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Structure-Function-Property-Design Interplay in Biopolymers: Spider Silk

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Abstract

Spider silks have been a focus of research for almost two decades due to their outstanding mechanical and biophysical properties. Recent advances in genetic engineering have led to the synthesis of recombinant spider silks, thus helping to unravel a fundamental understanding of structure-function-property relationships. The relationships between molecular composition, secondary structures, and mechanical properties found in different types of spider silks are described, along with a discussion of artificial spinning of these proteins and their bioapplications, including the role of silks in biomineralization and fabrication of biomaterials with controlled properties.

Keywords

Spider silk; proteins; secondary structure; self-assembly; genetic engineering

Introduction

The order Araneae (spiders) contains over 37,000 species that have ability to produce silk, a key protein for spider survival [1]. Spiders use silk for a variety of practical purposes including arresting a fall, swathing prey, building a web, lining burrows, or making egg cases [2, 3]. Diverse uses of Araneae silks originate from the outstanding physical properties that are tailored for specific needs, resulting in variation of mechanical properties. Mechanical properties of spider silks have not yet been duplicated by man-made materials. For instance, major ampullate silk (also known as dragline silk) of Caerostris darwini has a maximum strength (the strength needed to break the fiber) of up to 1.7 GPa that is in the range of that of high-tech materials. Nevertheless, man-made materials such as steel (1.5 GPa) and Kevlar (3.6 GPa) present higher strength and stiffness, the overall toughness (energy required to break the fiber) depends on the extensibility. In these terms, high-tech
fibers appear to be more brittle compared to the more extensible dragline threads making spider dragline silks the strongest material [4–6]. In addition to this, spider dragline silks are light-weight materials that can undergo supercontraction when hydrated [7]. On an equal weight basis, spider silk has a higher toughness (the amount of energy absorbed per volume before breakage) than steel and Kevlar due to extensibility [8, 9]. Moreover, the other types of silk demonstrate unique mechanical properties as well. For example, flageliform silk (also known as a capture spiral silk) is over ten times more extensible than dragline silk, on the other hand flageliform silk is neither strong as nor as stiff than dragline silk [10]. Little force is required to extend flageliform silk 200–300% of its original length until the fiber gradually stiffens under the increasing strain. The remarkable mechanical properties of spider silks originate from their unique amino acid sequences that have remained almost unchanged for more than 100 million years [11]. Moreover, spider silk primary amino acid sequences have been shown to be organized in block (or domain) copolymer-like arrangements, in which two basic sequences, crystalline and less crystalline polypeptides alternate. At least seven different types of silk proteins are known for one species of spider [12]. They differ in primary sequence, physical properties and functions [1]. The complex spinning process, allows spiders to transform water soluble silk proteins into solid fibers at ambient temperature and pressure, giving rise to an environmentally safe, biodegradable, and high performance material [13–15].

Diverse and unique biomechanical properties together with biocompatibility and a slow rate of degradation make spider silks excellent candidates to be studied as biomaterials for tissue engineering, guided tissue repair and drug delivery, for cosmetic products (e.g. nail and hair strengthener, skin care products), and industrial materials (e.g., nanowires, nanofibers, surface coatings) [14–18]. A complete understanding of the anatomy of spider glands, the silk spinning process, and the role of silk primary structural elements and their contributions to the physical and biological properties of the biopolymer are important in order to fully understand the materials. Several reviews have focused on the structure and properties of spider silks, with the main focus on dragline silks [3, 16–19]. The structure-function relationships in other types of silk polymers are not as fully understood and this gap in knowledge can result in limitations in the use of these materials. In this review article, the goal is to provide information on the current understanding of the structure and properties of various types of spider silks with the main focus on the interplay among structure, architecture, and function, and how these attributes are being used in biomaterial designs.

1. Anatomy and Physiology of the Spider Spinning Apparatus (*Nephila clavipes*)

Orb web spiders produce silk in seven distinct pairs of glands. However, it is thought that all spider glands evolved from a single type of gland which diverged in anatomy, morphology and luminal content [20]. The presence of different amino acids in the luminal contents within each gland is responsible for the secretion of defined silk fibers with specific functions. Today, most research is focused on the major ampullate gland, which produces dragline silks.

In *Nephila clavipes* the silk formation glands are located within the abdomen of the spider, and each type of gland occurs in pairs with bilateral symmetry [12]. The major ampullate gland can be schematically divided into four zones as shown in Figure 1: the tail zone, responsible for synthesis and secretion of spider silk proteins, the lumen (also known as the sac), involved in protein accumulation, the spinning duct responsible for the alignment of silk fibers, and the spigot for final fiber production [21].

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The gland represents a reservoir of soluble silk proteins that are synthesized in specialized columnar epithelium cells in the tail zone and secreted into the lumen, where the proteins are stored as a highly concentrated liquid crystalline solution [22]. According to Vollrath and Knight [23], the tail zone is named the A-zone and the first part of the lumen is named the B-zone, based on the presence of the single type of cells known as the tall columnar secretory epithelium. The spinning duct follows the lumen, which is folded into an S-shape and narrows to the end [23]. Within this duct, silk proteins are present as liquid dope [22]. The spinning duct consists of three limbs. In each limb spidroin molecules have a different orientation. In the first limb, silk proteins are anchored perpendicular to the cuticle lining but parallel to each other [23]. In the second limb of the duct, silk molecules are bent in such a way allowing the formation of layered discs made up of the amphiphilic rod-shaped molecules that are still connected to the ducts cuticle lining. Finally, in the third limb the silk dope pulls away from the cuticle lining forming a draw down taper [22]. At this moment, the liquid silk solution is converted into a solid thread surrounded by water and it is thought that β-sheet formation is initiated during this process by means of the rapid extension flow that pulls silk molecules close together and aligns them using hydrogen bonding into β-sheets crystals [24, 25].

During the silk dope journey through the spinning duct, a number of changes in the chemical environment take place. These include lowering of pH from 6.9 in the first limb to 6.3 in the third limb, decreasing sodium and chloride concentrations, and increasing potassium, phosphate, and sulphate concentrations. At the same time, the drop in pH together with ionic changes can facilitate the neutralization of repulsive negative charges aiding silk molecule alignment into β-sheets. The final part of the major ampullate gland is represented by the valve and the spigot. The valve is used as a clamping device to control spider dragline and as a pump to remove threads that are broken inside the spigot. The spigot (spinneret) is located at the very end of the gland from which the spider thread is drawn. It has been proposed that the spigot is also capable of further modifying the silk thread [22].

