Development of New Smart Materials and Spinning Systems Inspired by Natural Silks and Their Applications

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Silks produced by spiders and silkworms are charming natural biological materials with highly optimized hierarchical structures and outstanding physicomechanical properties. The superior performance of silks relies on the integration of a unique protein sequence, a distinctive spinning process, and complex hierarchical structures. Silks have been prepared to form a variety of morphologies and are widely used in diverse applications, for example, in the textile industry, as drug delivery vehicles, and as tissue engineering scaffolds. This review presents an overview of the organization of natural silks, in which chemical and physical functions are optimized, as well as a range of new materials inspired by the desire to mimic natural silk structure and synthesis.

Keywords: silk, protein primary sequence, spinning, hierarchical structure, self-assembly, biomimetic, silk-inspired functional materials

INTRODUCTION

Over the last 5000 years, silk has been widely studied and used in the textile industry; however, many lessons may still be learned from this outstanding natural biological material. A remarkable feature of natural silk is its outstanding toughness, which is superior to that of any other artificial fiber (Liu et al., 2005; Keten et al., 2010; Omenetto and Kaplan, 2010; Porter et al., 2013; Tokareva et al., 2014). Most researchers believe that the toughness of silk originates from the combined action of the alternating hydrophobic and hydrophilic pattern in the protein primary sequence and the unique spinning process used to prepare the silk fibers. Spiders and silkworms generate a variety of silk fibers. Among the known silk variants, the silk of Bombyx mori (B. mori) and the orb web spider dragline silks of Nephila clavipes and Araneus diadematus are the most widely used one. These particular silks have inspired researchers to develop new smart materials, which is the main topic of this review.

Although B. mori silks are not as tough as spider dragline silks, the high yield of these silks through domesticated production has facilitated the use of these silks in a variety of fields, from luxury fabrics to biomedicine, since their development in China thousands of years ago. Spider dragline silk is thought to be the toughest biological material yet identified; however, limitations on methods for mass production of spider silk have contributed to a gap between applications of silkworm silk and spider silk. Researchers recently developed a series of recombinant DNA technologies that enable spider silk production of the large scale (Tokareva et al., 2013). Recombinant technologies permit genetic modifications that can be imparted on the functionalities of spider silks in addition to their extraordinary mechanical properties (Schacht and Scheibel, 2014). Such advances in biotechnology
have led to a flourishing of spider silk applications that have narrowed the gap with silkworm silk applications (Schacht and Scheibel, 2014).

The hierarchical structures of natural silks are crucial to the silks’ strength. The design of materials with hierarchical structures similar to that of silk may contribute to the development of novel high-performance functional materials. Silk proteins are hierarchical amphiphilic block proteins composed of alternating hydrophobic and hydrophilic blocks. Hierarchical amphiphilic patterns drive silk proteins to self-assemble into micelles and play a critical role in silk protein storage inside the glands (Jin and Kaplan, 2003; Askarieh et al., 2010; Hagn et al., 2010; He et al., 2012; Schwarze et al., 2013). Therefore, the synthesis of silk proteins similar to the natural proteins is a key issue in material science. Another critical aspect of silk engineering is the spinning system used to prepare the fibers. Such systems are typically designed to mimic natural spinning mechanisms. The tapered spinning duct of silkworm or spider provides an elongational field during spinning process, which orients the silk protein micelles along the direction of the duct via a shear force, leading to the formation of β-sheet crystals. The fibers are then drawn away from the spinning ducts either by the legs of the spider or by the “figure eight” movement of the silkworm head to further orient the silk proteins and form β-sheets. The as-spun silk fibers are composed a hierarchical system of β-sheet crystal domains and α-helix/turn non-crystalline domains, which convey to the silk fibers non-linear material properties and outstanding mechanical properties. Although several reviews of synthetic silk protein morphologies have been published (Altman et al., 2003; Vepari and Kaplan, 2007; Kim et al., 2010, 2014; Rockwood et al., 2011; Tao et al., 2012; Kundu et al., 2014), new materials inspired by silk protein and various spinning methods mimicking spider and silkworm have not been reviewed, despite their importance and extensive applications. In addition, how silk fibers assemble into 3D hierarchical macroscale structures (cocoons and webs) to meet various needs by silkworms and spiders are not summarized until now.

This review presents an overview of the beautiful organizational structures of natural silks from the nanoscale (protein sequence), microscale (spinning) to macroscale (cocoons and webs) aspect and introduces new materials and processes inspired by nature. As shown in Figure 1, the natural silk is produced from the synthesis of silk protein, spinning by shear force and formation of web and cocoon, and understanding of sequence–structure–property interplay is critical in developing smart materials inspired by the spider and silkworm. Here, we described a variety of new materials and applications inspired by silk’s hierarchical structures and silk spinning process.

NEW SYNTHETIC NANOMATERIALS INSPIRED BY THE SILK PRIMARY SEQUENCE AND DIRECTED SELF-ASSEMBLY

Nanomaterials are potentially useful in a variety of applications (Zhang, 2003). As scientists have dreamed of designing nanoscale material macrostructures through bottom-up methods, they have studied mimics of the self-assembly processes used by natural systems, such as silkworms or spiders. The self-assembly of silk inside the gland and spinning duct is a marvelous process. Silk proteins are produced inside glands and stored in relatively high concentrations of up to ~50% w/w without aggregation. Self-assembly among silk proteins into highly ordered micelle cluster structures protects the silk proteins from aggregation. The silk proteins are amphiphilic and consist of alternating hydrophobic and hydrophilic domains with highly conserved C- and N-terminal domains. The hydrophobic interactions between hydrophobic domains constitute the main force driving self-assembly (Jin and Kaplan, 2003). The small hydrophilic domains may remain hydrated inside the micelles, and the terminal domains define the micelle edges, stabilizing the solubilized proteins (Hagn et al., 2010). The hydrophobic domains are dominated by a hydrophobic sequence (GA)n or An, which enables protein chain self-assembly to form β-sheet-rich crystal structure and gives silk fibers their outstanding mechanical properties. Stimuli-responsive C- and N-terminal domains play a role in fiber formation under shear force. The primary silk protein sequence has been introduced into artificial polypeptides/peptides and polymer–peptide hybrids in an effort to direct self-assembly and prepare supramolecular hierarchical architectures (Table 1). This section describes the design of diverse supramolecules based on silk proteins.

