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Journal

Biophysical Journal, 113(4)

ISSN

0006-3495

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Publication Date

2017-08-01

DOI

10.1016/j.bpj.2017.06.064

Peer reviewed

Article



α -Actinin Induces a Kink in the Transmembrane Domain of β_3 -Integrin and Impairs Activation via **Talin**

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ABSTRACT Integrin-mediated signaling is crucial for cell-substrate adhesion and can be triggered from both intra- and extracellular interactions. Although talin binding is sufficient for inside-out activation of integrin, other cytoplasmic proteins such as α -actinin and filamin can directly interfere with talin-mediated integrin activation. Specifically, α -actinin plays distinct roles in regulating $\alpha_{\text{IIIb}}\beta_3$ versus $\alpha_5\beta_1$ integrin. It has been shown that α -actinin competes with talin for binding to the cytoplasmic tail of β_3 -integrin, whereas it cooperates with talin for activating integrin $\alpha_5\beta_1$. In this study, molecular dynamics simulations were employed to compare and contrast molecular mechanisms of $\alpha_{\rm IIb}\beta_3$ and $\alpha_5\beta_1$ activation in the presence and absence of α -actinin. Our results suggest that α -actinin impairs integrin signaling by both undermining talin binding to the β_3 -integrin cytoplasmic tail and inducing a kink in the transmembrane domain of β_3 -integrin. Furthermore, we showed that α -actinin promote talin association with β_1 -integrin by restricting the motion of the cytoplasmic tail and reducing the entropic barrier for talin binding. Taken together, our results showed that the interplay between talin and α -actinin regulates signal transmission via controlling the conformation of the transmembrane domain and altering natural response modes of integrins in a type-specific manner.

INTRODUCTION

Cell-substrate signaling is critical for various mechano-biological functions of the cell, including rigidity sensing, migration, differentiation, and growth (1-3). More than 150 signaling and scaffolding molecules are involved in orchestrating mechanochemical communication of cells with their environment through focal adhesions (4,5). Early adhesions are formed by a few molecules but are transformed to mature focal adhesions as local forces increase (5–7). Mature focal adhesions are firmly attached to the cytoskeleton, converting the actin retrograde flow to traction forces exerted on the substrate (7).

Talin is a chief focal adhesion binding protein and initiates inside-out signaling by disrupting a key interaction between integrin subunits known as the inner membrane clasp (IMC) (8). However, other molecules such as kindlin, filamin, and α -actinin also associate with the integrin tail and directly couple it with the actin cytoskeleton (9-13). The effect of these molecules on the process of integrin activation depends

Submitted January 24, 2017, and accepted for publication June 28, 2017.

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Editor: Markus Buehler.

http://dx.doi.org/10.1016/j.bpj.2017.06.064

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strongly upon their interplay with talin (14). More specifically, these molecules can interfere with integrin activation through either competing or cooperating with talin (14). α -Actinin, one of the most prominent players in focal adhesions (5,15,16), binds to various molecules including integrin and vinculin and is directly involved in transmitting forces between the cytoskeleton and adhesion sites (17–23). β_3 -integrin is necessary for nascent adhesion formation, whereas β_1 -integrins localize to more mature adhesions. Roca-Cusachs et al. (15), in 2013, reported that α -actinin competes with talin for binding to β_3 -integrin and suppresses activation, whereas it cooperates with talin for β_1 -integrin activation.

Proline-induced kinks in the transmembrane domain (TMD) of integrins are shown to be functionally important and associated with diseases in some cases (24). Recent efforts have tried to recognize and reverse the effects of proline mutations. An important example is the A711P mutation, which can compensate for the integrin activation induced by the K716E mutation. The A711P mutation introduces a kink in the β_3 -TMD that breaks the continuity of the helix and replaces the IMC interaction (25). It should be noted that while they both play important roles in regulating the TMD topography (25), neither K716E nor A711P has



been recognized as a disease-causing mutation. However, such proline-induced kinks are evolutionarily significant, and can be used toward mutagenesis-based treatments. Other potential sources of TMD kinks and their impacts on the signal transmission across membrane receptors are still not as well understood.

