Elucidating silk structure using solid-state NMR

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An overview of solid-state NMR structural studies on various silk forms and analogs conducted by the authors’ research groups is presented. The well-studied silkworm and spider silks together with related silk peptides have a mixture of secondary structures including β-sheet, β-turn, helix and random coil that are difficult to analyze by X-ray diffraction and electron microscopy but conveniently investigated by solid-state NMR. Several newly developed solid-state NMR techniques and stable isotope labeling approaches of the silks were effectively used to characterize silk structure. The techniques discussed provide not only information on the secondary structure, but also on the hydrogen-bonding interactions present in the silks. Structural studies on other types of silk, silk peptide mimics and recombinant silk proteins are also discussed.

1. Introduction

Recently, considerable attention has been paid to silk by a range of scientists from textile engineers to biochemists and biomedical researchers. Silk is an attractive material because of its excellent mechanical properties, that is, the combinations of strength and toughness not found in today’s man-made materials together with excellent biocompatibility.1 These appealing physical properties originate from the silk fibroin structure and therefore, the atomic level information on the silk structure gives us the answer to why the silk has such excellent properties. X-ray and electron diffraction methods together with other spectroscopic methods have been applied for this purpose. However, silk fibers are inherently non-crystalline or semi-crystalline biopolymers and a mixture of secondary structures including β-sheet, helix, β-turn and random coil are present making silk difficult to analyze by X-ray diffraction and electron microscopy. The conformation-dependent NMR chemical shifts can determine the fraction of mixed structures and monitor the conformational change easily and selectively.2–4 The techniques discussed provide not only the information on secondary

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structures of the mixtures, but also on hydrogen-bonding. In addition, NMR covers both solution and solid state, including soft matter (gel state) and further includes looking at silk in living organisms (e.g., silkworms) by solution NMR. Furthermore, the atomic level structure can be obtained by using solution or solid-state NMR, by combining with stable $^2$H, $^{13}$C and/or $^{15}$N isotope labeling of silks and their sequential model peptides. The focus of this review is on the recent solid-state NMR studies used to elucidate the molecular structure of silk fibers and related model peptides.

2. Silkworm silk

Most recently, Zhou’s group obtained a single crystal of the N terminal domain of Bombyx mori silk fibroin and determined the crystal structure, revealing an entangled β-sheet dimer. However, the structure of the main part, the crystalline domain with the sequence, (AGSGAG)$_n$ is still the target of solid-state NMR. Two crystalline forms of B. mori silk fibroin, silk I and silk II, have been reported as the dimorphs of the silk fibroin. Here silk I is the structure of B. mori silk fibroin in the solid state obtained from the middle silk gland after drying. The silk II is the structure of B. mori silk fibroin fiber after spinning. The structure analyses of silk I and silk II reported by Asakura’s group during the past 10 years are reviewed in this section although the structure of another silk protein, silk sericin was also studied with NMR.

2.1 Structure of B. mori silk fibroin before spinning (silk I) in the solid state

Despite a long history of studying silk I, the structure remained poorly understood because any attempt to induce a macroscopic orientation of the silk sample for X-ray diffraction or electron diffraction analyses, readily caused a conversion of the silk I form to the silk II form. Therefore solid-state NMR is useful because the determination of the silk I structure is possible without orientation or crystallization. The structural features of B. mori silk fibroin are conveniently studied by using the synthetic peptide, (AG)$_n$, as a model for the crystalline region because the lack of Ser in the model peptide (AG)$_{15}$ does not make any difference in the $^{13}$C cross-polarization magic angle spinning (CP-MAS) NMR chemical shifts of the Ala and Gly residues in the repeated sequence (AGSGAG)$_n$ of the native silk fibroin.

