The critical role of water in spider silk and its consequence for protein mechanics

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Abstract

Due to its remarkable mechanical and biological properties, there is considerable interest in understanding, and replicating, spider silk's stress-processing mechanisms and structure-function relationships. Here, we investigate the role of water in the nanoscale mechanics of the different regions in the spider silk fibre, and their relative contributions to stress processing. We propose that the inner core region, rich in spidroin II, retains water due to its inherent disorder, thereby providing a mechanism to dissipate energy as it breaks a sacrificial amide-water bond and gains order under strain, forming a stronger amide-amide bond. The spidroin I-rich outer core is more ordered under ambient conditions and is inherently stiffer and stronger, yet does not on its own provide high toughness. The markedly different interactions of the two proteins with water, and their distribution across the fibre, produce a stiffness differential and provide a balance between stiffness, strength and toughness under ambient conditions. Under wet conditions, this balance is destroyed as the stiff outer core material reverts to the behaviour of the inner core.

Introduction

The mechanical properties of spider silk originate from the core of the fibre, which is protected by a series of 'skin' layers¹ (Figure 1). The outer core (≈15% of the radius) is thought to consist mainly of the highly ordered protein Spidroin I, with the inner core (≈80% of the radius) consisting of both Spidroin I and the less-ordered Spidroin II¹¹². The intrinsic disorder in Spidroin II occurs due to the presence of proline, which twists away from simple, ordered configurations³. This combination of proteins results in a distribution of approximately 40% ordered domains (two hydrogen bonds per amide group), 15% permanently disordered domains (one hydrogen bond per amide group) and 45% intrinsically disordered domains but with potential for order³⁴⁴. Although the degree of order has a considerable influence on the strength, stiffness and elasticity⁵, the interactions between the ordered and disordered domains are also critical⁴⁶⁷.

An important factor in the ordering, and therefore mechanics, of spider silk fibres is water. In addition to understanding the structure-function properties of the material, the change in structure and mechanics upon wetting is particularly important for biomedical applications of spider silk, which involves an inherently wet environment. A number of studies of the bulk response of spider silk to water have, therefore, been undertaken using techniques such as tensile testing⁸, often in combination with other techniques such as x-ray scattering⁹, vibrational spectroscopy¹⁰, neutron scattering¹¹ and nuclear magnetic resonance (NMR)¹². These bulk fibre studies have shown that water infiltrates disordered domains, decreasing the alignment of the ordered

domains and contracting the fibre, yet our understanding of the response of different components, and the distribution of this response across the spider silk fibre, is lacking.

The distribution of Spidroin I and II in the different layers of spider silk provide a means by which their separate roles and interactions can be studied on a level of hierarchy above that of single-protein unfolding¹³ and below that of bulk mechanical analysis¹⁴. Here, we probe the nanoscale mechanical properties of the core material in situ, using atomic force microscopy (AFM) to mechanically investigate the roles of the two proteins and the influence of water on their respective behaviours. X-ray scattering was then used to determine the relative influence of water on the ordered and disordered domains. We find that water plays a dominant role in the mechanics of fibre, even under ambient conditions.

Methods

Major ampulate spider silk was harvested from a single Nephila clavipes (golden orb weaver) web in Redland Bay, Australia, the morning after spinning. A bundle of approximately 30 fibres was mounted in paraffin, cut into 5 µm cross-sections by microtome (RM2155, Leica Mikrosysteme GmbH, Austria) and mounted on a platinum plated silicon wafer. The paraffin was then removed by a sequence of 3 dips in citrisolve followed by a citrisolve/ethanol mixture and a gradual rehydration in a series of distilled water/ethanol mixtures. The samples were left to equilibrate overnight at approximately 60% relative humidity and 24°C, leaving an array of

samples. The ability to remove paraffin from the samples after sectioning provides a significant advantage over other standard cutting media such as epon and LR-white¹, which penetrate the material and harden, thus altering its mechanical properties.

