The mechano-sensing role of the unique SH3 insertion in plakin domains revealed by Molecular Dynamics simulations

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The plakin family of proteins, important actors in cross-linking force-bearing structures in the cell, contain a curious SH3 domain insertion in their chain of spectrin repeats (SRs). While SH3 domains are known to mediate protein-protein interactions, here, its canonical binding site is autoinhibited by the preceding SR. Under force, however, this SH3 domain could be released, and possibly launch a signaling cascade. We performed large-scale force-probe molecular dynamics simulations, across two orders of magnitude of loading rates, to test this hypothesis, on two prominent members of the plakin family: desmoplakin and plectin, obligate proteins at desmosomes and hemidesmosomes, respectively. Our simulations show that force unravels the SRs and abolishes the autoinhibition of the SH3 domain, an event well separated from the unfolding of this domain. The SH3 domain is free and fully functional for a significant portion of the unfolding trajectories. The rupture forces required for the two proteins significantly decrease when the SH3 domain is removed, which implies that the SH3 domain also stabilizes this junction. Our results persist across all simulations, and support a force-sensing as well as a stabilizing role of the unique SH3 insertion, putting forward this protein family as a new class of mechano-sensors.
the role of the SH3 insertion in spectrin repeats has remained largely elusive, especially as in this case, the canonical binding site of the SH3 is inhibited through interactions with the preceding spectrin repeat. The initial suggestion was that the purpose of the SH3 insertion is to stabilize the junction of the two adjacent SRs, but the fact that SH3 domains are generally interacting with binding partners suggests an alternative (or additional) role, namely, that the SH3-SR interface can be opened and the SH3 domain thereby activated under force.

In fact, the hypothesis of force activation has some experimental support for at least two proteins featuring the unique SH3 insertion into spectrin repeats. For nonerythroid α-spectrin, a binding partner for the SH3 domain has been experimentally observed in the integrin signaling complexes, and similarly, the Ic isoform of plectin has been shown to interact with the microtubuli-associated protein 1 (MAP1) through the SH3 domain. While neither of these studies directly show that this process is force-induced, they provide evidence for that the SH3 domain is released at least part of the time and is capable of binding other proteins.

Here, we study the force response and putative mechano-sensing mechanisms of desmoplakin and plectin, two members in the plakin family. Both proteins have been structurally resolved and are known to have roles in tissue integrity: 25 different disease mutants of desmoplakin related to arrhythmogenic right ventricular cardiomyopathy (ARVC) have recently been identified, while the knockout of plectin has been shown to disturb adherens junctions (AJs) by abolishing the vimentin-actin crosslinks.

Desmoplakin is an obligate linker protein in the desmosome, linking the inner dense plaque (at the inner side of the cellular membrane) to intermediate filaments. Desmosomes are specific to epithelial and cardiac tissue. The central plakin domain of desmoplakin contains six spectrin repeat domains (called SR3 through SR8). The third spectrin repeat, SR5, contains the mysterious SH3 domain insertion that is inhibited by interacting with the previous spectrin repeat, SR4.

Figure 1. Plakin family proteins contain a curious SH3 domain insertion. (a) The three systems we study using force-probe molecular dynamics: (b) four spectrin repeats and an SH3 domain (only desmoplakin). (c) two spectrin repeats and an SH3 domain (desmoplakin and plectin), and (d) only two spectrin repeats with the SH3 domain removed (desmoplakin and plectin). The blue surface denotes the SH3-SR4 interface for cases. (e) The assignment of helices 4A-C and 5A-C in desmoplakin with the SH3 domain removed, for clarity.
Plectin is a cytolinker protein present at several kinds of cellular junctions: AJ, focal adhesions, and also hemidesmosomes. Plectin also contains a plakin domain with a very similar structure, including the SH3 domain insertion. The immediate proximity of this insertion, namely, the spectrin repeats SR4–SR5, has a high structural similarity to desmoplakin and both proteins have high sequence identity to other plakin family proteins (23–64%, see SI).

