Vimentin intermediate filaments undergo irreversible conformational changes during cyclic loading

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Abstract

Intermediate filaments (IFs) are part of the cytoskeleton of eukaryotic cells and are thus largely responsible for the cell's mechanical properties. IFs are characterized by a pronounced extensibility and remarkable resilience that enable them to support cells in extreme situations. Previous experiments showed that under strain, α -helices in vimentin IFs might unfold to β -sheets. Upon repeated stretching, the filaments soften, however, the remaining plastic strain is negligible. Here we observe that vimentin IFs do not recover their original stiffness on reasonable time scales, and we explain these seemingly contradicting results by introducing a third, less well-defined conformational state. Reversibility on a microscopic scale can be fully rescued by introducing crosslinkers that prevent transition to the β -sheet. Our results classify IFs as a material with intriguing mechanical properties, which is likely to play a major role for the cell's local adaption to external stimuli.

Introduction

The mechanical properties of biological cells are defined by the cytoskeleton, a composite network of microtubules, actin filaments and intermediate filaments (IF)^{1,2}. Although the exact division of labour among the three filament types is still not fully resolved^{1,2}, there is ample evidence that IFs are the load bearing elements when cells are subjected to external tensile^{3,4} or compressive⁵ stress. During embryogenesis and tissue formation, in particular, cells undergo dramatic changes in shape and size. In order to withstand these transitions, cells show reversible superelasticity, which is linked to their IF network³. In order to achieve the required material properties for IFs, nature applies design principles distinct from human engineering solutions and instead relies on self-organization and structural hierarchy. As a consequence, IFs stand out among the cytoskeletal filaments by their high flexibility^{6,7} and enormous extensibility^{8–11}.

Within the IF family, vimentin is typical for cells of mesenchymal origin¹². Like all cytoskeletal IFs, vimentin monomers comprise an α -helical rod domain with intrinsically unstructured head and tail domains¹³. The monomers assemble following a hierarchical pathway resulting in filaments with laterally and longitudinally arranged monomers (Fig. 1a)^{14,15}. Theoretical considerations^{10,11}, molecular dynamics simulations^{16,17} and X-ray diffraction studies¹⁸ have shown that the intriguing tensile properties of IFs originate from conformational changes on different levels of the hierarchical filament structure. Upon stretching, the α -helical parts of the vimentin monomers and, as a consequence, also the dimeric coiled coils unfold, leading to an elongation of the filament. There is evidence from studies on vimentin and other IFs that the unfolded strands at least partly form β -sheets¹⁶⁻¹⁸. As the measured forces during relaxation of the filament are greater than zero and we previously observed only little plastic deformation upon stretching and relaxing¹¹, it was concluded that this conformational change is reversible and that relaxed vimentin IFs eventually return to the α -helical state^{10,11}. In the same study it was observed that, when repeatedly stretched, vimentin IFs become softer with every cycle, indicated by a decrease of the slope in the force-strain curve (Fig. 1b)¹¹. Monte Carlo simulations suggest that this behavior is due to the presence of mixed conformational states of parallel monomers¹¹: Fewer and fewer α -helical monomers survive, rendering the filament softer. These observations raise the question if and on which time scales a complete recovery of the initial mechanical properties, connected to refolding into α -helices, occurs.

Here, we find that, surprisingly, the initial tensile behavior cannot be recovered on reasonable time scales suggesting that this reversibility on a microscopic scale is not achieved, although the filaments show negligible excess strain after relaxation. However, crosslinking of the filaments leads to a restoration of the initial tensile behavior. We present an attempt to disentangle the seemingly contradicting results towards a comprehensive understanding of the intricate mechanical behavior of IFs. We propose a consistent mechanistic model for the observed phenomena that requires to include a third, disordered, conformational state, and postulate that the native system cannot return to the α -helix, which becomes less populated with each loading cycle.

Results & Discussion

Initial stiffness does not recover on experimental timescales

In order to examine the mechanical recovery after stretching of vimentin filaments we perform a straight-forward optical trapping experiment: Between two stretching cycles the filament is kept

