

A Comparison of the Mechanical and Structural Properties of Fibrin Fibers with Other Protein Fibers

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Abstract In the past few years a great deal of progress has been made in studying the mechanical and structural properties of biological protein fibers. Here, we compare and review the stiffness (Young's modulus, E) and breaking strain (also called rupture strain or extensibility, ϵ_{\max}) of numerous biological protein fibers in light of the recently reported mechanical properties of fibrin fibers. Emphasis is also placed on the structural features and molecular mechanisms that endow biological protein fibers with their respective mechanical properties. Generally, stiff biological protein fibers have a Young's modulus on the

order of a few Gigapascal and are not very extensible ($\epsilon_{\max} < 20\%$). They also display a very regular arrangement of their monomeric units. Soft biological protein fibers have a Young's modulus on the order of a few Megapascal and are very extensible ($\epsilon_{\max} > 100\%$). These soft, extensible fibers employ a variety of molecular mechanisms, such as extending amorphous regions or unfolding protein domains, to accommodate large strains. We conclude our review by proposing a novel model of how fibrin fibers might achieve their extremely large extensibility, despite the regular arrangement of the monomeric fibrin units within a fiber. We propose that fibrin fibers accommodate large strains by two major mechanisms: (1) an α -helix to β -strand conversion of the coiled coils; (2) a partial unfolding of the globular C-terminal domain of the γ -chain.

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Introduction

Across biological systems, an array of intra- and extra-cellular protein fibers perform various mechanical functions. As is the case with all biological components, these fibers evolved structural and mechanical properties that are appropriate for these functions. Two important mechanical properties are the stiffness and the breaking strain (extensibility) of these fibers (detailed definitions of mechanical properties: see Section "Mechanical Measurements on

Single Fibrin Fibers and Fibrinogen Molecules”). A review of the literature reveals that stiff fibers are usually not very extensible. These fibers often have a regular, paracrystalline (nearly crystalline) structure and/or are crosslinked. Examples include actin filaments, microtubules, collagen, matrix-embedded keratin fibers, and the spokes of spider webs. On the other hand, softer fibers are often very extensible. The structure of these fibers is often amorphous and/or contains less crosslinking; though these fibers utilize different molecular mechanisms to achieve high extensibility. Examples of soft, extensible fibers include elastin, resilin, fibrillin, fibronectin, intermediate filament, myofibrils, mussel byssal fibers, and the catching thread of spider webs.

It was recently discovered that fibrin fibers, which are the major structural component of a blood clot, are extraordinarily extensible and elastic [1], and that they are relatively soft [2]. This was unexpected, because fibrin fibers have a regular, paracrystalline structure [3, 4] and crosslinked monomer units.

In this review we compare the mechanical and structural properties of fibrin fibers with those of other polymerized protein fibers. This comparison leads us to suggest feasible molecular mechanisms that allow for the large extensions of fibrin fibers despite their nearly crystalline structure.

This review is divided into four sections:

- (1) Fibrin(ogen) structure and fibrin fiber assembly.
- (2) Mechanical measurements of fibrin fibers and *single* fibrinogen molecules.
- (3) Stiffness (Young's Modulus) and breaking strain (extensibility) of fibrin fibers and other polymerized protein fibers.
- (4) Proposed molecular mechanisms for fibrin fiber extension.

Fibrin(ogen) Structure and Fibrin Fiber Assembly

Fibrinogen is a highly abundant, soluble plasma protein. Removal of two pairs of fibrinopeptides converts it into fibrin monomers, which polymerize into a meshwork of fibrin fibers, the basic structural component of a blood clot. Fibrinogen consists of six peptide chains (2A α , 610 residues; 2B β , 461 residues; 2 γ , 411 residues; human numbering is used throughout this article). The recently solved crystal structures of human fragment D [5], bovine fibrinogen [6] and *native* chicken fibrinogen [7] (Fig. 1) added much clarity to the structure of fibrinogen and corroborated many features that had been gleaned from previous experiments. Fibrinogen has an approximately centrosymmetric, trinodular, S-shaped structure and is 46 nm in length and 4.5 nm in diameter [9–11]. Two nodules

(D nodules) are at either end of the protein and one nodule (E nodule) is in the center of the protein. The nodules are connected via two, 17 nm-long coiled coils comprised of three α -helices, including residues α 51–161, β 85–197, and γ 33–143. The D nodule contains the globular β C-terminal domain (β 197–461; called β C) and the globular γ C-terminal domain (γ 143–411; called γ C), both of which consist of a β -sheet core flanked by a few small α -helices. The central, globular E-nodule contains all six N-termini and also fibrinopeptides A and B. The α C-terminal threads briefly through the D nodule, rejoins the coiled coils as a fourth helix (α 164–220) and ends with the segment called the α C domain (α 220–610) that stretches from the distal D-nodule back towards the central E-nodule. The α C domain is mobile, and contains little well-defined secondary structure, although there is some evidence that this region consists of a flexible connector region (221–391) and a globular unit (392–610) [12–14]. This segment is shorter in chicken fibrinogen, α 220–491 (chicken numbering). Although this segment is present in the crystals of chicken fibrinogen, the electron density for this region is too weak to resolve a structure, as expected for a segment with high mobility. Hence, these residues are not depicted in Fig. 1.

Disulfide Bonds

Fibrinogen contains 29 disulfide bonds that stabilize the molecule. Five disulfide bonds are within the central E nodule linking together the two halves of the molecule. Prominent features are the four “disulfide rings” in each fibrinogen molecule; there is one ring at each end of the two coiled coils. Each “disulfide ring” consists of three disulfide bonds, linking the α - to the β -chain, the α - to the γ -chain, and the β - to the γ -chain, thus forming a brace that keeps the three α -helices in the coiled coils together. Last, there are twelve intra-chain disulfide bonds, three in each of the two globular β C domains (D-nodules), two in each of the two globular γ C domain (D-nodules), and one in each of the two α C domains.

Fibrin fibers are assembled from fibrin monomers and the widely accepted sequence of events leading to fibrin fiber formation, starting with fibrinogen, is outlined in Fig. 1. Activated thrombin proteolytically removes fibrinopeptides A (16 residues) and B (14 residues) from the N-terminal region of the A α - and B β -chains, respectively, thereby converting fibrinogen into fibrin monomers and exposing binding sites (“knobs”) A (Gly-Pro-Arg-Val) and B (Gly-His-Arg-Pro). These positively charged “knobs” fit into negatively charged binding pockets “a” and “b” in the γ - and β -chains within the distal globular D domains, respectively, thus forming the A:a and B:b interactions. Pocket “a” in γ C is composed of four loops including key

