

3D structure of Alzheimer's amyloid-β(1-42) fibrils

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Alzheimer's disease is the most fatal neurodegenerative disorder wherein the process of amyloid- β (A β) amyloidogenesis appears causative. Here, we present the 3D structure of the fibrils comprising AB(1-42), which was obtained by using hydrogen-bonding constraints from quenched hydrogen/deuterium-exchange NMR, side-chain packing constraints from pairwise mutagenesis studies, and parallel, in-register β -sheet arrangement from previous solidstate NMR studies. Although residues 1-17 are disordered, residues 18-42 form a β -strand-turn- β -strand motif that contains two intermolecular, parallel, in-register β -sheets that are formed by residues 18-26 (β 1) and 31-42 (β 2). At least two molecules of $A\beta(1-42)$ are required to achieve the repeating structure of a protofilament. Intermolecular side-chain contacts are formed between the odd-numbered residues of strand β 1 of the *n*th molecule and the even-numbered residues of strand β 2 of the (n-1)th molecule. This interaction pattern leads to partially unpaired β-strands at the fibrillar ends, which explains the sequence selectivity, the cooperativity, and the apparent unidirectionality of $A\beta$ fibril growth. It also provides a structural basis for fibrillization

NMR | hydrogen/deuterium exchange

athological protein fibril formation can be observed in more than a dozen usually fatal human diseases, including Alzheimer's disease (AD) (1). Therefore, the fibrils' 3D structures are a focus of interest both for a molecular understanding of amyloidogenesis and for the development of innovative therapeutic and diagnostic approaches. Solid-state NMR studies have contributed substantially to the understanding of amyloid fibrils (2, 3). Only one 3D structure of an engineered amyloid fibril has yet been determined (4), owing in part to fibrils' noncrystalline, insoluble, and mesoscopically heterogeneous nature, which makes them difficult to access by established structural techniques such as x ray crystallography or solution state NMR.

The major component of AD-associated amyloid plaques is the 39- to 42-residue-long amyloid- β ($\Delta\beta$) peptide, which is generated from the amyloid precursor protein by the proteolytic activities of β - and γ secretase (5, 6). The $\Delta\beta$ (1-42) fragment is the dominant $\Delta\beta$ species in the amyloid plaques of Δ D patients, and, compared with $\Delta\beta$ (1-40), it displays a dramatically increased propensity to form amyloid fibrils in vitro (7-9). Several important structural characteristics of $\Delta\beta$ (1-42) fibrils have been determined (2), establishing that $\Delta\beta$ (1-42) fibrils form a cross β structure (10) that contains parallel, in-register β sheets (11). Here, we combined the available structural information on $\Delta\beta$ (1-42) fibrils with detailed amide hydrogen exchange measurements, pairwise mutagenesis, thiof lavin T (ThioT) binding, and high resolution cryoelectron microscopy to determine a 3D structure of $\Delta\beta$ (1-42) fibrils.

Materials and Methods

H/D Exchange. For the H/D exchange studies, recombinant 15 N labeled $A\beta(1-42)$ fibrils were used. The fibrils were resuspended in 10 mM Tris DCl (pH* = 7.7) containing 150 mM sodium ehloride and D_2O as the solvent. At suitable intervals, aliquots of the fibrillar suspension were sedimented at $100,000 \times g$ for 10

min. The pellet was frozen in pure D₂O on liquid nitrogen and lyophylized to quench H/D exchange. The fibrils were dissociated in DMSO containing 0.01% trifluoroacetic acid and, within 30 s, diluted 10 fold in DMSO. Immediately thereafter, a series of 80 2D [¹⁵N, ¹H] correlation spectra were measured for 4 h.

Structure Calculation. For the CYANA structure calculation of an Aβ(1 42) pentamer, we used the following upper limit restraints (UPLs): (i) two long range UPLs per monomer of $d_{\text{max}}(^{\text{F19}}\text{C}^{\alpha}, \text{G38}\text{C}^{\alpha}) = 8.0 \text{ Å}$ and $d_{\text{max}}(^{\text{A21}}\text{C}^{\alpha}, ^{\text{V36}}\text{C}^{\alpha}) = 8.0 \text{ Å}$ from pairwise mutagenesis data, (ii) the intermolecular salt bridge between residues D23 and K28 from pairwise mutagenesis was implemented by the intermolecular UPL of $d_{\text{max}}(^{\text{D23}}\text{C}^{\gamma,\text{K28}}\text{N}^{\zeta}) = 2.8 \text{ Å}$, and (iii) H bond constraints for residues 18 26 and residues 31 42 from H/D exchange were set to be consistent with intermolecular, parallel, in register β sheets (11). H bonds were implemented by two UPLs per H bond [$d_{\text{max}}(^{\text{HN}}, ^{\text{O}}) = 2.0 \text{ Å}$ and $d_{\text{max}}(^{\text{N}}, ^{\text{C}}) = 2.9 \text{ Å}$]. The H bonds also provided backbone torsion angle constraints ($-164 \le \Phi \le -74$ and $68 \le \Psi \le 158$) for residues 18-26 and residues 31-42.

