Mechanical Perturbation of Filamin A Immunoglobulin Repeats 20-21 Reveals Potential Non-equilibrium Mechanochemical Partner Binding Function

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Accessibility
Mechanical perturbation of filamin A immunoglobulin repeats 20-21 reveals potential non-equilibrium mechanochemical partner binding function

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The actin crosslinking protein filamin A (FLNa) mediates mechanotransduction, a conversion of mechanical forces into cellular biochemical signals to regulate cell growth and survival. To provide more quantitative insight into this process, we report results using magnetic tweezers that relate mechanical force to conformational changes of FLNa immunoglobulin-like repeats (IgFLNa) 20–21, previously identified as a mechanosensing domain. We determined the force magnitudes required to unfold previously identified structural organizations of the β-strands in the two domains: IgFLNa 20 unfolds at ~15 pN and IgFLNa 21 unfolding requires significantly larger forces. Unfolded domain IgFLNa 20 can exist in two different conformational states, which lead to different refolding kinetics of the IgFLNa 20 and imply a significant impact on the reformation of the domain pair at reduced force values. We discuss the relevance of the findings to force bearing and mechanosensing functions of FLNa.

Actin filament (F-actin) networks provide mechanical stability of cells, and their remodeling and contractility mediate mechanical events such as cell shape changes and movements. While chemical signals such as ATP hydrolysis and phosphorylation mediate remodeling and contraction of these networks, a reverse phenomenon, mechanotransduction, also occurs where mechanical forces elicit chemical changes that affect cell growth and survival1–3.

Filamins (FLNs), large actin-binding protein family, mediate these biological processes4–6. Since FLNs crosslink two actin filaments and have spring-like properties, they can potentially transmit and respond to mechanical forces applied to F-actin networks4,6,7. FLNs also bind and modulate multiple cellular components, and disruption of these interactions results in impairment of binding partner’s functions8–10. These findings suggest that FLN-partner interaction is at least in part regulated by mechanical force. Consistent with this hypothesis, externally imposed stress and internally generated contractile force to F-actin networks crosslinked by FLNa, which is the most abundant isoform among the vertebrate FLNs, regulate ligand binding to FLNa in vitro11.

FLNa is composed of two identical subunits, each containing an N-terminal actin-binding domain followed by 24 immunoglobulin-like (Ig) repeats (IgFLNa 1–24)12. The 24 Ig repeats can be divided into three segments: rod 1 (IgFLNa 1–15), rod 2 (IgFLNa 16–23), and Ig repeat 24 or dimerizing domain. Two short flexible hinge sequences separate these three segments. Repeats IgFLNa 1–15 in rod segment 1 form an extended linear structure without obvious inter-domain interactions. In contrast, rod segment 2 (IgFLNa 16–23) assumes a compact structure due to multiple inter-domain interactions. Domains 16–17, 18–19, and 20–21 form stable paired structures13,14. The compact rod 2 segment mediates most protein-protein interactions involving FLNa15.

Due to the unique domain-domain interactions among IgFLNa repeats 16–23, some binding interactions are considered auto-inhibitory. For example, the CD face of IgFLNa 21 provides the binding interface for the
cytoplasmic tail of the β subunit of integrin, a key membrane adhesion molecule and migfilin, a promoter of integrin activation, while also interacting with the A-strand of IgFLNa 20–21. When IgFLNa repeats 20–21 are in their closed conformation, the A-strand of IgFLNa 20 blocks the CD face of IgFLNa 21, thus preventing integrin binding. Fully atomistic steered molecular dynamics (MD) simulations have inferred that mechanical force applied to this domain can expose the cryptic integrin binding site, consistent with force-dependent binding of integrin peptide to FLNa in vitro17. Rognoni et al. recently reported that the disruption of the domain pair IgFLNa 20 and 21 occurs in the force range of 2–5 pN19. They estimated that domain pair disruption increases the binding affinity of β-integrin tail to IgFLNa 21 by ~17 fold. They also reported that at higher forces, the two domains mechanically unfold: IgFLNa 20 unfolding occurred at ~15 pN and that of IgFLNa 21 at >30 pN.