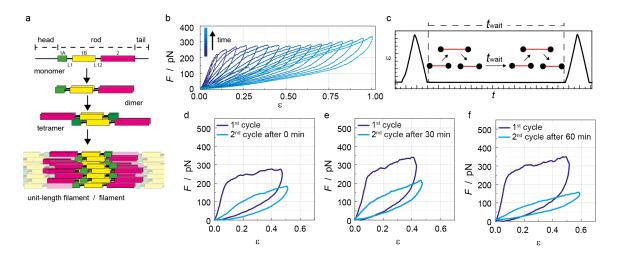


Fig. 1. Cycle experiments on untreated vimentin IFs. **a** Schematic overview of the vimentin assembly pathway from the monomer to the mature filament, via the parallel coiled-coil dimer, the half-staggered anti-parallel tetramer and the unit-length filament that on average consists of eight tetramers. **b** Vimentin filament stretched to increasing distances with each cycle (for a filament stretched for several times to the same distance see Supplementary Fig. S1), **c** Sketch of the experimental protocol for stretching cycles including waiting time t_{wait} . **d**, **e**, **f** Examples for force-strain data from experiments with different waiting times t_{wait} . **d** $t_{wait} = 0$, **e** $t_{wait} = 30$ min and **f** $t_{wait} = 60$ min. For examples of all values of t_{wait} and a box plot diagram of the input energy ratios see Supplementary Fig. S2.

at its initial contour length for a varying time t_{wait} (Fig. 1c). By this procedure, conformational changes caused by straining the filament may be reversed and the filament is allowed to return to its equilibrium conformation. The first stretching cycle shows the characteristic non-linear force response described previously¹¹ (Fig. 1d-f, dark blue), comprising an initial linear force increase, followed by a force plateau and eventually a pronounced hysteresis upon relaxation to zero strain. Note that in all experiments shown here, we pull the filament into the plateau region but not beyond. When the filament is stretched again (Fig. 1d-f, light blue), we observe softening, but almost reach zero strain at zero force. Furthermore, the hysteresis decreases, supporting that full recovery of the initial conformation is not achieved. However, the relaxation curve of the first cycle always lies below the stretching curve of the second cycle, indicating partial reversibility. The ratio of the input energies required to stretch the filament the first and second time shows that the degree of reversibility is independent of t_{wait} (Supplementary Fig. S2) even after 1 h. From these experiments, we conclude that the conformational changes occurring during the initial extension of the filament are microscopically not reversible on reasonable experimental time scales.

Crosslinking leads to reversibility on a microscopic scale

An easy way to limit the degrees of freedom upon unfolding is by introducing permanent crosslinkers, *i.e.* glutaraldehyde (GA, 5 C-atoms) and paraformaldehyde (PFA, 1 C-atom), and thereby increasing the refolding probability along the same path on which unfolding occurred. Both aldehydes may form covalent bonds with amine groups through imine formation, in particular targeting accessible lysine residues. The longer the crosslinker, the more links are expected to form.

Qualitatively, the force-extension curves are not altered by crosslinking and show the same characteristic force plateau as found for untreated filaments, albeit shifted to higher forces (Fig. 2a).

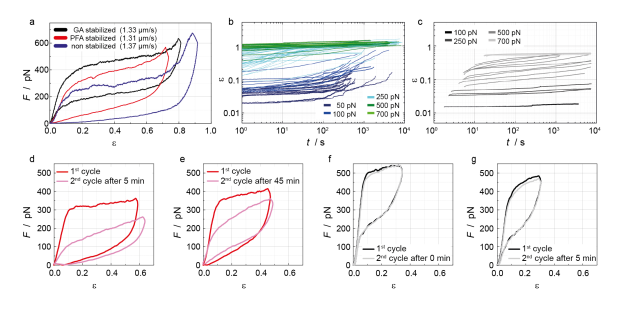


Fig. 2. Recovery experiments with crosslinked IFs. **a** Typical force-strain cycles of untreated (blue), PFA-crosslinked (red) or GA-crosslinked (black) filaments. **b** Force clamps of untreated vimentin IFs ranging from 50 to 700 pN, presented as a log-log plot of strain versus time.¹¹ **c** Force clamps of GA-crosslinked vimentin IFs ranging from 100 to 700 pN, presented as a log-log plot of strain versus time. **d**, **e**, Examples for recovery experiments performed with PFA-crosslinked vimentin IFs with $t_{wait} = 5 \min (d)$ and $t_{wait} = 45 \min (e)$. For examples of all values of t_{wait} and a box plot diagram of the energy ratios for PFA crosslinked vimentin IFs with $t_{wait} = 5 \min (d)$ and $t_{wait} = 0 \min (f)$ and $t_{wait} = 5 \min (g)$. For examples of all values of t_{wait} and a box plot diagram of the energy ratios for DFA crosslinked vimentin IFs with $t_{wait} = 5 \min (g)$. For examples of all values of t_{wait} and a box plot diagram of the energy ratios for PFA crosslinked vimentin IFs with $t_{wait} = 5 \min (g)$. For examples of all values of t_{wait} and a box plot diagram of the energy ratios for SI crosslinked vimentin IFs with $t_{wait} = 0 \min (f)$ and $t_{wait} = 5 \min (g)$. For examples of all values of t_{wait} and a box plot diagram of the energy ratios for GA crosslinked filaments, see Supplementary Fig. S4.