Polypeptide/Peptide Design Inspired by Silk Proteins

Pure Silk-Like Polypeptides with a Lower Molecular Weight

Recombinant forms of the major native ampullate dragline spidroin 1-like proteins, with a molecular weight range of 250–320 kDa, have been successfully overexpressed in the host Escherichia coli BL21 (DE3), and strong fibers have been fabricated using these synthesized proteins (Xia et al., 2010). However, the synthesis of silk-like proteins comparable in size to the native silk protein still poses significant challenges to many material research groups. The most famous spider silk protein, the dragline silk protein of A. diadematus fibroin and N. clavipes fibroin, has received its fame through intensive investigations and recombinant genetic engineering redesign. Spider dragline silk proteins are great models and guides for the design of novel polypeptide. The spider dragline silk of N. clavipes contains at least two proteins: the major ampullate dragline spidroin 1 (MaSp1) and the major ampullate dragline spidroin 2 (MaSp2). The spider dragline silk of A. diadematus also contains two major proteins: A. diadematus fibroins 3 and 4 (ADF-3 and ADF-4). These dragline spidroins are composed of two alternating peptide motifs: a crystalline [An or (GA)n] motif and a less crystalline (GGX or GPΓXX) motif (Scheibel, 2004). The An and (GA)n motifs form β-sheet crystal structures to provide a high tensile strength and stiffness in the dragline silk. On the other hand, the GPΓXX motif forms type II β-turn structures, and the GGX motif forms a 3₂-helix, both of which convey extensibility to the dragline fiber (Scheibel, 2004; Tokareva et al., 2013). With this understanding of the interplay between the sequence and the secondary structure...
properties, scientists have designed a series of spider silk-like polypeptides and peptides. The repeating motifs of ADF-3 and ADF-4 have been multimerized through recombinant DNA technology to obtain an ADF-like polypeptide (Huemmerich et al., 2004; Rammensee et al., 2006, 2008). The ADF-4-like polypeptide (called C16) with a molecular mass of 48 kDa self-assembles into nanofibers in the presence of methanol. These nanofibers then further self-assemble to form a hydrogel (Rammensee et al., 2006). The same artificial polypeptide may be used to fabricate both microspheres and nanofibers by inducing self-assembly in the presence of a kosmotropic agent, such as potassium phosphate (Slotta et al., 2007). C16 also can self-assemble to form a thin film at an oil–water interface and encapsulate active agents inside a water phase (Hermanson et al., 2007). C16 adsorbs onto oil–water interface, and a film or microcapsule is formed rapidly due to a secondary structure transition undertaken by C16 at the interface (Figure 2A). These encapsulates can be released rapidly by digesting the C16 film with proteinase K (Hermanson et al., 2007). The ADF-3-like polypeptide and ADF-3’s conserved non-repeating region at the C-terminus has been recombinantly produced in an effort to probe the natural spider spinning process (Rammensee et al., 2008; Hagn et al., 2010).

![FIGURE 1](image_url) | Overview the inherent properties of natural silks produced by silkworm and spider, including the structural motif pattern of protein primary sequence, unique spinning process, and macroscale 3D hierarchical structure. Spider image: reprinted with permission from Kang et al. (2011). Copyright 2011 Nature Publishing Group; Silkworm image: reprinted with permission from Chae et al. (2013). Copyright 2013 John Wiley and Sons.

**Silk Peptide Fusion with Other Functional Peptide Sequences**

Spider silk proteins have been fused with a variety of functional peptides, including the integrin-binding motif RGD (Leal-Egana and Scheibel, 2012; Leal-Egana et al., 2012; Wohlrab et al., 2012), the cell-binding peptide IKVAV (Graf et al., 1987; Grant et al., 1989; Widhe et al., 2013), antimicrobial peptides...
TABLE 1 | Summary of the self-assembled nano/microstructures based on silk peptide sequence and their potential applications.

| Sequencea–c | Structures | Media | Potential applications | Reference |
|-------------|------------|-------|------------------------|-----------|
| Peptide/polypeptide | GSSAAAAASGPYGGPGPGPSGPG | Nanofiber/hydrogel | Methanol/water | Tissue engineering | Rammensee et al. (2006) |
| | GYGGPGPGPGPGPGPGPG | Sphere | Potassium phosphate | Drug delivery | Slotta et al. (2007) |
| | GSSAAAAASGPYGPGPGPG | Capsule | Water/oil interface | Encapsulation | Hermanson et al. (2007) |
| | GYGGPGPGPG | Nanoparticle | Water | Gene delivery | Numata and Kaplan (2010) |
| | GSSAAAAASGPYGGPGPG | Nanoparticle | Water | Gene delivery | Numata et al. (2011) |
| | [GVGVP]4(GGGVP)(GVGVP)3 | Nanoparticle/ nanofiber | Water | Controlled release | Xia et al. (2011) and Xia et al. (2014) |
| | [GVGVP]3(GGGVP)(GVGVP)3 | Nanoparticle/ nanofiber | Water | Controlled release | Xia et al. (2011) and Xia et al. (2014) |
| | H10-[((GVGVP)4(GGGVP)(GVGVP)3 | Micelle | Water | Biosensor/drug delivery | Lin et al. (2014) |
| Peptide–polypeptide hybrid | (GVGVP)3(GGGVP)16 | Injectable hydrogel | Water/in vivo | Gene delivery | Megeed et al. (2002) and Megeed et al. (2004) |
| | (GVGVP)3(GGGVP)16 | Micelle/nanofiber/hydrogel | Water | Target delivery/tissue engineering | Martens et al. (2009), Beun et al. (2012), Beun et al. (2014), and Golinska et al. (2014) |
| | (GVGVP)3(GGGVP)16 | Nanotube/nematic lamellar/nanorod/nanofibril | Water | Nanofabrication/protein misfolding research/antibacterial peptide | von Maltzahn et al. (2003), Bucak et al. (2009), Liu et al. (2009), Wang et al. (2009), Castelletto et al. (2010), Chen et al. (2010), Cenker et al. (2011), and Cenker et al. (2014) |
| | (GVGVP)3(GGGVP)16 | Nanofiber | Water | Controlled release | Xia et al. (2011) and Xiao et al. (2014) |
| | (GVGVP)3(GGGVP)16 | Capsule | Water | Drug delivery | Slotta et al. (2007) |
| | (GVGVP)3(GGGVP)16 | Tape/nanofiber/interpenetrating supramolecular network | Nanofiber/hydrogel | Drug delivery/tissue engineering | Shah et al. (2010) |

