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Integrin structures and conformational signaling

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Abstract
Integrins are cell adhesion molecules that play critical roles in development, wound healing, hemostasis, immunity and cancer. Advances in the past two years have shed light on the structural basis for integrin regulation and signaling, especially on how global conformational changes between bent and extended conformations relate to the inter-domain and intra-domain shape shifting that regulates affinity for ligand. The downward movements of the C-terminal helices of the αI and βI domains and the swing-out of the hybrid domain play pivotal roles in integrin conformational signaling. Experiments have also shown that integrins transmit bidirectional signals across the plasma membrane by coupling extracellular conformational change with an unclasping and separation of the α and β transmembrane and cytoplasmic domains.

Introduction
Integrins are cell adhesion molecules that mediate cell–cell, cell–extracellular matrix and cell–pathogen interactions. They transmit signals bidirectionally across the plasma membrane and regulate many biological functions, including wound healing, cell differentiation and cell migration. Integrins contain two non-covalently associated, type I transmembrane (TM) glycoprotein α and β subunits with large extracellular domains, single-spanning TM domains and short cytoplasmic domains (Figure 1a). The structures of the extracellular fragment of integrin αVβ3 revealed an unexpected, compact, V-shaped conformation, with each leg bent (Figure 1b) [1,2]. Recently, an increasing number of studies have together established that the bent conformation represents the physiological low-affinity state, whereas priming and ligand binding induce a large-scale conformational rearrangement in which the integrin extends with a ‘switchblade’-like motion (Figure 1b and c) [3-5,6••]. In this review, we focus on recent progress on how signals are communicated between the ligand binding domains and the plasma membrane at the molecular and atomic level.

Integrin ectodomain crystal structures
The integrin β-subunits contain very sophisticated domain insertions: the βI domain is inserted in the hybrid domain, which is in turn inserted in the PSI (for plexins, semaphorins, and integrins) domain (Figure 1a) [6••]. These domain insertions play a critical role in integrin signal transmission. The βI domain directly binds ligands in integrins that lack αI domains, and indirectly regulates ligand binding by integrins that contain αI domains. The structure of the βI domain was first solved in the context of αVβ3 extracellular domains in the absence of ligand [1]. The βI domain is structurally homologous to integrin α subunit I domains, which have been solved only as isolated domains, and are described in more detail below. In αI domains, rearrangements in loops surrounding the metal-ion-dependent adhesion site (MIDAS) increase affinity for ligand, and are linked to downward displacement of the α7-helix. Soaking of a ligand-mimetic Arg-Gly-Asp-containing cyclic peptide into the integrin...
αVβ3 crystals revealed that the Arg binds the αV β-propeller domain and the Asp binds a metal ion held in the MIDAS of the β3 I domain [2]. Movements of residues near the MIDAS in the β1-α1 loop, α1-helix, and β6-α7 loop were seen that enabled ligand binding in the closed state (Figure 2a). However, downward displacement of the α7-helix was not seen (Figure 2a), and it was therefore suggested that the α I and the β I domains are activated by distinct mechanisms [2]. However, subsequent mutagenesis studies [7-9,10,11] and the structure of the αIIbβ3 headpiece co-crystallized with different ligands [6••] revealed downward α7-helix displacement in the open, high-affinity state of the β I domain (Figure 2a), and marked structural similarity between α I and β I domain allostery.

The β3 subunit I, hybrid, and PSI domains from the closed, low-affinity unliganded αVβ3 structure, the closed, low-affinity liganded (ligand soaked) αVβ3 structure, and the open, high-affinity liganded (ligand co-crystallized) αIIbβ3 structure are compared in Figure 2a. The liganded, high-affinity αIIbβ3 headpiece structure enables atomic-level understanding of the mechanism of integrin activation [6••]. In the high-affinity, liganded β I domain compared with the low-affinity, unliganded β I domain, there are concerted movements of the β1-α1 and β6-α7 loops surrounding the ligand-binding pocket and of the α1 and α7 helices (Figure 2). Coordination of the Met335 backbone carbonyl in the β6-α7 loop to the ADMIDAS (adjacent to MIDAS) Ca²⁺ in the low-affinity, unliganded conformation is broken in the high-affinity, liganded conformation. This enables the ADMIDAS metal, and residues in the β1-α1 loop that coordinate to both the ADMIDAS and the MIDAS metals, to shift markedly, remodeling the ligand binding site and increasing affinity for ligand. Movements of the α1-helix, β6-α7 loop and α7-helix are tightly coupled, so that reshaping to the high-affinity ligand-binding site is allosterically linked to downward movement of the α7-helix. This linkage is critical for propagation of conformational signals from the ligand-binding pocket to the other integrin domains and vice versa (Figure 2a).