Further details are provided in *Supporting Text*, which is published as supporting information on the PNAS web site.

Results and Discussion

Identification of \beta-Sheets. For the following structural studies, amyloid fibrils were produced from recombinant 35Mox AB(1 42) peptide that contains a methionine sulfoxide at position 35 (see Supporting Text). We first identified the sequence-specific positions of regular secondary structure elements, which are stabilized by hydrogen bonds of the peptide backbone. Using the quenched hydrogen-exchange NMR technique (12, 13) for $\frac{35\text{Mox}}{A}\beta(1-42)$ fibrils (see Supporting Text), the protonation state of most residues could be retained after dissociation into monomers (Fig. 1 C and D), thus allowing for the accurate determination of backbone amide exchange rates. Fig. 1 A and B show the [15N-1H]-correlation NMR spectra of 35Mox Aβ(1-42) monomers in DMSO corresponding to fully protonated and partially H/D-exchanged ^{35Mox}AB(1-42) fibrils, respectively. After exchange of the fibrils in a D₂O buffer for 1,990 h, ~30% of the peaks were absent from the spectrum (Fig. 1B; compare Fig. 1A). This observation is consistent with previous qHX (quenched hydrogen/deuterium exchange) mass spectrometry studies and the qualitative notion that 35MoxAB(1-42) fibrils contain a highly protected core structure (14, 15).

Fig. 1E shows the plot of the resulting exchange rates against the sequence of $A\beta(1-42)$. In the N-terminal segment of residues 1-17, approximately half of the population of each-backbone amide exchanged quickly ($\geq 10 \text{ h}^{-1}$); the other half exchanged

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Abbreviations: $A\beta$, amyloid- β ; AD, Alzheimer's disease; ThioT, thioflavin T; UPL, upper limit restraint.

Data-deposition: The coordinates of the fibrillar core structure of A β (1–42) have been deposited in the Protein Data Bank, www.pdb.org (PDB-ID-code-2BEG).

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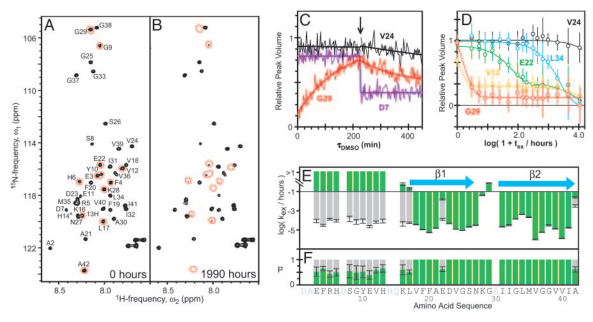


Fig. 1. Sequence-specific determination of regular secondary structure in $^{35\text{Mox}}$ A β (1–42) fibrils by quenched hydrogen exchange. (*A* and *B*) Fast heteronuclear multiple quantum correlation spectra of 15 N-labeled monomeric $^{35\text{Mox}}$ A β (1–42) in d₆-DMSO containing 0.01% d₁-trifluoroacetic acid corresponding to fully protonated (*A*) and partially exchanged (*B*) amyloid fibrils. Sequence-specific chemical shift assignment of peptide backbone amide cross-peaks are indicated. Red lines encircle cross-peaks that show a virtually complete loss of intensity between exchange times $t_{\rm ex} = 0$ and $t_{\rm ex} = 1,990$ h. The shown spectra are a sum of the first 10 spectra recorded at the given $t_{\rm ex}$. (C) intrinsic exchange curves of $^{35\text{Mox}}$ A β (1–42) in DMSO. The evolution of the peak volumes versus the time ($\tau_{\rm DMSO}$) after dissociation of partially hydrogen-exchanged amyloid fibrils into monomers is shown. The identities are indicated next to each curve. The arrow indicates the time point at which a solution of 50% (mol/mol) D₂O in H₂O was added to yield a final combined deuteron/proton concentration of 2.0 mol-liter⁻¹. Smooth, solid lines represent the monoexponential fits of the raw data. (*D*) Plot of peak volumes at $\tau_{\rm DMSO} = 0$ min versus logarithm of $t_{\rm ex}$ of $^{35\text{Mox}}$ A β (1–42) fibrils. Data for the residues V12 (yellow), E22 (green), V24 (black), G29 (red), and L34 (cyan) are shown. Smooth, solid lines represent fits of the data to a monoexponential equation (V24, L34, and G29) or to the sum of two monoexponential equations (V12 and E22). Error bars indicate the uncertainty in peak volume due to the noise level in each single measurement of initial peak volumes. (*E* and *F*) Plots of the observed exchange rates $k_{\rm ex}/h^{-1}$ (*E*) and the relative population *P* (*F*) against the amino acid sequence of $^{35\text{Mox}}$ A β (1–42). The blue arrows indicate 6 -strands.