The composition of proteins at the integrin tail during adhesion formation plays an important role in integrinmediated singling (14). The molecular mechanism of α -actinin and talin crosstalk is not yet understood, largely due to the limitations of experimental studies in capturing the molecular events at sufficient spatial and temporal resolution. In this study, we employed molecular dynamics (MD) models and simulations to investigate the order of binding events and mechanisms of regulating integrin activation through cytoplasmic interactions with talin and α -actinin.

MATERIALS AND METHODS

All molecular models were prepared using the Visual Molecular Dynamics software package (26) and simulations were performed in NAMD (27) using the CHARMM27 force field (28). The configuration of the system in the binding simulations included the full-length α -actinin (PDB: 1SJJ), the head domain of talin (F₀ to F₃ subdomains from PDB: 3IVF), and only the transmembrane and cytoplasmic parts of either $\alpha_{\text{IIb}}\beta_3$ (PDB: 2KNC) or $\alpha_5\beta_1$ integrins; α -actinin was excluded from the control simulations (Fig. 1). The $\alpha_5\beta_1$ integrin crystal structure is not available, thus the $\alpha_{\text{IIb}}\beta_3$ was submitted to the PHYRE2 server as the template for homology modeling. The output model was quite reliable, with the confidence level

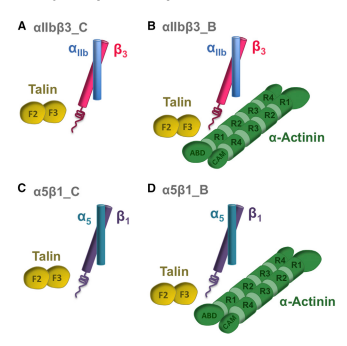


FIGURE 1 A schematic overview of simulations carried out in this work. (A) The interaction between $\alpha_{\text{IIb}}\beta_3$ and talin was modeled in a set of control simulations marked as $\alpha_{\text{IIb}}\beta_3$ _C. (B) α -Actinin, talin, and $\alpha_{\text{IIb}}\beta_3$ were all included in the binding simulations ($\alpha_{\text{IIb}}\beta_{3}$ B). (C) α -actinin was absent in the $\alpha_5\beta_1$ control simulations, marked as $\alpha_5\beta_1$ _C, whereas (D) it was included along with talin and $\alpha_5\beta_1$ in the binding simulations. For clarity, the lipid membrane is not shown. To see this figure in color, go online.

of 99.9% reported by PHYRE2. In the initial configuration of the binding simulations, the integrin binding sites on talin and α -actinin were positioned at equal distances from the integrin cytoplasmic tail, allowing a few water layers to form in the space between the molecules, whereas the TMD domain of the integrin heterodimer was embedded in the lipid membrane.

A triclinic box (size: $41.2 \times 18.2 \times 15.8 \text{ nm}^3$) was generated to account for the elongated structure of α -actinin and also to ensure that molecules did not interact with their periodic image. The TIP3P water model was used for solvating the system but water molecules were deleted from the hydrophobic region of the lipid membrane (29). The excessive charge of the system was neutralized and the overall ion concentration was set to 150 mM of KCl, mimicking the physiological conditions. The number of atoms in the presence of α -actinin reached 1,918,344 and 1,918,175 for the $\alpha_{\text{IIb}}\beta_3$ and $\alpha_5\beta_1$ simulations, respectively.

The SHAKE algorithm was applied to constrain bond lengths and geometry. The temperature was maintained at 310 K and the pressure was kept at 1 bar using Langevin dynamics and Langevin piston pressure control, respectively. Periodic boundary condition was applied in all three directions and the time-step of 2 fs was used in all simulations. The PME method was employed to account for the electrostatic interactions efficiently. The van der Waals interactions were modeled using a switching parameter for smoothly truncating the van der Waals potential at the cutoff distance. The initial configurations of simulations were minimized for 100,000 steps to relax the structures and remove all bad contacts. Minimized structures were then equilibrated for 1 ns after which energy, pressure, and the root mean square deviation were monitored to examine whether an equilibrium state had been reached. Fully equilibrated structures were then used in the final production simulations that each ran for 20 ns and was repeated three times to produce statistically significant results. Simulation parameters of the production runs were the same as those used in the equilibration simulations. The total simulation time performed for this work is 360 ns. From each simulation, 100 time frames were extracted for postprocessing the data, thereby, with three runs, 300 data points were available for each simulation set. The Visual MD package was used to visualize the results (26) and model mutations. The Bio3D package in R was utilized for analyzing the data (30). Normal mode analysis was done using the NOMAD-REF server (31).