The backbone structure of the silk I conformation for (AG)$_n$ was determined to be a repeated type II β-turn structure by combining several solid-state NMR techniques, that is, the quantitative use of the conformation dependent chemical shifts, 2D spin diffusion NMR under off magic angle spinning spectra and Rotational Echo Double Resonance (REDOR) experiments for selectively stable-isotope labeled peptides. Namely, by using the $^{13}$C chemical shift contour maps for various amino acid residues in the protein as a function of the $\Phi$ and $\Psi$ values, the $\Phi$ and $\Psi$ regions in the Ramachandran map that satisfy both the Cα and Cβ chemical shifts of the Ala residue of (AG)$_{15}$ with silk I form, are determined to be $\Phi = -80^\circ$ to $-20^\circ$, $\Psi = 90^\circ$ to $180^\circ$. Then the 2D spin-diffusion NMR spectra were observed for (AG)$_n$A[1-$^{13}$C]G[1-$^{13}$C]
A\textsuperscript{13}G(AG), and (AG)\textsubscript{15}(1-\textsuperscript{13}C)A\textsuperscript{13}F[\textsuperscript{1-13}C]G\textsuperscript{16}(AG), to determine the torsional angles of Ala\textsuperscript{15} and Gly\textsuperscript{16} residues. From the spectral simulation, the torsional angles were determined to be (\(\Phi = -60^\circ\) and \(\Psi = 130^\circ\)) for the Ala residue and (\(\Phi = 70^\circ\) and \(\Psi = 10^\circ\)) for the Gly residue. The REDOR experiments were performed to determine the atomic distance between two selectively \(1^\text{H}\) and \(1^\text{H}\) double labeled (AG)\textsubscript{15} were also performed and they supported the structural model for silk I. By adding the structural information from X-ray diffraction analysis of silk I, the final silk I structure forming intra- and inter-molecular hydrogen bonds was proposed as shown in Fig. 1.\textsuperscript{14} This structure was also supported by \(1^\text{H}-\text{H}\) REDOR experiments\textsuperscript{15} and \(1^\text{H}\) solid state NMR.\textsuperscript{16}

2.2 Structure of B. \textit{mori} silk fibroin after spinning (silk II)  

The structure of silk II has been proposed as a regular array of antiparallel \(\beta\)-sheets firstly by Marsh \textit{et al.}\textsuperscript{17} about a half century ago, based on a fiber diffraction study of native \textit{B. mori} silk fibroin fiber. Later, Fraser \textit{et al.}\textsuperscript{18} Lotz \textit{et al.},\textsuperscript{9} and Takahashi \textit{et al.}\textsuperscript{19} pointed out some intrinsic structural disorder in the silk II structure although they essentially supported the general features of this anti-parallel \(\beta\)-sheet model. Actually, the Ala C\beta peak in the \(1^\text{3}\)C CP-MAS NMR spectrum of [3-\textsuperscript{13}C]Ala \textit{B. mori} silk fiber in the silk II form is broad and asymmetric, reflecting the heterogeneous structure of the silk fiber (Fig. 2).\textsuperscript{20,21} Namely, in the crystalline domain with the sequence, (AGSGAG), (solid blue line: 56% of \textit{B. mori} silk fibroin), 18% distorted \(\beta\)-turn, 25% \(\beta\)-sheet A and 13% \(\beta\)-sheet B (each is shown as broken blue lines). On the other hand, in the amorphous region (thick broken red line: 44%), the distorted \(\beta\)-turn and distorted \(\beta\)-sheet are of the same fraction, 22% (each is shown as broken red lines). Takahashi \textit{et al.}\textsuperscript{19} proposed two kinds of anti-parallel \(\beta\)-sheet conformation with different inter-molecular arrangements by X-ray diffraction analysis. However, the two \(\beta\)-sheet models by Takahashi \textit{et al.} are unstable with high conformational energies. Thus, a further study is required. Yamauchi and Asakura\textsuperscript{22-24} developed a micro-probe with ultra-high speed MAS for high resolution solid-state \(1^\text{H}\) NMR and used this technique with advanced energy and chemical shift calculations\textsuperscript{26} to determine the two \(\beta\)-sheet structures with different intermolecular arrangements, A and B, in Fig. 2. The structural change from the aqueous solution to silk II was studied with Rheo NMR.\textsuperscript{25} The change from silk I film to silk II fiber by stretching was also studied with solid-state NMR\textsuperscript{23} as well as with MD simulation.\textsuperscript{26}