1Red-bolds denote silk protein sequence.
2Single amino acid abbreviations used: A, Ala; G, Gly; H, His; K, Lys; L, Leu; M, Met; N, Asn; P, Pro; Q, Gln; R, Arg; S, Ser; and V, Val.
3Non-functional amino acid sequence domains at N- and C-terminus are omitted for simplicity. GXY X and Y positions can be any residue but are frequently occupied by Pro and Hyp.

(Gomes et al., 2011), silver ion-binding peptides (Currie et al., 2011), and silica-binding peptides (Wong Po Foo et al., 2006). These hybrids may be applied as a film and coating material to improve the surface properties of biomaterials. Silk protein-based biomaterials prepared via top-down methods have been reviewed extensively elsewhere. Here, we discuss supramolecular assemblies formed by silk-functional peptide fusions. Numata and Kaplan (2010) reported the use of a silk–polylsine–ppTG1
fusion polypeptide as gene carrier. The consensus repeat sequence (SGRGGLGGQQAGAAAGGAGQGGYGLGSGGT) of the native MaSp1 was chosen to prepare the fusion polypeptide (silk6mer-30lysine-ppTG1), with 6 contiguous copies of MaSp1 repeating domains, 30 lysines, and a ppTG1 peptide domain fused at its C-terminus. The silk–polysine–ppTG1 protein trapped pDNA to form micelle-like nanoparticles with an average hydrodynamic diameter of 99 nm (Figure 2B), and this structure displayed a high transfection efficiency (Numata and Kaplan, 2010). The profile of the released genes could be further regulated by selectively controlling the β-sheet content of the silk domains in the gene carrier nanoparticle. The same group also fused this basic silk-30 lysine polypeptide to tumor-homing peptides to form gene delivery nanoparticles 150–250 nm in size. This gene delivery system showed significant target specificity to tumor cells (Numata et al., 2011). Silk crystalline protein domains have been genetically fused with other fibrous protein domains, including the elastin repeating hydrophobic domain (GVGVP) and the collagen repeating triplet domain. Silk–elastin polypeptides (SELPs) consist of silkworm silk protein domains (GAGAGS) and environmentally sensitive elastin domains (GXGVP, X could be any amino acid other than proline). SELPs undergo a two-step self-assembly process in aqueous solutions (Xia et al., 2011). The first assembly step is driven by the silk domains. Because the elastin domains are more hydrophilic below their transition temperature than the silk domains, the silk domains drive SELPs to self-assemble into micelles through hydrophobic associations among the silk domains in water. The second self-assembly step is driven by the elastin domains at higher temperatures (Cheng et al., 2013, 2014a). In the second step, the SELP micelles associate to form higher-order nanostructures, including hydrogels and nanofibers (Xia et al., 2011). These SELPs may be used to encapsulate the hydrophobic antitumor drug doxorubicin into the core of the SELP nanoparticles to promote significant uptake by the cells (Xia et al., 2014). SELPs are good candidates for use in the design of thermo-responsive photonic/electronic systems (Lin et al., 2014). Lin et al. (2014) designed a series of SELPs with a polyhistidine tag at the N-terminus, and these SELPs were shown to bind to the surfaces of nickel chelate nitrilotriacetate-functionalized gold nanoparticles through the nickel affinity of the polyhistidine tag. SELPs drive the gold nanoparticles to aggregate through hydrophobic interactions among the elastin domains at high temperatures, leading to a red shift in the Au nanoparticle surface plasmon resonance. The SELP-Au nanoparticle system may be used as a self-adaptive plasmonic nanodevice for biosensing. Megeed et al. (2002, 2004) designed SELPs-based injectable hydrogels for gene delivery applications. An SELP-47K was designed based on four silk domains and eight elastin domains. A large number of lysine amino acids were coded into elastin domains, and negatively charged DNA bound to the positively charged lysine groups to form a DNA/SELP complex. The SELP/DNA complex formed a hydrogel at body temperature and could be used for injectable gene delivery. Cohen Stuart’s group designed a series of stimuliresponsive silk collagen-like polypeptides \((\text{C}_{n} \text{S}_{m}^\text{a} \text{C}_{n}^\text{b})\) composed of a middle silkworm silk-like peptide domain \((S)\) flanked by two collagen-like peptide domains \((C)\) (Martens et al., 2009; Beun et al., 2012, 2014; Golinska et al., 2014). The middle silk domains consisted of the sequence GAGAGAGX, and pH-responsive amino residues were introduced at position X, resulting in the pH-triggered self-assembly of \(\text{C}_{n} \text{S}_{m}^\text{a} \text{C}_{n}^\text{b}\) into various nanomaterials (micelles, nanofibers, nanotapes, and hydrogels). The \(\text{C}_{n} \text{S}_{m}^\text{a} \text{C}_{n}^\text{b}\) sequence was shown to mimic the properties of natural collagen, lending its potential utility in tissue engineering applications.

