Structural Comparison of Mouse and Human α-Synuclein Amyloid Fibrils by Solid-State NMR

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Fibrillar α-synuclein (AS) is the major component of Lewy bodies, the pathological hallmark of Parkinson's disease. Mouse AS (mAS) aggregates much faster than human AS (hAS), although mAS differs from hAS at only seven positions in its primary sequence. Currently, little is known about the site-specific structural differences between mAS and hAS fibrils. Here, we applied state-of-the-art solid-state nuclear magnetic resonance (ssNMR) methods to structurally characterize mAS fibrils. The assignment strategy employed a set of high-resolution 2D and 3D ssNMR spectra recorded on uniformly [13C, 15N], [1-13C]glucose, and [2-13C]glucose labeled mAS fibrils. An almost complete resonance assignment (96% of backbone amide and 93% of all 13C nuclei) was obtained for residues from Gly41 to Val95, which form the core of mAS fibrils. Six β-strands were identified to be within the fibril core of mAS based on a secondary chemical shift and NHHC analysis. Intermolecular 13C,15N labeled restraints obtained from mixed 1:1 13C/15N-labeled mAS fibrils reveal a parallel, in-register supramolecular β-sheet arrangement. The results were compared in detail to recent structural studies on hAS fibrils and indicate the presence of a structurally conserved motif comprising residues Glu61–Lys80.

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Introduction

Parkinson's disease (PD) is a neurodegenerative pathology characterized by the loss of dopaminergic neurons in the substantia nigra region of the brain and the formation of Lewy bodies (LBs) whose major component is α-synuclein (AS) in the form of amyloid fibrils. The aggregation of the 140-residue-long cytoplasmic protein AS is thus believed to play an important role in the etiology of PD. AS has been classified as a “natively unfolded” or disordered monomeric protein, which can adopt α-helical structure in solutions that contain lipid-mimetic detergent micelles or in the presence of lipid vesicles. The primary sequence of mouse AS (mAS) differs from human AS (hAS) at seven positions (Fig. 1b). An in vitro study showed that mAS has a “natively unfolded” structure in solution.
similar to hAS, whereas at elevated concentrations, mAS forms amyloid fibrils with predominant β-sheet secondary structure much more rapidly than its human counterpart.9 Currently, this enhanced fibrillation is not well understood. Recently, an in vivo study indicated that endogenous hAS occurs physiologically as a folded tetramer with predominant helical secondary structure even without the addition of lipid vesicles.10 Although there could be a big difference of the soluble form between in vitro and in vivo studies, in vitro fibrils have been shown to possess a morphology that closely resembles the one found in LBs of patients with PD.11–13 Because LBs are a pathological hallmark of PD, extensive research has focused on the structural characterization of AS fibrils.11–18

Magic-angle spinning solid-state nuclear magnetic resonance (ssNMR) spectroscopy has emerged as a powerful tool for the characterization of noncrystalline and insoluble proteins,19 including membrane proteins,20–25 oligomeric assemblies,26–31 and amyloid fibrils.32–39 Extensive studies on hAS fibrils using ssNMR revealed that the central fibril core extends at least from about residue Leu38 to about residue Val95 with mostly β-sheet secondary structure.14–18 However, much less is known about the structural properties of mAS fibrils. Rochet et al. demonstrated the presence of β-sheet secondary structure in mAS fibrils, but the extent of the core region and site-specific structural information remained absent.9 Meanwhile, one recent study on the detailed sequence-dependent fibrillation differences between mAS and hAS revealed that the Ala53Thr substitution dominates the aggregation growth rates, while the combination of the Ala53Thr and Ser87Asn mutations affects the lag phase of the aggregation.40 Nevertheless, so far no structural details of mAS fibrils and their differences to hAS fibrils were available which motivated the present study.