slowly ($\leq 10^{-2}$ h⁻¹). This observation indicated structural heterogeneity in this segment, which is consistent with previous electron paramagnetic resonance studies (16). Conversely, in the segment of residues 18–42, the exchange was slow and almost exclusively monoexponential (Fig. 1 D and E), which suggests that this C-terminal segment constitutes the core structure of the $A\beta(1-42)$ fibril. Considering residues with exchange rates of $\leq 10^{-2}$ h⁻¹ as strongly hydrogen-bonded, the overall protection pattern (Fig. 1 E and F) suggests the presence of two β -strands, $\beta 1$ and $\beta 2$, which consist of residues $\approx 18-26$ and $\approx 31-42$, respectively.

Intersheet Side-Chain Packing. Fiber diffraction studies have established that AB fibrils contain a cross-B structure, in which β-sheets run parallel and β-strands run perpendicular to the fibril axis (10). It has previously been demonstrated by solidstate NMR that $A\beta(1-40)$ fibrils and $A\beta(1-42)$ fibrils consist of parallel, in register β sheets (11). Given the exact sequence location of the β -sheets as determined here by qHX-NMR (Fig. 1), this finding suggests two β -sheets that run along the fibril axis. Sheet β 1 is formed by residues \approx 18–26, and sheet β 2 is formed by residues ≈31–42, so that corresponding residues of adjacent monomers line up along the fibril. If fully extended, this structure would span ≈90 Å perpendicular to the fibril axis. However, from cryoelectron micrographs, we knew that the protofilaments contained dimensions of only $\approx 25 \times \approx 45 \text{ Å}$ (Fig. 4E). Therefore, it appeared plausible that the two β -sheets form a parallel β -sandwich that is stabilized by inter- β -sheet side-chain contacts. These geometric restraints reduced the search for the correct side-chain packing essentially to a two-dimensional problem of intersheet side-chain interactions. Considering (i) the nature of β -sheets with two distinct faces, (ii) the shortness of the loop that connects the two β -strands, (iii) the solvent inaccessibility of residue 19 (17), (iv) the experimental evidence that M35 is not part of the intersheet side-chain packing because it is accessible for intermolecular chemical cross-linking to residue 4 (18), and (v) parallel, in-register β -sheets and a salt bridge, which had previously been proposed for $A\beta(1-40)$ fibrils based on solid-state NMR data (19), the following intersheet side-chain interactions appeared likely in $A\beta(1-42)$ fibrils: intersheet contacts formed between residues F19/G38 and A21/V36 and a salt bridge between residues D23 and K28.

To test for the presence of these specific interstrand side-chain contacts in $A\beta(1-42)$ fibrils, we used the following pairwise mutagenesis strategy (Fig. 2A): The presumed side-chain interactions in the fibrillar core were disturbed by amino acid substitutions aimed to diminish or alter fibril formation. In a second step, compensating substitutions were introduced to restore the side-chain interaction pattern and, concomitantly, WT-like fibril formation. For these studies, we used the variant $^{35L}A\beta(1-42)$, which can be bacterially expressed and easily purified in substantial amounts (see *Supporting Text*; compare ref. 20). $^{35L}A\beta(1-42)$ forms fibrils with indistinguishable ultrastructural morphology, and its H/D-exchange pattern is similar to that of $^{35Mox}A\beta(1-42)$ (data not shown).