The orientation between the β I and hybrid domains appears to be the critical ‘translator’ converting global conformational change into local intradomain conformational changes that regulate affinity (Figure 2a). The piston-like displacement of the β I domain α7-helix in the high-affinity, liganded crystal structure results in complete remodeling of the interface between these domains, leading to the swing-out of the hybrid domain (Figure 1b, panel iii and Figure 2a) [6••]. Relative to the closed conformation, the hybrid domain swings out about 60°, causing the knees of the α and β subunits to separate by 70 Å

Structures of the β3 and β2 integrin PSI domains and β2 I-EGF1 domain [6••,12,13•] revised the connectivity of the previously identified long-range disulfide bond, which is now shown to link β3 Cys-13 to Cys-435 (or β2 Cys-11 to Cys-425). The structures show that this long-range disulfide bond occurs within the PSI domain, and therefore the hybrid domain is inserted in the PSI domain (Figure 1a). During the rearrangement of the headpiece between the closed and open conformations, there is no change in the hybrid/PSI domain interface. Therefore, this rigid interface, which is reinforced by the two polypeptide chain connections, nearby disulfide bonds and an Arg deeply buried in the interface, enables the PSI domain to amplify the leg separation triggered by the swing-out of the hybrid domain [6••,13•] (Figure 2a). The PSI and I-EGF1 domains are also shown to be intimately associated so that the hybrid and PSI/I-EGF1 domains move as a rigid unit [13•]. Some activating antibodies bind to the PSI domain and induce the high affinity state [14,15].

The knee of the β subunit is located between the PSI/I-EGF1 and I-EGF2 domains; the knee or genu of the α subunit is a small Ca²⁺-binding loop between the thigh and calf-1 domains. Work using an αL antibody that reports extension and maps to the inner face of the thigh domain, and which requires the genu and a Ca²⁺-coordinating residue donated by the calf-1
domain, suggests that integrin α-subunit extension occurs by movement of the thigh–genu linker (αV residues 594-595), a conclusion that is supported by structural inspection [16].

A large range of studies support the importance of integrin extension and hybrid-domain swing-out in integrin activation. Perhaps most definitive are the crystal structures of four independent examples of the αIIbβ3 head-piece, in two different crystal forms, with three different ligands or a pseudo ligand bound, all of which reveal similar hybrid domain swing-out [6••]. The swing-out of the hybrid domain necessitates the existence of the extended conformation, because the hybrid domain is central in the interfaces that are buried in the bent conformation; these interfaces are completely broken by hybrid domain swing-out [6••].

Integrin electron microscopy studies

Early electron microscopy (EM) studies revealed extended conformations. A later EM study revealed the bent conformation and showed it had low affinity for ligand and was stabilized by Ca2+ and close association of the α and β subunits near their junction with the membrane, and that activation with Mn2+ or breaking the juxtamembrane clasp favored extension [3]. In integrins on the cell surface, stabilizing the bent conformation with mutationally introduced disulfide bonds inhibits ligand binding [3]. Both closed and open headpiece conformations, i.e. with the hybrid domains swung in or out, respectively (Figure 1b), were seen in unclasped and Mn2+-treated extended integrins; however, cyclic Arg-Gly-Asp not only induced extension but also specifically stabilized the open headpiece conformation [3]. Recent studies with the I-domain-containing integrins αLβ2 and αXβ2 reveal the same three overall conformational states. Furthermore, Fab fragments of antibodies known to induce or report the active conformation bind exclusively to the extended conformation, with both open and closed headpiece conformations present, definitively establishing that extension is sufficient to activate integrins on the cell surface, and that physiologic agonists such as chemoattractants and agents such as phorbol myristate acetate and the talin head domain induce cell surface integrin extension [17••]. Electron tomography of negatively stained, active detergent-soluble αIIbβ3 purified on an Arg-Gly-Asp peptide affinity column reveals an extended conformation with >90% of particles showing an open headpiece structure that matches perfectly [18•] the open, liganded αIIbβ3 headpiece crystal structure [6••].

Two notable studies differ in their conclusions from those described above. Cryo-EM reconstructions of detergent-soluble αIIbβ3 molecules revealed a conformation that is intermediate between bent and extended conformations [19]. However, for particles the size of integrins, cryo-EM cannot distinguish between a particle in two different orientations or two different conformations [20]. Since preparations of integrins, including αIIbβ3, often contain a mixture of particles with different conformations [3], the intermediate αIIbβ3 conformation may have resulted from averaging together particles in extended and bent conformations. Other observations support this speculation, since to fit an atomic model into the intermediate αIIbβ3 EM density, marked changes in orientation at the β-propeller/thigh interface and calf-1/calf-2 interface were required [19] that are inconsistent with recent EM studies [3,4,17••,18•].