3. Spider silk

Spider silk is one of the toughest biopolymers known.\textsuperscript{27} Solid-state NMR has played a critical role in determining the molecular structure and dynamics of this proteinaceous fiber. \(1^\text{3}\)C CP-MAS was used to first determine that the poly(Ala) repeat units formed the nanocrystalline \(\beta\)-sheet domains in major and minor ampullate spider silks. Deuterium solid-state NMR was used to probe the molecular orientation and two-component nature of the \(\beta\)-sheet crystalline fraction.\textsuperscript{28} Two-dimensional (2D) proton driven spin-diffusion\textsuperscript{14} and DOuble Quantum Spectroscopy (DOQSY) solid-state NMR techniques illustrated that the glycine-rich, disordered regions form an approximate 3\textsubscript{10}-helical conformation. In addition to understanding the backbone conformation and orientation with respect to the fiber axis, solid-state NMR has provided considerable insight into the interaction between spider silk and water and the resulting supercontraction process.\textsuperscript{29-38} In this section, we detail the recent (past 5 years) solid-state NMR developments from the Holland and Yarger research groups to elucidate the spider silk structure. Emphasis is placed on the information gained about the spider silk conformational structure and the reader is directed towards a recent review that details the NMR methods used.\textsuperscript{4}

3.1 Secondary structure from two-dimensional (2D) solid-state NMR

NMR isotropic chemical shifts provide valuable information regarding the protein secondary structure and have been used since the early 1980’s to determine backbone conformation in
Although these one-dimensional (1D) $^{13}$C CP-MAS NMR methods provide valuable information about the structure, they often have limited resolution because of the broad linewidths observed for silks. The primary amino acid sequences of spider and silkworm silks are highly repetitive where a single amino acid can be present in multiple structural environments. This results in considerable chemical shift heterogeneity that broadens the lines for individual amino acid groups. Recently, two-dimensional homonuclear and heteronuclear solid-state NMR approaches have been used to characterize the secondary structure in $^{13}$C- and $^{13}$C/$^{15}$N-isotopically enriched spider silks.\textsuperscript{41–46} These multi-dimensional, multi-nuclear NMR techniques yield considerably more structural information compared to the earlier 1D approaches.

An example of a 2D $^{13}$C/$^{13}$C homonuclear correlation spectrum collected with dipolar-assisted rotational resonance\textsuperscript{47,48} (DARR) recoupling for $^{13}$C-alanine enriched spider dragline silk is presented in Fig. 3.\textsuperscript{44} These 2D spectra are collected with various DARR mixing times to probe short- and long-range through-space dipolar contacts between $^{13}$C spins. In our experience with moderately (15–50\%) $^{13}$C-enriched spider silks, intramolecular (within an amino acid) spin connectivity is established with 100–150 ms mixing times, while longer mixing times up to 1 s provide intermolecular (between amino acids) contacts. The 2D $^{13}$C/$^{13}$C correlation spectrum shown in Fig. 3 was collected with a longer recoupling period of 1 s where both intramolecular and intermolecular spatial contacts are observed. These 2D spectra provide two valuable pieces of information not present in the typical 1D $^{13}$C CP-MAS spectrum. The first is the ability to extract the distinct carbonyl isotropic chemical shift for the different amino acids by slicing through the low ppm region of the spectrum. The carbonyl isotropic chemical shift provides valuable information regarding the conformational structure and is completely masked in a 1D spectrum where a single, uninformative broad resonance is observed that spans nearly 10 ppm (see Fig. 3a, projection). The second piece of information that the 2D $^{13}$C/$^{13}$C correlation spectrum with a 1 s DARR mixing period provides is the ability to detect long-range intermolecular magnetization exchange between neighboring amino acids. This information can be used to assign specific NMR resonances to the repetitive amino acid motifs present in the spider silk proteins, major ampullate spidroin 1 and 2 (MaSp1 and MaSp2).\textsuperscript{49,50} This approach was used to determine that alanine and glycine are both present in two conformational environments, an ordered \(\beta\)-sheet and disordered \(3_{10}\)-helical structure (see Fig. 4a and b). The poly(Ala) and flanking poly(Gly-Ala) repeats in the spider silk proteins form the \(\beta\)-sheet domains while, the Gly-Gly-X motif is present in a \(3_{10}\)-helical structure.