Given the role of the SH3 domain as an adaptor protein, we can reasonably infer that the plakin family members’ SH3 domain can generally serve as a signaling agent once released from the spectrin repeat. We here set out to study this possibility using simulations, in particular, force-probe molecular dynamics (FPMD). FPMD, also known as steered MD, has been used to study the individual unfolding of both SRs and SH3, often in conjunction with experiments. In the following, we will apply this method to different constructs of desmoplakin and plectin (Fig. 1) to answer the following questions: (i) do the domains unfold simultaneously or in different ways for the two proteins. For desmoplakin, we solvate the protein so the initial conformation is at least at a 5.0 nm distance from its periodic image, which results in a box size of about 100 \( \times \) 10 \( \times \) 10 nm\(^3\) and a total number of 1.2 M atoms. For plectin, we use a two-step procedure due to a high variance of orientations during unfolding (see discussion below). First, we extract the central frames of the clusters obtained from the equilibrium simulations, keeping only the protein atoms. We align the protein in such a way as to have the x-axis parallel with the vector connecting the centers of mass of these two pulling groups. We then re-solvate the protein in an oblong cuboid box where the x-axis is long enough to contain the unfolded polypeptide. We do this in two slightly different ways for the two proteins. For desmoplakin, we solvate the protein so the initial conformation is at least at a 5.0 nm distance from its periodic image, which results in a box size of about 100 \( \times \) 10 \( \times \) 10 nm\(^3\) and about 800k atoms (for the 4 SR + SH3 simulations) or 200k atoms (for the other simulations). After solvation and the addition of ions, we first perform an energy minimization using the steepest-descent method with a step size of 1 pm, tolerance of 20 kJ/mol/nm, and a maximum of 500000 steps. Thereafter, we perform 500 ps of NVT and 500 ps of NPT equilibration, both with a harmonic restraint on the protein atoms of 1000 kJ/mol/nm\(^2\). After 300 ns of equilibrium simulation, we choose the top 10 clusters in a cluster analysis to obtain the starting frames for FPMD simulations. The rupture forces from FPMD show no significant correlation with the time at which the initial frames have been extracted from the equilibrium simulations. For the “hybrid” model of plectin, we perform an equilibration of 100 ns and a further 900 ns of production run, on which we do the previously mentioned cluster analysis to obtain the starting frames for FPMD simulations. We perform additional simulations of a plectin mutant as described in the supplementary information.

**Methods**

We use GROMACS version 5.0 with the Amber99SB-ildn force field and TIP3P water molecules. All bonds are frozen in our simulations using the LINCS procedure. We use the integration time step of 5 fs made possible by the use of virtual sites for all hydrogen atoms. Further Force-probe simulations (FPMD) show no significant correlation with the time at which the initial frames have been extracted from the equilibrium simulations. For the “hybrid” model of plectin, we perform an equilibration of 100 ns and a further 900 ns of production run, on which we do the previously mentioned cluster analysis to obtain the starting frames for FPMD simulations. We perform additional simulations of a plectin mutant as described in the supplementary information.

We use the Particle Mesh Ewald (PME) technique to treat long-range electrostatic interactions, with a grid spacing of 0.16 nm and cubic interpolation. Neighbor lists with a cutoff of 1 nm with the Verlet cutoff are updated every 25 time steps (0.125 ps).

We start our desmoplakin simulations from the crystal structure 3R6N, and those of plectin from the crystal structure 3PE0. The more recent 5J1H crystal structure features slight differences in one of the spectrin repeats but lacks the SH3 domain. We also repeat a part of the plectin simulations using an alternative model based on the 3PE0 (for the SH3 domain) and the recently published 5J1H crystal structures (for the SRs) as templates, which we denote as “hybrid” model in the following. We solvate the two structures in water and proceed to add Na\(^+\) and Cl\(^-\) ions with a concentration of 0.15 M.

