

N-terminal strands of filamin Ig domains act as a conformational switch under biological forces

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ABSTRACT

Conformational changes of filamin A under stress have been postulated to play crucial roles in signaling pathways of cell responses. Direct observation of conformational changes under stress is beyond the resolution of current experimental techniques. On the other hand, computational studies are mainly limited to either traditional molecular dynamics simulations of short durations and high forces or simulations of simplified models. Here we perform all-atom discrete molecular dynamics (DMD) simulations to study thermally and force-induced unfolding of filamin A. The high conformational sampling efficiency of DMD allows us to observe force-induced unfolding of filamin A Ig domains under physiological forces. The computationally identified critical unfolding forces agree well with experimental measurements. Despite a large heterogeneity in the population of force-induced intermediate states, we find a common initial unfolding intermediate in all the Ig domains of filamin, where the N-terminal strand unfolds. We also study the thermal unfolding of several filamin Ig-like domains. We find that thermally induced unfolding features an early-stage intermediate state similar to the one observed in force-induced unfolding and characterized by the N-terminal strand being unfurled. We propose that the N-terminal strand may act as a conformational switch that unfolds under physiological forces leading to exposure of cryptic binding sites, removal of native binding sites, and modulating the quaternary structure of domains.

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Key words: discrete molecular dynamics (DMD); filamin; Ig domain; thermal unfolding; forced unfolding.

INTRODUCTION

Filamin family members (A, B, and C) are composed of an actin-binding domain and subsequent 24 immunoglobulin(Ig)-like domains. Filamins have been shown to play an important role in maintaining the rheological properties of the actin cytoskeleton.¹ Filamin molecules can crosslink actin fibrils into actin-networks. As mechanical proteins, filamins localize to the cortex, stress fibers, and muscle Z-line. Besides their critical roles as mechanical proteins, filamins are also involved in cellular response to stress. Filamin A has been postulated to respond to cues from the stress environment of cells, as reported from its involvement in cell-fate determination,² activation of platelets,^{3–5} and mechanoprotection.^{6,7} How could filamin A transduce the stress state of cells? Some studies report that filamin A turnover and ability to bind actin can be regulated through its interactions with calmodulin and calpain, proteins that are activated by changes in cellular calcium level.^{2,8,9} Calcium levels in turn fluctuate with cellular stress levels.¹⁰ Other studies suggest that filamin A's role as a scaffold protein, binding over 70 proteins,¹¹ is modulated by its quaternary structure, which in turn could be regulated by either physical or thermal stress.^{12,13} Based on these studies, we hypothesize that stress-induced conformational changes of filamin A play a direct role in signaling either by disrupting existing interactions or by introducing new interactions. Therefore, it is important to investigate possible stress-induced conformational changes that could play a functional role.

Atomic force microscope (AFM) has been commonly used to study a protein's response to mechanical stress at the single molecule level. Several AFM studies on other proteins with Ig-like fold indicate that Ig domains can have force-induced metastable conformations. *D. discoideum* filamin (ddFLN), a six Ig-like domain homolog of mamma-

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lian filamin, has been reported to have a metastable conformation in domain 4 (ddFLN4), which lacks β strands A and/or B.^{14–16} The 27th Ig-like domain of titin (TNI27) has been reported to have a force-induced^{17–19} stable intermediate that lacks β strand A.²⁰ Fibronectin has been reported to have a forced-induced²¹ conformation in its fibronectin III domain 10 (FN-III₁₀) that lacks β strand A and B.²² Although AFM by itself has been used to detect the presence of intermediate states in these domains, complementary studies have been used to determine the atomistic conformation of the intermediate states. Mutational studies, such as amino acid inserts and Φ -value analysis, have been used to obtain structural insights into conformational changes in proteins under stress. These studies have been successful at proposing hypotheses about the conformation of intermediates of ddFLN4¹⁴ as well as identifying critical residues for the mechanical stability of N-terminal β strands of TNI27.²³ However, a limitation of these studies is that mutations can alter the conformational changes under stress. Thus, though existing biochemical and biophysical techniques are effective at generating hypotheses about stress-induced metastable states, they lack detailed direct structural information about these conformations.

Being able to provide a direct atomistic picture of conformational changes under forces, computational modeling has been used in the study of protein conformational changes under mechanical stress. However, the time-scales accessible by traditional molecular dynamic simulations are severely limited (~ 10 – 100 ns) and are not sufficient to study large-scale conformational dynamics of proteins (sub-microsecond to millisecond time-scales). Two major strategies have been used to overcome this limitation. The first approach is to use simplified proteins models, in which residues are represented as one or several pseudo-atoms having empirical structural-based interactions.^{24,25} For example, West *et al.* used a simplified model to study the thermal and forced unfolding of TNI27, and found distinct unfolding pathways.²⁶ The second approach often relies on simulating the system under extreme conditions, such as high temperatures or high forces, to promote rapid protein unfolding events.²⁷ For example, Fowler *et al.*, used traditional MD and a pulling protocol similar to AFM with pulling forces approximately two times the experimentally observed critical unfolding force (≥ 300 pN) for TNI27.¹⁹ Steered molecular dynamics was used by Lu *et al.* to understand the unfolding pathways of TNI27 with very high forces (~ 1000 pN).²⁸ Many other studies have been recently reviewed by Sotomayor *et al.*²⁹ One major limitation of these approaches is that unfolding pathways could be altered under extreme conditions.

To avoid the artifacts arising out of using unphysical simulation parameters and coarse-grained models in the study of force-induced filamin A conformational changes, we use all-atom discrete molecular dynamics (DMD)

simulations.^{30–35} DMD is similar to traditional MD in that it calculates the evolution of the coordinates of the system under study as a function of time and follows the same laws of conservation of energy and momentum as MD, but it affords much more rapid and diverse sampling. DMD solves the ballistic equations of motion rather than Newtonian ones, which reduces the simulation algorithm to an iterative search of the immediate collision events in the system, making it several orders of magnitude faster when compared with MD. The reduction in simulation time achieved by DMD gives us the ability to study conformational dynamics over physiologically relevant time-scales. DMD has been shown to have a higher sampling efficiency than traditional molecular dynamics and has been used to study protein folding thermodynamics and protein aggregation.³⁶ All-atom DMD features a transferable force field and has been successfully used to fold several small proteins *ab initio* (< 60 amino acids)³⁵ and to investigate the conformational dynamics of superoxide dismutase.³⁷