Six sections were imaged in tapping mode (see Figure 2) before being subjected to indentation with an atomic force microscope under ambient conditions (Enviroscope, Veeco Instruments Inc., USA), first in its equilibrated state at 60% relative humidity, then immediately after 10 minutes immersion in distilled water, and again after reaching equilibrium overnight. Indentation was performed to a depth of approximately 50 nm, with a tip of nominal radius 15 nm (VISTAprobes T300; ≈10 nm radius before testing and 20 nm after). The cantilever had a resonant frequency of 298 kHz, from which the spring constant was calculated to be 40 N m⁻¹, by interpolating empirical values supplied by the manufacturer. A possible limitation of this technique is that by cutting the fibres, any pretension in the axial direction will be lost, potentially affecting the mechanical behaviour at the cut surface.

A second group of approximately 100 fibres was cut into a 15 mm length and fixed to a cardboard frame relaxed under 20% RH, forming a bundle of approximately 200 µm diameter. Small and wide angle x-ray scattering patterns were taken with the sample mounted (meridonal direction horizontally) in a closed environmental box at <1, 20, 40, 60, 80 and >99% RH measured with a humidity sensor (Hydroclip IE-1, Rotronic AG, Bassersdorf, Switzerland), at the SAXS beamline of the ELETTRA synchrotron radiation facility, Trieste, Italy¹⁵. Humidity was

controlled using a mixture of dry and wet air. The sample was hydrated in steps, with each step held for 20 minutes before measurements were taken. This was repeated three times down the length of the fibre bundle.

The x-ray scattering patterns were collected from a spot size of $200x50~\mu m^2$ at each position. A Mar300 Image Plate detector with a circular active area of 300 mm in diameter was used to acquire the patterns, which were processed using FIT2D Software¹⁶ and IGOR Pro (WaveMetrics, Oregon, USA). One-dimensional (1-D) profiles were obtained using an azimuthal integration over the region containing the reflections. The data were corrected for dark counts and background, and normalized to the region 6 nm⁻¹ < Q < 7 nm⁻¹.

The a and b dimensions were determined from analysis of the [200] and [120] reflections, respectively. For an orthorhombic structure, the unit cell dimensions a, b and c relate to the lattice spacing d for a particular set of Miller indices (hkl) as:

$$\frac{1}{a_{hkl}^2} = \frac{h^2}{a^2} + \frac{k^2}{b^2} + \frac{l^2}{c^2}$$

a can be extracted directly from the position of the 200 reflection, as $a = 2 d_{200}$. Determination of b relies on both the value of a determined from (200) and the position of the (200) peak, since:

$$\frac{1}{a_{120}^2} = \frac{1^2}{a^2} + \frac{2^2}{b^2}$$

The FWHM of the [200] and [120] reflections, ΔQ_{hkl} , can be used to determine the corresponding crystallite sizes, L_{hkl} according to the Debye-Scherrer equation:

$$L_{hkl} = \frac{k4\pi}{\Delta Q_{hkl}}$$

in which k is a scaling constant taking into account the formfactor of the crystallites.

Results

Under ambient conditions, the mechanical behaviour varied with position on the cross-section (Figure 2, top). The highly ordered outer core exhibited a purely elastic behaviour, with the unloading curve closely following the loading curve. The inner core showed a distinct 'plastic' effect, whereby some of the applied energy was dissipated and the strain not recovered. Although the outer-core silk showed considerably more resistance to indentation than silk from the inner core (22%, given by the slope of the force-displacement curves in Figure 3), the first few nanometres of unloading, to which elastic properties are usually attributed¹⁷, were very similar (within 5%) in the two regions.

After immersion in water (Figure 2, bottom), the behaviour of the inner core changed little: a lower indentation stiffness (23±10%) and a slightly decreased elastic recovery (7±3%) during unloading, compared to ambient conditions. The outer core, however, showed a radically different load-unload profile, changing from a purely elastic response to a highly plastic behaviour, similar to that of the inner core under ambient conditions. It is also interesting to note that the slopes of the unloading curves were remarkably similar (within 3%) for the wet inner and outer and core material (Figure 3B).

