The Structure of a Full-length Membrane-embedded Integrin Bound to a Physiological Ligand*

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Background: Integrins undergo large conformational changes when ligand-bound.
Results: Using nanodisc technology and EM, we observed Mn2+-activated αIIbβ3 integrin both alone and fibrin-bound.
Conclusion: MnCl2-activated αIIbβ3 integrin alone has a compact conformation, becoming fully upright and open when fibrin-bound.
Significance: The first structure of membrane-embedded integrins bound to physiological substrate reveals the importance of integrin extension in macromolecular ligand binding.

Increased ligand binding to integrin (“activation”) underpins many biological processes, such as leukocyte trafficking, cell migration, host-pathogen interaction, and hemostasis. Integrins exist in several conformations, ranging from compact and bent to extended and open. However, the exact conformation of membrane-embedded, full-length integrin bound to its physiological macromolecular ligand is still unclear. Integrin αIIbβ3, the most abundant integrin in platelets, has been a prototype for integrin activation studies. Using negative stain electron microscopy and nanodisc-embedding to provide a membrane-like environment, we visualized the conformation of full-length membrane-embedded, full-length integrin bound to its physiological macromolecular ligand is still unclear. Integrin αIIbβ3, the most abundant integrin in platelets, has been a prototype for integrin activation studies. Using negative stain electron microscopy and nanodisc-embedding to provide a membrane-like environment, we visualized the conformation of full-length αIIbβ3 in both a Mn2+-activated, ligand-free state and a Mn2+-activated, fibrin-bound state. Activated but ligand-free integrins exist mainly in the compact conformation, whereas fibrin-bound αIIbβ3 predominantly exists in a fully extended, headpiece open conformation. Our results show that membrane-embedded, full-length integrin adopts an extended and open conformation when bound to its physiological macromolecular ligand.

Integrins are non-covalent heterodimers containing α- and β-subunits, both of which have a large extracellular (ectodomain), a single transmembrane (TM) α-helix and a small cytoplasmic domain (1). Integrins respond to cytoplasmic signals by increasing the affinity of their ectodomains for ligands through an allosteric rearrangement that requires the binding of talin to the β-subunit cytoplasmic domain, in a process termed “integrin activation.” Integrin activation plays an important role in development, immune responses, hemostasis, and cancer (2). Integrin αIIbβ3, the most abundant surface protein in platelets, mediates platelet aggregation and is critical in hemostasis, thrombosis, and inflammation (3). Mutations of αIIbβ3 are known to cause diseases such as Glanzmann thrombasthenia, a disease of abnormal hemostasis (4). The affinity of integrin αIIbβ3 must be precisely regulated to enable hemostasis responses upon platelet stimulation while preventing pathological thrombosis due to uncontrolled platelet aggregation. Thus, αIIbβ3 has been a prototype for studies of integrin activation in health and disease (5–8).

In integrins without an α-A domain, which includes αIIbβ3, the α-subunit β-propeller domain, and the β-subunit β-A domain, jointly form a ligand binding pocket (see Fig. 1B) (9–11). The ligand binding pocket, the thigh domain of the α-subunit, the hybrid domain, the PSI domain, and I-EGF1 of the β-subunit are referred to as the headpiece and can be studied as a separate fragment (11, 12). The rest of the ectodomain including the calf-1 and calf-2 domains from the α-subunit, I-EGF2–4, and β-tail domains from the β-subunit comprise the leg pieces (Fig. 1B). Integrin conformation has been studied by x-ray crystallography, electron microscopy (EM), NMR, and molecular dynamics simulation of integrin fragments and detergent-solubilized integrin (1, 13–16). The reported conformations include a compact shape with the headpiece pointing toward the membrane (Fig. 1A) (compact, head down) (9, 10), a compact form with the headpiece pointing away from the membrane (compact, head up) (17), an extended form with hybrid and calf-1 domain juxtaposed (extended, headpiece closed) (18), and an extended form with the hybrid domain swung out (Fig. 1, B and C) (extended, headpiece open) (12, 18, 19). Despite these efforts, membrane-embedded, full-length integrin bound to its physiological ligand has not been directly visualized, and its precise conformation is not clear.