For native, untreated vimentin IFs the origin of the force plateau was previously assigned to an increase of the contour length of the filament due to force-induced conformational changes^{10,11,17}, a mechanism that appears to be at least partially conserved in crosslinked filaments. The relaxation curves of GA-crosslinked filaments, however, show a distinctly different progression compared to the untreated filaments with a force plateau and an inflection point, similar to the extension curve, albeit at a lower overall force. Force-clamp data of untreated and GA-crosslinked filaments (Fig. 2b, c) confirm that a much higher force is needed to extend the crosslinked filaments. Whereas the untreated filaments can be stretched up to strains higher than 1.0 at 250 pN, GA-crosslinked filaments stay at strains below 0.1 even after two hours at this force level. The force needed to stretch GA-crosslinked filaments to strains of about 0.8 is most likely around 400 pN.

Repeated force-distance cycles of crosslinked filaments reveal striking differences to untreated IFs concerning the reversibility of unfolding. When PFA-crosslinked filaments are stretched again after $t_{wait} = 5$ min, they behave similar to untreated filaments (Fig. 2d). However, for $t_{wait} = 45$ min, there is a clear recovery towards the initial curve shape (Fig. 2e). By contrast, GA-crosslinked filaments show identical force-distance curves in the first and the second cycle even without any waiting time (Fig. 2f,g). These observations are confirmed by the input energy ratio as a function of t_{wait} (Fig. 3a). Notably, at low strain the GA-crosslinked filaments behave fully elastic, *i.e.* at small strains the relaxation curve reproduces the initial slope (Fig. 2f,g). Thus, crosslinking with GA remodels the energy landscape of IF stretching and crosslinked filaments return to the initial conformation.

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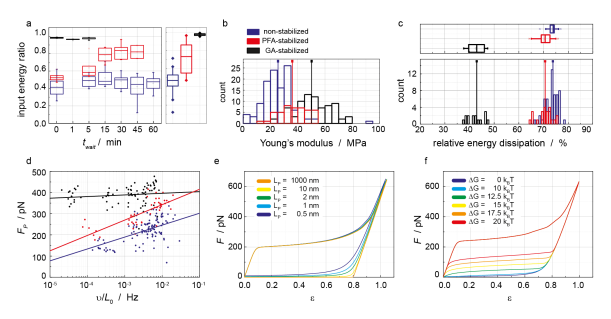


Fig. 3. Physical analysis and numerical simulations of vimentin IF mechanics. **a** Left: ratio of input energies between the first and the second stretching event versus t_{wait} . Right: distribution of input energy ratios for all t_{wait} . The input energy ratio for PFA-treated filaments shows a slow increase and reaches values of about 0.85 after 30 min (red). For GA-treated filaments the input energy ratio stays constant around 1 as the first and second force-distance cycles show almost identical progressions (black). **b** Histograms of Young's moduli of untreated, PFA- and GA-stailized IFs calculated from the initial slope (Supplementary Fig. S5) of the first force extension curve of each vimentin IF assuming a circular filament cross-section and a filament radius of 5 nm. Legend and color codes apply to a,c, and d as well. **c** Bottom: histograms of relative dissipated energies for untreated, PFA- and GA-crosslinked filaments. Top: distributions of dissipated energy. **d** Force at the onset of unfolding (*i. e.* the beginning of the plateau region) versus the normalized loading rate (Supplementary Fig. S5). **e** Simulation of force-strain curves according to Block et al.¹¹ with back-reaction for different ΔG .

Crosslinking alters the energy landscape of strained IFs

These results raise the question whether there exists a fundamental difference between the conformational changes during stretching for crosslinked and untreated filaments. The Young's modulus (Fig. 3c) already shows a clear trend. It increases from 26 MPa to 36 MPa and eventually to 50 MPa for untreated (blue), PFA-crosslinked (red) and GA-crosslinked IFs (black), respectively, indicating a strong effect caused by crosslinking. Fig. 3b shows histograms of the energy dissipated during the first stretching-relaxing cycle, normalized by the input energy. For PFA-treatment (red), similar values (\sim 70-80%) to untreated filaments (blue) are found¹¹, whereas GA treatment (black) drastically decreases the energy dissipation to only \sim 45%.