Spider’s threads are only produced when the spider physically moves away from one attachment point to another. A combination of elongation flow and wall shear within the gland creates tensile strain that induces β-sheet transitions in silk proteins as they pass through the fluid-to-solid transition zone (the taper). Higher tension imposes greater molecular alignment in the thread, making the silk stiffer, stronger and less extensible. Lower tension makes the silk less stiff, and weaker but more extensible. It should be noted that at higher temperatures, spiders move faster and therefore spin silk faster. Faster moving animals produce silk threads with different diameters and mechanical properties than compared to slower moving animals. When the silk protein is drawn from the spinning duct all the way down to the spigot, silk becomes progressively more dehydrated, slightly acidified, and birefringent, suggesting that the orientation and shape of silk molecules have been already programmed. Spiders can also apply a tension brake upstream from the valve that allows them to increase silk thread tension while the thread is still in the high humidity environment of the spigot. Figure 2 depicts the natural spinning process in spider.

2. Spider Silk Types and Their Modules

One spider is capable of producing up to seven different types of silks with varying mechanical properties. Silks are produced in specialized glands including the major and minor ampullate, tubuliform, flagelliform, aggregate, pyriform, and aciniform [21]. Spiders use their silk for a variety of practical purposes including: arresting a fall, swathing a prey, building a web, lining burrows, or making egg cases. Figure 3 depicts silk glands, silk types and silk functions of *Nephila clavipes*.
2.1 Major Ampullate Silk Proteins

The golden orb weaver spider, *Nephila clavipes*, produces dragline silk in the major ampullate gland [26]. The protein complex is composed of major ampullate dragline silk protein 1 (MaSp1) and major ampullate dragline silk protein 2 (MaSp2). In 1990 Xu and Lewis screened a cDNA library with a probe based on a short peptide isolated from acid hydrolyzed *N. clavipes* dragline silk and found the gene encoding MaSp1 [27]. The total length of MaSp1 transcript is 12.5 kb [28]. Analysis of the primary sequences of MaSp1 from different species revealed regular small peptide motifs that can be grouped into three categories: poly-alanine domains (poly(A/GA)), glycine-rich domains (GGX), and non-repetitive N- and C-termini [1]. Additional studies [28] demonstrated that the poly-alanine regions can contain between four to seven alanine residues and the X position in the GGX repeat can be occupied only by Y, L or Q. Figure 4 depicts the repeating protein units of MaSp1 revealed by Hu and Lewis [27].

MaSp2 possesses similar structural motifs compared to MaSp1. It is composed of poly(A) and poly(GA), GPGGX domains, and non-repetitive N-terminus and C-termini. The main difference between MaSp1 and MaSp2 is the presence of proline (P) residues accounting for 15% of the total amino acid content in MaSp2 [1]. MaSp1 is proline-free. By calculating the number of proline residues in *N. clavipes* dragline silk, it is possible to estimate the fiber content; 81% MaSp1 and 19% MaSp2 [29]. Different spiders have different ratios of MaSp1 and MaSp2. For example, a dragline silk fiber from the orb weaver *Argiope aurantia* contains 41% MaSp1 and 59% MaSp2 [30]. Such changes in the ratios of major ampullate silks can dictate the performance of the silk fiber [23].

The sizes of the mRNAs and genes encoding these proteins have been determined, with the mRNA sizes for MaSp1 and MaSp2 approximately 12.5 and 10.5 kb, respectively, generating a protein complex with the molecular weight of 350 kDa [1]. *N. clavipes* uses MaSp1 and MaSp2 to build web frames and safety-lines. Another spider, *Araneus diadematus*, produces analogous MaSp1 and MaSp2 proteins known as ADF-3 and ADF-4 that possess similar structural motifs and perform similar functions [9]. In addition, a genomic library for the black widow spider, *Latrodectus hesperus* (Theridiidae) was constructed in order to identify full-length silk genes and regulatory regions [31]. The gene sequences of *MaSp1* and *MaSp2*, as well as adjacent non-coding regions, were identified. The existence of higher-order repeat units that range from 70 to over 2,000 bp was revealed; and the authors showed that the repetitive sequences of MaSp1 were more homogeneous than those of MaSp2. In addition to the repetitive domains, dragline silks contain non-repetitive N and C-termini [27] [32]. The length and amino acid composition of the C-termini are conserved between major ampullate proteins from different spiders and the C-terminal region is important for solubilizing silk proteins in the highly concentrated spinning dope [1]. In contrast to the C-terminus, little is known about the N-terminal region of spider silks. The sequence possesses an enzymatic cleavage site, has an usual amino acid composition and may include a signal peptide [25].

2.2 Minor Ampullate Silk Proteins

Minor ampullate silk may be particularly interesting for biomedical applications since it is strong and does not supercontract in water (note: NMR data show some structural changes) [33, 34]. The minor ampullate silk from *N. clavipes* consists of two distinct proteins, the minor ampullate dragline silks 1 and 2 (MiSp1 and MiSp2), with transcripts of 9.5 and 7.5 kb, respectively, generating proteins of 320 kDa and 250 kDa [28]. The cDNA sequence representing the MiSp1 transcript from *N. clavipes* was published shortly after the initial discovery of the MaSp1 sequence [35]. Figure 5 shows the consensus repeats found in MiSp1 and MiSp2 from *N. clavipes*. 

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The minor ampullate dragline silk proteins possess similar structural domains to the major
ampullate silk proteins; however, there are some differences. The similarities include
the presence of GGX and poly(A) modules. However, poly(A) modules of the MaSps are
replaced by poly(GA) repeats [33]. The organization of modules is also similar to MaSps;
whereas, the number of repeats is different. Another difference is the presence in MiSps of a
conserved serine (S)-rich non-repetitive 137 amino acid sequence that is termed a spacer
region [35]. The spacer regions have a similar serine composition relative to the amorphous
regions of Bombyx mori and are highly conserved among different spider species. It is
hypothesized that the spacer plays a role in fiber formation by serving as a site of protein-
protein interactions. Recently, a full-length minor ampullate spidroin gene sequence was
obtained from the spider Araneus ventricosus using PCR [33]. Sequence alignment of spacer
regions was performed; however, the functional role of this spacer remains to be determined.

2.3 Flagelliform Silk Proteins

The major component of the capture spiral of a spider web is flagelliform silk (Flag), which
is produced by the flagelliform gland. Hayashi and Lewis retrieved a partial cDNA sequence
of flagelliform silk from the N. clavipes flagelliform gland [36]. The flagelliform gene
encodes an mRNA transcript of 15kb that generates a protein of ~500 kDa [1]. Flagelliform
silk has several distinctive features; a) GPGGX domain; b) highly conserved spacer with
charged and hydrophilic amino acids; c) nonrepetitive C-terminal region [37]. Such a
structural combination gives flagelliform silk high elasticity; it can stretch up to 200% of its
own length. The GPGGX domain is thought to be responsible for the elasticity of
flagelliform silk. This GPGGX domain was also found in MaSp2. Flagelliform silk has the
same GPGGX repeat but it occurs seven times more in flagelliform than in MaSp2.
Flagelliform silk exhibits seven times more elasticity than MaSp 2 [21]. The molecular
mechanism of such elasticity is the formation of a molecular spring (β-spiral), in which a
series of type II β-turns are linked together. The spacer region has no homology with known
proteins and the functions are unknown. Also unknown is the function of the C-terminal
region, which shows no similarity to the C-termini of MaSps and MiSp [32] [38].

