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Site-Specific Identification of Non- β -Strand Conformations in Alzheimer's β -Amyloid Fibrils by Solid-State NMR

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ABSTRACT The most well-established structural feature of amyloid fibrils is the cross- β motif, an extended β -sheet structure formed by β -strands oriented perpendicular to the long fibril axis. Direct experimental identification of non- β -strand conformations in amyloid fibrils has not been reported previously. Here we report the results of solid-state NMR measurements on amyloid fibrils formed by the 40-residue β -amyloid peptide associated with Alzheimer's disease (A β_{1-40}), prepared synthetically with pairs of ¹³C labels at consecutive backbone carbonyl sites. The measurements probe the peptide backbone conformation in residues 24-30, a segment where a non- β -strand conformation has been suggested by earlier sequence analysis, cross-linking experiments, and molecular modeling. Data obtained with the fpRFDR-CT, DQCSA, and 2D MAS exchange solid-state NMR techniques, which provide independent constraints on the ϕ and ψ backbone torsion angles between the labeled carbonyl sites, indicate non- β -strand conformations at G25, S26, and G29. These results represent the first site-specific identification and characterization of non- β -strand peptide conformations in an amyloid fibril.

INTRODUCTION

Amyloid fibrils are filamentous structures, with typical diameters of roughly 5-20 nm and lengths up to several microns, formed by numerous peptides and proteins with disparate sequences and molecular weights (Sunde and Blake, 1998). Current interest in amyloid fibrils arises from their occurrence in amyloid diseases (Sipe, 1992), including Alzheimer's disease, type-2 diabetes, Parkinson's disease, Huntington's disease, and prion diseases, and from fundamental questions regarding the molecular mechanism of amyloid formation and the nature of the intermolecular interactions that make the amyloid fibril a stable form for an extremely diverse class of polypeptides (Chiti et al., 1999; MacPhee and Dobson, 2000). Amyloid fibrils share certain defining properties, including an unbranched morphology in electron microscope (EM) images, pronounced and typically green optical birefringence after Congo Red staining, and an x-ray scattering pattern indicative of a "cross- β " structure, i.e., an extended β -sheet structure formed by intermolecular hydrogen bonding in which the β -strand segments run approximately perpendicular to and the hydrogen bonds run approximately parallel to the long axis of the fibril. Definitive evidence for the cross- β structure comes from x-ray diffraction studies (Kirschner et al., 1987; Inouye et al., 1993; Sunde et al., 1997) and more recently from EM (Serpell and Smith, 2000).

More detailed and site-specific structural information has been difficult to obtain in amyloid fibrils because of their inherently noncrystalline and insoluble nature. Recently,

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several groups have used solid-state NMR techniques to characterize the supramolecular organization of the β -sheets in amyloid fibrils, especially fibrils formed by the β -amyloid $(A\beta)$ peptide associated with Alzheimer's disease (Antzutkin et al., 2000; Antzutkin et al., 2002; Balbach et al., 2002; Petkova et al., 2002) and several A β fragments that serve as important model systems (Lansbury et al., 1995; Benzinger et al., 1998; Gregory et al., 1998; Balbach et al., 2000; Benzinger et al., 2000; Burkoth et al., 2000). One unanticipated result from solid-state NMR spectroscopy is that the β -sheets in full-length A β fibrils and in certain A β fragment fibrils have a parallel, in-register structure (Benzinger et al., 1998; Gregory et al., 1998; Antzutkin et al., 2000; Benzinger et al., 2000; Burkoth et al., 2000; Antzutkin et al., 2002; Balbach et al., 2002), whereas the β -sheets in other A β fragment fibrils have antiparallel structures (Lansbury et al., 1995; Balbach et al., 2000). Linewidths in solidstate NMR spectra of amyloid fibrils indicate a high degree of structural order in certain segments of the peptide sequence (Balbach et al., 2000; Ishii, 2001; Petkova et al., 2002). In the case of full-length A β fibrils, NMR linewidths and intermolecular dipole-dipole couplings indicate a disordered N-terminal segment of ~ 10 residues (Balbach et al., 2002; Petkova et al., 2002), consistent with proteolysis data (Roher et al., 1993; Saido et al., 1996; Kheterpal et al., 2001). ¹³C NMR chemical shifts have been used to identify specific peptide segments that form β -strands in amyloid fibrils (Balbach et al., 2000; Ishii, 2001; Laws et al., 2001; Petkova et al., 2002).

Although the cross- β structure is the predominant structural motif in amyloid fibrils, the dimensions of $A\beta$ fibrils suggest that the full-length peptide ($A\beta_{1-40}$, 40 residues; $A\beta_{1-42}$, 42 residues) adopts a structure that includes non- β -strand conformations. The thinnest $A\beta_{1-40}$ and $A\beta_{1-42}$ fibrils observed in EM images have diameters of 6 \pm 1 nm

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(Goldsbury et al., 2000; Antzutkin et al., 2002; Petkova et al., 2002). The length of a β -strand formed by M residues is approximately $M \times 0.34$ nm. Given a 10-residue disordered N-terminal segment, a minimum diameter of 10 nm would be expected for $A\beta_{1-40}$ and $A\beta_{1-42}$ fibrils if the remainder of the peptide formed a single, continuous β -strand. Circular dichroism spectra and secondary structure predictions based on the A β amino acid sequence (Kirschner et al., 1987; Hilbich et al., 1991) have suggested a β -turn in residues 23-30 (sequence DVGSNKGA). A β-turn (Lazo and Downing, 1998; George and Howlett, 1999; Li et al., 1999) or other non- β -strand conformation (Tjernberg et al., 1999) in this segment has been incorporated into several structural models. Measurements of enzymatic N-terminal proteolysis of $A\beta_{10-43}$ analogs with intramolecular disulfide linkages have been described as experimental support for a β -turn encompassing residues 26-29 (Hilbich et al., 1991).