**Small Surfactant-Like Peptide Based on the Silk Protein Domain**

The crystalline An domains of the native silk proteins drive self-assembly via hydrophobic interactions and stabilize the
assembly via inter- or intramolecular hydrogen bonding. This sequence has been attached to a head group bearing one or more charged amino acids [e.g., histidine (H), arginine (R), lysine (K), or aspartic acid (D)] to prepare surfactant-like small peptides (SLSPs). SLSPs have properties very similar to those observed in biological surfactant molecules and show particular promise in bionanotechnology and biomedicine applications. A variety of silk domain-based SLSPs have been reported, including A6H, A6D, AnK, and A12R2, and these SLSPs undergo self-assembly to form nanostructures in an aqueous environment.

The A6K family is the most extensively studied SLSP, where n may be 3, 4, 6, 8, 9, or 10. The self-assembly of A6K is concentration dependent. A6K was found to self-assemble above a critical concentration in aqueous solutions and form hollow nanotubes with a radius of 26 nm (Figure 2C), followed by a transition to a nematic lamellar phase with increasing peptide concentration (Bucak et al., 2009; Castelletto et al., 2010; Cenker et al., 2011). A6K could not self-assemble to form nanostructures. On the other hand, A6K and A6K self-assembled to form thin rod-like aggregates, possibly due to the increased hydrophobicity as more alanine amino residues were added (Cenker et al., 2014). The A6K peptides designed by Zhang et al. (von Maltzahn et al., 2003) and later by Zhao et al. (Qiu et al., 2009; Wang et al., 2009) were capped either at both ends or at the N-terminus. These peptides formed nanofibrils other than nanotubes, because the caps influenced the hydrophobicity of the peptides. The A6D, A6H, and A12R2 constructs are self-assembled to form various nanostructures, including nanotubes, nanovesicles, nanotapes (Figure 2D), nanosheets (Figure 2E), or twisted fibrils (Vauthey et al., 2002; Zhao et al., 2006; Hamley et al., 2013; Castelletto et al., 2014).

**Polymer–Peptide Hybrids Inspired by Silk Proteins**

The incorporation of silk’s β-sheet-forming peptide domain (GA)n or (A)n to form a synthetic polymer offers an effective approach to design supramolecular materials via self-assembly. Smeenk et al. (2005, 2006) reported the development of a triblock polymer consisting of a central silk β-sheet domain [(AG)3EG]n with one PEG block at both its N- and C-termini. The central silk β-sheet domain was overexpressed in *E. coli* and purified via nickel–nitrilotriacetate (Ni–NTA) affinity chromatography. One cysteine amino residue at each peptide terminus was used to selectively alkylate the maleimide-functionalized PEG. The PEG blocks prevented the central silk β-sheet domains from forming random macroscopic aggregates. Instead, the central β-sheet domains stacked together in an orderly fashion to form well-defined fibrils (Smeenk et al., 2005).

Zhou et al. (2006) reported a series of silk-like multiblock peptide–polymer hybrids containing A5 blocks and polyisoprene oligomers (PI). The PI oligomer improved the solubility of the hybrid polymer, just as glycine-rich domains do in natural silk proteins. The A5 blocks directed the hybrid polymer to self-assemble into micelles through the formation of an antiparallel β-sheet core in the presence of chloroform or 2-chloroethanol. A similar design concept was applied by Rathore et al., who replaced amorphous domains of the silk proteins with PEG. The remaining β-sheet-forming domain of the silk proteins were then included in the synthesized PEG–peptide copolymer, which self-assembled to form nanostructures (Rathore and Sogah, 2001a,b).

The GAGAG pentapeptide was conjugated to tetrathioene to form diblock or triblock peptide/polymer hybrids that self-assembled into supramolecular structures at the graphite surface and in organic solvents due to β-sheet formation among the peptide blocks (Gus’kova et al., 2008). The thiophene blocks further stabilized the supramolecule through the formation of π–π stacks, which conveyed conductivity to the supramolecular structures (Gus’kova et al., 2008).

Croiser et al. (2014) synthesized a series of peptide/poly(isobutylene) hybrid elastomers. In these hybrids, silk peptides were conjugated to one terminus or both termini of an poly(isobutylene) unit to form a diblock hybrid (monofunctional hybrid M) or a triblock hybrid (difunctional hybrid D). The silk peptide domains directed M and D to self-assemble into various nanostructures. The length of the silk peptide domain determined the morphology of the nanostructures formed through self-assembly. Both the M and D units comprising short peptide domains formed small aggregates. As the length of the peptide domain increased, single β-sheet tapes and stacked nanofibrils formed. Mixing M and D together gave rise to inherently reinforced thermoplastic elastomers and interpenetrating supramolecular networks (Croiser et al., 2014).

Self-assembly in the various silk-inspired peptide–polymer hybrids mentioned above required an organic solvent environment, and hydrogen bonds formed among the silk peptides to strengthen and direct the self-assembly process. Stupp’s group described the use of a series of well-known diblock polymers, the peptide amphiphiles (PA), consisting of a peptide segment and a more hydrophobic alkyl tail that self-assemble to form supramolecular nanofibers in an aqueous solution (Harterink et al., 2001, 2002; Claussen et al., 2003; Behanna et al., 2005; Shah et al., 2010). The silk GA domains were incorporated into the PAs to provide stability to the nanofibers by forming β-sheets in the fibers (Shah et al., 2010). The PAs provide a powerful tool for drug delivery and tissue engineering.

**NOVEL SPINNING PROCESSES AND MICROPLATFORMS INSPIRED BY NATURE**

The outstanding mechanical properties of silk fibers originate from both the silk protein’s unique primary sequence and the distinctive spinning process used during fiber fabrication. The success of spider silk spinning lies in the various silk proteins produced in its multiple glands, the extensional field formed inside its tapered duct, and the post-spin drawing action of the spider’s legs. This section discusses a variety of spinning methods using microfluidic platforms that provide excellent quality fibers through the control over multiphase flows in small amounts of liquid.