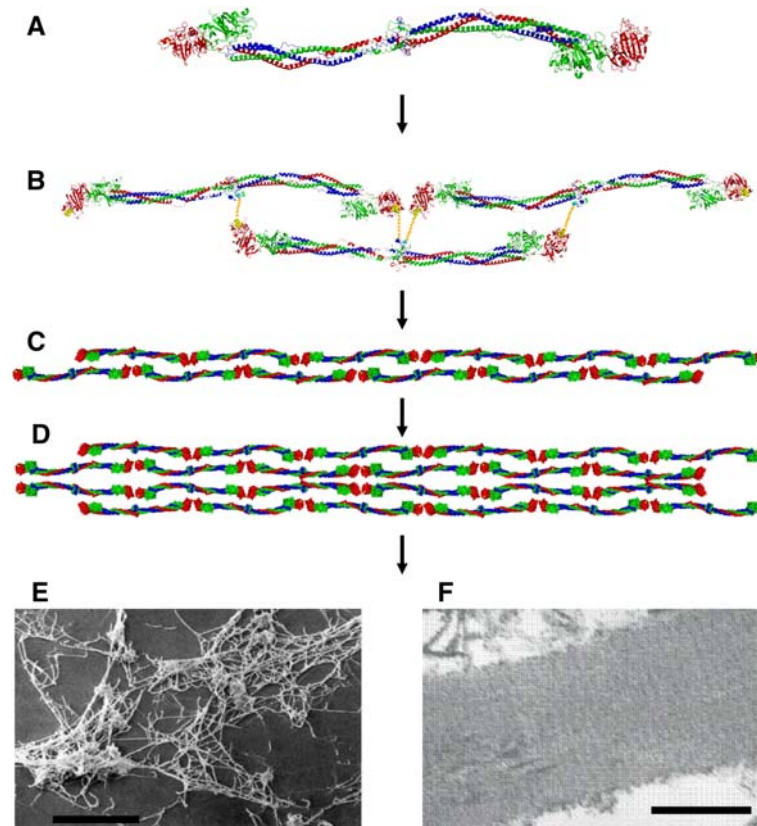


Fig. 1 Crystal structure of chicken fibrinogen [7] and fiber assembly. (A) The fibrinogen molecule is 46 nm long and 4.5 nm in diameter and consists of six polypeptide chains; two α -chains (blue), two β -chains (green), and two γ -chains (red). (B) The monomers assemble in a half-staggered fashion to form the two-stranded protofibrils (C); the A: a interactions are shown as yellow lines. (D) The protofibrils aggregate laterally to form thicker fibers, branching occurs and eventually a full clot (E) is formed in which fiber diameters range

from about 20 to 200 nm (E) shows an SEM image of a 50% lysed thrombus containing fibrin fibers and platelets; courtesy R. R. Hantgan; reproduced with permission from [8]; scale bar: 10 μ m). (F) A striated pattern with a periodicity of 22.5 nm (half the length of a fibrin monomer) is seen in the TEM image of a longitudinal cross section of a single fiber (Image: W. Liu, scale bar 200 nm). The striated pattern is due to the half-staggered arrangement of the monomers

residues γ Gln329, γ Asp330, γ His340, and γ Asp364 [5, 15, 16]. Pocket “b” in β C is composed of key residues β Glu397, β Asp398, and β Asp432 [17]. Fibrinopeptide A is cleaved first and upon the exposure of knob A, fibrin monomers assemble in a half-staggered fashion into two-stranded protofibrils [18]. This assembly also aligns the D:D interface of abutting fibrin monomers. The key D:D interfacial residues include γ Arg275, γ Tyr280, and γ Ser300 [7]. The assembly is illustrated in Fig. 1B, which depicts a fibrin trimer, i.e., a nascent protofibril. Upon growing to an average length of about 15 monomers [18, 19], approximately twice the length of the protofibril shown in Fig. 1C, the protofibrils aggregate laterally to form fibers (Fig. 1D) that branch into a three-dimensional network (Fig. 1E). Mature fibers have a diameter of about 100 nm [20], i.e., about 150 monomers per cross section (assuming 30% protein content in a fiber [21]). Release of fibrinopeptide B (exposure of knob B) occurs subsequent to release of fibrinopeptide A. The B:b interaction may aid lateral

aggregation; however, it is not required for lateral aggregation and is likely not a strong contributor to fiber stability. This statement is based on the observation that clotting reactions catalyzed with various snake venom enzymes, which only cleave Fibrinopeptide A (i.e., Batroxobin), produce mature and stable fibers [22].

Covalent Crosslinking

Fibrin fibers are stabilized by the formation of covalent crosslinks between specific glutamine and lysine residues (ϵ -amino(γ -glutamyl)lysine isopeptide bonds) [23]. The formation of these bonds is catalyzed by the transglutaminase factor XIIIa (FXIIIa). Initially, and concomitantly with protofibril formation, two reciprocal crosslinks are formed between the γ -chains of the fibrin monomers *within* a protofibril at residues Lys406 and Gln398/399, forming γ chain dimers. At about ten times slower rate than the γ - γ

crosslink formation, Lys-Gln crosslinks are also formed between several sites in the α C domains, forming α -chain oligomers [24]. The main potential acceptor and donor sites within the α -chain may include Lys 418, 448, 508, 539, 556, 580, 601 and Gln 221, 237, 328, 366 or a subset of those [25–27]. The α - α crosslinks may form between fibrin units of different protofibrils, i.e., the protofibrils are bound together within fibrin bundles by α chain crosslinking. There is also some crosslinking between α - and γ -chains; these crosslinks may occur internally within one fibrin molecule [28], such that they will not contribute much to fiber stability.

Although establishing exactly how fibrin molecules pack together within a fiber is still an active area of research, the major interaction sites have been determined. It is becoming clear that the main non-covalent stabilizing inter-fibrin interactions are the D:D interface interaction, the A:a interaction, and to a lesser extent the B:b interaction; the main covalent interactions are the γ - γ crosslinks and perhaps to a lesser extent the polymeric α -crosslinks. The putative D:D interface does not exhibit any specific hydrogen bonds or salt bridges; the apparent driving force is the burying of 7.5 nm² of surface area [5]. Thus, this interaction may not be very strong and the stability of fibrin fibers may mainly depend on the A:a interaction that supports protofibril formation and a large number of weak interactions, including D:D, that support the assembly of protofibrils into fibers [19]. The covalent cross-links, of course, greatly stabilize the noncovalent interactions. It is interesting to note that, except for the weak, hydrophobic D:D interaction, all other interactions involve flexible and mobile regions whose structures have not been discerned in any crystal [29]. This mobility may be functionally important. The A-knobs are attached to a mobile 10 residue-long tether (α 17–27); the B-knobs are attached to a mobile 48 residue-long tether (β 15–63). The reciprocal γ - γ links between Lys406 and Gln398/399 combine the two 19 residue-long, mobile C- γ -terminal domains which results in the formation of a 21 residue-long flexible γ - γ crosslink. The α - α crosslinks occur between the large, highly mobile α C domains.

It is not yet entirely clear how protofibrils aggregate laterally, though some putative interfaces on the γ -chains have been inferred from protein contacts within crystals, having given rise to the (speculative) multibundle model of fibrin packing within fibers [30]. In this model, fibrin molecules arrange longitudinally in the well-established, half-staggered fashion to form protofibrils, as outlined above. The two strands of a protofibril are 8 nm apart (Fig. 1B, C). It is suggested that the protofibrils then arrange in a complex, four-tiered, half-staggered fashion in which the tiers are apart by 4.75 nm, roughly the diameter of a protofibril (Fig. 1D). Such an arrangement would result in a

“pseudo-unit cell” with dimensions of 19 nm \times 19 nm \times 45 nm containing eight fibrin molecules. The longitudinal and lateral distances of this suggested unit cell are consistent with energy dispersive X-ray diffraction techniques [31], giving credence to this model. Nevertheless, the data show real fibrin fibers will not assemble in such a uniform, *perfectly* crystalline fashion, as uniform crystals would give extremely sharp peaks, rather than the much broader peaks observed in the X-ray studies. Moreover, branching would become difficult if the fibers assembled in perfectly crystalline fashion.

Electron microscopy images of negatively stained fibrin fibers show a striated pattern with a periodicity of 22.5 nm (half the length of a monomer) along the fiber axis (Fig. 1F; [3, 4]). This observation clearly supports the model of fibrin monomers assembling in a regular, half-staggered, nearly crystalline fashion.