RESULTS

Inspired by the previous studies (15), here, we investigated the mechanisms by which α -actinin binding to the β_3 -tail suppresses integrin activation, whereas colocalization of α -actinin with talin promotes $\alpha_5\beta_1$ activation. For that, we designed and performed four sets of simulations as shown in Fig. 1 to capture the effect of α -actinin binding on talin-induced activation of both $\alpha_{\text{IIb}}\beta_3$ and $\alpha_5\beta_1$. The simulations are labeled in Fig. 1, and are referred to as such throughout the text. Two other sets of simulations were performed to test particular mutations speculated to be important in the first four sets of simulations described above.

The competition between talin and α -actinin at the cytoplasmic tail of $\alpha_{\text{IIb}}\beta_3$ integrin

The presence of α -actinin at the β_3 -tail resulted in significantly weaker interaction of talin with β_3 in $\alpha_{IIb}\beta_3$ B simulations (Fig. 2 A). Conversely, talin engaged with the NPLY motif on the β_3 -tail in the absence of α -actinin in the control simulations (Fig. 2 B). This was in agreement with the previous studies that indicated a binding between the F₃ subdomain of talin and the first NPxY motif of the β_3 -tail (Fig. 2 C) (8). In $\alpha_{\text{IIb}}\beta_{3}$ B simulations, the NPLY motif was drawn toward the integrin binding region between R1 and R2 repeats along the α -actinin rod domain (Fig. 2 D).

The density plot of the average distance between talin and β_3 showed the effect of α -actinin in moving the β_3 -tail away from talin (Fig. 2 E). Specifically, the peak value of the talinintegrin distance in the $\alpha_{\text{IIb}}\beta_{3}$ _C simulation was at 11 Å, whereas the peak was shifted to 16 Å in the $\alpha_{\text{IIb}}\beta_{3}$ B simulations showing that the β_3 -tail was pulled away from talin toward α -actinin. The common peak at 14 Å indicated that the initial distance between talin and integrin was the same in the $\alpha_{\text{IIb}}\beta_{3}$ _C and $\alpha_{\text{IIb}}\beta_{3}$ _B simulations. The peak at 17 Å in the $\alpha_{\text{IIb}}\beta_{3}$ B curve occurred as β_{3} moved further toward α -actinin in the last 5 ns of the simulations.

The formation of a kink in the transmembrane domain of β_3 -integrin

Upon α -actinin binding, a kink was formed at residue A711 on the β_3 -TMD, whereas such deformation was not observed in $\alpha_{\text{IIIb}}\beta_{3}$ _C simulations (Fig. 3). The resolved crystal structure of β_3 with the A711P mutation (PDB: 2N9Y) shows a similar kink at P711. Moreover, we carried out a set of simulations in which residue A711 of the wildtype β_3 was mutated to proline (referred to as $\alpha_{\text{IIb}}\beta_3$ _IM simulations in Fig. 3). Our results confirmed the formation of a kink in both $\alpha_{\text{IIb}}\beta_3$ B and $\alpha_{\text{IIb}}\beta_3$ IM simulations, which most likely inhibit integrin signaling across the membrane as shown in Fig. 3.

The continuum model of the TMD backbone structure could be a simple rod, thus its first natural mode is expected to be an in-plane bending. The normal mode analysis confirmed that the first nonrigid motion of the integrin TMD is bending as shown by the dashed arrows in Fig. S1 A. However, the motions of the N- and C termini of TMD were decoupled after the kink was established (Fig. S1 B). The fluctuations of residues were smoothly coupled in the straight TMD, and residue A711 acted as a nodal point. Also, the N-terminus (N-TMD) on the extracellular side moved in phase with the C terminus (C-TMD). In contrast, the root mean square fluctuations (RMSFs) of the kinked structure was highly jagged and motions of the N-TMD and C-TMD went out of phase (Fig. S1 B and C).

