β-Sheet Core of Tau Paired Helical Filaments Revealed by Solid-State NMR

Venita Daebel,† Subbashchandrabose Chinnathambi,∇# Jacek Biernat,∇# Martin Schwalbe,⊥ Birgit Habenstein,† Antoine Loquet,† Elias Akoury,⊥ Katharina Tepper,∇# Henrik Müller,⊥ Marc Baldus,⊥§ Christian Griesinger,⊥ Markus Zweckstetter,⊥∇# Eckhard Mandelkow,⊥∇# Vinesh Vijayan,*⊥§ and Adam Lange*⊥§

†NMR-based Structural Biology, Max Planck Institute for Biophysical Chemistry, Am Fassberg 11, 37077 Göttingen, Germany
∇DZNE, German Center for Neurodegenerative Diseases, Ludwig-Erhard-Allee 2, 33175 Bonn, Germany
§CAESAR Research Center, Ludwig-Erhard-Allee 2, 33175 Bonn, Germany
©DZNE, German Center for Neurodegenerative Diseases, Grisebachstraße 5, 37077 Göttingen, Germany

ABSTRACT: One of the hallmarks of Alzheimer’s disease is the self-assembly of the microtubule-associated protein tau into fibers termed “paired helical filaments” (PHFs). However, the structural basis of PHF assembly at atomic detail is largely unknown. Here, we applied solid-state nuclear magnetic resonance (ssNMR) spectroscopy to investigate in vitro assembled PHFs from a truncated three-repeat tau isoform (K19) that represents the core of PHFs. We found that the rigid core of the fibrils is formed by amino acids V306 to S324, only 18 out of 99 residues, and comprises three β-strands connected by two short kinks. The first β-strand is formed by the well-studied hexapeptide motif VQIVYK that is known to self-aggregate in a steric zipper arrangement. Results on mixed [15N:13C]-labeled K19 fibrils show that β-strands are stacked in a parallel, in-register manner. Disulfide bridges formed between C322 residues of different molecules lead to a disturbance of the β-sheet structure, and polymorphism in ssNMR spectra is observed. In particular, residues K321–S324 exhibit two sets of resonances. Experiments on K19 C322A PHFs further confirm the influence of disulfide bond formation on the core structure. Our structural data are supported by H/D exchange NMR measurements on K19 as well as a truncated four-repeat isoform of tau (K18). Site-directed mutagenesis studies show that single-point mutations within the three different β-strands result in a significant loss of PHF aggregation efficiency, highlighting the importance of the β-structure-rich regions for tau aggregation.

INTRODUCTION

The microtubule-associated protein tau¹ is expressed in the adult human brain mainly in six different isoforms. Due to alternative splicing, two N-terminal inserts (N1, N2; Figure 1) and the second out of four repeats (R2) in the C-terminal microtubule-binding domain can be present or absent.²,³ The major physiological role of tau is to bind to and stabilize microtubules facilitating axonal transport. However, upon phosphorylation, tau detaches from microtubules and self-assembles into amyloid fibrils.⁴ In Alzheimer’s disease (AD) and other tauopathies, hyperphosphorylated tau aggregates into paired helical filaments (PHFs),⁵ which represent the major constituents of neurofibrillary tangles (NFTs).⁶ A progressive accumulation of NFTs is one of the neuropathological hallmarks of AD.⁷ Hence, a structural model of tau PHFs could be helpful in the development of small compounds that prevent tau aggregation.⁸,⁹

PHFs can be structurally divided into two regions: (i) a rigid amyloid core and (ii) a highly flexible fuzzy coat of more than 200 residues that transiently attaches to the amyloid core.¹⁰ Protease digestion and solvent accessibility studies indicated that the core of PHFs is mainly built from amino acids belonging to the repeats R2 (present only in four-repeat isoforms) and R3.¹¹,¹² Furthermore, circular dichroism (CD) spectropolarimetry and Fourier transform infrared (FTIR) spectroscopy, together with X-ray diffraction and selected area electron diffraction, have indicated the presence of cross-β structure, where β-strands run roughly perpendicular to the fiber axis.¹³–¹⁵ In addition, electron paramagnetic resonance (EPR) studies have suggested that, within tau fibrils, β-strands are arranged in-register and parallel.¹⁶,¹⁷

Despite these improvements in understanding the principles of tau fibril formation, a detailed insight into the residue-specific structural arrangement of β-strands formed by the amino acids in R2 and R3 is still not available. In this regard, magic-angle spinning (MAS) solid-state nuclear magnetic resonance (ssNMR) spectroscopy constitutes a powerful method to
obtain structural information on insoluble and noncrystalline biological systems including amyloid fibrils.