Here we report independent confirmation of these quantitative unfolding results. In addition, we demonstrate that unfolded domain IgFLNa 20 has two distinct states with different degrees of stability that impact the refolding kinetics of IgFLNa 20 and therefore the reformation of the domain pair at lower forces. As the experimentally determined force of ~15 pN required to unfold domain IgFLNa 20 falls within a physiological range, this finding may provide further insights to the mechanosensing function of the domain pair IgFLNa 20–21.

Results

We engineered and purified a recombinant protein construct consisting of IgFLNa 20–21: this pair of repeats was flanked by two identical “handles” comprised of IgFLNa 1–3 (Fig. 1a). As a control, we also generated a construct containing a single domain (IgFLNa 21) flanked by the same IgFLNa 1–3 “handles” (Fig. 1b). Both constructs featured a hexahistidine tag (His-tag) at the N-terminus, and a biotinylated C-terminus. In the experiments, a tether was formed by attaching the His-tagged N-terminus to an NTA-copper coated coverslip surface, whereas the biotinylated C-terminus was attached to a streptavidin-coated paramagnetic bead.

To validate our observations statistically, 29 independent protein tetters were stretched and a total of 139 unfolding or refolding events were analyzed. The results are summarized as scattered symbols of transition forces and step sizes in Fig. 2b. Three distinct data groups were revealed: two corresponding to unfolding events with a positive step size, and one corresponding to refolding events with a negative step size. From these data, histograms of the force (top) and the step size (right) were generated. The force histogram reveals the highest probability distribution peak at ~15 pN, followed by unfolding forces spread over large range up to ~120 pN. The step size histogram reveals three distinct groups: ~25 nm unfolding steps under relatively large forces (>30 pN), ~5 nm unfolding steps under ~15 pN, and around ~8 nm refolding steps under ~15 pN. Based on worm-like chain model of polypeptide, the unfolded contour length can be obtained from unfolding force and step size (Supplementary Information: Fig. S1).

To identify the unfolding and refolding events specific to the IgFLNa 20–21 domain pair, we employed a control protein construct in which the IgFLNa 20–21 domain pair was replaced with a single IgFLNa 21 domain (Fig. 2c). As with the protein construct containing the IgFLNa 20–21 domain pair, we observed ~25 nm unfolding steps with widespread, large unfolding forces (most are greater than 30 pN). By contrast, the ~8 nm unfolding and refolding steps at ~15 pN were not observed with the control construct (Fig. 2d). These results indicated that the unfolding and refolding events resulting from application of ~15 pN force, with a step size of ~8 nm, are specific to the IgFLNa 20–21 domain pair.

The nature of the unfolding signals can be compared with recent results obtained by Rognoni et al.17, which reported that (i) domain pair disruption occurred at ~2–5 pN, (ii) unfolding of IgFLNa 20 occurred at ~15 pN, and (iii) unfolding of IgFLNa 21 occurred at >30 pN. Therefore, our observations are fully consistent with their results by assigning the unfolding event at ~15 pN with step size of ~8 nm to the unfolding of IgFLNa 20. The unfolding of IgFLNa 21, which was merged with the unfolding of other Ig domains (IgFLNa 1–3) in the two handles, occurred at higher force range with a typical step size of ~25 nm. Likely due to the limitations in sampling rate of our instrument, signal of the domain pair disruption in ~2–5 pN was not observed in our experiments.

We obtained the results described above using a constant loading rate until the tether either unfolded completely or broke at high force. Since unfolding of IgFLNa 20 occurs under physiologically relevant force range, we carefully examined mechanical properties of IgFLNa 20 subjected to low forces that are sufficient to unfold this domain while keeping IgFLNa 21 intact. The curve depicted in black in Fig. 3a shows the extension change of a tether with a low loading rate of ~0.1 pN/second until the unfolding and refolding of IgFLNa 20 occurred (the spike in black curve), whereupon the force was kept constant at ~15.3 pN (the red colored curve). Rapid fluctuations with a rate of ~0.3 sec⁻¹ between two distinct extension values (red curve left to the blue vertical dashed line) were observed, 

Fig. 1 | Schematics of protein constructs. (a) Protein tether containing a pair of repeats IgFLNa 20–21 flanked by two handles consisting of repeats IgFLNa 1–3. The N-terminus is fused to a His-tag and is attached to a NTA-Cu coated cover glass surface, whereas the C-terminus is labeled with biotin and attached to a streptavidin-coated paramagnetic bead. (b) Control protein tether containing a single IgFLNa 21 domain spanned by the same handles.
presumably corresponding to transitions between the folded state N and unfolded states of IgFLNa 20.