Here, we aimed at the structural characterization of mAS fibrils using state-of-the-art ssNMR. As a major bottleneck in structural studies of fibrillar proteins by ssNMR remains the unambiguous and confident sequential resonance assignment that is often difficult due to poorly resolved and highly crowded ssNMR spectra. Furthermore, structural polymorphism can lead to the phenomenon that different sample preparations exhibit slightly different ssNMR spectra. The combination of a highly reproducible sample preparation, tailored isotope labeling schemes, and state-of-the-art ssNMR methods enabled us to obtain the near-complete sequential resonance assignment and a subsequent

Fig. 1. Initial comparison of mAS and hAS fibrils. (a) Electron micrograph of [U-13C/15N]-labeled mAS fibrils used for ssNMR experiments. (b) Primary sequence alignment of mAS and hAS. The seven mutation sites are indicated with green boxes. The six highly repetitive, imperfect KTKEGV motifs are shown in black boxes. (c) Comparison of 2D PDSD spectra of [U-13C/15N]-labeled hAS fibrils (black) and [U-13C/15N]-labeled mAS fibrils (red, reproduced from Ref. 46). Correlations of residue Ile88 for mAS and hAS are illustrated with broken and continuous lines, respectively. mAS and hAS fibrils were obtained under identical fibrillation conditions.
Comparison of Mouse and Human Amyloid Fibrils

structural characterization of the fibril core of mAS fibrils. For this purpose, we prepared uniformly \([1^{13}C, 1^{15}N] - [U-^{13}C/^{15}N] \) labeled mAS fibrils, and two complementary sparsely \(^{13}C\)-enriched variants of mAS fibrils. Bacterial growth in media containing \([1^{13}C]\)glucose (\([1^{13}C]\)Glc) or \([2^{13}C]\)glucose (\([2^{13}C]\)Glc) results in highly diluted \(^{13}C\) labeling in the proteins with only one out of six carbons labeled. Therefore, this leads to enhanced resolution and a reduced number of cross-peaks that facilitates the sequential resonance assignment considerably.

Highly confident assignment of 96% of backbone amide \(^{15}N\) and 93% of all \(^{13}C\) atoms of the detected residues from Gly41 to Val95 was obtained. Six \(\beta\)-strands were identified to be within the fibril core of mAS fibrils by secondary chemical shift and NHHC analysis. Additionally, by measurement of intermolecular \(^1H/\(^1H\) correlations using the NHHC scheme on a mixed \(1:1 \) \(^{13}C:\(^{15}N\) \([M-^{13}C/^{15}N]\) variant of mAS fibrils, we could deduce the supramolecular arrangement of mAS fibrils. We found that they are stacked parallel and in-register. This result is in agreement with our recent investigation of hAS fibrils prepared under identical fibrillation conditions to the mAS fibrils studied here and that were also found to adopt a parallel, in-register \(\beta\)-sheet arrangement.

Results and Discussion

Morphological characterization by EM and initial comparison to hAS fibrils

In this study, \([U-^{13}C/^{15}N]\), \([1^{13}C]\)Glc-, and \([2^{13}C]\)Glc-labeled mAS fibrils were prepared under identical conditions to obtain the sequential resonance assignment. The morphology of the three samples was then monitored by electron microscopy (EM). The mAS fibrils were all well ordered, and long straight fibrils with a diameter around 160 \(\text{Å}\) dominate all three samples (as an example, an EM micrograph of \([U-^{13}C/^{15}N]\)-labeled mAS fibrils is shown in Fig. 1a). Rarely, twisted fibrils were observed in \([U-^{13}C/^{15}N]\)-labeled mAS fibrils, and no twisted fibrils were observed in \([1^{13}C]\)Glc- or \([2^{13}C]\)Glc-labeled mAS fibrils. The highly similar morphology of the three samples indicates high reproducibility of our sample preparation, which is an essential requirement if different labeling schemes are to be used for ssNMR. However, both straight and helically twisted fibrils of mAS were observed by EM in the recent study by Kang et al. Therefore, the observed differences of the mAS fibril morphology might be due to different fibrillation conditions and suggests the existence of two types of mAS fibrils (straight and twisted). In the case of hAS fibrils, the overall morphology was demonstrated to be strongly dependent on the fibrillation conditions, and both straight and twisted fibrils were observed and characterized.

