Aqueous urea solution destabilizes $A\beta_{16-22}$ oligomers

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Edited by Harold A. Scheraga, Cornell University, Ithaca, NY, and approved August 27, 2004 (received for review June 25, 2004)

We use long multiple trajectories generated by molecular dynamics simulations to probe the stability of oligomers of A β_{16-22} (KLVF-FAE) peptides in aqueous urea solution. High concentration of urea promotes the formation of β -strand structures in A β_{16-22} monomers, whereas in water they adopt largely compact random coil structures. The tripeptide system, which forms stable antiparallel B-sheet structure in water, is destabilized in urea solution. The enhancement of β -strand content in the monomers and the disruption of oligomeric structure occur largely by direct interaction of urea with the peptide backbone. Our simulations suggest that the oligomer unbinding dynamics is determined by two opposing effects, namely, by the increased propensity of monomers to form β -strands and the rapid disruption of the oligomers. The qualitative conclusions are affirmed by using two urea models. Because the proposed destabilization mechanism depends largely on hydrogen bond formation between urea and the peptide backbone, we predict that high urea concentration will destabilize oligomers of other amyloidogenic peptides as well.

A n increasing number of diseases are linked to aggregation of proteins and peptides (1). Although proteins implicated in this class of diseases are known, the mechanisms of their aggregation in the amyloid structures with a characteristic cross- β -pattern are not fully understood (2). It is important to characterize the cascade of events in the assembly pathway of disease related proteins because of the suspicion that low molecular weight soluble oligomers and protofibrils are the primary cause of neurotoxicity (3, 4). The finding that a common antibody recognizes mobile oligomers formed from proteins with little or no sequence similarity (5) suggests that there is a limited number of scenarios for their formation (6). Furthermore, the formation of ordered amyloid aggregates appears to be a generic property of all polypeptide chains (7).

Understanding the factors that contribute to the stability and dynamics of oligomers of amyloid β (A β) peptides, which are cleaved in a variety of lengths from the membrane amyloid precursor protein, is necessary for devising methods to block their formation. The interest in A β peptides is associated with their ability to form amyloid oligomers and fibrils, which have long been considered to be the primary pathogenic agents of the Alzheimer's disease (8). The amyloidogenic pathway for $A\beta$ peptides is a complex cascade of molecular events, involving large conformational changes in the monomers, formation of soluble oligomeric intermediates, and gradual accumulation of protofibrils and fibril deposits. The hallmark of amyloid assembly is the emergence of ordered cross β -structure, in which the $A\beta$ peptides are oriented perpendicular to a fibril axis to form long β -sheets. Solid-state NMR experiments have begun to reveal the details of the internal architecture of amyloid fibrils formed by the wild-type A β peptides and their fragments (9–12). These experiments suggest that depending on length and sequence both parallel or antiparallel aligned in-registry organizations of A β peptides are possible. Much less is known about the kinetics of amyloid assembly. Experimental studies suggested that transient formation of α -helical structure may be involved in amyloid assembly (13). This result is striking, because α -helical conformations are absent in both monomer aqueous solution structures (14) and amyloid fibrils (12).

Because of the complexity and the generic nature of amyloid assembly the molecular mechanisms of their formation may be gleaned by the detailed study of aggregation of short peptide fragments (15, 16). Simulations have been used to propose plausible fibril structures of A β peptides of various lengths (17). Fibrillization of fragments of $A\beta$ peptides bear all of the characteristics of amyloidogenesis of the full-length wild-type peptides (11). Molecular dynamics (MD) study of the assembly of $A\beta_{16-22}$ oligomers in explicit water (16) and simulations of the fragment of Sup35 using implicit solvent (18) show that the initial formation of mobile oligomers is driven by side chain hydrophobic and electrostatic interactions. Here, we examine the effect of aqueous urea solution on the stability of $A\beta_{16-22}$ oligomers. We were motivated to undertake this study because interactions of urea with A\beta peptides can elucidate the mechanisms of amyloidogenesis. In addition, certain proteins, such as Ig light chains, assemble into fibrils in the presence of urea in *vivo*. We show that, although A β _{16–22} monomers prefer β-strand conformations in urea, the oligomers are destabilized at high urea concentrations.

Methods

MD Simulations. We performed MD simulations of $A\beta_{16-22}$ (KLVFFAE) monomers and oligomers in aqueous urea solution by using the MOIL package (19) with a protocol similar to that described in ref. 16. The terminals of $A\beta_{16-22}$ peptide are oppositely charged and are capped with neutral acetyl and amide groups. Starting with the hydrated peptide or oligomer system equilibrated at 300 K, we randomly replaced waters with urea molecules by taking into account that, at 8 M urea, the ratio of water to urea molecules is \approx 4.5 and the overall density of the solution is 1.18 g/cm³. Solvent molecules that cause steric clashes were removed. The resulting system was then energy minimized, heated, and equilibrated at 300 K in a (34.7 Å)³ cube with periodic boundary conditions (16).

