Lipid membrane templates the ordering and induces the fibrillogenesis of Alzheimer’s disease amyloid-β peptide

Eva Y. Chi,1 Canay Ege,1 Amy Winans,1 Jaroslaw Majewski,2 Guohui Wu,1 Kristian Kjaer,3,4 and Ka Yee C. Lee1*

1 Department of Chemistry, Institute for Biophysical Dynamics, and The James Franck Institute, The University of Chicago, Chicago, IL 60637
2 Manuel Lujan Jr. Neutron Scattering Center, Los Alamos Neutron Science Center, Los Alamos National Laboratory, Los Alamos, NM 87545
3 Max-Planck Institute of Colloids and Interfaces, Am Mühlenberg, Germany
4 Niels Bohr Institute, University of Copenhagen, Copenhagen, Denmark

ABSTRACT

The lipid membrane has been shown to mediate the fibrillogenesis and toxicity of Alzheimer’s disease (AD) amyloid-β (Aβ) peptide. Electrostatic interactions between Aβ40 and the phospholipid headgroup have been found to control the association and insertion of monomeric Aβ into lipid monolayers, where Aβ exhibited enhanced interactions with charged lipids compared with zwitterionic lipids. To elucidate the molecular-scale structural details of Aβ-membrane association, we have used complementary X-ray and neutron scattering techniques (grazing-incidence X-ray diffraction, X-ray reflectivity, and neutron reflectivity) in this study to investigate in situ the association of Aβ with lipid monolayers composed of either the anionic lipid 1,2-dipalmitoyl-sn-glycero-3-[phospho-rac-(1-glycerol)] (DPPG), the zwitterionic lipid 1,2-dipalmitoyl-sn-glycero-3-phosphocholine (DPPC), or the cationic lipid 1,2-dipalmitoyl 3-trimethylammonium propane (DPTAP) at the air-buffer interface. We found that the anionic lipid DPPG uniquely induced crystalline ordering of Aβ at the membrane surface that closely mimicked the β-sheet structure in fibrils, revealing an intriguing templated ordering effect of DPPG on Aβ. Furthermore, incubating Aβ with lipid vesicles containing the anionic lipid 1-palmitoyl-2-oleoyl-sn-glycero-3-[phospho-rac-(1-glycerol)] (POPG) induced the formation of amyloid fibrils, confirming that the templated ordering of Aβ at the membrane surface seeded fibril formation. This study provides a detailed molecular-scale characterization of the early structural fluctuation and assembly events that may trigger the misfolding and aggregation of Aβ in vivo. Our results implicate that the adsorption of Aβ to anionic lipids, which could become exposed to the outer membrane leaflet by cell injury, may serve as an in vivo mechanism of templated-aggregation and drive the pathogenesis of AD.

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Key words: protein aggregation; fibril formation; protein–lipid interactions; protein conformation; lipid monolayer; Alzheimer’s disease; grazing-incidence X-ray diffraction; X-ray reflectivity.

Abbreviations: Aβ, amyloid-β peptide; AD, Alzheimer’s disease; APP, amyloid precursor protein; CMC, critical micelle concentration; d62-DPPG, tail-deuterated DPPG; DLS, dynamic light scattering; DMSO, dimethyl sulfoxide; DMTAP, 1,2-dimyristoyl-3-trimethylammonium-propane; DPPC, 1,2-dipalmitoyl-sn-glycero-3-phosphocholine; DPPG, 1,2-dipalmitoyl-sn-glycero-3-phospho-rac-(1-glycerol); DPTAP, 1,2-dipalmitoyl 3-trimethylammonium propane (DPTAP) at the air-buffer interface. We found that the anionic lipid DPPG uniquely induced crystalline ordering of Aβ at the membrane surface that closely mimicked the β-sheet structure in fibrils, revealing an intriguing templated ordering effect of DPPG on Aβ. Furthermore, incubating Aβ with lipid vesicles containing the anionic lipid 1-palmitoyl-2-oleoyl-sn-glycero-3-[phospho-rac-(1-glycerol)] (POPG) induced the formation of amyloid fibrils, confirming that the templated ordering of Aβ at the membrane surface seeded fibril formation. This study provides a detailed molecular-scale characterization of the early structural fluctuation and assembly events that may trigger the misfolding and aggregation of Aβ in vivo. Our results implicate that the adsorption of Aβ to anionic lipids, which could become exposed to the outer membrane leaflet by cell injury, may serve as an in vivo mechanism of templated-aggregation and drive the pathogenesis of AD.

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Grant sponsor: Alzheimer’s Association; Grant number: IIRG-9901175; Grant sponsor: American Health Assistance Foundation; Grant number: A1999057; Grant sponsor: US Department of Energy; Grant number: W-7405-ENG-36; Grant sponsor: National Institute of Health; Grant number: AG025649; Grant sponsors: National Science Foundation Materials Research and Engineering Centers Programs; Grant number: DMR-0213745; Grant sponsors: Packard Foundation, Beckman Scholars Program; GW acknowledges the support of Burroughs Wellcome Fund Interfaces No. 1001774.

EYC and CE contributed equally to this work.

*Correspondence to: Ka Yee C. Lee, Department of Chemistry, The University of Chicago, 929 E. 57th Street, Chicago, IL 60637. E-mail: kayeelee@uchicago.edu

Received 27 July 2007; Revised 26 September 2007; Accepted 23 October 2007

Published online 10 January 2008 in Wiley InterScience (www.interscience.wiley.com); DOI: 10.1002/prot.21887
PROTEINS

INTRODUCTION

The misfolding, aggregation, and accumulation of the amyloid-β peptide (Aβ) into insoluble deposits have been implicated as the primary cause of neuronal death in Alzheimer’s disease (AD). In addition to AD, more than 20 degenerative disorders affecting either the central nervous system (e.g., Parkinson’s disease, Huntington’s disease, and transmissible prion diseases) or a variety of peripheral tissues (e.g., diabetes, liver cirrhosis, and degenerative eye diseases) share the same general pathology, each arising from the aggregation of an associated protein. Although proteins implicated in these diseases share no structural or sequence homology, one defining feature is that they all aggregate into highly ordered insoluble fibrils containing extensive β-sheet structures. The large body of research performed during the past 10 years has significantly increased our understanding of the pathways involved in Aβ aggregation and the mechanism of cellular toxicity. However, the molecular basis of the early events during the aggregation process and the nature of the structural fluctuations that triggers the misfolding and association of Aβ remain poorly understood.

Aβ is a 40- to 43-amino acid residue peptide derived from the proteolytic processing of the transmembrane amyloid precursor protein (APP) during regular cell metabolism. The peptide is amphiphilic, with a hydrophilic N-terminal region and a hydrophobic C-terminal region that belongs to the transmembrane domain of APP. The assembly of native Aβ into fibrillar deposits is accompanied by substantial changes to the peptide conformation. Under conditions that mimic the cell membrane, where the C-terminus of Aβ resides before it is cleaved from APP, Aβ exhibits a significant amount of α-helical structure. When assembled into fibrils, Aβ adopts the characteristic β-sheet structures found in amyloid fibrils. In physiological buffers at low Aβ concentrations, the peptide is largely a random, flexible chain, with some local structural motifs. Thus, regardless of the conformation the native Aβ adopts, the emergence of cytotoxic Aβ aggregates necessitates the formation of inter- and intra-molecular β-sheets.

In vitro, Aβ fibrillogenesis has been found to proceed through a nucleation-dependent polymerization mechanism characterized by an initial lag phase. Fibril formation occurs readily at concentrations above a “critical micelle concentration” (CMC) that has been found to range from 17.5 to 100 μM after which fibrils nucleate within these micelles. The lag phase, to a significant degree, reflects the energy barrier that must be overcome in order for the otherwise unfolded Aβ to adopt a β-sheet rich conformation and oligomerize into ordered intermediates or protofibrils. In vivo, the concentration of free Aβ in the cerebral spinal fluid is exceedingly low, in the subnanomolar concentration range. Since the Aβ concentration required for homogeneous nucleation differs from the Aβ concentration found in vivo by several orders of magnitude, the onset of Aβ fibrillogenesis in vivo likely proceeds by an alternative mechanism.

There is increasing evidence that the cell membrane plays a central role in mediating Aβ fibrillogenesis. It has been shown that membrane lipids extracted from human brain cells can accelerate Aβ fibril formation and that the composition of lipid membranes controls Aβ-lipid interactions and fibrillogenesis. One characteristic that has been well documented is the effect of lipid headgroup charge on Aβ fibril formation. Compared with zwitterionic lipids, anionic lipids have been shown to enhance Aβ association with and insertion into lipid membranes, inducing the formation of β-sheets, and promoting the formation of Aβ fibrils. The association of Aβ to negatively charged lipids is dominated by electrostatic attraction and the phosphate group on the lipid headgroup has been found to be essential for Aβ binding. In addition to mediating Aβ fibril formation, the interaction of the peptide with cell membranes may also serve as a pathway by which Aβ exerts toxicity. One proposed mechanism for Aβ neurotoxicity is membrane disruption and depolarization mediated by either ion-channel formation or an increase in overall membrane conductance, resulting in the alteration of ion homeostasis and disregulation of neuronal signal transduction, leading to cell death.

