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Induced β -Barrel Formation of the Alzheimer's A β 25–35 Oligomers on Carbon Nanotube Surfaces: Implication for Amyloid Fibril Inhibition

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ABSTRACT Recent experimental studies show that carbon nanotubes impact the aggregation process of proteins associated with neurodegenerative diseases. However, the details of molecular interactions between proteins and carbon nanotubes are still not well understood. In this study, we investigate the initial adsorption features and dynamics of the Alzheimer's amyloid- β peptide spanning residues 25–35 (A β 25–35) on a single-walled carbon nanotube (SWNT) surface using fully atomic molecular dynamics simulations (MD) in explicit solvent. The initial configurations of the A β 25–35 peptides consist of two preformed bilayer β -sheets, each with four or five β -strands in parallel or mixed antiparallel-parallel orientations. Our simulations show, for what we believe is the first time, that two disjointed A β 25–35 β -sheets with mixed antiparallel-parallel strands can assemble into β -barrels wrapping the SWNT. In contrast, both simulations of A β 25–35 without SWNT, and simulations of SWNT–A β 25–35 with purely parallel β -strands, lead to disordered aggregates. We find that A β 25–35 β -barrel formation involves at least two steps: i), curving of the A β 25–35 β -sheets as a result of strong hydrophobic interactions with carbon nanotube concomitantly with dehydration of the SWNT-peptide interface; and ii), intersheet backbone hydrogen bond formation with fluctuating intrasheet hydrogen bonds. Detailed analysis of the conversion shows that β -barrel formation on SWNT surface results from the interplay of dehydration and peptide-SWNT/peptide-peptide interactions. Implications of our results on amyloid fibril inhibition are discussed.

INTRODUCTION

Carbon nanotubes have drawn significant attention in the field of nanobiotechnology and nanomedicine due to their potential applications in biosensors (1), drug delivery (2), and templates for biomolecule assembly (3). Self-assembly of peptides is also attracting attention due to its applications in bio-nanomaterial design (4) and its implications in amyloidoses such as Alzheimer's disease (5,6). These diseases involve self-assembly of soluble proteins into insoluble fibrils through a nucleation-dependent polymerization process, during which the slow formation of a nucleus (nucleation phase) is followed by a very rapid elongation phase (7.8). There is experimental evidence that graphite sheets (9,10), collagen fibers (11), dichloromethane (12), nanogels (13), carbon nanotubes (14,15), and fullerenes (16,17) promote (prevent) amyloid formation depending on protein intrinsic stability by decreasing (increasing) the lag time for nucleation, but they all leave the elongation phase invariant, suggesting a surface-controlled nucleation mechanism (12,15,16,18). Although these experimental studies have greatly enhanced our understanding of the impact of surfaces on the kinetics of amyloid formation, the detailed adsorption dynamics remains, however, to be determined. Such a knowledge at an atomic level of detail has still not been explored by classical molecular dynamics (MD) simulations (19,20), recent MD focusing on folding and stability of α -helical peptides within single-walled carbon nanotube (SWNT) (21,22), encapsulation of various peptides into

SWNT (23,24) and the conformational change of the human serum albumin protein for 2 ns on SWNT surfaces (20).

The goal of this work is to study the interaction of the low molecular weight (LMW) oligomers of the Alzheimer's amyloid- β peptide spanning residues 25–35 (A β 25–35) with a SWNT of different diameters. This 11-residue fragment of sequence GSNKGAIIGLM, with a positively charged N-terminus and a hydrophobic C terminus, forms amyloid fibrils in vitro and retains the toxicity of the full-length A β peptide (25,26). Hydrogen/deuterium exchange NMR experiments indicate that A β 25–35 amyloid fibrils display a cross- β core formed from K28 to M35, with residues I31 and I32 being the most protected, and suggest two fibril models with out-of-register antiparallel and in-register parallel β -strands (27).

Because the formation of β -rich LMW oligomers from random states is out of reach using all-atom explicit-solvent MD simulations at physiological temperature (28,29), and the equilibrium ensemble of an octamer consists of various bilayer β -sheets and amorphous states (30–33), we study, as a first step, the interaction between two preformed $A\beta 25-35$ β -sheets and carbon nanotubes. By using allatom MD simulations in explicit solvent, we show that two disjointed A β 25–35 β -sheets with mixed antiparallel-parallel strands, can assemble into β -barrels wrapping the SWNT. In contrast, both simulations of A β 25–35 without SWNT, and simulations of SWNT/Aβ25-35 with purely parallel β -strands, lead to disordered aggregates. The conversion process of two β -sheets into β -barrels is studied in detail and the implications of β -barrel formation on amyloid fibril inhibition are discussed.