To test the force field dependence of the simulation results, we used two urea models. The first is the optimized potentials for liquid simulations (OPLS) urea model, which uses AMBER parameterization for covalent interactions (20, 21). The second model is the one developed recently by Weerasinghe and Smith (WS) (22). The WS model differs from OPLS mainly in the values of the partial charges on the urea oxygen, which is increased from -0.390e to -0.675e, and hydrogen, which is decreased from +0.333e to +0.285e. The WS model also uses a Lennard–Jones potential for urea hydrogens with $\sigma=0.158$ nm and $\varepsilon=0.021$ kcal/mol.

Simulation Details. For $A\beta_{16-22}$ monomers, we generated four 10-ns trajectories in 8 M OPLS urea and four 10.5-ns trajectories

This paper was submitted directly (Track II) to the PNAS office.

Abbreviations: $A\beta$, amyloid β ; MD, molecular dynamics; OPLS, optimized potentials for liquid simulations; WS, Weerasinghe and Smith; RC, random coil; ASA, accessible surface area; HB, hydrogen bond; FSS, first solvation shell; DO, disordered oligomers; OO, ordered oligomers.

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for 8 M WS urea. Two independent sets of MD simulations were also performed for $A\beta_{16-22}$ oligomers in 8 M urea solution. Starting with a disordered oligomer conformation (16), which is a precursor to the antiparallel β -sheet structure in $A\beta_{16-22}$ oligomer, we generated four 11-ns trajectories by using the OPLS urea model. The second set of simulations (WS model, four 10.5-ns trajectories) tested the stability of an ordered, antiparallel registry of peptides in the $A\beta_{16-22}$ oligomer. For this set of simulations, we started from the structure that is "close" to the ordered antiparallel conformation of the $A\beta_{16-22}$ trimer (16).

Probes of A β_{16-22} **Structure.** We used a number of quantities to probe urea-induced changes in A β_{16-22} monomers and oligomers (16). In addition to the time-dependent changes in the α -helix and β -strand contents, we also computed the distributions of A β_{16-22} structural states [i.e., fractions of β -strand, α -helix, and random coil (RC) conformations] as described in ref. 16. The integrity of A β_{16-22} oligomers was examined by using the distance between the centers of mass of peptides i and j, RM $_{ij}^{CM}(t)$, and the accessible surface area (ASA) A(t).

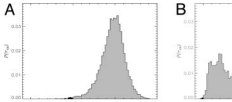
Probes of $A\beta_{16-22}$ Solvation. To probe peptide-solvent hydrogen bonds (HBs) we computed pair correlation functions $g_{\text{H-O}}(r)$, which report the density distribution of solvent along the distance between hydrogen (H) and oxygen (O) atoms. For reference, we used the backbone atoms, which are either carbonyl oxygen O_B, oxygen in Glu side chain O_E, amide hydrogen H_B, or hydrogens in Lys side chain H_K. The solvent atom is either water oxygen O_W and hydrogen H_W or urea oxygen O_U and hydrogen H_U. The pair correlation functions were calculated by averaging over all of the saved conformations and H-O pairs from all trajectories obtained in a given set of simulations. All g(r)functions are normalized to approach unity at $r \to \infty$. The first maximum in g(r) corresponds to the formation of the first solvation shell (FSS) around a backbone atom. Unless indicated otherwise, the FSS refers to an individual backbone atom. We also define a joint FSS, which combines all FSSs of identical atoms in a peptide. Backbone FSS is a union of O_B and H_B joint FSSs. We count only distinct solvent atoms in joint or backbone FSSs.

To verify that the functions g(r) accurately describe HBs, we computed the fractions of FSS solvent molecules that actually make a HB with a peptide. (A HB is formed if the distance between donor D and acceptor A is ≤ 3.5 Å and the angle D-H... A is $\geq 120^{\circ}$.) On average, >90% of FSS solvent molecules are hydrogen bonded to the peptide atoms. These calculations validate the analysis of hydrogen bonds based on g(r).