A negative stain study of integrin αVβ3 in 0.2 mM Mn2+ or 0.2 mM Mn2+ with a fibronectin fragment revealed a bent conformation [21], whereas a study of αVβ3 with 1mM Mn2+, or cyclic Arg-Gly-Asp in the presence of either 1 mM Mn2+ or 5 mM Ca2+, revealed predominantly extended conformations [3]. These differences might reflect the different ligands or Mn2+ concentrations used. Another important difference is the extensive aggregation present in the one field view shown in Adair et al. [21] but not in the eight field views shown in Takagi et al. [3], which led Adair et al. [21] to state that, “We cannot exclude the possibility that unsampled regions on the grid might have preferentially arisen from aggregated extended forms.”
It is interesting to note the differing methodologies employed in 3D reconstructions of negatively stained integrins [20,22]. In the random conical tilt [4] and tomography studies [18•], each particle was imaged at two or 23 different tilt angles, respectively, and 3D electron density maps were then computed independently of any crystal structure information. The EM density showed excellent agreement with crystal structures that were currently [4,18•] or only subsequently [4] available. In the angular reconstitution study [21], each particle was imaged at a single angle. Reconstruction used resolution-filtered crystal structures as starting models, and particles were automatically selected for use in reconstruction if they were similar to 2D projections of these models. The final models are similar to the starting atomic models, except that in the liganded model several of the fibronectin domains are no longer present and density for I-EGF domains 1 and 2, which was absent from the starting model, was acquired; however, in the unliganded model, density for I-EGF domains 1 and 2 remains missing in the final model.

Other studies on ectodomain conformation

Aside from structural work [3,4,6•,17•,18•,23] integrin extension and hybrid domain swing-out are supported by a wide range of other studies. Stabilizing the open head-piece by mutationally introducing an N-glycosylation site into the hybrid-β I domain interface increases ligand-binding affinity [24,25]. An allosteric β1 antibody that inhibits ligand binding has been shown by epitope mapping and EM to restrict the swing-out of the hybrid domain [25]. The functional properties of a β2 mAb suggest it also inhibits by blocking signal transmission at the β I-hybrid domain interface [26]. Activation-dependent mAbs that map to the inner face of the hybrid domain support conformational change between the β I and hybrid domains [9,27]. Epitope exposure suggests that ligand binding and a mutation of the β I domain α7 helix that stabilizes the high affinity state induce hybrid domain swing-out, confirming the relationship between movement of the α7 helix and hybrid domain swing-out [9]. Integrin extension on the cell surface was confirmed by studies using fluorescence resonance energy transfer (FRET) between fluorescent ligand-bound integrins and lipophilic probes [28].

As an alternative or supplement to integrin extension and hybrid domain swing-out, a ‘deadbolt’ model has been proposed in which interaction at a very small 60 Å² interface between the β-tail domain CD loop (the dead-bolt) and the β I domain α7 helix is critical for stabilizing integrins in the low affinity state [29]. Since hybrid domain swing-out requires β6-α7 loop and α7-helix displacement, mutation of the β-tail domain CD loop is required to test this model. We found that deleting β3 integrin CD loop residues 672–674 or mutating these residues to Ala has no effect on ligand binding or activation epitope exposure by integrins αIIbβ3 and αVβ3 (our unpublished data). Therefore, the β-tail CD loop does not function as a deadbolt.