In our equilibrium simulations, we use a dodecahedron box with at least 3.0 nm between periodic replicas of the protein, resulting in a system containing about 640k (for the 4SR + SH3 simulations) or 200k atoms (for the other simulations). After solvation and the addition of ions, we first perform an energy minimization using the steepest-descent method with a step size of 1 pm, tolerance of 20 kJ/mol/nm, and a maximum of 500000 steps. Thereafter, we perform 500 ps of NVT and 500 ps of NPT equilibration, both with a harmonic restraint on the protein atoms of 1000 kJ/mol/nm\(^2\). After 300 ns of equilibrium simulation, we choose the top 10 clusters in a cluster analysis on backbone motion with a cutoff of 0.09 nm on 6001 frames (extracted every 50 ps) as initial frames for the FPMD simulations. The rupture forces from FPMD show no significant correlation with the time at which the initial frames have been extracted from the equilibrium simulations. For the “hybrid” model of plectin, we perform an equilibration of 100 ns and a further 900 ns of production run, on which we do the previously mentioned cluster analysis to obtain the starting frames for FPMD simulations. We perform additional simulations of a plectin mutant as described in the supplementary information.

**Force-probe MD.** In all of our pulling simulations, we subject the atoms of the N- and C-terminal residues to a harmonic potential. The minimum of the harmonic potential is at the position of the center of mass of the residues at \( t = 0 \) and each spring is moving at velocities \( v = \pm \frac{1}{2} \mu_{\text{pull}} \) to reduce protein-water friction. The force constant is \( k = 500 \text{kJ/(mol \cdot nm)} \approx 830 \text{pN/nm} \) in all simulations.

After the equilibrium simulations, we extract the central frames of the clusters obtained from the equilibrium simulations, keeping only the protein atoms. We align the protein in such a way as to have the x-axis parallel with the vector connecting the centers of mass of these two pulling groups. We then re-solvate the protein in an oblong cuboid box where the x-axis is long enough to contain the unfolded polypeptide. We do this in two slightly different ways for the two proteins. For desmoplakin, we solvate the protein so the initial conformation is at least at a 5.0 nm distance from its periodic image, which results in a box size of about 100 \( \times \) 10 \( \times \) 10 nm\(^3\) and about 800k atoms (for the 4 SR + SH3 simulations) and then, when the width of the partially unfolded protein approaches 60 nm, we re-solvate the protein into a box size of 100 \( \times \) 8 \( \times \) 8 nm\(^3\). This involves a...
We define refer to the distance to the transition state, the spring stiffness (kept constant in our case), with $Q_1$, $Q_3$ being the first and third quartiles in the equilibrium increase the boxes to this size. To understand the effect of the SH3 domain on the unfolding trajectories, we repeat the same pulling simulations containing the SH3 domain: SR4 and SR5. For desmoplakin, we observe (Fig. 2b, left) a very clear order of intra- and inter-domain unfolding events: First, SR4 unfolds at extensions of ~15–25 nm, then, the SH3-SR4 interface is severed and the SR5 unfolds at approximately the same time of extensions of ~40–60 nm. Between the moment in which the SH3 domain is released (~45–50 nm) and when it starts unfolding (~70–75 nm), the SH3 domain is free to bind to partners. We do not observe SH3 domain unfolding occurring before or during “activation” in any of the simulations. This would have been a distinct possibility, since helix 3 C often unfolds or loses contact in the very first unfolding event and thereby the SH3 domain is directly under stress. The unfolding pathways of plectin (shown in Fig. 2b, right) are largely similar to those of desmoplakin. In particular, the unfolding of the SH3 domain still happens as the last event and, for at least an extension interval of 30 nm, it is intact and not autoinhibited. However, in 8 out of the 33 unfolding trajectories, it loses the interface with SR4 at even earlier moments in time, therefore being available for binding at an even earlier point. Further analysis on the residues with the interactions that last longest between the SH3 domain and the helix 4C5A reveals that this interface is lost in a very similar way in both proteins (see Supporting Information for details) and the most important residues involved are either identical or strongly similar.

To understand the effect of the SH3 domain on the unfolding trajectories, we repeat the same pulling simulations with the two spectrin repeats with the SH3 domain removed. Fig. 2c shows that without the SH3 domain, for both plectin and desmoplakin, the inter-domain interface is the very first feature to be lost during the unfolding pathways. Compared to the previous cases, where the SH3-SR4 interface was usually only lost after SR4 had unfolded, here, the two spectrin repeat domains immediately lose contact and unfold independently. In the case
of desmoplakin, there is a clear preference (about 90%): SR4 unfolds before SR5, while for plectin the order is random. We can estimate the difference between the unfolding barriers of the two spectrin repeats, $\Delta \Delta G = \Delta G_{SR5} - \Delta G_{SR4}$ in desmoplakin from the 3/33 ratio to be most likely $\Delta \Delta G = 1.2$ kcal/mol (SR4 having a lower barrier than SR5), with the 95% confidence interval between $\Delta \Delta G = 0.7 \ldots 2.0$ kcal/mol (see SI.