To examine the energy landscape associated with folding and unfolding, the force at the onset of the plateau is determined and plotted against the loading rate v normalized by the initial filament length L_0 (Fig. 3d). Whereas the plateau forces for PFA-crosslinked and untreated filaments show a similar velocity dependence, albeit with slightly higher plateau forces for the crosslinked filaments, the GA-crosslinked filaments show very high plateau forces around 400 pN that are almost independent of the loading rate. We have shown previously that the force-extension curve of vimentin can be modelled with an elastically coupled two-state-model¹⁰. This model predicts a logarithmic scaling of the plateau force with pulling velocity (see Methods, equation 5). For the untreated filaments we find a potential width $x_u = 0.17 \pm 0.02$ nm (mean \pm st.d.) and a zero-force reaction rate of $k_0 = (5 \pm 4) \cdot 10^{-7}$ s⁻¹ in good agreement with the parameters found

previously by directly fitting to individual force-strain curves¹⁰. Treatment with PFA leads to an only slightly decreased potential width of $x_u = 0.13 \pm 0.02$ nm and zero-force reaction rate of $k_0 = (2.5 \pm 2) \cdot 10^{-7} \text{ s}^{-1}$. By contrast, GA drastically alters the energy landscape. The reaction is slowed down dramatically, $k_0 = (1 \pm 8) \cdot 10^{-54} \text{ s}^{-1}$, which is accompanied by an enormous increase of the potential width, $x_u = 1.2 \pm 1$ nm. This huge change probably indicates that the molecular elongation mechanism is fundamentally different after GA-treatment. Therefore, it is remarkable that basic characteristics of the tensile behavior like the high extensibility and the force plateau are largely conserved after crosslinking.

We can only speculate about the molecular origin of the observed changes due to crosslinking. However, a previous crosslinking study employing disulfosuccinimidyl tartrate (DST) led to a dense network of crosslinks (12 possible crosslinking positions within the coiled coil and between dimers of fully assembled filaments) in vimentin IFs^{19,20}. It is expected that these crosslinks prevent the necessary rotation of the peptide chain for the α -helices to completely unfold, so we regard them as pinning points for the protein conformation, which ensure the presence of a reasonably close starting point for the reformation of the α -helices.

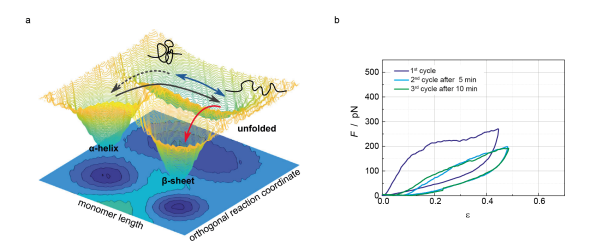
Molecular dynamics simulations have shown that coiled coils need a critical length of about 40 amino acids in order to be able to undergo an α - β -transition²¹, which is in good agreement with the dramatically increased stability we observed for the short α -helical segments between the crosslinks. Furthermore, we assume that, when the α -helices in GA-crosslinked filaments unfold, parallel helices have to unfold simultaneously since they are covalently connected. This is in contrast to the untreated filament, where computational models have predicted that parallel unfolding is initiated by stochastic opening of a few monomers which leads to loss of stability of all remaining parallel helices, similar to what has been observed in binding clusters^{22,23}. Using the analogy to binding clusters, this can be regarded as an extreme case of shared loading which might explain the huge increase of the potential width^{24,25}.

Impact of reversibility on the relaxation curve

The relaxation force curve of untreated vimentin IFs as shown in Fig. 2a, blue, decays almost linearly from a restoring force around 100 pN at a strain of 0.8 to zero force at strains below 0.1. This positive restoring force was earlier interpreted as a sign of reversibility of the underlying molecular process. This can be illustrated by modelling the force extension curve using a twostate-model for the filament elongation introduced earlier¹¹. If we switch off the back-reaction in the model (Fig. 3e) the force drops to zero already at large strains around 0.8 if we assume that the persistence length of the filament stays constant at 1 μ m upon stretching (orange). If, however, the persistence length is decreased dramatically upon stretching, a force progression closer to the experimental relaxation curve is recovered (green, blue, purple). This is plausible, since a thinning of IFs upon stretching has been observed^{8,9} and would lead to smaller persistence lengths. Furthermore, since coiled coils are particularly rigid, it is likely that the filament gets more flexible when being unfolded. Indeed, a similar mechanism with a drastic decrease in persistence length upon unfolding was proposed by Minin et al. for the α - β -transitions in simpler coiled-coil peptides²⁶. In contrast to an earlier interpretation, we conclude that entropic elasticity of the protein chain is sufficient to explain the restoring force upon filament relaxation for untreated vimentin filaments.

The force plateau and inflection point in the relaxation curves of GA-crosslinked filaments (Fig. 2a, black), are reproduced by the two-state-model when increasing the energy difference ΔG between the shorter and the longer state and thereby the reversibility of the transition (Fig. 3f).

This indicates that the positive restoring force of GA-crosslinked filaments originates from the refolding enthalpy of the α -helices. Interestingly, a similar increase in reversibility might also be achieved by a symmetrization of the energy landscape.