Here, we employed MAS ssNMR to investigate PHFs assembled in vitro from the 99 amino acid (aa) tau construct K19 (Figure 1). It corresponds to the juvenile htau23 isoform of tau, containing a three-repeat version of the microtubule-binding domain comprising R1, R3, and R4, but lacking the N- and C-terminal domains. A previous study on K19 PHFs using ssNMR has shown that these fibrils consist of a rigid core surrounded by regions of higher flexibility. The most rigid, water-inaccessible part of the fibrils was found to be formed by repeat R3, whereas R1 and R4 are more water-accessible. However, in this study a pronounced structural heterogeneity resulting in substantial line broadening hampered the determination of the exact arrangement of β-structure within the core. Based on an improved sample preparation and an extensive labeling strategy, we present here a detailed ssNMR analysis of the K19 core structure. Additionally, we conducted H/D exchange NMR experiments to probe the accessibility of the core residues in a site-specific manner. To assess whether the identified core residues of K19 are also relevant in longer tau constructs, we also studied PHFs formed by K18, a four-repeat construct (comprising R2; Figure 1), by H/D exchange. Further insight into PHF assembly was obtained by investigating the aggregation behavior of K19 variants mutated at specific sites in the identified core region of the fibrils.

RESULTS AND DISCUSSION

Flexible Regions of Tau K19 PHFs in R1 and R4. In order to detect the flexible segments of K19 PHFs which do not belong to the rigid core, we performed INEPT-based CC-TOBSY ssNMR experiments. In the resulting spectra only resonances of residues with increased mobility (on the ps to ns time scale) are observed.

The K19 sequence features three unique amino acids, A246 in R1, Y310 in R3, and F346 in R4 (Figure S1A), with distinct chemical shifts that can be easily identified. While the INEPT-CC-TOBSY spectra exhibit resonances with unambiguous random coil chemical shifts of alanine (Figure S1B) and phenylalanine (Figure S1C, black spectrum), tyrosine resonances are clearly absent (Figure S1C, dashed circles). To unequivocally validate that the signal in Figure S1C originates from phenylalanine, we repeated the experiment on the K19KFLV sample (for a detailed description of the different samples see SI). Since phenylalanine is not [13C,15N]-labeled, the corresponding peaks vanish in the spectrum.

The appearance of F346 and A246 as well as the absence of Y310 in the INEPT-CC-TOBSY spectra shows that R1 and R4 are much more flexible than R3. Furthermore, the absence of the Y310 signals implies that the resonances in the spectra do not originate from soluble, monomeric K19 protein. In this case, the INEPT-CC-TOBSY spectra would contain signals from all residues of the molecule.

ssNMR Studies on K19 PHFs Identify a Well-Defined Rigid Fibril Core. 13C−13C correlations for residues of the rigid fibril core were obtained using PDSD experiments with mixing times of 20, 150, and 500 ms to gain intraresidue, sequential, and medium- and long-range correlations, respectively (see, e.g., Figure S2A). Other experiments were performed (e.g., DREAM; see Figure S3), but the maximum number of spin systems could be observed in the PDSD experiments. Sequential assignment was also obtained by means of NCAXC and NCOXC experiments.

As expected from the previous study, the measured resonances cover only a subset of the 99 aa of the entire K19 sequence. With V306−S324 (see Table S1), in total 18 residues are assigned to the rigid K19 PHF core. These residues belong to the repeat R3, which corroborates the results from the INEPT-based experiments.

