Molecular structure of monomorphous peptide fibrils within a kinetically trapped hydrogel network

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Most, if not all, peptide- and protein-based hydrogels formed by self-assembly can be characterized as kinetically trapped 3D networks of fibrils. The propensity of disease-associated amyloid-forming peptides and proteins to assemble into polymorphic fibrils suggests that cross-β fibrils comprising hydrogels may also be polymorphous. We use solid-state NMR to determine the molecular and supramolecular structure of MAX1, a de novo designed gel-forming peptide, in its fibrillar state. We find that MAX1 adopts a β-hairpin conformation and self-assembles with high fidelity into a double-layered cross-β structure. Hairpins assemble with an in-register Syn orientation within each β-sheet layer and with an Anti orientation between layers. Surprisingly, although the MAX1 fibril network is kinetically trapped, solid-state NMR data show that fibrils within this network are monomorphous and most likely represent the thermodynamic ground state. Intermolecular interactions are not available in alternative structural arrangements apparently dictate this monomorphic behavior.

MAX1 fibrils are known to be polymorphous at the molecular structural level (7, 8, 12, 13), implying that structures within peptide and protein fibrils are generally not determined uniquely by amino acid sequences and do not necessarily represent thermodynamic ground states. A model for fibrils formed by the designed peptide RADA16-I has been proposed by Cormier et al. based on solid-state NMR data (14), in which RADA16-I monomers form single β-strands within a double-layered cross-β structure. Solid-state NMR spectra of this hydrogel-forming peptide also indicate coexistence of several distinct fibril structures, suggesting that polymorphism may also be a trait of designed sequences.

Here, we use solid-state NMR to develop a full structural model for MAX1 fibrils, including molecular conformation, β-sheet organization, and intersheet interactions, with experimental restraints on all levels of structure. We find that MAX1 self-assembles with high fidelity to form monomorphous fibrils with well-defined and uniform structures. This structural homogeneity suggests that although the evolution of the fibril network is governed by kinetics, the molecular structure within MAX1 fibrils most likely represents the thermodynamic ground state. Additionally, results described below represent, to our knowledge, the first complete experimentally based model for a cross-β fibril structure comprised of β-hairpins.

Results

MAX1 Fibrils Are Monomorphous. TEM images of nascent MAX1 assemblies (Fig. 2A, Left) reveal fibrils of uniform appearance, with diameters of ~3.5 nm. Along with previous atomic force microscopy (AFM) data, which indicate a fibril height of 2.5 nm (15), these dimensions are consistent with models in which β-hairpins assemble into a double-layered cross-β structure, i.e., a β-sheet bilayer with specific molecular conformations and intermolecular alignments within and between layers.


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MAX1 is a 20-residue peptide designed de novo to fold into an amphiphilic β-hairpin that self-assembles to form a fibrillar network within a self-supporting hydrogel (1). The MAX1 gel exhibits shear thin-recovery rheological behavior (2), is cytocompatible toward mammalian cells, yet is inherently antimicrobial (3) and thus has applications in tissue engineering and drug delivery. In addition to exploring the utility of the gel, we seek to understand the mechanism of gelation, the macroscopic morphology of its fibrillar network, and the underlying molecular structure of its fibrils.

MAX1 contains two segments of alternating lysine and valine residues, connected by a four-residue turn-forming segment. When initially dissolved in water, electrostatic repulsions among protonated lysine sidechains lead to an ensemble of monomeric random coil conformations (1). Peptide folding and self-assembly, leading to gelation (Fig. 1), can be triggered by attenuating electrostatic repulsions, by adjusting the solution pH and/or ionic strength. Increasing the solution temperature also drives hydrophobic collapse of valine sidechains, further favoring MAX1 assembly. According to circular dichroism (1), cryo-transmission electron microscopy (TEM) (4), small-angle neutron scattering (5), and dynamic light scattering coupled with rheological measurements (4), soon after the triggering event, peptides assemble into branched clusters of β-sheet-rich, semiflexible nanofibrils throughout the solution. Individual clusters contain dangling fibril ends that grow and interpenetrate neighboring clusters as the network evolves. Multiple particle tracking microrheology shows that the time at which the fibril network percolates the entire sample volume, defining the gel point, is less than 1 min at 1% (wt/vol) peptide (6). In this mechanism of gelation, the growing fibrils become kinetically trapped in the evolving network as they percolate the sample volume. Fibrils do not precipitate, but rather form a 3D random network that defines the gel state.