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1796 Fu et al.

MATERIALS AND METHODS

Our initial A β 25–35 states consist of two layers, each composed of four or five β -strands with the solid state NMR-derived parallel or mixed antiparallel-parallel out-of-register arrangements (27) and the N- and C-extremities charged for each chain (27). Unless specified and shown in Fig. 1, the two β -sheets are separated by intersheet C_{α} - C_{α} distances of at least 1.73 nm, i.e., free of any intersheet side chain-side chain contacts, are parallel to the central axis of SWNT (SWNT axis or z axis) and characterized by angles θ_1 (between strands d and e) and θ_2 (between strands a and h) varying between 90° and 180°. Note that most initial orientations are centered at 90° so that the system can freely evolve toward 0° and 180°. A series of armchair SWNT (n, n) are constructed with n = 3, 4, and 5 yielding the tube diameters of 0.404, 0.542, and 0.677 nm, respectively. All tube lengths are set to 4.25 nm to provide sufficient surface for A β 25–35 peptides to adsorb. The SWNT-A\beta 25-35 complexes are placed in a dodecahedron box of simple point charge water molecules (34) and the minimum distance between the oligomer and the water box wall is 1.2 nm.

All MD simulations are carried out in the isothermal-isobaric (NPT) ensemble at neutral pH using the GROMACS software package (35) and GROMOS96 force field (36). The solute and solvent are separately coupled to external temperature and pressure baths. The temperature is maintained close to 300 K by weak coupling to an external temperature bath (37) with a coupling constant of 0.1 ps. The pressure is kept constant at 1 bar with a coupling time of 1 ps (37). The SETTLE algorithm (38) is used to constrain the bond lengths and bond angles of simple point charge molecules. The bond lengths of the peptides are constrained by LINCS (39). This allows an MD integration time step of 2 fs. A twin-range cutoff 0.9/ 1.4 nm is used for the nonbonded interactions and a reaction-field correction with dielectric permittivity $\varepsilon = 78$ is used to calculate the long-range electrostatic interactions. Carbon atoms of SWNT are uncharged in accordance with Hummer et al. (40) and the Lennard-Jones parameters for the protein-SWNT and water-SWNT interactions are obtained using the Lorentz-Berthelot rule (41).

Because the amino acids G25-S26-N27 are mostly random during our simulations and are excluded from the cross- β core of the A β 25–35 fibrils (27), we use, unless specified, all amino acids for the potential energy analysis, but only the amino acids from K28 to M35 for all other analysis. These

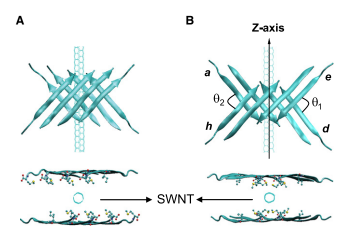


FIGURE 1 Top view and side view of the initial states. Cartoon representation of one of the starting disjointed $A\beta25$ –35 bilayer β -sheets with eight (A) purely parallel and (B) mixed antiparallel-parallel β -strands and (θ_1 and θ_2) values at (98.8°, 100.5°), and line representation of SWNT (the central axis of SWNT is set to be z-axis). *Upper*: side view; *lower*: top view. The side chains of hydrophobic residues facing SWNT, i.e., I31 and M35, are shown by CPK representation in the top view. The θ_1 and θ_2 are the angles between chains e and e0, between chains e1 and e1 are the angles defined by arc cosine of the dot product of the normalized K28-to-M35 (e1 careful in the cosmology of the product of the normalized K28-to-M35 (e2 unit vectors.

include the 2D free energy surface, backbone root mean-square deviation (RMSD) of the $A\beta$ oligomer from the final β -barrel structure, the number of intersheet and intrasheet backbone H-bonds, and the total radius of gyration (Rg) (including side chain and backbone). The hydrophobic residues radius gyration (Rg-HP) uses only the I31 and M35 side chains because the A30, I32, and Leu 34 side chains mostly point to water bulk during the simulations.