Figure 2. Unfolding events observed in all single force-probe simulations. Boxes extend from the start to the end of the loss in contact area, either within a single domain (SR3, SR4, SR5, SR6, and SH3) or at inter-domain interfaces (SR3-SR4, SH3-helix 4C5A, and SR5-SR6), see Methods for details. Each set of horizontally aligned boxes correspond to one FPMD trajectory. The unfolding events are represented as a function of inter-spring distance, which is very close to the net increase of the protein end-to-end distance. Three sets of systems are presented: (a) four spectrin repeats and an SH3 domain (only desmoplakin (20 runs)), (b) two spectrin repeats and an SH3 domain (left: desmoplakin and right: plectin (33 runs each)), and (c) only two spectrin repeats with the SH3 domain removed (left: desmoplakin and right: plectin (33 runs each)). The purple area in subplot a. shows that the domains SR3, SR6 and the associated interfaces are first to unfold (until $v_t \approx 60$ nm), while the green areas for subplots a, b show the minimum active area, i.e., the region of the unfolding trajectories after the SH3–4C5A interface is lost and before the beginning of the SH3 domain unfolding.
for the Bayesian arguments in support of this). For plectin, where the preference is 17/33 for SR5, the difference of the barriers has the mode at ∆∆G = −0.4 kcal/mol, and with the 95% confidence interval between ∆∆G = −0.04 ± 0.4 kcal/mol.

The SH3 insertion stabilizes the plakin domain. The fact that the SH3 domain remains intact until the very last part of the unfolding simulations supports the idea that the insertion has a force-sensing role. In none of the simulations did we observe the SH3 domain to unfold before it loses contact with the spectrin repeats. On the contrary, for a significant part of the unfolding time, it is both fully folded and at the same time its binding site fully solvent-exposed.

We next asked whether the SH3 domain also affects the rupture forces of a plakin domain. Fig. 3 summarizes how the rupture force observed for the two proteins depends on the pulling velocity (i.e., the loading rate) and the presence of the SH3 domain. Inserting the SH3 domain into the SRs of both plakin domain constructs increases the rupture force significantly along the whole range of loading rates. The rupture force increase amounts to 48% for desmoplakin and 31% for plectin on average. Clearly, one of the effects of the SH3 domain is the stabilization of this junction.

This consistent drop in rupture forces supports the notion that one main role of this SH3 domain is the mechanical stabilization of the plakin domain. Furthermore, it is noteworthy that the rupture forces for the loss of the two different SH3-SR4 interfaces probed here are significantly higher than any rupture force corresponding to the unfolding of any single domain of the five considered here (SR3-6, SH3). The stabilizing role of the SH3 domain is also corroborated by the fact that all of the simulated unfolding trajectories of the large construct for desmoplakin showed the outer two domains to fully unfold before the unfolding of the central two begins (see Fig. 3a). Thus, the SRs not in contact with the SH3 domain have a lower mechanical resistance than those with a direct interface with the SH3 domain.

Rupture forces show desmoplakin to be more robust than plectin. We observe that desmoplakin, on average, has rupture forces about 10–20% higher than plectin (Fig. 3). The higher stability of desmoplakin as compared to plectin is significant at the three fastest pulling velocities (where 10 pulling simulations per system have been performed), and is more pronounced when the SH3 domain is present. Interestingly, while the fit with the Bell model (see Eq. 1) is not entirely satisfactory, the observed slope for the four different considered systems is quite consistent: 200–270 pN/(order of magnitude in velocity). This suggests the higher robustness of desmoplakin to also hold at lower pulling velocities such as those probed experimentally.