In this paper, we report solid-state NMR data that demonstrate a non- β -strand conformation in residues 24-30 of $A\beta_{1-40}$ fibrils and permit estimation of the ϕ and ψ torsion angles, which define the peptide backbone conformation, at specific sites in this segment. The measurements were carried out on a series of samples in which pairs of ¹³C labels were introduced at consecutive backbone carbonyl sites. Three solid-state NMR techniques that place independent constraints on the ϕ and ψ values between the two ¹³C labels were employed, namely constant-time finite-pulse radiofrequency-driven recoupling (fpRFDR-CT) (Ishii et al., 2001), double-quantum chemical shift anistropy (DQCSA) spectroscopy (Blanco and Tycko, 2001), and two-dimensional magic-angle spinning (2D MAS) exchange spectroscopy (Tycko et al., 1996; Weliky and Tycko, 1996; Tycko and Berger, 1999). These techniques have previously been demonstrated on model compounds and applied in structural studies of a variety of peptides and proteins, including amyloid fibrils (Long and Tycko, 1998; Weliky et al., 1999; Balbach et al., 2000; Blanco et al., 2001; Antzutkin et al., 2002; Balbach et al., 2002). The results in this paper represent the first site-specific characterization of non-βsheet structure in an amyloid fibril.

MATERIALS AND METHODS

Peptide synthesis, purification, and fibrillization

Peptides with the human $A\beta_{1-40}$ sequence DAEFRHDSGYEVHHQKLVF-FAEDVGSNKGAIIGLMVGGVV were synthesized on an Applied Biosystems (Foster City, CA) Model 433A automated solid-phase peptide synthesizer, cleaved from the synthesis resin, and purified by high-performance liquid chromatography as previously described (Antzutkin et al., 2000; Balbach et al., 2002). $A\beta_{1-40}$ was fibrillized by dissolution in deionized water at ~ 1 mM concentration (estimated from lyophilized peptide weight), adjustment to pH 7.4 by dropwise addition of dilute NaOH (from an initial pH ≈ 3.3), addition of 0.01% NaN₃, and incubation at room temperature with gentle rocking for 10–20 days. Solutions typically became noticeably viscous after several hours and gelled within several days. Incubated solutions were lyophilized for solid-state NMR measurements.

The extent of fibrillization was assessed by one-dimensional (1D) 13 C NMR spectroscopy (see below). Samples found to be incompletely fibrillized were reincubated for an additional 10–20 days. Typical solid-state NMR samples were 10 mg.

Samples were prepared with 13 C labels at the backbone carbonyl sites of D23 and V24 (D23,V24-A β_{1-40}), V24 and G25 (V24,G25-A β_{1-40}), G25 and S26 (G25,S26-A β_{1-40}), K28 and G29 (K28,G29-A β_{1-40}), and G29 and A30 (sample G29,A30-A β_{1-40}). Amino acids with carboxyl 13 C labels were purchased from Cambridge Isotopes Laboratories (Cambridge, MA) and Isotec (Miamisburg, OH) either with protecting groups required for peptide synthesis or as the bare amino acids. Fluorenylmethoxycarbonyl (FMOC) and side-chain protection reactions were performed by Midwest Biotech (Fishers, Indiana) when required. Because 13 C-labeled asparagine could not be obtained, samples labeled at the backbone carbonyl site of N27 could not be prepared.

Protection of ¹³C-labeled aspartate, performed by Midwest Biotech, resulted in a mixture of the desired product (N- α -FMOC-L-aspartic acid β -tbutyl ester) and an undesired isomer (N- α -FMOC-L-aspartic acid α -t-butyl ester) that incorporates into the peptide as β -aspartate with a ¹³C-labeled side chain. The undesired isomer was present at a level of $\sim 20\%$ in D23,V24-A β_{1-40} after purification, based on ¹³C NMR (Fig. 1). Although substitution of β -aspartate for aspartate may introduce a local structural perturbation in peptide chains containing the undesired isomer in the D23,V24-A β_{1-40} sample, peptide chains containing this isomeric substitution did not contribute significantly to the fpRFDR-CT, DQCSA, and 2D MAS exchange measurements (because of the larger intramolecular ¹³C-¹³C distance and because of resolvable chemical shift differences) and hence did not affect any conclusions in this study regarding the conformation of $A\beta_{1-40}$ in amyloid fibrils. We emphasize that this impurity was present only in the D23,V24-A β_{1-40} sample, and that data for this sample are not the basis for our report of non- β -strand conformations in $A\beta_{1-40}$ fibrils (see below).