Nevertheless, the highly uniform morphology of mAS fibrils obtained by our preparation indicates high molecular homogeneity of our samples, and indeed, no detectable polymorphism was observed in our recorded ssNMR spectra (see below). The absence of polymorphism allowed us to obtain sequential resonance assignments of the core region of mAS fibrils with straight morphology.

The primary sequence of mAS differs from hAS at seven positions (Fig. 1b). Despite the highly conserved sequence, the aggregation kinetics of mAS and hAS are remarkably different. mAS aggregates about twice faster, and the lag phase is reduced by almost 1 order of magnitude. To characterize the structural differences between mAS and hAS fibrils at the atomic level, we recorded a set of high-resolution ssNMR spectra on mAS fibrils and compared it to existing data for hAS fibrils that were prepared under identical conditions. As an example, Fig. 1c shows the comparison of two-dimensional (2D) PDSD (proton-driven spin diffusion) spectra with a mixing time of 20 ms recorded on hAS (red) and mAS (black) fibrils. The hAS fibrils considered here exhibit the same morphology as mAS fibrils (i.e., straight). From the spectral comparison, a difference between mAS fibrils and hAS fibrils is obvious, as illustrated with the straightforward Ile88 assignment in Fig. 1c. Since the amino acid sequence of mAS differs from hAS at position 87 (Ser versus Asn) (Fig. 1b), different chemical shifts of Ile88 due to the disturbance of the chemical environment by the Ser87Asn mutation are expected. A more detailed comparison between mAS and hAS fibrils requires the sequential resonance assignment of mAS fibrils as described in the following section.

Sequential resonance assignment using high-resolution ssNMR spectra

The sequential resonance assignment strategy using 2D \(^{13}C/^{13}N\) and \(^{13}C/^{13}C\) correlation spectra recorded on \([U-^{13}C/^{15}N]\)-labeled mAS fibrils is illustrated in Fig. 2. The recorded NCAX, NCOCX, and 2D PDSD spectra with mixing times of 20 ms and 150 ms were well resolved, and the \(^{13}C\) and \(^{15}N\) linewidths were found to be \(\sim 0.5\) ppm and \(\sim 1.3\) ppm, respectively. As an example, the sequential resonance assignment of Gly41-Ser42-Lys43 is illustrated in Fig. 2. Additionally, three-dimensional NCAX and NCOCX spectra were recorded to confirm and obtain more unambiguous assignments (Fig. S1). Taken together, the combination of 2D and three-dimensional \(^{13}C/^{15}N\) and \(^{13}C/^{13}C\) spectra of \([U-^{13}C/^{15}N]\)-labeled mAS fibrils enabled us to obtain the sequential connections for the following stretches unambiguously:
Gly41–Val48, Ala56–Lys60, Asn65–Val70, and Thr72–Phe94.

The limited resolution and spectral overlap observed for the [U-13C/15N]-labeled sample are major obstacles obtaining the complete sequential resonance assignment. Furthermore, the full sequential resonance assignment for mAS fibrils is complicated not only due to the highly repetitive, imperfect KTKEGV motifs (Fig. 1b) but also due to the similar secondary structure adopted by most residues in the β-sheet arrangement. Another difficulty to complete the full assignment of mAS fibrils relates to the high abundance and sequential clustering of Val and Thr, for example, Val48–Val49, Val52–Thr53–Thr54–Val55, Val70–Val71–Thr72, Val74–Thr75, and Thr81–Val82, since the Cα and Cγ chemical shifts of Val and Thr are similar. In order to alleviate spectral overlap and enhance the resolution, we used sparse 13C labeling schemes. In the sample of [2-13C]Glc-labeled mAS fibrils, 13Cα is labeled for 17 out of 20 residues without simultaneous 13C labeling at Cβ. With this labeling scheme, not only the resolution is enhanced substantially due to the removal of one-bond 13C–13C J and dipolar couplings but also the sequential transfer of 13Cα(i)–15N(i) is facilitated by the 13C spin dilution.