In this work, the radius of gyration of A β 25–35 peptides is calculated with respect to the central axis of SWNT, unless specified. One H-bond is considered formed if the distance between N and O is <3.5 Å and the angle of N-H...O is >120° (36). We also calculate the angles θ_1 and θ_2 . An open (closed) β -barrel is considered formed if there are \geq 3 H-bonds between strands e and e3 and e4 and zero (\geq 3) H-bonds between strands e5 and e6 for Ae7. All the systems are displayed using the VMD program (42).

RESULTS

The setup details of 86 MD runs totaling 900 ns starting from two disjointed β -sheets are shown in Table 1. For each system, we give the number of runs and simulation time, the initial values of the angles θ_1 and θ_2 , and the number of MD runs in which a closed or open β -barrel is observed. Although these simulations do not test all possible (θ_1 , θ_2) couples, binding of A β oligomers with initial purely parallel β -strands on SWNT surfaces and interaction of A β oligomers in the absence of SWNT both lead to disordered aggregates within 10–30 ns. By contrast, adsorption of A β octamer and decamer with mixed antiparallel-parallel strands onto different diameter of carbon nanotubes often leads to β -barrel formation, independently of the starting (θ_1 , θ_2) values.

Binding of A β oligomer with purely parallel β -strands on SWNT surface leads to disordered aggregates

The runs (1-8, 54-57, and 63-67) show that neither open nor closed β -barrel forms on SWNT surface starting from parallel β -strand arrangements of A β . This is independent of the dimension of the carbon nanotube (SWNT (3,3), SWNT(4,4)) or the A β oligomer size (octamer and decamer). Looking at the SWNT(4,4)-octamer trajectories, we observe the two β -sheets quickly adsorb on the SWNT (4,4) surface; however the global structure of the octamer is disordered with two to three peptides in fully random coils after 10-30 ns. The A β assembly at 10 ns and the time evolution of the β -sheet content of each peptide in run 1 are shown in Fig. S1 in the Supporting Material. These results indicate that parallel β -strand arrangements do not favor β -barrel formation in a mixed hydrophobic-polar environment, in agreement with our survey of soluble and transmembrane protein structures (see http://www.rcsb.org/pdb/home/).

Adsorption of A β oligomer with mixed antiparallel-parallel β -strands on SWNT surface leads to β -barrel formation

For all $A\beta$ systems with mixed antiparallel-parallel β -strands, we observe the formation of open or closed β -barrels

TABLE 1 Setup details of all simulations

System	MD runs	Simulation time	$\theta_1,\theta_2(^\circ)$
$\overline{\text{SWNT (3,3)} + \text{octamer (P)}}$	run 1-run 8	10 ns × 8 (0)	98.8, 100.5
SWNT $(3,3)$ + octamer $(m A + P)$	run 9-run 53	$10 \text{ ns} \times 35 (9)$	98.8, 100.5
		$10 \text{ ns} \times 5 (1)$	120.6, 118.7
		$10 \text{ ns} \times 5 (1)$	180.0, 180.0
SWNT $(4,4)$ + octamer (P)	run 54-run 57	$10 \text{ ns} \times 4 (0)$	116.7, 115.9
SWNT $(4,4)$ + octamer $(m A + P)$	run 58-run 62	$10 \text{ ns} \times 5 (2)$	116.7, 115.9
SWNT $(4,4)$ + decamer (P)	run 63-run 67	$30 \text{ ns} \times 1 (0)$	117.8, 118.9
		$10 \text{ ns} \times 4 (0)$	
SWNT $(4,4)$ + decamer $(m A + P)$	run 68-run 75	$30 \text{ ns} \times 1 (1)$	117.8, 118.9
		$10 \text{ ns} \times 7 (3)$	
SWNT $(5,5)$ + decamer $(m A + P)$	run 76-run 78	$10 \text{ ns} \times 3 (1)$	117.8, 118.9
Octamer (m A $+$ P) without SWNT	run 79-run 83	$10 \text{ ns} \times 5 (0)$	98.8, 100.5
Decamer (m A + P) without SWNT	run 84-run 86	$10 \text{ ns} \times 3 (0)$	117.8, 118.9

Letters in parenthesis of the first column give the β -strand character (P, parallel; m A + P,: mixed antiparallel-parallel) of A β sheets, and the number in parenthesis of the third column gives the number of MD runs in which a β -barrel is observed. Note that the runs starting from a given (θ_1 , θ_2) pair use different initial velocities, and only the runs 49–53 start from antiparallel β -sheets.

