Multiscale simulations suggest a mechanism for integrin inside-out activation

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Integrins are large cell-surface adhesion receptors that can be activated to a high affinity state by the formation of an intracellular complex between the integrin β-subunit tail, the membrane, and talin. The F2 and F3 subdomains of the talin head play a key role in formation of this complex. Here, activation of the integrin αIIb/β3 dimer by the talin head domain was probed using multiscale molecular dynamics simulations. A number of novel insights emerge from these studies, including (i) the importance of the integrin αIIb subunit F992 and F993 residues in stabilizing the “off” state of the αIIb/β3 dimer, (ii) a crucial role for negatively charged groups in the F2-F3/membrane interaction, (iii) binding of the talin F2-F3 domain to negatively charged lipid headgroups in the membrane induces a reorientation of the β transmembrane (TM) domain, (iv) an increase in the tilt angle of the β TM domain relative to the bilayer normal helps to destabilize the α/β TM interaction and promote a scissors-like movement of the integrin TM helices. These results, combined with various published experimental observations, suggest a model for the mechanism of inside-out activation of integrins by talin.

Integrins are heterodimeric (αβ) cell-surface receptors; each subunit contains a large extracellular (ecto) domain, an α-helical transmembrane (TM) domain, and a cytoplasmic domain (1). Integrins are crucial for signal transduction events involved in cell adhesion, migration, and differentiation (2–5). In mammals, combinations of the 18 α and 8 β subunits may form at least 24 different integrins. TM signaling can occur via outside-in (6) and inside-out pathways. In inside-out activation, the cytoplasmic tail of the integrin β subunit forms a complex with a cytoplasmic protein, talin, at the membrane surface (7–14).

Talin consists of a flexible rod and a globular head that has four subdomains (F0 to F3) (15, 16). The F3 subdomain alone is sufficient for αIIbβ3 integrin activation (17), but interactions between a positively charged patch in the F2 surface (14, 18) and negatively charged groups in the membrane are also crucial for integrin activation and clustering (19). Structural and functional studies suggest that the F0-F1 subdomain also contributes to the activating membrane complex (16, 20), partly via a flexible loop in the F1 domain. PIP2 is also believed to play a role (19).

The inactive state of integrins is maintained by intersubunit associations in the TM (21–26) and cytoplasmic regions (27–29) of the two integrin subunits. Integrin activation is believed to involve dissociation or destabilization of the two integrin TM helices (29–33), followed by rearrangement of the ectodomain to an extended conformation (34–37). Recent NMR and modeling studies (24) revealed similar structures for the αIIb and β3 TM helices in a membrane-like environment. Two main interaction surfaces between the two TM helices of the αIIb/β3 complex were identified in the NMR structure. These were called an outer membrane clasp (OMC), in which close packing of the helices is facilitated by a GxG motif in αIIb, and an inner membrane clasp (IMC), which involves interactions of F992 and F993 in αIIb with the β tail together with a salt bridge between αIIbR995 and β3D723 (24).

Different models have been proposed for structural changes in the membrane region during activation, including “piston” and “scissors” movements of the TM helices (38). However, more recent studies (7, 23, 29, 32, 39, 40) have been interpreted in terms of a model in which the binding of talin to the integrin β cytoplasmic tail and the TM domain disrupts the OMC and IMC interactions between the two TM helices, leading to helix separation and subsequent switching of the integrins to a high affinity state (14). Resolution of the underlying mechanism of activation thus requires information about the conformational dynamics of integrin/talin/membrane interactions that is not readily available from experimental methods.

Molecular dynamics (MD) simulations (41) can play a key role in understanding the dynamic interactions of membrane proteins with their environment (42). They have been used to investigate the interactions of TM helices with one another in a bilayer environment (43) as well as the interactions of peripheral proteins with membranes (44–47). MD studies of the F2-F3 domain of talin (18) in a bilayer demonstrated a key role for electrostatic interactions between the protein and anionic lipid headgroups.

Here, we use a multiscale MD simulation approach (48, 49) to explore the conformational dynamics of the interaction of the integrin TM helix dimer with talin F2-F3 and a lipid bilayer. The aim was to understand the inside-out activation mechanism. On the basis of these studies, we suggest how talin can weaken the α/β TM association and promote an integrin conformation that corresponds to a high affinity state.