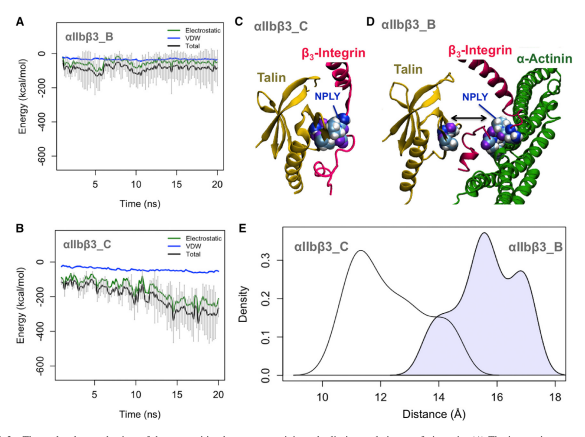


FIGURE 2 The molecular mechanism of the competition between α -actinin and talin in regulating $\alpha_{IIb}\beta_3$ integrin. (A) The interaction energy between talin and the β_3 -tail in the presence of α -actinin averaged over three trials is relatively weak. (B) This interaction energy in $\alpha_{IIIb}\beta_3$ -C simulations drops to -300 kcal/mol after 15 ns. (C) Talin (yellow) is strongly associated with the NPLY motif of β_3 -integrin (mauve) in the last frame of the $\alpha_{IIb}\beta_3$ _C simulations. (D) α -Actinin (green) pulls the NPLY motif away from talin toward its rod domain in $\alpha_{IIb}\beta_{3}$ _B simulations. (E) The density plot of the distance between the NPLY motif and talin for $\alpha_{\text{IIb}}\beta_{3}$ _C simulations (white) peaks at 11 Å, whereas for $\alpha_{\text{IIb}}\beta_{3}$ _B simulations (blue), the major peak occurs at 16 Å. To see this figure in color, go online.

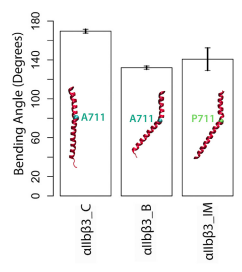


FIGURE 3 Formation of a kink in β_3 -TMD upon α -actinin binding. A kink centered at A711 formed in β_3 -TMD after α -actinin association with its cytoplasmic tail (middle) quantified by the bending angle of the TMD helix. Conversely, no such kink was observed in $\alpha_{\text{IIb}}\beta_3$ _C simulations, indicating that talin alone cannot induce such deformation. Mutating residue A711 to proline created a similar kink. To see this figure in color, go online.

The cooperative role of α -actinin in the binding between talin and β_1 -integrin

To contrast the role of α -actinin in regulating talin association to $\alpha_{\text{IIb}}\beta_3$ and $\alpha_5\beta_1$ integrins, the initial configurations of the two systems were designed to be similar. The interaction between talin and β_1 -integrin formed in both $\alpha_5\beta_1$ _C and $\alpha_5\beta_1$ _B simulations. In the first 15 ns of $\alpha_5\beta_1$ _B simulations, the interaction energy between talin and β_1 -integrin strengthened linearly, whereas this energy remained stable in $\alpha_5\beta_1$ _C simulations (Fig. 4, A and B). However, the talin-integrin interaction was slightly weakened in the second 15 ns of $\alpha_5\beta_1$ B simulations and the average energy was comparable to that in $\alpha_5\beta_1$ _C simulations (Fig. 4, A) and B). This demonstrated that the α -actinin molecule had a positive impact on the initial talin binding by aligning the β_1 -tail. As talin formed a stable interaction with β_1 , α -actinin binding remained intact. Thereby, talin and α -actinin molecules could simultaneously interact with opposite faces of the β_1 -tail.

The complex of talin, α -actinin, and β_1 -integrin formed several interactions at their binding interface (Fig. 4 C). To indicate the packing of the binding residues more clearly, the space in which no interaction occurred, also referred to as the zero interaction zone, was marked in blue. In the control simulation, talin and β_1 -integrin loosely interacted and the zero interaction zone was larger (Fig. 4 D). It should be noted that the zero interaction zone is not necessarily an empty space but it rather represents a region with no interaction between the molecules.