Full structural models for naturally occurring amyloid (7–10) and prion (11) fibrils have been developed from solid-state NMR data, but less is known about fibril structures within designed peptide hydrogels. Disease-associated amyloid and prion fibrils

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pair of ribbon-like β-sheets in which β-strands run approximately perpendicular to and hydrogen bonds run approximately parallel to the fibril growth axis. Solid-state NMR experiments were performed on lyophilized and lyophilized/rehydrated MAX1 hydrogels. After rehydration of a lyophilized hydrogel, TEM images show quite similar fibril morphology and diameters (Fig. 2, Right), indicating that MAX1 fibril structures are not perturbed during preparation of solid-state NMR samples.

Fig. 2B shows 2D solid-state $^{13}$C–$^{13}$C NMR spectra of MAX1 fibrils that were uniformly $^{15}$N, $^{13}$C-labeled at residues 8–13 and $^{13}$C-labeled at the carbonyl site of $^{13}$C-P10 (sample IX in Table 1). A single set of cross-peak signals is observed, corresponding to a single set of $^{13}$C chemical shifts for all uniformly labeled residues. Two-dimensional spectra of lyophilized and rehydrated gel samples are identical except that NMR linewidths are somewhat reduced in the rehydrated state (0.7–0.9 ppm versus 0.8–1.2 ppm full width at half maximum), presumably due to small-amplitude local motions that require hydration. Together with the TEM images, these data show that lyophilization does not perturb the molecular structure within MAX1 fibrils. Importantly, the 2D spectra indicate that no significant nonfibrillar material is present and that all MAX1 molecules within the fibrils have nearly identical conformations and structural environments. This rules out the possibility that MAX1 fibrils contain mixtures of the supramolecular structures shown in Fig. 1, strongly arguing against the possibility that MAX1 fibrils are inherently polymorphic, in contrast with disease-associated amyloids (7, 8, 12, 13).

MAX1 Adopts a β-Hairpin Conformation. Experimental restraints at three distinct levels of structure, depicted in Fig. 3, were obtained from a series of solid-state NMR measurements on samples with the labeling patterns in Table 1. The first step was to obtain restraints on the conformation of individual MAX1 molecules. Conformational information is contained in $^{13}$C and $^{15}$N chemical shifts (SI Appendix, Table S1), which were measured in 1D and 2D solid-state NMR spectra (Fig. 2 and SI Appendix, Figs. S1, S5, and S6). Backbone torsion angles were predicted from chemical shifts for certain sites in residues 9–13, using the TALOS+ program (16). $^{13}$C chemical shifts for backbone carbonyl sites of all valine residues in samples II–IX were found to be in the 173.3–173.9-ppm range, significantly upfield from the random-coil value of 176.3 ppm (17) and hence consistent with β-strand conformations at values in residues 3–9 and 16.

Additional conformational restraints were obtained from quantitative measurements of nuclear magnetic dipole–dipole couplings with the rotational echo double resonance (REDOR) and the $^{15}$N and $^{13}$C backbone recoupling ($^{15}$N- and $^{13}$C-BARE) techniques (18, 19). Examples of the data are shown in Fig. 3A. The complete data sets are shown in SI Appendix, Figs. S2 A–C. Each of these techniques includes a dipolar recoupling period, during which the relevant couplings are activated by an appropriate...
Radiofrequency (rf) pulse sequence, before the detection of solid-state NMR signals. Dipole–dipole coupling strengths are inversely proportional to the cubes of the distances between coupled nuclear spins. Shorter distances (i.e., stronger couplings) produce more rapid decays of the NMR signals with increasing recoupling period.

$^{15}\text{N}$–$^{13}\text{C}$ REDOR measurements on samples I–IV (Fig. 3A and SI Appendix, Fig. S2A) indicate distances of 4.3–4.5 Å between carbonyl $^{13}\text{C}$ labels at residues 1, 3, 5, and 7 and amide $^{15}\text{N}$ labels at residues 20, 18, 16, and 14, respectively. These distances are consistent with hydrogen bonding between $^{13}\text{C}$-labeled and $^{15}\text{N}$-labeled residues, as expected in a β-hairpin conformation. The intramolecular nature of these hydrogen bonds is supported by REDOR measurements on a MAX1 fibril sample in which molecules with the labeling pattern of sample II were diluted in unlabeled molecules (SI Appendix, Fig. S3). In particular, the REDOR data show that the β-hairpin structure extends to the N- and C-termini of MAX1, although the broadened carbonyl $^{13}\text{C}$ line for V1 (SI Appendix, Fig. S1B) and the REDOR data for sample I (SI Appendix, Fig. S2A) suggest somewhat greater disorder and a broader distribution of $^{15}\text{N}$–$^{13}\text{C}$ distances at the termini. Importantly, the intramolecular hydrogen bonds must involve pairs of valine residues, rather than pairs of lysine residues, a finding that places strong constraints on the turn conformation.