Results

Urea Enhances *β*-Strand Content in $A\beta_{16-22}$ Monomers. Our previous work showed that, in water, almost 70% of $A\beta_{16-22}$ monomers are in a RC state, whereas the *β*-strand conformations make up 29% (16). To probe the changes in the monomer conformations in 8 M aqueous urea, we generated four 10-ns trajectories. By classifying the peptide conformations as α -helix, β -strand, and RC (16), we find that the fraction of β -strand conformations (0.53) is larger than RC state, whereas the population of α -helix is negligible (0.01). Thus, 8 M aqueous urea promotes β -strand formation at the expense of RC conformations. Simulations of $A\beta_{16-22}$ monomers in 4 M urea also show the enhanced β -strand propensity, which is very similar to that at 8 M. The enhanced β -strand content in $A\beta_{16-22}$ monomers has implications for the stability of oligomers (see below).

Urea-induced changes in peptide conformations are reflected in the distribution of the end-to-end distance $P(r_{1N})$ (Fig. 1). The average of r_{1N} in 8 M urea (14 Å) is larger than $\langle r_{1N} \rangle = 12$ Å in water. More importantly, $P(r_{1N})$ in 8 M urea solution and water are drastically different (Fig. 1). In aqueous urea, $P(r_{1N})$ has a



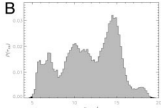


Fig. 1. The distributions of the end-to-end distance $P(r_{1N})$ for $A\beta_{16-22}$ monomer in aqueous OPLS urea (A) and water (B). Shift in $P(r_{1N})$ distribution toward larger r_{1N} indicates that $A\beta_{16-22}$ peptides adopt more extended, open-like structures in urea solution.

single peak at \approx 15 Å. In contrast, there are three maxima in water, two of which (at \approx 10 and 14 Å) contain a mixture of β -strand and RC conformations. The third maximum at \approx 6.5 Å represents only RC states. The broad distribution of $P(r_{\rm IN})$ in water is consistent with the large fraction of RC states and weaker propensity for β -strand structures. The increase in $\langle r_{\rm IN} \rangle$ in urea correlates with the changes in ASA, which grows from 1,159 Ų in water to 1,219 Ų in urea solution. Thus, various measures show that 8 M urea promotes β -strand formation in A β ₁₆₋₂₂ monomers.

Electrostatic Interactions Between Urea and the Peptide Backbone **Drive** β -Strand Formation in $A\beta_{16-22}$ Monomers. To probe the mechanism of enhanced β -strand propensity in 8 M urea solution, we analyzed the interactions between urea and the peptide by using a number of pair correlation functions for selected peptide and solvent atoms (Methods). To this end, we used four 10-ns trajectories for $A\beta_{16-22}$ monomer in 8 M OPLS urea. The interactions between the backbone amide hydrogens H_B and water O_W or urea O_U oxygens analyzed by using the functions $g_{\rm H_B-O_W}(r)$ (black) and $g_{\rm H_B-O_U}(r)$ (gray) show the preference of backbone amide groups to form HBs with urea as compared to water (Fig. 2). The maximum concentration of O_U in the FSS of backbone H_B (i.e., at the distance of r = 1.8 Å between H_B and O_U, which is typical for an optimal HB) exceeds the bulk value by a factor of 3.2, whereas the local water density exceeds the bulk value by only $\approx 30\%$. By integrating g(r) over the FSS, we find that the average number of urea and water molecules in the H_B FSS is 0.43 and 0.58, respectively. Consequently, the ratio of

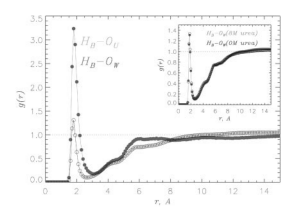


Fig. 2. The pair correlation functions $g_{\rm HB-OU}(r)$ (gray) and $g_{\rm HB-OW}(r)$ (black) probe the formation of HBs between backbone amides and urea or water molecules, respectively. Data are collected over 40,000 conformational snapshots of A $\beta_{\rm 16-22}$ monomers from four 10-ns trajectories in 8 M OPLS urea solution. There is a dramatic preference for backbone amides to form HBs with urea molecules. (*Inset*) The functions $g_{\rm HB-OW}(r)$ obtained for aqueous urea (gray) and water (black). Similarity between $g_{\rm HB-OW}(r)$ plots suggests that urea causes minimal perturbation in local water structure.

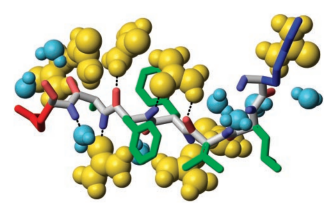


Fig. 3. Illustration of the mechanism of solvation of the $A\beta_{16-22}$ monomer with urea (yellow) and water (light blue). The picture shows solvent molecules in the backbone FSS, which combines H_B (pale blue) and O_B (pale red) FSSs. Lys, Glu, and hydrophobic side chains are colored in blue, red, and green, respectively. Stable HBs between urea and amide groups result in significant increase in urea concentration near peptide backbone. A single urea molecule is shown to cross-bridge amide and carbonyl backbone groups by making two, Ou-HB and H_U–O_B, HBs. Few representative HBs are indicated by dotted lines. Images were produced with MOLMOL (31).

water to urea oxygens in H_B FSS is 1.3, whereas it is 4.5 in the bulk (Methods). Therefore, the relative concentration of urea in H_B FSS increases by factor of 3.3. Urea also solvates backbone carbonyl groups better than water, but the preference is not as dramatic as for backbone amides.