The effect of cytoplasmic interactions on activating integrins $\alpha_{\text{IIb}}\beta_3$ and $\alpha_5\beta_1$

Integrin activation initiates as the TMDs of the integrin subunits dissociate in response to cytoplasmic interactions. We measured the association energy between α_{IIb} - β_3 and α_5 - β_1 subunits in all simulations and observed that the presence of α -actinin markedly decreased the strength of α_5 - β_1 interaction (Fig. 5 A), which makes it prone to activation.

Mechanical signal transmission along the β_3 -TMD strongly relies upon atomic motions. To find the correlation between high fluctuations of the interaction energy and actual physical movements of residues along TMDs, we calculated the RMSFs of β_1 - and β_3 -integrins in all simulations. The RMSF of every atom was obtained from averaging over 100 time frames, and the error bars were calculated by averaging over three runs for each simulation. The RMSFs of β_3 -TMD in $\alpha_{\text{IIb}}\beta_3$ _C simulations as shown in Fig. 5 B was relatively smooth, indicating a tight coupling of residues connecting the key interactions between TMDs, i.e., inner and outer membrane clasps. Conversely, the RMSFs of β_3 -TMD showed a relatively sharp change in $\alpha_{\text{IIb}}\beta_3$ B, which may demonstrate local decoupling events along the β_3 -TMD (Fig. 5 B). Specifically, fluctuations of residues 700, 701, and 705 differed statistically more significantly between $\alpha_{IIb}\beta_3$ B and $\alpha_{IIb}\beta_3$ C as the error bars did not overlap. These residues are all located above A711 toward the N-terminus of β_3 -TMD, whereas RMSFs of TMD residues below A711 was relatively smooth. This may suggest that the kink at A711 allosterically affects fluctuations of the N-terminus of TMD, and breaks the continuity of the TMD domain.

The RMSF of β_1 -integrin in $\alpha_5\beta_1$ _B simulation closely followed that in $\alpha_5\beta_1$ _C along the extracellular and transmembrane sides of the β_1 -TMD, whereas the $\alpha_5\beta_1$ _B RMSF curve gets smoother toward the cytoplasmic side (Fig. 5 C). The RMSFs of the cytoplasmic domains of both β_1 - and β_3 -tails were higher in control simulations (Fig. 5 C), indicating that α -actinin binding constrained the motion of the cytoplasmic regions of both integrins, most likely decreasing the unfavorable entropic barrier for talin binding. Also, the RMSF of the cytoplasmic region was generally higher than the TMD domain, due to the confinement effect of the lipid molecules (Fig. 5 C). Differences in the RMSFs of β_1 and β_3 integrins suggested that signal transmission along integrin is strongly type-specific.

A direct interaction between talin and α -actinin

We observed a very stable interaction between talin and α -actinin at the β_3 -tail, which is most probably an important factor in regulating β_3 -integrin activation as these molecules compete for the same binding site on the integrin tail (Fig. 6). Two important interactions between talin

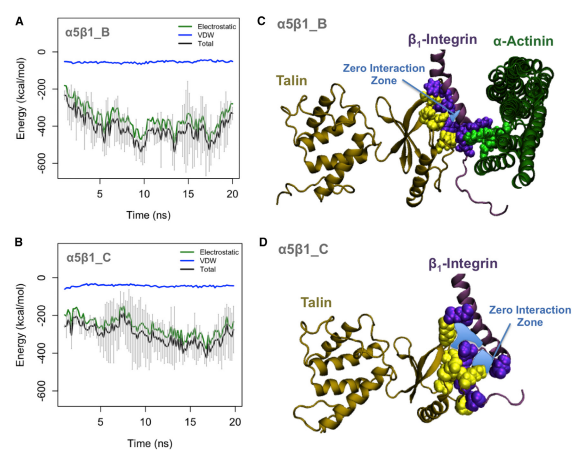


FIGURE 4 The cooperative role of α -actinin in the binding between talin and β_1 -integrin. (A) The average interaction energy of talin and the β_1 -tail in the presence of α -actinin drops to around -400 kcal/mol within the first 15 ns and becomes stable in the last 10 ns of simulations. (B) In $\alpha_5\beta_1$ _C simulations, a large energy drop in the talin- β_1 interaction is not observed and the enthalpy is reduced to ~ -250 kcal/mol. (C) α -Actinin (green) and talin (yellow) simultaneously interact with the β_1 -tail (purple), resulting in a closedly packed residues in the binding region. The zero interaction zone (the area shown in blue) is defined as a gap between atoms with no significant interaction between talin and the β_1 -tail. (D) In $\alpha_5\beta_1$ _C simulations, the zero-interaction zone is increased compared to $\alpha_5\beta_1$ B simulations, indicating the cooperative role of α -actinin in the binding between talin and the β_1 -tail. To see this figure in color, go online.