Results and Discussion

Conformation and Dynamics of the Integrin αIIb/β3 TM Helix Dimer

As a first step toward understanding how talin interactions may modulate the behavior of the integrin αIIb/β3 TM helix dimer, we simulated the conformational dynamics of the TM helix dimer. We started with the coordinates of the NMR structure [Protein data Bank (PDB) ID code 2K9J], which is probably a reasonable approximation of the low-affinity integrin “off” state. Coarse-grained (CG)-MD simulations were used to optimize the position of the TM helix dimer relative to a palmitoyl-2-oleoyl-sn-glycerol-3-phosphocholine/palmitoyl-oleoyl-phosphatidylglycerol (POPC/POPG) bilayer, while restraining the αIIb TM helix to be perpendicular to the bilayer. This orientation of αIIb is suggested by the NMR structures of the αIIb helix (50) and the αIIb/β3 dimer in bicelles (24). The resultant CG system (i.e., helix dimer plus bilayer) was converted to atomic resolution (49) to form the starting point for 3 × 10^6 ns simulations.

In these αβ1-AT (Table 1) simulations, the TM dimer remained stable (Fig. 1A). Both IMC interactions [i.e., the αIIbR995-β3D723 salt bridge and the αIIb(F992,F993)-β3 aromatic cluster interactions] and the OMC interaction, involving the αIIb GxG motif, were maintained throughout all three simulations.

Author contributions: A.C.K., I.D.C., and M.S.P.S. designed research; A.C.K. performed research; A.C.K., I.D.C., and M.S.P.S. analyzed data; and A.C.K., I.D.C., and M.S.P.S. wrote the paper.

The authors declare no conflict of interest.

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with a crossing angle of approximately $\alpha$ to tact, and helix packing in the OMC was not perturbed. Thus, the alteration of the conformational stability of the dimer, as the hydrophobic mutation disrupted the salt bridge but did not appear to greatly alter the aromatic cluster) were further explored by simulations of the OMC (magenta), and the phosphorus atoms of the POPC/POPG lipid bilayer (gray). (c) Helix crossing angle distributions for the $\alpha$-AT (blue), $\alpha$-F5 (i.e., $\alpha$llb995D; red), and $\alpha$-AT (i.e., $\alpha$llb992A,993A; green) simulations. A negative crossing angle corresponds to right-handed helix packing, a positive angle to left-handed packing.

Fig. 1. The $\alpha$llb/3 TM helix dimer. (A) Structure of the $\alpha$llb (blue) $\beta$ (red) TM helix dimer (PDB ID code 2K9J). Interacting residues in the OMC are in magenta, in the IMC class in orange, and the $\alpha$llb R995 to $\beta$ D756 salt bridge is in green. (B) Snapshot (100 ns) from the $\alpha$-AT simulation (i.e., WT; see Table 1 for details) showing the two TM helices, the interacting residues of the OMC (magenta), and the phosphorus atoms of the POPC/POPG lipid bilayer (gray). (C) Helix crossing angle distributions for the $\alpha$-AT (blue), $\alpha$-F5 (i.e., $\alpha$llb995D; red), and $\alpha$-AT (i.e., $\alpha$llb992A,993A; green) simulations. A negative crossing angle corresponds to right-handed helix packing, a positive angle to left-handed packing.

This suggests that release of the IMC alone may not be sufficient to disrupt all the $\alpha$llb/3 TM interactions.

The OMC is formed by a glycoporphin-like interaction (52), involving a GxG motif formed by residues G972 to G976 of the $\alpha$llb helix. Mutation of either G972 to serine ($\alpha$-G5-AT; Table 1) or of G976 to leucine ($\alpha$- AT; i.e., $\alpha$llb995D; red), and $\alpha$-AT (i.e., $\alpha$llb992A,993A; green) simulations. A negative crossing angle corresponds to right-handed helix packing, a positive angle to left-handed packing.

Interactions with Talin Perturb the Integrin TM Helices. We next explored how interactions with the F2-F3 domains of the talin head

### Table 1. Summary of simulations

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All simulations were of the protein complex in a POPC/POPG lipid bilayer. System sizes ranged from approximately 40,000 ($\alpha$-AT simulations) to approximately 100,000 ($\alpha$-F3 simulations). $\beta$- indicates a chimeric $\beta$ chain ($\beta$/1/1D) — see main text for details.