To further probe the preferential solvation of $A\beta_{16-22}$ backbone by urea, we monitored the number of water and urea molecules in the backbone FSS. If backbone groups (amides and carbonyls) do not have a preference to form HBs with urea, then the ratio $\langle N_{\rm fss}^{\rm W} \rangle / \langle N_{\rm fss}^{\rm U} \rangle \lesssim 4.5$, where $\langle N_{\rm fss}^{\rm W} \rangle$ and $\langle N_{\rm fss}^{\rm U} \rangle$ are the average number of water and urea molecules in backbone FSS, respectively. However, the simulations show that $\langle N_{\rm fss}^{\rm W} \rangle / \langle N_{\rm fss}^{\rm U} \rangle \approx$ 1.5, which implies that urea preferentially solvates the peptide backbone. The average number of solvent (water and urea) molecules in the joint H_B FSS is 5.3. Because there are seven H_B in $A\beta_{16-22}$ peptide, approximately one urea and one water molecules make simultaneously two HBs (per solvent molecule) with backbone amides (Fig. 3). Similar computations show that, on an average, one of six urea molecules solvating $A\beta_{16-22}$ backbone cross-bridges amide and carbonyl backbone groups. Interestingly, water is rarely engaged in such amide-carbonyl cross-bridging. Multiple HBs formed between single solvent molecule and $A\beta_{16-22}$ backbone lend additional stability to the expanded peptide conformations.

The solvated $A\beta_{16-22}$ monomer in urea solution is highly mobile with rapid exchange of solvent molecules near the backbone occurring on a picoseconds time scale. The average residence time of water in the H_B FSS is 9 ps, whereas the corresponding lifetime for urea-amide HB is 14 ps. These results are consistent with recent MD simulations of protein denaturation by urea (23). The HB lifetimes give another indication of the enhanced preference of urea over water for the solvation of the backbone of $A\beta_{16-22}$ peptide. Although the total number of urea and water molecules near the peptide's backbone is approximately constant, their individual values undergo large, sometimes highly collective, fluctuations. Over time intervals as short as 1 ns, the number of urea molecules in the joint H_B FSS changed from five to zero, whereas the number of waters increased from one to six.

 β -Strand Propensity Is Not Related to Solvation of Charged and **Hydrophobic Side Chains.** A β_{16-22} peptide has two charged side chains (K and E) and five contiguous hydrophobic residues. To

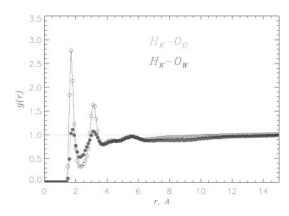


Fig. 4. The pair correlation functions $g_{H_K-O_U}(r)$ (gray) and $g_{H_K-O_W}(r)$ (black) characterize the solvation of positively charged lysine side chain by urea and water, respectively. Water solvates Lys better than urea. An ordered second hydration shell around H_K results in the second peak in g(r). Data were collected as for Fig. 2.

eliminate the possibility that solvation of the side chains by urea increases β -strand propensity, we monitored the solvation of K, E, and the hydrophobic residues. Comparison of pair correlation functions $g_{H_K-O_U}(r)$ and $g_{H_K-O_W}(r)$ shows that the amide group in the lysine side chain is better solvated by water than urea (Fig. 4). The formation of the second solvation shell indicates further structuring of water around Lys side chain. By computing the number of water and urea molecules in the H_K FSS we conclude that the relative (with respect to urea) concentration of water increases 50%. Similar effect is observed for the glutamic acid side chain (data not shown), in which OE oxygens accept most of HBs from water. The relative (with respect to urea hydrogens) concentration of water hydrogens in the O_E FSS increases by nearly 30%. Thus, water solvates charged residues Lys and Glu to a greater extent than urea.