Eight different variants of the $^{35L}A\beta(1-42)$ peptide were generated and allowed to form fibrils for 2 months (see *Supporting Text*) before they were examined by negative staining EM (Fig. 2 *B–J*). Whereas nonordered precipitates and spherical aggregates were observed for the two variants K28D and G38F (Fig. 2 *B* and *E*), rod-like fibrils were obtained for the variants D23K and F19G (Fig. 2 *C* and *F*). Thus, all of these single amino

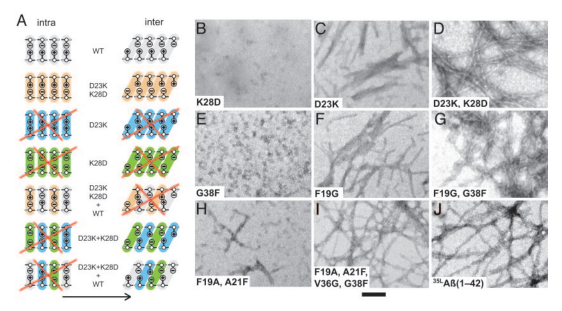


Fig. 2. Pairwise mutagenesis of $^{35L}A\beta(1-42)$ peptides. (A) Cartoon of intramolecular versus domain swapping-type interaction between monomers in the $A\beta(1-42)$ protofilament that consists of parallel, in-register β -sheets exemplified by the salt bridge formed between the charged residues D23 and K28. Individual $A\beta$ molecules are indicated as colored bars that correspond to a cross section along the protofilament axis through the C^{α} atom positions of one presumed interacting pair of amino acids. The identity of each variant Aβ peptide is indicated next to the corresponding schemes in rows 1-4. "intra" denotes the intramolecular scenario, and "inter" indicates the domain swapping-type scenario. Red diagonal crosses mark all scenarios that were considered to be incompatible with WT fibril formation. (B-J) Negative staining electron microscopy of $^{35L}A\beta(1-42)$ peptides. The variants are indicated on each image. All electron micrographs were recorded at a nominal magnification of \times 72,000. (Scale bar, 100 nm.)

acid variants formed aggregates that were readily distinguishable from WT $^{35L}A\beta(1-42)$ fibrils. In contrast, the complementary variant D23K-K28D formed fibrils that looked like WT (Fig. 2 D and J), suggesting that D23 and K28 form a salt bridge in $A\beta(1-42)$ fibrils. Likewise, the complementary variant F19G-G38F was capable of forming fibrils that looked similar to WT (Fig. 2 G and J), even though the majority of the observed aggregates had the typical morphology of premature fibrils (21, 22) (data not shown). Hence, the residues F19 and G38 appeared to form a hydrophobic interaction pair that connects the two β -strands.

To support the side-chain interaction pattern implied by the F19/G38 permutation, we then generated the double variant F19A-A21F. This peptide formed globular aggregates and some worm-like fibrils (Fig. 2H). The additional introduction of the compensatory substitutions V36G-G38V on strand β2 restored WT 35L A β (1–42) fibril formation (Fig. 2 *I* and *J*). Whereas this pairwise mutagenesis approach simultaneously exchanged two presumed interaction pairs along the β-strands, the F19/G38 cycle inverted a single interaction pair between strands. Thus, two independent sets of $A\beta(1-42)$ variants suggested a close spatial proximity between the pairs of residues F19/G38 and A21/V36. This approach established the register of side-chain contacts in the hydrophobic core of $A\beta(1-42)$ fibrils, which is formed by the odd-numbered residues of the β 1-strand and the even-numbered residues of the β 2-strand.

Intermolecular Arrangement. With the EM studies (Fig. 2), it is not possible to determine whether the side-chain interactions between the two β -strands are intramolecular or intermolecular. In a conformation with parallel, in-register β -sheet secondary structure, both scenarios could be envisioned (Fig. 2A). Because the fibril formation of peptide mixtures can distinguish between both scenarios (Fig. 2A, rows 5-7), we used a cofibrillization assay of binary $A\beta$ peptide mixtures measured by ThioT binding. A relative decrease of the ThioT fluorescence compared with pure $A\beta$ peptides indicates a different fibrillar structure or the presence of fewer fibrils (23).