Mechanical Measurements on Single Fibrin Fibers and Fibrinogen Molecules

Before reviewing recent measurements of the mechanical properties of fibrin fibers and their building block, fibrinogen, we will define some of the terminology of force measurements. When a force, F , is applied longitudinally to a uniform, isotropic fiber with cross-sectional area, A , and length, L , it will stretch by a distance ΔL . In an *elastic* deformation, the fiber will return to its original length upon the release of force. In a *linearly elastic* deformation, the strain $\varepsilon = \Delta L/L$ is proportional to the stress, σ (force per unit area; F/A): $\sigma = E \cdot \varepsilon$, where E is the *Young's modulus* (Fig. 2). The Young's modulus is a measure of fiber *stiffness*; the higher the Young's modulus, the stiffer the fiber. If a material is *non-linearly elastic*, the elastic modulus (slope of stress-strain curve) changes as a function of strain. The material may become stiffer (strain-hardening) or softer (strain-softening) with increasing strain and one may define the modulus either as the secant modulus ((total stress)/(total strain)) or the tangent modulus ((differential stress)/(differential strain)).

The *extensibility* (also called *breaking strain* or *strain at rupture*) of a fiber is defined as the strain, ε_{\max} , at which it will rupture. The *elastic limit* is the largest strain a fiber can sustain and still return to its original length upon the release of the applied force. This property is important, because fibers can recover without any permanent deformations from strains below the elastic limit, whereas strains above this limit cause permanent damage in the fiber.

Using an Atomic Force Microscope (AFM), Guthold et al. performed the first direct force measurements on single fibrin fibers as they determined the radius-dependence of the rupture force (breaking strength or ultimate

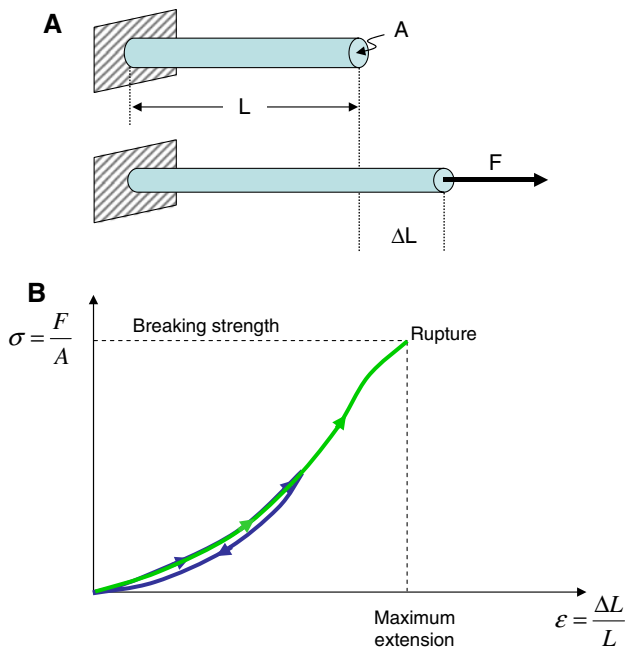


Fig. 2 Stress–strain curves of stretched fibers. **(A)** A force F is applied in the longitudinal direction to a fiber with length L and cross sectional area A . The fiber extends by an amount ΔL . **(B)** A schematic stress–strain curve of the stretching of a fiber. The slope of the curve corresponds to the stiffness of the fiber. In a linear, elastic model, the stiffness (slope) is called Young’s modulus of the material. The maximum extension at which the fiber ruptures is called *breaking strain* (or *extensibility*) of the fiber

strength) of fibrin fibers [32]. They found that the rupture force increases as $R^{1.3}$, where R is the radius of the fibers. This was unexpected, since the rupture force would increase as R^2 for a uniform cylindrical fiber with a solid cross section. This finding suggests that the density of fibrin fibers may be proportional to $R^{-0.7}$ (not uniform), i.e., the fibers may be denser in the center than on the periphery. These force measurements were complicated by the fact that the fibers were adsorbed onto mica, so friction with the surface added resistance. Moreover, these measurements were done on dried fibers, whereas under physiological conditions fibrin fibers contain 70–80% aqueous solvent [21].

Subsequently, Collet et al. used laser tweezers to determine the *Young’s modulus* of fibrin fibers in buffered saline analogous to plasma [2]. For small strains they found that uncrosslinked and crosslinked fibrin fibers have a Young’s modulus of 1.7 and 14.5 MPa, respectively. These values are of the same order of magnitude as those for other soft biological fibers; however, they are about a 1,000 times less than those of stiff biological fibers (Table 1). Thus, the data at small strains indicate fibrin fibers are “soft”.

Recently Liu et al. used a combined atomic force/fluorescence microscopy instrument to measure the breaking strain (extensibility) and elastic limit of

crosslinked and uncrosslinked, single fibrin fibers formed at low fibrinogen concentrations in buffered saline [1] (γ – γ crosslinking was 50–5% complete and α –crosslinking was 25–5% complete). In these experiments, fluorescently labeled fibrin fibers were suspended over 12- μm wide groves in a striated substrate. The AFM tip was used to stretch out these fibers, while the stretching process was observed from underneath with the fluorescence microscope (Fig. 3). These studies showed fibrin fibers have extremely large *extensibilities* and are able to recover *elastically* from very large strains.

Figure 4A–F shows movie frames of an experiment that measured *breaking strain* (extensibility). The AFM tip was brought in contact with the fiber and moved to the right, parallel to the ridges, until the fiber ruptured. The fiber did not break at the point where the AFM tip contacted the fiber, but instead the two segments stretching from the tip to the two anchoring points broke independently. This observation indicates that the fiber did not break due to bending stress at the tip, but due to stretching between the tip and the anchoring point. After rupture, the fiber arms contract to nearly their original length (Fig. 4F), indicating that the deformations were mainly elastic. Thus, significant elasticity was retained even after the large extension and ultimate rupture. The distribution of extensibilities and the average extensibilities of the different fiber types are shown in Fig. 4G, H. Uncrosslinked fibers formed with thrombin and batroxobin showed a breaking strain of $226 \pm 52\%$ and $226 \pm 72\%$, respectively (average \pm standard deviation). Cross-linked fibers formed with thrombin or batroxobin showed a breaking strain of $332 \pm 71\%$, and $265 \pm 83\%$, respectively. A breaking strain (extensibility) of 332% corresponds to a fiber that is stretched to 4.32 times its original length as is schematically depicted in Fig. 4H.

The *elastic limit* of fibrin fibers was tested by stretching them to a certain strain and releasing the applied force. In the experiment shown in Fig. 5, both elastic fiber deformation and deformation that induced permanent damage to the fiber were seen. First, the fiber was strained 80% after which it returned to its *original* length with no discernable permanent lengthening (Fig. 5A–D). The fiber was then strained 230% (Fig. 5E–G). When it snapped back under the tip, the fiber showed permanent damage with an increased length (Fig. 5G). A quantitative analysis (Fig. 5H) showed that the elastic limit of uncrosslinked fibers (batroxobin and thrombin) and crosslinked batroxobin fibers was at strains of about 120%; that is, these fibers were stretched to 2.2 times their original length and recovered completely elastically. Remarkably, crosslinked thrombin fibers were stretched to over 2.8 times their length (180% strain) and still recovered without any discernable permanent damage.