The tilt angles for the $\alpha$llb and the $\beta$ helices during the $\alpha$-AT simulations were approximately 5 and 30°, respectively, and the helix dimer maintained its right-handed packing with a helix crossing angle of approximately −35° (Fig. 1C).
modulate the TM helices in a lipid bilayer. If signaling from inside to outside is indeed mediated by a change in packing interactions of the TM helices, then one would anticipate that F2-F3 bilayer interactions (18) could perturb TM/TM and/or TM/bilayer interactions in the simulations. A short (30-ns) atomistic simulation of F2-F3 bound to the PC/PG bilayer revealed a small relative movement (approximately 6°) of the F2 subdomain relative to F3. This movement optimizes contacts with the lipids and is made possible by a flexible linker between the two subdomains. More detailed analysis of F2-F3/lipid contacts confirmed earlier results (18), showing a preference for the F2-F3 domain to interact with negative lipid headgroups.

We next investigated how the close association of the F2-F3 domain with an anionic bilayer might modulate the integrin TM domains, by performing simulations of F3 and of F2-F3 in complex with the β TM helix. The results were compared with the β TM helix alone (β-AT, β-F3-AT, and β-F2F3-AT in Table 1). The β helix was modeled on the structures of the αIIb/β3 TM helix dimer [PDB ID code 2K9J (24)] and of talin F2-F3 bound to the cytoplasmic tail of the integrin β1D [PDB ID code 3G9W (14)]. In the β-F2F3 simulations, there was a significant increase (approximately 15°) in the tilt angle of the β TM helix relative to that observed in either the β-AT or αβ-AT simulations (Fig. 2). Further analysis of this change revealed no significant movement of the F3 subdomain relative to the β helix. In contrast, the F2 subdomain rotates approximately 10° relative to F3, possibly to optimize the interactions with the bilayer. Examination of H-bonds and close contacts reveals that the F2-F3/bilayer interface in the β-F2F3-AT simulation is similar to that identified in simulations of F2-F3 and bilayer alone. A comparable movement of the β TM helix was seen in the β-F3-AT simulations. Again an increase in β TM helix tilt angle of approximately 15° was seen over the course of the simulation.

The β TM helix also adopted a similar tilted orientation (approximately 37°) in simulations (β-AT) of just the β TM in a bilayer. Taken together, these three simulations indicate that the β TM helix prefers to adopt a higher tilt angle relative to the bilayer than in the αIIb TM dimer. The drive for the β TM helix to a higher tilt angle may arise because the hydrophobic helix length (approximately 40 Å) is better accommodated; the tilted orientation also places the W715 sidechain in the lipid headgroup region and allows the K716 sidechain to interact with lipid phosphate groups. This tendency of the β TM helix to adopt a tilted orientation in the membrane is consistent with other studies of integrin TM domains (19, 55). The higher tilt angle of the β TM also allows the talin F2-F3 subdomain to optimize its interactions with the anionic bilayer surface.

Changes in Integrin TM Helix Packing Induced by Talin F2-F3. We next sought to explore the effect of the talin F2-F3 subdomain on TM helix packing and/or the β TM tilt. To do this, the NMR structure of the αIIb/β3 TM helix dimer [PDB ID code 2K9J (24)] and the X-ray structure of F2-F3 in complex with the cytoplasmic tail of integrin β1D [PDB ID code 3G9W (14)] were combined. The backbone coordinates of the helical region of β1D from the F2-F3-β1D structure and the β3 helix from αβ TM dimer structure were aligned. After the alignment, the β3 residues from T720 to F727 were removed, thus keeping the equivalent β1D residues in the positions found in the X-ray structure of F2-F3 in complex with the cytoplasmic tail of integrin β1D (PDB ID code 3G9W). This resulted in a chimeric structure (β3 residues 688–719/β1D residues 753–787) for the β subunit that combined the TM domain of β3 and the C-terminal tail of β1D. We note that the TM domains of β3 and β1D are very similar in sequence (90% similarity; 52% identity) and that the crucial salt bridge in the IMC, between β1D D759 and αIIb R995, is preserved.