wrapping SWNT in 18 of 61 runs (three representative β -barrels are shown in Fig. 2, A–C). The remaining 43 runs either lead to disordered aggregates or assemblies not satisfying our criteria of open or closed β -barrels. We start the analysis with the simulations of an A β octamer with a SWNT (3,3) surface. Among a total of 45 independent MD runs (runs 9–53) starting from different (θ_1 , θ_2) orientations or the same β -sheet orientation with various random seeds for velocity generation, 11 MD runs (runs 9–19) show a fast conversion of β -sheets into β -barrels.

Using the runs 9–19 of $A\beta$ -octamer-SWNT (3,3), we plot the free energy landscape as a function of the number of intersheet backbone H-bonds and the Rg-HP (Fig. 3). Note that our calculation does not provide the full free energy surface of $A\beta$ -SWNT, because the $A\beta$ peptides are not initially in randomly chosen conformations and orientations. We observe a first delocalized, continuous basin spanning (H-bond, Rg-HP) values of (0, 0.7 nm)–(6, 0.7 nm) and a second smaller basin centered at (11, 0.65 nm). These basins

correspond to compact assemblies and open β -barrels, and closed β -barrels of A β , respectively. The overall "L" shape of the free energy map with the structural description of the minima indicates that β -barrel formation is driven first by strong hydrophobic interactions of $A\beta$ with SWNT and then by intersheet H-bond formation. This is further illustrated in Fig. 4 by the time evolution of three structural properties averaged over all runs 9–19 (Fig. 4, red lines) or in the single run 9 (Fig. 4, black lines): Rg and Rg-HP (Fig. 4 A), and the number of intersheet H-bonds (Fig. 4 B). It is noted that run 19, which starts from θ_1 and $\theta_2 \sim 180^\circ$, is excluded from Fig. 4, E and F. In all simulations, the rapid reduction in Rg and Rg-HP is accompanied with a curving of the two β -sheets (see the snapshot at t = 0.2 ns in Fig. 4 G) and a small decrease of the angles θ_1 and $\theta_2 \sim 85^\circ$ (Fig. 4, E and F). Subsequently, the number of intersheet H-bonds increases and the open β -barrel forms concomitantly with a high plasticity of the two β -sheets (Fig. 4 C). An extreme plasticity is observed in run 9 (Fig. 4 C, black line) where the number of

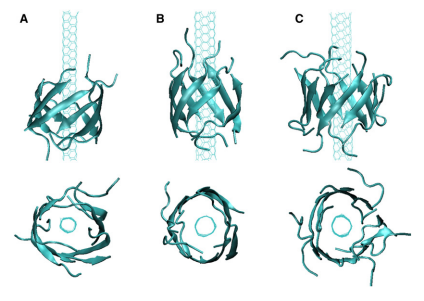


FIGURE 2 Cartoon representation of the MD-generated SWNT- β -barrels. (*Upper*) Side view; (*lower*) top view at t=10 ns for three different SWNT-oligomer systems: (A) SWNT(3,3)-octamer in run 9. (B) SWNT(4,4)-octamer in run 58. (C) SWNT(4,4)-decamer in run 68.

1798 Fu et al.

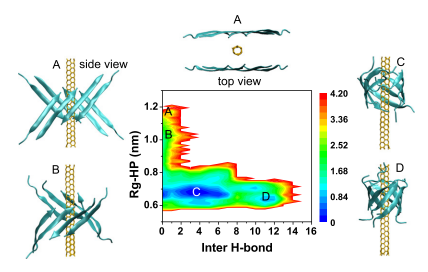


FIGURE 3 Free energy landscape of the $A\beta 25-35$ octamer on a SWNT (3,3) surface along with the structures of SWNT- $A\beta 25-35$ at four different locations. (*A*) One of the initial state (*top view* and *side view*). (*B*) SWNT-curved β -sheet. (*C*) SWNT-open β -barrel. (*D*) SWNT-closed β -barrel. The free energy (in Kcal/mol) is projected onto the number of intersheet backbone H-bonds and Rg-HP. The data are constructed using 11 independent MD runs leading to β -barrel formation.

intrasheet H-bonds drops from 31 to 22 in the first 4 ns (indicating a partial disruption of the initial H-bond network) followed by an increase from 22 to 27 H-bonds. Note that the final H-bond network differs from the initial one in 3 of 11 runs as seen in Fig. 5. Finally, we see a gradual increase of the angles θ_1 and θ_2 up to $140-160^\circ$, indicating an antiparallel arrangement of the newly joined strands, and a constant decrease in the RMSD (Fig. 4 D) as a function of time.