Figure 3 illustrates the sequential backbone resonance assignment strategy based on 2D 13C/13C and 13C/15N correlation spectra recorded on [2-13C]Glc-labeled mAS fibrils. The observed linewidths for 13C and 15N are ~0.3 ppm and ~0.5 ppm, respectively. The aliphatic and carbonyl regions of a 2D PDSD spectrum with long mixing time of 500 ms are dominated by 13Cα(i)–13CO(i−1) correlations, which can be easily connected to 15N(i)–13Cα(i) correlations in the NCA spectrum (Fig. 3c). The 13CO(i−1) resonances can be identified via the
$^{13}$C$_\alpha$ resonances in the carbonyl region of the 2D PDSD spectrum (Fig. 3a) and/or from the $^{15}$N($i$) resonances in the NCO spectrum (Fig. 3b). As an example, the broken arrows show the sequential connections from Gly41 to Thr44, and the continuous arrows show the intra-residue connections for residues Ser42 to Thr44 (Fig. 3). It is worth to note that $^{13}$C$_\alpha$ and $^{13}$C$_\beta$ are labeled simultaneously for Val using this labeling scheme, which enabled us to obtain and confirm the assignments of Val by sequential $^{13}$C$_\alpha$($i$)-$^{13}$C$_\alpha$($i\pm1$) correlations (Fig. 4). Also, the presence of $^{15}$N($i$)-$^{13}$C$_\beta$($i$) correlations for Val in the NCA spectrum due to the attenuation of dipolar truncation effects by the sparse $^{13}$C labeling scheme allowed for the unambiguous assignments of $^{15}$N($i$)-$^{13}$C$_\alpha$($i$) correlations of Val, although they heavily overlap with the ones of Thr (Fig. S2). In summary, the sequential walks involving $^{13}$C$_\alpha$($i$)-$^{13}$C$_\alpha$($i\pm1$) correlations and/or $^{13}$C$_\beta$($i$)-$^{13}$C$_\alpha$($i\pm1$) (for Val) correlations lead to sequential resonance assignments for the stretches Gly41–Val49, Gly51–Val52, Thr54–Lys60, Gln62–Val70, Thr72–Val74, and Thr75–Val95 without ambiguity (Fig. 4). In addition, doubling of resonances for the residues Gly84, Ala85, Gly86, Ala89, Gly93, and Phe94 was revealed by the sufficient resolution resulting from the sparse glucose labeling scheme (Fig. 4). A single set of resonances is observed for the other residues, indicating that no polymorphism exists in our samples, whereas the presence of the few doubled resonances may suggest local disorder of these.
residues. This correlates quite well with their location in the secondary structure of mAS fibrils: Gly84 and Ala85 are close to one loop that includes Gly86 and Ala89, while Gly93 and Phe94 are located at the end of the structured region of mAS fibrils (see the discussion about the secondary structure analysis below).

As a perfectly complementary labeling scheme, [1-13C]Glc-labeled mAS fibrils were prepared to identify and/or confirm both backbone and side-chain assignments of Thr, Asn, Asp, Gln, Glu, Lys, and Ile. Figure S3 shows the assignment of Thr and Val based on the combination of a PDSD spectrum with a mixing time of 700 ms recorded on [2-13C]Glc-labeled mAS fibrils and PDSD (mixing time: 400 ms), NCO, and NCA spectra recorded on [1-13C]Glc-labeled mAS fibrils. Broken lines in Fig. S3 illustrate the unambiguous...
assignment of the critical residue Thr53, one of the two variant sites between hAS and mAS located in the fibril core.