To fully understand the effect of lipid headgroup charge on Aβ-membrane interactions, we used lipid monolayers at the air-water interface, which model the outer leaflet of the cell membrane, to probe Aβ interaction with lipid membrane and to characterize Aβ-induced alterations in membrane morphology and lipid packing. In an earlier study, we reported that monomeric Aβ spontaneously inserted into lipid monolayers composed of either the anionic lipid 1,2-dipalmitoyl-sn-glycero-3-phospho-rac-(1-glycerol) (DPPG) or the cationic lipid 1,2-dipalmitoyl-3-trimethylammonium propane (DPTAP) at bilayer-equivalent lipid densities whereas the insertion of Aβ into the zwitterionic lipid 1,2-dipalmitoyl-sn-glycero-3-phosphocholine (DPPC) occurred at surface pressures well below that found in a lipid bilayer. Structures of these different lipids are shown in Figure 1(B). Phospholipids in the cell membrane usually contain one tail that is unsaturated. The saturated lipids were chosen for our monolayer studies because they undergo phase transition to form condensed domains when compressed to 30 mN/m, thus allowing us to use membrane morphology to gauge the extent of lipid-peptide interaction upon the introduction of Aβ in the subphase. Using fluorescence microscopy (FM), we found that the association and insertion of Aβ into lipid monolayers can induce a near complete disruption of monolayer morphology and that in some cases, inserted Aβ colocalized with the disordered region of the lipid monolayer. These results confirmed that the ele-
trostatic interaction between Aβ and the lipid headgroup strongly mediates Aβ-membrane interactions and implicated that charged lipids may play a role in Aβ fibrillogenesis and toxicity.

Although pressure-area measurements and FM imaging of Aβ association with lipid monolayers gave us valuable information on Aβ-lipid interactions and micron-scale monolayer morphology, they offered limited insights regarding the molecular-scale structural details of the protein–lipid films. Our initial attempt to obtain these structural details has led us to apply X-ray scattering techniques to examine Aβ-phospholipid monolayers and our data revealed an intriguing templating effect of the anionic lipid DPPG on Aβ in water. In this current study, we have expanded upon this earlier work by incorporating several complementary surface sensitive X-ray and neutron scattering techniques, including grazing-incidence X-ray diffraction (GIXD), X-ray specular reflectivity (XR), and neutron specular reflectivity (NR), to investigate in situ Aβ-lipid films at the air-subphase interface on either water or the more physiologically relevant solution condition of phosphate buffered saline (PBS, pH 7.4, 10 mM sodium phosphate, 3 mM potassium chloride, and 138 mM sodium chloride). All experiments were carried out at 30°C and an Aβ concentration of 250 nM that is well below its documented CMC to better mimic in vivo Aβ concentration. GIXD measurements provide structural information on the in-plane (i.e., in the plane of the monolayer) crystalline (hence diffracting) portion of the film, while XR measurements yield information about the out-of-plane (z-direction) monolayer structure, laterally averaged over both crystalline and amorphous portions. Additionally, NR experiments were carried out to provide complementary information on out-of-plane structures. Combining these scattering techniques allowed for the first complete, angstrom-scale, characterization of Aβ’s association with different charged lipids.

Our results confirmed that the anionic lipid DPPG uniquely templated the folding and ordering of Aβ into β-sheet crystalline structures at the lipid surface. To evaluate whether these templated structures can trigger the aggregation of Aβ with a more physiological membrane system, the peptide was incubated with lipid vesicles comprised of mono-unsaturated and more fluid lipids. Vesicles were made up of either the zwitterionic lipid 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine (POPC), or POPC containing either 30 mol% of the anionic lipid 1-palmitoyl-2-oleoyl-sn-glycero-3-[phospho-rac-(1-glycerol)] (POPG), or 30 mol% of the cationic lipid 1,2-dimyristoyl-3-trimethylammonium-propane (DMTAP). Charged lipids were used in combination with the neutral lipid POPC because charged lipids alone do not form stable vesicles. Size distributions of the vesicles, the adsorption of Aβ to vesicles, and Aβ fibril formation and morphology were monitored with incubation time using a number of analytical techniques.

**METHODS**

**Materials**

The lipids DPPC, DPPG, DPTAP, palmitic acid (PA), tail-deuterated DPPG (d62-DPPG), POPC, POPG, and DMTAP were purchased from Avanti Polar Lipids (Alabaster, AL). PA and Thioflavin-T (ThT) were purchased from Sigma-Aldrich (St. Louis, MO). PBS was purchased from Invitrogen (Carlsbad, CA). All water used was filtered through a Milli-Q Ultrapure water purification system from Millipore (Bedford, MA).

Aβ40 peptide (amino acid sequence of DAEFRHDS-GYEYHHQKLVFAEDVGSKGAILGLMVGGV) used
for X-ray and neutron scattering experiments was purchased from AnaSpec (San Jose, CA) and used without further purification. Purity was reported to be 95%. The molecular weight of the peptide was confirmed by matrix-assisted laser desorption/ionization (MALDI) mass spectrometry. Aβ was stored at -20°C in lyophilized form until use. Aβ40 used for the incubation experiments was synthesized using 9-fluorenylmethoxycarbonyl chemistry on an Applied Biosystems 433A Peptide Synthesizer (Foster City, CA). Peptides were purified by reverse phase-HPLC (RP-HPLC) on a preparative Zorbax C18 column at 60°C. Peptide sequence was confirmed by a mass spectrometry-based protein identification method where purified Aβ was first digested with trypsin and followed by RP-HPLC and mass spectrometry/mass spectrometry analysis. Peptide mass was confirmed by MALDI mass spectrometry and peptide purity was determined by analytical RP-HPLC to be 90% or higher. Purified Aβ was lyophilized and stored at -20°C until use.

To ensure the complete dissociation of Aβ into monomers, Aβ was first dissolved in dimethyl sulfoxide (DMSO) to yield a stock solution of 5 mg/mL. Importantly for our study, DMSO did not exhibit any surface activity. The stock Aβ solution was subsequently diluted with an appropriate buffer for X-ray scattering experiments or a vesicle solution for incubation experiments.

### X-ray scattering experiments

All X-ray scattering experiments were carried out on the BW1 beam line at the HASYLAB synchrotron source (DESY Deutsches Elektronen-Synchrotron, Hamburg, Germany) with a dedicated liquid surface diffractometer. A temperature-controlled Langmuir trough was mounted on the diffractometer. The trough was equipped with a Wilhelmy plate balance for measuring the surface pressure (σ) and a Teflon barrier for changing the surface area. All experiments were carried out at 30°C. The X-ray beam that illuminates the sample had a wavelength (λ) of ~1.3 Å and a power of ~0.3 mW.

After calibrating the Wilhelmy plate pressure sensor at 30°C, 240 mL of aqueous subphase was decanted into the trough. To damp mechanically excited long wavelength waves on the liquid surface, a glass block was placed in the trough such that the water thickness above the glass block was less than 1 mm. To minimize the adsorption of Aβ to glass surface, the glass block was coated with polyethylene glycol. In between X-ray experiments, the glass block was cleaned by sonication in a dilute soap solution followed by a thorough rinse with Milli-Q water.

DPPC and DPTAP dissolved in chloroform (HPLC grade, Fisher Scientific) and DPPG dissolved in chloroform containing 10% methanol (Fisher Scientific) were spread at the air–water interface. The system was allowed to equilibrate for 15 min to ensure the complete evaporation of the organic solvent, after which the lipid monolayer was compressed to 23 or 30 mN/m. Once the desired surface pressure was reached, pressure was kept constant via a feedback loop. An aliquot of Aβ in a matching solution was then injected into the subphase of the trough using a L-shaped syringe underneath the barrier to avoid disturbing the lipid film [Fig. 1(A)]. Aβ concentration in the subphase was 250 nM for all experiments. After injecting Aβ, trough area was recorded, from which changes in the effective area per lipid molecule (ΔA/σ = (Afinal - Ainitial)/Ainitial) compared with those before Aβ injection was calculated. To prepare lipid monolayers for X-ray measurements, the trough canister was purged for 30–40 min with helium to reduce background scattering from the gas phase and prevent oxidative beam damage during X-ray scans. Oxygen level was kept lower than 1%. To further reduce beam damage, the trough was translated by 0.025 mm in the horizontal direction after every step during GIXD scans and by 2 mm during XR scans.

To maximize the surface sensitivity of GIXD measurements, the incidence angle of the X-ray beam (αi) was 0.85αc, where αc ~ 0.13° is the critical angle for total external reflection.53 The footprint of the incoming X-ray beam on the surface was ~2 × 50 mm². Diffracted intensities were collected using a one-dimensional position-sensitive detector (PSD, OEM-100-M, Braun, Garching, Germany) as a function of the vertical scattering angle and has a measuring window of 0 < 4σ < 1.3 Å⁻¹. A Soller collimator was mounted in front of the PSD, which gave a horizontal resolution of the detector of Δqxy = 0.0084 Å⁻¹. Scattering intensity was measured by scanning over a range of the horizontal scattering vector component qxy and simultaneously recording the entire range of qz of the PSD.

For XR measurements, an additional slit is used to exclude diffusely scattered background around the reflected beam. This slit, together with a scintillation detector having a thin vertical measuring window (with a qz resolution of ~0.1qz, where qz is 0.022 Å⁻¹), is mounted on an elevator situated on a diffractometer arm, which is pivoted around a vertical axis through the sample center. A Soller collimator was not used for XR measurements.