**Spinning and the Mimicking of Multiple Glands**

Individual spiders can spin a variety of silks. For example, the female orb-weaving spiders spin up to seven different types of
silks to meet different needs using seven different glands. The multigland design of the orb-weaving spider was implemented by our group to continuously spin fibers with spatiotemporal variations in chemical composition and morphology using polydimethylsiloxane (PDMS) microfluidic chips (Figure 3A) (Kang et al., 2011). The microfluidic chip consisted of several individually controllable inlets that mimicked the multiple glands of the spider. Switching among these inlets was precisely controlled using a computer-controlled pneumatic valve that enabled the preparation of fibers with spatially organized morphologies and compositions. Multiple compositions and topographies, including gas microbubbles, nanoporous spindle knots, and a variety of cells, could be coded into a single fiber (Kang et al., 2011). Similar design concepts were implemented by Qin et al. to prepare hybrid microfibers with bamboo-like or pearl-like morphologies by combining the droplet microfluidic technique with microfluidic wet spinning (Yu et al., 2014). Hydrophobic droplets containing polymer microspheres or multicellular spheroids were encapsulated in a continuous alginate hydrophilic phase using “T” junction channels. The Gu’s group replaced the spheroids were encapsulated in a continuous alginate hydrophilic phase using variations on these chip designs (Shin et al., 2007; Hwang et al., 2008, 2009; Lee et al., 2009, 2010). Stupp’s group used a tapered pipet to prepare noodle-like strings in which peptide amphiphile nanofibers aligned and were bundled along the longitudinal direction of the string (Zhang et al., 2010). The strings could be used to encapsulate cells and direct the orientations of the cells in a 3D environment. Human mesenchymal stem cells (hMSCs) were found to remain viable during the string preparation process. Both the cell bodies and filopodia were aligned along the peptide amphiphile nanofibers due to contact guidance by the oriented matrix (Zhang et al., 2010). Takeuchi’s group designed a series of coaxial microfluidic glass capillaries with tapered orifices to control the assembly of diverse small structures under flow (Kiriya et al., 2012a,b; Hirayama et al., 2013; Onoe et al., 2013). In the elongational flow, small nanofibers formed the core flow and were oriented along the flow stream as the sheath flow velocity exceeded the core flow (Kiriya et al., 2012b). The elongational flow overcame the hydrophobic/hydrophilic interactions and oriented the lipid-type monomers to form meter-long fibers. The fibers were then used as templates to synthesize polyaniline conductive nanofibers with a high conductivity and potential utility as sensor materials (Kiriya et al., 2012a). Fast sheath flows and slower core flows can create an elongational field and such fields have been used to induce alignment of carbon nanotubes (CNTs) (Davis et al., 2009). Single-wall CNTs in pure chlorosulfonic acid at a concentration of 8.5 vol% were spun into a faster-flowing 96% sulfuric acid sheath flow to form CNT fibers uniformly composed of oriented CNT fibrils. These fibers showed great mechanical properties and a high conductivity (Davis et al., 2009).

Our group has fabricated PDMS-only cylindrical coaxial flow channels with a tapered orifice to mimic the spider’s tapered spinning duct (Figure 3B) (Kang et al., 2010, 2011). The PDMS-based mold was prepared using soft lithography techniques. A thin PDMS membrane was subsequently bonded to the PDMS base mold, and a negative pressure was applied to form a concave hemicylindrical channel. The hemicylindrical channel was replicated with SU-8, and a convex hemicylindrical SU-8 master was obtained. The convex hemicylindrical SU-8 master was used to prepare a concave hemicylindrical PDMS channel. The PDMS chip used to prepare the fibers was constructed by aligning and bonding two concave hemicylindrical PDMS channels using oxygen plasma treatment (Kang et al., 2010, 2011). This PDMS chip was used to continuously prepare monodisperse microfibers for cell encapsulation and tissue engineering (Kang et al., 2012, 2014; Jun et al., 2013, 2014a,b).

**Post-Spin Drawing Process**

During the natural silk spinning process, the fibers are drawn out of the spinning duct either using the legs of the spider or through the “figure eight” movements of the silkworm head (Ormenetto and Kaplan, 2010). This drawing process orients the silk proteins along the longitudinal axis of the fibers and improves the fibers’ mechanical properties. The post-spin drawing process used by silkworms and spiders has been replicated in conventional processes to improve the mechanical properties of the artificial
fibers. Chen et al. prepared strong artificial silk fibroin fibers by continuously drawing the fibers through a series of plastic rollers with different rotating speeds, the post-spun drawn fibers showed mechanical properties comparable to those of the natural silkworm silk fibers (Figure 3C) (Zhou et al., 2009; Yan et al., 2010). The same spinning process was applied by Chen’s group...
to a spinning dope composed of a silk fibroin–CNT composite to prepare strong artificial fibers with breaking energies comparable to that of the spider dragline silk fibers (Fang et al., 2015). Lewis’s group used post-spin drawing to prepare strong artificial spider silk fibers (An et al., 2011). Post-spin drawing significantly increased the fiber tensile strength due to the improved protein molecule rearrangement and β-sheet formation in the polyalanine region (An et al., 2011). We recently successfully spun microfibers using silk fibroin and alginate, and the mechanical properties of the fibers were significantly enhanced by the post-spin drawing process (unpublished data).