To probe the missing ~80% of the K19 sequence, we conducted PDSD experiments (t_mix = 20 ms) at variable temperatures of ~4, 5, and 15 °C (see Figures 3A, S4, and S5). While the spectra recorded at 5 and 15 °C resemble each other in peak intensity and resolution, spectra recorded at ~4 °C exhibit significantly increased signal intensity and broader lines as well as a large number of additional peaks.

Thus, when aggregated into PHFs, ~80% of the K19 residues are highly dynamic and become sufficiently rigid to be observed in dipolar-based ssNMR experiments only when the sample is frozen.

This result is reminiscent of observations by Jaroniec and co-workers who found that signals from approximately 100 residues (~80% of all residues) of human PrP(23−144) amyloid fibrils are not detected above approximately ~20 °C in dipolar-based ssNMR experiments.

Spectra of K19 PHFs Reveal Two Conformations for Residues K321−S324. Surprisingly, in PDSD spectra of a sample, where only cysteine, tyrosine, and leucine are [13C,15N]-labeled (Figure S2B), we unambiguously detected two cysteine resonances, despite the fact that only one, C322, is present in the K19 sequence. Both resonances (in the following: C322 and 2C322) exhibit chemical shifts typical for oxidized cysteines involved in disulfide bond (DSB) formation.

Figure 1. Tau isoforms and constructs K18 and K19. (A) Tau is expressed in the adult human brain in six isoforms with htau40 as the longest and htau23 as the shortest one. While K18 is a truncated form of htau40, K19 originates from htau23. Repeat domains common to all constructs are shown in dark gray, whereas the alternatively spliced R2 is depicted in light gray. The highlighted amino acids (aa) in K19 are one of the repeats and have unique chemical shifts that make them easily identifiable in ssNMR spectra.

Figure 2. ssNMR analysis of the K19 core region of the fibrils. (A) Comparison of INEPT-CC-TOBSY spectra of htau40 and htau23, which contain only resonances that originate from soluble nanomolar quantities of K19 and K19, respectively. The highlighted amino acids (aa) in K19 are also relevant in longer repeats. (B) Amino acid sequence of the constructs K19 and K18. Amino acids in bold letters are exclusively present in the K19 core region of the fibrils.
The presence of two conformationally different cysteines in the fibril core could be confirmed by PDSD experiments on a sparsely labeled K19glyc sample (Figure 2), where we observed two sets of resonances for residues K321−S324 (set 1, K321−S324 shown in green; set 2, 2K321−2S324 shown in blue). Notably, not only sequential and medium-range connections within the sets (Figure 2A) were observed, but five long-range correlations between the two sets as well (C322−2K321, C322−2C322, C322−2S324, 2C322−2S324, S324−2G323; Figure 2B). Furthermore, both sets are sequentially connected to the preceding residues—that are observed only once—indicating a structural difference only in the C-terminal part of the PHF core.

To unequivocally ensure that the splitting of residues K321−S324 is due to the presence of an intermolecular DSB, we mutated C322 to alanine. The NCA spectrum of K19 C322A PHFs clearly reveals a simplified resonance pattern compared to that of wild-type (WT) K19 (Figure 3B) with two interesting findings: (i) Instead of two cysteine resonances, one strong alanine signal is observed, with chemical shifts indicative of β-sheet structure. Also residues K321−S324 do not give rise to two sets of signals with equal intensity any more. Nevertheless, peak doubling still occurs to a small extent, as seen, e.g., by a second weak alanine signal (2A322 in Figure 3B). (ii) In the WT K19 spectra, some peaks appear that do not show sequential correlations and could therefore not be assigned (marked with asterisks in Figures 2C and 3B). These
resonances may arise from additional peak doubling due to multiple conformations in the K19 WT. In contrast, they are absent or occur only weakly in spectra of K19 C322A PHFs. The observed spectral simplification suggests that the study of C322A K19 PHFs could be a promising approach for future ssNMR studies that aim at structure determination of tau fibrils. In future studies we will analyze in detail chemical shift changes between the two forms and perform experiments on C322A K19 PHFs that yield long-range distance information.