### Neutron reflectivity measurements

NR experiments were carried out on the horizontal reflectometer beamline (NG7) at the National Institute of Standards and Technology in the Center for Neutron Research (Gaithersburg, MD). Using a 4.75Å wavelength neutron beam from a reactor source, the reflectivity of the lipid film as a function of qz values from 0.01 to 0.20 Å⁻¹ was collected. Scanning time for this qz range was 3–4 h. The reflected neutrons were counted using an Ordeala position-sensitive 3He detector (Ordeala, Oak Ridge, TN).
**Aβ incubation with large unilamellar vesicles**

To investigate the effect of the different charged lipids on Aβ fibril formation, the peptide was incubated with vesicles composed of POPC, POPC containing 30 mol% POPG, or POPC containing 30 mol% DMTAP in either water or PBS at 25°C. To prepare lipid vesicles, desired amounts of lipids were first dissolved in chloroform, dried under a stream of nitrogen, and further dried under vacuum overnight. The lipids were then hydrated with water, put through five freeze-thaw cycles, and extruded 19 times using an extruder with 100-nm pore size polycarbonate membranes (Avanti Polar Lipids, Alabaster, AL) to yield large unilamellar vesicles.

Stock Aβ40 dissolved in DMSO was added to each vesicle solution to yield incubation mixtures containing 100 μM Aβ with 1:20 peptide to lipid mole ratio. Samples were then incubated at 25°C without agitation. Control samples of Aβ alone and vesicles alone were also prepared and incubated. At each time point, an aliquot of each sample was taken and the size distribution of vesicles, the amount of soluble Aβ remaining, and the presence of fibrils were determined using a number of analytical methods.

To evaluate changes in vesicle size during incubation with Aβ, size distribution of samples and controls were measured by dynamic light scattering (PD2000DLS Detector, Precision Detectors, Franklin, MA). To assess the extent of soluble Aβ depletion from solution because of adsorption onto vesicles and formation of fibrils, the amount of soluble Aβ remaining in each sample was analyzed by size exclusion-HPLC (SE-HPLC). Samples were first centrifuged at 15,000 rpm for 10 min to remove insoluble materials. Supernatant was then injected onto a BioSep-SEC-S2000 column (Phenomenex, Torrance, CA) and chromatograms were collected as UV absorbance at 280 nm. Peak area in the chromatogram was used to quantify the amounts of soluble Aβ in the samples. The amount of protein present was calculated by dividing the measured peak area by the peak area for an unincubated control sample.

To monitor the formation of Aβ fibrils, ThT fluorescence of incubated samples was measured on a Fluoromax-3 spectrofluorometer (Horiba Jobin Yvon, Edison, NJ). At each time point, 20 μL of a sample incubated in either water or PBS was mixed with 2 mL of 10 μM ThT in either pH 5.5 10 mM sodium acetate with 140 mM sodium chloride or PBS. Fluorescence emission intensity of the sample was immediately recorded for 10–30 min (λ_{ex} = 446 ± 3 nm, λ_{em} = 490 ± 10 nm).24,49 ThT fluorescence of control samples, Aβ alone and vesicles alone, were also collected to enable proper background subtraction.

To image the morphology of Aβ aggregates formed, transmission electron microscopy (TEM) was used. Samples were diluted five times and applied to a glow-discharged carbon-coated support film, washed with water, stained with 1% uranyl acetate for 30 s, and then dried. Micrographs were recorded using a FEI Tecnai F30 scanning transmission electron microscope (FEI Company, Hillsboro, OR).

**RESULTS**

**Analysis and interpretation of X-ray and neutron scattering data**

From XR data, detailed information on the electron density distribution [ρ(z)], in the z-direction of the lipid membrane laterally averaged over the entire film (xy plane) [Fig. 1(A)], can be obtained from modeling the ratio, R/R_p, between the measured reflectivity (R) and Fresnel’s reflectivity (R_p) calculated for a perfectly sharp interface between air and a pure subphase.46,47,54 The simplest, and physically reasonable, model to fit R/R_p is a stack of homogeneous slabs, where each slab represents a chemically distinct moiety of a molecule (e.g., a lipid headgroup) with constant electron density ρ and a characteristic thickness (L).50–52 To account for the vertical roughness, or diffuseness (σ), of the interface between the different slabs, ρ(z) was “smeared” in the z-direction by convolution with a Gaussian function of the root-mean-square with σ. A σ value of ca. 3 Å for most lipid monolayers mainly stems from thermally excited capillary waves on the liquid surface.55,56 Fitting the slab model to R/R_p data yields parameters characterizing the vertical structure of a 2D film—ρ and L of each molecular layer and σ of the inter-layer interface. Because each slab introduces three parameters, fitting can easily become over-parametrized as the number of slabs is increased to model the physical system. A criterion used in our fitting was to produce the best fit with the fewest parameters possible. In the cases where multiple slabs were needed, e.g., lipid head group, lipid tails, adsorbed protein, and inserted protein, as many parameters as possible were fixed with known values to minimize the problem of over-parameterization.

Analogous to XR, NR yields angstrom-level information along the z-direction of a thin film. Instead of the scattering of X-ray by electrons, neutrons are scattered by atomic nuclei.48 Modeling of NR data gives the scattering length density (SLD) distribution of atomic nuclei in the z-direction. NR data were also analyzed using the slab model and three parameters were extrapolated from each slab: SLD, L, and σ.

In GIXD diffracted intensity from an incident X-ray beam that is totally externally reflected is collected over a range of the horizontal scattering vector component |q_{xy}| ≈ (2π/λ) [1 + cos^2(α_ε) − 2 cos(α_ε) cos(2θ_{xy})]^{1/2} and of the vertical scattering component |q_z| ≈ (2π/λ) sin(α_ε), where 2θ_{xy} is the horizontal scattering angle and α_ε is the vertical exit angle. Integrating the scattering intensity I(q_{xy},q_z) over q_z yields Bragg peaks and integrating over
The Bragg peak profile is used to calculate the repeat distances (d-spacings, \(d = 2\pi/q_{xy}\)) for the 2D crystalline structure and the full width at half maximum [FWHM(\(q_{xy}\))] of the Bragg peaks yields the coherence length \(L_{xy} = 0.89 \cdot 2\pi/\text{FWHM}(q_{xy})\), that is, the average size of diffracting crystals. Bragg rod profile can be used to extract important structural information on the molecular crystalline arrangement in a monolayer, such as the tilt angle and tilt direction of lipid tails around the surface normal. Additionally, FWHM of a Bragg rod profile can be used to estimate the length of the molecule that scatters coherently.

**Aβ adsorbed to the air-water interface**

The amphiphilic Aβ40 readily adsorbed to the air–water interface when injected into the aqueous subphase of a Langmuir trough. At 250 nM, Aβ equilibrated to final surface pressures of 12 and 15 mN/m in water and PBS, respectively. The XR data for surface-adsorbed Aβ are shown in Figure 2(A). A one-slab model, the simplest physically reasonable model, was used to fit both data sets; fitting parameters are summarized in Table I. For comparison, the reflectivity of a bare air–water interface with a 3 Å roughness (Fresnel curve) (dotted line) was also plotted. The solid lines represent fitted reflectivity curves. Corresponding electron density profiles of Aβ in water (lower plot) and PBS (upper plot) obtained from one-slab model fitting of XR data (B). Both unsmearred (solid line) and smeared (dotted line) electron density profiles were plotted. Depth at 0 Å corresponds to the air-Aβ interface. XR data and electron density profiles have been vertically offset for clarity.

![Figure 2](image-url)

*Figure 2*  
XR data and fit of Aβ in water (○) and PBS (▲) adsorbed to the air-subphase interface at 30°C (A). For comparison, the reflectivity profile of a bare air-water interface with a 3 Å roughness (Fresnel curve) (dotted line) was also plotted. The solid lines represent fitted reflectivity curves. Corresponding electron density profiles of Aβ in water (lower plot) and PBS (upper plot) obtained from one-slab model fitting of XR data (B). Both unsmearred (solid line) and smeared (dotted line) electron density profiles were plotted. Depth at 0 Å corresponds to the air-Aβ interface. XR data and electron density profiles have been vertically offset for clarity.

The q_{xy} position of a Bragg peak is used to calculate the repeat distances (d-spacings, \(d = 2\pi/q_{xy}\)) for the 2D crystalline structure and the full width at half maximum [FWHM(\(q_{xy}\))] of the Bragg peaks yields the coherence length \(L_{xy} = 0.89 \cdot 2\pi/\text{FWHM}(q_{xy})\), that is, the average size of diffracting crystals. Bragg rod profile can be used to extract important structural information on the molecular crystalline arrangement in a monolayer, such as the tilt angle and tilt direction of lipid tails around the surface normal.

Additionally, FWHM of a Bragg rod profile can be used to estimate the length of the molecule that scatters coherently.
Table I

Fitting Parameters for XR Data of Aβ and Single Component Lipid Films at the Air-Subphase Interface at 30°C

<table>
<thead>
<tr>
<th>Sample</th>
<th>Water subphase</th>
<th>PBS subphase</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Thickness L (Å)</td>
<td>Density ρ water</td>
</tr>
<tr>
<td>Aβ</td>
<td>18.6 ± 1.6</td>
<td>12.6 ± 0.01</td>
</tr>
<tr>
<td>DPPC</td>
<td>23</td>
<td>6.6 ± 0.7</td>
</tr>
<tr>
<td>30</td>
<td>9.1 ± 0.5</td>
<td>1.34 ± 0.03</td>
</tr>
<tr>
<td>30°</td>
<td>9.8 ± 0.2</td>
<td>1.30 ± 0.007</td>
</tr>
<tr>
<td>DPPG</td>
<td>30</td>
<td>9.3 ± 0.4</td>
</tr>
<tr>
<td>DPTAP</td>
<td>30</td>
<td>5.2 ± 1.1</td>
</tr>
</tbody>
</table>

*Electron density normalized by the electron density of water, ρ water = 0.334 eÅ⁻³.