To explore the stability of the IMC αIIb R995/β1D D759 salt bridge in this model of αβ-F2-F3, four atomistic simulations, each of 20-ns duration, were performed for the model in a POPC/POPG bilayer. Rapid rearrangement of the salt bridge was observed, due to the interactions of the αIIbR995D sidechains with a positively charged loop (residues M322–L328) in the talin F3 subdomain. An additional control was performed, using a second model with a slightly different β chain chimera (β3 residues 688–727/β1D residues 764–787). In this model, the homologous salt bridge is formed between the sidechains of β3 D723 and αIIb R995. A similar rapid reorganization of the salt bridge was again observed in the presence of F2-F3, resulting in interactions of the αIIb R995 with the charged loop (residues M322–L328) on the F3 subdomain.

In order to explore the possibility of longer timescale (>0.1 μs) changes in the conformation of the αβ-F2-F3 complex, a multiscale approach was taken, combining CG simulations to explore longer timescale events with subsequent atomistic MD simulations to refine the resultant models. A CG representation of the αβ-F2-F3 model structure was embedded in a POPC/POPG bilayer using our self-assembly protocol (56). Elastic network model restraints between the αIIb subunit and the β/F2-F3 complex were removed. Five simulations (each of 5 μs) were performed. These simulations suggested that talin F2-F3 destabilizes the αIIb/β1D dimer: the number of contacts between the αIIb and β helices was greatly reduced at the dimer interface with both the IMC and OMC clasps being disrupted. In particular, the close packing of the two helices via the αIIb GxG motif and the interactions of the αIIbF992, F993 aromatic cluster and the αIIbR995 with the β tail were disrupted. The αIIb helix maintained its orientation perpendicular to the bilayer (tilt angle approximately 10°), whereas the β helix tilt increased (tilt angle approximately 28°).

Analysis of the movements of the β and F2-F3 domains relative to the α TM helix suggests how F2-F3 may perturb the αIIb/β interactions. Each of the three CG simulations showed similar
persistence of the αIIb/β dimer. The movements of β, F2, and F3 relative to α are shown in Fig. S1.) A 25° ± 4° rotation of F2 in a plane perpendicular to the bilayer normal (i.e., parallel to the lipid–water interface) is observed, along with a 22° ± 1° rotation of F3 in the same plane. This, in turn, induced a 22° ± 1° rotation of the β subunit perpendicular to the membrane in the same direction as for F3 (Fig. S2). This rotation resulted in disruption of the αIIb/β TM helix dimer interactions in both the IMC and OMC regions and the formation of a new interface between the two helices (see Fig. 3A).

Three representative CG structures were selected for detailed investigation by atomistic simulations. The starting structures for the three simulations were: (i) αII/F2F3c-AT, a structure with the IMC and OMC interactions preserved, taken to represent a “closed” (c) model; (ii) αII/F2F3p-AT, a structure with the OMC disrupted but the IMC interaction intact, taken to represent a “partially” (p) disrupted model; and (iii) αII/F2F3o-AT, a structure with both the OMC and IMC disrupted taken to represent an “open” (o) model. All three of these models were converted back to atomistic resolution using our CG2AT protocol (49) and were then subjected to 100-ns simulations (Table 1).

In simulation αII/F2F3c-AT, the helix dimer was observed to stay closed, but the β helix tilt angle increased to approximately 35° relative to the bilayer normal. This increase in the β TM tilt angle and the preservation of the tight interactions between the TM helices thus increased the αIIb helix tilt angle relative to the bilayer normal to approximately 20°. In simulation αII/F2F3p-AT, the αIIbR995D–βD759 salt bridge was disrupted but the interactions between the αIIb/F992/F993 residues and the β tail remained, in good agreement with results from simulations of mutations in the αIIb/β3 dimer, discussed above.