The interactions between urea and the hydrophobic side chains are probed by the pair correlation functions $g_{\text{CM}_{\text{H}}\text{-CM}_{\text{S}}}(r)$ between the centers of mass of hydrophobic side chains CM_H (H = L,V,F,A) and solvent molecules CM_S (S = W,U). There is an enhancement of urea concentration in the vicinity of Phe side chains relative to its bulk value, whereas the concentration of water barely changes (Fig. 5). Qualitatively similar results were obtained for other hydrophobic residues as well. To ascertain

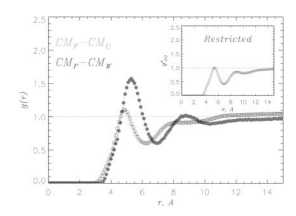


Fig. 5. The pair correlation functions $g_{CM_H-CM_S}(r)$ probe the solvation of Phe hydrophobic side chain. CM_E is the center mass of Phe side chain and S is either water W (black) or urea U (gray). The excess of urea concentration in Phe FSS is entirely due to the HBs between urea and the peptide's backbone. (Inset) When $g_{CM_F-CM_U}(r)$ is restricted to the urea molecules not bound to the backbone, the excess of urea concentration is no longer observed. Data are collected as for Fig. 2.

whether the well formed FSS of urea around Phe side chain is merely a consequence of the HBs between urea and $A\beta_{16-22}$ backbone and not due to favorable interactions with hydrophobic residues, we computed the restricted pair correlation functions $g_{CMH^-CM_S}^R(r)$, which exclude urea molecules interacting with the peptide backbone. In this case, the maximum urea concentration in the Phe FSS barely reaches the bulk value (Fig. 5 *Inset*). Therefore, formation of HBs between urea and peptide backbone automatically increases the concentration of urea around hydrophobic residues. Taken together these results show that the enhanced β -strand propensity in $A\beta_{16-22}$ peptides is caused by direct hydrogen bond formation between urea oxygens and backbone amide hydrogens (21, 24–26).

Water Structure Is Not Perturbed by Urea. Water-backbone HBs are not altered by urea. For example, the functions $g_{\rm H_B-O_W}(r)$ (Fig. 2 Inset) for $A\beta_{16-22}$ monomers in water and aqueous 8 M urea are nearly identical. Although the average number of water molecules in the joint $\rm H_B$ FSS decreases from 4.6 in water to 3.1 in urea solution, the characteristics of peptide backbone hydration by water remain unchanged. Similar observations were made earlier (27). This result is striking, because, on an average, there are 2.2 urea molecules in the vicinity of backbone amide groups.

Electrostatically Induced Enhancement of β -Strand Propensity in Urea Is Generic. To ascertain that the proposed mechanism of increased β -strand propensity is not an artifact of OPLS urea potentials, we used the recently developed WS urea model (22) to generate four 10.5-ns trajectories for $A\beta_{16-22}$ monomers. Comparison of the results using the two models leads to the following conclusions: (i) The urea-induced propensity for β -structure is more pronounced for the WS model. The fraction of peptides in the β -strand state grows from 0.53 (OPLS) to 0.68 (WS), whereas the RC fraction decreases from 0.46 to 0.32. (ii) The solvation of $A\beta_{16-22}$ peptides by WS urea is greatly enhanced. The maximum of the $g_{H_B-O_U}(r)$ function increases by a factor of 2 compared to the OPLS urea model. This results in a 5.5 times increase in the local urea concentration with respect to water in the H_B FSS. The enhanced stability of urea-backbone amide HBs increases the average residence time of urea molecules in the H_B FSS from 14 ps (OPLS) to 25 ps (WS). (iii) As a consequence of the enhancement of solvation of peptide backbone with WS urea, the solvation of hydrophobic residues is also increased. On an average, the WS model adds one urea molecule and eliminates about four water molecules from the joint FSS of hydrophobic residues.

Despite the quantitative differences both models are in qualitative agreement regarding the mechanism of solvation of $A\beta_{16-22}$ monomers in aqueous urea. The difference between WS and OPLS models arises because of the variations in the partial charges on the urea and the treatment of Lennard–Jones interactions for urea hydrogens (*Methods*). The improvement in the solvation of backbone and additional enhancement of β -structure propensity for the WS urea model are exclusively traced to these modifications. Therefore, the main factor in the urea interaction with $A\beta_{16-22}$ monomers is electrostatic interactions, which are responsible for the formation of HBs between urea and peptide backbone. The increase in the β -structure content is a consequence of interactions between backbone amides and urea molecules.