To study force-induced unfolding of filamin A using DMD, we need an atomistic structure of filamin A. However, structures of only five of the 24 Ig-like domains of filamin A have been experimentally determined.^{12,38–40} Hence, in this study, we use the all-atom structure of filamin A derived from comparative modeling in an earlier study (Kesner, *et al.* submitted). The average sequence similarity of all the Ig-like domains is $> 40\%$. Therefore, we expect the homology-derived structures to be close to their native structures.⁴¹ In that study, we observed that there are two classes of Ig-like domains in filamin: six-stranded (lacking the N-terminal strand) and canonical seven-stranded. Domains 16, 18, and 20 are six stranded. In a recent crystal structure of domains 19–21, it was observed that the putative first strand of domain 20 interacts with domain 21 instead of forming a β sheet with the second strand. This interaction has been proposed to inhibit domain 21's interaction with other binding partners.¹² Based on this quaternary structure, it is possible that β strand A of an Ig-like domain could form complementary strands with adjacent Ig-like domains. Hence, we hypothesize that unraveling of the N-terminal strand of Ig-like domains could initiate interdomain interactions through that strand. We study the six-stranded Ig-like domains from two perspectives. First, we compare domains 16 and 18 to that of domain 20 (whose structure has been determined)¹² to study if the six-stranded domains have similar physical characteristics. Second, we compare these domains to the seven-stranded domains to understand the role of the missing strand.

For each of the 24 Ig-like domains that make up filamin, we perform constant-force simulations by applying a set of physiologically relevant ranges of forces (0–315 pN).⁴² Based on these systematic DMD simulations, we estimate the critical force of unfolding of each individual

domain and characterize the conformational changes under mechanical stress. We find that the critical forces are heterogeneous among the 24 domains due to the sequence differences. We also find that there is no significant difference between the critical unfolding forces of domains with and without N-terminal strands. We find that under small forces up to ~ 35 pN, which correspond to the physiological forces,⁴³ most Ig-like domains remain in the native states. As the force increases up to the intermediate force levels of ~ 70 pN, large conformational changes take place. Interestingly, Ig-like domains feature a common initial conformational change, where the first β strand unfurls. Domains that lack their N-terminal β strands appear similar to one another, maintaining their native-like conformation under low forces. At intermediate forces, they unfold to a heterogeneous population of intermediate states similar to seven-stranded domains.

Additionally, the effect of temperature versus mechanical stress on unfolding pathways has been the subject of much debate.⁴⁴ The general hypothesis is that thermally induced and force-induced unfolding pathways are independent from one another because the thermal fluctuations are exerted on the protein globally, whereas the effect of mechanical forces is localized and nonhomogeneous. To examine whether the thermally and force-induced unfolding of filamin Ig-like domains are different from each other, we also perform thermal unfolding of three Ig-like domains, 14, 21, and 24. Interestingly, we find that thermally induced unfolding shares the same initial unfolding intermediate state as forced unfolding, where the first strand unfolds. This observation suggests that filamin evolved to feature the intermediate state of unfolded N-terminal strand(s). Given the observation that the putative first strand of domain 20 interacts with domain 21 instead of forming a β sheet with the second strand, we propose that the N-terminal strand may act as a conformational switch that unfolds under stressed physiological conditions leading to exposure of cryptic binding sites, removing of native binding sites, and modulating the quaternary structure of domains.

MATERIALS AND METHODS

All-atom DMD

A detailed description of the DMD algorithm can be found elsewhere.^{35,45} Briefly, interatomic interactions in DMD are governed by square-well potential functions. Neighboring interactions along the sequence (such as bonds, bond angles, and dihedrals) are modeled by infinitely deep square-well potentials to model the geometry of residues and peptide bonds. During a simulation, an atom's velocity remains constant until a potential step is encountered, during which it changes instantaneously according to the laws of conservation of energy, momen-

tum, and angular momentum. Simulations proceed as a series of such collisions, with a rapid sorting algorithm used at each step to determine the next collision.

The all-atom DMD method uses a united atom protein model, where heavy atoms and polar hydrogen atoms are explicitly modeled.³⁵ We include van der Waals, solvation, and environment-dependent hydrogen bond interactions. For solvation, we adopt the Lazaridis-Karplus solvation model^{46,47} and use the fully-solvated conformation as the reference state. Because of the strong screening effect of solvent, distant charges have weak polar interactions. For salt-bridges, we expect the hydrogen bonds to partially account for their polar interactions.

Protein models

As described in the introduction, filamin A is composed of an ABD followed by 24 Ig-like domains. Ig-like domains are seven-stranded β sandwich folds [Fig. 1(A)]. Loops with conserved residues GPG and SPF bisect the first and last strands. We refer the resulting subdomains as A, A', G', G. Subdomain A forms a β sheet with B (A-B region) and in some domains A' and G (A-G region) form a parallel β sheet. We use the homology models of human filamin A Ig-like domains [Fig. 1(A)] that were derived from an earlier work (Kesner *et al.*, submitted). Briefly, closest matching homology templates were derived from a structural alignment of homologous filamin domains in the protein data bank (PDB).⁴⁸ *Insight II* (Accelrys, CA) was then used to perform homology modeling of each filamin A Ig-like domain. Domain 16, 18, and 20 have only six complete strands since the best-match template lacks all (16), or substrand A (18, 20), of β strand A.

Each model is validated using the sequence-structure compatibility score from the *Verify-3d* module of *InsightII* (Accelrys Inc.). This analysis measures the compatibility of modeled residues with their structural environment.⁴⁹ Quality scores of greater than 0.1 indicate valid structural models⁴⁹ and typical biochemical structures have a score of 1. The mean *Verify-3d* score obtained for Ig-like domain models was 0.89 with standard deviation 0.30. IgFLNa18 is the only model to produce a low score (0.11). However, IgFLNa18's homology model has high sequence identity (64% id), and a low RMSD (0.23 Å) when compared with its IgFLNb18 (PDB ID 2dmc) template, suggesting that the IgFLNa18 model was correctly built based on the experimental template. Furthermore, the experimental structure of the template itself has a low quality score of 0.56. Overall, Ig-like domains have high quality scores indicative of their high accuracy.