The controversy in the current literature necessitates the present focus on the conformation of membrane-embedded, full-length integrin bound to its physiological ligand fibrin. Activated αβ3 integrin ectodomain in the compact, head-down conformation bound stably to a fragment of fibronectin (20), whereas in a comparable study using αβ3, integrin only the extended, headpiece-open conformation was observed when bound to the same protein ligand (12). In a similar con-
troverty, crystals of αβ3 integrin ectodomain bound to a cyclic RGD peptide, which typically activates integrins, showed the compact, head-down conformation (21), whereas in an EM study utilizing both αβ and αINHβ3 only the extended, headpiece-open conformation was observed in the presence of an RGD peptide (22). Adding to this controversy are the reports that both the compact and the extended-open conformation of integrin ectodomain fragments can be in the activated form (9, 20, 22–25).

The focus of more recent studies (17, 25, 26) has been on how well these observations made with ectodomain or ectodomain fragments correspond to the conformation of integrins embedded in the membrane. Observations such as a surprising compact, head-up conformation for inactive αINHβ3 integrin in nanodiscs (17), a stable integrin αIβ2 conformation with intermediate affinity in cells (26), and conflicting inferences on the conformation of different integrin isoforms in the cell membrane drawn from fluorescence studies (25, 27–29) further complicate this issue.

We have previously reported that the talin head domain (THD), the integrin-activating fragment of talin, more than doubles the percentage of extended conformation of full-length integrin αINHβ3 embedded in islands of phospholipid bilayers formed in nanodiscs (30). However, we were unable to determine the conformation of activated integrin bound to macromolecular ligands such as fibrin because we could not observe the THD in the proposed talin-integrin-fibrin complex. Because of these uncertainties, we lacked independent verification of the activation state of each individual integrin, i.e. we were unable to pinpoint the activation state of the individual integrin in the complex without using the extended conformation as the proxy. An alternative way of ensuring uniform and high percentage of integrin activation is Mn2+ treatment (31), which is often used as a positive control for integrin activation. Mn2+ replaces all Ca2+ and occupies 100% of the integrin metal ion sites (21), thus producing a homogeneously activated integrin population.

Here we reconstituted purified full-length αINHβ3 integrin into phospholipid nanodiscs, activated these αINHβ3 nanodiscs with Mn2+, and visualized them in complex with fibrin by negative stain EM. Our EM results show that fibrin-free, Mn2+-activated integrin adopts a compact conformation. However, integrins bound to fibrin, a physiological macromolecular ligand, predominantly adopt a fully extended, headpiece-open conformation.

Experimental Procedures

Preparation of Integrin Nanodiscs and Two-dimensional Fibrin Complexes—Full-length αINHβ3 integrin was purified from outdated blood platelets following published procedures (30, 32, 33). Integrin nanodiscs were assembled using a protocol adapted previously (34, 35). Briefly, detergent-solubilized 1,2-dimyristoyl-sn-glycero-3-phosphocholine, 1,2-dimyristoyl-sn-glycero-3-phospho-(1′-rac-glycerol), membrane scaffold protein, integrin, and other solutes were combined, and the detergent was removed with SM-2 Biobeads. The assembled integrin nanodiscs were then purified with a size exclusion column.

The lipid monolayer technique previously described was employed to produce fibrin-integrin complexes suitable for EM (30). We used a rigid lipid monolayer consisting of dioctadecylmethyl ammonium bromide and dipalmityl phosphatidylcholine. This lipid combination is in a gel phase, i.e. it is not fluid, at 4 °C, thereby preventing in-plane diffusion of the integrin nanodiscs after they bind the lipid. In the present paper, MnCl2 instead of THD was used to activate the integrin. We preferred Mn2+ over THD as the activator because it will activate all of the integrins, whereas THD at achievable concentrations in vitro will not (30). Preformed integrin nanodiscs were activated by incubation for 2 h at room temperature in 20 mM Tris, 50 mM NaCl, 50 mM MnCl2, pH 7.6. After injection under the monolayer the final Mn2+ concentration was 5 mM. Between 1.0 and 1.5 μl of the activated integrin solution was injected by Hamilton syringe into the buffer phase under the fibrin-lipid monolayer samples, all done at 4 °C. After a 24-h incubation, the lipid monolayer samples were evaluated by negative stain EM to verify the presence of an adequate concentration of integrin bound to the fibrin before data collection.