Urea Destabilizes A\beta_{16-22} Oligomers. Our previous work showed that, in water, A β_{16-22} trimers can rapidly form disordered oligomers (DO) (16). DOs are stabilized by favorable interpeptide interactions between hydrophobic residues. Because the DO structures typically do not have the required interpeptide salt bridges, they do not form antiparallel β -sheets. The low energy

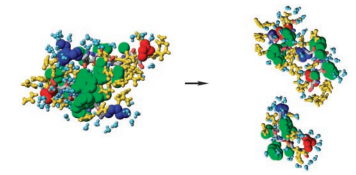


Fig. 6. The disintegration of DO in one of the trajectories for 8 M OPLS urea solution is illustrated by using the same color scheme as in Fig. 3. The intact DO on the left is stabilized by interpeptide hydrophobic interactions. Ten nanoseconds later, DO is completely disrupted, when all interpeptide side chain contacts are broken and urea penetrates between $A\beta_{16-22}$ peptides. Water and urea molecules form FSS around peptides. Images were produced with MOLMOL (31).

structures of $A\beta_{16-22}$ trimers are ordered oligomers (OO), which are stabilized not only by interpeptide hydrophobic contacts, but also by the salt bridges between Lys⁺ and Glu⁻ side chains. In the OO $A\beta_{16-22}$ peptides are arranged in antiparallel β -sheets (16). To probe the mechanism of destabilization of $A\beta_{16-22}$ oligomers in 8 M urea we used both, the DO and a structure, which rapidly converts to OO, as initial conformations.

We used DO initial structure for the $A\beta_{16-22}$ trimer to generate four independent 11-ns trajectories to investigate the effect of 8 M urea solution on $A\beta_{16-22}$ oligomers. There is a heterogeneity in the time scales and pathways of unbinding of peptides from the oligomer. In one of the trajectories the distances between centers of mass $R_{13}^{\rm CM}(t)$ and $R_{12}^{\rm CM}(t)$ increase >2-fold in 11 ns. The growth in $R_{ij}^{\rm CM}(t)$ reflects the separation of the peptide 1 from the oligomer. By the end of the trajectory (>8 ns), the remaining dimer consisting of the peptides 2 and 3 also disintegrates because all interpeptide contacts are disrupted (Fig. 6). As the oligomer breaks apart the ASA grows from $\approx 2,700$ to 3,500 Å² (a 30% increase). Another trajectory shows that the distances $R_{13}^{\rm CM}(t)$ and $R_{23}^{\rm CM}(t)$ more than double after 2 ns because peptide 3 breaks away from the oligomer. Consequently, the ASA grows from $\approx 2,700$ to 3,300 Å². These results are in sharp contrast with the dynamics of $A\beta_{16-22}$ oligomers in water, in which the distances between centers of mass of peptides do not change on the same time scale (16).

The distances between peptides centers of mass, $\langle R_{ij}^{\rm CM}(t) \rangle$, and the ASA, $\langle A(t) \rangle$, averaged over four trajectories indicate that, in 11 ns, $\langle R_{ij}^{\rm CM}(t) \rangle$ increases $\approx 50\%$, whereas $\langle A(t) \rangle$ changes from <2,800 Ų to 3,200 Ų (14% change). In water, the ASA remains approximately constant fluctuating between 2,400 Ų and 2,600 Ų. The orientational order in the A β_{16-22} oligomer is significantly reduced by urea. The variance in the orientational order δd (16) averaged over four trajectories in urea solution is 0.26, which is larger than $\delta d=0.16$ for the DOs in water. These results show that urea dramatically reduces the stability of A β_{16-22} oligomers.

To verify that destabilization of $A\beta_{16-22}$ oligomers in aqueous urea does not depend on the specific urea model or initial conditions, we also generated four trajectories by using the WS urea model. In these simulations the starting structure is the one which favors the formation of the OO in water (16). The OO simulations with WS urea also demonstrate dramatic increases in the average ASA and the distance between peptides centers of mass, which are similar to those observed for DO. In three of four trajectories at least one of the peptides completely breaks away from the oligomer. However, the time scale of OO disruption is larger than for DOs, which is consistent with enhanced

stability of OO. For example, within 10.5 ns, OOs are partially dissolved only in two trajectories, whereas in the third trajectory it takes almost 20 ns for the peptide 3 to separate from the oligomer. By the end of WS simulations most of the interpeptide salt bridges in OO are broken. These results indicate gradual unraveling of the $A\beta_{16-22}$ OOs.

The dynamics of urea-induced unraveling of the OOs is different from the disruption of the DOs. On a relatively short time scale urea transiently stabilizes the OO by promoting the interpeptide Lys–Glu salt bridges. The average probability of finding ordered dimer structures is $P_{\rm d} \approx 0.30$ on the time scale of 10.5 ns. For comparison, $P_{\rm d}$ computed from the simulations of $A\beta_{16-22}$ OO in water, on comparable time scale, is less than 0.01. However, on a larger time scale (exceeding ≈ 20 ns), the OO structures are still disrupted. Thus, we conclude that urea-driven eventual denaturation of $A\beta_{16-22}$ oligomers does not depend on the urea model or initial conditions. However, the pathways of the disintegration of $A\beta_{16-22}$ oligomers critically depend on the starting structures.