Secondary Structure Analysis of the K19 PHF Core Residues. In Figure 4A, secondary chemical shifts (SCSs)—indicative of the secondary structure—of residues V306–S320 (gray) as well as the two sets K321–S324 and 2K321–2S324 are depicted. Based on our chemical shift assignment, and in line with the previous ssNMR study, the fibril core is predominantly found in β-strand conformation. This is further corroborated by backbone dihedral angle predictions obtained from the program TALOS+ (Figure 4C; predictions for all residues were classified as “good”, except for 2G323). Exceptions occur for residues P312, S316, and G323, as well as 2C322 and 2G323. The SCSs of these residues indicate the presence of kinks that disrupt the β-strands (Figure 4A) and
might change the characteristic inside–outside order of side chains observed in a fully extended conformation.

Previous studies have shown that the hexapeptide \(306VQIVAL311\) in R3 displays \(\beta\)-sheet propensity in disordered full-length tau and is furthermore able to self-assemble into fibrils by itself.\(^{9,47–49}\) Consistently, SCs identify the first \(\beta\)-strand from V306–K311. At P312 a kink is probably formed, because proline is a known \(\beta\)-strand breaker and the \(\alpha\)t chemical shift of P312 is larger than the random coil value, indicating a non-\(\beta\)-strand conformation. P312 is followed by a second, short \(\beta\)-strand, V313–L315, and another kink located at residue S316. In molecules belonging to set 1, the third strand extends from K317 to C322, followed by a further kink at G323. In set 2 molecules, the third \(\beta\)-strand comprises only residues K317–2K321, and a larger kink is formed by 2C322 and 2G323.

Previously it was shown that a spin label attached to C322 in PHF tau can be quenched by a reducing agent, indicating that the side chain of C322 is not fully buried in the fibrillar core\(^{10}\) in line with formation of a kink at C322.

Downstream of this kink, S324 as well as 2S324 chemical shifts indicate \(\beta\)-strand structure. However, as there is no unambiguous assignment for the consecutive residues, it is not clear whether a fourth \(\beta\)-strand follows.

While deletion of the C-terminus of full-length tau tends to enhance fibril formation, the deletion of residues D314–S320 abrogates fibrillation,\(^{50}\) probably as it eliminates strands \(\beta2\) and \(\beta3\). Interestingly, \(314^{DLSK317}\) shows conformational propensity for a turn in natively unfolded tau in solution as well as in chemical denaturant, as observed by SCs and residual dipolar coupling analysis.\(^{48,51}\) This indicates that the turn conformation might be a precursor to fibril conformation similar to the hexapeptide motif.

H/D Exchange Experiments Reveal a Protective Environment in the Identified Core Region. To probe the solvent accessibility of the residues forming the core of PHFs, we conducted H/D exchange experiments, using solution-state NMR.\(^{52}\) As the fibrillar state is not accessible by solution-state NMR, solvent-protected amide protons were identified as described previously (for a detailed description see SI). In short, after an initial forward exchange of PHFs to 99.9% \(D_2O\), the PHFs are dissolved in a strongly denaturing buffer with 50% \(D_2O/50%\) H\(_2O\). Thus, the H/D exchange rates can be determined by solution-state NMR, and with these rates the residual protonation after forward exchange is back-calculated. Solvent-protected sites will display a higher residual protonation relative to solvent-exposed sites.

In agreement with the ssNMR results, the most protected region in K19 PHFs belongs to residues I308–V313. With a certain degree of protection visible for residues D314–K321 as well (Figure 4D). In contrast, the two other repeats, R1 and R4, are more solvent-exposed, with the majority of residues displaying protonation levels below or equal to the average value (Figure S6). A short, consecutive stretch in R1 from D252–K257 also displays above average protonation levels similar to D314–K321, suggesting some degree of protection for this region. However, this is not apparent for the same region in K18 (see below). Residues D252–K257 showed a lower intensity relative to other residues, in particular in R3, and, as the error bars indicate, the above average protonation may be the result of a poor fit.