In addition to $^{13}$C/$^{13}$C through-space homonuclear correlation experiments, 2D $^{13}$C through-bond double quantum/single quantum (DQ/SQ) correlation spectra collected with the refocused INADEQUATE [(incredible natural abundance double quantum transfer experiment) solid-state NMR pulse sequence\textsuperscript{51} have also been used to characterize $^{13}$C-enriched spider silks.\textsuperscript{43–46} The DQ/SQ refocused INADEQUATE experiment greatly improves the resolution by doubling the chemical shift dispersion in the indirect, DQ dimension and has been shown to provide considerable resolution enhancement for disordered solids.\textsuperscript{52} For spider silks, the INADEQUATE approach has been used to structurally characterize Tyr in Gly-Gly-X motifs as random coil\textsuperscript{45} and the Pro-rich Gly-Pro-Gly-X-X regions unique to MaSp2 as elastin-like, type II \(\beta\)-turns\textsuperscript{46}

![Fig. 3](image-url)  
**Fig. 3** The 2D $^{13}$C/$^{13}$C correlation spectrum collected with DARR recoupling for $^{13}$C-labeled *Nephila clavipes* spider dragline silk. The spectrum was collected at 800 MHz with 40 kHz MAS, a 1 ms CP contact time and 1 s DARR mixing period with \(\tau_{RF} = \tau_{1R} \). The (a) high ppm, carbonyl and (b) low ppm regions of the spectrum are shown along with the projections in both dimensions. Intramolecular and intermolecular amino acid contacts are indicated with dashed and solid lines, respectively.

![Fig. 4](image-url)  
**Fig. 4** Secondary structures for the repetitive amino acid motifs found in the spider silk proteins, MaSp1 and MaSp2. Poly(Ala) and poly(Gly-Ala) repeat units present in both proteins form (a) \(\beta\)-sheet structures, the repetitive Gly-Gly-X motif of MaSp1 in a (b) \(3_{10}\)-helical structure and the (c) elastin-like type II \(\beta\)-turn structure for the Gly-Pro-Gly-X-X motif unique to the MaSp2 protein.
(see Fig. 4c). Heteronuclear 2D $^{13}$C–$^{15}$N solid-state NMR spectra collected with double CP (DCP) were used to obtain further structural information from the $^{15}$N dimension and confirm that the Pro-rich motifs in spider silk display nearly identical $^{13}$C and $^{15}$N chemical shifts to those observed for elastin mimics. The presence of elastin-like Pro-containing regions in spider dragline silk helps explain the silk’s elongation properties and overall high toughness.

3.2. Quantitative correlation between protein sequence and secondary structure

Utilizing NMR chemical shift assignments and correlations from the 2D through-space $^{13}$C–$^{15}$N correlation spectrum and 2D through-bond $^{13}$C–$^{15}$N refocused INADEQUATE experiments, significant progress has been made in determining the secondary structures of various amino acid motifs that together make up dragline spider silk. Most dragline spider silks are very similar in amino acid composition and overall structure. We have chosen to focus on Black Widow (Latrodectus hesperus) dragline (major ampullate) silk (BW MA) for quantitative correlation because it is the first and only spider silk that has been fully DNA sequenced (rather than just partial cDNA sequenced, which is available for numerous spider silks). For BW MA silk, alanine was determined to be in three distinct secondary conformations, a $\beta$-sheet, 3$_{10}$-helix, and $\alpha$-helix. Proline was ascribed to a $\beta$-turn structure. Glycine was found to be in two discrete secondary structures, a $\beta$-sheet and a 3$_{10}$-helix. Additionally, the Gly in a helical conformation was determined to be spatially close to both Ala in a $\beta$-sheet and Ala in a helical structure indicating that the helical portion of Gly is an intermediate between the two. Finally, Ser was found to also be in three distinct structures, a $\beta$-sheet, helix, and turn. As an example, we show the 2D $^{13}$C–$^{15}$N INADEQUATE NMR spectrum that was used to identify the three Ser structures in Fig. 5.