The rapid fluctuations lasted for only ~18 seconds, after which the protein tether stabilized in the unfolded state with longer extension (red curve right to the blue vertical dashed line). After ~30 seconds without observing a transition to the folded state with the shorter extension, the force was gradually decreased to ~10 pN. However, the refolding transition did not occur over the time scale of ~40 seconds.
at these smaller forces. These results strongly suggest the existence of two distinct unfolded states of IgFLNa 20 with different stabilities under ~15 pN force: a less stable unfolded state U₁ experiencing rapid transitions with the folded state N with a lifetime of around three seconds, and another more stable conformation U₂ that remains unfolded even at decreased stretching forces of ~10 pN.

To see how the selection between the states U₁ and U₂ may be regulated by force, we studied the duration τ during which the rapid dynamic transitions between N and U₁ states were observed before switching to the stable U₂ state at constant forces in the range of 16.1 – 19.5 pN. At each force, after switching to the U₂ state, the tether was refolded at low forces (~2 pN). The time traces shown in Fig. 3b clearly indicate shorter duration τ at larger forces. The reproducibility of such force-dependent duration τ was confirmed by experiments on multiple independent protein tethers, which also showed an overall trend of shorter duration τ at larger forces (Supplementary Information: Fig. S2).

The existence of two different unfolded states of IgFLNa 20 is also supported by the level of hysteresis in extension recorded between force-increase scan and the subsequent force-decrease scan. Fig. 4a show two force-extension curves recorded in two cycles of such force scans on the same tether over a force range around 2–18 pN at a loading rate of ~0.5 pN/second in force-increase scan and ~0.5 pN/second in force-decrease scan, respectively. In both cycles, the unfolding transition was observed in 15–18 pN (black lines in Fig. 4a) during the force-increase scans. However, the force-extension curves recorded in the force-decrease scans are different between the two cycles. In the first force-cycling scan, the refolding transition occurred at ~16 pN; whereas in the second force-cycling scan, refolding of IgFLNa 20 was not observed at a force of >8 pN. Such bimodal refolding transitions obtained on different tethers are also shown to demonstrate their reproducibility (Supplementary Information: Fig. S3). These results suggest that a simple two-state model cannot explain the transitions between the folded and unfolded states of IgFLNa 20. The observation of the two different refolding transitions associated with distinct refolding forces again indicates the existence of two unfolded states U₁ and U₂ with different degrees of stability.

The force-dependent selection between the states U₁ and U₂ was also investigated by force-cycling scanning experiments. Fig. 4a and Fig. S3 have shown that scanning up to ~18 pN results in nearly equal probability of the selection between U₁ and U₂. Fig. 4b shows five successive force-cycling scans obtained from another independent experiment over a wider force range up to ~22 pN. In all the five force-decrease scans, rapid refolding and unfolding transitions were not observed, indicating only the U₂ state was selected at the higher force of ~22 pN during the force-increase scans. After the U₂ state was selected, even during force-decrease scans to a very low force range of ~2.4 to 4 pN under which the domain pair IgFLNa 20–21 may form[17], the stable unfolded IgFLNa 20 conformation U₂ still had significant probability (~50%) of being unable to refold at a time scale of 50 seconds (Fig. 4b and Supplementary Information: Fig. S4).