Evidence from solid-state NMR for β-hairpin conformations in fibrils formed by the related peptide MAX8 has been reported previously by Leonard et al. (20). Their data only define distances from V3 to K17 and V18. Further restraints on the conformation in residues 8–12 of MAX1 were obtained from $^{15}\text{N}$- and $^{13}\text{C}$-BARE measurements on sample IX. As previously described (19), $^{15}\text{N}$-BARE measurements probe $^{15}\text{N}$–$^{15}\text{N}$ dipole–dipole couplings of the backbone amide nitrogen of residue $k$ to amide nitrogens of residues $k-1$ and $k+1$, which depend on backbone torsion angles $\psi_k-1$, $\phi_k$, and $\psi_k$. (When only residue $k-1$ or $k+1$ is $^{15}\text{N}$-labeled, the data depend only on $\psi_k-1$ or $\psi_k$, or equivalently on one $^{15}\text{N}$–$^{15}\text{N}$ distance.) $^{13}\text{C}$-BARE measurements probe $^{13}\text{C}$–$^{13}\text{C}$ dipole–dipole couplings that provide experimental restraints on internuclear distances in MAX1 fibrils. (A) $^{13}\text{C}$–$^{13}\text{C}$ REDOR data from sample II and $^{15}\text{N}$-BARE and $^{13}\text{C}$-BARE data from sample IX support a β-hairpin conformation. (B) $^{13}\text{C}$ PITHIRDS-CT data from samples II and VII indicate a $\psi$-sheet alignment of MAX1 β-hairpins within each β-sheet. The relevant residues and labels (C for carbonyl $^{13}\text{C}$, N for amide $^{15}\text{N}$) are indicated in each plot. The detected sites in BARE data are underlined. Solid lines in all plots are averaged simulations using atomic coordinates from the final ensemble of 20 MAX1 fibril structures, with dashed lines indicating 1 SD. Error bars on experimental data points represent uncertainties due to the root-mean-squared noise in the solid-state NMR spectra. (C) Two-dimensional solid-state $^{13}\text{C}$–$^{13}\text{C}$ NMR spectrum of MAX1 sample X as a rehydrated gel, with a 500-ms RAD mixing period, obtained at 100.8-MHz $^{13}\text{C}$ NMR frequency with 9.00-kHz MAS. Arrows indicate long-range interresidue cross-peaks that serve as restraints on supramolecular structure in MAX1 fibrils.
couplings of the backbone carbonyl carbon of residue k to carboxyl carbons of residues k — 1 and k + 1, which depend on backbone torsion angles $\phi_k$, $\psi_k$, and $\phi_{k+1}$. Experimental $^{15}$N-BARE and $^{13}$C-BARE curves for sample IX (Fig. S4 and SI Appendix, Fig. S2 B and C) show unique signal decays for each labeled residue, consistent with a $\beta$-turn conformation in residues 8–12. Thus, MAXI adopts a well-defined $\beta$-hairpin conformation in the fibrillar solid state, without competing alternative folds that could lead to polymorphism.

**Evidence for Syn Alignment Within $\beta$-Sheets.** As illustrated in Fig. 1, several supramolecular structures with different $\beta$-hairpin orientations are consistent with information derived from scattering and microscopy experiments. In both Syn/Syn and Syn/Anti structures, $\beta$-hairpins within a given $\beta$-sheet have their turns on the same edge of the $\beta$-sheet. In the Syn/Syn structure, the two $\beta$-sheets have their turns on opposite sides. An ideal Syn/Syn structure has twofold axes of rotational symmetry perpendicular to the fibril growth axis and parallel to the $\beta$-sheet plane, whereas an ideal Syn/anti structure has approximate twofold symmetry about the fibril growth axis (see below). In both Anti/Syn and Anti/Anti structures, $\beta$-hairpins within a given $\beta$-sheet have their turns on alternating edges of the $\beta$-sheet. In the Anti/Syn and Anti/Anti structures, $\beta$-hairpins from the two $\beta$-sheets have their turns on the same side of the fibril, whereas in the Anti/Anti structure, facially associated $\beta$-hairpins have their turns on opposite sides. An ideal Anti/Anti structure has twofold symmetry axes perpendicular to the fibril growth axis and parallel to the $\beta$-sheet plane, whereas an ideal Anti/Anti structure has approximate twofold symmetry about the fibril growth axis.