Furthermore, we probed using PHFs formed by K18 how repeat R2 contributes to the fibril core (Figures S6–S8). The H/D exchange behavior is similar for K19 and K18 (Figure S7), and the most protected region in both K19 and K18 is repeat R3 (Figures S6 and S8). The protective environment provided by the fibril core in R3 might explain why residue Y310 is the only tyrosine in tau that is nearly resistant to chemical modification by site-specific nitration\(^{53}\) and that gives the most extensive shift changes in fluorescence spectra even in full-length tau during PHF aggregation.\(^{11}\) K18 residues of repeat R2 are also protected from exchange with solvent (Figures S6 and S8). In particular, the hexapeptide \(275^{VQINK380}\) at the N-terminus of R2 is highly protected, indicating its involvement in the rigid fibril core of K18 PHFs.

Aggregation Behavior of K19 Single Mutants Supports the NMR-Derived Structural Data. The \(\beta\)-strand \(\beta1\) of the PHF core corresponds to the hexapeptide \(306^{VQIVYK311}\) that has been studied extensively.\(^{14,47,49,54}\) It was shown that \(\beta\)-strand-breaking proline point mutations within this motif, e.g., I308P, abrogate PHF formation.\(^{14}\) On the other hand, the influence of strands \(\beta2\) and \(\beta3\) (corresponding to V313–C322/ V313–2K321) on fibril formation and stability is largely unknown. Hence, in the current study we introduced single-point mutations in this part of the K19 core and followed the aggregation behavior of the mutant proteins. The kinetics of aggregation was monitored fluorimetrically using the thioflavin S (ThS) assay (Figure 5A–C). The presence and morphology of PHFs were monitored by electron microscopy (EM, Figure SD).

First, K19 proline mutants were designed, which were expected to be inhibitory for aggregation by disrupting the \(\beta\)-strands /\(\beta1, \beta2, \text{and } \beta3\), respectively (Figure 5A,D). As expected, the mutations V313P and T319P disrupt the \(\beta\)-strands and prevent fibril formation. The most pronounced inhibition was observed for V313P, which leads to nearly complete abrogation of aggregation, comparable to the previously described I308P mutation, furthermore, we generated the mutations L315E within \(\beta2\) and S320E as part of \(\beta3\), but also L325E, which is just outside of the assigned region (Figure 5B). These mutations disrupt PHF formation to different degrees. The farthest C-terminal mutation, L325E, shows the least drastic effect.

With S320V, a hydrophilic serine is replaced by a hydrophobic valine. Figure 5C illustrates the aggregation kinetics of S320V, which is even faster than of the WT.

Figure SD summarizes the structures observed by negative stain EM. The control sample of K19 WT PHFs shows abundant long filaments, predominantly with twisted substructure. The morphology of K19 S320V fibrils is indistinguishable from PHFs of K19 WT. In the remaining micrographs we can distinguish between two different morphologies: (i) fragmented short filaments formed by the mutants S320E and L325E and (ii) amorphous deposits without filaments for the mutants I308P, V313P, L315E, and T319P.

In summary, the aggregation behavior of the proline mutants provides evidence that the \(\beta\)-structure is highly important for fibril formation in the overall region of residues 306–325.

A hydrophobic influence on the stability of the PHF core was probed by mutating hydrophobic to charged residues. These mutations abrogate WT-like fibril formation. In contrast, replacing S320 by a more hydrophobic valine actually leads to an enhancement of fibrillation, indicating hydrophobic interactions to be a substantial stabilizing factor for K19 PHFs.

Intermolecular Stacking of Tau K19 PHFs Is Parallel and In-Register. To elucidate the intermolecular arrangement of tau K19 molecules to form the cross-\(\beta\) structure, a two-
dimensional $^{15}$N–$^{13}$C PAIN-CP spectrum of a mixed labeled K19$_{1:1}$ sample was measured (Figure 6, blue). These PHFs were aggregated from an equimolar mixture of $^{15}$N-labeled and $^{13}$C-labeled K19 protein. Since no $^{15}$N–$^{13}$C-spin pairs are present within one K19 molecule of such mixed-labeled fibrils, only intermolecular $^{15}$N(i)–$^{13}$C$\alpha$(i) correlations are obtained in the PAIN-CP spectrum$^{56}$ (Figure 6B). The spectrum overlays well with an NCA spectrum of the K19uni sample (Figure 6A, black) revealing a parallel, in-register arrangement of the $\beta$-strands along the fibril axis, in line with previous EPR studies.$^{16,17}$

Notably, no intermolecular $^{15}$N(i)–$^{13}$C$\alpha$(i) correlations are detected for residues K321–S324 (and 2K321–2S324; orange circles in Figure 6A), indicating either that this part is not stacked parallel and in-register or that set 1 and set 2 molecules are stacked in an alternating arrangement (see also discussion in the next paragraph).