Solid-state NMR measurements

1D ¹³C NMR spectra were obtained with cross-polarization, magic-angle spinning (MAS), and high-power proton decoupling, using a Varian (Palo Alto, CA) Chemagnetics Infinity spectrometer operating at a proton NMR frequency of 599.1 MHz and a homebuilt probe with a 2.5-mm Varian/Chemagnetics MAS module. Cross-polarization contact times were 1.5 ms. Decoupling fields were 120 kHz.

DQCSA and fpRFDR-CT NMR measurements were performed on Varian/Chemagnetics Infinity spectrometers operating at proton NMR frequencies of 400.6 MHz and 399.2 MHz, using Varian/Chemagnetics 3.2 mm MAS probes. DQCSA data were acquired as described by Blanco and Tycko (2001), using an MAS frequency of 5.0 kHz, DQ preparation and mixing periods of 6.4 ms, 13 C π pulse lengths of 15.0 μ s, and proton decoupling fields of 110 kHz during the RFDR sequence in these periods. DQ preparation and mixing periods were deliberately kept short to minimize possible effects of intermolecular couplings on the DOCSA data. The fpRFDR-CT data were acquired as described by Ishii et al. (2001), using an MAS frequency of 20.0 kHz, 13 C π pulse lengths of 10.0 μ s and proton decoupling fields of 110 kHz during the fpRFDR sequence, and with the parameters K = 3 and M + 2N = 96, in the notation of Ishii et al. Pulsed spin locking was used to enhance sensitivity in both fpRFDR-CT and DOCSA measurements, as described by Petkova and Tycko (2002). Two-dimensional MAS exchange measurements were performed on a Chemagnetics CMX spectrometer operating at a proton NMR frequency of 599.1 MHz, using the 2.5-mm MAS probe. Two-dimensional MAS exchange data were acquired as described by Weliky and Tycko (Tycko et al., 1996; Weliky and Tycko, 1996), using an MAS frequency of 3.78 kHz and 500-ms exchange periods. These data were processed in the manner of Hagemeyer et al. (Hagemeyer et al., 1989; Tycko and Berger, 1999). Radio-frequency pulses were actively synchronized with an MAS tachometer signal in fpRFDR-CT, DQCSA, and 2D MAS exchange measurements. Total

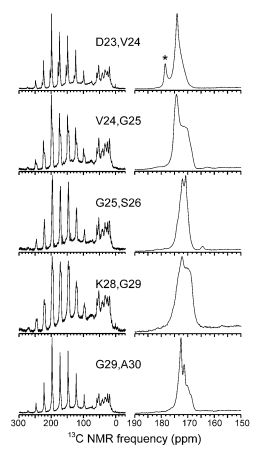


FIGURE 1 Solid-state 13 C NMR spectra of $A\beta_{1.40}$ fibril samples with 13 C labels at the backbone carbonyl sites of the indicated residues, obtained with MAS, cross polarization from protons, and high-power proton decoupling. Spectra on the left were obtained at a 3.75-kHz MAS frequency. Strong spinning sidebands between 90 ppm and 250 ppm for the carbonyl signals indicate a rigid peptide backbone. Signals between 10 ppm and 60 ppm arise from natural-abundance aliphatic 13 C nuclei. Spectra on the right, showing only the carbonyl region, were obtained at a 15-kHz MAS frequency. Asterisk in the D23,V24-A $\beta_{1.40}$ spectrum indicates a labeled side-chain carboxylate signal from a β -aspartate impurity at residue 23 (see Materials and Methods).

experiment times were typically 8 h, 20 h, and 8 days for fpRFDR-CT, DQCSA, and 2D MAS exchange measurements, respectively.

NMR data analysis

DQCSA, fpRFDR-CT, and 2D MAS exchange data were analyzed by comparison with numerical simulations, performed with Fortran programs written specifically for this purpose (Tycko et al., 1996; Blanco and Tycko, 2001). Simulations of fpRFDR-CT data were carried out for a four-spin system (two carbonyl labels on two parallel chains), using an explicit evaluation of the quantum mechanical evolution operator for the full pulse sequence under MAS and including typical carbonyl CSA parameters as well as all dipole-dipole couplings. Four-spin fpRFDR-CT simulations for a full 5° grid in ϕ and ψ with 864 powder orientations required ~110 h of processor time (Pentium III, 1 GHz). Six-spin simulations (three carbonyl labels on three parallel chains) were approximately six times slower and produced fpRFDR-CT curves that were not significantly different. Simulations of DQCSA and 2D MAS exchange data were carried out for

a two-spin system (two carbonyl labels on one chain). DQCSA simulations also used an explicit evaluation of the evolution operator for the full pulse sequence under MAS. Two-dimensional MAS exchange simulations used numerical evaluations of integral expressions for the crosspeak intensities (Tycko et al., 1996; Tycko and Berger, 1999). Effects of intermolecular couplings on 2D MAS exchange and DQCSA data were ignored for the following reasons: 1), 2D MAS exchange data are sensitive only to the relative orientations, and not the displacements, of the labeled carbonyl groups. Crosspeaks arise only from exchange between sites with different orientations. Assuming that each peptide chain is labeled at a site A and a site B and that neighboring chains in the cross- β amyloid fibril structure are related by translational symmetry along the fibril axis, then exchange between sites A or sites B of neighboring chains generates no crosspeak intensity, whereas exchange between site A of one chain and site B of a neighboring chain generates the same crosspeak intensities as intramolecular exchange; 2), DQCSA data depend on the excitation of DQ coherences between carbonyl labels by an RFDR sequence (Gullion and Vega, 1992; Bennett et al., 1998a). The RFDR sequence only recouples two ¹³C sites under MAS if they have different isotropic shifts or different CSA tensor orientations. Again assuming translational symmetry, no DQ coherence can be directly excited between sites A or sites B of neighboring chains. Although DQ coherences can be excited between site A of one chain and site B of a neighboring chain, numerical four-spin simulations show that the contributions of such coherences to DQCSA signals are negligible under the conditions of our experiments.