Electron Microscopy—Samples were stained with 2% uranyl acetate following previous procedures (30). Projection images were collected using low dose protocols on an FEI CM120 TEM (FEI, Hillsboro, OR) at 120 keV using a TVIPS F224 2K × 2K CCD camera (Tietz; TVIPS GmbH, Germany). The magnification was 60,000×; the defocus was ~2.5 μm. The final pixel size was 0.222 nm. A total of 600 micrographs were collected.

Data Processing—Most image processing was performed with the I3 package (36), which utilizes modulation analysis and hierarchical ascendant classification.

Integrin nanodiscs could be readily identified because of their distinctive shape and size. Images with good density of both free and fibrin-bound integrin nanodiscs were picked manually for further processing. A total of 1515 fibrin-free integrin nanodisc particles were normalized, corrected for the
contrast transfer function by Wiener filtering, and low passfiltered to 1.78-nm resolution. For each of the selected particles, the nanodisc density was placed in the center and first aligned by translation only. A global average of all extracted particles was used as the initial reference for translational alignment. For the free integrin nanodiscs, rotational alignment was used to place the integrin density at the top of the nanodisc density. The procedure comprised the following: a randomly picked particle with clear integrin density was used as the initial reference, and subsequent cycles used the class averages as the reference for alignment of all the raw particles. When the integrin density was aligned well at the top of the nanodisc, multivariate data analysis and multi-reference alignment using only class averages, known as alignment by classification (37), were used to focus further alignment on the integrin density.

For the fibrin-bound integrin nanodiscs, we selected a total of 4,656 particles. Once the nanodisc density was centered well with the integrin density positioned in the bottom half of the image, a raw particle with clear, straight horizontal fibrin density at the bottom of the image was chosen as the initial reference. Rotational alignment was then applied to all raw particles to make the fibrin density horizontal as the second point of reference. In this way the integrin density was concentrated between the centered nanodisc and the horizontal fibrin. A trapezoidal shaped Boolean mask covering the space between the nanodisc and the fibrin was used for focused classification to separate fibrin-bound integrin conformations. This procedure minimizes the possibility of reference bias in the integrin class averages because no alignment is done on the integrin class itself; all the alignment was done to center the nanodisc density and to place the fibrin fiber at the bottom.

The majority of class averages showed well defined nanodisc and fibrin density, but the integrin density was uninterpretable or ambiguous (more than one interpretation for leg pieces and head piece). We then examined montages of the class membership systematically rejecting class members of poor quality based on visibility of headpiece and legs (i.e. preservation in negative stain) thereby reducing the number of fibrin-bound integrin nanodiscs to 1,057.

Alignment on only the nanodisc and fibrin for this subset of images generated interpretable class averages that incorporated only 23 integrin nanodiscs, suggesting that the integrins were broadly distributed across the gap between the nanodisc and fibrin as well as conformationally variable. Consequently, we did further alignment based on the integrin position. For the first cycle we did a single reference alignment of raw particles against a class average having the integrin facing vertically down. After this first cycle, we used multireference alignment by classification but masked the images on the integrin density. In this way we obtained four additional interpretable class averages incorporating a further 245 raw nanodisc images.