The conversion of two tetrameric β -sheets into a β -barrel on SWNT (3,3) surface can also be monitored by counting as a function of time the number of water molecules within successive cylindrical shells of 0.5 nm height (z direction) and 1.0 nm radius from z axis. In Fig. 6, we show the results of run 9, but a similar picture emerges from runs 10–19. The number of water molecules at the SWNT/A β interface in the region of z=-1.0–1.0 nm is \geq 15 at 0 ns and decreases gradually with time reaching approximately zero at 1.0 ns (Fig. 6 A). The four snapshots at time of 0, 0.24, 0.48, and 1 ns in Fig. 6 B show a clear dehydration process of SWNT-A β interface.

To understand further the physical driving forces underlying the conversion of two tetrameric β -sheets into a β -barrel, we plot the time evolution of the potential energy contribution of SWNT-water, SWNT-A β , A β -water, A β -A β , and waterwater interactions from run 9 (Fig. 7). Given the fact that the simulation timescales are too short to estimate the entropic contributions, our analysis is only qualitative and indicative. We see that the SWNT-A β , A β -A β , and water-water interactions suffice to compensate the energy cost to repel water from the SWNT-A β interface and the energy lost associated with SWNT-water and A β -water interactions. Because simulations without SWNT do not lead to β -barrels (octamer: runs 79–83, decamer: runs 84–86; see Fig. S2), our energy decomposition along with the observation of dehydration at the SWNT(3,3)-A β octamer interface suggest that A β β -barrel formation on SWNT surface results from an interplay of dehydration, $A\beta$ -SWNT, and $A\beta$ -A β interactions.

β -barrel formation with larger A β oligomers and SWNT surfaces

To determine whether β -barrel formation occurs for various SWNT-A β sizes, we carry out MD simulations of an A β octamer with SWNT (4,4) and SWNT (5,5) and simulations of an A β decamer with SWNT (5,5). Although all MD simulations of A β oligomers with parallel strands (octamer: runs 54–57 and decamer: runs 63–67) result in amorphous aggregates, seven among 15 runs with mixed antiparallel-parallel strands lead to β -barrels, indicating a significant effect of β -sheets with mixed antiparallel-parallel strands on the formation of β -barrels. Fig. 8 and Fig. S3 follow the formation of open and closed β -barrels in run 58 of SWNT(4,4)-A β octamer and in run 68 of SWNT(4,4)-A β decamer. The time evolution similarity of Rg and the number of intersheet and intrasheet H-bonds between SWNT(4,4)-A β octamer (Fig. 8), SWNT(4,4)-A β decamer (Fig. S3) and SWNT(3,3)-A β octamer (Fig. 4) is striking. Taken together with the dehydration process in SWNT(3,3)-A β octamer (Fig. 6) and SWNT(4,4)- $A\beta$ decamer (Fig. S4), we can assess that the formation of open/closed β -barrels involves two steps, independently of the SWNT diameter and A β oligomer size: 1), curving of the A β β -sheets as a result of strong hydrophobic interactions of peptides with SWNT and dehydration of the SWNTpeptide interface; and 2), formation of intersheet backbone H-bonds along with the disruption and reformation of intrasheet H-bonds.

DISCUSSION AND CONCLUSIONS

The formation of $A\beta 25-35$ β -barrels from two four- and five-stranded β -sheets with mixed parallel/antiparallel strands is not systematic and we find that 30% of the simulations lead to β -barrel surrounding the carbon nanotube. To explore the impact of the initial configuration of SWNT-A β β -sheets on β -barrel formation, we carry out five additional MD