Recovery of initial contour length indicates third conformational state

Fig. 4. Introduction of a third conformational state during vimentin IF stretching. **a** Energy landscape illustrating the proposed model. Besides the two well-defined α -helical and β -sheet conformations, there exists a less defined unfolded conformation along an orthogonal reaction coordinate. When a force is applied, the α -helix can unfold (black arrow) and the unfolded random coil can be stretched (blue arrows). **b** An untreated vimentin IF is cycled three times with $t_{wait}=5$ min between the cycles. Whereas the force-strain behavior of the second cycle differs considerably from the first one, the data for the third cycle are almost identical to the second one.

The strongest argument for the proposed reversibility of vimentin elongation was the observation that repeated cycles always start at nearly the same strain (Fig. 1a) and that only very little residual plastic strain of vimentin was observed (Supplementary Fig. S6). The softening of filaments without residual strain was explained by mixed states – α -helical and elongated – in the parallel monomers. However, the observed force plateau requires all parallel monomers in a unit length filament (ULF) to elongate in a cascading manner¹¹. Thus the softening with repeated cycles can only be explained when there is a way for elongated monomers to return to a shorter effective length, which seems to contradict the non-reversibility we observe here.

To resolve the contradiction, we propose a model for which the transition between the shorter and the longer state is not a one-step process as assumed in the two-state-model proposed previously¹⁰, but consists of three distinct states (Fig. 4a). When a force is applied to the filament, the α -helical coiled coils unfold (black arrow) and the polypeptide chains assume a random coil. This unfolding can be described as a transition along a reaction coordinate orthogonal to the applied force. Due to its high energy cost this reaction does not occur spontaneously but only at high loads around 200 pN, *i.e.* the plateau force. The unfolded chain can elongate in response to the applied strain leading to a reversible extension of the monomer (blue arrow). This elongation has presumably not only an entropic cost to extend the semiflexible polypeptide chain, but also an enthalpic component due to the breaking and formation of hydrogen bonds, rendering the energy landscape of the unfolded state rather rough. This roughness of the energy landscape might indeed give rise to the viscoelastic relaxation behavior investigated earlier, where we find a power law behavior¹¹. When the random coil is elongated, hydrogen bonding between parallel polypeptide chains can lead to the formation of β -sheets (red arrow). This process is presumably rather slow, since the polypeptide chains need to assume the exact right geometry, and irreversible, due to the high energy difference. When the filament is subsequently relaxed, only the peptide chains that were not able to form a β -sheet will return to a shorter contour length. Since the relaxed unfolded polypeptide chain can assume many possible conformations, reformation of α -helices is highly unlikely (dashed arrow).

Our model is supported by an experiment, where we cycle an untreated filament three times with a waiting time of t_{wait} = 5 min between the cycles. Whereas the data for the second cycle (light blue) differ from the first one (dark blue), in line with the data shown in Fig. 1, the third cycle (green) progresses in an identical manner to the second one. In our model energy land-scape (Fig. 4a) we interpret this as follows: upon first stretching, the α -helices are unfolded into random-coils, which are subsequently elongated. Relaxation and repeated stretching leads to "shuffling" between the short and the long random-coil state and thus to identical force-strain curves.

This model is sufficient to justify why the filament always returns to essentially the same contour length, while also explaining the observed irreversibility. Apparently, PFA crosslinked polypeptide chains are kept close enough to the starting conformation that slow refolding of the α -helices is observed, leading to a slow recovery of the initial force extension curve. By contrast, the energy landscape of the GA-crosslinked filaments is strongly altered, and rather resembles the idealized two-state-model, with an energetically heavily favored shorter state and a spontaneous back-reaction from the elongated state. From our measured force-strain cycles (Fig. 2a, black) and the proposed model, one may draw an analogy to pseudoelastic (also referred to as superelastic) shape-memory alloys, which, when heated after being strained, return to a previous shape²⁷. Of note, however, in our system, we do not observe a phase transition, as required for pseudoelasticity.

Conclusion

By employing controlled *in vitro* experiments, we are able to precisely tune mechanical strainresponse of vimentin IFs to applied stress. Untreated IFs can be stretched to high strains and largely recover their original length upon relaxation, *i.e.* only neglectable residual strain is observed. As the filaments soften with each subsequent cycle and the original stiffness is not recovered on reasonable experimental time scales, we examine the energy landscape of strained IFs in detail. Our experiments lead to the conclusion that besides the α -helix and the β -sheet a third – presumably random coil – conformational state must exist, and that subunits are not able to return to their initial state after stretching. This irreversible transformation is avoided when introducing permanent crosslinkers into the system and thereby reducing the degrees of freedom upon stretching.