Taken together, the near-complete sequential resonance assignment for residues from Gly41 to Val95 is obtained with all the recorded spectra. In total, 96% of backbone amide $^{15}$N and 93% of all $^{13}$C resonances (97% of the backbone and 89% of the side chains) are obtained for the detected residues from Gly41 to Val95 (Table S1). To the best of our knowledge, our study reports the sequential resonance assignments of mAS fibrils for the first time and with more complete information compared to hAS fibrils.

![Diagram](attachment:Figure5.png)

**Fig. 5.** Secondary structure analysis of the rigid core of mAS fibrils and the comparison to hAS fibrils. Secondary chemical shifts ($\Delta C^\alpha - \Delta C^\beta$) are shown as a function of residue for (a) mAS fibrils as studied here and hAS fibrils as investigated by (b) Gath et al.,\textsuperscript{18} (for the sake of clarity, the values for residues Met1–Leu38 are omitted), (c) Comellas et al.,\textsuperscript{17} and (d) Heise et al.\textsuperscript{14} (B-Form hAS fibrils with straight morphology). For hAS fibrils, no $^{13}$C$^\beta$ assignment was available for Val66 and Phe94 [in (b)] and for His50, Lys60, Lys96, and Lys97 [in (c)]. Non-assigned residues are marked with a cross. The secondary structure of the rigid core of mAS fibrils is illustrated in (a). $\beta$-Strands are indicated by white arrows (slashed bars in the white arrow represent possible kinks), non-$\beta$-strand regions (kink, loop, or turn) are shown as a curve, and non-assigned amino acids are shown as a broken line. The Ala53Thr and Ser87-Asn variant sites in the rigid core are colored red. The conserved region of AS fibrils are revealed by the comparison of mAS and hAS fibrils studied by Gath et al.\textsuperscript{18} and Comellas et al.\textsuperscript{17} is highlighted in gray.
Secondary structure analysis of the fibril core of mAS fibrils and detailed comparison to hAS fibrils

The secondary structure of the rigid core of mAS fibrils was analyzed based on secondary chemical shifts \[ (\Delta C^\beta - \Delta C^\alpha) = [\Delta C^\alpha \text{(obs)} - \Delta C^\alpha \text{(rc)}] - [\Delta C^\beta \text{(obs)} - \Delta C^\beta \text{(rc)}] \] and is illustrated in Fig. 5a. For all the residues apart from Gly, three or more continuous negative values of \((\Delta C^\beta - \Delta C^\alpha)\) are indicative of \(\beta\)-strand conformation, while positive values point to \(\alpha\)-helical structure, and values close to 0 indicate a random-coil conformation. As an independent and complementary indicator, backbone \(^1\text{H} / ^2\text{H} \) distances were probed indirectly using the NHHC scheme \(^{14}\) to characterize the secondary structure of mAS fibrils (Fig. S4). Longitudinal \(^1\text{H}/^2\text{H} \) mixing times of 150 \(\mu\)s and 175 \(\mu\)s for [U-\(^{13}\text{C}/^{15}\text{N}\)- and [2-\(^{13}\text{C}\)]Glclabeled mAS fibrils, respectively, were used. The enhanced resolution of the NHHC spectrum of [2-\(^{13}\text{C}\)]Glclabeled mAS fibrils compared to the spectrum of [U-\(^{13}\text{C}/^{15}\text{N}\)-labeled mAS fibrils facilitated the site-specific identification of secondary structure. Observed \(^{15}\text{NH}(i+1)-^{13}\text{C}^\alpha\text{H}(i)\) correlations with stronger intensities compared to \(^{15}\text{NH}(i)-^{13}\text{C}^\beta\text{H}(i)\) correlations or the exclusive presence of \(^{15}\text{NH}(i+1)-^{13}\text{C}^\beta\text{H}(i)\) correlations, as seen, for example, for Ser42 and Lys43, are indicative of \(\beta\)-sheet conformation. In contrast, the clear absence of \(^{15}\text{NH}(i+1)-^{13}\text{C}^\beta\text{H}(i)\) correlations or the presence with weaker intensities compared to \(^{15}\text{NH}(i)-^{13}\text{C}^\alpha\text{H}(i)\) correlations, for example, observed for Lys58 and Thr59, is indicative of loop or turn structure. Together with the secondary chemical shifts, six \(\beta\)-strands were identified to be within the fibril core of mAS fibrils: Gly41–Val49, Val52–Glu57, Lys60–Val66, Gly68–Gln79, Thr81–Ala85, and Ala90–Val95. Possible kinks in the \(\beta\)-strands are marked by slashed bars (Fig. 5a).