We then took the remaining integrin images supplemented with previously rejected images, a total of 4,388 particles, and repeated the analysis twice using a conventional single particle approach with alignment on the integrin nanodisc, only with the fibrin masked out. In each cycle we rejected 20% of particles with poor correlation. Both single and multireference alignment strategies were used reducing the particle count to 2,384 images. However, no new conformations were identified, and those that were found were substantially inferior in quality to the ones shown. Finally, we repeated the analysis using the EMAN software package (38). We computed 40 class averages, but they were no more interpretable than those produced using I3 and suggested no new conformations. We conclude from this that our unconventional image analysis was better suited than conventional methods to the analysis of these micrographs.

Model Building—A density map of a bent-over integrin was generated using the crystal structure of the ectodomain of integrin αIIbβ3 with the TM domain (PDB code 4CAK; Ref. 17) (Fig. 1A). The density map of active integrin (shown in Fig. 1, B and C) was generated mainly using the crystal structure of integrin αIIbβ3 with the open headpiece (PDB code 3FCU; Ref. 9). Projections of the electron density volume were generated using the Spider software package (39). Projections of these maps were generated in a grid covering the three Euler angles in increments of 10°.

Results

Structure of Fibrin-free Integrin Nanodiscs—Inactive integrin is believed to adopt a bent-over (referred to here as compact, head-down) conformation as shown by crystal structures of the αIIbβ3 ectodomain (9, 10). This structure has been confirmed for the α-I integrin αIIbβ3 (40). However, whether activated integrin is extended before engaging its ligand is still not clear. To study integrin conformation in a homogeneously activated integrin population, we investigated Mn2+-activated full-length integrin αIIbβ3 embedded in an island of membrane formed by our previously developed αIIbβ3 nanodiscs (30, 35). We picked, classified, and averaged 1,515 particles of integrin nanodiscs unbound by ligand from images of samples where we added integrin nanodiscs to preformed fibrin (Fig. 2). The fibrin-free integrin nanodiscs lie in the background of the fibrin-bound integrin nanodiscs so that the solution conditions and other variables

FIGURE 2. Fibrin rafts decorated with αIIbβ3 integrin nanodiscs. Electron micrograph of fibrin rafts formed on a positively charged lipid monolayer and was treated with αIIbβ3 integrin nanodiscs previously treated with Mn2+ to achieve an activated state with high affinity for fibrin. In this sample the nanodiscs density is disk-shaped; the filamentous structure is fibrin. The density connecting the nanodiscs with the fibrin is the integrin. The pointers locate some fibrin-bound integrin nanodiscs. Bar, 100 nm.
are identical for both populations. The general shape of the unbound integrin nanodiscs had two distinctive but connected densities: one large and circular and the other smaller and irregular (Fig. 3). The large moiety at 12–13-nm diameter was close to the size of nanodiscs predicted from the length of the membrane scaffold protein and observed in our previous study (30). Thus, the large moiety is assigned to the nanodisc density. The nanodiscs lay nearly exclusively “en face” on the surface of the lipid monolayer, an orientation that is likely to be due to the negative charge of the phospholipids and the positive charge of the lipid monolayer.

The small moiety is around 9–10 nm in length in its longest dimension. Its size and shape resemble integrin density observed previously (30) and is assigned to the integrin ectodomain. The size of the integrin density is consistent with previous studies (17, 33, 41). The size is slightly smaller than previously observed for detergent-solubilized integrin (41), which is about 13 nm. However, the TM and cytoplasmic domains and the surrounding detergent micelle may increase the observed size in detergent. In our preparations using integrin nanodiscs, the ectodomain nearly always lay over the edge of the nanodiscs. The shape and length of the ectodomain is consistent with a bent-over conformation or other non-extended conformation (17, 33, 41). Computation of class averages and visual inspection of class members as well as raw micrographs revealed only a small fraction of fibrin-free integrin nanodiscs in the extended conformation. Classes with integrins clearly in the compact conformation comprised 1,394 of 1,515 fibrin-free integrin nanodiscs with the best morphological detail. Numbers in the upper right hand corner are the number of class members within each class average.