Fig. 5 depicts a hypothetical three-state model to describe the unfolding and refolding transitions of IgFLNa 20. The model includes a folded state N, an intermediate unfolded state U₁, and a more stable unfolded state U₂. The two unfolded states likely exhibit different conformations, but the difference in extension is too small to be resolved by the magnetic tweezers used in our experiments. In the absence of tension, the folded state N is the most stable. As force increases, the free energy landscape tilts, and, at ~15 pN, the state N and U₁ have comparable free energies, and rapid transitions between the two states occur (assuming a small energy barrier between the two states). When the tether is held for sufficiently long time under force >15 pN, a transition from the intermediate U₁ state to the more stable U₂ state occurs. A larger energy barrier separating the U₁ and U₂ states may account for the slower transition rate between U₁ and U₂. As U₂ is more stable than U₁, the U₂-to-N transition will not occur under the same force of ~15 pN. The possible physical pictures of the two unfolded states U₁ and U₂ will be discussed in the Discussion section.

**Discussion**
Our results demonstrated that unfolded domain of IgFLNa 20 can exist in two different conformational states U₁ and U₂, which lead to different refolding kinetics of the IgFLNa 20. Though the two

![Figure 4](https://www.nature.com/scientificreports/images/1642_4a.png)  ![Figure 4](https://www.nature.com/scientificreports/images/1642_4b.png)

**Figure 4** | Unfolding and refolding of IgFLNa 20 during force-cycling scans. (a) Two successive force-extension curves obtained from another protein tether recorded at a constant loading rate of 0.5 pN/sec in the force-increase scan (black), followed by ~0.5 pN/sec in the force-decrease scan (grey), over a force range up to ~18 pN. The curves are shifted in extension for clarity. In the first force-cycling scan, two-state fluctuations occurred at force ~17 pN, and remained at a state with longer extension at the end of force-increase scan. In the force-decrease scan, unfolded IgFLNa 20 refolded at force ~16 pN indicated by a step-wise extension reduction. The similar transition forces of the unfolding in the force-increase scan and refolding in the force-decrease scan indicate a reversible transition between the two states with negligible hysteresis. In the second force-cycling scan on the same tether, similar observations of the transitions in 16–18 pN between unfolding and refolding of IgFLNa 20–21 were recorded in the force-increase scan. However, refolding of IgFLNa 20 was not observed in the same force range in the force-decrease scan. All shown curves were smoothed using a 0.1 second time window from the raw data. (b) Five force-cycling scans on another independent protein tether of IgFLNa 20–21 were also performed over a wider force range up to 22 pN by changing the distance between the magnets and bead d (Supplementary Information: Magnetic tweezers setup) at a constant speed of 20 μm/sec. Unlike the constant loading rate force-scanning, this constant speed force-scanning procedure has an advantage of allowing protein refolding at low force for a longer time (Supplementary Information: Fig. S4). In all the force-decrease scans, refolding steps under the forces greater than 10 pN were not observed. Two of the four force-increase scans (excluding the first scan for the initial native protein tether) showed unfolding signals in 13–18 pN, indicating refolding probability of ~50% at low forces in our force-scanning procedure.
U2 state then represents a state where the “seed” is unfolded. In this roughly proportional to a factor $\text{Exp}^l$ picture, the “nucleation” rate under force clear, we consider a possibility that the seed is a small but stable Although the exact physical picture of the nucleated “seed” is not might associate with each other in the U2 state, leading to a larger energy barrier for refolding. However, the possibility of this event in single-molecule stretching experiments is small because 1) there are no free proteins in solution, and 2) the self-association of the unfolded domains is unfavorable under large tension. We instead propose a second possibility that the folding from the unfolded state to the native state N requires a nucleation state U1 which contains a tiny folded “seed” that adsorbs a small peptide with length of l. The U2 state then represents a state where the “seed” is unfolded. In this picture, l should be shorter than our instrumentation spatial resolution of ~2 nm since we did not directly observe it in experiments. Although the exact physical picture of the nucleated “seed” is not considered, we possibly make a possibility that the seed is a small but stable $\beta$-hairpin structure that requires looping of a short random coiled peptide chain. In this picture, the “nucleation” rate under force f is roughly proportional to a factor $\text{Exp}(-fl/k_BT)$, where $k_B$ is the Boltzmann constant, and T is the absolute temperature. At a force ~15 pN where the U1 to U2 transition was observed and with l ~2 nm, this factor is small (~10$^{-3}$), which can explain why refolding from U2 is much slower than refolding from U1 states. A possible analogy is the zipping of an unzipped DNA hairpin with a random coiled single-stranded end-loop. By choosing the sequences of the DNA hairpin and the size of the end-loop, we were able to produce a free energy profile similar to that sketched in Fig. 5b (Supplementary Information: Fig. S5).