Experimental restraints on $\beta$-sheet organization in MAXI fibrils were obtained from measurements of intramolecular $^{13}$C–$^{13}$C dipole–dipole couplings among carbonyl $^{13}$C labels in samples I-V. Figure 3A shows a representative alignment of $\beta$-hairpins deduced from 2D-1D dipole–dipole coupling (PITTHIRDS-CT) technique (21). Examples of the data are shown in Fig. 3B. The complete data sets are shown in SI Appendix, Fig. S2 D and E. In PITTHIRDS-CT measurements on samples I–IV, signals decayed by 25–40% as the recoupling period increased from 0 to 76.8 ms. These signal decays indicate $^{13}$C–$^{13}$C distances greater than 8 Å (21) and are similar to decays observed for natural-abundance $^{13}$C signals in unlabeled samples (22). The absence of shorter distances rules out Anti alignments of $\beta$-hairpins with all possible registries of intermolecular hydrogen bonds, because any Anti alignment would imply intermolecular $^{13}$C–$^{13}$C distances of ~6.1 Å at least one of several samples I–IV (SI Appendix, Fig. S4).

To test for a Syn alignment within individual $\beta$-sheets, samples V–VIII were prepared from 1:1 mixtures of molecules with a carbonyl $^{13}$C label at V16 and molecules with carbonyl $^{13}$C labels at V1, V3, V5, or V7. Use of 1:1 mixtures eliminated effects of intramolecular couplings in each sample. PITTHIRDS-CT measurements revealed more rapid signal decay for sample VII than for samples V, VI, or VIII (SI Appendix, Fig. S2E). For sample VII, the decay by 42% at a recoupling period of 57.6 ms indicates an intermolecular distance of ~6.4 Å, after taking into account the random distribution of V5-labeled and V16-labeled molecules within sample VII. To produce an intermolecular carbonyl–carbonyl distance that is ~6.4 Å in sample VII, with longer distances observed in samples V, VI, and VIII, intermolecular hydrogen bonds must connect lysine residues 2 and 19, 4 and 17, 6 and 15, and 8 and 13 of neighboring peptides. This result supports an in-register Syn alignment of $\beta$-hairpins within each $\beta$-sheet (SI Appendix, Fig. S4D).

**Evidence for Anti Alignment Between Two $\beta$-Sheet Layers.** Prior work suggests that individual MAXI fibrils contain two $\beta$-sheet layers that interact through hydrophobic amino acid sidechains (15, 23–25). The distances between backbone atoms of the two layers must exceed 8 Å to accommodate amino acid sidechains in the hydrophobic interior of the fibril. Such long distances cannot be measured reliably in selectively $^{15}$N- or $^{13}$C-labeled samples with REDOR, PITTHIRDS-CT, or similar dipolar recoupling techniques because the corresponding dipole–dipole coupling strengths are less than 15 Hz. To distinguish between Syn and Syn/Anti structures, we performed 2D $^{13}$C–$^{13}$C rf-assisted diffusion (RAD) measurements (26, 27). When acquired with long mixing periods, 2D RAD spectra of samples that contain uniformly labeled residues typically show interresidue cross-peaks between $^{13}$C NMR signals of residues whose sidechains are within 6–8 Å of one another (7, 8, 28). Hence, sample X was designed with uniformly labeled residues at opposite ends of the MAXI $\beta$-hairpin, with V1 and V20 being more than 20 Å away from P11 and T12. Given the Syn alignment within a single $\beta$-sheet, any 2D RAD cross-peaks between V1/V20 and P11/T12 must arise from interlayer proximities of these labeled residues, not from intralayer proximities.

Experimental 2D RAD spectra of sample X are shown in Fig. 3C and SI Appendix, Fig. S5. With a 500-ms mixing period, cross-peaks are clearly observed between the $^{13}$Cα chemical shift of V20 and the T12 $^{13}$Cα chemical shift as well as the P11 $^{13}$Cα and $^{13}$Cβ chemical shifts. These cross-peaks are absent from the 2D RAD spectrum with a 25-ms mixing period, as expected for interlayer $^{13}$C–$^{13}$C distances (SI Appendix, Fig. S5A). Assignment of the 60.3-ppm chemical shift to $^{13}$Cα of V20 (rather than V1) is confirmed by data for a sample in which V1 was not labeled (SI Appendix, Fig. S6).