Recently, Choi et al. (17) reported that full-length integrin \( \alpha_{IIb}\beta_3 \) in nanodiscs adopts a compact, head-up conformation with the headpiece pointed away from the membrane and not toward the membrane as is commonly thought. We, therefore, generated density maps from three compact structures, PDB codes 3IJE, 3FCS, and 4CAK (9, 25). We then computed projection maps covering the whole of Euler space and aligned each with 18 class averages displaying the best-defined integrin density (Fig. 3). The projections from the Choi et al. (17) model (PDB code 4CAK) appeared to be most similar to our class averages (Fig. 4) due to the fact that the membrane-proximal part of the integrin molecule was always adjacent to the nanodiscs. The best alignment of the other two structures (PDB codes 3IJE and 3FCS) placed the membrane-proximal domains adjacent to the nanodiscs <50% of the time. Such a placement is unlikely in the integrin nanodiscs specimen. The difference in correlation coefficient between our map and the three-modeled maps was small (~20%), but the Choi (17) model produced consistently better correlation. The significant difference between this study and that of Choi et al. (17) is the shape of the
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nanodisc in projection. In our experiment we used a positively charged lipid monolayer to support the nanodiscs, whereas Choi et al. (17) used glow-discharged carbon-coated grids, a process that makes the carbon film generally negatively charged if done in air. The negative phospholipid in the nanodiscs would be attracted to the positively charged lipid monolayer making the nanodiscs orient en face, thereby presenting a circular shape. This would not be the case with glow discharging, which is likely to bind the integrin to the carbon film rather than the lipids in the nanodiscs. Choi et al. (17) found that the orientation of the nanodiscs varies considerably, thus presenting ellipsoidal shapes rather than orienting en face as in our case, consistent with this interpretation.

Structure of Fibrin-bound Integrin Nanodiscs—To facilitate the visualization of $\alpha_{IIb}\beta_3$-ligand interaction, we chose fibrin, the product of fibrinogen polymerization after thrombin cleavage, as the ligand for $\alpha_{IIb}\beta_3$. Fibrin fibers were formed on a lipid monolayer to enforce a two-dimensional arrangement of the sample, which simplifies interpretation (42, 43). Fibrin and fibrinogen, the physiological ligands of $\alpha_{IIb}\beta_3$, have been used in numerous studies of platelet and integrin activation (1, 3, 44 – 49). The binding of these ligands has been regarded as a measure for integrin activation. Furthermore, no direct observation of the molecular structure of $\alpha_{IIb}\beta_3$ in complex with fibrin has been reported previous to this study.

In micrographs of the negatively stained fibrin complexes (Fig. 2), the fibrin fibers exist as a two-dimensional bundle of varying width. Decoration of $\alpha_{IIb}\beta_3$ nanodiscs along the edge of the fibrin filaments can be clearly seen in the form of round disks generally at a uniform distance from the filament and connected to it by integrin densities. Because ligand binding is the measure of integrin activation, these $\alpha_{IIb}\beta_3$ nanodiscs decorating the fibrin filaments are, by definition, the activated form. We processed images of the fibrin-attached integrin nanodiscs, as we have done with ligand-free integrin nanodiscs. Starting with 4,656 single particle images, the nanodisc density was placed in the center of the boxed image, which was large enough to include both the integrin and a few strands of the fibrin raft. During the processing, the fibrin filament density was rotated to be oriented parallel to the bottom of the box as much as possible. This gave a frame of reference consisting of the centered integrin nanodiscs and the horizontally oriented fibrin strands, which made it easier to visualize the variable shape of integrin and fibrin. We could also focus the multivariate data analysis on the smaller area between the nanodiscs and the fibrin fiber where the integrin is presumed to lie. Because of the complicated geometry and variable size of the fibrin bundles, we were unable to obtain a unique structure of fibrin; each averaged fibrin filament is essentially featureless. Single, double, and even triple strands of fibrin were found in the class averages. To further reduce the heterogeneity problem, the extracted fibrin-bound integrin nanodiscs were further separated based on the number of fibrin strands within the boxed area. Because the angle between integrin and fibrin filament in the raw images shows considerable variation (Fig. 2), classification by multivariate data analysis grouped particles with similar conformation and similar integrin/fibrin angle into separate classes, resulting in considerably smaller population per class.