In all simulations, a standard carbonyl 13 C chemical shift anisotropy (CSA) tensor orientation was assumed, with the δ_{33} axis perpendicular to the carbonyl plane and the δ_{22} axis at 130° to the peptide C-N bond, as supported by experimental (Oas et al., 1987; Teng et al., 1992; Asakura et al., 1998) and ab initio quantum chemical (Walling et al., 1997) studies. CSA principal values were determined experimentally for each doubly labeled sample (see below).

The χ^2 deviation plotted in Fig. 4 has the general form:

$$\chi^{2}(\phi,\psi) \equiv \frac{1}{\sigma_{\text{rms}}^{2} + \sigma_{\text{sim}}^{2}} \sum_{i=1}^{N} [E_{i} - \lambda_{1}(\phi,\psi)S_{i}(\phi,\psi) - \lambda_{2}(\phi,\psi)]^{2},$$

where $\{E_i\}$ are the experimental data values, $\{S_i(\phi,\psi)\}$ are the simulated values for the given torsion angles, N is the number of values, $\sigma_{\rm rms}$ is uncertainty in the experimental data due to the root-mean-squared noise in the experimental spectra, and σ_{sim} is uncertainty in the simulations. For DQCSA and 2D MAS exchange simulations, $\sigma_{\rm sim}$ was taken to be negligible compared with $\sigma_{\rm rms}$. For fpRFDR-CT simulations, $\sigma_{\rm sim}$ was taken to be 5% of the maximum experimental data value, reflecting the high signal-to-noise of the experimental spectra and uncertainty about effects of intermolecular couplings beyond the four spins included in these simulations. $\lambda_1(\phi,\psi)$ is a scaling factor calculated to minimize χ^2 for the given torsion angles, required because the intensity of the NMR signals is not measured on an absolute scale (Tycko et al., 1996). $\lambda_2(\phi,\psi)$ is an offset parameter, also calculated to minimize χ^2 , that was included only in fpRFDR-CT analyses to account for contributions of natural-abundance 13C nuclei and residual unfibrillized $A\beta_{1-40}$ to the data. Good agreement between simulations and experiments corresponds to the condition $\chi^2 \sim N$.

RESULTS

One-dimensional ¹³C NMR spectroscopy

Fig. 1 shows 1D 13 C MAS NMR spectra of the five doubly labeled A β_{1-40} fibril samples. As previously demonstrated (Antzutkin et al., 2000; Antzutkin et al., 2002; Balbach et al., 2002), nearly complete fibrillization of A β_{1-40} is indicated by the relative sharpness of the natural-abundance aliphatic 13 C

NMR signals (10–60 ppm spectral region), especially the pronounced splitting of the α -carbon signals into peaks at 52 ppm and 58 ppm. Two-dimensional ^{13}C - ^{13}C exchange spectra of $A\beta_{1-40}$ samples with uniformly labeled residues (Petkova et al., 2002) indicate that the peak at 58 ppm arises from the α -carbon signals of valine and isoleucine residues, which converge to this NMR frequency when β -strands form concomitantly with fibrillization.

Spectra at MAS frequency $\nu_{\rm R}=3.75$ kHz (*left* side of Fig. 1) show the spinning sideband patterns expected for carbonyl $^{13}{\rm C}$ labels. CSA principal values determined from these spectra (Herzfeld and Berger, 1980) are given in Table 1. CSA principal values in Table 1 represent average values for the two labels in each sample except in the case of V24,G25-A β_{1-40} , where signals from the two labels could be deconvolved to permit determination of the principal values of the individual sites. All CSA values are consistent with the absence of large-amplitude motions of the carbonyl groups, i.e., a rigid peptide backbone.

Significant variations in the carbonyl ¹³C lineshapes are observed (right side of Fig. 1). Due to overlap, it is not possible to extract the lineshape for each individual ¹³Clabeled site from these spectra. Certain sites (e.g., the A30 carbonyl) apparently have 13 C NMR linewidths of ~ 1.5 ppm, whereas other sites (especially the G25 and G29 carbonyls) have larger linewidths or exhibit multiple, partially resolved lines. These lineshapes appear to indicate a mixture of local structural uniformity at some sites and structural heterogeneity at other sites in the segment from D23 through A30. For comparison, carbonyl ¹³C NMR linewidths for V12, L17, F20, V24, L34, and V39 in $A\beta_{1-40}$ fibril samples with single ¹³C labels have been found to be 1.3-3.0 ppm, with both F20 and L34 exhibiting two partially resolved carbonyl lines in ¹³C MAS spectra (Balbach et al., 2002). We associate the observation of multiple ¹³C NMR lines for single sites with the observation of multiple fibril morphologies in typical amyloid fibril preparations (Goldsbury et al., 2000; Jimenez et al., 2002). The degree of structural heterogeneity at the molecular level, especially the