and α -actinin were: 1) a salt bridge between residues K360 on talin and E464 on α -actinin, and 2) E375 on talin and Q460 on α -actinin as shown in Fig. 6 A. These residues were mutated to alanine (Fig. 6 B) and tested in a set of simulations referred to as $\alpha_{\text{IIb}}\beta_{3}$ TM simulations. It should be noted that the α -actinin-talin interaction was not observed at the β_1 tail (Fig. 6 C). Also, the double mutation on talin completely disrupted its interaction with α -actinin (Fig. 6 C). The binding between talin and β_3 was also correlated with the presence of α -actinin (Fig. 6 D); the less engaged talin became with α -actinin, the more strongly it associated with β_3 .

DISCUSSION

Dynamic bidirectional signaling through integrins is required for regulating cell adhesion in both physiological and pathological conditions. Integrin-talin interaction provides the initial physical coupling essential for signal transmission to the extracellular matrix. As forces are increased at these nascent adhesions, other molecules are recruited to reinforce the extracellular matrix-cytoskeleton linkage and macromolecular complexes with distinct compositions are formed at integrin tails. Thereby, the interplay of cytoplasmic molecules with talin is essential for regulating integrin-mediated adhesions. Talin associates with the cytoplasmic tail of β -integrin and breaks the IMC, which leads to global extension of the integrin ectodomain (32). Other proteins such as filamin, α -actinin, kindlin, and FAK can also interact with the integrin tail and interfere with the activation process. The order in which these proteins bind to integrin and their potential impacts on talin's function can play an important regulatory role in adhesion formation and maturation. α -Actinin is an interesting molecule as it plays opposite roles in regulating $\alpha_{\text{IIb}}\beta_3$ versus $\alpha_5\beta_1$ integrins and thus comparing its interplay with talin and integrin may reveal fundamental differences in type-specific regulatory mechanisms. Here, we investigated talin-mediated signal transmission from cytoplasm to the TMD domains of $\alpha_{\text{IIb}}\beta_3$ and $\alpha_5\beta_1$ integrins in the presence and absence of α -actinin.

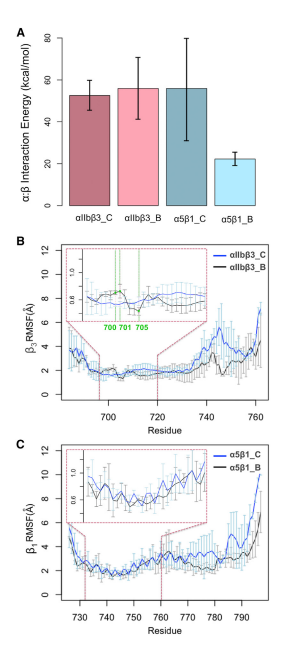


FIGURE 5 The effect of cytoplasmic interactions on integrin activation. (A) The average energy between α_5 - β_1 subunits in $\alpha_5\beta_1$ _B simulations is significantly decreased compared to that in $\alpha_5\beta_1$ _C simulations. Within the same simulation time, no change in the α_{IIb} - β_3 energy is observed upon α -actinin binding. (B) Atomic fluctuations are limited in the transmembrane domain relative to the cytoplasmic end of integrins in all simulations. The TMD regions are magnified in the insets. The RMSF of β_3 -tail is different between $\alpha_{\text{IIb}}\beta_3$ _C and $\alpha_{\text{IIb}}\beta_3$ _B simulations. Specifically, the error bars of atomic fluctuations do not overlap at residues 700, 701, and 705, which are all above residue 711 toward the extracellular side of TMD, showing a notable difference between the RMSF curves at those residues. (C) The RMSF of β_1 -TMD is similar between $\alpha_5\beta_1$ _C and $\alpha_{\text{IIb}}\beta_3$ _B simulations. This most likely indicates that signal transmission through the β_1 -TMD is not significantly altered in the presence of α -actinin. To see this figure in color, go online.