Table 1 lists the stiffnesses (Young's modulus, E); breaking strain (extensibility ϵ_{\max}); and the proposed molecular mechanisms of extensibility for various protein fibers

Material	E (MPa)	ϵ_{\max} (%)	Proposed extensibility model	References
<i>High extensibility, soft fibers</i>				
Crosslinked fibrin fibers	1–10	332	$\alpha \leftrightarrow \beta$ transition, domain unfolding, protofibril sliding	[1, 2, 33]
Uncrosslinked fibrin fibers	1–10	226	$\alpha \leftrightarrow \beta$ transition, domain unfolding, protofibril sliding	[1, 2, 33]
Spider silk (Araneus Flag silk)	3	270	Modular composition; large extensible amorphous or β -spiral regions connected by stiff crystallites	[34–39]
Elastin (Bovine ligament)	1	150	Compact, amorphous, hydrophobic domains (random or β -spiral), which are crosslinked together, entropic, rubber-like elasticity	[40–46]
Resilin (Dragonfly tendon) ^a	1–2	190 ^a	Compact, amorphous, hydrophobic domains, which are crosslinked together, entropic, rubber-like elasticity	[40, 46, 47] ^a
Cloned resilin ^b		313 ^b		Cloned [48] ^b
Matrix-free Intermediate filament (mammalian ^c , hagfish ^d)	6–300	160 ^c –220 ^d	$\alpha \leftrightarrow \beta$ transition and/or fibril sliding	[49–53]
Fibrillin	0.2–100	> 185	Unstacking of pleated domains	[54–58]
Myofibrils (sarcomere) Titin (Connectin)	1	200	Unfolding of PEVK and Immunoglobulin domains in Titin	[59–61]
Mussel byssus (distal)	10–500	109	Block copolymer: collagen-like central domain (stiff) flanked by crystalline β -sheets (stiff) and amorphous or β -spiral motifs (elastic)	[62–64]
Fibronectin	–	200–300	Extension of bent and looped molecules and/or domain unfolding	[65–67] [68]
<i>Low extensibility, stiff fibers</i>				
Spider silk (Araneus MA silk)	10, 000	27	Modular composition; small, extensible amorphous or β spiral regions, connected by stiff crystallites	[34–39]
Uncrosslinked, self-assembled collagen-I	–	24–68	Sliding of collagen fibrils	[69, 70]
Cross-linked, self-assembled collagen	5,000–7,500	12–16	Highly regular, paracrystalline structure, small, reversible molecular deformations	[69, 70]
Tendon collagen (mammalian tendon)	160–7,500	12	Highly regular, paracrystalline structure, small, reversible molecular deformations	[71–74]
Actin	1,800–2,500	≤ 15	Highly regular, paracrystalline structure	[75–80]
Microtubules	1,000–1,500	≤ 20	Highly regular, paracrystalline structure	[76, 81–84]
Wet, hard α -keratin in high-sulfur matrix (hair, wool)	2,000	45	$\alpha \leftrightarrow \beta$ transition of keratin (embedded in crosslinked, high-sulfur matrix)	[50, 85–90]

There are two reasons for the range of values in the Young's modulus of a given fiber: (1) Authors report a range of values. (2) Strain hardening/softening; the fibers become stiffer/softer as the strain increases. The table includes microscopic fibers (diameter: order of micrometer or less) and macroscopic fibers (diameter: order of several micrometers or larger). Microscopic fibers are fibrin fibers, elastin fibers, resilin fibers, intermediate filaments, fibrillin, myofibrils, fibronectin, actin filaments, microtubules; macroscopic fibers are spider silk, mussel byssus, collagen fibers, wool fibers

^a Dragonfly tendon

^b Cloned resilin

^c Mammalian, matrix free intermediate filament

^d Hagfish threads, a model fiber for matrix-free intermediate filament

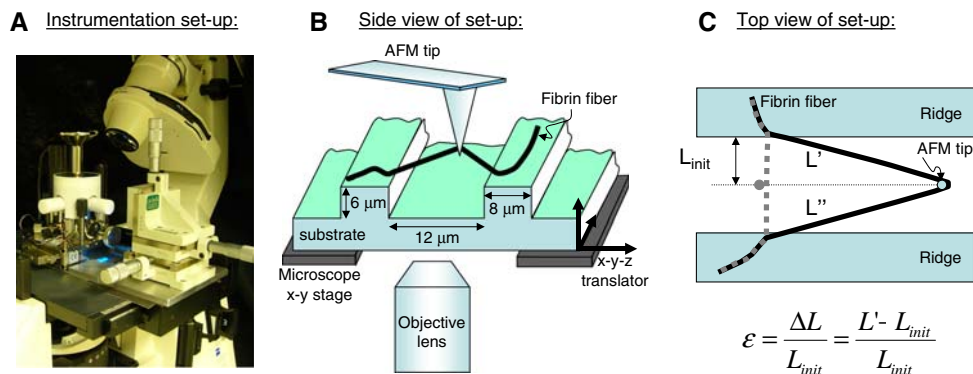
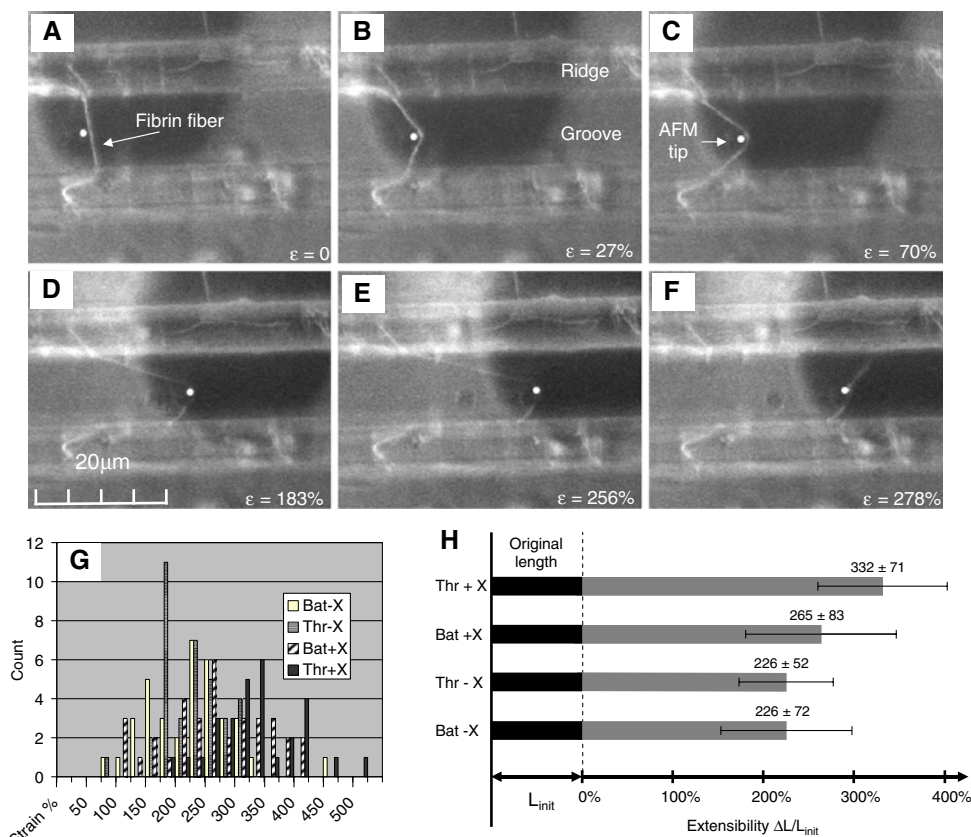


Fig. 3 Experimental set-up to determine fibrin fiber breaking strain (extensibility) (Figure adopted from [1] with permission). The experimental set-up to stretch single fibrin fibers is depicted photographically in Fig. 2A and schematically in Fig. 2B. The atomic force microscope (AFM) was used to stretch fibers that were suspended across ~12 μm-wide channels; the fluorescence microscope was used to record movies of this stretching process. This set-up has several advantageous features for stretching out fibers. Somewhat fortuitously, we found that the fibers were very well anchored on the ridges of the striated substrate as the anchoring points

rarely changed; even at the most extreme fiber extensions. This implies that the observed deformations are *not* due to fiber slipping on the ridges. We selected fibers that bridged the grooves at a right angle with respect to the ridge edge. Thus, our set-up yielded a well-defined, easy-to-analyze geometry. By suspending the fibers over grooves, the substrate did not interfere with the measurement. Being able to record movies of the stretching process allowed us to accurately determine the lengths of the fibers and ascertain that the fibers did not slip at the anchoring points. The technique may also be applied to other fibers

Fig. 4 Extensibility of fibrin fibers (adopted from [1] with permission and modified). (A–F) Snapshots from a fluorescence microscopy movie of a fiber extensibility experiment (crosslinked batroxobin fiber). A fluorescently labeled fiber is suspended between two ridges that appear as two horizontal bands. The AFM cantilever is seen from underneath as a dark, ~35 μm-wide rectangle; the location of the AFM tip is indicated by a white dot. The AFM tip is moved to the right, thus stretching the fiber; the lower segment of the fiber breaks at 183% strain and the upper segment of the fiber breaks at 278%. Histogram (G) and bar graph (H) representations of the extensibilities of the four different types of fibrin fibers; Thr = thrombin, Bat = Batroxobin, X = crosslinking



Force-Induced Fibrinogen Unfolding

In an very recent study, Brown et al. investigated the force-induced unfolding of single fibrinogen molecules [33].