Following homology modeling, each structure is relaxed using 1000 steps of steepest descent energy minimization as in AMBER⁵⁰ 9.0. In addition, each model is minimized using the Medusa modeling suite.^{51–53}

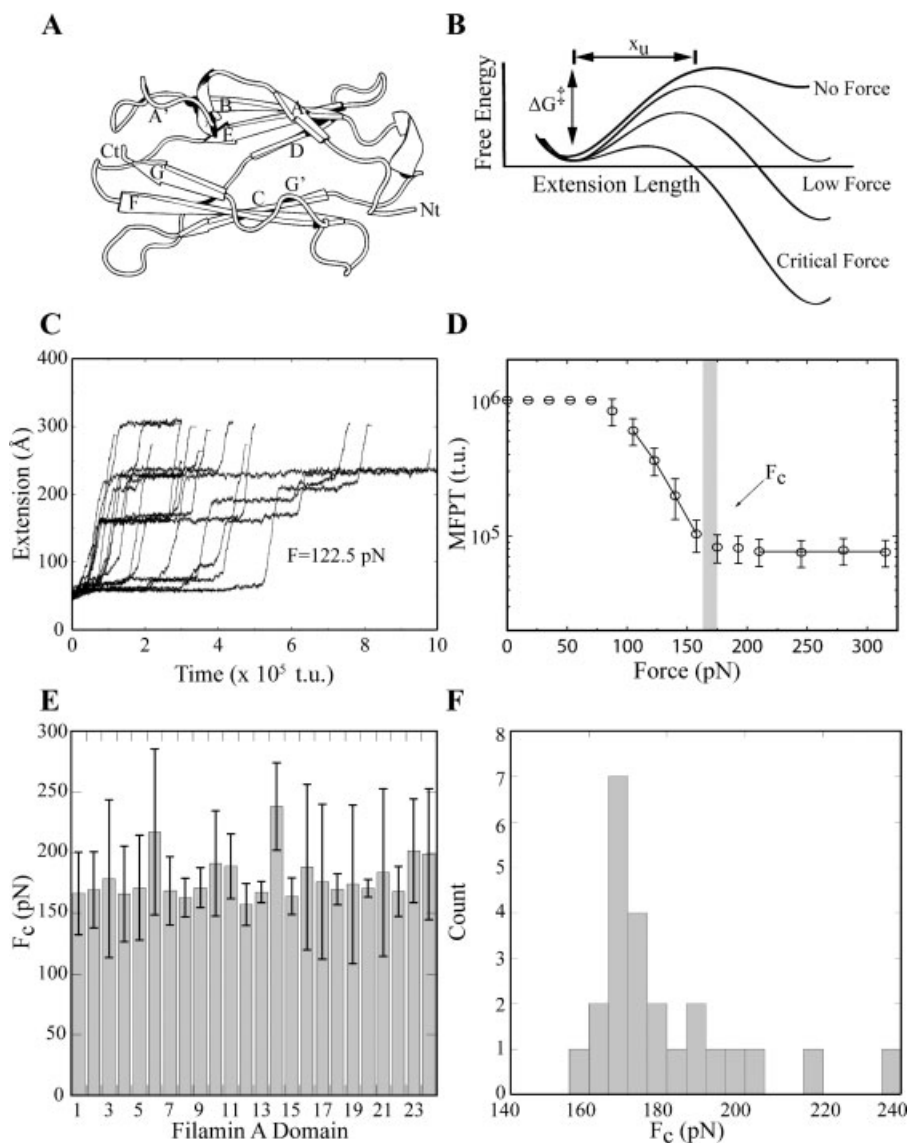


Figure 1

Critical unfolding parameters of filamin Ig-like domains. (A) Seven-stranded Ig-like domain with subdomain termini and β strands A-G labeled. (B) Graph depicting the free energy landscape along the extension length as the reaction coordinate at different levels of constant-force pulling. F_c is the critical unfolding force and x_u is the distance to the center of the unfolding energy barrier. (C) Unfolding behavior of a domain pulled apart with a constant force (122.5 pN) is described by plotting the extension length as a function of time. Replicates with differing initial velocities vary in their unfolding behavior. (D) A linear natural log plot of the pulling force versus mean first passage time. F_c is the critical unfolding force. A linear fit of forces is shown for the barrier (left of F_c) and barrierless (right of F_c) regimes. (E) Critical unfolding forces observed for each domain. The error bars are determined as described in Methods. (F) Histogram of mean F_c .

Constant-force unfolding

For each domain, we start the constant-force simulations from the homology-derived native state. The dimension of a cubic simulation box is 500 Å and the periodic boundary condition is used. We use an Anderson thermostat⁵⁴ to maintain constant temperature at the room temperature (300 K). Constant-force pulling is achieved by applying a discretized step-function with a constant energy jump, dE , at the distance step of

$dR(1 \text{ \AA})$ between the N- and C-termini. The pulling force equals to dE/dR . For example, dE of 0.5 kcal/mol is equivalent to ~ 35 pN. The applied forces range from zero to 315 pN. From zero to 210 pN, we sample the force values with an interval of 17.5 pN. From 245 to 315 pN, the interval is 35 pN. For each force, we perform 10–20 realizations of pulling simulations, each of which start with randomized velocities. We perform 10 simulations for pulling simulations at low forces (70 pN or less), because there is a low probability of unfolding dur-

ing the simulation time. For higher forces, we perform 20 realizations.

Derivation of critical unfolding forces

Under external force F , the unfolding time under pulling forces can be approximated as $\tau = Ae^{(\Delta G^\ddagger - Fx_u) \times \beta}$. Here, A is the Arrhenius prefactor, ΔG^\ddagger is the zero force activation energy barrier, x_u is the distance to the center of the activation energy barrier, and β is the inverse of kbT , where k_b is the Boltzmann constant, and T is the temperature. We use the mean first passage time (MFPT) of unfolding as the approximation of the unfolding time. We use 200 Å as the cutoff distance to assess whether a protein is unfolded. This cutoff was determined based upon observations that after an extension of 200 Å most domains lack metastable states, although the minimum full extent for an unfolded domain is 240 Å (Supporting Information Fig. 1). If the protein has not unfolded within the maximum simulation time (one million time units), we use this time as the first passage time.

We plot the $\ln(\text{MFPT})$ versus force and obtain linear fits in the barrier and barrierless unfolding regions. The intersection of the two linear fits determines the critical force of unfolding (F_c). We use the error bar of the linear regression to estimate the error bar of the estimated critical forces with a simple Monte Carlo method. In each Monte Carlo step, linear lines constructed based on the Gaussian distribution of error bars (derived from the linear regression) is used to determine a new intersection. The mean and standard deviation of the critical unfolding force is derived from the set of intersections.