Our results showed that talin binding to β_1 -integrin was notably enhanced by including α -actinin in the simulations (Fig. 4). Specifically, α -actinin stabilized talin's initial

engagement by decreasing atomic motions, as indicated by lower RMSF of the β_1 -tail after α -actinin association, which most likely decreases the entropic barrier for talin binding (Fig. 5 *C*). Furthermore, the simultaneous interaction of α -actinin and talin lowered the binding energy between the $\alpha_5\beta_1$ integrin subunits, elucidating the cooperative role of α -actinin.

Control simulations featuring β_3 -integrin and talin consistently showed successful association of talin with the NPLY motif of β_3 -tail, which was in agreement with previous studies (33,34). However, this interaction was markedly lowered upon introducing α -actinin, as it pulled the β_3 -tail away from talin. As a result, α -actinin bound exclusively to β_3 -integrin and prevented talin from competing any further by directly associating with it. Although the mechanisms of integrin activation have been previously examined (33,35), we report for the first time to our knowledge, the emergence of a striking kink along the β_3 -TMD upon α -actinin binding. The β_3 -TMD forms a $\sim 25^{\circ}$ tilting angle with the α_{IIb} -subunit, thus integrin monomers cross each other inside the lipid membrane. Talin binding increases the tilting angle of β_3 -integrin, resulting in the disruption of the IMC (24). The location of the kink observed in our simulations was near the crossing point of integrin subunits and occurred at the position of residue A711. We strongly hypothesize that the conformational change of the TMD domain is a critical factor in inhibiting integrin-mediated signaling. Previous studies also showed that the continuity of the integrin TMD is key to integrin activation (24). Interestingly, it was indicated that the A711P mutation also resulted in a flexible kink that broke the continuity of the β_3 -TMD, decoupling the tilting motion of integrin via talin (36). This supports our hypothesis on the mechanism by which α -actinin association suppresses integrin activation and signaling because the kink induced by α -actinin in the β_3 -TMD greatly resembles the effect of A711P mutation.

Normal mode analysis performed on both kinked and straight conformations of the β_3 -TMD indicated that the first bending mode of the TMD domain was significantly altered upon α -actinin binding (Fig. S1). Interestingly, the position of A711 after the appearance of the kink acted as a decoupling joint such that motions of the N-TMD and C-TMD went out-of-phase as illustrated by the gray arrows in Fig. S1, A and B. The RMSFs of C_{α} atoms in the first bending mode of the straight β_3 -TMD resembled a cosine wave function that most likely allowed for smooth transfer of mechanical signals along the TMD domain (Fig. S1 C). However, a jagged-looking RMSF plot of the first bending mode of the kinked structure suggests that random fluctuations of the TMD atoms most likely result in an inefficient transfer of mechanical signals across the membrane after α -actinin association. It should be noted the RMSF of β_3 -TMD atoms in the bending mode (Fig. S1 C) does not have to match that RMSF extracted from the MD

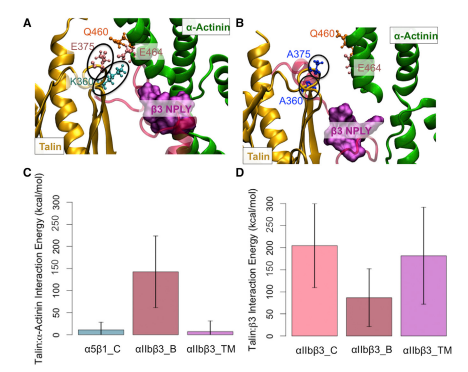


FIGURE 6 The binding between talin and α -actinin is a possible mechanism for competition. (A) Strong interactions between α -actinin and talin are exclusively formed at the β_3 -tail in $\alpha_{IIb}\beta_3$ _B simulations, mainly between residues E375 and K360 on talin, and residues Q460 and E464 on α -actinin. (B) Mutating both E375 and K360 to alanine changes the position of talin relative to both the NPLY motif and α -actinin. (C) The barplots indicate that the absolute value of interaction energy between talin and α -actinin is significantly reduced upon talin's double mutation. It is also evident from the plot that α -actinin and talin do not associate at the β_1 -tail. (D) The interaction between talin and β_3 is increased upon talin mutations. To see this figure in color, go online.

simulation (Fig. 5, B and C) since this mode was not excited in the simulation. However, our analysis shows that the conformational changes in the TMD domain significantly affect affected the natural modes of the β_3 -TMD.