![Fig. 5 2D $^{13}$C–$^{15}$N through-bond DQ/SQ refocused INADEQUATE NMR spectrum of U–$^{13}$C/$^{15}$N-serine enriched Latrodectus hesperus major ampullate (dragline) silk. The three distinct serine resonances are assigned to serine incorporated in three distinct secondary structures, a $\beta$-sheet, helix, and $\beta$-turn.](image)

These secondary structural assignments were then quantitatively correlated and related to the primary amino acid sequences for both MaSp1 and MaSp2 in BW MA silk. Utilizing solid-state NMR, this study determined that 88% of Ala, 40% of Gly, and 42% of Ser in BW MA silk is incorporated into a $\beta$-sheet secondary structure. These values are in good agreement with the predicted values from the primary amino acid sequences of 88%, 42%, and 41% for Ala, Gly, and Ser, respectively. In total, solid-state NMR shows that 47% of all BW MA silk is incorporated into a $\beta$-sheet secondary structure. Further comparisons between quantitative NMR and amino acid sequences are underway as this first study to quantitatively correlate the protein sequence and secondary structure in BW MA silk has been very encouraging.

3.3 Hydrogen-bonding from solid-state NMR

Hydrogen-bonding is the primary interaction responsible for the secondary and tertiary structure of proteins. Solution and solid-state NMR spectroscopy have played a considerable role in detecting and quantifying hydrogen-bond lengths in small molecules and biopolymers. The advent of very fast (35–40 kHz) and ultra-fast (>60 kHz) MAS solid-state NMR probes has allowed for improved resolution in the 1H dimension by averaging the strong 1H–1H dipolar interactions in solids with rapid MAS. Sufficient resolution in the 1H dimension is obtained with these rapid MAS rates when the experiments are conducted at high magnetic field strengths (>600 MHz). Recently, the experimental results from fast and ultra-fast MAS 1H NMR spectra have been used in conjunction with theoretical 1H chemical shift calculations to determine hydrogen-bond lengths from amide proton chemical shifts for silk model peptides.

2D 1H–$^{13}$C heteronuclear correlation (HETCOR) solid-state NMR spectra collected with fast MAS were obtained for $^{13}$C-labeled spider dragline silk to probe the backbone conformation and hydrogen-bonding from the 1H dimension. The greater chemical shift dispersion of the $^{13}$C dimension is required to extract the 1H spectrum in an amino acid specific manner for the spider silk proteins. The 1H spectra extracted for the two Ala C$\beta$ components and the Gly C$\alpha$ resonance are shown in Fig. 6.

At the intermediate (1 ms) CP contact time used in the 1H–$^{13}$C HETCOR experiment all proton spins are observed for a given amino acid (i.e. Hz, H$\beta$ and NH). The 1H dimension for the two Ala C$\beta$ components displays different Hz chemical shifts where the slices extracted at 21.0 ppm and 17.4 ppm in the $^{13}$C dimension display Hz shifts of 5.1 and 4.2 ppm, respectively (see Fig. 6a and b). The observed 1H shifts for the Ala environments agree with previous 1H solid-state NMR on polypeptides where $\beta$-sheet structures had shifts of 5.1–5.5 ppm, while helical structures displayed lower shifts of 3.9–4.0 ppm. Thus, the Hz resonance can be used in combination with $^{13}$C chemical shifts to characterize the conformational structure in spider silks.

The 1H spectra extracted for both Gly and Ala display two distinct amide proton chemical shifts indicating two
hydrogen-bonding environments for both residues (see Fig. 6). The amide proton chemical shift can be dependent on both the hydrogen-bond strength and the backbone conformation. A series of density functional theory (DFT) NMR chemical shift calculations were conducted on model poly(Gly) peptides having β-sheet, 3_10-helical and α-helical structures with varying hydrogen-bond lengths to discern the contribution of conformation and hydrogen-bonding. The results of these calculations are presented in Fig. 7. From the DFT calculations, the amide proton chemical shift trend could be determined for β-sheet and helical structures as a function of hydrogen-bond length and the following equations were extracted from the calculated data,

\[ \delta_{\text{NH}} = 26.8d^3 + 4.7, \beta\text{-sheet} \]  
\[ \delta_{\text{NH}} = 25.3d^3 + 4.1, \alpha\text{-helix} \]  

where \( \delta_{\text{NH}} \) is the amide \(^1\text{H} \) chemical shift and \( d \) is the NH···OC hydrogen-bond distance. Using these equations the hydrogen-bond strength was determined to be equivalent for β-sheet and helical structures in spider silk with a 1.83–1.84 Å NH···OC hydrogen-bond distance. This hydrogen-bond is shorter compared to those previously reported for silk peptide mimics. In addition, this hydrogen-bonding is consistent with inter-strand interactions and provides some of the first pieces of evidence for intermolecular interactions in spider silk.