**DEVELOPMENT OF NEW SMART MATERIALS AND APPLICATIONS INSPIRED BY COMPLEX HIERARCHICAL ARCHITECTURES CONSTRUCTED BY SILKWORM AND SPIDER**

The advanced functions of most natural materials rely on the complex hierarchical structures of the materials (Sanchez et al., 2005). Recent studies have revealed that the strength of silk fibers also relies on its hierarchical structure (Cranford et al., 2012; Koski et al., 2013; Qin and Buehler, 2013; Silva and Rech, 2013; Qin et al., 2015). The structure of a spider’s web is hierarchical in two aspects: the protein secondary structure inside a single silk fiber is hierarchical and the geometric structure of the whole web is hierarchical. Together, these structures contribute to the outstanding properties of natural spider webs (Cranford et al., 2012; Qin et al., 2015). The hierarchical protein secondary structure (semi-amorphous and β-sheet nanocrystals) in a single spider silk fiber leads to non-linear stress–strain behaviors of the silk fiber, resulting in small web deformations under the application of an external force to a single fiber (Cranford et al., 2012). Orb web spiders produce up to seven different types of silk fibers using their different glands to weave a hierarchical web net, and any insects trapped in this thin fiber net have no chance of escape. The framework of the web is constructed using dragline silk produced by the major ampullate gland (MA gland). The capture spiral is jointly constructed by minor ampullate spidroin silk fibers produced in the minor ampullate gland (MI gland) and by flagelliform silk fibers produced in the flagelliform gland. The auxiliary spiral is first constructed using ampullate spidroin silk fibers to stabilize the emerging web structure and to serve as a template in support of the construction of the capture structure using the flagelliform silk fibers. The whole web structure is fixed by a gluey substance called attachment cement, which is produced in the piriform gland (Heim et al., 2010).

Spiders spin a variety of silk fibers to meet their different needs, such as prey capture, prey swathing, or egg protection. Although the silkworm *B. mori* produces only one type of silk fiber to meet its exclusive need for cocoon weaving as a defense mechanism, the single silkworm silk fibers and the cocoon are hierarchical structural materials. Silkworm silk fibers consist of a fibroin core and a sericin shell, and the cocoon wall is a multilayer hierarchical architecture randomly woven using the silk fibers.

**Cocoon Silk’s Core/Shell Composite Structure Has Inspired the Development of Strong Fibers**

Cocoon silk fibers consist of a fibroin core and a sericin coating layer. The sericin layer allows for the efficient transfer of an external loading force to the fibroin core and improves the mechanical properties of the silk fiber. Inspired by this core/shell design, Jiang’s group enforced lotus fibers by applying a poly(vinyl alcohol) (PVA) coating layer (Figure 4A) (Wu et al., 2014). In addition to improving the mechanical properties, the PVA layers acted as a glue to hold the individual fibers together as a fiber bundle. The mechanical properties of the fibers were further improved by introducing cross-links through heat treatment and glutaraldehyde.

**Spider Silk-Inspired CNT-Based Fiber Design**

The outstanding mechanical properties of spider silk rely on the plentiful intra- and intermolecular hydrogen bonds among the structural proteins. These features suggested a simple approach to strengthening artificial fibers. CNTs are extremely stiff and strong nanomaterials and have been widely used in nanocomposites to improve the mechanical properties of the materials. Scientists and engineers believe that the best approach to fully utilizing the outstanding mechanical properties of CNTs is to spin them into continuous fibers on the macroscale (Ericson et al., 2004; Zhang et al., 2005; Peng et al., 2008). The continuous spinning of CNTs has remained challenging due to the atomically smooth surfaces and cylindrical profile of the CNTs, which provide a low shear load transfer and facilitate sliding among aligned CNTs (Kis et al., 2004; Peng et al., 2008). Inspired by the silk fibers, hydrogen bonding functional groups were introduced onto the pristine CNT surfaces to improve the shear load transfer and prepare continuous CNT fibers. In addition to small functional groups, such as OH, NH₃, may be introduced onto the CNT surfaces to form bridging hydrogen bonds among CNTs. Ultra-short polymers may also be covalently conjugated to the CNT surfaces to create a microstructural environment similar to that present in natural spider silk. Naraghi et al. (2013) functionalized CNT bundles with poly(methyl methacrylate)-like oligomers using chemical vapor deposition methods (Figure 4B). The van der Waals interactions increased the shear strength of the bundle junctions by an order of magnitude compared to the shear strength of the pristine CNT bundles.

**Spider Silk Adhesion Glue**

The viscid fibers of web-weaving spiders are coated with glycoprotein-based glue droplets that facilitate prey capture (Figure 4C) (Opell and Hendricks, 2009, 2010; Sahni et al., 2010; Stellwagen et al., 2014). The properties of the glue droplets depend on the rate at which they are extended. The glue droplets act as a viscoelastic material under rapid extension, for example, when snared by a fast-flying insect, or as a viscoelastic material under slow extension, for example, during the attempts by trapped prey to escape (Sahni et al., 2010). The adhesion of the spider’s glue droplet is...
FIGURE 4 | Development of new smart materials by mimicking silk's complex hierarchical structure. (Continued)
humidity responsive and increases with humidity. These properties are exactly opposite the properties displayed by synthetic adhesive systems. Although glue droplets appear to consist of a glycoprotein core and a viscous aqueous salt coating (Vollrath et al., 1990; Opell and Hendricks, 2009, 2010; Sahni et al., 2010; Stellwagen et al., 2014), an in-depth examination of the relationship between the chemical composition and the physical properties has not yet been conducted. Synthetic adhesive systems that display adhesion properties similar to those of spider silk have not yet been developed.

**Wet Assembly Fibers**

Aside from their outstanding mechanical properties, the water-collecting properties of spider silk can provide critical insights into the design of advanced materials (Wen et al., 2015). The water-collecting properties of silk result from the hierarchical spindle knot-joint structure formed during wetting in humid air (the upper panel in Figure 4D) (Zheng et al., 2010). Dry spider silk consists of puffs separated by joints. As the dry spider fibers encounter humid air, tiny water drops condense on the semitransparent puffs, and the puffs transform into periodic spindle knots as water condensation continues (Zheng et al., 2010). The spindle knots and joints are composed of random and aligned nanofibrils, respectively, which results in a surface energy gradient and a difference in the Laplace pressure between the spindle knots and the joints (Zheng et al., 2010). As a result, water continuously condenses and collects around the spindle knots.