2.4 Aciniform Silk Proteins

Aciniform silks are used by araneoid orb-weavers, including Nephila clavipes, to wrap and
immobilize prey, build sperm webs and for web decorations [1]. Aciniform silk contains
only one protein, aciniform spidroin 1 (AcSp1), with 14 highly homogeneous repeats, with
each ensemble repeat consisting of 200 amino acid domains [39]. AcSp1 has a low
percentage of G and A relative to dragline silk and possess a C-terminal domain that has
little similarity with major ampullate, minor ampullate or flagelliform silks. Moreover, the
repetitive sequences predicted from the AcSp1 cDNA do not resemble iterations found
within other spider silk proteins [39]. In terms of mechanical performance, aciniform silks
have a lower tensile strength relative to dragline silks but higher extensibility and toughness.
Based on the lack of these similarities aciniform silks likely represent divergent members of
the silk gene family.

2.5 Tubuliform (Eggcase) Silk Proteins

Tubuliform silk (TuSp1) is produced by female spiders during the reproductive season to
make egg cases. This silk type is secreted by the tubuliform (cylindrical) gland and has a
distinct amino acid composition which is responsible for the unusual mechanical properties.
Tubuliform silk has a relatively high tensile strength with a fairly low elasticity that makes it
different from other silk types [40]. Protein sequences from three orb-weaving spiders,
Arsiope aurantia, Araneus gemmoides, and N. clavipes revealed that TuSp1 rarely show the
four common spider silk motifs discussed earlier, GPGXX, GGX, poly-A, spacer. Instead,
TuSp1 has a more complex architecture with different amino acid motifs such as Sn, (SA)n, (SQ)n, and GX (X represents Q, N, I, L, A, V, Y, F, or D) [41, 42]. The interesting feature is that tubuliform silk is also capable of forming β-sheets as a result of the structural domains, such as AAQAASAA, AAQAA, and AASQAA [42]. The presence of side chains in the silk modules (Q and S) directs larger spacing among the formed β-sheets relative to the poly-A/(GA)ₘ model; this makes the tubuliform silk more hydrophilic [40]. The tubuliform C-terminal domain has no similarity with other silks and it is assumed that TuSp1 is evolutionarily an ancient silk [42].

Only recently, two other proteins from egg case core fiber of L. hesperus were discovered [43]. They are called egg case protein 1 and 2 (ECP-1 and ECP-2). Analysis of the primary sequence of ECPs indicate similarity to dragline silks based on the presence of poly-A/(GA) modules [43]. As a result, it is now assumed that the core fiber of egg case silk is a trimeric complex consisting of TuSp1, ECP-1 and ECP-2, where TuSp1 serves as a core protein and ECPs function as intermolecular crosslinkers, with a potential structural role in an egg case fiber.

2.6 Fiber Coating (Glue) and Pyriform Silk Proteins

Little is known about fiber coating peptides and pyriform (alternative spelling: piriform) silks. Recently, Hu and coworkers have isolated the cDNAs of two distinct fiber coating proteins, which were named spider coating peptide 1 (SCP-1) and spider coating peptide 2 (SCP-2), respectively [44]. Secreted forms of SCP-1 and SCP-2 contain 36 and 19 amino acids, respectively, and their primary sequences display no similarities to the ensemble repeat units from traditional spidroins. Quantitative real-time PCR analyses showed that mRNAs of SCP-1 and SCP-2 are produced by the aggregate gland. In addition to this, extensive expression studies have also revealed that L. hesperus, commonly referred to as the black widow spider, is capable of producing large amounts of SCP-1 in the flagelliform gland[45, 46]. Biochemical studies demonstrated that the SCP-1 peptide has intrinsic metal binding properties, suggesting a role of peptide-metal ion interactions with the fiber constituents to enhance thread performance [44]. In L. hesperus the aggregate gland has also been shown to produce two different proteins, aggregate gland silk factor 1 (AgSF1) and aggregate gland silk factor 2 (AgSF2) [47]. These proteins were found to be co-localized to the connection joints of both webs and wrapping silks spun from the black widow spider. Biochemical analysis revealed that AgSF1 and AgSF2 have different protein architectures compared to traditional spider silk family members. In case of AgSF1, the amino acid sequence analysis revealed the presence of pentameric QPGSG iterations that are similar to conserved modular elements within mammalian elastin. In case of AgSF2, the consensus novel repetitive sequence, NVNVN, was determined. The amino acid block repeats were found to be embedded in a glycine-rich matrix of AgSF2 [47]. Adhesive spider silk proteins (glues) are important members of the silk family that allow spiders to perform a variety of tasks, such as web construction, prey capture, and locomotion. More extensive studies are needed to reveal the primary structure, properties and functions of these silks among different spider species.

Pyriform (piriform) silk proteins are relatively newly discovered members of the spider silks family. They are components of the attachment discs, which lashes the joints of the web and attaches dragline silk to surfaces [48]. Recently, a novel member of the silk gene family, pyriform spidroin 1, was discovered in attachment discs of the black widow spider, L. hesperus [49]. Alignment of the PySp1 amino acid sequence to other fibroins revealed conservation in the non-repetitive C-terminal region of the silk family. Scanning electron microscopy studies revealed the presence of small diameter fibers embedded in a glue-like cement that formed a continuous network with large diameter dragline silk threads,
producing a strong, adhesive material. The pyriform silk is produced by the pyriform gland and it is used for attachments and for joining spider threads. The amino acid composition has been determined using luminal gland materials from orb-weaver spiders and it was found that pyriform silk contains very polar amino acids, such as S, D, E, L, T, and N [20]. The precise molecular identities of the proteins involved in both silks remain to be determined. Recently, pyriform spidroin 2 (PySp2) was discovered within the attachment discs of N. clavipes by Geurts and coworkers [50]. Biochemical studies revealed that PySp2 contains internal block repeats that consist of two subrepeat units: SQA-rich and P-rich. Artificial spinning of recombinant PySp2 truncations shows that the SQA-rich subrepeat is sufficient for the assembly of polymeric subunits and subsequent fiber formation. These studies support that both orb- and cob-weaving spiders have evolved highly polar block-repeat sequences with the ability to self-assemble into fibers, suggesting a strategy to allow fiber fabrication in the liquid environment of the attachment discs[50].

In conclusion, the relationships between the core silk sequences and all of their roles in spider survival remain unclear. Additional studies are required to elucidate the silk sequences for a variety of orb weaver and cob weaver spiders. Table 1 summarizes the domains and core units found in different spider silk types. This table also provides information regarding the structure of aciniform (AcSp1), tubuliform (TuSp1), and pyriform (SCP1 and 2) silks.