The similar slopes from the Bell fit can also be interpreted as similar conformational changes at the rate limiting step, supporting the notion of highly resembling unfolding mechanisms of the two plakin domains investigated here. Extrapolating these trends across several orders of magnitude, however, would result in very high unfolding rates at zero force (∼1 µs⁻¹). We refrain here, however, from using more complicated rupture force-loading rate relations to avoid overfitting our data that is clustered along the high loading rate region and only use the fits to compare the stabilities of the four considered systems.
Alternative models of plectin have no major effect on rupture forces. The higher stability of desmoplakin over plectin is not easily comprehensible given the high structural homology of the two plakin domains. The two major structural differences between the considered desmoplakin and plectin fragments are: (i) a small ruptured helix at the N-terminus of the helix 5B and (ii) a different salt bridge network between the 4A-4B loop and the upstream SH3 loop. To see whether the significant differences in rupture force between desmoplakin can be accounted for by either of these observations, we repeat (at the highest loading rates only) the pulling simulations of plectin on suitably modified models. As Table 1 shows, however, the rupture forces do not significantly change compared to the ones of wild-type plectin. This implies that the details of the structural model have little influence and that the differences in stability of the two proteins have a different source.

Differences in force profiles correlate with a small cluster of sequence variations. Paying closer attention to the force profiles of the two considered proteins, we can notice that for desmoplakin, 32/33 of the trajectories feature at least three force peaks, while only 25/33 from plectin do (Fig. 3a). This behavior appears to correlate very closely with a tilting motion of the partially unfolded proteins we observe more frequently for plectin. While in many (or most as for desmoplakin) trajectories the SR helices remain oriented along the pulling direction (Fig. 4a), some unfolding events feature a pronounced tilting of the SRs away from the pulling direction, resulting in a 'tearing' or 'unzipping' of the remaining α-helices of the SRs (Fig. 4b). We can quantify this by measuring the maximum angle created by the vectors drawn by (i) the α-carbons of N375-P342 (for plectin, these are N858-P825) and the pulling axis and (ii) N375-Y403 (for plectin, N858-Y886) and the pulling axis (Fig. 4c). We denote these tilting angles α1 and α2, respectively. Fig. 4c shows that the magnitude of the maximum tilting very well predicts whether an unfolding happens at a high or low force in the second peak. In fact, the highest of the two maximum tilting angles has a receiver-operating characteristic of the high/low force of rupture events with an area under curve of 0.919 (see Supplementary Information for details). More tilting leads to unfolding at lower forces since tilting exposes the SH3-SR4 interface directly to the external force. Tilting also allows the gradual unzipping of SRs from as opposed to a shearing mechanism which requires higher forces, much like in the case of the β-barrel of GFP.

The higher prevalence of the tilting for plectin rather than desmoplakin can be rationalized by a small patch of sequence variations at the N-terminal of the helix 5 C (the orange patch in Fig. 4a). For plectin, this helix

| model                  | F_r (pN) | ε (pN) |
|------------------------|----------|--------|
| desmoplakin            | 659.5    | 13.5   |
| plectin 3PE0           | 534.3    | 16.6   |
| plectin 3PE0 + 5J1H    | 526.0    | 18.3   |
| plectin 3PE0 + mutations | 535.8    | 19.5   |

Table 1. Comparison of rupture forces and standard errors on the mean ε observed at 10 replicas each pulled at a velocity of 1 m/s for three different plectin models and desmoplakin.

Figure 4. The two different ways the SH3-SR4 interface can be lost: (a) “shearing” and (b) “tearing” (representative snapshots with red = SR4, green = SR5, blue surface = residues at helix 4 C in contact with the SH3 domain, and orange highlight: the N-terminal domains of helix 5 C, whose unfolding triggers the tilt and therefore the “tearing” topology), and (c) the maximum tilt angle observed for the 33 unfolding trajectories of desmoplakin (left) and plectin (right). Red points refer to force profiles with three distinct peaks and blue points show a lack of a middle peak. See also the ROC curve based on this plot in the Supplementary Information.
unfolds more often and allows the tilting to occur (since the helix loses contact to helices 5 A and B), while for desmoplakin, even in the cases where helix 5 C loses its secondary structure, it retains contacts with the rest of the spectrin repeat. As a consequence, the SH3 domain of plectin is activated with significantly less work. In addition, plectin unfolding and activation requires a lower rupture force in general (Fig. 3b), independent of the described tilting, further stressing its lower resilience against the external pulling force.