Urea Decreases the Lifetime of α-**Helical Intermediate in A** β_{16-22} **Oligomers.**We have previously shown that the dominant pathway during the early stages of the assembly of $A\beta_{16-22}$ DO in water is $RC \to \alpha \to \beta$ transition in which there is a transient accumulation of α-helical intermediate. Because there is a propensity in $A\beta_{16-22}$ monomers to form β-strands in urea, we expect that urea should affect the lifetime and stability of the α-helical intermediate. The α-helix $\langle H(t) \rangle$ and β-strand $\langle S(t) \rangle$ contents averaged over four trajectories for $A\beta_{16-22}$ oligomers in 8 M urea show that $\langle H(t) \rangle$ is initially larger than $\langle S(t) \rangle$. However, within 1 ns, $\langle H(t) \rangle \approx \langle S(t) \rangle$ and, on larger time scales, β-structure content rapidly grows and levels off at ≈ 0.4 , whereas the α-helical content decreases and stays constant at ≈ 0.1 .

These results are qualitatively similar to the assembly dynamics in water (16). However, comparison of $\alpha \to \beta$ transitions in water and 8 M urea demonstrates that urea reduces the amount of transiently accumulated α -helix structure ($\langle H(t=0) \geq 0.4$ in water, but ≤ 0.3 in 8 M urea). More importantly, α -helical intermediate converts into β -strand structure faster in urea. It takes <1 ns for $\langle H(t)\rangle$ to fall below $\langle S(t)\rangle$ in aqueous urea, as opposed to ≈ 5 ns in water (16). Simulations of $A\beta_{16-22}$ OO using the WS urea model are in qualitative agreement with those based on the OPLS model. The WS urea model leads to even faster $\alpha \to \beta$ transition and smaller accumulation of an α -helical intermediate than predicted by the OPLS model.

Hydrogen Bond Formation Between Urea and Peptide Backbone **Destabilizes A** β_{16-22} **Oligomers.** To probe the molecular mechanism of the disruption of $A\beta_{16-22}$ oligomers by urea, we compute several pair correlation functions during the first and the last 1-ns segments of 11-ns MD trajectories for DOs. The function $g_{\rm H_B-O_W}(r)$ remains virtually unchanged on the 11-ns time scale (Fig. 7 Inset, data in black), which implies that water does not play an important role in the oligomer denaturation. In contrast, the interactions of A β_{16-22} oligomers with urea show strong time dependence. Comparison of the solvation of peptide backbone amides by urea during the first and last 1-ns intervals (Fig. 7) shows that the maximum concentration of urea near amide hydrogens H_B increases by a factor of 2.4. Backbone solvation with urea also enhances the solvation of hydrophobic residues. Accordingly, the concentration of urea in the FSS of Phe increases by 50% in 11 ns.

The radial distributions of water and urea molecules with respect to the center of mass of the oligomer $g_{\text{CM}_0-\text{CM}_S}(r)$ (S = W,U) offer a striking illustration of the urea-driven penetration of solvent into $A\beta_{16-22}$ DO (Fig. 8). In the first nanosecond, the $A\beta_{16-22}$ oligomer interior is essentially dry (Fig. 8 *Left*). The concentrations of urea and water approach their bulk values at

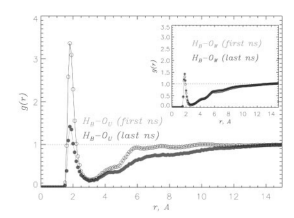


Fig. 7. The pair correlation functions $g_{\rm H_B-O_U}(r)$ (for the first and last nanoseconds in gray and black, respectively) are computed by using 11-ns simulations of $A\beta_{16-22}$ DOs in 8 M OPLS urea solution. The plots offer dramatic illustration of the "invasion" of urea molecules into $A\beta_{16-22}$ oligomers. The concentration of urea near peptide's backbone amides more than doubles in 11 ns. In contrast, the hydration of peptide backbone illustrated by $g_{\rm H_B-O_W}(r)$ in *Inset* is virtually unchanged.

 $r \gtrsim 12$ Å, which roughly corresponds to the dimensions of the oligomer. After 11 ns, a "wave" of urea molecules penetrates into the oligomer structure (Fig. 8 *Right*). The exclusion of water from the oligomer interior observed in the simulations of A β_{16-22} oligomers in water (16) (solid curve in Fig. 8) shows no time dependence. Therefore, the interior of A β_{16-22} oligomer in water remains dry. The crucial difference between the simulations in water and 8 M urea is related to a dramatic solvent (largely, urea) "invasion" into A β_{16-22} oligomers in aqueous urea. These results show that the mechanism of oligomer destabilization is similar to the urea solvation of A β_{16-22} monomer, which leads to enhanced β -structure content.