The equilibrium ThioT-binding capacity of cofibrillized mixtures of WT $^{35L}A\beta(1-42)$ with one of the single-point variants D23K, K28D, F19G, and G38F, or the double variant F19A-A21F, was significantly reduced (Fig. 3 A, C, and E). These data further support the results from the EM studies (Fig. 2 B–J) that these variants are incapable of forming WT-like fibrils and of being incorporated into the final WT structure. In particular, ThioT binding of F19G and D23K was only slightly affected by the presence of a small amount of WT $^{35L}A\beta(1-42)$, whereas WT $^{35L}A\beta(1-42)$ peptide was strongly decreased by these mutants. This finding suggests that the F19G and D23K variants form fibrils of a different molecular structure, which is consistent with the observed morphological differences between the corresponding fibrils (Fig. 2 C, F, and J).

To find clues about the intermolecular interaction pattern, the sequence-permutated variants D23K-K28D, F19G-G38F, and F19A-A21F-V36G-G38V were mixed with 35LAB(1-42). However, even though these peptides formed WT-like fibrils, they did not readily coaggregate with $^{35L}A\beta(1-42)$ (Fig. 3 B, D, and F), suggesting the presence of intermolecular side-chain packing, which forbids the formation of mixed fibrils (Fig. 2A, row 5). To explore this possibility further, we coaggregated the complementary single variants G38F and F19G, or, alternatively, the variants D23K and K28D, none of which had formed WT-like fibrils (see Fig. 2). At equimolar mixing ratios, a local plateau of ThioT binding is observed for a mixture of G38F and F19G, and a maximum is observed for a mixture of D23K and K28D (Fig. 3 G and H), indicating cooperative fibril formation. Negative staining EM of the equimolar mixtures subsequently confirmed WT-like fibril formation (Fig. 3 *J* and *K*; compare Fig. 2*J*). Thus, this result is consistent with an intermolecular interaction scheme (Fig. 2A, row 6). We then reasoned that, despite the strong inhibitory effect of D23K or K28D on WT $^{35L}A\beta(1-42)$ fibril formation (Fig. 3A), a mixture of all three peptides should coaggregate, provided that D23K and K28D are present in

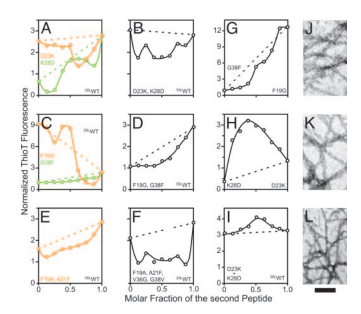


Fig. 3. Phase behavior of binary 35L A β (1–42) peptide mixtures. (A–F) Plots of the normalized ThioT fluorescence against the mixing ratio with 35L A β (1–42), as observed after 2 months of equilibration at room temperature. (G–f) Plots of the normalized ThioT fluorescence against the mixing ratio of two variant A β peptides. In A–f, the identities of the peptides present in the mixtures are indicated next to the corresponding y axis on each diagram. Plots corresponding to variants of the β 1-strand are shown in orange, plots corresponding to variants of the β 2-strand or its preceding loop are shown in green, and plots corresponding to double variants are shown in black. Dotted lines in the corresponding colors indicate the ThioT fluorescence that would be expected if the peptides in the mixture would not interact and thus aggregate independently of each other. The ThioT signal was normalized to the fluorescence signal of the single tyrosine in $A\beta$ (1–42). (f–f) Negatively stained electron micrographs corresponding to the samples at a molar fraction of 0.5 in G–f, recorded at a nominal magnification of ×72,000. (Scale bar, 100 nm.)

equimolar amounts (Fig. 2A, row 7). Indeed, our data confirmed this prediction (Fig. 3 I and L). In summary, these mixing experiments yielded a consistent data set, which suggests that $A\beta(1-42)$ fibrils are stabilized by intermolecular domain swapping-type side-chain interactions.

The direction of the observed intermolecular side-chain interactions is ambiguous, because each molecule in the A β fibril has two immediate neighbors. In a β -sheet, all H bonds involving the odd-numbered residues are on one side of an individual β -strand, and those of the even-numbered residues are on the opposite side. Taking the β 1-strand of a given A β (1–42) molecule as a reference, the "odd" neighbor extends the H-bonding network involving the odd-numbered residues of the β 1-strand (Fig. 6A, which is published as supporting information on the PNAS web site), and the "even" neighbor extends the H-bonding network involving its even-numbered residues (Fig. 6B). Aβ16-22m is a synthetic in vitro inhibitor of fibril extension (24) that contains residues 16-22 of A β , in which the backbone amide groups of L17, F19, and A21 have been methylated. Binding to the protofilament can therefore only occur on the odd side of the protofilament (Fig. 6). Tight binding of the inhibitor is straightforward for an odd intermolecular side-chain interaction pattern (Fig. 6A), because extensive hydrophobic contacts of the inhibitor with the protofilament can be formed. Thus, strand β 2 most likely forms side-chain contacts with strand β 1 of the odd neighbor.