*The first value represents σ_Aβ-water and the second value represents σ_Aβ-air.

*Parameters at this pressure were obtained from a nonlinear least-square regression method.

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**Table Legends:**
- **Water subphase**
  - **Headgroup region**
  - **Tail region**
  - **Thickness L (Å)**
  - **Density ρ water**
  - **σ (Å)**
- **PBS subphase**
  - **Headgroup region**
  - **Tail region**
  - **Thickness L (Å)**
  - **Density ρ water**
  - **σ (Å)**
Aβ association with DPPC monolayers

XR data for DPPC monolayers at 23 and 30 mN/m on both water and PBS were collected; fitting parameters are summarized in Table I. Pure DPPC monolayers on water and PBS were fit with the simplest two-slab model, one slab for the headgroup and one slab for the tail group, with a single interfacial roughness parameter (Table I and Fig. 4).52 During the refinement of the fit, the number of electrons in the tail region was set to 242 (calculated from the chemical structure for DPPC) and the area per molecule was set to be the measured value to reduce the number of fitting parameters. The fitted reflectivity of DPPC on water showed deviations from the measured reflectivity at high $q_z$ values ($q_z/q_c > 26$) [Fig. 4(A), solid line], which could stem from the inhomogeneity of the electron density of the large headgroup [Fig. 1(B)], where a constant $q$ value, or a single slab, to model the headgroup might be inadequate. However, when a three-slab model was used, fitting became over-parameterized. Two additional fitting techniques, nonlinear regression and a model independent technique, were also used to fit the reflectivity data. Neither method significantly improved fitting of the high $q_z$ data. The dashed line in Figure 4(A) shows the best fit to the reflectivity data using a nonlinear least square regression method with a Levenberg-Marquardt minimization algorithm.61 The fit showed improvements in the first minimum and maximum of the reflectivity curve compared with the two-slab model fit; fitting parameters are summarized in Table I. As shown, fitting parameters were close to those obtained from the two-slab model. The improved fit gave smaller errors for both roughness and length parameters.

Aβ did not insert into DPPC monolayers at a bilayer equivalent pressure of 30 mN/m on water or PBS32 and no detectable changes in $R/R_f$ with the addition of Aβ were observed compared with those of the lipid alone (data not shown). At a lower pressure of 23 mN/m, Aβ inserted into the DPPC monolayer, resulting in $\Delta A/A$ of 4 and 6% on water and PBS subphase, respectively (Table III).32 XR profile of DPPC on water after Aβ injection was identical to that of DPPC alone at 23 mN/m (data not shown). On the PBS subphase, however, injection of Aβ induced small changes in the XR profile of DPPC at 23 mN/m [Fig. 4(B)]. Fitting of the XR data yielded two slabs with $q$ and $L$ values nearly identical to that of the pure lipid film and an additional slab with a $q$ that was slightly higher than that of water [Fig. 4(B’), Tables I and III]. These results indicate that a diffuse layer (or low concentration) of Aβ was adsorbed underneath the DPPC monolayer at 23 mN/m and 30°C on PBS.

The Bragg peaks and Bragg rod profiles of DPPC at 30 mN/m on water and PBS are shown in Figure 5(A,B). At 23 mN/m, DPPC was just below its phase transition pressure and no in-plane ordering was detected. At 30 mN/m and 30°C on a water subphase, two Bragg peaks at $q_{xy} = 1.38$ and 1.46 Å$^{-1}$ [Fig. 5(A), filled squares] were resolvable, indicative of a distorted hexagonal (or centered rectangular) unit cell. When compressed to 30 mN/m at 15°C, DPPC has been shown to give rise to three resolvable Bragg peaks, indicative of oblique packing

Figure 3
Bragg peaks (A) and Bragg rod profiles (B) from GIXD data of Aβ in water adsorbed to the air-water interface at 30°C. Solid lines correspond to model fits. Bragg rod profiles for the two Bragg peaks have been vertically offset for clarity.
of acyl chains. At 30°C the d-spacings of the DPPC Bragg peaks in Figure 5(A) were \(d_{11} = 4.56\ \text{Å}\) and \(d_{02} = 4.29\ \text{Å}\), corresponding to a rectangular unit cell with dimensions \(a_1 = 5.39\ \text{Å}, b_1 = 8.58\ \text{Å}\), and \(\gamma = 90°\). In the distorted hexagonal packing representation, the dimensions of the unit cell were \(a_b = b_h = 5.07\ \text{Å}\) and \(\gamma = 116°\), where \(a_b\) and \(b_h\) are the magnitude of the unit cell vector and \(\gamma\) is the angle between the base vectors of the unit cell (Table II). Analysis of the Bragg rod profile [Fig. 5(B), filled squares] showed that the hydrocarbon tails of DPPC were tilted by 27° from the surface normal. The tilt (azimuthal angle) was 19° from nearest neighbor direction of \(a_h + b_t\) (vectors) (Table II). Addition of Aβ resulted in small changes in the unit cell dimensions as evident by the shift towards the smaller \(q_{xy}\) values of the first, broad, Bragg peak [Fig. 5(A), open squares] and significant decreases of the integrated intensities of the diffraction peaks indicated a decrease in the amount of ordered material in the footprint of the beam. On PBS, a third Bragg peak was resolvable for DPPC at 30 mN/m [arrow in Fig. 5(A)]. The three peaks arise from an oblique unit cell, where \(a_h \neq b_t\) and \(\gamma \neq 120°\). With the addition of Aβ, the third peak was no longer discernable [Fig. 5(A), open triangles]. Overall, the presence of Aβ in the subphase did not result in significant changes in the lateral ordering of the DPPC film for all pressures (23 and 30 mN/m) and subphase conditions tested.

**Aβ association with DPPG monolayers**

XR data and fit of DPPG at 30 mN/m and 30°C on water and PBS are shown in Figure 6(A,B); the corresponding \(\rho(z)\) profiles are shown in Figure 6(A', filled circles) and (B', filled triangles). Fitting of XR data on both water and PBS yielded similar \(\rho(z)\) profiles; the headgroup thicknesses on water and PBS were 9.3 and 9.5 Å and the tail group thicknesses on water and PBS were 17.9 and 17.8 Å (Table I), giving rise to total lipid thicknesses on water and PBS of 27.2 and 27.3 Å, respectively. \(\rho/\rho_{\text{water}}\) values of the headgroup and tail group on both subphase conditions were ~1.4 and 0.97, respectively.

Aβ spontaneously inserted into DPPG at 30 mN/m and 30°C on water, with a \(\Delta A/A\) of 14%. On PBS, however, the electrostatic repulsion between the overall negatively charged Aβ and the negatively charged lipid head groups resulted in negligible insertion, with a \(\Delta A/A\) of less than 2% (Table III). It is thus not surprising that with the addition of Aβ, the \(R/R_0\) did not change significantly for DPPG at 30 mN/m on PBS (data not shown). On a water subphase, however, a reproducible shoulder

### Table II

Parameters Obtained from GIXD Data of Aβ, Single Component Lipids, and Aβ-Lipid Films at the Air-Subphase Interface at 30°C

<table>
<thead>
<tr>
<th>In-plane Bragg peaks</th>
<th>Out-of-plane Bragg rods</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample</td>
<td>d-Spacing</td>
</tr>
<tr>
<td></td>
<td>(a, b) (Å)</td>
</tr>
<tr>
<td><strong>Water subphase</strong></td>
<td></td>
</tr>
<tr>
<td>Aβ</td>
<td>12 Peak 1</td>
</tr>
<tr>
<td>DPPC</td>
<td>23 No in-plane diffraction</td>
</tr>
<tr>
<td>DPPG</td>
<td>30 Peak 2</td>
</tr>
<tr>
<td>DPTAP</td>
<td>30 No in-plane diffraction</td>
</tr>
<tr>
<td>Aβ + DPPC</td>
<td>30 No in-plane diffraction</td>
</tr>
<tr>
<td>Aβ + DPPG</td>
<td>30 Peak 1</td>
</tr>
<tr>
<td>Aβ + DPTAP</td>
<td>30 No in-plane diffraction</td>
</tr>
<tr>
<td><strong>PBS subphase</strong></td>
<td></td>
</tr>
<tr>
<td>Aβ</td>
<td>15 No in-plane diffraction</td>
</tr>
<tr>
<td>DPPC</td>
<td>23 No in-plane diffraction</td>
</tr>
<tr>
<td>DPPG</td>
<td>30 No in-plane diffraction</td>
</tr>
<tr>
<td>DPTAP</td>
<td>30 No in-plane diffraction</td>
</tr>
<tr>
<td>Aβ + DPPC</td>
<td>23 No in-plane diffraction</td>
</tr>
<tr>
<td>Aβ + DPPG</td>
<td>30 No in-plane diffraction</td>
</tr>
<tr>
<td>Aβ + DPTAP</td>
<td>30 No in-plane diffraction</td>
</tr>
<tr>
<td>Aβ + DPPG</td>
<td>30 No in-plane diffraction</td>
</tr>
</tbody>
</table>

*Values calculated assuming a hexagonal packing to estimate the area occupied by each Aβ at the air-water interface.

Parameter was fixed during refinement.