A direct comparison between mAS and hAS fibrils has to take into account the existence of multiple sets of chemical shifts, incomplete sequential resonance assignments, and the presence of different morphologies of hAS fibrils. For example, 48 residues (form A, twisted fibrils) and 36 residues (form B, straight fibrils) from Leu38 to Val95 were assigned and identified to be within the fibril core of hAS for two distinct fibril morphologies by Heise et al. \(^{14}\) Recently, sequential resonance assignments for 85% of residues from Met1 to Lys97 were achieved by Gath et al. \(^{18}\) and 91% of the \(^{15}\text{N} \) and \(^{13}\text{C} \) resonances of residues from Leu38 to Lys96 were reported by Comellas et al. \(^{17}\) Collectively, hAS fibrils seem to recruit more residues into the \(\beta\)-sheet fibril core than mAS fibrils. Residues from Gly41 to Val95 were sequentially assigned and identified to be within the fibril core of mAS fibrils by our current work. The assignment of Val40 was obtained ambiguously. Unambiguous assignments for residues of Leu38 and Tyr39 could so far not be obtained due to a lack of sequential correlations. This might be because these two residues are at the beginning of the structured region, and the peak intensities may thus be too weak to be observed. On the other hand, there are clearly no detectable sequential connections from Val95 to Lys96 in our spectra. However, we were able to find residues outside the identified fibril core in PDS spectra, such as Ser (Ser9 or Ser129), Pro (Pro108, Pro117, Pro120, Pro128, or Pro138), and Asp (Asp2, Asp98, Asp115, Asp119, or Asp135). However, we could not detect any sequential connections to assign them unambiguously. Importantly, the Ser and Pro resonances are absent in a DREAM (dipolar recoupling enhancement through amplitude modulation) spectrum (see Fig. S5), which may indicate that they are relatively flexible. This is corroborated by the observation that the Ser and Pro signals have random-coil chemical shifts, whereas the unassigned Asp exhibits \(\beta\)-strand-like chemical shifts (See Table S1).