With a 500-ms mixing period, strong 2D RAD cross-peaks also occur between chemical shifts of P11 and T12, as expected for sequentially labeled residues. For example, all P11 chemical shifts have cross-peaks with $^{13}$Cβ of T12. Cross-peaks between Cβ chemical shifts of V1 and V20 are also observed (SI Appendix, Fig. S5B), consistent with the expected intramolecular and intermolecular $^{13}$C–$^{13}$C distances of ~5 Å between V1 and V20 sidechains within each $\beta$-sheet layer. Taken together, the data support an Anti alignment of $\beta$-hairpins between the two $\beta$-sheet layers.

**Structural Model for MAXI Fibrils.** Structural restraints from solid-state NMR, listed in SI Appendix, Table S2, were included as potential energy functions in two stages of structure calculations, performed with Xplor-NIH (stage 1) and NAMD (stage 2). Structure calculations included eight copies of MAXI to form a bilayer of $\beta$-sheets composed of four monomers each. Restraints were included in Xplor-NIH calculations by standard methods (29) and in NAMD calculations by a Monte Carlo algorithm (Materials and Methods and SI Appendix). NAMD calculations also included explicit hydration and full electrostatics (30). Conformational and translational symmetry restraints were imposed during Xplor-NIH calculations, but not during NAMD calculations. Fig. 4 shows several representations of the final structural model of a MAXI octamer (specifically, the final frame from NAMD calculations). Overall dimensions are consistent with the apparent fibril diameters in Fig. 24, providing further support for a structure that consists of two $\beta$-sheet layers. A bundle of structures (SI Appendix, Fig. S7C), representing 20 frames from the latter half of the NAMD calculations, has been deposited in the Protein Data Bank, PDB ID code 2N1E. Values of backbone torsion angles in residues 1–9 and 12–19 are typical of $\beta$-strands (average $\phi$ angles between −144° and −120° from K2 through K9 and from K13 through K19; average $\psi$ angles between 119° and 140° from V1 through V9 and from V12 through V18). Backbone torsion angles are $\phi = 29° \pm 4°$, $\psi = −118° \pm 9°$ for P10 and $\phi = −87° \pm 16°$, $\psi = 12° \pm 35°$ for P11. These values are similar to the torsion angles of an ideal type II $\beta$-turn (31).

As shown in Fig. 4 A and B, $\beta$-strands in one $\beta$-sheet layer are displaced along the fibril growth axis relative to $\beta$-strands in the other layer, i.e., $\beta$-hairpins across the bilayer do not directly oppose one another. This displacement allows rows of valine sidechains from the two layers to interdigitate. Interdigitation in the direction perpendicular to the fibril growth axis is also present, as shown in Fig. 4E, producing a tightly packed, dry, hydrophobic interface between $\beta$-sheet layers. No water molecules
Discussion

Significance of Structural Homogeneity. Kinetically trapped fibril networks play important biological roles, for example in cytoskeletal networks of actin (32, 33) and functional amyloid such as curli-derived protein matrices important in bacterial colonization (34). The formation of kinetically trapped networks is often accompanied by local structural heterogeneity at the nanoscale. Amyloid fibrils formed in vitro are generally polymorphic unless special fibril growth protocols are used, such as repeated seeding (8) or long periods of incubation with agitation to promote convergence to a single structure with lowest free energy (35). Prior solid-state NMR studies of the gel-forming peptides RADA16-I and MAX8 suggested polymorphism or incomplete molecular structural order within the hydrogels (14, 20).

Surprisingly, we find that the molecular structure within kinetically trapped MAX1 fibril networks is homogeneous and most likely represents the thermodynamic ground state for local assembly. The Syn/Anti arrangement of MAX1 hairpins allows critical intermolecular interactions that may dictate the monomorphic character of MAX1 fibrils. These interactions are not accessed in other competing structures (Fig. 1). For example, the Syn alignment of hairpins within a given β-sheet permits van der Waals and hydrophobic contacts between the turn regions of neighboring hairpins, resulting in an approximate 20% decrease in solvent-exposed surface area compared with an Anti alignment within β-sheets. The in-register nature of the Syn alignment within β-sheets maximizes intermolecular hydrogen bonding. The Anti alignment of hairpins across β-sheet layers effectively packs the N- and C-terminal valine sidechains of one monomer into the puckered interface formed by the diproline turn residues of its bilayer partner. In the competing Syn alignment, opposing β-turns would project away from the hydrophobic core and become solvent-exposed. Added over the length of the fibril, the additional favorable interactions provided by the Syn/Anti arrangement represent a significant thermodynamic driving force for the formation of monomorphic fibrils.