**Intermolecular Disulfide Bonds Stabilize K19 PHFs.** All investigated samples were prepared under conditions in which DSB formation can occur. The chemical shifts of the two assigned cysteines (C322 and 2C322) show that this happens indeed. Importantly, five long-range restraints correlating the two sets were detected in the PDSD 500 ms spectrum of the K19$_{1:1}$glyc sample.

Because cysteine is a unique amino acid within the K19 sequence, the DSB must be formed intermolecularly (Figure 7). In principle, two possibilities are conceivable: (i) an intermolecular DSB within one protofibril along the fibril axis (Figure 7A), which is possible because of the parallel, in-register arrangement$^{57,58}$ or (ii) an inter-protofibril DSB or rather a DSB formed by symmetrically nonequivalent monomers in a lateral dimer within one protofibril (Figure 7B)$^{59,60}$

Aggregation assays on K19 molecules under reducing conditions using DTT investigated by ThS and EM reveal that K19 is still able to form PHFs, but at a much slower rate.$^{61}$ This strongly supports the notion that K19 intermolecular DSBs are an important factor for a fast and stable aggregation behavior, not least because DSB-linked tau dimers are known to successfully seed PHF aggregation.$^{35}$

**CONCLUSION**

As tau aggregation into PHFs is one of the hallmarks of AD, it is important to understand structural features of the fibril core as well as the fibrillation process itself. Our site-specific ssNMR study on K19 PHFs reveals a well-defined rigid core (V306–S324; from R3 only) composed of three $\beta$-strands. Our data unambiguously disclose a parallel, in-register supramolecular arrangement, including the presence of intermolecular disulfide bonds between the tau monomer units.

Furthermore, our results support that the PHF core region is essential not only in K19 but in longer constructs, such as K18, as well. Ongoing ssNMR studies on the K19 C322A mutant...
will be helpful to obtain unambiguous long-range distance restraints required for a high-resolution model of tau PHFs.

**MATERIALS AND METHODS**

WT and mutant K19 proteins were prepared and aggregation of tau was performed as described previously. ssNMR experiments were conducted on aggregated tau K19 samples (protein quantities of 10–15 mg) using 2 or 3.2 mm triple-resonance (1H, 13C, 15N) MAS probes at static magnetic fields of 20.18.8, and 14.1 T (Bruker Biospin, Karlsruhe, Germany). Further details are given in the SI.

**ASSOCIATED CONTENT**

Supporting Information

Detailed materials and methods; supplemental figures regarding assignment strategy and H/D exchange data; chemical shift assignment table. This material is available free of charge via the Internet at http://pubs.acs.org.

**AUTHOR INFORMATION**

Corresponding Author

adla@nmr.mpibpc.mpg.de; vinesh@iisertvm.ac.in; mandelkow@dzne.de

Present Addresses

1Forschungszentrum Jülich GmbH, 52425 Jülich, Germany
2Utrecht University, Padualaan 8, 3584 CH Utrecht, Netherlands
3IISER Thiruvananthapuram, India

Notes

The authors declare no competing financial interest.

**ACKNOWLEDGMENTS**

We are grateful to Dr. E.-M. Mandelkow for discussions and I. Lindner and B. Angerstein for excellent technical assistance. We thank the Max Planck Society, the DFG (DFG ZW 71/7-1 and Heisenberg scholarship DFG ZW 71/2-2 to M. Zweckstetter, Emmy Noether fellowship to A. Lange), and the European Molecular Biology Organization (long-term fellowship to A. Loquet) for financial support. V. Daebel was supported within the scope of the Graduiertenkolleg GRK 782 of the DFG.

**REFERENCES**


(43) Sharma, D.; Rajarathnam, K. J. Biomol. NMR 2000, 18, 165.