### 3. Secondary Structure Elements of Silk Proteins

The outstanding physical properties of major ampullate dragline silks originate from the protein primary sequences, where hydrophilic terminal domains flank large central hydrophobic domains [51]. Spider silk proteins and their genes are only partially characterized; however, it is generally concluded that most of the spider silks are made up from four different motifs: (i) 3\textsubscript{10}-helix forming repeats GXX; (ii) crystalline β-sheet rich poly(A)/poly(GA) motifs, (iii) an elastic beta-spiral region, also known as proline-rich region, composed of multiple GPGXX motifs; and (iv) a spacer region with unknown functions. These motifs appear to be conserved among different species and are found in orthologs of cDNAs from other Araneus genera [52]. The silk protein secondary structure arises from the interplay between spider silk key elements, namely A-rich and GA-rich motifs. This interplay leads to the formation of both helical structures (3\textsubscript{10}-helices, α-helices, PP II helices) or β-sheet structures that are responsible for the physical properties of spider silks as well as the explanation of the rapid fiber formation [9, 22].

The majority of studies on spider silks have been focused on the major ampullate dragline silks due to a unique combination of elastic and tensile properties. The studies established that dragline silks contain extended, ribbon-like β-sheets that are oriented parallel to the fiber axis, β-turns and helices [39]. Poly(A) and poly(GA) regions are responsible for the fiber tensile strength due to the formation of crystalline β-sheets. Poly-A repeats form a β-sheet conformation in which each A residue is placed on alternative sides of a backbone [28, 38]. The hydrophobic interactions that arise from such conformation connect poly(A) chains together by protruding methyl groups occupying the void space near the alpha carbon of a residue on a neighboring chain. As a result, β-sheets are poorly hydrated, have no void space, and are impenetrable to water. The poly(GA) regions form a similar structure, but with a different hydrophobic pattern. The glycine side chain is unable to form the same hydrophobic interactions as the A side chain resulting in fewer links in the β-sheet structure. Figure 6 depicts the comparison between poly(A) and poly(GA) domains.

Computer models for two antiparallel strands of the poly(A) and poly(GA) motifs are shown. Note a large vacant area in the poly(GA) when compared with the poly(A) strands.
A difference in hydrophobic pattern in the poly(GA) β-sheets compared to poly(A) β-sheets results in lower tensile strength of the minor ampullate silks (poly(GA) module) relative to the major ampullate silks (poly(A) module). The primary structure dictates the strength of interactions between β-sheets, which in turn determines in part the tensile strength of the silk fibers generated from these proteins. Poly(A) sequences have been shown to form a Poly(P) II helix in the unfolded state [12, 53], whereas in the folded state they form β-sheet aggregates [8]. These findings can explain the spider silk liquid state in the silk gland and its transition to a solid fiber.

NMR studies conducted by Holland and colleagues showed that water affects the mechanical properties of a silk fiber [34]. MaSp 1 and MaSp 2 have different interactions with water. After hydration significant chain motion occurs in the GGX regions and the alanine regions which have a tendency to form a helix (Figure 6). It is thought that the mobile A regions observed by NMR are able to participate in the connections between the crystalline and amorphous domains in a silk fiber [34, 54, 55]. The appearance of the mobile regions can also facilitate crystal reorientation in the silk fiber that is observed by fiber X-ray diffraction. It is thought that the GAG repeats that terminate the poly(A) runs are these mobile A regions [54]. The mobility of GAG repeats together with GGX repeats is an explanation for the increased extensibility and decreased stiffness observed for wet, supercontracted, dragline silks. It has been postulated that the mobility of these repeats is a reversible process [34]. The reversible conversion of GGX protein segments between the β-sheet and helical conformation provides the driving force for the supercontraction in MaSp1 silks [56]. However, in other silk types where GPGXX sequences dominate over GGX sequences, supercontraction is likely due to the recruitment of kinetically free (i.e. rubber-like) domains made from the GPGXX repeats. When β-sheets are formed, they have a certain size that also influences the mechanical properties of silks. Spin-diffusion NMR experiments estimate that the average poly(A) crystalline domain is about 6 ± 2 nm [34], in agreement with WAXS data on dry silks [21]. Additionally, the size of the β-sheet crystallites does not change after contact with water due to the nonpolar and hydrophobic nature of poly(A) that reinforce crystal stability by preventing water diffusion between stacks of β-sheets [57].

The second element, the GGX segment, has been proposed to form either a β-sheet or α-helix structure (Hayashi et al., 1999). FTIR and NMR data support the helical conformation of the GGX repeats [58]. The GGX segment is a tight three amino acid repeat that assumes a poly(G)PII helix based on molecular modeling [58]. Figure 7 depicts the computer model of the GGYGGGLGGQGGA repeat [12].

The hydrophobic core of this structure is composed of methylene groups, while tyrosine (Y) and glutamine (N) residues form the hydrophilic shell through hydrogen bonding. By forming the hydrophobic core, the G residues provide stabilization to fiber formation. The GGX helices can also serve as a link between crystalline β-sheets within the protein and between neighboring GGX helices in adjacent protein molecules facilitating fiber alignment [59]. The GPGXX segment found in MaSp2 is proposed to be responsible for the extensibility of dragline silks, forming β-turns that participate in the formation of a β-spiral [28]. Figure 8 depicts a computer model of the silk β-spiral. These proline-rich pentapeptide blocks contribute to the extensibility of spider silks and are largely responsible for the high toughness of MaSp silks. More studies are needed to fully understand the secondary structure of GPGXX blocks and its role in silk’s extensibility.

The β-spiral spring serves as the elastic mechanism in the silk fiber; namely, by simple extension and contraction of the β-spiral, a silk fiber exhibits elasticity. Hayashi et al. (1999) and Lewis (2006) suggest two key features of this model. First, the presence of P residues
provides focal points for the retraction of energy after stretching. Thus, when forcing the P
bonds to torque in response to extension, a large retraction force is generated. The second
key feature is that the positions of S and Y hydroxyls allow them to form hydrogen bonds
with downstream G residues, where the longer Y side-chains stabilize the tight β-turns and
the shorter S side-chains stabilize the layers of coils. The importance of these bonds is
supported by the strong tendency for Y and S residues to regularly alternate in flagelliform
silk. Only two silk types possess the GPGXX segment, the major ampullate spidroin 2 and
flagelliform silks [60]. Further support of the role of the GPGXX segment is provided by the
correspondence between the number of GPGXX repeats in one molecule and the different
extensibilities of the two silks. The wet major ampullate silk has up to 35% extension when
β-turns occur nine times in a row before being interrupted by another motif [5]. Flagelliform
wet silk has 200% extensibility and has at least 43 contiguously linked β-turns in its spring-
like spirals [36]. More experimental and computational studies are needed to fully
understand the secondary structure of GPGXX blocks and its role in silk’s extensibility.
Little is known about the structures formed by the spacer regions of the minor ampullate and
flagelliform silks. The only available information is that spacers are long, highly conserved
among spiders, non-repetitive and rich in serine [22]. Figure 9 summarizes the secondary
structure and core modules found in orb-weaver spider silks.

