in designing mutants (6) | Closed state structures used in designing mutants (4) | Current open state structures (9) | Current closed state structures (5) |
|------------|-----------------------------------------------------|------------------------------------------------------|----------------------------------|-----------------------------------|
| | Leu<sup>1B8</sup>: 9.7 | Leu<sup>1B8</sup>: 0.0 | Leu<sup>1B8</sup>: 3.65 | Leu<sup>1B8</sup>: 0.3 |
| | Glu<sup>20E</sup>: 4.1 | Glu<sup>20E</sup>: 1.5 | Glu<sup>20E</sup>: 0.87 | Glu<sup>20E</sup>: 1.5 |
| | Ser<sup>24E</sup>: 2.5 | Ser<sup>24E</sup>: 10.2 | Ser<sup>24E</sup>: 2.13 | Ser<sup>24E</sup>: 10.2 |
| | Arg<sup>352</sup>: 0.3 | Arg<sup>352</sup>: 5.0 | Arg<sup>352</sup>: 0.22 | Arg<sup>352</sup>: 7.3 |
| | Lys<sup>417</sup>: 0.8 | Lys<sup>417</sup>: 2.9 | Lys<sup>417</sup>: 8.42 | Lys<sup>417</sup>: 1.0 |
destabilize the closed state with minimal disruption of the open state. In the closed state, this partially buried serine side chain faces an area of this side chain near the hinge region was also found to increase upon switching from the closed to open state (Table 1). A detailed examination of the environment of each mutated residue, in the closed (a, c, e, g, and i) and open (b, d, f, h, and j) conformations. Mutated residues were as follows: Arg352 (a and b), Ser243 (c and d), Leu138 (e and f), Lys417 (g and h), and Glu206 (i and j). This figure was generated using PyMOL (14).

occluded from solvent and forms a series of hydrogen bonds (Fig. 2a). After the switch to the open state, however, the α7-helix unwinds just enough that this side chain becomes largely exposed and forms a single surface-exposed hydrogen bond that caps the C terminus of the α7-helix (Fig. 2b). In order to destabilize the closed state with minimal disruption of the open state, a glutamate was introduced at this position.

Ser243 (β1-Hybrid Interface)—The solvent-accessible surface area of this side chain near the hinge region was also found to increase upon switching from the closed to open state (Table 1). In the closed state, this partially buried serine side chain faces the α7-helix and occupies a small cavity below the plane of the side chain of Arg352 (described above) (Fig. 2c). This cavity is sterically bounded by Tyr110 and Phe421, shown in the foreground of Fig. 2c (in stick representation). The relocation of the α7-helix and release of Arg352 associated with the transition to the open conformation, meanwhile, expose this side chain to solvent (Fig. 2d). Additionally, Tyr110 and Phe421 have shifted to where they are no longer visible in the view presented (Fig. 2d). In order to take advantage of the tight steric constraints of the low affinity state, aspartate and glutamate were each introduced in place of this serine.

Leu138 (β1 Domain)—The SASApack value computed for this leucine side chain in the closed conformation, about zero, showed that this residue was packed in an environment similar to that of an average leucine in the Protein Data Bank (Table 1). After the transition to the open conformation, however, this side chain was much less well packed than the Protein Data Bank average for leucine (Table 1). Examination of the surrounding side chains shows that packing considerations are responsible for this difference. In the closed state, this inward facing leucine side chain fits snugly into a pocket that exactly complements the shape of the side chain (Fig. 2e). Upon transition to the open conformation, however, the α1-helix slides relative to the rest of the β1 domain, such that this side chain fits into a pocket formed by a different set of side chains. This pocket exhibits much less shape complementarity toward the leucine side chain (Fig. 2f); in particular, a void remains near the leucine C-β atom, which would be filled by substitution of a β-branched amino acid at this position. To improve the packing of the open conformation while disrupting the packing of the closed conformation, this position was mutated to isoleucine.

Lys417 (Hybrid Domain)—A difference in SASA between the open and closed conformations was observed for this lysine side chain (Table 1), which is located on the outward facing side of the βF-strand of the hybrid domain. In the closed state, the βF-βG loop adopts a noncanonical conformation, allowing it to pack against the α5-helix of the β1 domain. The backbone carbonyl groups in this loop face inward, allowing one of these to form a hydrogen bond to the side chain of Lys417. This lysine side chain also forms a hydrogen bond to Asn203, which in turn acts as a “capping residue” on the C-terminal end of the α5-helix (Fig. 2g).