### 4. Other silks

While the vast majority of molecular level structural characterization and NMR studies of silk has been done on silkworm and spider silks, silk is a natural protein fiber that is produced by numerous insects and arthropods, both in terrestrial and aqueous environments. Understanding the molecular level differences in silks across a range of species and environments is a critical area where NMR is starting to play a major role. To date, NMR has been used to discern the secondary structure in a number of less studied silks including embiid, caddisfly, hornet, bee, and praying mantis based silk. Most of the NMR to date has been simple \(^1\text{H}–^{13}\text{C} \) CP-MAS solid-state NMR. More informative but complicated NMR can be accomplished, but typically requires isotopic enrichment of \(^1\text{H}, ^{13}\text{C} \) and/or \(^{15}\text{N} \), which presents a lot of individual technical challenges for each species. As an example of silk other than silkworm or spider silk that has been well characterized by NMR, we present a brief summary of NMR data on caddis y larval silk, which is spun under water. \(^1\text{H}–^{13}\text{C}–^{31}\text{P} \) DCP-MAS solid-state NMR was used to elucidate the molecular protein structure of caddisfly larval silk from the species Hesperophylax consimilis. Results provide strong evidence for a structural model in which phosphorylated serine repeats (psX)_n complex with divalent cations (Fe\(^{2+}\), Ca\(^{2+}\) and Mg\(^{2+}\)) to form rigid nanocrystalline β-sheet structures. Furthermore, \(^{13}\text{C} \) NMR data indicates that both phosphorylated serine and neighboring valine residues exist in a β-sheet conformation, while glycine and leucine residues common in GGX repeats reside in random coil structures. Additionally, \(^{31}\text{P} \) chemical shift anisotropy (CSA) analysis shows that the phosphates on phosphoserine residues are doubly ionized, and are charge-stabilized by divalent cations. The resulting model for the molecular level structure and architecture of caddis y larval silk is a critical area where NMR is starting to play a major role.
the nanocrystalline β-sheet structures in caddisfly silk is shown in Fig. 8.

Another example of an interesting silk that is being studied by NMR is embiid (webspinners) silk. Individual embiid silk fibers are typically 50–80 nm in diameter and do not contain poly-A or poly-GA motifs like in spider silk and silkworm silk, respectively. In this silk, we find serine-based motifs in nanocrystalline β-sheet domains. Furthermore, this silk is waterproof and the high surface area allows direct observation of the lipids found on the surface of this silk. As NMR has developed as the premier technique for molecular level structural characterization in silkworm and spider silks, it is likely to also become an important characterization tool in all types of terrestrial and aquatic silk materials.

Wild silkworm silks are also interesting. The amino acid composition of silk fibroin from a wild silkworm, S. c. ricini and A. pernyi, is considerably different from that of B. mori silk fibroin. The proportion of Gly residues is greater in B. mori silk fibroin, while the content of Ala residues is greater in S. c. ricini silk fibroin. The solution structure of S. c. ricini silk fibroin has been studied with solution 13C and 15N NMR. The fast exchange in the NMR time scale between helix and coil forms of the poly(Ala) region has been observed during the helix-to-coil transition with changing temperature. Moreover, solid-state NMR analysis of the 13C-labelled appropriate model peptides leads to the precise silk structure before spinning, where the poly(Ala) sequence takes a typical α-helix pattern with a tightly wound helical structure at both terminal regions of the poly(Ala) sequence. Namely, the 2D spin-diffusion NMR spectra were observed for several [1-13C] double labeled Ala residues in the model peptides. The agreement between observed and simulated spectra is good and therefore the structural model of poly(Ala) sequence is proposed as shown in Fig. 9. The silk fiber after spinning is in a β-sheet structure as determined from the 1H-13C CP-MAS NMR spectrum.