The primary sequence domains found in spider silks determine the biological, chemical and
physical properties of silk fibers. By knowing the role of each structural element, it is
possible to predict properties of silks. For example, by increasing the number of poly(A)
repeats in a given peptide the resulting fiber will likely have higher tensile strength than
those peptides containing poly(GA) repeats. The presence of the GPGXX motif give rise to
a β-spiral spring that is responsible for the extensibility of silk-like polymers. Finally, GGX
regions can be present as transition domains between different types of structural regions
facilitating self-assembly. In summary, it can be concluded that a few amino acid motifs or
domains found in spider silks give rise to the tremendous diversity of silks that has evolved
through the modification and rearrangement of these motifs. These few amino acid motifs
directly determine the complex array of mechanical properties exhibited by silk proteins.

4. Mechanical Properties of Spider Silks

The orb-weaving spiders produce a variety of high-performance fibers with mechanical
properties that outperform synthetic materials in the combination of strength and elasticity.
Dragline silks form extremely strong fibers. Dragline silk tensile strength is comparable to
Kevlar and it has extensibility which is seven times greater than the extensibility of Kevlar
[18, 61]. This combination of features makes major ampullate silk one of the toughest
materials known [5, 62]. Minor ampullate silk is similar to major ampullate silk in tensile
strength but has limited elastic behavior. However, flagelliform silk has outstanding
extensibility, stretching as much as 200% and, at the same time, it is a strong fiber [38].
Table 2 summarizes the strength, elasticity and energy to break values for known spider
silks along with selected biomaterials and man-made materials [63].

Spider silks are a useful fiber model to study relationships between sequence chemistry,
secondary structure and mechanical properties in protein-based materials and are excellent
candidates for biomaterials.

5. Spider Silk Processing – Artificial Modes to Mimic Nature

The development of genetic recombination techniques has opened the possibility to
reproduce the silk spinning process in an in vitro environment. However, many of the
desirable mechanical properties of the various spider silks are a result of the higher order
protein structures [64] that result from specific processing of the primary protein sequences that occurs in the silk gland. Therefore, reconstitution of primary spidroin protein sequences into functional fibers has emerged as a topic of great interest for many materials and tissue engineering applications where the high strength, extensibility and toughness intrinsic to natural silk are ideal. This challenge has led to efforts to adapt processing techniques to mimic the natural spinning process in order to promote improved protein configuration in the secondary structure, which primarily refers to the formation and alignment of β-sheets. Techniques that simulate the mechanistic changes that occur during the spinning process are largely performed by rate-controlled extrusion directly through a syringe. Figure 10 depicts a light microscope bright field image of fibers spun from recombinant spider silk. Important elements that improve overall fiber formation and β-sheet quantity and alignment are common among the many variations of this basic extrusion method.

First, initial solvents of the spidroin proteins need to exhibit strong hydrogen bonding characteristics to improve solvent-protein interactivity and solubility, making distilled water and hexafluoroisopropanol (HFIP) good candidates. Improved solubility allows for higher concentrations of dope, which has shown greater success in fiber formation [65, 66]. Next, shearing of the silk dope with the wall of the tubing through which the dope is flowed initiates protein alignment, which has been shown to be a major factor in the spider major ampullate glands [67]. Then, extrusion into a miscible organic solvent, such as methanol or isopropanol, has been shown to help induce fiber formation through dehydration of the silk sequence [68, 69]. And lastly, drawing the newly formed fiber to cause strain elongation has shown to induce higher β-sheet content and improve overall mechanical properties of the silk including elastic modulus, toughness, elongation to failure, and ultimate strength [70, 71]. Current applications using these techniques have produced fibers with diameters in the range of 10–60 μm and ultimate strengths between 60–160 MPa, which is still lacking in comparison against native major ampullate fibers with diameters in the range of 2.5–4 μm and failure strengths typically in the 1–2 GPa range.

While these drawn synthetic fibers exhibit improved mechanical properties for synthetic sequences, these values are still well below the measured properties of the native structures, which encourages the discussion of ways to improve the structural integrity of synthetically derived spidroin sequences. Some strategies have included elongating the polymeric repeat structure of the hydrophilic and hydrophobic blocks in synthetic constructs to promote greater chain-chain interactivity. Increasing hydrophobic chain length has been modeled and reported to increase β-sheet crystallinity at the molecular level [72] and experimental evidence has been presented that show stronger materials resulting from longer polymeric sequences [70]. Other strategies include the incorporation of the characteristic pH gradient of the natural spinning duct using microfluidics [71]. This offers the added advantage of microfluidic control over final fiber dimensionality based on laminar flow parameters of the spinning device, which additionally mimic the volumetric tapering of the spinneret. However, this technique must be balanced against the loss of shearing at the boundary of the channel that is important in initial fiber formation.

Apart from techniques to mimic spinning gland mechanisms in an effort to generate biomimetic micron-scale fibers, there is also interest in using silk as a functional surface itself. This is accomplished through electrospinning the purified spidroin or fibroin dopes onto a surface, which generates a nonhomogeneous fiber mesh comprised of submicron features [73]. This technique is appealing for biomaterial applications because of the feasibility of electrospinning and the fiber mesh customization that is possible. Figure 11 depicts electrospun fibroin.
Modifications, such as the addition of other proteins into the silk dope, can affect the bulk characteristics of the mesh. Some reported examples include the addition of carbon nanotubes in order to increase fiber elastic modulus [74], addition of poly(lactic acid) (PLA) and poly(glycolic acid) (PGA) to incorporate more hydrolysable motifs [75], and incorporation of various growth and/or chemical factors for applications in neuron [76, 77], cardiovascular [78], skin [79], and bone [80] engineering to name a few. Fibers generated in this fashion range in diameter from 0.2–2.5 μm, which are a closer approximation dimensionally than extruded fibers, but proportionally lack the extruded fibers’ tensile strength with failures occurring in the 0.5–4 MPa range.

To summarize, fiber generation can be tuned based on the scaling and properties desired with fiber diameter ranges from hundreds of nanometers to tens of microns. The former is yielded in an unwoven mesh of nano- to micro-scale fibers that has a variable, collective behavior based on spinning composition. The latter is yielded in discrete, individual fibers that are able to be manipulated by hand. Many of these techniques are universally applicable to the known varieties of silk; however, more investigative effort has been invested in optimizing techniques that employ Bombyx mori fibroin proteins. While there are strong correlates and results that have bearing for spider fibroin and spidroin-based fiber processing applications, it may be necessary to focus specifically on the unique mechanisms that differentiate the two silk-spinning organs and organisms in order to maximize the properties of the processed materials for ongoing and future synthetic spidroin-based fibers and fiber networks.