Forced unfolding pathways analysis

For each domain and each force, we combine trajectories of all realizations and compute the histogram of extension lengths. Gaussian curve fitting of the histogram is used to identify metastable states. Only the states that are present for more than 1% of the total simulation time are considered stable. To determine the representative structure of each metastable state, we first extract all conformations having extension length within the mean ± 2 SD of the corresponding Gaussian fit. Then, we compute a contact frequency map for each extracted ensemble. Finally, we select the representative structure as the one that has the least contact distance from the average contact frequency of the ensemble.

We characterize the unfolding pathway using a directed graph. The nodes of the graph are formed by the states identified from the extension length histogram and the edges are the transitions between the different states. Occupancy of each node corresponds to the number of conformations associated with that state normalized by the total number of conformations. Edges have an associated transition probability determined by counting the num-

ber of transitions between a node and its neighbor and then dividing by total number of transitions to all its neighbors in the same direction. The graph representation of the unfolding pathway is visualized using Cytoscape.⁵⁵

Thermal unfolding

Equilibrium simulations are performed at a wide range of temperatures: 300, 330, 360, 390, and 420 K. We use the percentage of native contacts, Q , to approximate the reaction coordinate. We define a contact when the distance between two C β atoms (C α for Glycine) is less than 8.5 Å. We use a similar approach to identify the representative structure of each state, where the Q -value rather than the extension distance is used as the approximation of the reaction coordinate. To determine the localization of loss of contacts during thermal unfolding simulations, we calculate a subdomain average Q -value. Subdomains are delineated based upon a strand-by-strand analysis of each Ig-like domain homology model. Average Q -values for an ensemble are obtained by dividing the marginal sum of native contacts from the residue-by-residue contact frequency map by the marginal native contacts from the reference structure contact map.

RESULTS

We model constant-force unfolding by applying a pulling force between the C α atoms of the N- and C-terminal residues (Materials and Methods). The force is modeled by assigning a corresponding potential between the corresponding atom pair. For each force value, we perform 10–20 independent simulations with randomized initial velocities. Each simulation lasts a million DMD time units.

Computationally identified critical unfolding forces of filamin Ig-like domains agree with AFM experiments

Most single domain proteins, such as the Ig-like domains in filamin, feature a two-state folding/unfolding dynamics without external forces. At room temperatures, a protein stays folded and its native state corresponds to the lowest energy state in the free energy landscape [Fig. 1(B)]. Unfolding requires overcoming a free energy barrier (the folding/unfolding transition state) separating the folded and unfolded states, ΔG^\ddagger . The unfolding time is proportional to the exponential of $\Delta G^\ddagger/k_B T$. As the force (F) applied to the N- and C-termini of a protein increases, the unfolded states with large end-to-end extensions are favored [Fig. 1(C)], and the corresponding free energy barrier decreases ($\Delta G^\ddagger - Fx_u$ [Fig. 1(B)]). Here, x_u denotes the difference of end-to-end distances between the transition and native states. Assuming that

x_u does not change for different external forces, the natural logarithm of the unfolding time is linearly dependent on the external force. As the force reaches the critical value (F_c), the barrier will disappear and activation driven unfolding dynamics is reduced to barrierless unfolding. As a result, the dependence of the unfolding time with respect to the external force changes near the F_c [Fig. 1(D)]. We estimate the critical unfolding forces based on the change of the scaling of the unfolding time with respect to external forces (see Methods). For each applied force, we compute the mean first passage time (MFPT) over 20 trajectories to estimate the unfolding time. We use a cutoff distance of 200 Å to assess whether a protein is unfolded. We estimate the critical force as the cross-over point in the plot of $\ln(\text{MFPT})$ as a function of external forces [Fig. 1(D)]. We compute the critical unfolding forces for each domain [Fig. 1(E)]. The mean critical unfolding force over all domains is 179 ± 19 pN. We find that the distribution of F_c values features a peak around 155–180 pN with a long tail from 185–205 pN [Fig. 1(F)]. Therefore, most of the Ig-like domains of filamin A can withstand the stretching forces up to 150 pN with a few domains being able to resist higher forces. For example, domains 6 (217 ± 68 pN) and 14 (238 ± 36 pN) have F_c values greater than 205 pN. Our calculated critical forces are consistent with the experimental measurements (50–220 pN as identified by AFM pulling study of Filamin A).⁴²

Sequence alignment during the homology modeling of domains 16, 18, and 20 suggested that these domains lack the N-terminal β strand of the canonical Ig fold. Such a deviation from the canonical Ig-like domain suggests a unique biological function for these domains. Interestingly, the absence of these structural features did not significantly change the mechanical stability of these domains with respect to other domains. The two domains, 18 and 20, have F_c of 170 ± 18 pN, and domain 16 has an F_c of 188 ± 68 pN. These data suggest that the mechanical stability of Ig domains is independent of the N-terminal β strand and likely to be caused by sequence difference in the core of the protein.

Recent *in vitro* pulling experiments on filamin-actin networks suggest that filamin A Ig-like domains are subjected to equilibrium forces up to 80 pN.⁴³ Hence, we analyze the conformational changes of each of the 24 Ig-like domains under 35 and 70 pN of constant pulling force next (see Forced Unfolding Pathways Analysis in Materials and Methods).

Under the 35 pN pulling force, most filamin Ig-like domains remain in a native-like state with a few domains featuring intermediate conformations that could potentially play a role in stress signaling

We analyze the population of conformations of each Ig-like domain under the 35 pN pulling force, which we expect to be physiologically relevant. We find that most

of the domains stay in their near-native conformations as shown for domain 21 [Fig. 2(B)]. The average end-to-end extension of these conformations is increased by ~ 10 Å, when compared with the extension length seen in zero force extension [Supporting Information Fig. 2(A,B)]. This is the result of pulling out leading N-terminal residues [Supporting Information Fig. 3(A,B)], which make little contacts in the native state. The rest of the structure remains intact [Supporting Information Fig. 3(A)].

Interestingly, we find that at the 35 pN pulling force a few domains feature a partially unfolded intermediate state in addition to a near-native conformation. This is exemplified by partially unfolded state of domain 13 [Fig. 2(A)], which has the interaction between strands A and B lost while maintaining A'-G region contacts [Fig. 2(A); the 55 Å conformation]. This conformation is analogous to changes hypothesized for TNI27 when it was pulled at much higher forces.^{19,20} The loss of β -strand A contacts while other β strands are left intact is common to most of the intermediate states of these domains, although domain 3 is unique in having a large loss of contacts in the G' region [Supporting Information Fig. 3(B)]. We do observe one exception: domain 15 loses its entire β strand A (Supporting Information Fig. 4). This major conformational shift occurs in a small number of ensembles and produces an extension length of ~ 86 Å [Supporting Information Fig. 2(B)]. Thus, while several domains have a modest conformational shift pulled at the 35 pN force, at least one domain has a major conformational shift that could serve as a potential signal of relatively low levels of cellular stress.