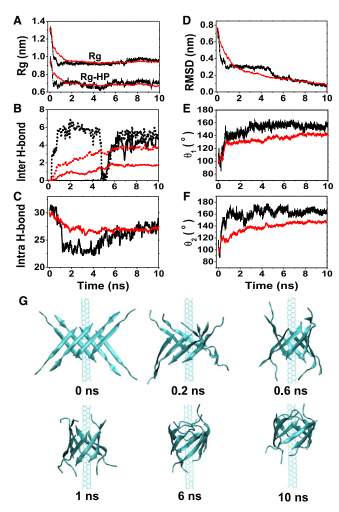


FIGURE 4 Detailed analysis of the assembly process of two tetrameric β -sheets into β -barrel induced by SWNT (3,3) surface. Black curves display run 9, whereas red curves online (or *gray curves* in print) are the averaged properties using runs 9–19 for A–D and runs 9–18 for E and E. Time evolution of: (A) total Rg and Rg-HP of the octamer; (B) number of intersheet backbone H-bonds (*solid line*: between strands e and e; (E) number of intrasheet backbone H-bonds; (E) number of intrasheet backbone H-bonds; (E) backbone RMSD of the octamer with respect to the e-barrel structure generated at 10 ns; (E) angle e₁; and (E) angle e₂. (E) Six representative snapshots of SWNT(3,3)-Ae0 octamer in run 9.

simulations starting from a different orientation of the two β -sheets relative to SWNT-axis (Fig. S5). Again, we observe the formation of β -barrel around SWNT. To determine the stability of the β -barrel, we also extend the MD run 10 to 80 ns. In Fig. S6 we see that all the conformations explored after t=1 ns deviate by <3.0 Å RMSD from the β -barrel formed at 80 ns. Taken together, these results indicate that the β -barrel is likely to be a true free energy minimum for A β 25–35 in the presence of carbon nanotubes. However, replica exchange molecular dynamics are needed to get the β -barrel populations with high accuracy.

Similarly, further studies are needed to understand the dependence of β -barrel formation on SWNT diameter and peptide amino acid sequence. We expect, however, based

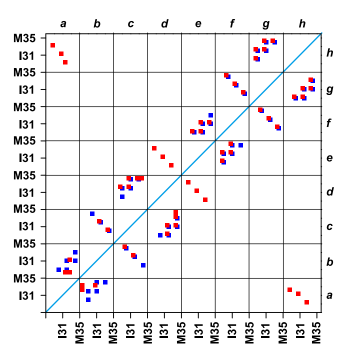


FIGURE 5 Backbone H-bond network map involving residues from K28 to M35 in the structures at t=0 (blue squares online or black squares in print) and t=10 ns (red squares online or gray squares in print) in run 9. The letters a-d are the labels of the four strands in one β -sheet and e-h the labels of the four strands in the other β -sheet. The two hydrophobic residues I31 and M35 in each strand are labeled. The red squares online (or gray squares in print) along the diagonal (from left top to right bottom) are the intersheet H-bond network and the others intrasheet H-bond network.

on our findings that strong hydrophobic interactions of $A\beta$ with SWNT and water expulsion dominate in the early steps of β -barrel formation, and the rate of conversion can be improved by using ideal amino acid patterns with high β -sheet preferences (43). Alternatively, there exists certainly a critical ratio of hydrophobic to hydrophilic, charged to neutral amino acids or unfavorable amino acid patterns, at which the binding of amyloid peptides to the carbon nanotube surface is very weak, therefore hindering the formation of β -barrel.

An important finding of this study is that initial structures composed of parallel β -strands do not form β -barrels in a mixed polar/apolar environment. To understand the microscopic reasons of the orientation preference of the chains, we carry out five runs starting from an idealized barrel containing only parallel β -strands (P-barrel) and five runs starting from the perfect barrel with antiparallel-parallel β -strands generated in run 10 (AP-barrel) at 300 K for 10 ns.

The time evolution of the backbone RMSD, Rg and percentage of initial inner $A\beta$ - $A\beta$ side chain-side chain contacts, and $A\beta$ - $A\beta$ potential energy averaged over the five runs are shown in Fig. 9. Although the RMSD of AP-barrel remains <0.2 nm during 10 ns, the RMSD of P-barrel increases to 0.7 nm after 1 ns (Fig. 9 A). Similarly, the Rg of AP-barrel remains at 0.63 nm, whereas the Rg of P-barrel fluctuates ~1.1 nm and this comes mainly from the rapid increase of the M35 Rg (Fig. 9 B). The percentage of