6. Spider Silks in Biomaterial Design

6.1 Spider Silk Block Copolymers

Block copolymers are a fascinating family of polymeric materials usually prepared by the covalent bonding of two or more thermodynamically incompatible polymeric or oligomeric chains that give rise to a variety of nanostructure morphologies in solution and in bulk [81]. Nanostructure formation by synthetic block copolymers is the focus of many studies, while biologically derived systems have been less studied [82]. Silks are natural block copolymers with a characteristic crystalline β-sheet secondary structure encoded by hydrophobic peptide blocks. β-sheets form through natural physical crosslinks of amino acid sequences, which in spider and silkworm silks consist of multiple repeats of mainly alanine, glycine-alanine, or glycine-alanine-serine, as described in some detail earlier in this review. The hydrophilic, non-crystalline regions of silk commonly consist of: (i) β-spirals similar to a β-turn composed of GPGXX repeats (where X is mostly glutamine) and (ii) helical structures composed of GGX [12]. Inspired by the extraordinary material properties of spider silks, a family of spider silk block copolymers was designed and bioengineered based on the assembly of individual spider silk domains, in particular poly-alanine (A) and glycine-rich (B) domains, that display different phase behavior in aqueous solution [83, 84]. The interplay between the silk block copolymer sequence, composition, length and self-assembly behavior was assessed to build materials with precise structural and morphological features [83]. The A block consisted of one polyalanine/polyglycine repeat (GAGAAAAAGGGAG) that is responsible for β-sheet formation. The B block was composed of four GGX repeats, separated by the GSQGSGR sequence. The GGX repeats adopted a helical conformation and served as a hydrophilic link between crystalline β-sheet regions as well as neighboring GGX helices in adjacent protein molecules that helped reinforce fiber alignment. Secondary structure and morphology were assessed by FTIR and SEM, respectively. Based on these analyses, trends in block copolymer assembly behavior into specific morphologies were observed as a function of the number of hydrophobic blocks (A) and solvent effects. In terms of structure, as the size of the hydrophobic block increased, the content of β-sheet increased in the silk block copolymers. In terms of morphological features, the increase in

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hydrophobicity (increase in the number of A blocks) was connected with a transformation from thin films to micelles (1–3 μm) and finally to large compound micelles (50 μm) in water. In terms of solvent selection, when 2-propanol was the solvent, a transformation from thin films to nanofibers (50–200 nm) and large compound micelles was observed (Figure 12). Further studies characterized the thermal properties and demonstrated the self-assembly of monolayers of spider silk-like block copolymers [85]. At high surface pressure, spider silk block copolymers with minimal hydrophobic domains (BA) formed oblate structures, whereas block copolymers with a 6-fold larger hydrophobic domain (BA6) formed prolate structures [86]. The study provided a basis for future designs of biomaterials based on spider silk chemistry, with controlled structure-architecture-function relationships. In addition, the observed nanofiber and compound micelle architectures are potentially useful for biomedical applications and could be employed as drug delivery reservoirs, tissue scaffolds and environmental friendly polymers.

Recently, a pH-responsive silk-like block copolymer with a sequence of [(GA)_3GE(GA)_3GL]_{28} was produced in the yeast *Pichia pastoris* at the g/L level [87]. As a model peptide, a silk-like peptide ([GA]_{3}GE([GA]_{3}GE)_{24}) known to be water soluble above the isoelectric point and formed insoluble stacks of β–sheets in a dried state [88]. Every second octapeptide had a hydrophobic L residue in place of E. This change made the peptide more amphiphilic [87]. The block copolymer formed fibers upon crystallization in 70% (v/v) formic acid under vapor diffusion of methanol, as shown by AFM, with the average fibril height of 2.7 nm and width of 49 nm. The CD spectrum of the negatively charged polymer in aqueous solution at high pH indicated a dominance of random and extended helical (silk III-like) structures, whereas the spectrum of a coating prepared from such a solution showed a conformation rich in β-turns. Given the biocompatibility of silk-like materials, the polymer may be of interest for biomedical applications, such as the coating of surgical implants or pH-responsive controlled drug release.

The construction of a biosynthetic multiblock protein polymer based on the sequences of dragline silk from *A. diadematus* [(AEAEAKAK)_2AG(GPGQQ)_6GS]_{9} was also reported [89]. The block copolymer spontaneously formed self-supporting macroscopic films via rearrangement of segments within the polypeptide from α-helices to β-strands. The primary structure of the polymer consisted of the amphiphilic peptide, (AEAEAKAK)_2, that was capable of adopting a β-strand conformation over a wide range of pHs and temperatures. The glycine-rich segment, GPGQQ, was derived from *A. diadematus* dragline silk. The central PG unit in the glycine-rich block had a high propensity for the formation of type II β-turns. High resolution FESEM of cryo-immobilized, water-swollen membrane revealed a network of fibrils, approximately 10–20 nm in diameter, interspersed within a less structured matrix.

The formation of β-sheet structures induced irreversible aggregation of [(AEAEAKAK)_2AG(GPGQQ)_6GS]_{9} polypeptide into a hydrogel network through interstrand hydrogen bonding interactions between chain segments [89]. The influence of the polypeptide structure on the physical properties of silk-like block copolymers using gel electrophoresis under denaturing conditions was investigated. The authors observed that the number of repetitive units, as well as side chain modifications (i.e., glycosylation, phosphorylation), significantly affected gel mobility. For example, negatively charged and carbamylated peptides showed low gel mobility then compared with unmodified peptides, positively charged and neutral polypeptides [90]. Ultimately, this block copolymer approach can provide a foundation for the design of highly engineered biomaterials for a range of important applications.

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6.2 Spider Silk/Silkworm Fibroin Hybrids

Silk-based biomaterials have attracted interest due to their mechanical and self-assembling properties in the biomaterial field. A unique combination of high elasticity, toughness, and mechanical strength, along with biological compatibility and biodegradability make silks advantageous candidates [59, 91, 92]. As mentioned earlier in this review, dragline silk from orb-weaving spiders is a copolymer of two large proteins, major ampullate MaSp1 and MaSp2. The ratio of these proteins is known to have large variation across different species of orb-weaving spiders resulting in different mechanical properties of silk fibers. For example, *N. clavipes* major ampullate silk contains 80% of MaSp1, whereas major ampullate silk of *A. aurantia* is comprised of only around 40% MaSp1 [3, 29]. The question of how the ratio of MaSp1 and MaSp2 influence fiber formation and its mechanical properties has been studied, including using engineered MaSp1/MaSp2 mixed fibers as well as chimeric silk fibers from MaSp1 and MaSp2 sequences in a single protein [93]. Spun fibers were stretched in a 75% isopropanol/water solution, as the as-spun fibers were semi-soluble in water. The proteins with a higher molar ratio of the polyalanine motif formed fibers with higher tenacity and Young’s modulus (about 10 MPa higher in tenacity, 0.4 GPa higher in Young’s modulus, for stretched and as-spun fibers). The protein that contained more GPGXX motifs were more extensible [93].