Structure Calculation of A\beta(1–42) Protofilament. The calculation of protein structures from NMR-derived UPLs is a well established technique (25). Here, we apply it toward the determination of a

3D structure of the A β (1–42) protofilament (see Table 1 and Supporting Text, which are published as supporting information on the PNAS web site). The UPLs are derived from the presented secondary and tertiary structure information and the previously established structural properties of A β (1–42) fibrils (see Supporting Text) [i.e., 4.9-Å intermolecular distance (10, 11) and parallel, in-register β -sheets, which were demonstrated by solid-state NMR (10, 11)], including (i) H-bond restraints identified from H/D-exchange studies, (ii) an intermolecular salt bridge restraint between residues D23 and K28 derived from the mutagenesis studies, (iii) two intersheet distance restraints between F19 and G38, and A21 and V36, as derived from the pairwise mutagenesis studies, and (iv) intermolecular polymer restraints that take the repetitive nature of the protofilament into account. Based on the restraints, the structure calculation was performed for an A β (1–42) pentamer. Of 100 randomized starting conformers, the 10 best final structures had minimal constraint violations and small deviations from ideal geometry. The middle molecule of the pentamer had a backbone rms deviation of 1.18 Å relative to the mean coordinates for the peptide segment 18-42 (Fig. 7A, which is published as supporting information on the PNAS web site). It might appear surprising that only three intersheet constraints should suffice to yield a well defined A β (1–42) protofilament structure. However, a number of additional test calculations showed that the repetitiveness of the fibril restricts the conformational space of each individual A β (1–42) molecule to a narrow range of possible conformations (Table 1 and Supporting Text). Therefore, we concluded that the overall 3D structure of the A β (1-42) protofilament is defined unambiguously.

The 3D A β (1–42) Protofilament Structure. The 3D structure of the $A\beta(1-42)$ protofilament consists of two stacked, intermolecular, parallel, in-register β -sheets that perpetuate along the fibril axis (Fig. 4 A and B). The residues L17, F19, and A21 of sheet β 1 mediate the hydrophobic intermolecular contacts with the evennumbered residues of sheet β 2. The loop region of residues 27–30 is connected to sheet β 1 by means of the intermolecular salt bridge D23-K28, which also forms contacts to residues I32 and L34 of sheet β 2 (Fig. 4C; see also Fig. 8, which is published as supporting information on the PNAS web site). These tertiary interactions are distinct from a previously proposed model of $A\beta(1-40)$ fibrils (19). There, the side-chain interactions are intramolecular, and their register deviates by ≈10 Å. As a consequence, the hydrophobic side-chain packing in the core of the proposed $A\beta(1-40)$ protofilament structure is entirely different (Fig. 9, which is published as supporting information on the PNAS web site; for a detailed comparison see Supporting

In cryoelectron micrographs of the $^{35\text{Mox}}A\beta(1-42)$ fibrils, the protofilaments show dimensions of ~25 × ~45 Å, with an average spacing of \approx 45 Å (Fig. 4E). To test whether our protofilament structure can explain the morphology of $A\beta(1-$ 42) fibrils, we assembled 1,200 Aβ molecules into a model fibril (Fig. 4D Upper) by using MOLMOL (26). In this model, the 44-Å-wide nodes correspond to the length of the β -strands, and the 24-Å thickness of the protofilament corresponds to the β -sheet sandwich as shown in Fig. 4. Manipulation of Fig. 4D Upper by adding noise and blurring to a resolution of 2 nm resulted in an image (Fig. 4D Lower) that is strikingly similar to the experimental cryoelectron micrographs of frozen-hydrated $^{35\text{Mox}}\text{A}\beta(1-42)$ (Fig. 4E). Hence, the dimensions predicted by the protofilament structure are in good agreement with mesoscopic experimental observations. In contrast to previous model predictions for $A\beta(1-40)$ (19), these data suggest the presence of only one $A\beta(1-42)$ molecule per protofilament unit length.