When pulling on short oligomers of γ - γ crosslinked fibrinogen molecules with an AFM tip, they observed the typical sawtooth pattern of unfolding proteins. The observed peak-to-peak distance of 23 nm between

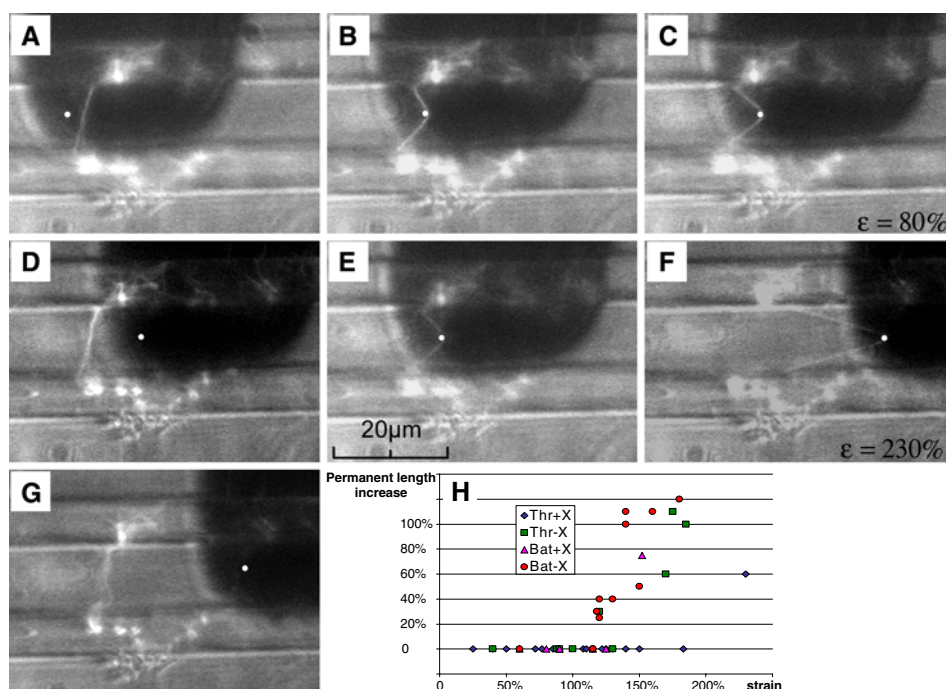


Fig. 5 Elastic limit of fibrin fibers (adopted from [1] with permission and modified). (A–D) A fibrin fiber (thrombin, crosslinked) was strained 80% (C), from which it snapped back to its original length (D) without any permanent lengthening. (E–G) The same fiber was strained to 230% (F); at this strain it did suffer some permanent damage as the fiber is significantly deformed in image (G). (H) Plot

of the amount of permanent deformations (% length increase) upon the release of force versus strain. Data points along the horizontal axis of 0%-permanent length increase indicate elastic deformations. Crosslinked thrombin fibers could be strained over 180% and the other fiber types about 120% without suffering permanent lengthening. None of the fibers analyzed here were ruptured

unfolding events suggests that application of about 100 pN of force caused the unfolding of the 17-nm long α -helical coiled coils into 40-nm long extended β -strands. Since each fibrinogen molecule has two α -helical coiled coils (Fig. 1A) such a α -helix to β -strand conversion would extend the molecule by $2 \times 23 \text{ nm} = 46 \text{ nm}$. This lengthening corresponds to a strain of about 100%, since the initial length of a fibrinogen molecule is 46 nm.

Stiffness (Young's Modulus) and Breaking Strain (Extensibility) of Fibrin Fibers and Other Polymerized Protein Fibers

Table 1 lists the stiffness (Young's modulus, E), maximum breaking strain (extensibility, ϵ_{max}) and putative extensibility mechanism of many important, polymerized protein fibers. Following the table we summarize the major, general features of the fibers listed in Table 1. An overview of the table reveals that high extensibility fibers ($\epsilon_{\text{max}} > 100\%$) have a lower Young's modulus, usually on the order of 1–10 MPa. Conversely, low extensibility fibers are usually much stiffer with a Young's modulus on the order of 1,000–10,000 MPa. By way of contrast to well-known man-made materials, elastic rubber bands have a Young's

modulus of about 1–10 MPa and nylon fishing line of about 1,000–10,000 MPa.

Despite the fact that detailed structural information is missing for most high extensibility fibers (many contain unstructured or high mobility motifs which hamper crystallization), there is reasonable experimental evidence to support a specific extension mechanism. The following mechanisms, correlated with each protein's structural features, have been proposed for high extensibility fibers: (i) Extension of amorphous, unstructured or β -spiral regions (spider silk, elastin, resilin); (ii) α -helix to β -strand transition (intermediate filament (keratin), hair (keratin), fibrin fibers) (iii) unstacking of intramolecular pleats (fibrillin fibers); (iv) unfolding of immunoglobulin or other protein domains (myofibrils (titin in sacromers), fibronectin); (v) straightening of bent or looped molecules (fibronectin); (vi) sliding of fibrils past each other (uncrosslinked collagen). Among these fibers, several contain coiled coil motifs [91]. For example, fibrinogen and the intermediate filament proteins, keratin and vimentin, and hair fibers (matrix-embedded keratin) have a long α -helical coiled coil, which may convert to an extended β -strand conformation upon the application of force.

The common theme for most *low* extensibility fibers (actin, microtubules, collagen) is a highly regular, nearly

crystalline arrangement of monomer units without regions that can extend, change secondary structure or unfold.

At this point we should also mention that in the text below ‘crosslinks’ usually though refers to covalent bonds; the exception is spider silk in which ‘crosslinks’ refer to bonds formed by crystallites.

High Extensibility Fibers

Fibrin Fibers

Fibrin fibers are very stretchable with an average breaking strain (extensibility) of 332% and 226% for crosslinked and uncrosslinked fibers, respectively (1). To our knowledge, crosslinked fibrin fibers display the largest extensibility of all protein fibers. They are also rather soft, with a Young’s modulus of 1–10 MPa (2). This is unusual, because the fibrin monomers within fibrin fibers are arranged in a relatively regular, half-staggered, almost crystalline pattern along the fiber axes [18, 19, 30], Fig. 1). As will be outlined in Section “Proposed Molecular Mechanisms for Fibrin Fibers Extension,” where we propose molecular mechanisms, these seemingly conflicting properties may be partly reconciled by assuming that the fibrin *monomers* within a fiber are very extensible *while* at the same time maintaining the most important fibrin–fibrin interactions.

Spider Silk

Of the many different types of spider silk fibers, the most relevant ones for examining extensibility and stiffness are those excreted by the major ampullate (MA) gland and the flagelliform (*Flag*) gland of *araneoidea* (web-spinning spiders) [34]. MA fibers, which form the frame and spokes of the web, are much stiffer and less extensible than *Flag* fibers, which form the catching threads spiraling around the frame. MA fibers have a maximum breaking strain of 27% and a Young’s modulus of 10,000 MPa. *Flag* fibers have a maximum breaking strain of 270% and an initial (low strain) Young’s modulus of 3 MPa [35, 36].