TABLE 1 Chemical shift anisotropy parameters for 13 C-labeled carbonyl carbons in A β_{1-40} fibrils, determined from analyses of spinning sideband intensities in spectra as in Fig. 1

Sample	δ (ppm)	η
$D23,V24-A\beta_{1-40}$	80.4	0.68
V24,G25-A β_{1-40} (V24 carbonyl)	86.9	0.70
$V24,G25-A\beta_{1-40}$ (G25 carbonyl)	78.3	0.80
G25,S26-A β_{1-40}	78.4	0.77
$K28,G29-A\beta_{1-40}$	79.3	0.85
G29,A30-A β_{1-40}	79.2	0.75

Except in the case of V24,G25-A $\beta_{1.40}$, values are averages for two labeled sites with unresolved 13 C NMR signals. $\delta \equiv \delta_{iso} - \delta_{33}$ and $\eta \equiv (\delta_{11} - \delta_{22})/(\delta_{iso} - \delta_{33})$, where δ_{11} , δ_{22} , and δ_{33} are the principal values of the chemical shift anisotropy tensor and $\delta_{iso} \equiv (\delta_{11} + \delta_{22} + \delta_{33})/3$. Uncertainties in δ and η are approximately $\pm 4\%$.

ranges of ϕ and ψ values, can not be determined in quantitative terms from these data.

Qualitative interpretation of structural data

As depicted in Fig. 2, ¹³C-labeling of two consecutive backbone carbonyl sites, at residues i-1 and i, permits the acquisition of solid-state NMR data that place constraints on the ϕ and ψ torsion angles of residue i. The fpRFDR-CT technique measures the ¹³C-¹³C magnetic dipole-dipole coupling, which depends on the ¹³C-¹³C distance and therefore on the ϕ angle. The DQCSA and 2D MAS exchange techniques measure the relative orientation of the two ¹³C CSA tensors, which depends on the relative orientation of the carbonyl groups and therefore on both ϕ and ψ . For all residues in any β -strand, the ϕ and ψ values are approximately $-140^{\circ} \pm 20^{\circ}$ and $140^{\circ} \pm 20^{\circ}$, respectively. Because of the uniformity of the ϕ and ψ values in a β -strand, if one prepares a series of doubly carbonyllabeled peptide samples with 13 C labels in a β -strand segment, one expects all samples in the series to yield nearly the same fpRFDR-CT, DQCSA, and 2D MAS exchange data. At a qualitative level, significant differences in these data from different samples would indicate a non- β -strand conformation.

Fig. 3, a and b, present fpRFDR-CT and DQCSA data for the five doubly labeled $A\beta_{1-40}$ fibril samples described above. Significant differences in the data from different samples are indeed observed, indicating that the ϕ and ψ values in the segment from D23 through A30 are not uniform in $A\beta_{1-40}$ fibrils. In fpRFDR-CT measurements, one records the decay of the 13 C NMR signal from the labeled sites with

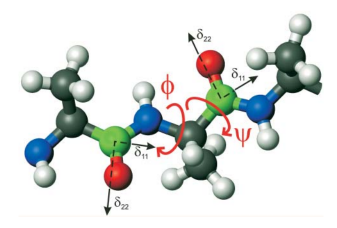


FIGURE 2 Illustration of the use of double carbonyl 13 C labeling to determine polypeptide backbone torsion angles ϕ and ψ . A segment of a polyalanine chain is shown, with labeled sites in green. Directions of the principal axes of the carbonyl CSA tensors are shown. The δ_{33} principal axis is perpendicular to the δ_{11} and δ_{22} axes. The fpRFDR-CT solid-state NMR technique primarily measures the 13 C- 13 C distance, which depends on ϕ . The DQCSA and 2D MAS exchange techniques measure the relative orientation of the two CSA tensors, which depends on both ϕ and ψ .

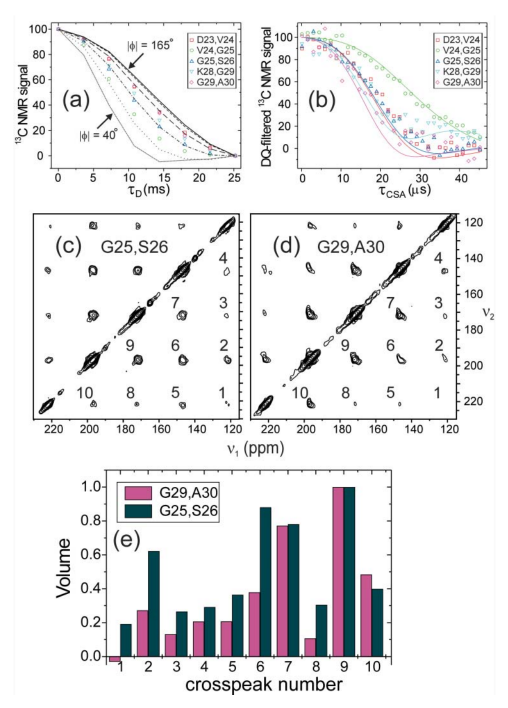


FIGURE 3 (a) Experimental fpRFDR-CT data for the five doubly labeled $A\beta_{1-40}$ fibril samples. Curves are simulations for $|\phi|$ between 40° and 165°, in 25° increments. (b) Experimental DQCSA data. Color-coded curves are simulations for the ϕ and ψ values in Table 2. (c and d) Experimental 2D MAS exchange spectra for two of the samples, represented as contour plots with contour levels increasing by factors of 1.6. Crosspeaks connecting MAS sidebands are numbered. (e) Crosspeak volumes from the 2D MAS exchange spectra, after symmetrization, normalized to the volume of the strongest crosspeak. The significant differences among fpRFDR-CT data, DOCSA data, and 2D MAS exchange data for different samples indicate significant variations in the ϕ and ψ values, demonstrating the presence of non- β -strand conformations at certain sites in the segment from D23 to A30.