Mechanistic insights gleaned from the water collection properties of spider silk have informed the development of spider silk-like artificial fibers based on a range of materials. Water collection properties similar to those of natural spider silk have been achieved. Zheng et al. prepared artificial spider silk-like fibers using a simple dip-coating method. Uniform nylon fibers were immersed into a poly(methyl methacrylate)/N,N-dimethylformamide/ethanol (PMMA/DMF-EtOH) solution and then horizontally drawn out quickly (Zheng et al., 2010). After drying, the polymers formed periodic spindle knots on the surfaces of the nylon fibers (Zheng et al., 2010). The artificial polymer spindle knots were composed of random pores and artificial joints prepared from the stretched pores. The differences between the surface morphologies of the artificial spider knots and joints gave rise to a surface energy gradient and difference in the Laplace pressure, yielding directional water collection around the spindle knots (Zheng et al., 2010).

Coaxial electrospinning methods offer a good approach to preparing spider silk-like fibers with periodic spindle knot/joint patterns (Tian et al., 2011; Dong et al., 2012). A viscous polystyrene (PS) solution and a less viscous PMMA or poly(ethylene glycol) (PEG) solution were used as the inner and outer solutions, respectively, in a coaxial system. Under a high-voltage electric field, the inner PS solution spun to form a thin thread, whereas the outer PMMA or PEG solutions formed a thin film on the surface of the PS thread. The PMMA or PEG films broke to form periodic droplets and solidified into spindle knots on the surface of the PS thread (Tian et al., 2011; Dong et al., 2012). These electrospun fibers demonstrated capacities for directional water collection that resembled the corresponding capacity of natural spider silk fibers.

Our group used a multigland spider-mimicking microfluidic chip to prepare artificial spider silk-like fibers containing a spindle knot/joint pattern that directionally transported water toward the spindle knots (Kang et al., 2011). This microfluidic chip continuously created microfibers with strictly coded morphological and chemical features using a digital programmable flow controller. An alginate solution containing salt was injected into one of sample channels at a high flow rate to construct porous spindle knots (the lower left image in Figure 4D). An alginate solution without salt was injected into another sample channel at a low flow rate to construct joints (the lower left image in Figure 4D). As the salt leached from one flow to the other through diffusion, nanoporous spindle knots were generated. These spindle knots provided a higher surface energy than the joints. The tapered morphologies of the spindle knots generated a difference in the Laplace pressures of the spindle knots and joints. As expected, water droplets were directionally collected around the spindle knots under the combined action of the surface energy and the Laplace pressure differential (the lower right image in Figure 4D).

The relationship between directional water transport and differences in the surface energies and Laplace pressures was used to prepare artificial fibers with the capacity to transport water droplets in the reverse direction, that is, from the spindle knots toward the joints. The relative wettability of the spindle knots and joints was tuned by changing the spindle knots’ chemical composition or the roughness (Bai et al., 2010). Interestingly, artificial fibers were prepared to enable in situ switching of the directional water droplet movement by introducing a stimulus-responsive material onto the spindle knots (Feng et al., 2013; Hou et al., 2013). Thermo-responsive copolymer poly(methyl methacrylate)-b-poly(N-isopropylacrylamide) (PMMA-b-PNIPAAm) or photosensitive azobenzene polymers have been introduced into spindle knots. PMMA-b-PNIPAAm-modified spindle knots displayed a wettability that could be tuned by adjusting the environmental temperature below or above the lower critical solution temperature (LCST) of PNIPAAm. This stimulus switched the direction...
of the water droplets' movement (Hou et al., 2013). Photosensitive azobenzene polymer-modified spindle knots permitted control over the wettability and, therefore, the droplet movement direction via UV–vis irradiation, which transformed the molecular configuration of the azobenzene polymer from a less hydrophilic to a more hydrophilic state (Feng et al., 2013).

Silk-Inspired Thermal Materials

Silkworms weave cocoons to protect the pupae from extremely cold weather. The cocoon wall is a multilayer hierarchical fiber-based structure that traps air within the cocoon to provide thermal insulation. The still air is retained within the cocoon wall, even in the presence of wind, by depositing a calcium oxalate crystal layer on the outer surface of the cocoon wall. This layer reduces wind penetration (Zhang et al., 2013; Tao et al., 2015). With this understanding in mind, Wang et al. (2013) fabricated thermally insulating hollow biomorphic fibers using silkworm silk as the templates (the left image in Figure 4E). The silk fibers were immersed in an AlCl₃ solution and dried at 70°C over 24 h. An alumina layer formed on the surfaces of the silk fibers during an oxidation reaction. The silk fibers themselves were removed by heating, yielding free-standing alumina fibers with a hollow core (Wang et al., 2013). The fibers showed better thermo-insulating properties than traditional alumina fibers due to their hollow structures and infrared absorbing properties (Wang et al., 2013).

In general, bulk polymers have a very low thermal conductivity owing to strong photon scattering within the material (Chae and Kumar, 2008). Wang et al. recently found that dragline silk is a thermally conductive material with a thermal conductivity of 340 W·m⁻¹·K⁻¹ in its relaxed state (0% strain). The thermal conductivity may be further improved to 416 W·m⁻¹·K⁻¹ simply by stretching the fiber by 19.7% (Huang et al., 2012). The inherently high thermal conductivity of dragline silk is largely attributed to its crystal/non-crystal hierarchical internal structure. The nanofibrils are intensely packed in the dragline silk fibers with few defects, leading to a high thermal conductivity. The silk's thermal conductivity enhances after stretching due to an increase in the alignment of both the β-sheet crystal structure and the helical non-crystal structure (the right image in Figure 4E), thereby further enhancing photon conduction (Huang et al., 2012).

Prior to Wang's work, researchers recognized the effects of polymer chain orientation on the thermal conductivity of the polymer fibers (Shen et al., 2010). Chen et al. fabricated high-quality ultra-drawn polyethylene nanofibers with a thermal conductivity as high as ~10³ W·m⁻¹·K⁻¹, larger than the thermal conductivities of half of the pure metals, including platinum, iron, and nickel (Shen et al., 2010). The high thermal conductivity was attributed to the reorientation of the polyethylene chains to form large crystals along the drawing direction after stretching, which decreased the defect density in the fibers and made the fibers ideal single crystalline fibers (Shen et al., 2010).