The small (SAXS) and wide (WAXS) angle x-ray scattering experiments (Figures 4a, 4b, 5) showed no change in crystal size with increasing hydration. Based on the conventions set forth by Warwicker¹⁸, the orthorhombic unit cell of the crystallites is designated to have its c-axis along the direction of the fibre, with its a- and b-axes distributed randomly around the fibre axis.¹⁹ As we measured only the [120] and [200] reflections, only the dimensions along the side chains (a-axis) and along the hydrogen-bonds between the pleated sheets (b-axis) were investigated. It is in these directions, however, that swelling is expected to occur.

Figure 4 shows the average of the three data sets collected in the SAXS/WAXS experiments. The data were collected along the equatorial direction, and exhibit the expected peaks corresponding to the [120] and [200] reflections (inset). The positions and shapes of the peaks are not significantly sensitive to humidity; however, humidity dependent changes can be observed in both the low-Q (SAXS) region and the very high-Q region, as well as in the intensity of the peaks. In the case of the SAXS data, the curves exhibit broadening and increased intensity above 80% humidity, which indicates an increased contrast between the amorphous and crystalline zones consistent with water uptake in the amorphous component.²⁰ In the case of the high-Q data (Q > 16 nm⁻¹), the intensity increases systematically with humidity. As in previous studies^{20,21}, we attribute the intensity at high Q values to scattering from disordered water, and accordingly fit this contribution with an additional Gaussian peak profile. The [200] and [120] reflections were fit simultaneously using Gaussian peak profiles. When freely fit simultaneously with the [200] and [120] reflection, this peak is consistently located at $Q = 16.3 \text{ nm}^{-1}$.

Plots of the humidity-dependent data are presented in Figure 5. The unit cell parameters (Figure 5a) are insensitive to humidity and in agreement with previous studies on supercontracted spider silk.^{20,22} The average cell parameters determined from the [200] and [120] reflections are a = 1.008±0.008 nm and b = 0.966±0.001 nm, respectively. The Debye-Scherrer crystallite sizes along both [120] and [200] are similarly insensitive to humidity (Figure 5b), though a small, statistically insignificant swelling trend was observed in the direction along the hydrogen-bonds between the pleated β -sheets. It is also interesting to note the increase in unit cell parameter a from dry to 20 % RH. We found minimum average crystallite sizes of 3.1 and 7.3 nm along these respective directions, consistent with previous reports of anisotropic crystallites²⁰. This size is smaller than the radius of the AFM tip and the displacements applied during indentation.

Discussion

This study aimed to investigate the mechanical effect of water on different regions of the spider silk fibre, and therefore on the two major proteins, using AFM indentation and x-ray scattering. We found that the inner core region, rich in spidroin II, retains water due to its inherent disorder, arguing that this dissipates energy through the breaking of a sacrificial amide-water bond and increasing its order under strain. The outer core, rich in spidroin I, is more ordered under ambient conditions and is inherently stiffer and stronger, yet does not on its own provide high toughness. When the distribution of these effects is considered across the fibre, we propose that

the tough inner core and stiff outer core combine to produce a highly resilient material under ambient conditions. In wet conditions, however, the outer core behaviour changes to that of the inner core, weakening the fibre as a whole.

We interpret the 'plastic' effect in Figures 2 and 3 as resulting from the level of disorder, which is known to be both intrinsic to the Spidroin II in the inner core due to the presence of proline³, and forced by immersion of the fibre in water²³ (see the change in ordered fraction and crystallite swelling in Figures 4 and 5). The induced strain from the indentation, when transferred to the protein level, will push/pull some of the disordered domains into an ordered configuration, preventing the material from recovering elastically. The amount of reconfiguration is difficult to quantify due to the complex nanoscale structure of spider silk, but for a given structural arrangement and constant indentation conditions, is proportional to the area between the loading and unloading curves, and therefore proportional to the amount of energy dissipated during the indentation. This energy is roughly equal to the product of the energy per hydrogen bond and the number of hydrogen bonds broken, with the weaker bonds breaking first. With the correct (tight) packing conditions, a similar effect is likely to be observed under tension³.