increasing dipolar dephasing time $\tau_{\rm D}$, due to the $^{13}{\rm C}^{-13}{\rm C}$ dipole-dipole couplings (Ishii et al., 2001). A more rapid decay indicates a shorter $^{13}{\rm C}^{-13}{\rm C}$ distance, implying a smaller value of $|\phi|$. In Fig. 3 a, the most rapid fpRFDR-CT decay is observed for V24,G25-A β_{1-40} fibrils, whereas the slowest decays are observed for D23,V24-A β_{1-40} fibrils and G29,A30-A β_{1-40} fibrils. In DQCSA measurements, one records the decay of the double-quantum filtered $^{13}{\rm C}$ NMR signal with increasing CSA evolution time $\tau_{\rm CSA}$. The decay rate of DQCSA signals depends on ϕ and ψ in a complicated

manner (Blanco and Tycko, 2001). According to simulations, the slowest DQCSA decays occur near $\phi = -60^{\circ}$ and $\psi = -65^{\circ}$ (or $\phi = 60^{\circ}$ and $\psi = 65^{\circ}$). The most rapid decays occur near $\phi = \pm 180^{\circ}$ or 0° and $\psi = \pm 180^{\circ}$ or 0° . In Fig. 3 b, the slowest DQCSA decay is observed for V24,G25-A β_{1-40} fibrils, whereas the most rapid decay is observed for G29,A30-A β_{1-40} fibrils.

Fig. 3, c and d, show 2D MAS exchange spectra for the G25,S26-A β_{1-40} and G29,A30-A β_{1-40} fibril samples. In these spectra, the intensities of crosspeaks that connect

spinning sideband signals of the two labeled carbonyl sites are complicated functions of both ϕ and ψ (Tycko et al., 1996; Tycko and Berger, 1999). Significant differences in crosspeak intensities for these two samples are apparent, with greater intensity in crosspeaks further from the diagonal in the case of G25,S26-A β_{1-40} fibrils.

Quantitative analysis of structural data

We analyze the fpRFDR-CT, DQCSA, and 2D MAS exchange data by simulating the data for all possible ϕ,ψ pairs (in 5° increments) and plotting the dependence of the χ^2 deviation between experimental data and simulations on ϕ and ψ . As previously demonstrated for DQCSA and 2D MAS measurements on model peptides of known structure (Tycko et al., 1996; Weliky and Tycko, 1996; Blanco and Tycko, 2001), these simulations are sufficiently accurate that the absolute minimum value χ^2_{\min} occurs within $\pm 10^{\circ}$ of the correct ϕ and ψ values. In χ^2 plots for DQCSA and 2D MAS exchange data, additional local minima χ^2_{loc} are typically observed at other ϕ and ψ values where the simulated data are similar to data for the correct ϕ and ψ values. When χ^2_{\min} $\approx \chi^2_{\rm loc}$, for example due to insufficient signal-to-noise in the experiments, the correct ϕ and ψ values can be determined by combining the results of independent techniques, which may then exhibit only a single common χ^2 minimum at the correct values (Bennett et al., 1998b; Weliky et al., 1999; Balbach et al., 2000).

Representative χ^2 contour plots, for G25,S26-A β_{1-40} fibrils and G29,A30-A β_{1-40} fibrils, are shown in Fig. 4. Ideally, the best-fit (i.e., minimum χ^2) regions in the contour plots for fpRFDR-CT, DQCSA, and 2D MAS exchange data for a given sample would overlap in a single small area of the ϕ, ψ plane, which would then indicate the correct values of ϕ and ψ . As can be seen in Fig. 4, precise overlap of the best-fit regions for the DQCSA and 2D MAS exchange data is not observed. We attribute the absence of precise overlap to two factors: 1), Conformational heterogeneity in the non- β strand segment of fibrillized $A\beta_{1-40}$, as suggested by the relatively broad carbonyl ¹³C NMR lines for G25 and G29 discussed above. If a distribution of ϕ and ψ values were significantly populated, one would expect χ^2_{\min} for different techniques to shift in different directions, reducing or possibly eliminating the overlap; 2), Necessary approximations in the simulations, including the use of average CSA principal values for carbonyl 13C pairs with unresolved NMR signals, the assumption of a standard CSA tensor orientation for all carbonyl sites, and the neglect of intermolecular ¹³C-¹³C couplings (see Materials and Methods). Again, one would expect these approximations to produce different shifts in χ^2_{\min} for different techniques.