At 70 pN, filamin domains have a diverse set of metastable states. The unfolding pathway of all domains is initiated by the loss of contacts in the N-terminal β strands

Under 70 pN pulling force, all Ig-like domains undergo large conformational changes [Fig. 3 and Supporting Information Fig. 2(C)]. The most populated states are those with end-to-end extension within 20 Å of native state, where β strands A and/or G has been pulled out slightly leaving the remaining structure native-like (Fig. 3). The stretched conformations with longer extensions (>20 Å) have one β strand sliding past another [Fig. 3(A)] and/or lose contacts between strand A' and G [63 Å extension in Fig. 3(B); and 72 Å extension in Fig. 3(C)]. Interestingly, TNI27 is reported to have similar initial unfolding pathways which features loss A-B contacts, loss of contacts in the A'-G region, and then quickly unfolding.^{19,20} However, at 70 pN stretching force filamin Ig-like domains do not traverse directly to the unfolded state, instead they get trapped in several metastable states [Supporting Information Fig. 2(C)]. These states include domains in which strand A [130 Å

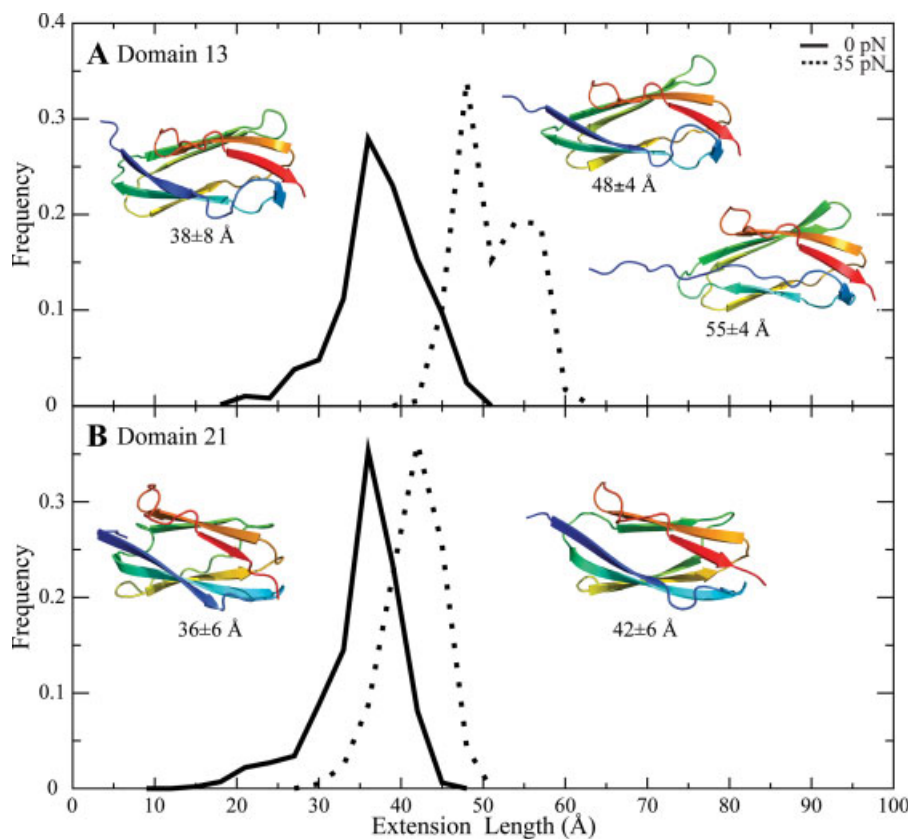


Figure 2

Equilibrium stable intermediate states of domains at 35 pN of constant-force pulling. Histogram of extension lengths for domains 13 (A) and 21 (B). A representative conformation for each state is shown. The solid line represents the distribution from simulations at 0 pN pulling and the dotted line represents the distribution from simulations at 35 pN pulling.

extension in Fig. 3(B)] or strands A and B have been unfolded [150 Å extension in Fig. 3(A), and 149 Å extension in 3(B)] from the rest of the domain resulting in conformations such as a single β sheet [150 Å in Fig. 3(A)] or a β sandwich [149 Å in Fig. 3(B)]. These conformations are analogous to those observed in the metastable intermediate of ddFLN4 when unfolded using forces in the order of 60 pN.^{14,15} Also, FN-III₁₀ has been observed to lose N-terminal β strands when forced to unfold using \sim 100 pN.^{21,22} Domain 21 has a proclivity to also lose β strand G entirely [Fig. 3(B) 149 Å], which could potentially be related to its role in the previously mentioned interdomain conformation involving domain 20. Most other domains only lose β -strands at the N-terminus.

The most stable domains at 70 pN unfolding force are 3, 9, and 22, which have metastable states close to that of their native state (extension length < 86 Å) [Supporting Information Fig. 3(C)]. The frequent interconversion between intermediate states of these domains suggests a low energy barrier between successive extension length states [e.g., Fig. 3(C) domain 3]. Domains 1, 6, 12, 13,

and 14 also feature the near-native state being highly populated. However, these domains also feature an intermediate state missing the N-terminal β strands. Interestingly, we found other domains (e.g., 21) [Supporting Information Fig. 3(C)] are fully unfolded using a force \sim 70 pN, suggesting a lesser stability of these domains. Domains in the range 9–15 are considered secondary sites for binding actin, and having unusually stable domains in this range could be critical to the structural role of filamin in cells.⁵⁶ We hypothesize that domains near the actin-binding boundary could have added stresses in the cell because they could act as the actin-binding fulcrum. It is also conceivable that domains near the end of the actin-binding range could require additional flexibility. Thus domains 13 and 14 having stable intermediates lacking β strands could meet this potential requirement.