The B-Form hAS fibrils studied by Heise et al. are straight, while the A-Form has a twisted morphology. \(^{14}\) However, the fibril morphology was not reported in the publications by Gath et al. \(^{18}\) and Comellas et al. \(^{17}\) on hAS fibrils. Because the ssNMR spectra are virtually identical for hAS fibrils as reported by Gath et al. \(^{18}\) and by us (see Fig. 1, straight fibrils, produced under the same conditions as the mAS fibrils with straight morphology reported in the current study), \(^{46}\) we assume that Gath et al. also obtained straight fibrils. However, no EM micrograph is shown in Gath et al. \(^{18}\) The fibril morphology for Comellas et al. is not clear. In the published EM micrograph (Comellas et al., \(^{17}\) Fig. 8), the fibrils appear straight as well. Interestingly, we see strong differences between mAS and hAS in terms of secondary chemical shifts and extent of \(\beta\)-strand regions if we consider hAS prepared under the same fibrillation conditions as mAS (with assignments for this form reported by Gath et al., \(^{18}\) see Fig. 5b). The most significant difference relates to residues Met1–Val37 that are only observed in spectra of hAS. In contrast, residues Thr44–Glu57 are only observed for mAS. Furthermore, for several residues in the core, secondary chemical shifts \((\Delta C^\beta - \Delta C^\alpha)\) with opposite sign for mAS and hAS were observed: Gly41, Lys58, Thr81, Ala89, and Ala91. In contrast, much higher similarity was observed between mAS fibrils and hAS fibrils studied by Comellas et al. (Fig. 5c), \(^{17}\) regarding both the length and the location of the \(\beta\)-strands. The only residues that adopt a significantly different secondary structure (i.e., opposite sign of the secondary chemical shift) in mAS and hAS fibrils studied by Comellas et al. \(^{17}\) (Fig. 5c) are Lys58, Thr59, and Ala85. The systematic structural comparison between mAS fibrils and the two different
forms of hAS fibrils reveals that residues Glu61--Lys80 constitute the structurally most conserved region of AS. Interestingly, the mutation sites of Ala53Thr and Ser87Asn that are responsible for the differences in aggregation kinetics and lag phases between mAS and hAS are outside this region. The conserved core of Glu61--Lys80 might play an important role in the initiation of the aggregation of AS fibrils. Consistent with this notion, it was previously reported that residues Glu61--Ala78,49 Gly68--Ala78,50 and Val71--Val8251 within the hydrophobic region of AS could form aggregates by themselves in vitro and that residues Gly68--Ala7850 constitute the shortest fragment sufficient for AS fibril assembly.

Supramolecular structure of mAS fibrils elucidated by means of [M-13C/15N]-labeled mAS fibrils

Previous ssNMR studies on heterogeneously labeled samples have yielded structural insight into the supramolecular arrangement of amyloid fibrils.33,43,52–58 Here, we have recorded an NHHC spectrum with a 1H/1H mixing time of 500 μs on [M-13C/15N]-labeled mAS fibrils to elucidate the stacking of molecules in the fibrils. With the chosen mixing time, cross-peaks between 13C and 15N sites that correspond to intermolecular (1H/1H) distances of about 3.5 Å can be observed.45 Since 15N and 13C spins in the [M-13C/15N]-labeled mAS fibrils are not labeled within the same molecule, all the observed cross-peaks in the NHHC spectrum are exclusively due to intermolecular transfer. As shown in Fig. 6a, most of the cross-peaks can be assigned to either inter-strand 15N(i)-13C"(i) or inter-strand 15N(i)-13C"(i−1) correlations. These contacts are summarized in Fig. 6b. The sensitivity of the intermolecular NHHC experiment is compromised due to (a) the three involved CP transfer steps; (b) the probability of one pair of neighboring molecules to be 15N/13C labeled, which is only 25%; (c) the long intermolecular distances involved; and (d) the distribution of magnetization during the 1H/1H transfer to protons that are not involved in the last transfer step. Still, the unambiguous intermolecular correlation pattern, for example, for Lys43, Glu46, and Glu57 (Fig. 6b), as well as the overall good overlay with the NCA spectrum recorded on [U-13C/15N]-labeled mAS fibrils (Fig. 6a), identifies a parallel, in-register arrangement of the β-strands. This result is in agreement with a similar recent study by our group showing that the hAS fibrils studied by us are stacked parallel and in-register.46

Conclusions

We recently reported a strategy for simplified and complete resonance assignment using the 13C spin dilution approach, demonstrated with the de novo assignment of the type three secretion system needle protein PrgI in the polymerized state, in which most residues adopt α-helical conformation.42 In the present work, as the second example, the substantially improved spectral resolution observed for [1-13C]Glc- and [2-13C]Glc-labeled mAS fibrils turned out to be highly useful for the near-complete sequential assignment of the β-sheet-rich amyloid fibrils formed by mAS.