To extract estimates of ϕ and ψ from the χ^2 contour plots, the following procedure was therefore adopted: 1), A range of allowed ϕ values was determined from the fpRFDR-CT contour plot, given by the condition $\chi^2 < 10$; 2), The χ^2

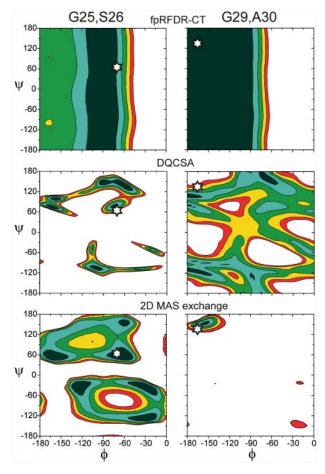


FIGURE 4 Contour plots of the χ^2 deviation between experimental data and simulations for two of the doubly labeled $A\beta_{1-40}$ fibril samples (see Materials and Methods for precise definition of χ^2). For fpRFDR-CT measurements, dark blue regions indicate $\chi^2 < 5$, white regions indicate $\chi^2 > 25$, and contour levels increase in units of 5. For DQCSA measurements, dark blue regions indicate $\chi^2 < 70$, white regions indicate $\chi^2 > 150$, and contour levels increase in units of 20. For 2D MAS exchange measurements, dark blue regions indicate $\chi^2 < 40$, white regions indicate $\chi^2 > 120$, and contour levels increase in units of 20. White stars indicate the estimated ϕ and ψ values for S26 (*left side*) and A30 (*right side*), obtained from a consensus fit to the three measurements (see text).

minima in the DQCSA and 2D MAS exchange contour plots that lie within this ϕ range and were closest together in the ϕ , ψ plane were identified; 3), The average of the ϕ and ψ values at these two minima was taken as an estimate of the correct values. Estimates of ϕ and ψ determined with this procedure are summarized in Table 2. Backbone torsion angles for G25, S26, and G29 are found to be significantly different from β -strand values, whereas torsion angles for V24 and A30 are consistent with a β -strand conformation. These results are sensible in light of evidence from ¹³C NMR chemical shifts for β -strand segments formed by residues 12-24 and 30-36 (Petkova et al., 2002). Uncertainty in ϕ and ψ due to conformational heterogeneity, suggested by the 1D ¹³C NMR spectra as discussed above, cannot be determined quantitatively from the data obtained to date.

For reasons of symmetry (Tycko et al., 1996), fpRFDR-CT, DQCSA, and 2D MAS exchange data are invariant to the substitution $\phi, \psi \to -\phi, -\psi$. Therefore, $\chi^2(\phi, \psi) = \chi^2(-\phi, -\psi)$ and only negative values of ϕ are plotted in Fig. 4. For V24 and A30, the choice of signs for ϕ and ψ in Table 2 is appropriate for a β -strand conformation. For G25, S26, and G29, the sign choices in Table 2 were selected after molecular modeling (see below). Solid-state NMR data in this paper do not rule out the opposite sign choices for G25, S26, and G29.

DISCUSSION

Molecular modeling

The data reported above do not permit a unique determination of the backbone structure of residues 24-30 in $A\beta_{1-40}$ fibrils, both because the signs of ϕ and ψ are indeterminate and because these data do not constrain ϕ and ψ of N27 and K28. To permit visualization of the structural consequences of the non- β -strand conformations at G25, S26, and G29, Fig. 5 shows structural models for residues 20-33, assuming typical β -strand ϕ and ψ values for residues 20-23, 27, 28, and 31-33. ¹³C chemical shift data support a β -strand conformation at K28 (Petkova et al., 2002). Values for V24, G25, S26, G29, and A30 are taken from Table 2. Eight models are shown, generated with all possible sign choices for ϕ and ψ of G25, S26, and G29. No energy minimization or other optimization of these models was performed.

Although all models in Fig. 5 are equally consistent with data in this paper, they are not all consistent with constraints imposed by supramolecular structure. ¹³C chemical shift data have established that the segments flanking residues 24-30 form β -strands in $A\beta_{1-40}$ fibrils (Petkova et al., 2002). Multiple quantum ¹³C NMR (Antzutkin et al., 2000) and ¹³C-¹³C dipolar recoupling data (Balbach et al., 2002) have established that these two β -strands form separate parallel β -sheets through intermolecular hydrogen bonding. To be compatible with a cross- β fibril structure, the two β -strands must be aligned so that all intermolecular hydrogen bonds are nearly parallel to a single axis (namely, the long axis of the fibril). Thus, all backbone carbonyl C-O bonds and amide N-H bonds in the β -strands must be nearly parallel to one

TABLE 2 Backbone torsion angles (degrees) in $A\beta_{1-40}$ fibrils, estimated from solid-state NMR data on doubly labeled samples as described in the text

Residue	ϕ,ψ
V24	-145,115
G25	-70,-40
S26	68,-65
G29	-120, -125
A30	-165,133

Data are invariant to the substitution $\phi, \psi \to -\phi, -\psi$. Choice of signs here corresponds to the model in Fig. 5 a.