For the phospholipids, area per molecule indicates area per one acyl tail.
appeared on the second hump of the reflectivity profile [arrow in Fig. 6(A)]. Fitting $R/R_F$ with a two-, three-, or four-slab model did not yield acceptable fits. Fitting using a five-slab model provided satisfactory results. However, the five-slab model introduced 11 parameters ($\rho$ and $L$ for each slab plus $\sigma$) and the fitting became over-parameterized. To resolve the problem, two constraints were used. The insertion of Aβ resulted in an area increase of 14% per DPPG molecule; we therefore constrained the electron densities of the tail group (slab 5) and headgroup with inserted Aβ (slab 3) to be the weighted averages between DPPG and Aβ. Specifically, $\rho_5/\rho_{\text{water}}$ was set to be 0.83 (a weighted average of $\rho_{\text{tail}}/\rho_{\text{water}} = 0.97$ and $\rho_{\text{air}}/\rho_{\text{water}} = 0$) and $\rho_3/\rho_{\text{water}}$ was set to be 1.40 (a weighted average of $\rho_{\text{head}}/\rho_{\text{water}} = 1.44$ and $\rho_{\text{Aβ}}/\rho_{\text{water}} = 1.26$). The $\rho(z)$ profile obtained using the constraints is shown in Figure 6(A'), along with a schematic depicting the molecular composition of each layer. As shown, in addition to being adsorbed underneath the DPPG monolayer, Aβ also penetrated into the lipid monolayer, with a portion of the peptide associating with the hydrophobic tail groups. The resulting total thickness of 39.6 Å of the Aβ layer (adding slabs 1–4) was roughly two times the thickness of pure Aβ adsorbed to the air–water interface, indicating that multiple layers of Aβ were associated with the DPPG monolayer on water.

The Bragg peaks and corresponding Bragg rod profiles of DPPG in water and PBS at 30 mN/m are shown in Figure 7. Pure DPPG on water showed a single Bragg peak at $q_{\text{xy}} = 1.50 \text{ Å}^{-1}$ with FWHM = 0.013 Å$^{-1}$ [Fig. 7(A), filled circles], indicative of a hexagonal unit cell with a six-fold degeneracy where the three scattering planes of the unit cell had identical $d$-spacings. The lattice dimensions of the ordered lipid tails were $a_h = b_h = 4.83 \text{ Å},$ 

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**Figure 4**

XR data and fit for DPPC on water at 23 mN/m (A). DPPC on PBS at 23 mN/m before (D) and after (A) Aβ addition at 30°C (B). The solid lines represent fitted reflectivity curves from slab models. Dashed line in A represents fitted reflectivity from a nonlinear least-square regression method. Corresponding electron density profiles of DPPC on water (A) and on PBS before and after Aβ addition (B) obtained from fitting XR data. Both unsmoothed (solid line) and smeared (dotted line) electron density profiles are plotted. Schematics of the different molecular slabs that gave rise to the electron density profiles are also included (A' and B'). XR data and electron density profiles have been vertically offset for clarity.
### Table III

**Fitting Parameters for XR Data of Aβ-Lipid Films at the Air-Subphase Interface at 30°C**

<table>
<thead>
<tr>
<th>Subphase/Air Insertion</th>
<th>Thickness $L_i$ (Å)</th>
<th>$\sigma_i$ (Å)</th>
<th>$\rho_i$ (mN/m)</th>
<th>$\Delta\beta_i$</th>
<th>$\Delta z_i$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water subphase</td>
<td>23</td>
<td>1.86 ± 0.01</td>
<td>14.0 ± 0.01</td>
<td>0.92 ± 0.01</td>
<td>0.93 ± 0.00</td>
</tr>
<tr>
<td>Aβ + DPPG</td>
<td>30</td>
<td>1.18 ± 0.00</td>
<td>9.1 ± 0.3</td>
<td>1.75 ± 0.00</td>
<td>1.93 ± 0.00</td>
</tr>
<tr>
<td>Aβ + DPPG</td>
<td>30</td>
<td>1.32 ± 0.01</td>
<td>12.6 ± 0.1</td>
<td>0.56 ± 0.03</td>
<td>0.53 ± 0.04</td>
</tr>
<tr>
<td>PBS subphase</td>
<td>23</td>
<td>2.7 ± 2.0</td>
<td>1.04 ± 0.03</td>
<td>1.29 ± 0.01</td>
<td>0.91 ± 0.01</td>
</tr>
<tr>
<td>Aβ + DPPG</td>
<td>30</td>
<td>3.7 ± 3.3</td>
<td>17.8 ± 0.7</td>
<td>0.36 ± 0.01</td>
<td>0.36 ± 0.01</td>
</tr>
<tr>
<td>Aβ + DPPG</td>
<td>30</td>
<td>11.7 ± 5.0</td>
<td>1.04 ± 0.01</td>
<td>4.7 ± 2.0</td>
<td>0.79 ± 0.02</td>
</tr>
</tbody>
</table>

$\gamma = 120^\circ$, and an area per acyl tail of the molecule ($A = 20.2$ Å² (Table II)). Upon peptide injection, the hexagonal unit cell changed into a distorted hexagonal unit cell, as the single Bragg peak split into two peaks [Fig. 7(A), open circles]. The $q_{xy}$ values of these new peaks were smaller compared with that of the single peak of the pure lipid, indicating an expansion of the unit cell. With inserted Aβ, unit cell dimensions were $a_h = b_h = 4.94$ Å, $\gamma = 118.7^\circ$, and $A = 21.4$ Å². Furthermore, the presence of Aβ gave rise to an additional Bragg peak at $q_{xy} = 1.32$ Å⁻¹ [see arrow in Fig. 7(A)], identical to the location of the β-sheet peak observed from the adsorbed Aβ layer at the air–water interface [Fig. 3(A)]. Moreover, the intensity of the Aβ peak associated with the DPPG monolayer was ~2.5 times higher and the coherence length was roughly five times longer (Table II), indicating that the presence of DPPG greatly enhanced the interfacial folding and ordering of Aβ into protofibrils. Note that no Aβ peak was observed for Aβ associated with either DPPC or DPTAP monolayers. Time-dependent GIXD scans taken during the insertion of Aβ revealed that the loss of the hexagonal lipid peak was accompanied by the appearance and growth of the peptide peak over a period of a few hours (see Fig. 8). The Bragg rod profile of the pure DPPG monolayer showed a minimal tilt of the lipids [Fig. 7(B), filled circles], with a calculated tilt angle of 4°; with the addition of Aβ, the tilt angle increased to 17° [Fig. 7(B), open circles, Table II]. The change in tilt angle was accompanied by decreases in the integrated intensities of the Bragg peaks and significant decreases of the in-plane coherence lengths, indicating a reduction of the ordered phase of the monolayer.

On PBS, the pure DPPG monolayer at 30 mN/m gave rise to two Bragg peaks at $q_{xy} = 1.48$ and 1.49 Å⁻¹, indicative of a distorted hexagonal unit cell [Fig. 7(A), filled triangles]; lattice dimensions of the unit cell were $a_h = b_h = 4.88$ Å, $\gamma = 119^\circ$, and $A = 20.9$ Å² (Table II). The addition of Aβ led to a slight expansion of the unit cell, resulting in lattice dimensions of $a_h = b_h = 4.90$ Å, $\gamma = 119^\circ$, and $A = 21.0$ Å² [Fig. 7(A), open triangles]. However, these changes were very small. Thus, in addition to exhibiting minimal insertion, Aβ had negligible effect on DPPG packing on PBS. No changes were observed with the injection of Aβ on the Bragg rod profile of DPPG in PBS [Fig. 7(B), open triangles].

### Aβ association with DPTAP monolayers

XR data and fit of DPTAP at 30 mN/m and 30°C in water and PBS are shown in Figure 9(A,B). The corresponding $\rho(z)$ profiles are shown in Figure 9(A',B'). A two-slab model was adequate in fitting the XR data for the lipid. Compared with DPPC at 30 mN/m, the headgroup thickness for DPTAP was approximately half the size [Table I, Fig. 1(B)].
**Figure 5**

Bragg peaks (A) and Bragg rod profiles (B) from GIXD data of DPPC on water at 30 mN/m before (●) and after (○) Aβ addition and DPPC on PBS at 30 mN/m before (▲) and after (△) Aβ addition at 30°C. Dashed lines are fits of individual peaks and the solid lines correspond to the overall fit. Data have been vertically offset for clarity.

**Figure 6**

XR data and fit for DPPG on water at 30 mN/m before (●) and after (○) Aβ addition (A) and DPPG on PBS at 30 mN/m (B, ▲) at 30°C. The solid lines represent fitted reflectivity curves. Corresponding electron density profiles of DPPG monolayer on water before and after Aβ addition (A') and on PBS without Aβ (B') obtained fitting XR data. Both unsmeared (solid line) and smeared (dotted line) electron density profiles were plotted. Schematics of the different molecular slabs that gave rise to the electron density profiles were also included (A' and B'). XR data and electron density profiles have been vertically offset for clarity.
The injection of Aβ into the subphase of the DPTAP monolayer on water resulted in a ΔA/A of 35% (Table III). The XR profile for the lipid film could still be fit by two slabs [Fig. 9(A, open circles) and (A', top)]. However, the slab parameters were different from those of the pure lipid. $q_2$ (slab at the air interface) was lower than $q_{\text{tail}}$, the $L_1$ (slab at the water interface) was larger than $L_{\text{head}}$, and the $r$ value was larger than that of pure DPTAP (Tables I and III). These parameters indicated that the lipid tails (slab 2) were at a lower packing density (due to Aβ insertion) than that of pure lipid and the thick slab at the water interface (slab 1) encompassed lipid headgroups and inserted peptides with part of the peptide protruding into the subphase [schematic in Fig. 9(A')]. Because of the similarity between $q_{\text{Aβ}}$ and $q_{\text{head}}$, DPTAP values (Table I), a single slab representing head/\(\alpha\)β and \(\alpha\)β underneath the monolayer yielded the best result in fitting the XR data with a larger $r$, stemming from the more inhomogeneous and diffuse slab.