During spinning, most of the water is expelled from the aqueous protein solution (70% water)^{24,25}; the highly hydrophobic, ordered domains expel all of their water but, as inferred from IR spectra²⁶, some water remains in the less hydrophobic, disordered domains. The higher shear stresses in the outer fibre during extrusion will arguably lead to the formation of more ordered domains compared to the inner

regions, similar to the increased order and orientation resulting from the higher stresses during fast extrusion²⁷. Concurrently, water would be forced out of the structure in these highly stressed outer regions, but less so from the inner regions. The disorder due to water results from it acquiring hydrogen bonds between the molecular chains in the fibre's oriented amorphous domains²³, which are easier to penetrate than the hydrophobic, crystalline domains such as β -sheets²⁸.

The integrated intensity of the [120] and [200] peaks (Figures 5c) is related to the degree of crystallinity of the fibre. Water is known to significantly reduce the volume fraction of the crystalline component in spider silk due to uptake in the amorphous domains²⁹, yet bulk cyclic loading experiments and modelling suggest that the ordered phase of a specific fraction of the peptide segments can be manipulated by exposure to water³. The decrease in the integrated intensities of the [120] and [200] reflections, and the increase in the intensity of the amorphous peak (Figures 4b, 5c), confirm a decrease in the volume fraction of ordered domains with humidity.

Through the same mechanism as for intrinsic disorder described above, the disordering of the structure caused by the infiltration of water (Figure 5c) would be reversed, and forced back into an ordered configuration under an applied load. This raises some interesting questions; particularly whether the water in the disordered domains plays an equal or even superior part to that of the proline³⁰ and if so, whether the main mechanical function of the proline is to enable the action of water.

We propose that under ambient conditions, the bound water between protein chains in the inner core will be pushed out under an applied load, allowing reconfiguration due to sacrificial bonding. Once the load is withdrawn, however, the now unbound water can reacquire some of the hydrogen bonds, allowing a partial recovery. This is essentially a self-repair mechanism, much like the repair of hydrogen bonded rubber³¹, but is controlled by water. As more bonds are taken up by water, a greater reconfiguration is possible, resulting in softening and a decreased elastic recovery (Figure 3). The observed softening, we propose, is due to the amide-water bonds being weaker than the amide-amide bonds that they replace²⁴. The decreased recovery will result from the combination of decreased order and the lower stress required for reconfiguration. Water-controlled recovery will be slower than the entropy-driven elastic recovery as it will take time for the water to infiltrate the more-ordered structures. The effect of this slow recovery can be seen when scanning 30 minutes after indentation, at which time the residual indentations are no longer observed. The timescale of water-driven recovery is in good agreement with stress relaxation data from bulk fibres¹⁴, which show >95\% relaxation after 30 minutes, and full relaxation at approximately 80 minutes.

The role of water in the intrinsically disordered inner core structure leads us to consider the behaviour of the outer core which, in the absence of proline, appears to be highly water dependent (Figure 3). A full elastic recovery was observed under ambient conditions due to the high degree of order, and therefore strength at a local scale stemming from the cooperative rupture of hydrogen bonds described above and in^{32,33}. The reversible disorder caused by immersion in water (Figure 5) leads to a plastic effect, similar to that of the inner core. That is, the mechanical behaviour of the outer core is affected by disorder in the same way as the inner core, but the

outer core does not possess the inherent disorder of the inner core under ambient conditions. The changes in behaviour after controlled humidification suggests that the disorder in the inner core acts as a reservoir, providing binding sites for the water, and thus creating a mechanism for the high toughness observed in bulk mechanical investigations¹⁴. The outer core does not provide this mechanism and, rather than contributing to toughness, is more likely to play a dominant role in the stiffness of the fibre, and is responsible for the linearity of the stress-strain behaviour in the elastic region. We speculate that the inner core region will exhibit a 'J' shaped stress-strain curve under ambient conditions, similar to that observed in bulk studies of supercontracted fibres^{14,34}.

Taking up nearly a third of the fibre's volume¹, the very different behaviour of the outer core is likely to play an important protective role, with its higher hardness and stiffness complementing the tough, impact absorbing inner core similarly to differentially hardened metal in gears, knives and swords, and has been synthetically introduced to increase toughness in biological systems^{35,36}. These interpretations of the action of water in spider silk mechanics are limited to small strain behaviour. It is difficult to extrapolate to large-strain behaviour, particularly as the role of water after the conversion to a rubber-like elasticity during yield and later strain hardening^{5,37} is likely to change considerably.