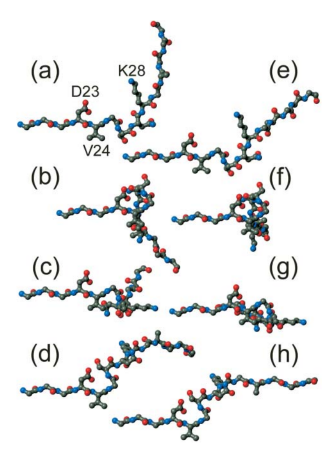


FIGURE 5 Possible conformations for residues 20-33 of $A\beta_{1-40}$ in amyloid fibrils, based on the ϕ and ψ values for V24, G25, S26, G29, and A30 estimated from solid-state NMR measurements. Typical β -strand values ($\phi = -140^\circ$, $\psi = 140^\circ$) are assigned to all other residues. The eight models represent all possible sign choices for ϕ and ψ of the non- β -strand residues G25, S26, and G29, given that the NMR data are invariant to the substitution ϕ , $\psi \rightarrow -\phi$, $-\psi$. Only nonhydrogen atoms are shown. Side chains of residues 20–22 and 31–33 are omitted for clarity. To facilitate comparison, all models are shown with residues 20–24 at the same orientation. Side-chain conformations are not constrained by the data. (a) Conformation resulting from the values and sign choices in Table 2. (b) Sign reversal for G25. (c) Sign reversal for S26. (d) Sign reversal for G25 and S26. (e) Sign reversal for G29. (f) Sign reversal for G25, S26, and G29. (g) Sign reversal for S26 and G29. (h) Sign reversal for G25, S26, and G29. (Figure created with MOLMOL (Koradi et al., 1996))

another. Only the models in Fig. 5, a and h, satisfy this requirement. Of these two, only the model in Fig. 5 a produces the net bend in the direction of the peptide backbone required by the experimentally observed diameters of $A\beta_{1-40}$ fibrils (see Introduction). For these reasons, the conformation in Fig. 5 a was used as a starting point for the recent development of a full structural model for $A\beta_{1-40}$ fibrils by constrained energy minimization (Petkova et al., 2002). In addition, the backbone conformation in Fig. 5 a permits close contacts between the side-chain carboxylate and amino groups of D23 and K28, as supported by experimental measurements of $^{15}N_{-}^{13}C$ dipole-dipole couplings that indicate internuclear distances on the order of 0.4 nm for these groups (Petkova et al., 2002).

Comparison with earlier structural models

Several models for full-length A β fibrils have included a β hairpin with a turn in residues 25-28 (George and Howlett, 1999; Li et al., 1999) or 24-27 (Lazo and Downing, 1998). A true β -hairpin requires intramolecular hydrogen bonding between β -strands that flank the turn. However, ${}^{13}\text{C}$ - ${}^{13}\text{C}$ dipolar recoupling data indicate intermolecular distances of 0.48 ± 0.05 nm (as required by intermolecular hydrogen bonding) between carbonyl or β -carbon sites of V12, L17, F20, A21, V24, A30, L34, and V39 in $A\beta_{1-40}$ fibrils (Balbach et al., 2002). Multiple quantum ¹³C NMR data have shown that the β -carbons of both A21 and A30 must form groups of at least four with interatomic distances of ~ 0.5 nm (Antzutkin et al., 2000). These data rule out a true β -hairpin with intramolecular hydrogen bonding. Thus, we believe that the non- β -strand conformations in residues 24-30 simply create a bend or hinge between two separate parallel β -sheets in $A\beta_{1-40}$ fibrils (Petkova et al., 2002), reminiscent of the bends between β -strands in β -helical proteins (Yoder et al., 1993; Emsley et al., 1996) (although our data do not imply that A β fibrils necessarily have the β -helical structure suggested for amyloid fibrils by other groups (Lazo and Downing, 1998; Perutz et al., 2002; Wille et al., 2002)). A similar structural role for these residues occurs in A β fibril models proposed by Tjernberg et al. (1999) and by Ma and Nussinov (2002).

Concluding remarks

One might expect the presence of a non- β -strand segment in the middle of the $A\beta_{1-40}$ sequence to have implications for the effects of amino acid substitutions on fibril structure or stability. Although a number of mutagenesis studies have been reported, most of these deal with effects of substitutions outside the region addressed by our measurements (Fraser et al., 1992; Hilbich et al., 1992; Pike et al., 1995; Esler et al., 1996; Murakami et al., 2002) or with amyloid fibrils formed by $A\beta$ fragments that may not be representative of the fulllength peptide (Kirschner et al., 1987; Fraser et al., 1994; Wood et al., 1995). A recent study by Morimoto et al. (2002) has shown that substitutions of proline at residues 24 and 26 in $A\beta_{1-42}$ suppress aggregation and fibrillization, consistent with the fact that proline cannot adopt the ϕ and ψ angles given for V24 and S26 in Table 2. Mutations associated with familial Alzheimer's disease, including A21G, E22Q, E22K, E22G, and D23N, occur near the non- β -strand segment identified by our measurements. It is not yet clear whether kinetic, structural, thermodynamic, or purely biological factors lead to the association of these mutations with genetic predisposition to Alzheimer's disease.

Finally, the solid-state NMR data reported above demonstrate in a quantitative and site-specific manner that amyloid fibrils have structural complexity at the molecular level beyond the simple requirement of a cross- β motif.

Determination of the full molecular structures of amyloid fibrils entirely from experimental data remains an important and difficult challenge. Nonetheless, we believe that this challenge can be met through a combination of solid-state NMR measurements that probe supramolecular organization, identify β -strand segments, characterize the conformation of non- β -strand segments, and establish intermolecular and intramolecular contacts, with assistance from EM and other physical techniques.

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