In contrast, simulations based on the open structure, αII/F2F3o-AT, revealed more complex behavior. The interactions in both the OMC and IMC clasp remained disrupted, and the αIIb/β3 dimer remains destabilized (Fig. 3B, C). Alignment of the αIIb helix at the start and end of these simulations revealed that the F2 domain had rotated perpendicular to the bilayer normal. This, in turn, causes rotation of F3 in the same plane but in the opposite direction, whereas the β tail rotated in the same direction of F3. Interestingly, the dimer destabilization in αII/F2F3o-AT allowed further dissociation of the helices in the OMC clasp region. In particular, the distance between the αIIb and the β helices in this region is increased to approximately 20 Å, and therefore the angle between the two TM helices also increased. However, the IMC region remained relatively intact, and thus a “scissor” movement of the two helices was observed, with the center of the scissor in the region of the IMC. The new interface in this contact region involves αIIb L978, L979, I982, A986, V990 and the β I707, G708, A711, T715, M718 residues (see Fig S3).

We note that during these movements the interactions of the F2-F3 and of the β C-terminal tail with the bilayer were preserved. In particular, the F2–F3 domains interacted preferentially with the negatively charged lipids in the membrane. This suggests that protein/lipid interactions must be considered when analyzing possible activation mechanisms of integrins.

**Suggested Mechanism.** Structural studies (7, 14, 16, 24, 25, 57) have greatly increased our understanding of integrin activation from inside the cell. However, static structures fall short of revealing dynamic mechanisms of activation. Using multiscale MD simulations, we have revealed further details of the mechanism whereby the talin head domain, the lipid bilayer membrane and the αβ TM helix dimer together interact to enable inside-out activation of integrins. In particular, we propose a mechanism whereby talin F2–F3 interactions lead to a conformational change in the αβ TM helix dimer, which may act as a trigger for inside-out activation of the integrin ectodomain. In this model (see Fig. 4A) the first stage is when F2–F3 binds to anionic lipids via electrostatic interactions (18). This is determined by the F2 subdomain, which forms basic sidechain/anionic headgroup interactions. Talin F3 binds to the β integrin C-terminal tail in three areas: the membrane distal NPxY motif, the β1D W775 (β3 W739) residue, and the β tail membrane-proximal region. Previous experimental studies have shown that a tyrosine to alanine mutation in the distal NPxY motif or the β3 R358A mutation inhibits integrin activation (7, 8, 58–60). Thus, F2–F3 associates with the αβ TM helix dimer via an interaction between F3 and the cytoplasmic tail of β. A recent NMR study (61) suggested that, in the resting integrin off state, the β TM region is bent at residue R724, allowing the C-terminal cytoplasmic β tail to interact with the membrane via two helices. Promotion of a helical structure in the R724 region would extend the β helix and allow binding of F3. This, together with F2–F3 interactions with the membrane bilayer, then leads to a scissoring motion of the αβ TM helix dimer about the β2 TM region such that the OMC is disrupted, and the N-termini (extracellular) separation of the TM helices is increased (Fig. 3A, Dataset S1, and Dataset S2).

The scissoring motion that was revealed in the αII/F2F3o-AT simulations reflects the tendency of the β TM helix to be more tilted in a lipid bilayer than it is in the off state. This tilting motion may be seen quite clearly by monitoring the distances between the α and β TMs. As shown in Fig. 4A and B, distances in the IMC region (d3, d4) are maintained, whereas the distances (d1, d2) at the N termini of the helices increase by about 6 Å from start to end of the simulation.
the role of the IMC and OMC clasps and explain much of the activation (23).

Suggest that OMC and IMC interactions are required for integrin activation. There is ample experimental data to support this suggestion that disruption of the F3/β tail interface or on the talin F2-F3 interface with the membrane reduce the ability of talin to activate integrins (7–14). Fluorescent resonance energy transfer (FRET) studies (29) using αL and β integrin subunits tagged with fluorescent proteins suggested significant reduction in the FRET signal in the presence of talin, and this was interpreted in terms of tail separation. A scissor movement around the IMC might be expected to separate the scissor handles as well as the blades. Fig 4B (d4) does not indicate an increase in separation, but the αLβ995-ββ723R is broken and the largely cytoplasmic α and β tails are flexible. So increased tail separation is not inconsistent with our model.

Recent structures of integrin ectodomains show that the membrane-proximal C-terminal regions of the two ectodomain subunits are in close proximity (26, 36, 67). It is therefore plausible that the scissor movement that causes dissociation of the OMC clasp promotes a rearrangement of the integrin ectodomain.