The structural comparison between mAS fibrils and two different forms of hAS fibrils reveals that residues Glu61--Lys80 constitute the structurally most conserved region of AS. While the Glu61--Lys80 conserved core appears to play a critical role as an initiator of aggregation, a subtle modulatory effect might be exerted on the AS aggregation pathway by mutations located inside the extended fibrillogenic region formed by residues Gly41--Val95. Indeed, it was reported recently that the Ala53Thr substitution dominates the aggregation growth rates, while the combination of the Ala53Thr and Ser87Asn mutations affects the lag phase of the aggregation of mAS compared to hAS.40

Fig. 6. Supramolecular arrangement of mAS fibrils. (a) Comparison of NCA spectrum (in black) recorded on [U-13C/15N]-labeled mAS fibrils and NHHC spectrum (in red) recorded on [M-13C/15N]-labeled mAS fibrils. 15N(i)-13C"(i) correlations and 15N(i)-13C"(i−1) correlations between adjacent molecules were labeled as, for example, Val63 and Val63--Gln62, respectively. (b) Summary of the detected intermolecular contacts. Continuous lines represent unambiguous assignments, and broken lines represent ambiguous assignments due to spectral overlap.
It should be worthwhile to determine structures of mAS and hAS fibrils in order to gain further insights into the largely different aggregation kinetics. Currently, we are exploiting the improved resolution and reduced dipolar truncation in the sparsely labeled mAS samples, to collect long-range distance restraints with the aim of an atomic model of mAS amyloid fibrils.

Materials and Methods

Sample preparation and morphology characterization

mAS samples labeled with [U-13C/15N], [1-13C]Glc, and [2-13C]Glc were produced using [U-13C]glucose, [1-13C]glucose, and [2-13C]glucose, respectively, as the sole carbon source in the bacterial growth medium. These three samples were uniformly 15N labeled. [M-13C/15N]-labeled mAS fibrils were prepared by aggregating an equimolar mixture of 13C-labeled and 15N-labeled mAS. Fibrillation of mAS was achieved according to the protocol previously developed for hAS.14 In short, mAS solutions were incubated with protein concentrations of 200 μM in 25 mM Tris–HCl buffer, pH 7.5, containing 0.02% NaN₃ at 37 °C with continuous stirring with micro stir bars at 300 rpm until the concentration of fibrillized protein reached a steady state according to an optical assay.59 The morphology of the mAS fibrils was monitored by negative-stained EM at room temperature.

ssNMR experiments and analysis

All ssNMR experiments were conducted using 4-mm and 3.2-mm triple-resonance (1H, 13C, 15N) probe heads at static magnetic fields of 20, 18.8, and 14.1 T, corresponding to 850 MHz, 800 MHz, and 600 MHz. 1H resonance frequencies (Bruker BioSpin, Germany). The chemical shifts of 13C and 15N were calibrated either with 2,2-dimethyl-2-silapentane-5-sulfonate as an internal reference or with adamantane as an external reference. Sample temperatures were determined by the position of the water 1H resonance using the relation δ(H2O)=7.83 – T/96.9. All experiments were carried out at a sample temperature of around +278 K and magic-angle spinning rates between 11 kHz and 18 kHz. An initial ramped cross-polarization was used to transfer the magnetization from 1H to 13C or 15N with contact times between 400 μs and 1000 μs, and 15N-to-13C transfer was achieved using SPECIFIC cross-polarization with contact times of 3–6 ms. SPINAL-64 high-power proton decoupling was applied during evolution and detection periods with radiofrequency amplitudes of 69–83 kHz. DREAM spectra employed a recoupling period of 4 ms. Double-quantum/single-quantum correlation spectra were recorded with the SPSCS scheme. 13C/15C correlation experiments were conducted using PDSD or DARR (dipolar assisted rotational resonance) with different mixing times. 1H/1H distances were probed indirectly using the NHHC scheme. All spectra were processed with Topspin and analyzed in SPARKY version 3.1 (T. D. Goddard and D. G. Kneller, University of California).

Accession numbers

Chemical shift assignments have been deposited to the BioMagResBank for release upon publication (BioMagResBank entry number 18232 for mAS fibrils).

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Supplementary Data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.jmb.2012.04.009

References