The main functional difference between the two proteins, Spidroin I and II, appears to be the ability of Spidroin II to retain water (see also Creager $et\ al^{12}$). Once the effect of water has been nullified either by saturation (Figure 3b) or expulsion by

applied stress (unloading curves, Figure 3a and b), they have very similar deformation mechanisms at small strains, with a small contribution from the proline in the Spidroin II. The effect of proline on the mechanical behaviour can be decoupled from that of water by considering the loading curves of the wet samples (Figure 3b). Here, the water-saturated, proline-rich inner core is only slightly softer than the water-saturated, proline-deficient outer core, with the difference in area between the loading and unloading curves for the two regions showing the dissipation of energy resulting from the presence of proline. The difference in energy dissipated by the proline appears small (21%), relative to that dissipated by the action of water in the inner core under either wet or ambient conditions.

The similar behaviour of the wet inner and outer cores (Figure 3b) indicates that the action of bound water is critical in load carriage, providing sacrificial amide-water bonds that can be broken to dissipate energy, allowing local strains as the protein chains are pulled into a more ordered configuration and the sacrificial bond is replaced by a stronger and more direct amide-amide bond. Once restructuring has occurred, through disordering by immersion in water or ordering by applied stress, the two proteins are virtually identical from a mechanical perspective. This sacrificial bonding has been observed in the highly disordered capture silk of Araneus spiders¹³ and in recombinant dragline silk³⁸, using AFM to apply tensile loads to individual proteins. Although these studies provide an interesting insight into silk protein unfolding, the single molecule tests that they employed do not retain the packing conditions existing in situ and consequently, are unable to relate their findings to water or the self-repair of hydrogen bonds as the material is restructured

under stress. The strength of these sacrificial bonds can, however, be related to the stiffness of the fibre and the softening observed after saturation with water⁸.

The critical role of water in the small strain mechanics of spider silk, the vastly different behaviours of inner and outer core silk under ambient conditions and their similarity under wet conditions, provide some insight into the stress-processing and energy dissipation mechanisms that underlie its remarkable mechanical properties. It may also build on our current understanding of this and other biological fibres such as collagen, elastin and keratin. In this context, our results indicate that the selective binding of water to proteins controls their mechanical behaviour by dictating the level of order and therefore the potential for energy dissipation, restructuring and self-repair.

Building upon this conclusion, we speculate on the impact of the x-ray scattering and indentation results for the development of mechanically robust synthetic silks as biomaterials, which will require compensation for the effects of water³⁹. Rather than synthesizing a material to recreate the properties of spider silk, we argue that it is better to synthesize a stiffer, more hydrophobic material that will restructure in wet conditions to provide the water-dominated sacrificial bonding, energy processing and self-repair mechanisms of the inner core material, yet retain the stiff, elastic outer core behaviour that is lost in natural fibres. By looking beyond spider silk mechanics to consider the ordered structures that provide strength and stiffness in wet fibres such as collagen and elastin, it may be possible to recreate, in vivo, the delicate

balance of properties that make spider silk such a formidable material in ambient conditions.

Outside of the biological environment, the sacrificial toughening mechanisms in the inner core material, and the complementary matching of inner and outer core properties may provide a target for biomimicry. In this respect, a composite can be designed, using a stiff material to carry 'design' loads coupled with a high toughness sacrificial material for one-off extreme loads in seatbelts, bullet proof vests, structural cables in airplanes, construction for earthquake resistance, and other safety applications.

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Figures

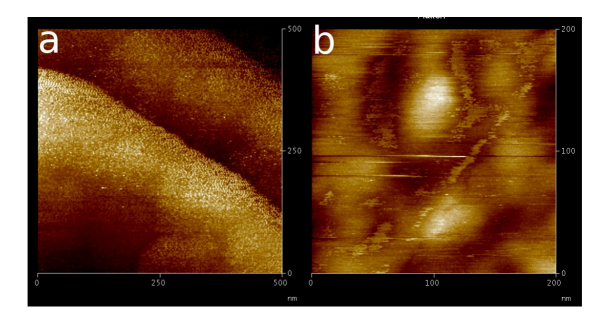


Figure 1: AFM height image of spider silk fibre cross-section. (a) two skin layers, with the centre of the fibre towards the bottom-left of the image; (b) core region showing globular morphology.