The irreversibility of the conformational changes we observed here might indicate that vimentin filaments that were strained in the past are not as capable to protect the cellular integrity. This implies that cells need a mechanism to "repair" strained vimentin. This reversibility may be achieved by either subunit exchange along the filaments^{28–30}, which is, however, very slow, or by disassembly and reassembly of filament structures as observed *e.g.* for Keratin networks^{31,32}.

Interestingly, the microscopic reversibility and the increased stiffness we observed as a result of GA-fixation seem to be advantageous mechanical properties for cells that are heavily subjected to external strains. For keratin IFs in maturating keratinocytes, controlled covalent crosslinking of subunits by disulfid bond formation is indeed observed and might be a possible avenue for cells to tune their resistance to external forces^{33,34}. Interestingly, vimentin was identified as a substrate for proteins introducing covalent crosslinks between different monomers, specifically transglutaminases, in two human tissues where structural integrity is paramount, arteries and lens^{35,36}. Like the crosslinkers used in our work, transglutaminases target primary amines suggesting they might alter the energy landscape in a way similar to what we show in our *in vitro* system.

Methods

Vimentin production and purification

The production and purification of recombinant human vimentin C328A with additional amino acids GGC at the C-terminus for force clamp and multiple-cycle-measurements was performed as described previously¹¹. For all other experiments the protein purification was performed similar to the protocol described previously¹¹ but with several changes to the buffers used, as described in the following.

Escherichia Coli bacteria (100 μ L) were thawed on ice, mixed with 2 μ L plasmid solution (concentration of 10 ng/ μ L) per 50 μ L bacteria solution and cultured on an agar plate¹¹. A single bacteria colony was picked and transferred into 5 mL terrific broth medium (TB, 47.6 g/L (#T0918, Sigma-Aldirch), containing glycerol (8 mL of 99.5 % glycerol (#3873, Sigma-Aldrich) per 1 L TB medium) and ampicillin (Sigma-Aldrich, added at a concentration of 1:1000) and incubated at 37 ° C at 220 rpm for four to six hours. Afterwards, the bacteria solution was transferred to 50 mL TB medium and allowed to grow for another four to six hours. Finally the bacteria were transferred to 1 L of TB medium and cultured over night at 37 °C at 150 rpm.

Bacteria were pelleted with a Beckmann Centrifuge (Beckmann Coulter Avanti J-26 XP, rotor JS5.3) at 5000 *g* at 4 °C for 25 min. For purification, the pellet was transferred to a cooled Douncer and kept on ice for all subsequent steps. The pellet was dissolved in 16 mL lysis buffer (50 mM TRIS (Carl-Roth GmbH), pH 8.0, 25 % w/v saccharose, 1 mM EDTA (Carl-Roth GmbH), 1 mM Pefabloc SC[®] (Carl-Roth GmbH), cOmpleteTM tablets (1 per 50 mL, Sigma-Aldrich)). 4 mL lysozym (Roche Diagnostics) solution (10 mg/mL in lysis buffer) were added, the mixture was homogenized and incubated on ice for 30 min. 200 μ L of 1 M MgCl₂ (Sigma-Aldrich), 20 μ L RNAseA (10 mg/mL in TRIS buffer, pH 7.5, Roche Diagnostics), 2 μ L benzonase (#71206-6, Novagen) and 400 μ L 10 % NP-40 (Roche Diagnostics) were added, the solution was again homogenized and incubated for 10 min. 40 mL detergent buffer (200 mM NaCl (Carl-Roth GmbH), 1 % NP40, 1 % DOC (sodium-deoxycholat, Sigma-Aldrich), 20 mM TRIS, 2 mM EDTA, 1 mM Pefabloc SC, cOmpleteTM tablets (1 per 50 mL)) were added, the mixture was again homogenized for 10 min, and then centrifuged for 30 min at 10,000 *g* and 4 °C using a Beckmann centrifuge (J26XP, rotor JLA 16.250).

The supernatant was discarded and the pellet washed by homogenization using 4 °C cold GII-buffer (10 mM TRIS, 0.1 Vol% TritonX-100 (Carl-Roth GmbH), 5 mM EDTA, pH 8.0) complemented by 40 μ L 1 M DTT (1,4 dithiothreitol, Carl-Roth GmbH) and 100 μ L Pefabloc SC. The solution was incubated on ice for 10 min and then centrifuged for 30 min using the same parameters as in the step before.

The supernatant was discarded and the pellet washed by homogenization using 4 °C cold GII-KCl-buffer (1.5 M KCl (Carl Roth GmbH) in GII-buffer) complemented by 40 μ L 1 M DTT and 100 μ L Pefabloc SC. The solution was incubated on ice for 30 min and then centrifuged for 30 min using the same parameters as in the step before.