The hinge motion involved in the transition to the open conformation results in a complete loss of the interactions between the βF-βG loop and the β1 domain, exposing the outward facing side chains on the βF-strand of the hybrid domain (including Lys417). The conformation of the βF-βG loop relaxes, allowing the backbone to form hydrogen bonds to solvent (Fig. 2h). Given the relative unimportance of the detailed interactions in the open conformation (further supported by the higher crystallographic B-factors in this state), “charge reversal” mutations of Lys417 to aspartate and glutamate were introduced to disrupt the hydrogen bonds stabilizing the βF-βG loop in the closed conformation.

Glu206 (β1 Domain)—Although the most obvious structural rearrangement of the β1 domain takes place in the α1- and α7-helices, a very subtle twist occurs in the α2-helix as well (8). Comparison of the quality of packing in both states via SASApack (see “Materials and Methods”) brought to light a subtle difference in this region (Table 1).
transfectants. In all experiments measuring binding of the ligand-mimetic antibody PAC-1 or soluble fibrinogen, binding was expressed relative to αIIbβ3 surface expression to correct for differing amounts of cell surface expression by the mutants. In the presence of Ca\(^{2+}\), wild type αIIbβ3 bound very little ligand-mimetic PAC-1 antibody, whereas the presence of Mn\(^{2+}\) and activating mAb PT25-2 greatly increased binding (Fig. 3B). In contrast to wild type, five of the seven mutants, L138I, E206T, S243E, K417E, and K417D, bound PAC-1 in Ca\(^{2+}\) (Fig. 3B). Binding in Ca\(^{2+}\) was comparable with that in activating conditions and with activated wild-type.

We have previously observed that mutations that stabilize integrins in the high affinity, extended conformation lead to decreased cell surface expression, presumably because the bent conformation is more efficiently processed and transported to the cell surface (9). Consistent with the tendency of activating mutations to decrease surface expression, three of five activating mutations, E206T, K417E, and K417D, decreased surface expression, whereas the two nonactivating mutations, E206I and S243D, had no effect on surface expression (Fig. 3). However, the L138I and S243E mutations were activating while having no or little effect on surface expression.

To further characterize the five high affinity mutants, stable CHO-K1 cell transfectants were established, and clones with similar expression of wild-type and mutant αIIbβ3 were selected. Soluble PAC-1 (Fig. 4A) and fibrinogen (Fig. 4B) binding showed that wild type αIIbβ3 bound ligands only when stimulated by Mn\(^{2+}\) and/or PT25-2. By contrast, the five mutants bound ligands with significant higher capability in the presence of Ca\(^{2+}\), confirming that all five mutations activated integrins for ligand binding (Fig. 4, A and B). In the presence of Mn\(^{2+}\) and PT25-2, all mutations and wild-type bound ligands maximally.

The affinity state of the wild type and mutants was further assessed by cell adhesion assays on immobilized fibrinogen. In contrast to soluble ligand binding, at coating concentrations above 5 μg/ml, wild type cells mediated efficient adhesion to immobilized fibrinogen even in the absence of activation (Fig. 4C). All mutant cells adhered to immobilized fibrinogen at lower coating concentrations than wild-type, and two mutants, K417E and K417D, even adhered to the substrate at 1 μg/ml coating concentrations.

**Mutations That Enhanced Ligand Binding Stabilized Integrins in the Extended Conformation**—Activation and/or ligand binding change the conformation of αIIbβ3, resulting in the exposure of neoepitopes called LIBS. We investigated the conformation of the mutants with three different LIBS antibodies, which bind to various sites in the β3-subunit (Fig. 5). Under basal conditions in Ca\(^{2+}\), all mutants showed increased expression of the D3 (anti-β3 residues 422–490) (19) and LIBS1 (anti-β3 residues 422–690) (20) epitopes; and, with the exception of S243E, showed increased expression of the AP5 (anti-β3 residues 1–5) (20) epitope. Thus, the equilibrium is shifted to a more extended conformation in the five mutant receptors with higher affinity for ligand.

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Although the aliphatic portion of the Glu\(^{206}\) side chain packs snugly against Val\(^{193}\) and Leu\(^{194}\) in the closed conformation (Fig. 2i), this interaction is lost in the open conformation (Fig. 2j). This small decrease in packing density led to the hypothesis that introduction of a β-branched amino acid in place of this glutamate could be accommodated in the open conformation but would prove greatly destabilizing to the closed conformation. Mutations to isoleucine and threonine were therefore selected. Soluble PAC-1 binding to 293T transfectants was determined in the presence of 5 mM Ca\(^{2+}\) (open bars) or 1 mM Mn\(^{2+}\) plus 10 μg/ml mAb PT25-2 (black bars) and is expressed as the MFI of PAC-1 staining as a percentage of MFI of staining with AP3. Data are mean and S.D. of three independent experiments.