We also find that the mechanical stability of some Ig-like domains does not depend entirely on N-terminal β strands. We modeled domains 16 (N-terminal residues up to and including β strand A), 18 and 20 (N-terminal residues up to and including β A) lacking N-terminal β

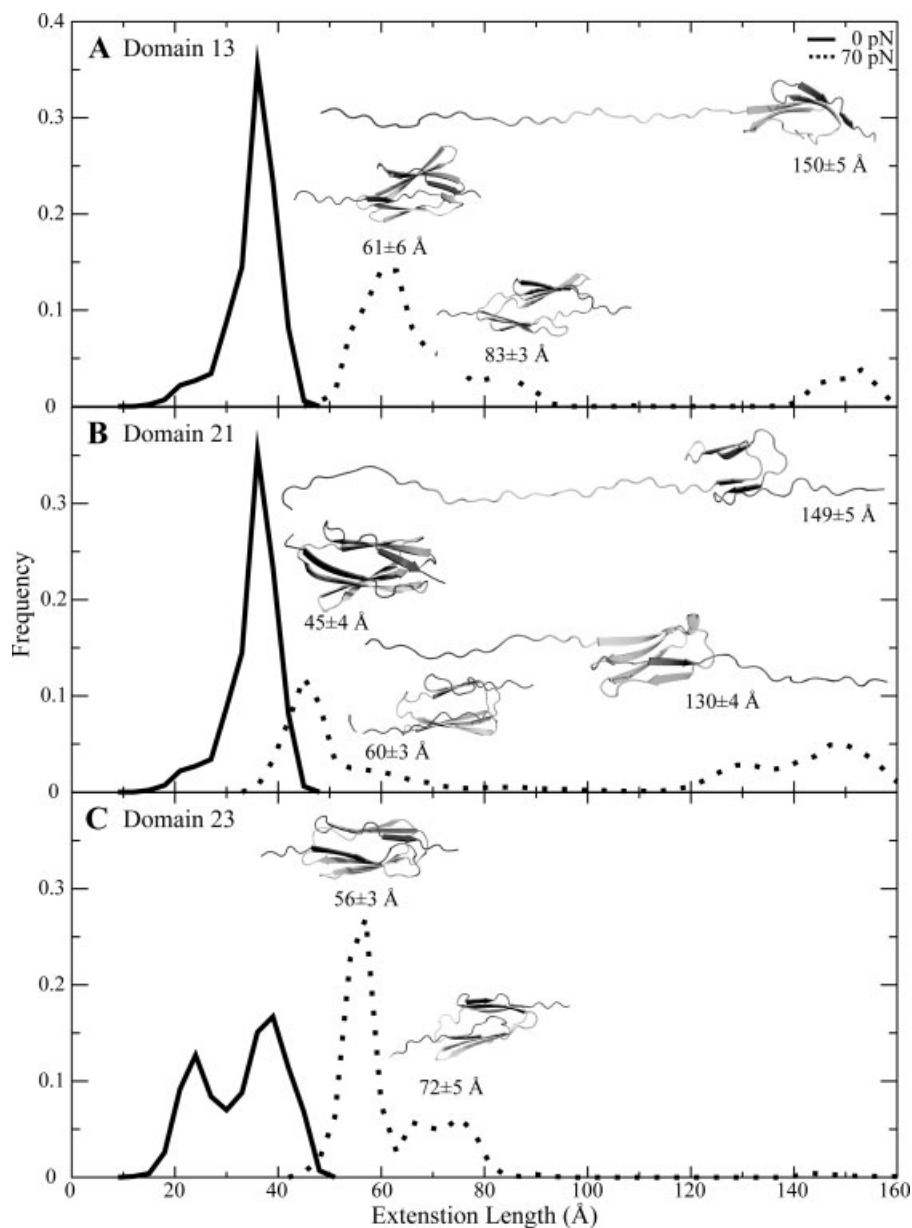


Figure 3

Equilibrium stable intermediate states of domains at 70 pN of constant-force pulling. Histogram of extension lengths for domains 13 (A), 21 (B) and 23 (C). A representative conformation for each state is shown. The solid line represents the distribution from simulations at 0 pN pulling and the dotted line represents the distribution from simulations at 70 pN pulling.

strands (Kesner, et al., submitted) as their templates lacked β strand integrity in this region. We find that at both 35 and 70 pN, these domains have similar unfolding pathways and metastable conformations as seven-stranded domains. Domain 20 maintains a highly populated native-like conformation featuring an A'-G region β sheet at 35 pN [Supporting Information Figs. 3(A) and 4(B)] and an unfolding pathway where the most populated state is native-like at 70 pN similar to seven-stranded domains. Domain 18 at 35 pN features two

metastable states and does not unfold entirely. The most populated state is native-like with few of the native contacts lost [Supporting Information Fig. 3(A)]. However, domain 18 also features a lowly populated state with a conformation that has an unfurled N-terminal β strand B at 35 pN [Supporting Information Fig. 4(B)], thus making it somewhat distinct from seven-stranded domains in which we observe the unfolding of β strand B only at 70 pN. At 70 pN, domain 18 has a similar unfolding pathway as that of domain 24, featuring highly

populated intermediate states that have extension lengths much greater than that of their native states [Supporting Information Fig. 2(C)]. Domain 16 features the greatest differences in the unfolding pathway when compared with seven-stranded domains: it has an unfolding pathway that is distinct from that of domains 18 and 20 and most seven-stranded Ig-like domains unfolding from the C-terminus at 35 pN [Supporting Information Fig. 4(B)]. However, domain 16 is very stable because at 70 pN, its most populated state is a metastable and not the fully unfolded state. Thus, all three domains lacking N-terminal β strands are highly stable at 35 and 70 pN of force, although domain 18 and 20 are more similar to each other than to domain 16, which unfolds from the C-terminus.