These findings have biological implications for the mechanosensing function of FLNa. As the ~15 pN force required to unfold IgFLNa 20 approximates the force generated by several myosin motors (12), unfolding of IgFLNa 20 may occur in vivo. We showed that the mechanically unfolded domain IgFLNa 20 could have two different conformations with different stabilities. The less stable unfolded IgFLNa 20 can reversibly refold at force ~15 pN, while the more stable unfolded IgFLNa 20 exhibits a much slower refolding kinetics even at low force of smaller than 4 pN. According to the reports achieved by the authors with a control to identify signals specific to IgFLNa 20–21. Both protein constructs were labeled with His-tag at N-terminal and Avi-tag at C-terminal which was biotinylated. The DNA constructs were transfected into siF9 insect cells and recombinant proteins were purified as previously described (13, 14).

A magnetic tweezers setup was developed and used to apply force to the protein constructs (Supplementary Information: Magnetic tweezers setup). The N-terminal His-tag of the protein construct was bound to the NTA-copper coated surface of glass coverslip, whereas the C-terminal biotin was bound to a streptavidin-coated paramagnetic bead (Dynal M-270 bead, Invitrogen). Using permanent magnets placed above the sample, stretching forces were applied through the paramagnetic bead, and forces were controlled by changing the distance between the sample and the magnets. The extension of the protein construct was measured by comparing the image of the paramagnetic bead with pre-stored images of the same bead at a series of different focal planes. Drift was canceled by a reference polystyrene bead stuck on the coverslip surface.

When the magnet’s position is fixed, the force applied to the protein is a constant. With the pre-calibrated force as a function of the distance d between the permanent magnets and the sample, the magnets can approach the sample continuously to increase force exponentially or linearly with time (15).

Methods
A protein construct of IgFLNa repeats composed of IgFLNa 20–21 flanked by two handles consisting of IgFLNa 1–3 was designed. In addition to separating the domain pair IgFLNa 20–21 from the surfaces of coverslip and paramagnetic bead, the handles also provide signature of regular unfolding steps to avoid non-specific tether (which lacks the unfolding step) or multiple protein tethers (which deflects the bead in the focal plane) formed between the bead and the surface (16). A protein construct with the IgFLNa 21 flanked by the same handles was expressed as a control to identify signals specific to IgFLNa 20–21. Both protein constructs were labeled with His-tag at N-terminal and Avi-tag at C-terminal which was biotinylated. The DNA constructs were transfected into siF9 insect cells and recombinant proteins were purified as previously described (13, 14).