It is thought that the mechanical properties of these fibers originate from the *modular* composition of the silk fibroin proteins comprising each fiber. The fibroin proteins are composed of two main domains: (i) poly-alanine repeats, which form crystallites of β -pleated sheets, and (ii) glycine-rich domains which form amorphous domains. The crystallites may act as “cross-links” between the amorphous regions, which can unravel and extend upon the application of stress [37–39]. The proportion of the fibroin protein devoted to each structural module affects the fibers’ mechanical behavior. Crystalline regions confer tensile strength and stiffness; and the amorphous regions provide

high extensibility. The stiff and less extensible MA fibers have a high proportion of crystallite domains, whereas the *Flag* fibers contain few crystallites and consist mainly of large amorphous domains [39].

Elastin Fibers

Elastin fibers are found in the extracellular matrix of many mammalian tissues, such as skin, lung, and large blood vessels (reviews [44–46]). Tropoelastin, the precursor of elastin is deposited into the extracellular matrix, where it is cross-linked at its lysine residues, thus rapidly forming elastic fibers with the help of several scaffolding microfibril proteins, such as fibrillin. Elastin fibers have a Young’s modulus of 1 MPa and a large breaking strain (extensibility) of 150% [41].

Due to the extensive cross-linking, elastin is nearly insoluble, and thus, structural information for elastin fibers remains limited. Elastin probably consists of two alternating domains: (i) a hydrophilic domain, rich in lysine and alanine, which forms cross-links; (ii) a hydrophobic domain, rich in glycine, proline, and valine, which forms repeats of poly (VPGVG) and poly (VGGVG) and endows the fiber with its elasticity and large extensibility. The structure of the hydrophobic domain may be a compact amorphous structure of random chains (similar to rubber) [42] or β -spirals [43] (review [44]). Both the models (random chain or β -spiral) could explain the observation that elastin fibers display entropic elasticity, similar to a typical rubber.

Resilin Fibers

Resilin is a protein found in the cuticles of many insects. It forms fibers that display elastic properties similar to elastin fibers [40, 46, 47]. Recently, Elvin et al. expressed resilin from *Drosophila melanogaster* in *E. coli* and formed resilin fibers with the purified recombinant protein [48]. These fibers have a breaking strain of up to 313% and a Young’s modulus of 1–2 MPa.

Though there is still ongoing debate on the structural basis for the elastic properties of these fibers, there seems to be consensus that spider silk, elastin, and resilin fibers consist of an amorphous, hydrophobic region of random chains or β -spirals, which can unravel, and an ordered region that links these amorphous regions together.

Intermediate Filament

Intermediate filaments are one of the three cytoskeletal fibers (in addition to actin filaments and microtubules) and they are prominent in cells that are exposed to mechanical

stresses. A number of proteins, including keratin, vimentin and desmin, polymerize into intermediate filaments. These proteins exhibit a characteristic “tripartite” structure consisting of a central elongated, α -helical domain, flanked by a globular head (N-terminal), and a tail (C-terminal) domain [92]. Monomers twist around each other to form an about 45 nm long and 2–3 nm wide dimer such that the two α -helices form a coiled coil. The dimers assemble, in a not yet fully understood geometry, into the 8- to 12-nm wide intermediate filament. These filaments then form bundles. Kreplak and Fudge recently wrote an excellent review about the mechanical properties of intermediate filaments [53].

Fudge and Gosline have measured the breaking strain of hagfish threads to be 220%. These fibers can be considered bundles of *matrix-free* intermediate filaments [49] (as opposed to matrix-embedded keratin-fibers, see below). These authors also showed that the Young’s modulus of these filaments is 6 MPa for small strains; when stretched further, the fibers become stiffer (strain hardening) displaying an instantaneous stiffness (or tangent modulus = slope of the stress–strain curve) of 300 MPa for strains over 100% [49]. The lower initial value (6 MPa) is consistent with the persistence length of 1 μm that was determined for single intermediate filaments from AFM images [52]. The persistence length, P , and Young’s modulus, E , are related for isotropic materials; $E \cdot I = k_B \cdot T \cdot P$, where I is the areal moment of inertia, k_B is the Boltzmann constant and T is the temperature [93]. Assuming the filament is cylindrical such that $I = 0.25 \cdot \pi \cdot R^4$, $E \sim 8$ MPa for a filament with persistence length of 1 μm and radius $R = 5$ nm.

The extensibilities of three different types of intermediate filaments (desmin, keratin, and neurofilaments) were determined by Kreplak et al. [51], by rupturing these filaments laterally with an AFM tip. They found an average breaking strain of 160%, and a maximum value of 260%. Guzman et al. used an AFM tip to bend single vimentin filaments, which were suspended over holes in a substrate. They determined a bending modulus of 300 MPa for uncrosslinked vimentin fibers [94].

Fudge and Gosline proposed that the large extension in intermediate filaments is made possible by a transition of the central, α -helical coiled coil rod to an elongated β -strand structure [50].

Fibrillin fibers

Fibrillin fibers are widely distributed in elastin-containing tissues, such as blood vessels, lung, and skin, where they act as a lattice for elastin deposition during elastin fiber formation. Fibrillin-rich fibrils also occur in tissue that does not contain elastin, such as the ciliary zonules, where they hold the lens of the eye in dynamic suspension [95]. Fibrillin

fibrils resemble beads-on-a-string with a periodicity of 56 nm and a diameter of 10–14 nm. They can be extended to a periodicity of at least 160 nm, which amounts to a strain of at least 186% [54]. Using molecular combing techniques, Sherratt et al. found a Young’s modulus of 78–98 MPa for fibrillin-rich microfibrils (zonular filaments) [56]. Megill et al. report a Young’s modulus of 0.9 MPa for fibrillin-containing fibers in the mesoglea of the hydromedusa *Polychorda penicillatus*. Megill et al. argue that the stress–strain curve for fibrillin, similar to hagfish threads, is J-shaped with an initial low value, in the 1 MPa range, at low strains and a much higher value, up to 100 MPa, at higher strains (strain hardening) [57]. Values of 0.19–1.88 MPa were also obtained for the zonular filaments from cows [58].

Though fiber assembly is still poorly understood, the model that best accounts for the observed data is the intramolecular pleating model [54, 55]. Each fibrillin molecule consists of rod-like regions that fold upon themselves, forming pleats. Upon stretching the pleats become undone.

Myofibrils

Myofibrils, which consist of concatenated structures called sarcomeres, are the stretchable fibers of striated muscle. Myofibrils are a highly ordered assembly of three myofilaments: the thick filament, which is mainly myosin, the thin filament, which is mainly actin, and titin, also called connectin. Sarcomeres have a large, 200% breaking strain [53]. A Young’s modulus of about 1 MPa has been determined for myofibrils from the *Drosophila* indirect flight muscle; this report argues that other muscles should have similar stiffness [61]. Titin is the myofilament that is mainly responsible for the large extensibility of myofibrils in relaxed muscle fibers. Titin is a giant, single-peptide-chain protein (~ 0.6 – 3.7 MDa, depending on isoform) that stretches half the length of the sarcomere (for excellent review see [96]). Titin consists of four different regions (Z-disc region, I-band, A-band, M-line region), but only the I-band, consisting of immunoglobulin repeats and the PEVK region (rich in proline, glutamate, valine, and lysine) is functionally extensible. Extension of the sarcomere first causes straightening of the compact, relaxed I-band section of titin. Further extension induces unfolding of the PEVK region and the immunoglobulin domains of titin. Titin refolds upon the release of force [60].

Mussel Byssal Fibers

Mussels are tethered to solid supports by a byssus composed of collagen-like threads. The byssal threads have

three serially arranged parts: a corrugated proximal segment, a smooth distal segment and an adhesive plaque. The breaking strain of the distal and proximal segment are 109% and 200%, respectively [62]. The Young's modulus for byssal threads ranges from 10 to 500 MPa [64]. The exact secondary structure of byssal fibers is not well known, but it appears to be a block copolymer consisting of a central collagen-like domain flanked by more elastic regions [63]. Some of the flanking domains have sequence motifs similar to the crystalline β -sheets and amorphous domains in spider silk and the β -spiral motifs of elastin.