Between the protofilaments, a 15-Å-wide area with low electron density is observed that appears to extend along the

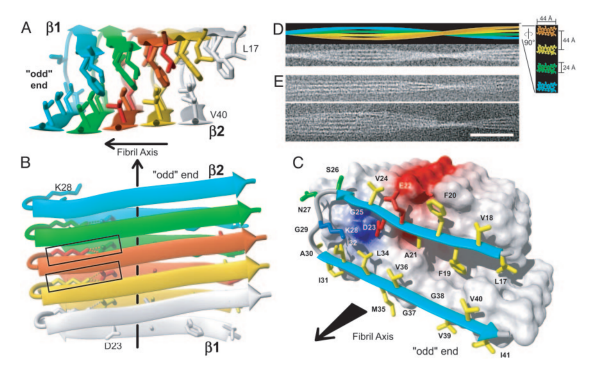


Fig. 4. The 3D structure of a 35MoxAβ(1-42) fibril. (A and B) Ribbon diagrams of the core structure of residues 17-42 illustrating the intermolecular nature of the inter- β -strand interactions. Individual molecules are colored. For example, the monomer at the odd end is shown in cyan. The β -strands are indicated by arrows, nonregular secondary structure is indicated by spline curves through the C^{α} atom coordinates of the corresponding residues, and the bonds of side chains that constitute the core of the protofilament are shown. In B, the intermolecular salt bridge between residues D23 and K28 is indicated by dotted lines, and the two salt bridges formed by the central $A\beta(1-42)$ molecule are highlighted by rectangles. (C) van der Waals contact surface polarity and ribbon diagram at the odd end of the $^{35\text{Mox}}A\beta(1-42)$ protofilament comprising residues 17-42. The β -sheets are indicated by cyan arrows, and nonregular secondary structure is indicated by gray spline curves. The hydrophobic, polar, negatively charged, and positively charged amino acid side chains are shown in yellow, green, red, and blue, respectively. Positively and negatively charged surface patches are shown in blue and red, respectively, and all others are shown in white. The direction of the fibril axis is indicated by an arrow pointing from even to odd. (D) (Upper) Simulation of a 35MoxAβ(1-42) fibril that consists of four protofilaments colored individually. Lower shows the same fibril in a noisy gray-scale image, which has been blurred corresponding to a resolution of 2 nm. In Right, a ×5-magnified cross section perpendicular to the fibril axis is shown, using the same color code. Dimensions are indicated. To match the experimental twist of the protofilamentof the fibril shown in E, a twist angle of 0.45° per molecule was used. (E) Two examples of cryoelectron micrographs of single $^{35\text{Mox}}A\beta(1-42)$ fibrils. (Scale bar,

entire fibril (Fig. 4E). The molecular details that lead to this regular spacing are currently unclear. However, it has been reported that residue 4 can form specific intermolecular crosslinks with methionine 35 in $A\beta(1-42)$ fibrils (18), suggesting that the structurally heterogeneous peptide segment 1–16 (see Fig. 1 E and F) is involved in interprotofilament binding and organization.

Fibril extension most likely occurs at the tip of the protofilaments. The domain swapping-type interactions between A β (1– 42) molecules imply the presence of distinct surfaces at the opposing ends: In contrast to the even end (Fig. 10, which is published as supporting information on the PNAS web site), a hydrophobic cleft is formed by strand $\beta 1$ and strand $\beta 2$ of the template at the odd end of the protofilament (Fig. 4C). An incoming $A\beta$ monomer could initially bind by means of the contiguous hydrophobic stretch of residues 17–21. This complex could be further stabilized by the formation of the intermolecular salt-bridge and backbone H bonds. However, the newly added monomer could still dissociate, because the \(\beta 2 \)-strand has substantially fewer interactions with the template than β 1. Hence, the last added A β monomer would be stabilized permanently only by the addition of the next monomer, suggesting a sequence-selective, cooperative mechanism of A β fibril extension that follows first-order kinetics. Recently, apparent unidirectional $A\beta(1-40)$ fibril growth with a constant slow rate was directly observed (27). Therefore, key experimental observations on fibril growth are in good agreement with the molecular mechanism suggested here, based on the structure of the A β (1– 42) protofilament. It appears evident that this mechanism also provides a rationale for the activity of A β peptide analog inhibitors comprising the A β peptide segment of residues 17–21 (24, 28). They would presumably bind to the hydrophobic stretch of residues 17-21 (Fig. 4C), thereby blocking this site and preventing incoming $A\beta$ molecules from extending the fibril (see above and Fig. 6).