**Expression and soluble ligand binding of integrin αIIbβ3 on 293T cells.** A, 293T transient transfectants were immunofluorescently stained with mAbs 7E3 (open bars), AP3 (gray bars), and 10E5 (black bars). B, soluble PAC-1 binding to 293T transfectants was determined in the presence of 5 mM Ca\(^{2+}\) (open bars) or 1 mM Mn\(^{2+}\) plus 10 μg/ml mAb PT25-2 (black bars) and is expressed as the MFI of PAC-1 staining as a percentage of MFI of staining with AP3. Data are mean and S.D. of three independent experiments.

**Expression of the Mutant αIIbβ3 in 293T Cells**—To determine whether these mutant β3 integrins could be expressed with the αIIb-subunit, we transiently transfected the eight αIIbβ3 mutants into 293T cells. Two anti-β3 antibodies, 7E3 and AP3, and an αβ complex-dependent anti-αIIb antibody, 10E5, were used to monitor expression on the cell surface. All three antibodies gave similar results (Fig. 3A). Three mutants (L138I, E206I, and S243D) were expressed at wild-type levels. Four mutants (E206T, S243E, K417E, and K417D) were expressed at levels lower than that of the wild type. The remaining mutant, R352E, was not expressed at detectable levels.

**Five Mutants Have Enhanced Ligand Binding Affinity**—The seven expressed mutants were tested for ligand binding in 293T
DISCUSSION

We computationally designed mutations that energetically favor the open, high affinity conformation of the β3 integrin headpiece. Five of the eight designed mutations bound ligands much better than wild type. More interestingly, these mutants were found to be in the more extended conformation than wild type, suggesting that the conformational change at the ligand binding headpiece was propagated to the legs of the integrin.

Rationalization of Unsuccessful Mutations—As noted earlier, locking the integrins into the open conformation through the introduction of a glycan wedge in the interface region led to a significant decrease in expression levels (9). It is therefore not entirely surprising that one of the mutations that may be expected to most destabilize the closed conformation (R352E) essentially abolished expression.

Unlike this unsuccessful design, however, two of the mutants, S243D and E206I, were expressed but failed to show a significant increase of ligand binding capability. For-tuitously, another mutation tested at each of these sites showed increased ligand-binding; this allows for a direct retrospective comparison.

In the case of substitutions at Ser243, RosettaDesign (21) was used to build both Asp and Glu side chains at this position of each crystal structure, keeping the backbones fixed. Examination of these structural models (Fig. 6) suggests an explanation for the different behavior of these two mutants.

In the closed conformation, both Asp and Glu may be sterically accommodated in place of Ser243; in both cases, however, the geometry of the substituted side chain is not suitable to form a hydrogen bond, leaving this charged group unsatisfied (Fig. 6,a and c). The models of the open conformation, however, show a marked difference between Asp and Glu. Although the aliphatic part of the glutamate side chain is long enough that the carboxyl group may be fully exposed to solvent (Fig. 6b), this is not true of the aspartate side chain; the Asp side chain is buried by its environment in a manner that precludes formation of a hydrogen bond in the open conformation (Fig. 6d).

Based on these models, we therefore conclude that introduction of the S243D mutation destabilizes both the open and closed conformations to a similar extent, leading to ligand affinity comparable with that of the wild type. By contrast, S243E...
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Whereas the possible design strategy of relieving strain specific to the open conformation did not seem promising, a related approach, designing new favorable interactions specific to the open conformation, might also be expected to fail for the same reasons. Such an approach has recently been used in the thermostabilization of an enzyme (18), demonstrating that function can be maintained after incorporation of new stabilizing interactions, but the structure of the target protein used in this study was available at a resolution of 1.14 Å (24).

Here, we have demonstrated that crystal structures at about 3 Å resolution are sufficient for rationally designing mutations that will preferentially destabilize one conformation, by using “robust” measures to select sites for mutation. This, in turn, raises the interesting possibility of using high quality homology models as the basis for selecting sites for mutation. The usefulness as a search model for solving the crystallographic “phase problem” via molecular replacement represents a very stringent test of the accuracy of a homology model (25). Recent progress in structure prediction has greatly loosened the requirement of sequence similarity for building models of sufficient accuracy for molecular replacement (26). This, together with design strategies that specifically focus on yielding robust results from structures at relatively poor resolution, offers the tantalizing prospect for rational manipulation of protein states for which high quality crystal structures are not yet available.