In PBS, where there was strong charge–charge attraction between the negatively charged Aβ and positively charged lipid headgroups, the addition of Aβ resulted in a ΔA/A of only 4% (Table III). The low insertion resulted from XR data of Aβ associated with the DPTAP monolayer was best-fit with a three-slab model [Fig. 9(B, open triangles) and (B', top)]. Slab 1, which was underneath the DPTAP monolayer and had a $\rho$-value that was slightly higher than $\rho_{\text{water}}$ was composed of a diffuse layer of Aβ. Slabs 2 and 3 were almost identical to those of the pure DPTAP monolayer, representing the headgroup and tail groups, respectively. $\rho_2$ was slightly lower than $\rho_{\text{head}}$ of pure DPTAP.
DPTAP, indicating that a small fraction of the peptide may be inserted into the headgroup region [Fig. 9(B')].

The Bragg peaks and the corresponding Bragg rod profiles of DPTAP on water and PBS are shown in Figure 10(A,B). For both subphase conditions, pure DPTAP gave rise to two Bragg peaks, indicating a distorted hexagonal cell with $a_h = b_h = 5.06 \, \text{Å}$, $\gamma = 116^\circ$, and $A = 23.1 \, \text{Å}^2$ for water, and $a_h = b_h = 5.01 \, \text{Å}$, $\gamma = 117^\circ$, and $A = 22.5 \, \text{Å}^2$ for PBS. The analysis of the Bragg rod profiles showed that the hydrocarbon tails of DPTAP lipids were tilted by $28^\circ$ and $22^\circ$ on water and PBS, respectively.

Contrary to the unit cell expansion found during the association of Aβ to a DPPG monolayer on water, the association of Aβ to DPTAP caused a contraction of the DPTAP unit cell, where the Bragg peak shifted to a higher $q_{xy}$ value [Fig. 10(A), open squares], yielding unit cell dimensions of $a_h = b_h = 4.98 \, \text{Å}$, $\gamma = 118^\circ$, and $A = 21.9 \, \text{Å}^2$ (Table II). A decrease in the total integrated intensities of the Bragg peaks was also observed. Aβ association also significantly reduced the tilt angle of the DPTAP tails, from $28^\circ$ to $18^\circ$. On PBS, the injection of Aβ also caused a contraction of the DPTAP unit cell. However, the change was smaller compared with that observed on water.

**Aβ association with binary DPPC-PA monolayers**

To determine whether the ordered Aβ peak induced by DPPG was a lipid-specific effect, we used binary lipid
monolayers composed of 1:1 (by mole) and 1:2 DPPC and the negatively charged PA to mimic a DPPG monolayer, providing different packing and charge density in the film. The observed Bragg peaks of the two mixtures are shown in Figure 11(A). The 1:1 mixture gave rise to two peaks, indicative of a distorted hexagonal unit cell. With the injection of Aβ, no insertion was observed; the unit cell dimensions of the lipids did not change and no ordered Aβ peak was detected [Fig. 11(A)]. The 1:2 DPPC:PA mixture gave rise to a single Bragg peak, indicative of hexagonal packing. The location of the peak was almost identical to the location of the DPPG peak on water (Table II), signifying that the 1:2 mixture was a good mimic of DPPG in terms of unit cell dimensions. However, the binary lipid mixture did not induce any ordering of Aβ. No significant differences in Bragg rod profiles and the total integrated intensity were observed before and after injecting Aβ for both binary lipid mixtures [Fig. 11(B)].

**Neutron reflectivity**

NR measurements were made for tail-deuterated DPPG (d62-DPPG) monolayer at the air-D2O interface before and after Aβ injection. The experimentally measured reflectivity profiles and fits for d62-DPPG at 30 mN/m and 30°C on D2O before and after peptide injection are shown in Figure 12(A); the corresponding SLD profiles are shown in Figure 12(B). The error bars on the data represent the statistical errors of the measurement and the uncertainty in the \( q_z \) resolution and were nearly constant in the measured range.

A two-slab model yielded an excellent fit of the NR data of d62-DPPG [Fig. 12(A), filled circles], with the first slab representing portions of the lipid headgroup nearest to the subphase and the second slab representing the deuterated tail group and the carboxylic ester groups on the lipid headgroup. The thickness values of the tail group and headgroup were larger and smaller (Tables IV and I), respectively, compared with those obtained from XR because the designation of the slabs was different. For NR, the carboxylic esters exhibit SLD similar to the deuterated tail groups and were included in the tail group slab, whereas for XR, the carboxylic esters were included as a part of the headgroup. The overall thickness of d62-DPPG obtained from NR was 26.5 Å, which is in good agreement with the overall thickness of 27.2 Å obtained from XR.

With the injection of Aβ, the NR of d62-DPPG changed significantly [Fig. 12(A), open circles]. A two-slab model still produced an excellent fit to the data. However, the first slab became a combination of peptide...
adsorbed underneath the monolayer and peptide partially inserted into the monolayer [Fig. 12(B), top]. The SLD of this layer was slightly larger than that of the lipid headgroup (Table IV) because the peptide brought with itself D₂O molecules as it inserted into the headgroup region.⁶³ We note that the thickness obtained for the peptide layer (41.5 Å) was in good agreement with the result from XR measurements [39.6 Å, Fig. 6(B)]. The SLD for the tail group was slightly smaller, by 3%, compared with that of pure lipid, indicating that the insertion of the peptide into the tail region was less than expected based on insertion results. This could be partly due to the differences in the phase behaviors between deuterated and undeuterated lipids,⁶⁴ which in turn can
affect how Aβ interacts with the tail groups. Additionally, the deuterated solvent, D₂O, may also exert some effects on the insertion of Aβ at the lipid interface.

**Effect of lipids on Aβ fibril formation**

To evaluate the effect of Aβ-lipid interactions on the aggregation of Aβ, 100 μM Aβ was incubated with large unilamellar vesicles (100-nm diameter) made up of either the zwitterionic lipid POPC, POPC containing 30 mol% anionic POPG, or POPC containing 30 mol% cationic DMTAP. The monounsaturated lipid POPC and POPG lipids and the 14-carbon acyl chain lipid DMTAP lipid were used instead of their saturated, 16-carbon acyl chain counterparts used in monolayer studies because they are more physiologically relevant. Additionally, these lipids have lower main transition temperatures (Tₘ), so that at the incubation temperature of 25°C, the vesicles were more fluid (Tₘ values of the saturated lipids DPPC, DPPG, and DPTAP are 41, 41 (Avanti Polar Lipids), and 45°C, respectively). Aβ, first dissolved in DMSO at 5 mg/mL, was incubated at 100 μM with each type of lipid vesicles in either water or PBS at 25°C. Control samples of Aβ incubated alone in water and PBS were also prepared. A number of analytical techniques were used to characterize the samples. The amount of Aβ adsorbed to the vesicle surface, vesicle size distribution, formation of fibrils, and the morphology of the incubated species were monitored using SE-HPLC, DLS, ThT binding fluorescence assay, and TEM, respectively.

At 100 μM, Aβ incubated alone in pH 7.4 PBS showed a lag phase of ~4 days, followed by large increases in ThT fluorescence, indicating a rapid fibril growth phase [Fig. 13(A), filled circles]. ThT fluorescence profiles of Aβ incubated with different vesicles, at a peptide to lipid mole ratio of 1:20, exhibited the same kinetics characteristics—a lag phase followed by rapid fibril growth phase. Aβ incubated with POPC or POPC/POPG vesicles yielded ThT fluorescence signal that was similar compared with that of Aβ incubated alone [Fig. 13(A), open

**Table IV**

<table>
<thead>
<tr>
<th>Sample</th>
<th>π (mN/m)</th>
<th>Thickness Lₘ (Å)</th>
<th>ρhᵃ</th>
<th>Thickness Lₜ (Å)</th>
<th>ρt</th>
<th>σ (Å)ᵇ</th>
<th>χ²</th>
</tr>
</thead>
<tbody>
<tr>
<td>d-62 DPPG</td>
<td>30</td>
<td>6.5</td>
<td>3.8 E–06</td>
<td>20</td>
<td>7.2 E–06</td>
<td>3</td>
<td>1.0</td>
</tr>
<tr>
<td>d-62 DPPG+Aβ</td>
<td>30</td>
<td>42</td>
<td>4.6 E–06</td>
<td>19</td>
<td>7.0 E–06</td>
<td>3</td>
<td>1.5</td>
</tr>
</tbody>
</table>

ᵃρ of D₂O is 6.38 E–6.
ᵇRoughness value (σ) was held constant at 3 Å because fitted parameters were not sensitive to small changes in σ values due to a limited qₖ range of 0.2 Å.
circles and open triangles]. Aβ incubated with POPC/DMTAP vesicles exhibited the earliest fibril growth phase [Fig. 13(A)], open inverted triangle]. This was likely due to the strong charge–charge attraction between the negatively charged Aβ at pH 7.4 and the positively charged DMTAP that resulted in the adsorption of Aβ to the vesicles surface, effectively increasing the local Aβ concentration that led to a higher fibril nucleation rate, which was reflected in the shorter lag phase.