Spider Web Network-Inspired Materials

Buehler's group examined the mechanism by which the nonlinear mechanical characteristics of single silk fibers contribute to the integrity and performance of a spider web through web deformation experiments and atomistic simulations (Figure 4F) (Cranford et al., 2012). The non-linear mechanical response of strain derives from the inherent semi-amorphous/β-sheet nanocrystal hierarchical molecular structure of the silk fibers. Under relatively low stress, semi-amorphous domains permit entropic unfolding, which softens the stress–strain curves. Once unfolding is complete, stress is transferred to the rigid β-sheet crystal domains, leading to a sharp increase in the stress. These unique mechanical properties of single silk fibers localize load-induced deformations by sacrificing the loaded fibers while maintaining the integrity of the web. This model offers a good explanation of why most natural webs are not intact in structure but remain functional for a spider's use (Cranford et al., 2012). The sacrifice of individual silk fibers to avoid damaging the whole web system has provided crucial inspiration for the design of new advanced 3D structure.

Recently, the same group prepared PDMS-based artificial spider webs using 3D printing to study how the geometrical arrangement of the silk fibers in a web influenced the web's mechanical performance (Qin et al., 2015). They found that the ratio of the radial silk diameter to the spiral silk diameter played a key role in tuning the mechanical response of the whole web to load. A homogeneous distribution of fibers with a radial/spiral ratio of ~1 tended to localize loading, whereas a heterogeneous distribution with much thicker radial silk fibers tended to distribute loading. This finding agrees well with the designs implemented naturally by spiders. Small webs built by garden spiders consist of radial and spiral silk fibers that are similar in size, which provides the maximum web strength against point loading by small prey. By contrast, giant webs built by spiders in rainforests are used to catch much larger prey; therefore, the spiders weave their webs with much thicker radial silk fibers to maximize the web strength against distributed loading (Qin et al., 2015).

Non-Web Spider Silk Materials

Besides aforementioned web silk, spiders also produce other types of silk, including pyriform silk, aciniform silk, and tubuliform/cylindriform silk for different uses, which also provide a lot of inspiration to fabricate new materials. Spiders produce attachment disks spun from pyriform silk to anchor their webs to diverse substrates. Cobweb spiders produce two types of attachment disks with distinct architectures using the same pyriform silk: “staple-pin” and “dendritic” disks (Sahni et al., 2012). “Staple-pin” disk firmly attaches the dragline silk to the substrate and “dendritic” disk weakly attaches the gumfoot silk to the substrate, the reason behind the difference in adhesion strengths of the two disks is the peeling angles. Small peeling angle of “stable-in” disk is the secret of high adhesion strength; on the other hand, dendritic disk is peeled at much higher angles, resulting in its much lower adhesion strength. With this understanding, synthetic adhesive attachment disks inspired by spider have been prepared. Jain et al. (2014) fabricated artificial attachment disk with a similar architecture to spider's “stable-pin” disks using polyurethane electrospun fibers. The adhesion energy and peeling forces can be controlled by adjusting the spacing of fibers and the surface energy of the substrate (Jain et al., 2014).

Spiders fabricate egg case using tubuliform/cylindriform silks to protect the egg from predator, because of their high
toughness. At least three different proteins have been found in tubuliform silk: tubuliform spidroin 1 (TuSp1), egg case protein 1 and 2 (ECP-1 and ECP-2). Tutuliform silks is the second most studied spider fiber following the dragline silk and the primary sequence and structure of tubuliform silk proteins have been clearly demonstrated (Lin et al., 2009, 2013; Gnesa et al., 2012). Lin et al. (2013) produced TuSp1 consisting of both repetitive and conserved terminal domains using recombinant DNA technology, and the artificial fibers spun from the recombinant TuSp1 showed outstanding mechanical properties.

Spiders produce aciniform silk to wrap and immobilize prey because aciniform silks possess the highest toughness out of all the spider silks including the famous dragline silk. Acriniform silk is also used to form the soft inner layer of egg case. Tremblay et al. (2015) found that the high toughness of aciniform silk is related to the primary sequence of spidroin and the distinctive mixture of α-helical, β-sheet, and non-canonical secondary structures. We expect that well understanding of the hierarchical architecture about these silks will inspire to the designs and development of advanced materials.

**Artificial Silk Fiber Prepared by Biomimetic Spinning**

Artificial spinning platforms have been designed to mimic the geometric construction of silkworm/spider spinning ducts, and artificial silk fibers have been prepared. Rammensee et al. (2008) designed a biomimetic microfluidic device in which ions and pH gradients were established along the elongational flow. Silk formation of a MaSp2 analog has been achieved using this device under similar condition to natural spider spinning duct. Kinahan et al. (2011) designed a microfluidic chip with a cross channel to prepare strong artificial silk fibroin fiber. PEO solution was under similar condition to natural silk. We expect that a much deeper understanding of the outstanding properties of silks will inspire researchers to design novel functional materials and to apply in diverse fields.

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**CONCLUSION AND OUTLOOK**

This review has outlined the sequence–structure–property interplay observed in natural silks and overviewed how smart materials are fabricated by mimicking natural silks. Significant progress in understanding the silk proteins’ self-assembly, natural spinning process, and structure–property interplay has been achieved over the past decade, and many hidden secrets lie behind silk production and structure have been unveiled. This progress enabled the development of new materials inspired by natural silk. We expect that well understanding of the outstanding properties of silks will inspire researchers to design novel functional materials and to apply in diverse fields.

**AUTHOR CONTRIBUTIONS**

Prof. S-HL is the corresponding author and contributed to writing. Dr. JC contributed to data collecting and writing.

**FUNDING**

This work was supported by National Research Foundation of Korea (NRF-2015R1A2A1A09004998), Republic of Korea.
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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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