Mutational Studies of Mechanism of Allosteric Switch—Interestingly, the five mutants with enhanced ligand binding affinity were found to be in a conformation more extended than wild type, suggesting that the conformational change at the ligand binding headpiece was propagated to the integrin legs. Previously, we showed that introducing a bulky glycan group to the interface between the β1 or β3 I domain and hybrid domain caused integrins to adopt a high affinity and extended conformation (9, 10). Other studies showed that mutations introduced in the β1 integrin I domain increased ligand binding and shifted the equilibrium toward more extended conformation (27). However, introducing disulfides to the β3 I domain C-terminal α7-helix or deleting four residues of the β2 I domain α7-helix increased integrin ligand binding, whereas the overall conformation of the receptors was still in the bent conformation unless ligands were added (28, 29), probably because these mutations circumvented allosteric communication of the β3 I domain through its α7-helix with the hybrid domain (30). Here, our studies demonstrate that single-residue substitutions both in the β3 I domain and in the hybrid domain can stabilize this integrin in the high affinity, extended conformation. The results suggest that integrins are pliable to environmental change; subtle changes in the structure, either by mutation or by binding of other proteins, can readily shift the equilibrium from one conformation to the other.

Preferential stabilization of one integrin conformation represents an attractive method for understanding integrin function; not surprisingly, several approaches have been used to engineer integrins locked into either the high or low affinity state.

Rationally designed amino acid substitutions have previously been applied to selectively stabilize both the open and closed states of a different integrin, Mac-1 (αMβ2, or CD11b/CD18).

A similar structural explanation may underlie the difference in the observed behavior of the E206I and E206T mutants. Although the E206T mutant demonstrates that disruption at this site is indeed a viable method for enhancing ligand binding affinity, isoleucine was not a good choice at this position. The energetic details underlying the differences between E206I and E206T are more complex, however, stemming from the chemical as well as steric differences between isoleucine and threonine.

Strategy for Selection of Mutation Sites—Although not all of the mutations led to the desired increase in ligand binding affinity, each site identified from the computational comparison led to at least one mutant with altered behavior. This underscores the robustness of the strategy of selecting mutation sites based on differences in SASA and packing. This robust approach was developed specifically to cater to this design problem because of the relatively poor resolution of crystal structures of each state (Bragg spacings for the structures of the closed conformation range from 3.1 to 3.3 Å and from 2.7 to 3.1 Å for the structures of the open conformation). This had two implications in selecting a design strategy.

First, it has long been known that crystal structures of poor resolution suffer larger deviations in torsion angles (22) as well as irregular packing (23). Since both of these features are captured in the full RosettaDesign energy function (21), it was not clear that residues scoring poorly with this energy function were indeed subject to energetic strain; rather, observed differences between the two conformations could be due solely to slight inaccuracies in atomic positions. For this reason, assessment of residue environments focused on using features that would be more robust toward such inaccuracies, such as solvent-accessible surface area. The use of six structures to represent the open conformation was an additional measure to alleviate these concerns.
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Several single- and double-point mutants in the α-I domain have been described (31–33), which lead to modest increases in the ligand binding affinity (34). Much larger effects on ligand binding affinity were obtained through the complete computational redesign of the α-I domain core (35). By using crystal structures of the open and closed conformations of this domain, redesigned domains were proposed and tested. These redesigned domains, which required four mutations to stabilize the closed conformation and 8–13 mutations to stabilize the open conformation, induced a stronger shift toward the desired conformation than a single-point mutant included for comparison (35).

Although it is expected to be generally true that the complete redesign of a protein core is likely to induce a larger energetic shift than most single-point mutants, such an approach was not applied to αIIbβ3, for two reasons. First, the crystal structures of the Mac-1 α-I domain were available at better resolution than the structures of the β3 headpiece. Second, carefully chosen single-point mutants are more likely to preserve subtle details of the structure, which can be important for maintaining long range propagation of the conformational changes as well as binding surfaces for other proteins. Both of these are preserved in the series of mutations presented here, evidenced by binding of anti-LIBS antibodies and the fact that mutations in the hybrid domain affect ligand binding in the β I domain.

CONCLUSIONS

Integrins represent attractive pharmaceutical targets for a variety of human diseases; abciximab, eptifibatide, and tirofiban all target integrin αIIbβ3 and have been approved by the United States Food and Drug Administration for the treatment of thrombosis (36, 37). Whereas these compounds block the interaction of integrins with the receptor, the conformational changes they induce can themselves initiate unwanted signals. In fact, it has been shown that these three antagonists can induce thrombocytopenia in 1–5% of patients (37). Due to the importance of allostericity in regulating integrin function, it is critical to extend our detailed understanding of the relationship of integrin conformation with ligand binding.

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