Aβ incubated in water exhibited a much longer lag phase—no increases in ThT fluorescence was observed for up to 26 days of incubation [Figs. 13(B) and 16, filled circles]. In the presence of lipid vesicles (1:20 Aβ to lipid mole ratio), ThT signal for Aβ incubated with POPC and POPC/DMTAP vesicles in water remained low during 7 days of incubation [Fig. 13(B)]. No increases in ThT fluorescence were detected from these samples after 22 days of incubation (data not shown). In contrast, Aβ incubated with POPC/POPG vesicles in water [Fig. 13(B), open triangle] showed an immediate increase in ThT fluorescence and the level of fluorescence stayed unchanged during 7 days of incubation. The origin of the ThT fluorescence signal from Aβ incubated with POPC/POPG vesicles, however, was unclear. Since the signal increase was immediate after the addition of Aβ to vesicles, it is unlikely that the signal was due to the binding of ThT to mature fibers. Moreover, the vesicles solution was observed to turn cloudy within 10 min of adding Aβ, indicating that the adsorption of Aβ to vesicle surface had caused the vesicles to agglomerate and precipitate. The aggregation of vesicles induced by the adsorption of a number of amyloid disease-linked peptides had been reported previously.66 Analyzing the sample with SE-HPLC, after the precipitates had been removed by centrifugation, showed that no soluble Aβ was left in the solution (see Fig. 14). Thus, at a protein to lipid ratio of 1:20, all Aβ became associated with the POPC/POPG vesicles. The moderate increase in ThT fluorescence observed during the incubation of Aβ with POPC/POPG vesicles [Fig. 13(B), unfilled triangles], therefore, was likely due to a weak binding of ThT to Aβ adsorbed to vesicle surface, not due to the formation of mature fibrils. Additionally, because all Aβ became associated with vesicles and precipitated out of solution, we were unable to evaluate the effect of POPC/POPG vesicle-associated Aβ on Aβ fibril formation.

To resolve the effect of POPC/POPG vesicles on Aβ fibrillogenesis in water, Aβ (100 μM) was incubated with lower concentrations of lipids, at 1000 and 200 μM, or at protein to lipid mole ratios of 1:10 and 1:2. Within 10 min of adding Aβ to vesicles, size distribution of vesicles measured by DLS broadened and shifted to larger sizes in a lipid concentration-dependent manner (see Fig. 15). SE-HPLC chromatographs of the samples showed that the level of soluble Aβ decreased with increasing lipid concentration (Fig. 14, solid lines), where at a 1:20 Aβ to lipid ratio, no soluble Aβ was present as all Aβ became associated with lipid vesicles. After 11 days of incubation, the amount of soluble Aβ remaining decreased in all samples, including Aβ incubated alone (Fig. 14, dashed lines). ThT

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**Figure 14**
Size exclusion-HPLC chromatographs of Aβ incubated with different levels of POPC vesicles containing 30 mol% POPG after 0 day (solid line) and 11 days (dashed line) of incubation at 25°C in water.

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**Figure 15**
Size distribution of POPC vesicles containing 30 mol% POPG, measured by dynamic light scattering, before (●) and immediately after the addition of Aβ. All samples were prepared in water and Aβ concentration was 100 μM. Lipid concentration increased from 200 (△) to 1000 (□) and 2000 (○) μM, yielding protein to lipid ratios of 1:2, 1:10, and 1:20.
fluorescence signal of Aβ incubated with lower lipid concentrations are shown in Figure 16. As shown, ThT fluorescence of Aβ incubated alone (filled circles) remained low during 26 days of incubation. Fluorescence signal of Aβ incubated with 1000 μM lipids, or 1:10 Aβ to lipid ratio (Fig. 16, open circles), showed an immediate increase in ThT fluorescence that matched the increase observed for 1:20 Aβ to lipid ratio sample [Fig. 13(B), open triangles] followed by no significant increases in ThT fluorescence during 26 days of incubation. As shown in Figure 14, after 11 days of incubation, no soluble Aβ remained for the 1:10 Aβ to lipid ratio sample. Thus, similar to the 1:20 Aβ to lipid ratio sample, adsorption of Aβ to vesicles and the subsequent agglomeration of vesicles were the primary events that took place in the 1:10 Aβ to lipid ratio sample; Aβ fibril formation did not occur to a significant extent with incubation. Aβ incubated with the lowest lipid concentration tested, 200 μM, or 1:2 Aβ to lipid ratio, showed a low level of initial adsorption to vesicles (see Fig. 14) and steady increases in ThT fluorescence with incubation time (Fig. 16, filled inverted triangles). After 26 days of incubation, ThT fluorescence signal was at the same level as those observed during Aβ incubation in PBS [Fig. 13(A)], indicating a near complete conversion of Aβ into fibrils.

TEM images of Aβ incubated alone and Aβ incubated at a 1:2 protein to lipid ratio are shown in Figure 17. Samples were diluted five times, adsorbed on glow-discharged carbon-coated support films, stained with 1% uranyl acetate, and dried before imaging. Aβ alone at the start of incubation did not show significant morphology, except for a few small circular features on the TEM micrograph [Fig. 17(A1)]. These could be small oligomers or artifacts from the drying process during sample preparation. After 4 days of incubation, small protofibril-like structures were observed [Fig. 17(A2)] and after 18 days of incubation, these structures were longer and more uniform in length [Fig. 17(A3)]. These protofibril-like structures likely accounted for the loss of soluble Aβ observed during incubation from SE-HPLC results (see Fig. 14). However, these aggregates did not yield ThT fluorescence characteristic of mature fibrils and did not induce the formation of mature fibrils. Aβ incubated with lipid vesicles with a 1:2 protein to lipid ratio at the beginning of incubation showed individual vesicles that were ~100 nm in diameter. Note that vesicles alone, in the absence of any added Aβ, either did not adsorb to the TEM grid or became completely disrupted during drying (data not shown). Thus the vesicles shown in Figure 17(B1) were those associated with Aβ. After 6 days of incubation, fiber-like structures that extended from the vesicles were observed, concomitant with a small increase in ThT fluorescence (see Fig. 16). After 18 days of incubation, a network of fibrils encompassing vesicles...
was observed [Fig. 17(B3)]. The fibrils exhibited twists that are characteristic of mature amyloid fibrils and yielded strong ThT fluorescence signal (see Fig. 16).

**DISCUSSION**

The principle finding of this study is that the negatively charged lipid DPPG uniquely templates the folding and assembly of native Aβ that is monomeric and unfolded into ∼500 nm β-sheet-like crystalline structures, likely protofibrils, at the lipid membrane-water interface. This finding elucidates the structural dynamics, on the molecular-level, of the early events that may trigger the aggregation of Aβ in vivo. Furthermore, we show that these templated Aβ protofibrils at the lipid membrane interface, formed from a sub-μM concentration and on 1-h time scale, are capable of inducing the aggregation of Aβ into mature fibrils. This study also provides the direct evidence that the association of Aβ with lipid membrane disrupts lipid packing, implicating that membrane disruption by Aβ as a possible toxicity pathway. Details of these findings are discussed below.

**Mode of Aβ association depends on lipid composition and subphase condition**

The molecular scale structural details of Aβ associated with DPPC, DPPG, or DPTAP monolayers at 30°C on water and PBS subphases were resolved in this study using a combination of complementary X-ray and neutron scattering techniques. The degree of Aβ penetration into the lipid membrane, the extent of Aβ association with lipid headgroup vs. tail groups, the effect of Aβ association on lipid packing, and the ordering of Aβ at the lipid interface are illustrated in Figure 18. Two cases were not depicted, Aβ associated with DPPC monolayer on water and Aβ associated with DPPG monolayer on PBS, because neither Aβ association nor insertion was detected and no changes in lipid packing were observed. When associated with a lipid monolayer, Aβ exhibited three different modes of interaction: (1) adsorption underneath the lipid headgroups, (2) insertion into the lipid headgroup region, and (3) penetration into the lipid tail region. Since the lipids tested in this study have identical tail groups, the mode of Aβ interaction thus depends on the lipid headgroup and subphase condition, both of which can modulate the electrostatic interactions between Aβ and the lipid headgroup.

The estimated net charge on Aβ40 based on its amino acid sequence at pH 5.5 (water) and 7.4 (PBS) are 0.2 and −2.8, respectively (Protein Calculator v3.3, The Scripps Research Institute). Thus, the electrostatic contribution to Aβ-lipid interactions ranges from strongly attractive [charge–charge attraction (e.g., Aβ-DPPG on PBS)] to moderately attractive [dipole–charge (e.g., Aβ-DPPG on water) and dipole–dipole (e.g., Aβ-DPPC on water)] to strongly repulsive [charge–charge repulsion (Aβ-DPPG on PBS)]. The effect of these interactions is largely reflected in the extent of Aβ insertion into lipid monolayers compressed to a bilayer-equivalent surface pressure, where attractive interactions led to peptide insertion and repulsive interactions prevented Aβ from inserting into the lipid monolayer (Table III). One case, however, does not follow the trend. The insertion of Aβ into a DPTAP monolayer on PBS, where Aβ and DPTAP exhibit strong charge–charge attraction, results in a ΔA/A of only 4%, compared with a ΔA/A of 35% for Aβ insertion into a DPTAP monolayer on water where Aβ and DPTAP exhibit a weaker charge–dipole interaction. We have hypothesized that the low level of peptide insertion was due to the trapping of the Aβ at the lipid headgroup region from the strong electrostatic interactions. X-ray results from this study provide a direct proof for the hypothesis—the peptide is indeed adsorbed underneath the lipid headgroup [Fig. 18(D)]. As a result, possible stabilization resulting from peptide insertion into the tail region of the lipid, which would give rise to an area increase, did not contribute to the overall Aβ-lipid interaction. In contrast, the weaker dipole–charge interactions between Aβ and lipid headgroup, led to not only adsorption of Aβ to the headgroups, but also insertion into the headgroup region and penetration into the tail groups [Fig. 18(B,C)] that likely stabilizes the hydrophobic C-terminus of the peptide. Moderate interaction with the lipid headgroup thus attracts Aβ to the membrane surface and at the same time, allows additional modes of interaction, such as insertion into the lipid headgroup region and association with lipid tail groups, to occur.