This systematic analysis of our *in silico* pulling studies of filamin Ig-like domains reveals metastable intermediate states as observed in AFM experiments. Our simulations are able to reproduce experimentally derived critical unfolding forces. The conformational changes of all the Ig-like domains feature a common intermediate state where the N-terminal strand is unfolded, which could play a role in the signaling pathway of cell responses to mechanical stress. Next, we study the thermally induced unfolding of the filamin domains to test whether the intermediate states observed in forced pulling are also observed in thermally induced unfolding pathways. We restrict our study to three Ig-like domains, 14, 21, and 24. These domains are selected due to distinct kinetic properties and biological significance: domain 14 has the highest critical force, domain 21 has well-characterized thermodynamics,¹³ and domain 24 is involved in dimerization.³⁸

Thermal unfolding of filamin Ig-like domains features a common initial unfolding of the N-terminal β strand

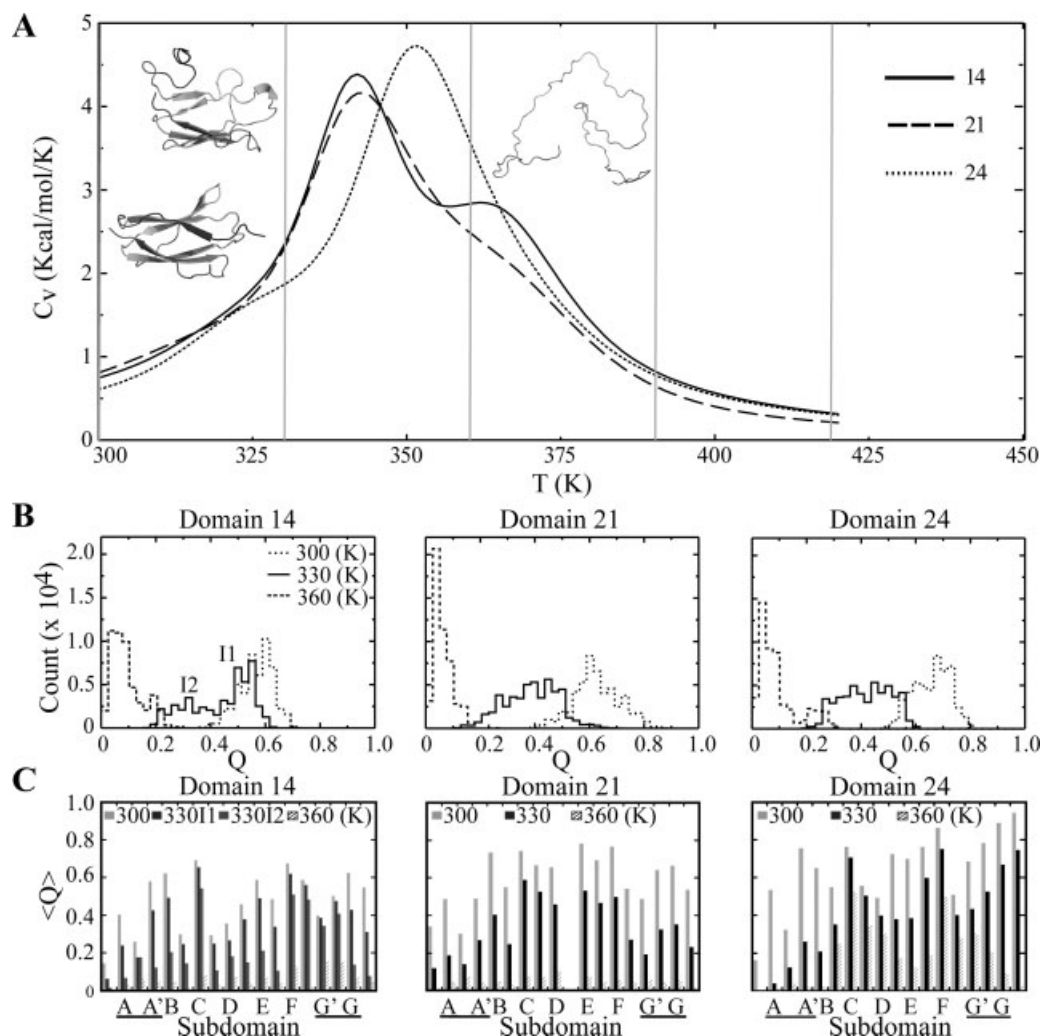
We perform equilibrium DMD simulations of Ig-like domains 14, 21, and 24 over a wide range of temperatures (300–420 K). We use the weighted histogram analysis method (WHAM)³⁵ to compute the specific heat [Fig. 4(A)]. The highest peak in the specific heat corresponds to the unfolding transition. The shoulders in the specific heat plot suggest the existence of an intermediate state. Our results indicate that domain 14 and 21 have similar unfolding temperature of ~ 340 K. Domain 24 is the most stable of these domains with unfolding transition temperature $\sim 10^\circ$ higher than either 14 or 21 [Fig. 4(A)]. The increased stability of domain 24 is possibly the result of its well-characterized role in dimerization of filamin monomers.³⁸ Domain 14 has a prominent shoulder at temperature ~ 370 K that is absent in domain 21, suggesting the existence of a thermally stable intermediate state.

To further investigate the conformational states of these domains, we use the fraction of native contacts (Q) as the approximation of the reaction coordination to analyze the conformational state at different temperatures [Fig. 4(B)]. We also compute the Q -value of each of segments in the native state [Figs. 1(A) and 4(C)]. At the low temperature of 300 K, all domains have near-native conformation with high Q -values [Fig. 4(B)]. As the temperature increases to 330 K, we find domain 14 features two states, I1 and I2, while domains 21 and 24 have only one. We examine the Q -values of each secondary structure segment at 330 K [Fig. 4(C)]. For domain 14, both states I1 and I2 have large losses in A–B region contacts, I2, however, has more contacts lost in A', B, and G than does I1. Furthermore, the conformation in I1 is a closed β sandwich, whereas the conformation of I2 is an open β sandwich [Supporting Information Fig. 5(A)]. Thus it is likely that A'–G region contacts are involved in maintaining the closed conformation of I1. Domain 21 has lost significant amount of native contacts in both the N- and C-terminal β strands. Domain 24 loses contacts of mainly in β strand A. The existence of two states of domain 14 is consistent with the shoulder in the specific heat. At high temperature of 360 K, the most populated state of each domain is the unfolded conformation. Consistent with the high thermal stability, the β strands CDFG of domain 24 still retain partial native interaction at 360 K [Fig. 4(C)].

Jiang *et al.*, using an NMR based approach determined the unfolding temperature of domain 21 to be 319 K. They also reported that different filamin Ig-like domains have different thermal stabilities. Domain 19 and 17 are fully folded at 333 K while domain 21 is unfolded.¹³ They have also reported a partially unfolded fraction of domain 21 under mild conditions (298 K). It is possible that the differences between domains observed were the result of β strand A's displacement that we observed in domain 21 in our simulations at 330 K [Supporting Information Fig. 5(B)] because the residue they used as a folding marker is adjacent to β strand A. Our results are quite distinct from those of other reported simulations of thermal unfolding of Ig-like domains. For example, Paci *et al.* simulated the unfolding of TNI27 at 450 K and found unfolding pathways in which the last strands to unfold were the terminal β strands.⁵⁷ Given the high temperatures used, it is possible that these observations were the result of nonequilibrium simulations.