Fibronectin

Fibronectin fibers are found in embryonic tissue, in healing wounds and in the extracellular matrix. Fibronectin fibers can be stretched to a strain of 200–300% as they are being assembled on the cell surface [66, 97]. The Young's modulus of fibronectin fibers has not been determined and it is also unknown how fibronectin molecules are arranged to comprise a fibril. Agreement on the stretching mechanism has not yet been achieved. There are models proposing that fibronectin domains may unfold upon stretching [68, 98]; these models are based on FRET experiments [68] and NMR and steered molecular dynamics simulations [98]. However, there are also models proposing that fibronectin may not unfold upon stretching [65, 67]. In those models fibronectin molecules are bent and looped into a compact conformation. Stretching then pulls the molecules into an extended conformation, but protein domains would remain folded.

Low Extensibility Fibers

Collagen

There are over 20 different types of collagen, some of which form fibrils (e.g., types I, II, III, V, and X) that provide the structural framework of many tissues. These fibrils are the major stress-bearing component in connective tissues such as bone, teeth, cartilage, tendon, ligament and the fibrous matrices of skin and blood vessels. Given the many different types and functions of collagen fibers, their properties are varied, though none are very extensible. All the fibril-forming collagens self-assemble into cross-striated fibrils with a characteristic 67 nm repeat [70]. The major structural feature in these collagens is an about 300 nm long, right-handed "collagen triple helix" consisting of three left-handed proline helices (*not* α -helices) wrapped around each other. The collagen triple helix should not be confused with the coiled coils consisting of α -helices. The

collagen triple helix is designed to *not* be very extensible as the rise per residue is 0.29 nm, vs. 0.34 nm for a β -strand. The rise per residue in an α -helix is 0.15 nm.

Type-I collagen molecules self-assemble into quarter-staggered arrays of fibrils 20 to several hundred nanometers in diameter. Bundles of fibrils pack into collagen fibers and higher order tissue structures.

The breaking strain (extensibility) of self-assembled, cross-linked and uncrosslinked type-I collagen (tendon is predominantly composed of crosslinked type I collagen) was found to be 10% and 50%, respectively [69, 70]. The authors attribute the larger extensibility of uncrosslinked collagen to sliding of collagen fibrils past each other. The total stretch modulus (including viscous components) of type-I collagen fibers is on the order of 300–500 MPa, as determined for both mineralized turkey tendons and for 6-month aged, cross-linked, self-assembled type-I collagen fibers. The Young's modulus (elastic part of the total stretch modulus) is on the order of 250–400 MPa for these fibers [71]. This study employed incremental stress–strain curves to separate the elastic and viscous components of the total stretch modulus [71]. These numbers were then corrected for the facts that fibrils slide within a fiber and that fibers are porous and a Young's modulus of 5,000–7,500 MPa was obtained for "solid" collagen fibers that do not slide. Values of 160 and 1,660 MPa, and 2,900 MPa were reported for newborn and mature pig digital tendons [72] and for bovine Achilles tendon [73], respectively.

Actin

Soluble G-actin polymerizes into helical actin filaments (F-actin). Actin- filaments have a variety of roles in cells that include forming a scaffold to maintain cell shape, being part of the mechanism for cell motility, and serving as "rails" for myosin in muscle contraction (the thin filaments of muscle (see myofibrils above) are composed of filamentous actin). Actin often interacts with a number of other structural proteins resulting in the formation of fibers with somewhat different properties. The breaking strain of actin filaments is small, 15% or less (Figs. 5 and 6 in [77], [78, 79]). The Young's modulus of actin filaments is on the order of 1,800–2,500 MPa [75–77, 80].

Microtubules

Microtubules are *hollow* fibers whose cylindrical shells have a diameter of 25 nm. They provide the framework that organizes the cell and guides the movement of organelles within cells. They are formed of alternating, noncovalently joined helices of globular α - and β -tubulin with 13

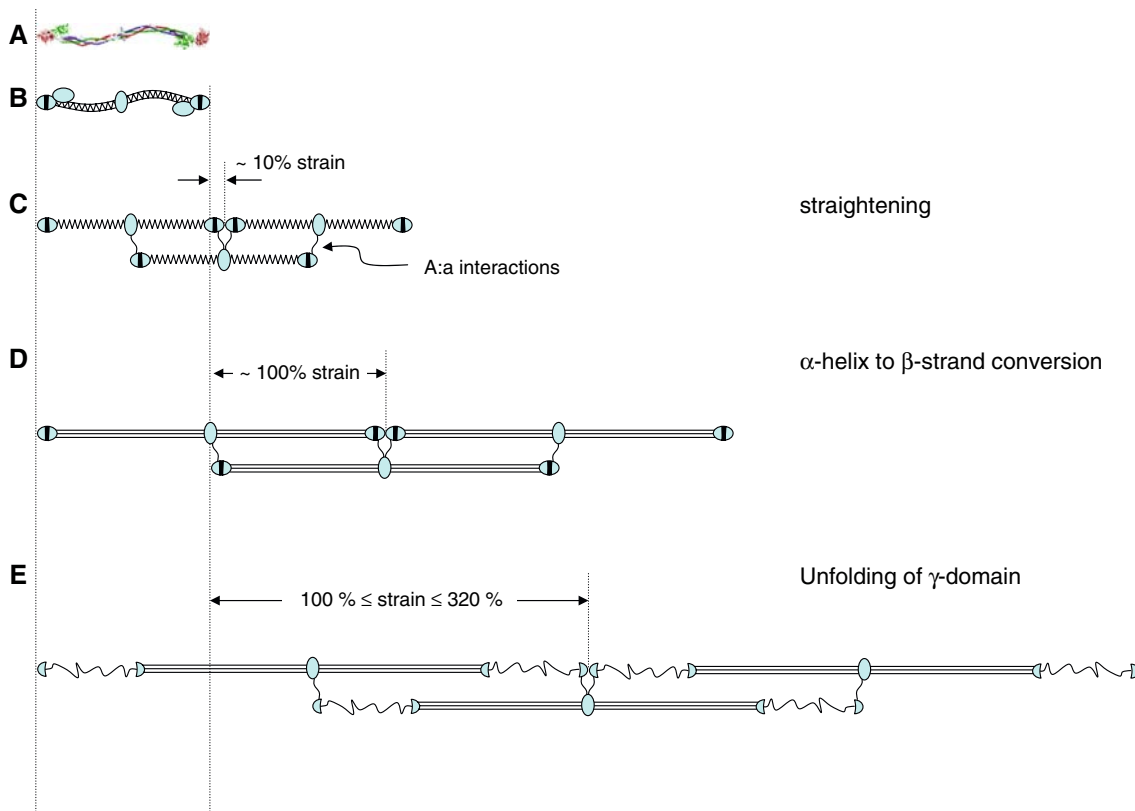


Fig. 6 Schematic molecular mechanism models for fibrin fiber extensions. **(A)** Crystal structure and **(B)** schematic model of fibrinogen. **(C)** Half-staggered assembly of three fibrin monomers; molecules are rotated 90° from view in **(B)**; β C domain is no longer depicted; straightening and slightly stretching the molecule may

accommodate 10% strain. **(D)** An α -helix to β -strand conversion of the coiled coil and a slight straightening and alignment of the molecules could accommodate ~100% strain. **(E)** Higher strains, up to 320% could be accommodated by a partial unfolding of the γ C domains; 230% strain is depicted

staggered α - β tubulin heterodimers per turn. Microtubules have a very regular, paracrystalline structure, and a breaking strain of less than 20% [81] and a Young's modulus of 1,000–1,500 MPa [76, 82, 83] as determined from the flexural rigidity of microtubules and in the 100 MPa range or higher as determined by bending suspended microtubules with an AFM tip [84].