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1. Brown, A. E. & Discher, D. E. Conformational changes and signaling in cell and matrix physics. Current biology : CB 19, R781–789 (2009).
2. Moore, S. W., Roca-Cusachs, P. & Sheetz, M. P. Stretchy proteins on stretchy substrates: the important elements of integrin-mediated rigidity sensing. Developmental cell 19, 194–206 (2010).
3. Schwartz, M. A. Integrins and extracellular matrix in mechanotransduction. Cold Spring HarbPerspect Biol 2, a005066 (2010).
4. Stossel, T. P. et al. Filamins as integrators of cell mechanics and signalling. Nat Rev Mol Cell Biol 2, 138–145 (2001).
5. Janmey, P. A. & McCulloch, C. A. Cell mechanics: integrating cell responses to mechanical stimuli. Annual review of biomedical engineering 9, 1–34 (2007).
6. Razinia, Z., Makela, T., Ylanne, J. & Calderwood, D. A. Filamins in mechanosensing and signaling. Annual review of biophysics 41, 227–246 (2012).
7. Furukue, S., Ito, T. & Yamazaki, M. Mechanical unfolding of single filament A (ABP-280) molecules detected by atomic force microscopy. FEBS Lett 498, 72–75 (2001).
8. Gawecka, J. E., Griffiths, G. S., Ek-Rylander, B., Ramos, J. W. & Matter, M. L. R-Ras regulates migration through an interaction with filamin A in melanoma cells. PLoS ONE 5, e11269 (2010).
9. D'Addario, M., Arora, P. D., Ellen, R. P. & McCulloch, C. A. Interaction of p38 and Sp1 in a mechanical force-induced beta 3 integrin-mediated transcriptional circuit that regulates the actin-binding protein filamin-A. The Journal of biological chemistry 277, 47541–47550 (2002).
10. Nakamura, F., Stossel, T. P. & Hartwig, J. H. The filamins: Organizers of cell structure and function. Cell Adh Migr 5 (2011).
11. Ehlicher, A. J., Nakamura, F., Hartwig, J. H., Weitz, D. A. & Stossel, T. P. Mechanical strain in actin networks regulates FilGAP and integrin binding to filamin A. Nature 478, 260–263 (2011).
12. Nakamura, F., Osborn, T. M., Hartemink, C. A., Hartwig, J. H. & Stossel, T. P. Structural basis of filamin A functions. J Cell Biol 179, 1011–1025 (2007).

Figure 5 | Hypothetical model of two different unfolded IgFLNa 20 conformations. (a) A folded state N and two hypothesized unfolded states: an intermediate state U1 and a more stable state U2. The transition between states N and U1 is reversible under force of ~15 pN, while a subsequent transition from U1 to a more stable conformation U2 is irreversible. (b) Hypothesized free energy landscape of IgFLNa 20. At zero force, the folded state N is the most stable. Under ~15 pN of force, states N and U1 have similar free energies and are separated with small energy barrier, resulting reversible transition between the two states. In contrast, transition from the U1 state to the more stable U2 state is irreversible, requiring overcoming a large energy barrier.
13. Heikkinen, O. K. et al. Atomic Structures of two novel immunoglobulin-like domain pairs in the actin cross-linking protein filamin. *Journal of Biological Chemistry* **284**, 25450–25458 (2009).
14. Lad, Y. et al. Structure of three tandem filamin domains reveals auto-inhibition of ligand binding. *EMBO J* **26**, 3993–4004 (2007).
15. Nakamura, F., Stossel, T. P. & Hartwig, J. H. The filamins: Organizers of cell structure and function. *Cell Adh Migr* **5**, 160 (2011).
16. Pentikainen, U. & Ylanne, J. The regulation mechanism for the auto-inhibition of binding of human filamin A to integrin. *J. Mol. Biol.* **393**, 644–657 (2009).
17. Rognoni, L., Stigler, J., Pelz, B., Ylanne, J. & Rief, M. Dynamic force sensing of filamin revealed in single-molecule experiments. *Proc Natl Acad Sci U S A* **109**, 19679–19684 (2012).
18. Chen, H. et al. Differential mechanical stability of filamin A rod segments. *Biophys J* **101**, 1231–1237 (2011).
19. del Rio, A. et al. Stretching single talin rod molecules activates vinculin binding. *Science* **323**, 638–641 (2009).
20. Chen, H. et al. Improved high-force magnetic tweezers for stretching and refolding of proteins and short DNA. *Biophys J* **100**, 517–523 (2011).
21. Fu, H. et al. Transition dynamics and selection of the distinct S-DNA and strand unpeeling modes of double helix overstretching. *Nucleic Acids Res* **39**, 3473–3481 (2011).
22. Gesse, C. & Croquette, V. Magnetic tweezers: micromanipulation and force measurement at the molecular level. *Biophys J* **82**, 3314–3329 (2002).

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**Author contributions**
H.C. and S.C. performed the experiments. M.P.S., T.P.S., F.N. and J.Y. conceived the research. H.C., F.N. and J.Y. designed the experiments and interpreted the data. H.C., T.P.S., F.N. and J.Y. wrote the paper.

**Additional information**
Supplementary information accompanies this paper at http://www.nature.com/scientificreports

**Competing financial interests:** The authors declare no competing financial interests.

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