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Intermolecular Alignment in β_2 -Microglobulin Amyloid Fibrils

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Abstract: The deposition of amyloid-like fibrils, composed primarily of the 99-residue protein β_2 -microglobulin (β_2 m), is one of the characteristic symptoms of dialysis-related amyloidosis. Fibrils formed in vitro at low pH and low salt concentration share many properties with the disease related fibrils and have been extensively studied by a number of biochemical and biophysical methods. These fibrils contain a significant β -sheet core and have a complex cryoEM electron density profile. Here, we investigate the intrasheet arrangement of the fibrils by means of ¹⁵N-¹³C MAS NMR correlation spectroscopy. We utilize a fibril sample grown from a 50:50 mixture of ${}^{15}N$, ${}^{12}C$ - and ${}^{14}N$, ${}^{13}C$ -labeled β_2m monomers, the latter prepared using 2-13C glycerol as the carbon source. Together with the use of ZF-TEDOR mixing, this sample allowed us to observe intermolecular ¹⁵N-¹³C backbone-tobackbone contacts with excellent resolution and good sensitivity. The results are consistent with a parallel, in-register arrangement of the protein subunits in the fibrils and suggest that a significant structural reorganization occurs from the native to the fibril state.

 β_2 -Microglobulin (β_2 m) is a 99-residue protein that forms amyloid fibril deposits associated with dialysis-related amyloidosis (DRA).¹ Under acidic conditions (pH = 2.5) and low salt concentration, the protein can also form amyloid fibrils in vitro through a nucleation-dependent mechanism.^{2,3} These fibrils are long, straight, and unbranched in appearance (Figure S1) and share many properties with the fibrils isolated from tissues of DRA patients. including the same characteristic amide I' band in FTIR spectra.⁴ It has been shown that the fibrils themselves, and not the prefibrillar oligomeric species formed in the lag phase of assembly, can disrupt model membranes and are toxic to cells.⁵ While an atomic structural model for these fibrils is not yet available, structural details emerged first through methods like limited proteolysis,^{6,7} hydrogen exchange,^{8,9} and more recently by magic angle spinning (MAS) NMR,10 electron paramagnetic resonance (EPR),¹¹ and cryo-electron microscopy (cryoEM).¹² In particular, analysis of the chemical shifts of 64 assigned residues of β_2 m fibrils has shown that the protein contains a rigid fibril core with substantially more β -sheet character than the native protein.¹⁰ CryoEM maps revealed a complex picture of the fibrils, where non-native globular β_2 m monomers pack in "dimer-of-dimers" building blocks that associate asymmetrically into crescent-shaped units.¹² In addition, site-directed EPR spin labeling suggested that the major building block consists of six β_2 m polypeptide chains, arranged in a parallel, in-register manner.11

In the experiments described here, we investigate the tertiary structure of β_2 m amyloid fibrils with ¹⁵N-¹³C MAS NMR correla-



Figure 1. (a) ¹³C CP spectrum of mixed $2-\beta_2$ m fibrils, 512 scans; (b) ZF-TEDOR spectrum obtained with $\tau_{mix} = 1.76$ ms, 512 scans;, (c) ZF-TEDOR with $\tau_{mix} = 18$ ms, 5120 scans.

tion spectroscopy. MAS NMR has been successfully used to obtain information about the inter- and intramolecular interactions that form the β -sheet core of amyloid fibrils, including $A\beta(1-40)$,¹³ a 22-residue fragment of β_2 m,¹⁴ Het-s(218–289),¹⁵ and curli amyloid.¹⁶ Various sample preparation techniques and experiments have been employed to achieve that end, including methods that rely on the incorporation of single labels^{17,18} or proton-mediated transfer.¹⁹ Here, we use ZF-TEDOR (z-filtered transferred echo double resonance) mixing^{20,21} to obtain intermolecular ¹⁵N–¹³C correlations that establish that the protein subunits in long, straight β_2 m fibrils formed at pH 2.5 are arranged as parallel, in-register β -sheets.

Our experiments utilize fibrils formed from a 50:50 mixture of ¹⁵N,¹²C- and ¹⁴N,¹³C- labeled β_{2m} monomers, the latter half being prepared using [2-¹³C]-glycerol as the carbon source. This sample, referred to as "mixed 2- β_{2} m", offers improved resolution in the ¹³C dimension^{22–24} (Figure 1a) as well as potential gains in experimental transfer efficiency due to the significantly reduced number of directly bonded ¹³C atoms.²⁵ The absence of ¹³C *J*-couplings and the elimination of strong (intramolecular) dipolar ¹⁵N–¹³C couplings as a result of the mixed nature of the sample improve the efficiency of ZF-TEDOR.^{20,26,27}

In a 100% uniformly ¹⁵N,¹³C labeled β_2 m sample, the experimental one-bond ¹⁵N–¹³C transfer efficiency after 1.76 ms of ZF-TEDOR mixing is typically ~20% of the ¹³C CP signal. In the mixed 2- β_2 m sample, after such a short mixing time, no significant buildup of ¹³C polarization is observed, as shown in Figure 1b. This is due to the absence of ¹³C nuclei in the ¹⁵N,¹²C-labeled monomers, which were prepared using ¹³C-depleted glucose (99.9% purity) to eliminate contributions from natural abundance. In particular, signals from one-bond ¹⁵N–¹³C interactions are not detected. On the other hand, longer ZF-TEDOR mixing times lead to the buildup of ¹³C intensity, which reaches a maximum at 18 ms (Figure 1c and Figure S2) and is consistent with ¹⁵N–¹³C