Most of these classes were uninterpretable, necessitating visual editing of the class membership according to image quality, thereby reducing the number of usable single particle images by ~75%. At the end of the processing, only four classes incorporating a total of 23 raw particles of the remaining 1,057 images showed both distinct headpiece and leg densities (Fig. 5A). Although few in number, these classes represent the most objective structures as no alignment has been done on the integrin itself. Based on the class averages, we measured the length of the extended integrin to be $20 \pm 1$ nm, consistent with that calculated from extended molecular models and that observed with full-length integrin $\alpha_{IIb}\beta_3$ in detergent (41). Both class averages and the raw images in the class clearly show that integrin headpieces are in an open conformation and the two leg pieces are discernible and separated (Fig. 5, A and B). They generally do not appear to be crossed as seen in talin-activated but ligand-free $\alpha_{IIb}\beta_3$ nanodiscs (30). The leg densities from both chains at the edge of the nanodiscs remain separated but not by much. The largest leg separation is about 4.5 nm. The angle between the hybrid domain and thigh domain is $84^\circ \pm 12^\circ$, consistent with a hybrid domain in a swung-out position after ligand engagement.

Using the remaining 1,034 raw particles, we did another alignment this time using the integrin density. We obtained an additional 4 class averages encompassing 245 integrin nanodiscs that showed clear integrin headpiece and leg density, but in this case the fibrin density was blurred out (Fig. 5C). Thus, if the fibrin was well aligned, few of the integrins were well aligned, and when the integrins were aligned, very little of the fibrin was aligned. The other class averages either did not show both the headpiece and leg-piece densities clearly or were
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Comparison of Fibrin-free and Fibrin-bound Integrin—The class averages clearly showed that in the fibrin-bound state, the integrins are extended, the leg pieces of $\alpha$- and $\beta$-subunits are well separated (although they sometimes cross near the edge of the nanodisc), and the head-piece is open, whereas in the Mn$^{2+}$-activated but ligand-free state, the integrins are bent over, and their leg pieces are juxtaposed and indiscernible. To directly show that ligand engagement induces the extension of integrin, a typical class average of activated but ligand-free integrin nanodiscs was superimposed with ligand-bound integrin nanodiscs by bringing their nanodiscs densities into register (Fig. 6). This superimposed image clearly demonstrates that the length of activated but fibrin-free integrin is much shorter than that in the fibrin-bound integrin nanodiscs (B).

Discussion

Our previous papers showed no dramatic change on the length of the ectodomain when the full-length $\alpha_{IIb}\beta_3$ integrin was activated by 1 mM Mn$^{2+}$ when incorporated into small unilamellar vesicles (33). The average length is around 11 nm, which means there is no large conformational change when activated by Mn$^{2+}$ alone. The presence of a compact conformation in detergent after Mn$^{2+}$ activation was later confirmed (41). Here we further confirmed that integrin in a membrane bilayer environment after Mn$^{2+}$ activation adopts primarily a compact conformation with no dramatic length change. In Eng et al. (41), 60% of the integrin population adopted a compact head-down conformation after Mn$^{2+}$ activation; even after the addition of another ligand mimetic L-739758, 20% of the integrins remain in the compact head-down conformation. In our result the ratio of compact to extended conformation for the unbound integrin nanodisc is close to 19:1, which is much higher than the 2.3:1 observed in the above-mentioned experiment. The higher ratio in our result could be due to the membrane environment, in which the integrin transmembrane domain interactions are stabilized by formation of inner and outer membrane clasps (50, 51), thereby maintaining the compact conformation.