A summary of the various unfolding possibilities is given in Fig. 5. It is easy to see that plectin exhibits much more variation in unfolding pathways and that all force-resistance of the protein is lost when the SH3-SR4 interface is lost.

Conclusions

The location of an SH3 domain, an adaptor protein, in the middle of spectrin repeats, known as force-buffering structural elements is striking the eye. Remarkably, it is a conserved feature of the plakin family as well as α-spectrin. What is the evolutionary advantage of inserting a highly conserved and widespread protein-protein interaction domain, even more so in its auto-inhibited conformation, right into spectrin repeats, which elastically deform under tension? To address this question, we here analyzed the force response of the central plakin domain of two representatives of the plakin family, namely, desmoplakin and plectin. For both proteins, we identified a likely mechano-sensory role of the SH3 insertion: under force, this domain is invariably freed and can thereafter interact with binding partners (Fig. 5). We have not observed any trajectories in which the domain even partially unfolds before it is released from the SH3-SR interface.

For plectin, in fact, there is some experimental evidence that shows that the SH3 domain is occluded under normal circumstances but binds when free: a recent study, using co-immunoprecipitation, showed that when a construct similar to our considered system (exons 16–24 roughly form SR4-6) interacts with the protein MAP1, it has less affinity to it than when the SH3 domain alone is interacting. For desmoplakin, on the other hand, no SH3 binding partners have been identified, but our data speaks, by similarity to plectin, for a binding-ready SH3 domain of desmoplakin in tensed desmosomes. We speculate that desmoplakin's SH3 domain by itself, or with adjacent but unfolded spectrin repeat fragments, might be the protein unit interacting favorably with substrates, and could be the construct of choice when searching for binding partners.

The two proteins investigated in this work (desmoplakin and plectin) exhibited similar unfolding scenarios across two orders of magnitude of pulling velocities. Given that the sequence identity between the two proteins of 42% is comparable to the sequence identities within the family (23–64%, see SI), we speculate that all plakin family members as well as the related α-spectrin follow a highly similar picture of SH3 mechano-sensing. However, there are also some significant differences between the two proteins. In particular, in the case of plectin, the SH3 domain can lose contact with the rest of the plakin domain with relatively reduced resistance due to a different topology of pulling characterized by a net tilting of the entire protein structure. We attribute this change in behavior to a relatively small pocket of nonconservative sequence variations at the beginning of the SCα-helix.

Extrapolating our conclusions to the physiological case is quite difficult, given how dense desmosomes or hemidesmosomes are. In particular, the length scale of the full unfolding in our simulations (about 100–150 nm) seems very large, yet should not be per se excluded to occur. Also, the interactions with neighboring structures could tether the protein in such a way as to concentrate the stress on this particular interface. Similarly, the levels of force acting on single desmoplakin or plectin proteins in the cell remain to be quantified such as with
in-cell force sensors.

It is interesting to note that force-induced SH3 activation in plakin domains as found here requires force levels similar to those needed for talin activation, a reminiscent force-sensing mechanism of focal adhesion sites. Furthermore, a wide range of proteins (including the structurally similar alpha-spectrin) are known to mechanically unfold when cells are stretched.

There is strong evidence that the plakin domain and in particular the SR-SH3 interface play a key role in the stability or function of the proteins, therefore under stress in physiological conditions. For desmoplakin, 16 out of the known 25 disease mutations are in the central plakin domain (9 of which are in the simulated fragment SR4-5 - see Fig. S11 for their location). Despite these difficulties, the order of unfolding observed here is valuable knowledge as these insertions are observed in other proteins as well.

Direct validation with force spectroscopy experiments, preferably including ones with very high loading rates, would further help understanding the force response of these proteins. These and other future efforts will shed further light onto the role of the SH3 domain insertion in tissue integrity and stress response.

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**Author Contributions**

C.D. and K.K. performed simulations. C.D. and F.G. wrote the manuscript.

**Additional Information**

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