1800 Fu et al.

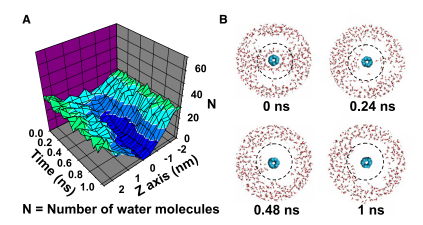


FIGURE 6 Dehydration process of SWNT-A β octamer interface observed in run 9. (A) Time evolution of the number of water molecules within each cylinder along SWNT-axis. The cylinder is 1.0-nm thick from SWNT-axis and 0.5 nm high. (B) Four snapshots of SWNT(3,3)-octamer-water (view parallel to SWNT-axis) generated within the first 1 ns. To visualize the dehydration process clearly, a radius of 1 nm from the SWNT-axis is drawn in dotted line, the A β octamer is omitted and only the water molecules within 2 nm from the SWNT-axis are displayed.

initial inner (M35-M35, N27-N27, I31-I31) contacts in Fig. 3 C decreases to (15%, 30%, and 95%) in P-barrel versus (80%, 90%, and 95%) in AP-barrel, indicating that the geometric constraints imposed by the bulky M35 side chains play a dominant role in the collapse of P-barrel, although the contribution of N27 cannot be ignored. Finally the substantial A β -A β potential energy difference between AP-barrel and P-barrel in Fig. 9 D, which cannot be compensated by the other potential energy terms involving water and SWNT, explains why our simulations do not form P-barrels.

Recent experimental studies showed that the influence of nanoparticles on fibrillation is protein-dependent and varies with particle characteristics. Copolymeric NiPAM/BAM nanoparticles (18) and fullerene (16) retard full-length $A\beta 1$ –40 fibrillation, whereas NiPAM/BAM nanoparticles and carbon nanotubes accelerate the nucleation phase of the 99-residue human β 2m-microglobulin (15). The different effects on fibrillation were attributed to the intrinsic stability of the systems: β 2m-microglobulin is a partially unfolded protein, whereas $A\beta 1$ –40 is mostly unstructured peptide in solution (18). By using surface plasmon resonance experiments, Linse and collaborators (15) could not distinguish between binding of monomeric and oligomeric $A\beta 1$ –40 forms on NiPAM/BAM nanoparticles (18).

Although our simulations do not start from disordered $A\beta$ peptides and do not explore how carbon nanotubes influence the nucleation process, they provide strong evidence that $A\beta$ 25–35 oligomeric species with β -barrel and more disor-

dered characteristics form easily around SWNT. These newly $A\beta$ LMW aggregates therefore block backbone amide sites for further monomer or oligomer addition with β -sheet structure and reduce the population of monomers and small oligomers available for the next steps of aggregation. Whether there are multiple binding sites between full-length peptides and carbon nanotubes remains to be determined, $A\beta1$ –40/1–42 also contains two other hydrophobic cores spanning residues 17–21 and 36–40/42. However, the propensity of the residues 28–35 to form barrels around SWNT is likely to reduce the loop or bent formation of the region Ser²⁶-Ala³⁰ proposed in the $A\beta1$ –42 fibrillar (44) and $A\beta17$ –42 annular (45) models, and this would also enhance the lag time for $A\beta$ nucleation in the presence of SWNT.

The construction of molecules inhibiting $A\beta$ aggregation is a very active field. The designed green fluorescent protein variant P13H with a β -barrel structure was found to inhibit $A\beta$ 1-42 fibrillation (46). Although the interaction sites between $A\beta$ 25–35–SWNT and $A\beta$ 1-42–P13H are different, $A\beta$ 1-42 and $A\beta$ 25–35 both bind to a cylindrical substrate (P13H or SWNT) through noncovalent interactions. Recent studies also showed that fullerene C60 inhibited the fibrillation of $A\beta$ 1-40 (16) and $A\beta$ 25–35 (17), and carbon nanotubes prevented the aggregation of protein (14). Based on the previous experimental studies (14,16,17,27) and the finding of the β -barrel formation of $A\beta$ 25–35 oligomers on SWNT surfaces, we propose that carbon nanotubes are likely to be potent inhibitors of $A\beta$ 25–35 fibril formation. Whether

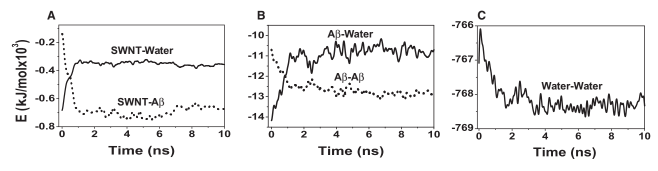


FIGURE 7 Time evolution of the potential energy components in Run 9. (A) SWNT-water and SWNT-A β . (B) A β -water and A β -A β . (C) Water-water.