In summary, our results suggest a mechanism for integrin inside-out activation that explains the role of the talin head domain, the membrane and the α/β TM interactions. We have shown that the α/β TM helix dimer is conformationally stable in a lipid bilayer on a 100-ns timescale, that the β TM helix has a propensity to tilt relative to the lipid bilayer and that this tilt motion is resisted by the presence of the α TM helix. Binding of talin destabilizes interactions between the TM helices and permits a scissor-like motion in part driven by the propensity of the β helix to tilt. This model shows how protein/membrane interactions can disrupt the TM helix interactions of complex receptors and potentially lead to structural changes in ectodomains. It also demonstrates the strength of multiscale simulations for developing models of the conformational dynamics of a complex membrane receptor.

Materials and Methods

CG-MD Simulations. CG-MD simulations were performed using a local variant (68) of the Martini force field (69) with an elastic network applied to backbone particles using a cutoff distance of 7 Å, and the protein inserted in a POPC/POPG bilayer. The system was solvated, energy minimized for 200 steps, and equilibrated for 5 ns with the protein backbone particle restrained. After that, five production simulations were carried out for 5 μs each.

All CG-MD simulations were performed using GROMACS 3.3.3 (www.gromacs.org) (70, 71). A Berendsen thermostat (72) (coupling constant of 1.0 ps; reference temperature 310 K) and barostat (coupling constant of 1.0 ps; compressibility value of 5.0 x 10^-6 bar^-1; reference pressure 1 bar) were used. The integration time step was 40 fs. Lennard-Jones and Coulombic interactions were shifted to zero between 9 and 12 Å, and 0 and 12 Å, respectively. CG to atomistic conversion used a fragment-based approach (49).

Atomic MD Simulations. Atomic MD simulations were performed using the GROMOS96 43a1 force field (73). The Parinello–Rahman barostat (74) and Berendsen thermostat (72) the LINCS algorithm was used to constrain bond lengths (75), and particle mesh Ewald particle mesh was used to model electrostatics up to 10 Å. A 10-Å cutoff distance was also used for van der Waals interactions. Systems were equilibrated for 2 to 5 ns with the protein Ca atoms restrained, followed by unrestrained MD simulations as in Table 1. Analyses used Gromacs, VMD (76), and locally written codes.

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**Supporting Information**

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Fig. S1. Simulations of the αIIbβ3 transmembrane (TM) helix dimer in complex with talin F2-F3. (A) Relative movements of the β, F2, and F3 domains in the αIIb-F2F3-CG simulation. The displacements of F2 (cyan), F3 (yellow), and the β TM helix (red) over the course of the simulation are shown as arrows mapped, the lengths of which represent the displacements. For this calculation, the αIIb TM coordinates were superimposed. (B) Initial configurations of the atomistic simulations of αβ-F2F3c-AT ("closed"; see Table 1 for details and nomenclature used for simulations), αβ-F2F3p-AT ("partial"), and αβ-F2F3o-AT ("open"); these configurations were obtained by clustering the coarse-grained–molecular dynamics simulations and converting them to AT resolution. The color scheme is outer membrane clasp (magenta), inner membrane clasp (orange and green), F2 (cyan), F3 (yellow), β TM (red), and αIIb TM (blue).

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Fig. S2. Alignment of the αIIb subunit from the initial F2-F3 + αIIb/β3 model (blue) and at the end of the αβ-F2F3o-AT (red) simulation, demonstrating the rotation of the β subunit perpendicular to the bilayer normal.
**Fig. S3.** Interface between the αIIb (blue) and the β3 (red) TM domains at (A) the beginning and (B) the end of the αβ-F2F3o-AT (open) simulation. The residues that make up the interaction surface in both subunits are shown in van der Waals format. The F2-F3 domains are omitted for clarity.

**Fig. S4.** Superposition of the β TM region (residues 697–726) from the αβ-F2F3p-AT simulation (blue) with the αIIβ/β3 (2KNC, orange) and the αIIβ/β3 (2K9J, red) structures. The alignment shows that the αIIβ/β3 (2KNC) structure is similar to the partial disrupted state (the αβ-F2F3p-AT simulation) and could therefore correspond to an intermediate state. The F2-F3 domain is omitted for clarity.

**Other Supporting Information Files**
Dataset S1 (TXT)
Dataset S2 (TXT)