DISCUSSION

Mounting experimental evidence suggests that filamin A molecules act not only as structural and mechanical proteins that cross-links actin fibrils but also as scaffold proteins that play a critical role in cellular response to external mechanical stress.^{2–7} We hypothesize that conformational changes of filamin A due to mechanical

**Figure 4**

Thermal unfolding of filamin domains. (A) Plot of the specific heat as a function of temperature for domains 14, 21, and 24 calculated using the WHAM. Representative conformations of the metastable states of domain 21 at temperatures 300, 330 and 360 are also shown. (B) Histogram of Q-values for compiled replicated trajectories at each temperature (300, 330, 360 K) for domains 14, 21 and 24. Metastable states of domain 21 at 330 K are labeled I1 and I2. (C) Average Q-value for metastable state for each subdomain (Fig. 1A) at each temperature (300, 330, 360 K).

stress are important for signaling, where stress-induced conformational changes can either disrupt existing interactions or introduce new interactions. We systematically studied the conformational dynamics of forced unfolding of filamin Ig-like domains using DMD. We developed a novel technique to derive the critical unfolding force from constant-force trajectories that is based upon finding the intersection between two unfolding regimes with respect to pulling forces. As a validation of our methodology, our computationally estimated forced unfolding dynamics of filamin A Ig-like domains agree with AFM experiments. For example, our simulation suggests the average critical unfolding force is 150–240 pN, which is consistent with the experimental measurements of 50–220 pN. Our analysis of the force-induced unfolding tra-

jectories suggests the existence of multiple intermediate states corresponding to partially unfolded β sandwich of Ig-like domains. Additionally, an intermediate state observed in the simulations common to most filamin A Ig-like domains is characterized by the unfolding of the first N-terminal β strand. This intermediate state is consistent with the intermediate state observed in the force-induced unfolding of *D. discoideum* filamin domain 4, 27 Ig-like domain of titin, and fibronectin type three domain 10.^{14,16–22} Thus, our stretching simulations using all-atom DMD is able to recapitulate the conformational dynamics of Ig-like domains under force.

Our simulation results suggest that at low force (35 pN) most of the Ig-like domains remain in the native state with only a few domains featuring a weak popula-

tion of a partially unfolded N-terminal strand. As the force increases (up to ~ 70 pN, which is close to the maximal forces observed *in vivo*),⁵⁸ we observe an increased population of intermediate states. Most of the domains still remain in their native or near-native conformations because these forces are still below the F_c . We find a large heterogeneity in both the population of the force-induced intermediate states and the transitions between them. However, we do find a common initial unfolding intermediate common to all Ig-like domains, where the N-terminal strand unfolds, similar to those observed in AFM experiments on TNI27.^{18,20}

Studies with TNI27 have suggested that contacts within A'-G region are important for maintaining the stability of its metastable state.²³ We observe a similar phenomenon in domain 20's metastable states. However, the stability of other six-stranded domains appears to be independent of this region. Domain 18 loses A'-G region contacts during unfolding, and yet does not completely unfold. Also, domain 16, which lacks subdomain A', has a similar critical unfolding force as other domains. These three domains might not be representative of all domains because in their native-like state they are stable absent N-terminal β strands. On the other hand both thermally and forced unfolding pathways of most seven-stranded Ig-like domains feature metastable states that lack any A' contacts. Thus, it is likely the A'-G region plays a minor role in filamin Ig-like domain stability.

Our results from thermal unfolding of Ig-like domains suggest that there is no correlation between thermal and mechanical stability. For example, domain 14, which has the highest critical unfolding force, is thermally less stable than domain 24. However, there does appear to be a correlation between thermal unfolding and force-induced unfolding pathways. In the case of both low force unfolding and low temperature thermal unfolding, unfolding is mostly initiated with the loss of contacts in N-terminal β strands. In the few cases where we observe initiation of forced unfolding at both the N- and C- terminus (domain 21), we also observe loss of contacts at both termini due to thermal denaturation. Thus, as evidenced by both thermal and force-induced unfolding pathways, Ig domains of filamin have a similar initial unfolding intermediate state. This similarity may be the result of evolutionary optimization of filamin due to its role as a structural and scaffold protein.

Filamins interact with a wide range of cell signaling proteins, and the conformational changes upon stress can either disrupt or promote these interactions. Additionally, the exposure of partially unfolded strands can promote modifications such as phosphorylation, which in turn can activate downstream signaling pathways. Therefore, we postulate that intermediates with their N-terminal strands unfolded could be involved in the cellular response to external mechanical stress. Further experimental studies are required to verify this hypothesis.

CONCLUSION

Stress levels in cells can be very large, especially in the cortex where the amount of stress depends on its compliance to the surrounding surface.⁵⁹ In this study, we find that under forces between 150–240 pN filamin domains readily unfold. However, under lower forces, domains unfold in stages producing a heterogeneous population of metastable states lacking N-terminal β strands. Our simulations of thermal unfolding reveal similar metastable states. Furthermore, we observe that the domains that were modeled lacking all or part of the N-terminal β strand (16, 18, and 20) have similar forced unfolding characteristics as canonical Ig-like domains. Thus, it appears that the stability of Ig-like domains is independent of N-terminal β strands. Domain 20 has been shown to have an N-terminal β strand playing an inhibitory role for protein-protein interactions involving filamin A.¹² Because domain 18 has a similar forced unfolding pathway as domain 20, it is possible that domain 18's β strand A has a similar biological function. Moreover, it is possible that unraveling of N-terminal β strands as observed during forced unfolding leads to interdomain interactions that could produce the inhibitory behavior that Lad *et al.*¹² observed. Other possible ramifications to having exposed β strands include exposure of cryptic binding sites or post translational modification sites; higher degree of connectivity between filamin monomers as more β strand edges are exposed; a higher turnover rate as more residues are exposed to potential cleavage; forming of new actin structures (cross-links vs. bundles); attraction of heat shock proteins to refold domains as stress increases; allowing Ig-like domains to more quickly refold back to their native fold when stresses and then relaxed; or allowing multiple domains to fold properly.⁶⁰ For example, proteomic screens revealed two phosphoserines (S¹⁰⁸¹ and S¹⁰⁸⁴)^{61,62} in the loop between β strands A and B of domain 9 of filamin A. Thus, we postulate that the stress-induced unfurling of the N-terminal β strands of filamin domains could play an important cellular role as a conformational switch.

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