Anaphe form a very large single common silk shell in which individuals then form their own small cocoons. The sequence predominantly contains a mixture of (AAG)1 or (AG)n. The solution and solid state structure were examined with 13C solution and solid state NMR, respectively. The characteristics of the NMR behavior of Anaphe silk were between those of B. mori and S. c. ricini silks.

5. Peptides with silk sequence and recombinant silk protein

The peptides with silk-like sequences are (AG)n or (AGSGAG)n for the crystalline domain of B. mori silk fibroin and poly(Ala) for S. c. ricini and spider silks. In this section, the lamellar structure was proposed for the former peptides, (AG)n or (AGSGAG)n. For the latter peptides, details of the poly(Ala) β-sheet structure with different inter-molecular structures were discussed. The recombinant silk protein based on these silk sequences and their structural analysis with solid-state NMR were reported. These studies have been performed by Asakura’s group for 10 years and applied to biomaterials.

5.1 Lamellar structure of (AG)15 with silk II structure

When (AG)15 is dissolved in formic acid and then dried, it adopts a silk II antiparallel β-sheet structure. A lamellar structure has been proposed, based on changes in the intensities of asymmetric Ala 13Cβ peaks in the 13C CP-MAS NMR spectra coupled with selective 13C labeling of different Ala methyl carbons. As shown in Fig. 10A, the relative intensities of the peaks at 16.7 ppm which were assigned to the distorted β-turn change largely depending on the labeled position of the Ala residue. Therefore the relative intensity was plotted against the residue number of the labeled [3-13C]Ala in Fig. 10B. The plot of the fraction indicates two maxima at the positions, 9 and 19. This implies the appearance of the folded lamellar structure with a β-turn at these positions. Panitch et al. reported that
eight amino acid residues contribute to the β-sheet structure for (AG)n, which was studied by X-ray diffraction. This is in agreement with the NMR result. Statistical treatment for the appearance of lamellar structure in (AG)15 was also performed. A slightly modified lamellar structure was also proposed for the peptide (AGSGAG)n.

It is interesting to design new silk materials for bone repair scaffold based on the NMR study of the lamellar structure. The silk-like peptides based on AG repeated sequences with a lamellar structure and Asp as a Ca binding site at the turn part were designed and produced in E. coli. The lamellar structure formation was confirmed for the model peptide with different 13C labeling positions by solid-state NMR. The surfaces of the recombinant silk protein with the lamellar structure are responsible for calcium binding and accelerated formation of hydroxyapatite. Another B. mori silk-like protein, (E)n(AGSGAG)4 where n = 4–8, mimics the primary structure of B. mori silk fibroin and were designed and then structurally characterized with 13C solid-state NMR. The balance of hydrophilic and hydrophobic characters of the peptide was controlled by changing the relative length, n, of (E)n from 4 to 8. Then, genetically modified silk fibroin containing a poly-glutamic acid site for mineralization was produced as fibers by transgenic silkworms, indicating a mineralization-accelerating material for bone repair.

5.2 Polyalanine
Poly(Ala) sequences with β-sheet structure appear frequently in the crystalline regions of spider dragline silk or wild silkworm silks, S. c. ricini or A. pernyi. The poly(Ala) domains seem to be the origin of the high strength of these silk fibers. Because (Ala)3 can be crystallized in both parallel (P) and antiparallel (AP) β-sheet structures, and the atomic resolution structure was reported by X-ray diffraction studies, this peptide was used to show the usefulness of solid-state NMR including 13C Radio frequency driven recoupling (RFDR), high-field 1H and 17O solid-state NMR and 13C–17O Rotational Echo, Adiabatic Passage, Double Resonance (REAPDOR). Then a systematic NMR analysis of AP poly(Ala) longer than (Ala)3 was performed. In the 13C CP-MAS NMR spectra of (Ala)3, (Ala)4, (Ala)5 and (Ala)6 there is a single central Cβ peak at 20.4 ppm, surrounded by smaller peaks from the two termini (see Fig. 11). In the (Ala)6 spectrum there is also a broad peak at 17.0 ppm, assigned to disordered residues. For (Ala)7 and higher, 13C NMR spectra are markedly different from those for short poly(Ala). NMR shows two Cβ peaks at 22.7 and 19.6 ppm. Line fitting indicated that there is also a third broad peak at about 21.0 ppm, plus the broad peak at 17.0 ppm seen for (Ala)6. It is therefore clear that there is a ‘long form’ packing, which is different from the ‘short form’. With further details from DARR and REDOR experiments for (Ala)7, it is possible to propose a model for the long form. The main difference from the short form is that the packing of chains in adjacent planes is staggered rather than rectangular. High field 1H MAS and 15N CP-MAS NMR were also applied to analyze the structure and hydrogen-bond formation.