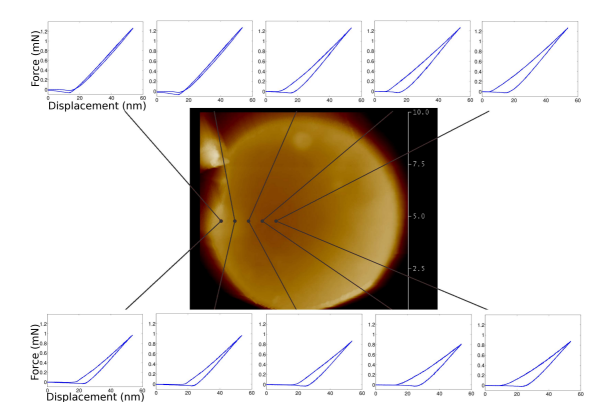


Figure 2: AFM image of cross-sectioned fibre with the profile of force-displacement curves from the outer to inner regions shown under ambient (top) and wet (bottom) conditions. The scale bar on the right of the AFM image is in microns, and the axes of the force-displacement graphs are constant with y-axes from 0 to 1.2 mN and x-axes from 0 to 60 nm.

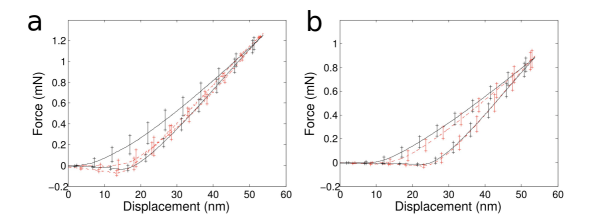


Figure 3: Average indentation force-displacement curves under ambient (a) and dry (b) conditions. Black lines represent inner core material; red, dashed lines represent outer core material.

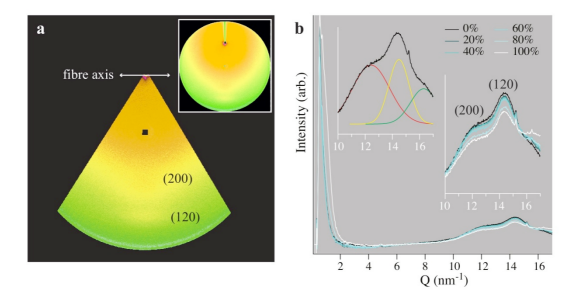


Figure 4: (a) Masked scattering data, indicating the integrated region used to obtain SAXS/WAXS profiles. The (200) and (120) peaks are indicated, along with the fibre axis direction and location. The inset shows the entire scattering pattern. (b) Integrated average SAXS/WAXS profiles for RH varying from 0-100%. The inset at right shows an enlarged view of the WAXS region, with the (200) and (120) peaks indicated. The inset at left shows a typical set of Gaussian fits, comprising the (200) peak (red), the (120) peak (yellow) and the amorphous water peak (green).

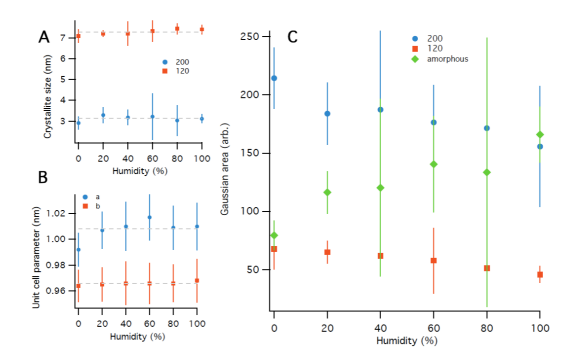


Figure 5: Fits of the data in Figure 3. Crystallite size calculated from the Debye-Scherrer equation (a), unit cell parameters (b), and integrated intensity of the peaks (c).