In a similar study two chimeric proteins based on the sequences of Flag and MaSp2 silks of *N. clavipes* were produced [14]. The designed was based on dragline MaSp2, but (GPGQQ GPGGY)2 was replaced with one of two Flag native variants. These chimeric Flag/MaSp2 chimeric proteins provided interesting self-assembly abilities in aqueous solution, triggered by heat and/or shear, depending on the nature of the Flag structural motif. Both of these proteins formed synthetic fibers in organic or aqueous conditions by extrusion or fiber-drawing from self-assembled films.

In a recent effort to develop tough fibers, transgenic silkworms encoding chimeric silkworm/spider silk proteins were produced using piggyBac vectors [94]. The vector that was previously used by the Tamada group [95] included the *Bombyx mori* fibroin heavy chain promoter and enhancer, a gene sequencing encoding a 78 kDa synthetic spider silk protein, and an enhanced green fluorescent protein (EGFP) tag. Strong EGFP signals were observed under a fluorescence microscope (Figure 13). The composite fibers were tougher than the parental silkworm silk fibers and as tough as native dragline spider silk fibers.

These results demonstrate that silkworms can be engineered to manufacture composite silk fibers containing stably integrated spider silk protein sequences, which significantly improved the overall mechanical properties of the silk fibers.

6.3 Functionalization of Spider Silks with Fusion Peptides

Biominalization is the process found in nature by which biological organisms produce mineralized tissues such as bones, diatoms, shells and teeth [91, 96]. In biological organisms organic molecules exert remarkable level of control over the process of mineral formation, including the hierarchical organization and functional properties [97]. Through the use of recombinant DNA technology, a variety of novel materials can be engineered with properties not present in nature [81]. Since spider silks have desirable self-assembly characteristics, they can be used as molecular building blocks for the core structure together with desirable inorganic-binding peptides tailored for practical applications. Silk-based materials have been used as organic scaffolds for biominalization of hydroxyapatite [98, 99], silver [100], silica [96, 101–103], and titanium [104].
Detailed control of silica morphology and distribution on the surface of the silk films was achieved by fusing *N. clavipes* dragline sequences and the silaffin-derived R5 peptide of the diatom (*Cylindrotheca fusiformis*) [96, 101, 102, 105]. Three-dimensional porous networks were formed with clustered silica nanoparticles and single silica nanoparticles on the surface of chimeric spider silk films. The silk served as the organic scaffolding to control material stability. The silk/silica biomaterials are suitable for different tissue regenerative applications. For example, human bone marrow derived mesenchymal stem cells subjected to osteogenic differentiation were used to evaluate the influence of the silk–silica composite systems on osteogenesis [106]. The silica in these silk films influenced osteogenic gene expression, with upregulation of several osteogenic markers (i.e., alkaline phosphatase, bone sialoprotein, and collagen type I). Furthermore, calcium containing deposits were observed on silk films treated with silica as a strong indication of bone formation [105]. Benton et al. (2012) investigated the influence of the silk/chimera ratio through the adjustment of the type and number of silicifying domains) together with the effect of pH (3–9) on reactivity. The driving force for fast silicification was likely the water reduced environment of spider silk protein domains, making the removal of water from the silica condensation reactions energetically more favorable and resulting in the rapid formation of large silica particles (800 nm and 2 μm, dispersity of ±10%) in the presence of the silk fusion proteins [103].

Recently, Currie *et al.* reported the production of new recombinant silk-silver binding chimeric proteins capable of forming silver nanostructures from a AgNO₃ solution on silk films [100]. Two silver binding peptides, Ag-4(NPSSLFRYLPSD) and Ag-P35(WSWRSPTPHVVT), were previously identified by phage display [107]. The peptides were fused with recombinant spider silk 6-mer and 15-mer proteins originated from MaSp1 of *N. clavipes*. The formation of silver nanostructures (nanoparticles and nano-rods) from a silver nitrate solution was observed with the average particle size of 35.5 nm ± 8.5 nm, average rod diameter of 57.2 nm ± 17 nm and length 226 nm ± 82 nm for 6-mer Ag-4 and 15-mer Ag-4, respectively. The antimicrobial capacity of the silver treated protein films was also examined by incubation with both Gram-positive and Gram-negative bacteria. The silver treated silk films significantly inhibited the growth of both bacterial species. The incorporation of an inorganic phase through mineralization of a silk based substrate establishes new materials which incorporate the mechanical strength, biocompatibility and aqueous processing of silk with added functions due to the chimeric mineralizing domains encoded in the new bioengineered proteins. In a different approach, silk fibroins from *B. mori* were chemically grafted with titanium binding peptide to increase adsorption of these chimeric proteins to metal surfaces [104], indicating utility for new composite materials.

Spider dragline silk of *N. clavipes* was fused with a hydroxyapatite binding peptide from dentin matrix protein I (DMP1) via a genetic approach [99]. Calcium hydroxyapatite (HA), Ca₁₀(PO₄)₆(OH)₂, is central in bone and dentin formation. Both collagenous macromolecules and non-collagenous proteins such as phosphoproteins, osteonectin and dentin matrix protein I are involved in controlling the nucleation and growth of HA crystals in these tissues. DMP1 has two acidic domains ESQES (residues 386–390) and QESSEQDQS (residues 414–422) that control Ca²⁺ binding and play a critical role in HA nucleation [108]. The resulting silk-DMP1 chimeric protein demonstrated successful nucleation of HA crystals on the surface of silk films confirmed by FESEM coupled with energy dispersive X-ray spectroscopy.

A different approach was taken by Lee and coworkers (2009), where multiple pulsed vapor-phase infiltration was used to improve the mechanical properties of spider dragline silks by direct metal incorporation into inner regions of the protein fiber structures. The result was the formation of a metal-incorporated protein matrix. The authors demonstrated that metals can be intentionally infiltrated through multiple pulsed vapor-phase processes performed...
with equipment conventionally used for atomic layer deposition. Pulsed vapor-phase incorporation of zinc, titanium and aluminum into spider dragline silks increased the strength and toughness of the material [109].