The observed interaction between talin and α -actinin, to our knowledge, has never been reported in the previous literature and could serve as a new target for future experimental studies. Our results suggested that one of the mechanisms by which α -actinin impairs β_3 -integrin-mediated signaling and competes with talin, is by forming simultaneous interactions with the β_3 -tail and talin head, which effectively prevents talin from associating with the integrin tail. Interestingly, α -actinin did not form any interaction with talin at the β_1 -tail, which confirmed our results on the exclusive occurrence of α -actinin-talin binding during competition.

Previous studies have discussed the effect of other cytoplasmic interactions at the integrin tail. For instance, it has been observed that the filamin competition with talin results in impaired integrin activation (11). On the contrary, kindlin binding promotes adhesion formation and reinforcement (24). Major approaches for examining the effect of these proteins are mostly limited to fluorescence imaging, celllevel assays (15,36), which can only provide macroscopic information about the significance of cytoplasmic molecules in the integrin activation. This calls for new computational efforts to shed light on the mechanistic details and the factors involved in integrin activation (37–39). Recent crystallography studies provided a greatly valuable insight into the interhelical modifications within the $\alpha_{\text{IIb}}\beta_3$ heterodimer upon the A711P mutation, however such static snapshots provides no evidence on the dynamics of the system and potential allosteric regulations.

Taken together, our results suggest that α -actinin binding induces conformational changes in the β_3 -TMD, which can substantially modify mechanical signal transmission across the lipid membrane (Fig. 7). Furthermore, proline-induced kinks are shown to be evolutionarily important and critical for regulating interhelical packing of integrin receptors. However, to our knowledge, this is the first study that associates kink formation with the cytoplasmic interactions and explains its impact on the mechanism of signal transmission, which in turn cause macroscopic changes in cell adhesion and spreading.

SUPPORTING MATERIAL

One figure is available at http://www.biophysj.org/biophysj/supplemental/ S0006-3495(30751-8).

AUTHOR CONTRIBUTIONS

H.S. and M.R.K.M. designed the experiments, analyzed the data, and wrote the manuscript. H.S. performed the experiments. M.R.K.M. contributed materials, and analysis tools.

ACKNOWLEDGMENTS

We thank Zainab Haydari and Zeinab Jahed and all other members of the Molecular Cell Biomechanics Laboratory for their fruitful discussions and feedbacks throughout this work.

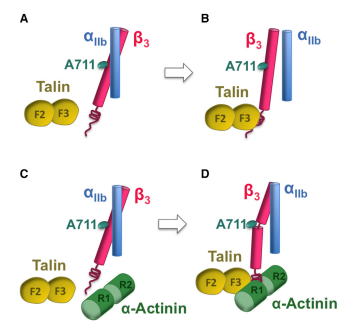


FIGURE 7 Possible molecular mechanism by which α -actinin suppresses integrin-mediated signaling. (A) In the inactive conformation of integrin, the TMD domains of integrin subunits directly interact. (B) As talin associates with the cytoplasmic tail of β_3 -integrin, the IMC interaction between integrin subunits is broken, changing the distance between α_{IIb} and β_3 subunits without altering the TMD conformation. (C) Talin and α -actinin compete for binding to the β_3 -tail. For clarity, only R1 and R2 repeats of α -actinin are shown. (D) α -Actinin directly engages with both the β_3 -tail and talin head and prevents talin from binding to integrin. It also induces a kink at A711 in the β_3 -TMD, which most likely impairs integrin signaling. For clarity, the full-length α -actinin is not shown but it was included in all simulations. To see this figure in color, go online.

This study was supported by National Science Foundation (NSF) grant CBET-0829205 and CAREER Award CBET-0955291 to M.R.K.M. In addition, this work used the Extreme Science and Engineering Discovery Environment (XSEDE), which is supported by the National Science Foundation (NSF) (grant ACI-1053575).

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