Because the fibrin-free integrin nanodiscs were quantified in the presence of ligand, it could be argued that the low percentage of activated integrins in the extended but ligand-free state compared with the 40% predicted by previous studies in detergent (41) was due to fibrin removing the upright, activated integrins from the pool of activated integrins. The fact the Mn$^{2+}$-treated nanodiscs were incubated with the fibrin for 24 h before EM specimens were made argues against this possibility. If fibrin indeed removed all the extended integrins, it seems likely that the prolonged incubation would have allowed the remaining fibrin-free nanodiscs to reach a new equilibrium even at 4°C. Furthermore, the rigid characteristic of the lipid monolayer used would have prevented in-plane diffusion of the lipid-bound nanodiscs and limited the possibility that fibrin may concentrate around itself the active, extended integrin nanodiscs among the integrin nanodiscs attached to lipid monolayer. Integrins in the extended or bent conformation, if they did not happen to hit the fibrin when first absorbed onto lipid monolayer, will therefore not "see" fibrin during the entire experiment due to nonfluidity of the underlying monolayer. These integrin nanodiscs away from fibrin would presumably remain bound in whatever conformation they were in and stay at a locus when they first contacted the monolayer. Thus, we feel that the internal control of ligand-free nanodiscs provides a reasonable measure of the proportion of extended integrins in solution.

Integrin $\alpha_{IIb}\beta_3$ extends to 18–20 nm in length upon binding to fibrin. This length is very close to the 23-nm maximum length of intact integrin $\alpha_{IIb}\beta_3$ in detergent (41). Considering the TM and cytoplasmic domains are visible in detergent-solubilized integrins but invisible in the negatively stained integrin nanodiscs, the length of the ectodomain in our study is reasonable. Although we cannot identify the TM domain directly, it is reasonable to expect that separation of the TM domains of the $\alpha$- and $\beta$-subunits has occurred based on the degree of separation between the adjacent leg densities. The moderate size of the lower leg separation, 4.5 nm, is consistent with the recent observation that TM domains from the two subunits are still close to each other after headpiece opening (41), although the size of the nanodiscs may have constrained the degree of separation. The fibrin-bound integrin adopts an extended, headpiece-open conformation as confirmed by the angle between the adjacent leg densities.

Therefore, full-length $\alpha_{IIb}\beta_3$ integrin embedded in a phospholipid bilayer is in an extended, headpiece-open conforma-
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tion when bound to its macromolecular ligand. Our results can be reconciled with previous reports on both sides of the controversy (12, 19, 20, 22). It is possible that the presence of a lipid bilayer would restrict a macromolecular ligand or protein ligand from accessing the membrane-facing ligand pocket in a bent integrin conformer even if it is activated. Thus, truncating the transmembrane domain and removal of the lipid bilayer would enable such an interaction with an activated but bent integrin conformer, as seen previously (20).

Recently, Choi et al. (17) reported that the headpiece of inactive integrin $\alpha_{\text{IIb}}\beta_3$ points away from the membrane (compact, head-up) rather than toward it as depicted in most current models. We, therefore, tested three models for the compact conformation. The model from Choi et al. (17) was the better fit but largely in terms of consistently providing a realistic orientation of the membrane proximal domains with respect to the nanodiscs. Nevertheless, due to the preferred orientation and possible flattening due to drying, our projections themselves are insufficient to conclude that the headpiece points away from the membrane.

To summarize, we have investigated the conformation of integrin in a physiological-like condition. Our study used full-length integrin $\alpha_{\text{IIb}}\beta_3$ incorporated into nanodiscs to mimic the interaction between the TM domain and the membrane environment. $\text{Mn}^{2+}$-activated but ligand-free integrin in a lipid bilayer remains in the non-extended conformation, but integrin nanodiscs predominantly adopt an extended, headpiece-open conformation when bound to its macromolecular ligand. We captured the images of both conformations and presented the first direct observation of global conformational change of membrane-embedded integrin upon binding to its physiological ligand.

Author Contributions—A. D. performed the data collection and image analysis, F. Y. prepared and characterized the integrin nanodiscs, D. W. T. prepared the specimens used for electron microscopy, G. H. carried out preliminary studies investigating feasibility, A. D., F. Y., D. W. T., M. H. G., and K. A. T. wrote the paper, and M. H. G. and K. A. T. supervised the experimental work in their respective laboratories.

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