Integrins containing an α I domain

Compared to the integrins lacking an α I domain, conformational regulation of integrins containing an α I domain requires the additional step of transmission of allostery from the β I domain to the α I domain (Figure 1c). Crystal structures of α I domains reveal three distinct conformations, namely closed, intermediate and open [30,31]. They differ not only in the coordination of residues with the MIDAS, but in the structure of surrounding loops and in the positions of the β6-α7 loop and α1 and α7 helices (Figure 2b). Introducing pairs of cysteines to stabilize the β6-α7 loop in the intermediate and open conformations led to 500- and 10,000-fold higher affinity to ICAM-1, respectively [31]. Molecular dynamic studies showed that the intermediate conformation was on the pathway from the closed to the open conformation of the αL and αM I domains, but not the α1 and α2 I domains [32]. The study provides strong support for the idea that the intermediate conformation could be of physiologic importance for fine regulation of integrin affinity.
EM studies of αXβ2 and αLβ2 integrins reveal no activation-dependent change in α I domain orientation relative to the β propeller domain analogous to that observed between the β I domain and hybrid domain [17••]. This is consistent with a proposal that the α I domain α7-helix transmits allostery between the α I MIDAS and the β I MIDAS. That is, it is proposed that in the active state, downward movement of the α I domain α7-helix enables an invariant Glu residue that is present a few residues after the α7-helix to act as an ‘intrinsic ligand’ and engage the β I MIDAS [33,34•]. Yang et al. showed that individual mutation of the αL linker residue Glu-310 or β2 MIDAS residues Ala-210 or Tyr-115 to cysteine abolishes I domain activation, whereas the double mutations of αL-E310C with either β2-A210C or β2-Y115C form disulfide bonds that constitutively activate ligand binding [34•]. The activation effect of the disulfide mutant is susceptible to small molecule antagonists that bind underneath the I domain α7-helix and certain allosteric antagonistic antibodies. This study provides direct evidence for an activating interaction between αL residue Glu-310 and the β2 MIDAS (Figure 1c, panel iii), and suggests that the α7-helix and its linker are better modeled as a pull spring than a bell rope [34•].

Conformational change in integrin cytoplasmic and transmembrane domains

Recently, the basis for integrin activation across the plasma membrane has also been studied. Separation of integrin legs results in integrin activation [5,35,36,37•], suggesting that association of the integrin TM and cytoplasmic domains stabilizes integrins in the low affinity state. FRET shows that in the resting state the integrin α and β subunit cytoplasmic domains are close to one another, but undergo significant spatial separation upon inside-out activation induced by G-protein-coupled receptors, phorbol ester or talin head domain, or upon outside-in signaling induced by ligand binding [5]. NMR studies of the integrin cytoplasmic tails suggest that their association is weak, with significant differences between published structures of associated cytoplasmic domains [38•,39,40], or with association between α and β subunit cytoplasmic domains being undetectable [41]. How the talin head domain and filamin bind to the integrin β cytoplasmic domain and activate integrins has been revealed by NMR structures [42-45].

Disulfide scanning of the exofacial portions of the TM domains showed a specific α-helical interface between the α and β? TM domains in the resting state [37•]. The two TM domains separate rather than rearrange after activation of integrins from inside the cell. Introduction of disulfide bridges to prevent or reverse separation abolished the activating effect of cytoplasmic mutations [37•]. Several other mutagenesis studies also suggest that a specific interface stabilizes integrins in the resting state [46,47,48•]. Modeling of the integrin TM domain interface, with or without experimental data, has resulted in models with different interfaces [37•,48•,49,50]. Further experimental data and more comprehensive analysis are required. Homomeric TM domain association following heterodimeric TM dissociation has been proposed [51], but in subsequent studies it has been shown that this does not occur as a consequence of integrin activation by α and β subunit TM separation, although it might occur after binding to multivalent ligands [47,52]. Thus, numerous studies from different labs suggest that integrin bidirectional signaling across the plasma membrane is accomplished by coupling extracellular conformational change to an unclasping and separation of the α and β TM and cytoplasmic domains.

Conclusions

Recent structural, biochemical and biophysical studies have greatly advanced our understanding of the mechanisms underlying integrin bidirectional signaling across the plasma membrane. We should always consider that integrins are in dynamic equilibrium among many different conformational states, rather than locked in one specific state. As reviewed above,
intracellular signals and ligand binding act by shifting the equilibrium and altering the population of the different conformational states. It will be of great interest to use biophysical methods to probe the dynamics of integrin signaling under physiological conditions.

Acknowledgements

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References and recommended reading

Papers of particular interest, published within the 2-year period of review, have been highlighted as:

• of special interest
•• of outstanding interest

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Figure 1.
Integrin architecture and conformational changes associated with affinity regulation. (a) Organization of domains within the primary structures. (b,c) Conformational change of integrins lacking an I domain (b) or containing an $\alpha$ I domain (c). The domains are shown with the same color scheme as in (a).
Figure 2.
Conformational regulation in integrin headpiece domains. (a) Overview of the movements of the βI, hybrid, and PSI domains. Non-moving segments of the βI backbone are shown as a grey worm. Moving segments are color-coded. The downward movement of the α7 helix is coupled to the swing-out of the hybrid domain, which in turn plays a critical role in transmitting signals between the ligand-binding headpiece and the integrin legs. (b) Conformational change of the αI domain. Non-moving segments of the backbone are shown as a grey worm. The moving segments, shown as Cα-traces, of the closed (gold) and open (cyan) αM I domains and their MIDAS metal ions are shown, and direction of movement is shown with arrows. The downward movement of the α7 helix plays a critical role in transmitting signals between the αI domain and the βI domain.