5.3 Recombinant silk protein
B. mori silk fibroin can be used for various biomaterial applications. One application is coating the surface of cell culture plates. However, the cell-adhesive ability is weaker than collagen or fibronectin, which are currently used for coating cell culture plates. To increase the biocompatibility of the silk,
recombinant silks that possessed partial collagen or fibronectin sequences, that is, [GERGLDPQGIAQGRGTVG(GER),]$_2$-GAS$_2$GP GCCGGG or [TGRGDPSA], respectively were produced by transgenic silkworm. The two types of recombinant silk films possessed a much higher cell-adhesive activity than unmodified silk. Thus, by introducing the cell-adhesive sequence, RGD from fibronectin into silk fibroin by bioengineering techniques, excellent biomaterials have been developed. To clarify the origin of such a high cell adhesion character and to design new recombinant silk protein with higher cell adhesion ability, the structure and dynamics of the RGD moiety in a recombinant silk-like protein consisting of the repeated silk fibroin sequence ([AGSGAG])$_3$, and the sequence ASTGRGDSPA including the RGD moiety, were studied using NMR. A $^{13}$C solid-state NMR study on the $^{13}$C selectively labeled model peptide indicates that a random coil state of the RGD moiety is maintained in aqueous solution, and also in both labeled and unlabeled model peptides. An NMR study of the RGD moiety included the RGD sequence, that is, [GERGDLGPQGIAGQRGVV(GER)]$_3$ and the sequence GASS$_8$GGPPGPCCGGG or [TGRGDPSA]$_8$, respectively were introduced between di-GAS and di-ASTGRGDSPA including the RGD moiety, were studied using NMR. The two types of recombinant proteins with characteristic sequences from the silk fibroin of Anaphe, and the cell adhesive region, including the RGD sequence were also designed and produced from E. coli. They showed higher cell adhesion activity than Anaphe silk fibroin without the RGD sequence. In addition, the activities were very similar to that of collagen, which acted as a positive control. Thus, it is demonstrated that the primary structure of Anaphe silk fibroin, which is composed largely of Ala and Gly residues, can be used as a platform for the basic structures of silk-like cell adhesive proteins. The structural characterization of the silk-like recombinant proteins was performed with $^{13}$C CP-MAS NMR. The calcium-binding site of the pearl oyster (Pinctada fucata) nacreous layer matrix protein MSI60 and calbindin D$_{9k}$ was introduced between different Ala-Gly repeating regions derived from the primary sequences of several silk fibroins. The local conformations of the flanking Ala-Gly repeating regions as well as the calcium-binding motif, MSI60, were determined by $^{13}$C CP-MAS NMR spectroscopy. The recombinant proteins of silk-elastin and B. mori silk-syder silk were also produced and characterized by $^{13}$C solid-state NMR.

Acknowledgements

T.A. acknowledges support from Grant-in-Aid for Scientific Research from Ministry of Education, Science, Culture and Supports of Japan (23245045,25620169) and Ministry of Agriculture, Forestry and Fisheries of Japan (Agri-HealthTranslational Research Project). GPH and JLY acknowledge support by grants from the Department of Defense (DOD) Air Force Office of Scientific Research (AFOSR) under Award no. FA9550-10-1-0275, the Defense University Research Instrumentation Program (DURIP) under Award no. FA2386-12-1-3031 DURIP 12RSL231 and the National Science Foundation (NSF) Division of Materials Research (DMR) under Award no. DMR-1264801.

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