**Aβ disrupts lipid packing**

Aside from understanding the different modes of Aβ association with lipid monolayers, the effect of Aβ on lipid ordering is also elucidated. GIXD data show that Aβ disrupts lipid ordering. For all systems studied, with the exception of DPPC on PBS at 30 mN/m where Aβ shows minimal interactions, the presence of Aβ decreases the integrated intensities and increases the FWHM values of the Bragg peaks compared with those of pure lipid films. Such changes result from reductions in the size and number of coherently scattering lipid domains, indicating overall decreases in lipid ordering. For the DPPG monolayer, the coherence length of the 2D lipid crystals is reduced from 600 Å for the hexagonal phase of the pure lipid to 110 and 250 Å in the two crystallographic directions [(1,0), (0,1)] and (−1,1), respectively, with the addition of Aβ. For the DPTAP monolayer, addition of Aβ results in a decrease of coherent length from 400 to 280 Å in the (−1,1) direction, while coherence length in the [(1,0), (0,1)] direction remained unchanged at 90 Å. The reduction in Bragg peak intensities is also in agree-
ment with FM observations where Aβ insertion reduces the area of the condensed phases of the same lipid films from our earlier study. Disruption of the lipid condensed phase by Aβ insertion is a general phenomenon for all lipids studied, but the peptide exhibits different effects on the unit cell geometry of the lipids. For the DPPG monolayer on water, Aβ causes the unit cell to expand and the lipid tails to tilt when it inserts into the monolayer. However, Aβ insertion into DPTAP monolayer on water and DPPC monolayer on PBS (at 23 mN/m) causes the lipid unit cell to contract. Such opposing effects can be partially explained by the location of the inserted Aβ in these lipid films. From dual probe fluorescence experiments, Aβ is found to associate with the fluid phase in a DPPG monolayer, whereas for DPTAP, although Aβ interacts with both condensed and fluid phases, its association is predominantly with the condensed phase. For the DPPG monolayer, insertion of Aβ caused the condensed domains in the DPPG monolayer on water to disappear over time and concomitantly, a new gray-color phase, intermediate between dark condensed and bright fluid phases, emerged. X-ray results from this study are in agreement with these findings, where Aβ association with DPPG monolayer led to decreases in DPPG ordering that correlated with increases in Aβ ordering (see Fig. 8). These effects are illustrated in

Figure 18
Schematic of the different modes of Aβ association lipid monolayers. All experiments were carried out at 30°C. The length scales shown in the figure are in Å. The shade of the lipid tail groups depicts ordering, where darker shade corresponds to more ordering and lighter shade corresponds to less ordering. The tilt of the lipid molecules with respect to the normal and lipid tail group spacing are also illustrated.
DPPG induced folding and templated assembly of Aβ into protofibrils

Our X-ray and neutron scattering data clearly show that folding and assembly can occur for Aβ at a low concentration of 250 nM on a time scale of 1 h. It is clear that Aβ adsorbs to an air–water interface and orders into 2D crystallites. The packing of the peptide is identical to the inter-β-sheet distance found in amyloid fibrils by NMR68 and X-ray fiber diffraction58 techniques. In vitro, amyloid fibrils need days to form at monomer concentrations orders of magnitude higher than the concentration used in this study. The β-sheet ordering observed at the low peptide concentration and the short time scale observed thus is not expected. In addition, the ordering of Aβ can be enhanced by the negatively charged lipid DPPG. The intensity of the ordered Aβ peak is higher when the peptide is associated with a DPPG monolayer than the intensity of the peak obtained from the peptide adsorbed to a bare air–water interface. Moreover, the FWHM of the Bragg peak of Aβ associated with DPPG is narrower, indicating that the ordered Aβ structures are larger, more than double in this case, compared with Aβ structures formed at the bare air–water interface. The enhanced peptide ordering by a negatively charged lipid may serve as the molecular basis for the early events during Aβ fibril formation in vivo. Physiologically, anionic lipid-induced fibrillogenesis can occur if these lipids, which normally reside only on the inner leaflet of the cell membrane, become exposed to the outer leaflet due to membrane damage (e.g., oxidative stress).69

Additionally, the total thickness of the associated Aβ layer also varied with different lipids. The thickness of Aβ associated with a DPPG monolayer on water is ~40 Å. In contrast, Aβ associated with monolayers composed of the lipids DPPC and DPTAP had thickness ranged from 7.3 to 12.7 Å (Table III). Compared with other lipids, DPPG induced not only lateral folding and ordering of Aβ, but also the formation of multi-layers.

To further understand the templating effect of DPPG on Aβ ordering, we tested Aβ association with binary DPPC/PA monolayers. The goal was to create a DPPG-like monolayer by adding negatively charged PA lipid molecules to the neutral DPPC monolayer. In addition to introducing negative charges to the DPPC monolayer, PA, which has a small headgroup compared with its tail group, can act as a “wedge” molecule to the DPPC, which has a larger headgroup compared with its tail, and help the DPPC molecules to pack more optimally with a smaller tilt of the tail.50 DPPG has similarly sized (in cross-section) head and tail groups, giving rise to a cylindrical molecule that packs well. Two DPPC/PA compositions were tested: 1:1 and 1:2. The first system still has a larger cross-sectional area for the head region and a lower charge distribution than DPPG: one negative charge for every three acyl chains. The second system mimics the DPPG monolayer well, both in tail group packing (Figs. 7 and 11) and charge density (two negative net charges per four acyl chains). However, neither system induced Aβ ordering, indicating that the templating effect of the DPPG monolayer may be related to specific interactions between DPPG and Aβ or that the negative charge on PA is not sufficiently exposed due to the screening effect of the bulky DPPC headgroups.

Vesicles containing POPG induced Aβ fibrillogenesis

The characterization of Aβ-lipid interactions and the elucidation of molecular scale structural details give us insights into the nature, strength, and mode of soluble Aβ interaction with lipid monolayers. The aggregation of Aβ, however, seems to be a necessary step for the pathogenesis of AD. One on-pathway product of Aβ aggregation is insoluble fibrils.1–3 To test the effect of Aβ-lipid interactions on Aβ fibril formation, Aβ was incubated with lipid vesicles composed of POPC, POPC containing 30 mol% POPG, or POPC containing 30 mol% DMTAP. At 100 μM, Aβ exhibited a 4-day lag phase followed by rapid fibril growth in PBS. In contrast, Aβ did not undergo fibrillogenesis for up to 26 days in water. Aβ exhibits minimal interaction with DPPC lipids in monolayer studies in either water or PBS, where no insertion was observed at 30 mN/m.32 The addition of POPC vesicles has no effect on Aβ fibrillogenesis in water and shifts the rapid fibril growth phase 1 day earlier in PBS [Fig. 13(A)]. Aβ exhibits strong electrostatic attraction with DPTAP in PBS and when incubated with POPC/DMTAP vesicles in PBS, the adsorption of Aβ to vesicle surface leads to fibril growth that occurred 2 days earlier than Aβ incubated alone [Fig. 13(A)]. In water, where Aβ associates and inserts into a DPTAP monolayer, incubating Aβ with POPC/DMTAP vesicles does not result in fibril formation. Aβ interacts weakly with DPPG in PBS and incubating with POPC/POPG vesicles does not significantly influence Aβ fibril formation. However, the unique templated β-sheet structures observed on a short time scale (e.g., 1 h) for a low concentration of Aβ at the DPPG monolayer surface on water is found to induce Aβ fibril formation. POPC vesicles containing POPG lipids entirely abolished the lag phase (26 days and longer) and seeded Aβ fibril formation in water. Thus, for all the lipids and solution conditions tested, the only case where the lipid vesicles clearly induced Aβ fibril formation
matched the condition where monomeric Aβ was induced into β-sheet conformation. This study provides the first molecular scale characterization of the early structural fluctuation and assembly events that may trigger the misfolding and aggregation of Aβ in vivo.

ACKNOWLEDGMENTS

We gratefully acknowledge beamtime at the BW1 beam line at the Deutsches Elektronen-Synchrotron in Hamburg, Germany and the NIST Center for Neutron Research. We are grateful to Dr. Sushil Satija for his help while we conducted experiments at the NIST Center for Neutron Research and Steven Danaukas for his assistance in fitting some of the X-ray reflectivity data. We thank Hélène Miller-Auer and Stephen C. Meredith of The University of Chicago for their help with peptide synthesis and Yimei Chen at the Electron Microscopy Facility at The University of Chicago for her help with imaging fibrils. We acknowledge the Biophysical Core Facility at The University of Chicago for the use of the Fluoromax-3 spectrophotometer.

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