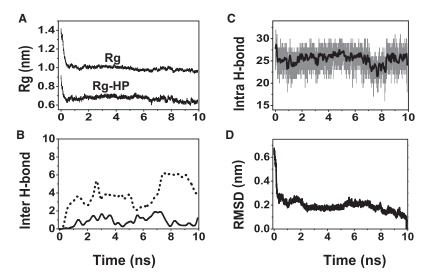


FIGURE 8 Analysis of the assembly process of two tetrameric β -sheets into a 8-stranded β -barrel structure on a SWNT (4,4) surface observed in run 58. Time evolution of: (*A*) total Rg and Rg-HP of the octamer; (*B*) number of intersheet backbone H-bonds; (*C*) number of intrasheet backbone H-bonds (the black curve is a smoothed line over 20 data points); and (*D*) backbone RMSD of the octamer with respect to the β -barrel structure generated at 10 ns.

the A β 25–35–SWNT aggregates and β -barrels are toxic remains to be determined, but a recent study suggests that SWNTs of short lengths <5–10 μ m (i.e., larger than our SWNT length) are not (47).

In summary, we have investigated the interactions of two $A\beta25-35$ oligomeric β -sheets with a SWNT in explicit solvent. Our all-atom MD simulations of multiple SWNT-water- $A\beta$ and water- $A\beta$ oligomer systems show, for what we believe is the first time, that adsorption of $A\beta25-35$ peptides with mixed antiparallel-parallel β -strands on carbon nanotubes can lead to $A\beta25-35$ open/closed β -barrels wrapping the SWNT. Our 2D free energy surface and MD trajectories indicate that β -barrel formation is driven first by strong hydrophobic interactions of $A\beta$ with SWNT simultaneously with water expulsion and then the formation of intersheet hydrogen bonds. Note that this first step was also observed

during the assembly of A β 16–22 protofilaments in solution (48). We also find that the prestructured oligomers display substantial intrasheet H-bond fluctuation during the conversion process, in agreement with simulations on monomer addition to preformed assemblies (49). By contrast, both simulations of A β 25-35 without SWNT, and simulations of SWNT/A β 25–35 with purely parallel β -strands, lead to disordered aggregates. Detailed analysis of the interactions between A β oligomers, water and SWNT show that β -barrel formation on the SWNT surface results from an interplay of dehydration, A β -SWNT, and A β -A β interactions. Collectively, these results help understand the interactions between $A\beta 25-35$ oligomers and SWNT at an atomic level of detail. The formation of β -barrels may offer the prospect of using carbon nanotubes as a new type of therapeutic agent to retard/prevent A β 25–35 fibrillation.

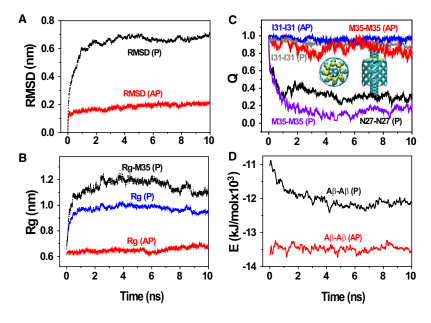


FIGURE 9 Analysis of the destabilization of an ideal β -barrel with purely parallel β -strands (P-barrel) coaxial with a SWNT (3,3) averaged over five independent 10-ns MD runs. Time evolution of: (*A*) backbone RMSD of all residues; (*B*) Rg of the side chains located in the inner side of the β -barrel (i.e., N27, I31, and M35) and Rg of the bulky Met³⁵ side chains; (*C*) fraction of the initial inter-side chain contacts located in the inner side of β -barrel; and (*D*) the A β -A β potential energy. For comparison, the same analyses are reported for mixed antiparallel-parallel β -barrel (AP-barrel). We also show in *C* the top view and side view of the constructed perfect P-barrel with the bulky side chains of M35 in VDW representation.

SUPPORTING MATERIAL

Six figures and a reference are available at http://www.biophysj.org/biophysj/supplemental/S0006-3495(09)01235-1.

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