Biological Activity of $A\beta$. During the presented structural investigation, morphological variants of $A\beta(1-42)$ aggregates were obtained that all share a virtually identical amino acid sequence. Because all deviations from WT morphology can be rationalized based on the presented structure, this set of variants appeared suitable for a structure–function study of A β (1–42). The 3-(4,5dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromideformazan assay (MTT assay) is a widely used assay of cell viability and cellular response to $A\beta$. To determine the cytotoxicity of the $^{35L}A\beta(1-42)$ variants, we used this assay in the CNS cell line B12, which is susceptible to A β toxicity (29) (see also Supporting Text). Table 2, which is published as supporting information on the PNAS web site, summarizes the results: All variants displayed a detectable neurotoxic response. With the exception of the variant K28D, the most toxic peptides all formed fibrils (Fig. 2 B-J and Table 1) and displayed toxicity similar to synthetic $A\beta(25-35)$, which was used as a reference. A lower but still significant toxicity was observed for the variants F19G and

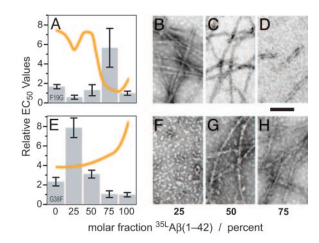


Fig. 5. Morphology and neurotoxicity of $A\beta(1-42)$ peptides. (A and E) Bar diagrams of the relative EC₅₀ values of binary mixtures of F19G (A) and G38F (E) with $^{35L}A\beta(1-42)$ as obtained by the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide assay using B12 cells. The orange lines indicate the relative ThioT fluorescence of the corresponding samples. (B-D and F-H) Negative stained electron micrographs of the individual peptide mixtures in A and E are shown to the right of the corresponding bar diagrams. The molar fraction of WT $^{35L}A\beta(1-42)$ is indicated below. (Scale bar, 100 nm.)

G38F, which form rod-like fibrils and seed-like (21) oligomers (Fig. 2). Finally, the variants F19A–A21F and F19G–G38F, which form mostly premature fibrils, displayed the lowest toxicity. These data suggest that mature $A\beta$ fibrils, as well as some types of oligomers, are neurotoxins.

To further investigate the structure–toxicity relationship of $A\beta$ aggregates, we next asked how mixing of structurally incompatible peptides might affect toxicity. Fig. 5*A* shows the relative EC₅₀ toxicity values for various ratios of F19G and WT ^{35L} $A\beta$ (1–42). Two of the ratios displayed near WT toxicity. However, at 75% WT ^{35L} $A\beta$ (1–42), the EC₅₀ value was increased 6-fold

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compared with pure WT peptide, indicating a significantly reduced toxicity. This sample contained very few short fibrils (Fig. 5D), which supports the previous notion that fibrils are toxic. In contrast, the exclusively oligomer-forming variant G38F possesses a significant toxicity, which can be abolished by the presence of a small amount of WT peptide (Figs. 5E and 2D). These data suggest that a specific oligomer tertiary structure is required for G38F toxicity. At higher fractions of WT peptide, fibrils (Fig. 5 G and H) were observed, which were again more toxic.

The structure–toxicity relationship indicates that $A\beta$ amyloid fibrils contain an intrinsic toxicity that correlates with their morphology (e.g., more regular and longer fibrils tend to be more toxic). This observation, which is in agreement with previous observations on various aggregation stages of $^{35\text{Mox}}A\beta(1-42)$ (21), supports the concept that neuronal degeneration in AD is caused by the deposition of $A\beta$ amyloid (30). However, our data also suggest that oligomers that are similar to small diffusible ligands (ADDLs, or $A\beta$ -derived diffusible ligands) (31) or protofibrils (32) are toxic and may play a central role in AD. Such oligomeric structures, which seem to be involved in a variety of human amyloid diseases, appear to have common structural features that are distinct from amyloid fibrils (33).

The presented structure of $A\beta(1-42)$ protofilament explains the sequence selectivity, the cooperativity, and the unidirectionality of $A\beta$ fibril growth and also provides a structural explanation for the working mechanism of current $A\beta$ fibrillization inhibitors. Furthermore, it shows a correlation between neurotoxicity and various conformational entities of $A\beta$ that might be involved in the amyloidogenesis of AD. These studies may therefore open an avenue toward a comprehensive molecular understanding of amyloidogenesis of AD and structure-assisted development of potent anti-AD drugs and AD diagnostic markers.

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