Future Outlook

The majority of knowledge about structure-function relationships in spider silks comes from three types of spider silk proteins: major ampullate spidroins, minor ampullate spidroins, and flagelliform silks. It has been possible to relate protein sequences to molecular structures and to the physical properties of native fibers. Currently much effort has been made to unravel the features of other types of silk such as aciniform, tubuliform, and pyriform. In addition, the majority of work has been focused on orb-weaving spiders. Hence, it is necessary to broaden attention to other types of spiders (e.g. cob-weaving spiders), and other members of the class Arachnida. A recent discovery of a novel silk system in the tube-building corophioid amphipod Crassicorophium bonellii is an example of these possibilities [110]. Natural silks have potential for a variety of material science and biomedical applications [111]. Over the last decade there has been considerable progress in understanding the genetic organization encoding spider silks. Cloning and expression for spider silk genes have improved, and the self-assembly and processing of spider silk into many material formats is now better understood. Just recently a native-sized (285 kDa) recombinant protein of the spider Nephila clavipes was produced and spun into a fiber displaying mechanical properties comparable to those of the native silk, indicating a breakthrough in standard recombinant production of spider silks [112]. Moreover, transgenic silkworm/spider silk production systems have been developed to produce tough fibers. It is possible to mix and match key modules via recombinant approaches, providing additional insights into the role of individual modules and effects of neighboring elements on properties. This approach should lead to the development of custom structures built from specific silk elements. Future challenges will include scaling up silk production. In addition molecular-level analysis of the nanostructure of silk assemblies using in silico approaches should play an important role to understand structure-property-function relationships.

Acknowledgments

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References


Figure 1.
(a) Adult female Nephila Clavipes (golden-orb) spider provided by the Miami Metrozoo, Florida. (b) (A) Dissected major ampullate (MA) gland of the spider. The ~1 μl blob (B) protruding through a rupture of the gland wall near the spinning canal (C) was used for the rheology experiments. Modified with permission from [113].
Figure 2.
The natural spinning process. (a) Illustration of a spider’s spinning gland divided into four parts. (b) Schematic model of the silk fiber assembly mechanism occurring along the spinning apparatus. Reproduced with permission from [3].
Figure 3.
Silk glands, silk types and silk uses of *Nephila clavipes*. Reproduced with permission from [114].
Figure 4. *Nephila clavipes* MaSp1 repeating units. Reproduced with permission and adapted from [27].
Figure 5.
Consensus amino acid sequence of minor ampullate silk protein 1 and 2 from *N. clavipes*. Adapted with permission from [115] and GenBank accession no. AAC14589.

\[
\text{MiSp 1} \\
\left[\text{GAGGAGGYGRGAGAGAAGAGAGAGGYYGGYGGGAGAGAGAAAAAG} \right]_n \text{Spacer} \\
\text{Core Units: } \left(\text{A}\right)_n, \left(\text{G/A}\right)_n \text{ GGX, spacer}
\]

\[
\text{MiSp 2} \\
\left[\text{GAVGAAGAAAGFAAGAGGAGGYR} \right]_n \text{Spacer} \\
\text{Core Units: } \left(\text{A}\right)_n, \left(\text{G/A}\right)_n \text{ GGX, spacer}
\]
Figure 6.
Computer models of the poly(A) and poly(GA) segments; Reprinted with permission from [28].
Figure 7.
Computer model of the GGX repeat region; Reprinted with permission from [28]. The model is a space-filling energy-minimized antiparallel two-strand GGS region. The starting configuration was a glycine II helix for both strands.
Figure 8.
Computer-generated model of a silk β-spiral; Reprinted with permission from [28] and 2012.igem.org. Computer-generated model of a pair of GPGQGPGY repeats is shown. Hydrogen bonds (shown as dashed lines) occur between residues in the same β-turn and between different β-turns.
Figure 9.
Structural motifs and their secondary structures. Reproduced with permission from [116]. The empty box marked ‘?’ indicates that the secondary structures of the ‘spacer’ motifs are unknown. MaSp1 or MaSp2: major ampullate spidroin 1 or 2; MiSp: minor ampullate spidroin; Flag: flagelliform protein.
Figure 10.
Representative light microscope bright field image of fibers spun from recombinant spider silk in both demonstrating the effect of drawing on fiber dimensions. Scale bar is 30 μm. Reprinted with permission from [70]. Copyright 2011 American Chemical Society.
Figure 11.
Electrospun fibroin demonstrating the nanoscale and interconnected mesh generated. Adapted with permission from [73]. Copyright 2002 American Chemical Society.
Figure 12.
Diverse molecular assemblies of spider silk-like block copolymers in water (A, C) and 2-propanol (B). A block is represented by poly(A) repeats and B block is composed of GGX repeats.
Figure 13.
Expression of the chimeric silkworm/spider silk/EGFP protein in (A) cocoons, (B and C) silk glands, and (D) silk fibers from spider 6-GFP silkworms. Reproduced with permission from [94].
Table 1

Comparison of the ensemble repeats and core units in spider silk proteins. Only one representative internal repeat unit for each fibroin type is illustrated from either orb- or cob-weavers. Color code is based on the hydrophobicity of the amino acids.

<table>
<thead>
<tr>
<th>Species</th>
<th>Protein</th>
<th>Ensemble Repaets</th>
<th>Core Units</th>
<th>References</th>
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</thead>
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<td>N. clavipes</td>
<td>MaSp1</td>
<td>(A)n, (G/A)n, GGX</td>
<td>(A)n, (G/A)n, GGX, GPGXX</td>
<td>Xu, 1990</td>
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<tr>
<td>N. clavipes</td>
<td>MaSp2</td>
<td></td>
<td></td>
<td>Hinman, 1992</td>
</tr>
<tr>
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<td>(A)n, (G/A)n, GGX, spacer</td>
<td>(A)n, (G/A)n, GGX, spacer</td>
<td>Colgin, 1998</td>
</tr>
<tr>
<td>N. clavipes</td>
<td>MiSp1</td>
<td></td>
<td></td>
<td>Colgin, 1998</td>
</tr>
<tr>
<td>N. clavipes</td>
<td>Flag</td>
<td>GGX, GPGXX, spacer</td>
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<td>AcSp1</td>
<td>(S)n, TGPSG</td>
<td></td>
<td>Hayashi, 2004</td>
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<tr>
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<td>HHX</td>
<td>Tian, 2006</td>
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<td>L. hesperus</td>
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<td>HHX</td>
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<td>L. hesperus</td>
<td>SCP2</td>
<td></td>
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Table 2

Comparison of mechanical properties of spider silks. Modified with permission from [12] and [63].

<table>
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<th>Material</th>
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<th>Elasticity (%)</th>
<th>Energy to break Jkg⁻¹</th>
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<td>35</td>
<td>1×10⁴</td>
</tr>
<tr>
<td>Minor ampullate silk</td>
<td>1×10⁹</td>
<td>5</td>
<td>3×10⁴</td>
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<tr>
<td>Flagelliform silk</td>
<td>1×10⁹</td>
<td>&gt;200</td>
<td>1×10⁵</td>
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<tr>
<td>Kevlar</td>
<td>4×10⁹</td>
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<td>3×10⁴</td>
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<td>Rubber</td>
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<td>Nylon, type 6</td>
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