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Figure 2. Comparison of the ${}^{15}N{-}{}^{13}C\alpha$ region of correlation spectra obtained with ZF-TEDOR mixing for two differently labeled β_2m fibril samples. (a) $2{-}\beta_2m$, $\tau_{mix} = 1.6$ ms, 12 mg of sample, 2 days of experimental time. Labels correspond to intramolecular N_i-C α_i transfer, unless otherwise noted, while labels in gray denote cross-peaks that appear only in the $2{-}\beta_2m$ spectrum. (b) Mixed $2{-}\beta_2m$, $\tau_{mix} = 16$ ms, 16 mg of sample, 9 days of experimental time. Labels correspond to intermolecular N_i-C α_i or N_i-C α_{i-1} transfer, unless otherwise noted.

distances of ~5.0–5.5 Å. The maximum bulk transfer efficiency for the C α region is ~3%, which is better than the experimental transfer efficiencies observed for uniformly ¹³C labeled samples (<1% for similar distances).²⁰

In order to obtain site-specific information regarding the origin of the ¹⁵N-¹³C intermolecular contacts in mixed 2- β_2 m, we recorded a 2D ZF-TEDOR experiment with $\tau_{mix} = 16$ ms (Figure S3). This spectrum presents excellent resolution (13 C line widths \sim 50 Hz) and sufficient sensitivity after a long acquisition period, which was facilitated by the robustness of the TEDOR sequence. Overall, the positions of the observed cross-peaks in this mixed $2-\beta_2 m$ spectrum correspond exactly with the positions of cross-peaks in one-bond (Figure 2a) or two-bond TEDOR spectra (data not shown) of a β_2 m fibril sample prepared from 100% ¹⁵N, 2-¹³C glycerol labeled material $(2-\beta_2 m)$. The majority of the cross-peaks in the mixed $2-\beta_2$ m sample could be readily assigned based on known chemical shifts of long, straight β_2 m fibrils,¹⁰ and they correspond exclusively to intermolecular $N_i - C\alpha_i$, $N_i - C\alpha_{i-1}$, $N_i - CO_i$, or $N_i - CO_{i-1}$ transfer (Figures 2b and S3). In particular, the following residues giving rise to intermolecular contacts in the mixed $2-\beta_2 m$ sample were assigned: H31-S33, N42, G43, R45, I46, V49, H51-F62, P72, T73, and Y78-V82. While P32, S33, G43, F56, S57, K58, and F62 are part of well-ordered loops in the fibrils, the remainder of the residues represent all of the currently assigned fibril β -strands.

Some cross-peaks in Figure 2b (mixed $2-\beta_2$ m) do not presently have assignments. Conversely, not all of the strong cross-peaks shown in Figure 2a ($2-\beta_2$ m) appear in the mixed $2-\beta_2$ m spectrum. This includes G18, G29, E44, H84, and V85 (shown in gray in Figure 2a) among others. This is most likely due to differences in



Figure 3. (a) Crystal structure of native monomeric β_2 m (PDB ID: 1DUZ)²⁸ showing the antiparallel β -sheet arrangement of the strands (labeled A to G). (b) Residues that form β -strands in fibrillar β_2 m painted onto the native fold. β -Strands in the fibrils¹⁰ are shown as thick tubes, and the residues giving rise to assigned intermolecular N_i-C α_i cross-peaks are shown in black. The structures were prepared using the Chimera software.²⁹

local dynamics and relaxation whose effects are exacerbated at long mixing times, resulting in large variations in the cross-peak intensities.³⁰

The data presented above suggest that long, straight β_2 m fibrils grown at pH 2.5 and low salt concentration form parallel, in-register β -sheets. In such a case the average distances for intermolecular N_i-C α_i and N_i-C α_{i-1} contacts are ~5 and ~5.5 Å respectively (Figure S4), which is consistent with the bulk ZF-TEDOR buildup (Figure S2). In order to accommodate such an arrangement, substantial reorganization of the native antiparallel β -sheet structure³¹⁻³³ is required, indicating that the structure of the monomers within the fibrils must be highly non-native. Figure 3 highlights two clear pieces of evidence for the non-native structure of β_2 m within fibrils: first, residues involved in loops/turns in native β_2 m (Figure 3a) reorganize to form ordered β -strands in the fibrils (Figures 3b and S5), and second, while all β -strands form antiparallel β -sheet contacts with residues distant in sequence in native β_2 m, β -strands in the fibrils are parallel and in register.

The parallel arrangement of the β -strands in β_2 m fibrils was predicted initially by FTIR experiments^{34,35} and is in agreement with data obtained by site-directed spin labeling and EPR.¹¹ The results described here verify and expand upon the latter, which indicates that spin labels attached to cysteine-substituted residues S33, S55, S61, and T73 among others give EPR spectra indicative of immobile, parallel, and in-register stacked spin labels (Figure S5). Stacks of six β_2 m monomers arranged in that manner are then required to fulfill the electron density maps obtained by cryoEM.¹² The site-specific information regarding the intermolecular arrangement of β_2 m fibrils presented here provides an important step toward a full molecular model of the fibrils. Additional experiments, particularly aimed at determining the quaternary fold of the fibrils, are in progress and should shed light on how this tertiary fibril arrangement fits into such a complex cryoEM electron density profile.

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Supporting Information Available: Sample and experimental details; EM image of the fibrils; 1D ZF-TEDOR buildup; full 2D ZF-TEDOR spectrum; expected intermolecular distances in a parallel, inregister arrangement; summary of the available sequence-specific structural information for the fibrils. This material is available free of charge via the Internet at http://pubs.acs.org.

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