Discussion and Conclusions

Here, we report the MD simulations of $A\beta_{16-22}$ peptides and oligomers in aqueous 8 M urea solution. By using $A\beta_{16-22}$ monomers and oligomers, different initial structures, and two urea models, we have obtained a number of results regarding the effect of urea on $A\beta_{16-22}$ monomers and oligomers.

Because of preferential (with respect to water) solvation of the peptide backbone by urea, $A\beta_{16-22}$ monomers undergo RC \rightarrow β -strand transition in aqueous urea solution. Compared to water, in which RC state dominates, the majority of conformations of $A\beta_{16-22}$ monomers in urea are classified as β -strand. Formation

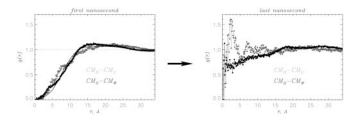


Fig. 8. The pair correlation functions $g_{\text{CM}_0-\text{CM}_W}(r)$ (black) and $g_{\text{CM}_0-\text{CM}_U}(r)$ (gray) describe solvent penetration into $A\beta_{16-22}$ DO in 8 M OPLS urea solution (CM $_{\text{O}}$, CM $_{\text{W}}$, and CM $_{\text{U}}$ are the centers of mass of the oligomer, water, and urea molecules, respectively). During the first nanosecond, $A\beta_{16-22}$ oligomer is generally devoid of solvent (*Left*, average over four trajectories). Ten nanoseconds later (*Right*, data for one of the trajectories), the wave of urea and water molecules penetrates the $A\beta_{16-22}$ oligomer, compromising its stability. Simulations of $A\beta_{16-22}$ oligomers in water show no signs of solvent penetration as the oligomer remains dehydrated during 10.7-ns MD trajectories (solid curve at *Left*).

of HBs between urea and backbone amides increases the relative (with respect to water) concentration of urea near peptide backbone by 3- to 4-fold with respect to the bulk level (depending on the specific urea model). This effect, which is electrostatic in nature, is the dominant contribution to urea–protein interactions. Because the increase in urea concentration near hydrophobic residues is linked to the formation of HBs between urea and the peptide backbone, the urea-induced change in hydrophobic interactions plays a secondary role. These results are in accord with earlier experimental and theoretical findings (21, 24–26).

Urea molecules penetrate the interior of $A\beta_{16-22}$ oligomers by making HBs with the peptide's backbone. As a consequence, urea coats hydrophobic side chains, increases the solvent exposure of oligomers, and compromises their stability. Partial or complete disintegration of $A\beta_{16-22}$ oligomers is accompanied by a dramatic growth in the ASA. The dynamics of $A\beta_{16-22}$ oligomers disruption is complex and involves a series of structural changes. Urea accelerates the $\alpha \to \beta$ transition and destabilizes the transient α -helical intermediate, which is an on-pathway species emerging before the antiparallel β -sheet formation in water (16). Urea-induced β -strand bias temporarily elevates the population of ordered antiparallel dimer structures that are stabilized by salt bridges. On longer time scales, the oligomers, regardless of the initial structures, disintegrate, because $A\beta_{16-22}$ monomers solvated with urea have lower free energy than the intact oligomers. As shown in ref. 16, electrostatic interactions

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alone cannot maintain the stability of $A\beta_{16-22}$ oligomer when hydrophobic interactions are compromised.

Our results show that urea has two opposing effects on A β_{16-22} oligomers. Urea destabilizes $A\beta_{16-22}$ oligomers, but it also increases the β -strand content in the monomer. At small concentration, urea is likely to accelerate the amyloid deposition, because the weak denaturing effect is not sufficient to destabilize the oligomers. However, at high urea concentrations, $A\beta_{16-22}$ oligomers are destabilized, which, in turn, blocks amyloid fibril formation. Because of the opposing effects, there must be an optimal concentration of urea, at which the rates of amyloidogenesis are maximum. This conclusion is consistent with the prediction of a "turnover" in the rates of amyloid assembly as a function of denaturant concentration (28), which was based on the kinetic partitioning mechanism for monomeric folding (29). The "turnover" was also observed in the recent experimental study of Hamada and Dobson (30), who examined the effect of urea on the rates of amyloid deposition for lysozyme. Because the mechanisms of promotion of β -strand propensity in A β_{16-22} monomers and the destabilization of oligomers are likely to be generic, amyloid deposition rates should, in general, show nonmonotonic dependence on urea concentration, with a maximum occurring near the concentration of equilibrium between monomeric and oligomeric species.

This work was supported in part by National Institutes of Health Grant IR01 NS41356-01.

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