Hard α -keratin

Hard α -keratin is a tough *composite* material that forms structures such as hair, hooves, and claws in mammals. The composite consists of keratin microfibrils, (very similar in structure to the intermediate filament), *embedded* in a sulfur matrix. The breaking strain of hard, wet α -keratin fibers, such as hair and wool, is about 45% and their Young's modulus is about 2,000 MPa [86]. It has long ago been suggested [87, 88] and experimentally verified [89, 90], that these fibers undergo an α -helix to β -strand transition, when stretched. The increased stiffness and decreased extensibility of hard α -keratin fibers as compared

to intermediate (keratin) filaments is mainly due to the matrix.

Proposed Molecular Mechanisms for Fibrin Fibers Extension

The molecular mechanisms that allow fibrin fibers to accommodate large and reversible strains are unknown. Nevertheless, utilizing the crystal structure of native chicken fibrinogen [7], studies of the other proteins listed in Table 1, and the single molecule stretching experiments by Brown et al. [33], it is possible to suggest several likely mechanisms (Fig. 6). These mechanisms are not mutually exclusive, such that they may all contribute to the overall mechanical properties.

1. Small strains of about 10% may be accommodated by small, reversible structural changes in the fibrin monomer, protofibril and protofibril arrangement within a fiber. Fibrin monomers may straighten their slight S-shape (Fig. 6C). Protofibrils are shaped like a wavy ribbon [30] and the major interactions which hold the two strands of a

protofibrils together utilize flexible linkers [29]. Upon stretching, the wavy shape of the protofibrils may straighten and the flexible linkers may align along the stretch direction, increasing the distance between monomer units. Though it is still not known how protofibrils aggregate laterally to form fibrin fibers, the open structure (fibrin fibers contain 70–80% aqueous solvent [21]) suggests that the interactions between protofibrils are mediated through long linkers. Upon stretching, those linkers could also align along the stretch direction.

2. At larger strains, the two 17 nm α -helical coiled coils may be converted to extended β -strands, resulting in an increased rise per residue from 0.15 to 0.34 nm (Fig. 6D). Completely converting the coiled coils would result in a 43-nm length increase of the 46-nm long fibrin molecule ($\sim 93\%$ strain). This mechanism is supported by early wide-angle X-ray scattering experiments that indicated an α to β conversion in stretched fibrin clots [99, 100]. Additional support for this mechanism comes from recent experiments showing that force-induced unfolding of fibrinogen monomers results in a fibrinogen length increase that is consistent with the α to β conversion model [33].

A similar α to β -conversion has been observed in stretched keratin fibers. This conversion is *reversible* in hydrated, hard keratin (e.g., wool) [86, 89], though it appears to be *irreversible* in hagfish threads, a model for uncrosslinked, matrix-free intermediate (keratin-like) filaments [49]. Interestingly, coils of protein helices are a structural feature in several fiber-forming proteins that are exposed to stress, such as fibrinogen, intermediate filament, keratin (α -helical coiled coils), collagen (not α -helical) and other load-bearing proteins, such as tropomyosin and laminin. The detailed structural features of the coiled coils in those proteins are different. In fibrinogen the coiled coil is left-handed (about one turn) and consists of three right-handed α -helices [7]. In keratin, the coiled coil is also left-handed but consists of only two α -helices. In collagen, the “collagen triple helix” coil is right-handed and consists of three left-handed helices (not α -helices); this coil is not very extensible. It is intriguing to think of coiled coils as mechanical springs in proteins [101].

Scenarios 1 and 2 might account for the 120% elastic limit of uncrosslinked fibrin fibers [1].

3. At extreme strains (larger than the combined strains of scenarios 1 and 2) one or more of the globular domains might partially unfold (Fig. 6E). The γ C domain in the D-nodule would be the most likely candidate, because the stress forces are transmitted through the D:D contacts, A:a interactions and the γ - γ crosslinks within the protofibril. The unfolded domain might involve a stretch of about 140 residues (from γ Glu 183 to γ Asn 325) in the globular γ -domain, which does not contain key residues of the A:a interaction (γ Gln329, γ Asp330, γ His340, γ Asp364) or γ - γ

crosslinks (γ Lys406 γ Gln398/399). Since there are no disulfide bonds in this segment, it can be fully extended, resulting in a 50-nm length increase for each of the two γ C domain; this unfolding could account for 220% additional strain. Including the 100% from scenarios 1 and 2, a total of 320% strain could be accommodated. Similar, *reversible* unfolding has been observed for the immunoglobulin-like (and PEVK) domains in titin [60], and reversible unfolding has been proposed as a possible mechanism in fibronectin extensions [65].

4. If there is little lateral crosslinking between them, protofibrils might slide past each other [102, 103] and fibrin monomers within a protofibril might slide past each other. Without a restoring mechanism which pulls those units back, sliding will *not be reversible*, though. From studies on whole clots, it has been proposed that the protofibrils within a fibrin fiber [102, 103] and the fibrin monomers within a protofibril may slide past each other [104].

5. Several protein fibers with very large, mainly *reversible* extensibilities (spider silk, elastin fibers, resilin fibers) have amorphous, unstructured regions, which unravel under stress. Fibrinogen has a large unstructured domain, the 170 residue flexible connector region of the α C-domain. As the α C-domain may have a role in fibrin fiber assembly [105], stress forces may unravel this unstructured domain upon stretching. Since the length of this domain varies across species, one would anticipate that extensibility would vary among species if unraveling of the α C domains is important.

Scenarios 1 and 2 are likely reversible and could account for elastic deformations of up to $\sim 120\%$; however, it cannot entirely account for the 180% elastic limit of crosslinked fibrin fibers. In this case, scenario 3 might have to be invoked. We also noted that the broken arms of the *ruptured* fibrin fibers contract back to nearly their original length (see Fig. 4). This implies that even in the ruptured fibers most of the deformations are reversible, which argues against significant sliding and for scenarios 1, 2, and perhaps 3.

The rupture force of one A:a interaction was recently determined with laser tweezers and found to be 130 pN [106]. The force to rupture *both* A:a interactions in each half-staggered fibrin dimer is thus on the order of 260 pN. This is one of the largest rupture forces for protein–protein interactions. The unfolding force (as determined by AFM force spectroscopy) of T4 lysozyme (mainly α -helical) and the titin Ig-domain (mainly β -structure) are 64 pN [107] and 150–300 pN [60], respectively. Thus, judging by these forces, it is certainly possible that the α to β transition, which occurs at 100 pN, and perhaps the (partial) unfolding of globular domains occur *before* the A:a interactions break, giving credence to scenarios 2 and 3 above. Here, it should be mentioned that the rupture force depends on the

pulling speed; thus some of the force values may not be directly comparable if they were obtained at different pulling speeds. Nevertheless, the dependence of the rupture force, F_{rupt} , on the pulling speed, v_{pull} , is weak ($F_{\text{rupt}} \propto \ln v_{\text{pull}}$) [108] and these experiments are often carried out at similar pulling speeds. Thus, the comparison of the rupture force values is most likely warranted.

Two years ago it was unknown how far fibrin fibers can extend and from how large a strain they can recover elastically. Due to this lack of knowledge, the prevalent assumption was that the main mechanism by which a whole clot responds to stress is by an alignment of its fibers in the stress direction. Now we know that even uncrosslinked fibrin fibers can be strained 120% and still recover elastically. Most likely, this is important in a physiological context, where fibrin fibers are subjected to the stresses of flowing blood and retracting clots [109]. Large extensibilities and elastic recovery endow fibrin fibers with toughness, so they can absorb a great deal of energy before breaking.

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