The supernatant was discarded and the pellet was washed by homogenization using $4 \degree C$ cold TE-buffer (10 mM TRIS, 0.1mM EDTA, pH 8.0) complemented by 40 μ L 1 M DTT and 100 μ L Pefabloc SC. The mixture was incubated on ice for 10 min and then centrifuged for 25 min using the same parameters as in the step before.

The supernatant was discarded and the pellet solubilized in hand-warm urea buffer. For the urea buffer 12.125 mL 9.5 M urea solution was complemented by $125 \,\mu$ L 1 M TRIS, $125 \,\mu$ L 0.5 M EDTA and $125 \,\mu$ L 1 M DTT, pH 7.5, and the pellet solubilized in as less buffer as possible. After homogenization the solution was finally centrifuged at 20 °C and 100,000 *g* for 60 min, using a Beckmann ultracentrifuge (Beckmann Coulter Optima L90K, rotor Ti70). The supernatant that contains the vimentin was in some cases stored at -80 °C.

The final purification by anion and cation exchange chromatography was performed as described in Block et al.¹¹ without any changes.

Vimentin and bead functionalization

Vimentin was labeled, reconstituted, and assembled according to previously published protocols. Labeling with ATTO647N (ATTO-Tech GmbH) and biotin-maleimide (Jena BioSciences GmbH) was performed according to Block et al.¹⁰ and Winheim et al.³⁷. Reconstitution and assembly were performed as described by Block et al.¹¹.

Carboxylated polystyrene beads were functionalized according to Janissen et al.³⁸ as described in Block et al.¹¹.

Optical tweezers measurements

Optical tweezers measurements were performed using a commercial instrument (C-trap, Lumicks, Amsterdam, Netherlands), combining optical tweezers with a microfluidic setup and confocal microscopy (Supplementary Fig. S7), similar to what was described previously¹¹. In short, a fresh pair of beads was captured for each experiment in the bead channel. A vimentin IF was captured in the vimentin channel and bound to the second bead in the buffer (100 mM KCl in 2 mM phosphate buffer, pH 7.5) channel. Parts of the measurements were performed with filaments bound to the beads by maleimide-chemistry and parts of the experiments with filaments bound via biotin-streptavidin. The covalent attachment of the filaments via maleimide-groups enables us to introduce waiting times between the filament stretching cycles. For crosslinking experiments the captured vimentin IF was moved to channel 4 (Supplementary Fig. S7) filled with buffer containing either PFA (0.12 % (v/v)) or GA (0.5 % (v/v)). Before a force-distance curve was recorded, the filaments were always moved back to the channel that only contains buffer (channel 3, Supplementary Fig. S7).

Following the measurement protocol sketched in Fig. 1b, we stretched the filament beyond the linear regime into the characteristic plateau region, where unfolding of the α -helices occurs. Subsequently, we relaxed the filament and, after a defined time $t_{wait} \leq 1$ h, stretched the filament again to the same trap position as in cycle one. The length of the filament and the applied force were recorded over the full experiment.

Data Sets

For the untreated filaments 57 individual recovery experiments were performed. The total number of recovery measurements for GA-crosslinked filaments was 15 and for PFA-crosslinked filaments it was 23. For the force-clamp experiments of untreated filaments about 100 measurements were performed. The number of force-clamps of GA-crosslinked filaments was 18.

Data analysis

Calculation of the energy ratio

All data were converted to ASCII format with the measuring software (TWOM, Lumicks). The bead diameter was subtracted from the distance and the time and force data were used as measured. In order to prevent the filaments from wrapping around the beads during the waiting time, a prestrain of 5 pN was imposed for all recovery experiments. Before the actual measurement, this value was set to 0 pN and this prestrained length of the filament was used as initial length L_0 to calculate the dimensionless strain during stretching.

To calculate the energy for the first cycle, the area beneath the stretching curve up to the maximum strain was calculated by integration. As we can only control the position of the trap and not the position of the bead, the maximum strain of the second stretching is slightly higher than the maximum strain for the first stretching cycle, due to the softening of the filament. To compare the input energy that was needed to strain the filament to the same length again, the energy needed for the second stretching was calculated by integrating the area beneath the second stretching curve but only up to the maximum strain of the first stretching curve. The two energy values were then divided to obtain the input energy ratio.

Determination of the Young's modulus

In order to determine the Young's modulus, the initial slope of the force extension curve was determined by a linear fit. Only data points with forces larger than 10 pN and smaller than 80% of the plateau force and at positive strains where considered.