The β-hairpin design of MAX1, which restricts the conformational freedom of the monomer, may also contribute to the monomorphic nature of MAX1 fibrils. However, even with the β-hairpin conformation, multiple supramolecular structures are conceivable, as discussed above and shown in Fig. 1.

Comparison with Amyloid Fibril Structures. Like MAX1 fibrils, disease-associated amyloid and prion fibrils contain cross-β structures. Early models for amyloid-β (Aβ) fibrils suggested that β-hairpins might be building blocks for amyloid structures (36, 37). However, subsequent solid-state NMR studies (7–10, 35) revealed in-register, parallel cross-β structures comprising peptide conformations in which N-terminal and C-terminal β-strand segments form separate β-sheets that contain only intermolecular hydrogen bonds, with sidechain–sidechain contacts between these β-sheets.

Recent work on Aβ aggregation intermediates by Härd and coworkers suggests that β-hairpins may be present in certain
intermediates (38, 39). Metastable protofibrils formed by the “Towa mutant” of Aβ have been found to contain antiparallel β-sheets, but still without β-hairpins (28). Laganowsky et al. have reported crystal structures of peptide oligomers derived from Aβ crystallin, called cylindrins (40). Cylindrins comprising tandem repeats of the Aβ crystallin peptide contain β-hairpins, but are not cross-β structures. The same tandem repeat peptides also form fibrils, but the peptide conformation within these fibrils has not been determined. Although cross-β structures comprising β-hairpins have not yet been demonstrated conclusively in disease-associated and naturally occurring fibrils, our results for MAX1 fibrils show that such structures certainly exist for appropriate amino acid sequences.

Concluding Remarks. We have shown that the MAX1 hairpin is capable of assembling with high structural fidelity, affording monomorphic fibrils within a kinetically trapped hybrid network. Knowledge of the exact arrangement of peptide molecules in the fibrillar state will facilitate future structure-based design of highly functional materials.

Materials and Methods
A series of MAX1 samples with isotopic labeling patterns in Table 1 was synthesized and purified by standard methods. Purified MAX1 was dissolved in water to afford a 4 wt % peptide solution. Hydrogel formation was initiated by addition of an equal volume of 250 mM bis-Tris propane buffer, pH 9, with 20 mM NaCl, followed by incubation at room temperature for 2 h. For solid-state NMR, lyophilized hydrogels were washed twice with deionized water and lyophilized again. Solid-state NMR samples contained 4 mmol of MAX1. Solid-state NMR data were acquired with Varian InfinityPlus, Bruker Avance III, and Varian Infinity spectrometers, operating at 1H NMR frequencies of 399.2, 400.9, and 746.6 MHz, respectively, using a Varian 3.2-mm magic-angle spinning (MAS) probe at the lower frequencies and a 1.8-mm MAS probe from the laboratory of Agos Samoson at the highest frequency. Sample temperatures during MAS were maintained near 24 °C with cooled nitrogen gas.

The structural model for MAX1 fibrils was generated in two stages. In the first stage, starting from a set of eight well-separated copies of MAX1 with randomized conformations, Xplor-NIH was used to generate an initial structural model, using restraints summarized in SI Appendix, Table S2. In addition, symmetry restraints were used to ensure identical conformations for all MAX1 molecules, translational symmetry within each β-sheet, and overall twofold rotational symmetry. Artificial distance and torsion angle restraints were also included to enforce hydrophobic contacts between β-sheet strands. In the second stage, the lowest-energy structure from 40 independent Xplor-NIH runs was solvated in a 80.7-nm3 box containing 1,654 water molecules, 77 Cl− ions, and 5 Na+ ions (for roughly 150 mM NaCl and charge neutrality). Molecular dynamics simulations were then run at 37 °C with the NAMD, using a Monte Carlo algorithm to enforce experimentally based structural restraints during these simulations. Fig. 4 shows the final structure from an 8.0-ns molecular dynamics/Monte Carlo simulation. SI Appendix, Fig. S7 compares structures from the ends of stages 1 and 2, as well as a superposition of structures from 20 equally spaced time points in the final 4.0 ns of the NAMD simulations (PDB ID code 2N1E). PyMOL and Accelrys Discovery Studio were used for molecular graphics.

Further details of sample preparation, NMR measurements, and structure calculations are given in SI Appendix.

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