Determination of the plateau force

The force at the onset of the plateau was determined by connecting the first and the last data point of an extension curve with a straight line and subtracting this straight line from the force curve. This effectively tilts the force curve, such that the beginning of the plateau can be determined as the maximum of the tilted curve.

Analytical solution for the two-state-model

Here we use an approximate analytical solution of the two-state-model for filament stretching. The full theory and its applicability have been described elsewhere^{10,11,39}. In brief, every ULF in the filament is assumed to be able to adopt a shorter and a longer state with a force dependent transition between them. These states do not necessarily need to be identical to defined secondary structures like the α -helix or the β -sheet, but can be interpreted as a two-dimensional projection of the higher dimensional energy landscape shown in Fig. 4a. This model gives us the force dependent contour length L_C of the filament that can be combined with elastic and entropic contributions to calculate the shape of the force-distance curve. The approach for the analytical solution is mostly adapted from Burte and Halsey³⁹. The key step for the analytical solution is to ignore elastic and entropic effects such that the filament length equals the contour length $L = L_C$. When we neglect the back-reaction to the shorter conformation, which is a good

approximation for the extension curve, an independent reaction of the separate helices in the vimentin monomer rate equation reads

$$\frac{\mathrm{d}N}{\mathrm{d}t} = (N_0 - N)k_0 \exp\left(Fx_u/k_BT\right),\tag{1}$$

with the number of ULFs in the longer state N, time t, the total number of ULFs in the filament N_0 , the zero force reaction rate between the shorter and the longer state k_0 , the force F, the potential width x_u , Boltzmann's constant k_B and the temperature T. Using the length change of a ULF during the conformational change $\Delta l = dL/dN$ we obtain:

$$\frac{dL}{dt} = \Delta l(N_0 - N)k_0 \exp\left(Fx_u/k_BT\right)$$
⁽²⁾

$$= L_0(\Delta l/l - \varepsilon_{\text{approx}})k_0 \exp\left(Fx_u/k_BT\right)$$
(3)

with the length of an ULF *l*, the initial filament length L_0 and the strain $\varepsilon_{approx} = N\Delta l/L_0$ (excluding elastic and entropic contributions). If a filament is stretched at a constant velocity, and elastic and entropic contributions are ignored, we can set dL/dt = v. Solving for the force gives us

$$F = \frac{k_B T}{x_u} \ln \left(\frac{v}{k_0 L_0 (\Delta l/l - \varepsilon_{\text{approx}})} \right)$$
(4)

Elastic contributions play a significant role in the force response of filaments and can be introduced via an elastic correction $\varepsilon_{\text{corr}} = \varepsilon_{\text{approx}} + F/k_{\text{eff}}L_0$ using the effective spring constant of the filament k_{eff} . Comparing this approximate solution to the numerical solution shows good agreement at low to intermediate strains (Supplementary Fig. S8).

A simple scaling law can be derived from equation 4 for the force F_P at the onset of the force plateau when the strain is completely caused by elastic deformation, *i.e.* $\varepsilon_{approx} = 0$. We find

$$F_P = \frac{k_B T}{x_u} \left(\ln(v/L_0) - \ln(k_0 \Delta l/l) \right)$$
(5)

This equation can now be used to independently determine x_u and k_0 by performing stretching experiments at different velocities and plotting F_P as a function of the normalized velocity v/L_0 and estimating ε_{max} based on structural data or from the length of the force plateau.

It has been shown previously that the velocity dependence of vimentin stretching could only be reproduced if independent reactions of subsections of the peptide chain were assumed. When we have multiple subunits (for example the three helices that comprise the vimentin monomer) the differential equation 3 becomes a sum of their respective contributions L_i and we get

$$\frac{\mathrm{d}L}{\mathrm{d}t} = \sum_{i}^{3} \frac{\mathrm{d}L_{i}}{\mathrm{d}t}.$$
(6)

This equation cannot be solved without further constrains. However, we can expect that the scaling in equation 5 still holds, since due to the exponential force dependence the element with the largest x_u will dominate at the onset of the plateau. Therefore, equation 5 was fitted to the plateau force as a function of the velocity normalized by the initial filament length v/L_0 . To estimate the uncertainty of the parameters, a bootstrapping algorithm⁴⁰ with 10000 iterations was used.

Data availability

The data that support the findings of this study are available from the corresponding authors, at sarah.koester@phys.uni-goettingen.de, or andreas.janshoff@chemie.uni.goettingen.de, upon reasonable request.

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Author contribution

SK conceived the project. JF and JK each performed parts of the experiments and the data analysis. HW performed parts of the data analysis, developed and implemented the models and performed all simulations. AJ and SK supervised the project. All authors wrote the manuscript.

Additional Information

Supplementary Information accompanies this paper at *URL inserted by publisher*.

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