

ARTICLE

# Synergistic Interactions between Repeats in Tau Protein and A $\beta$ Amyloids May Be Responsible for Accelerated Aggregation via Polymorphic States

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Supporting Information

**ABSTRACT:** Amyloid plaques and neurofibrillary tangles simultaneously accumulate in Alzheimer's disease (AD). It is known that  $A\beta$  and tau exist together in the mitochondria; however, the interactions between  $A\beta$  oligomers and tau are controversial. Moreover, it is still unclear which specific domains in the tau protein can interact with  $A\beta$  oligomers and what could be the effect of these interactions. Herein, we examine three different  $A\beta$ -tau oligomeric complexes. These complexes present interactions of  $A\beta$  with three domains in the



tau protein; all contain high  $\beta$ -structure propensity in their R2, R3, and R4 repeats. Our results show that, among these,  $A\beta$  oligomers are likely to interact with the R2 domain to form a stable complex with better alignment in the turn region and the  $\beta$ -structure domain. We therefore propose that the R2 domain can interact with soluble  $A\beta$  oligomers and consequently promote aggregation. EM and AFM images and dimensions revealed highly polymorphic tau aggregates. We suggest that the polymorphic tau and  $A\beta$ -tau aggregates may be largely due to repeat sequences which are prone to variable turn locations along the tau repeats.

n the Alzheimer brain, the two hallmark pathological lesions, first described by Dr. Alzheimer in 1906, are senile plaques<sup>1</sup> and neurofibrillary tangles (NFTs).<sup>2</sup> In Alzheimer's disease (AD), senile plaques and NFTs simultaneously aggregate. These two aggregation processes appear to occur independently because the senile plaques are extracellular deposits which develop around nerve endings and consist primarily of A $\beta$  oligomers, while the NFTs are intracellular lesions which develop in neuronal cell bodies and comprise primarily of aggregates of the microtubule (MT)-binding protein tau. Yet, recently it has been shown *in vivo* that  $A\beta$  binds to tau to form soluble stable complexes, and this binding can promote aggregation.<sup>3</sup> Importantly, the presence of both A $\beta$  and tau cause a rapid dissociation of tau from the microtubules and a collapse of axonal structure leading first to the malfunction of the synapses and ultimately to the neurons' death.<sup>4</sup>

Further studies have shown synergy between the tau and  $A\beta$  pathology in the mitochondria,<sup>5–9</sup> and it was suggested that  $A\beta$  may accelerate tau NFT formation.<sup>10–12</sup> Recent experiments led to four different mechanisms through which  $A\beta$  can promote the tau pathology:<sup>13,14</sup> (1) *in vitro* neuronal exposure to  $A\beta$  induces activation of GSK3 $\beta$  which increases tau phosphorylation,<sup>15–17</sup> thus triggering cell death by destabilizing the microtubule

network; (2) the inflammatory response may provide a link between A $\beta$  and tau pathology in AD;<sup>18</sup> (3) the increased levels of A $\beta$  may lead to proteasomal impairment within the brain and the degradation of tau;<sup>19,20</sup> and finally (4) transport deficits may lead to tau pathology as readily as tau pathology may lead to transport deficit.<sup>21,22</sup>

These studies emphasize that to understand the mechanism through which  $A\beta$ -tau complexes lead to AD pathology the interactions between  $A\beta$  and tau should be probed. Tau is a natively disordered protein which binds and stabilizes micro-tubules and shares structural features with amyloid fibrils. Amyloid oligomers, amyloid fibrils, and fibrils of tau protein<sup>23</sup> have a predominantly cross- $\beta$  pleated sheet conformation in which the polypeptide backbone is perpendicular to the fibril axis. The longest human tau isoform htau40 was observed to exist in the central nervous system. It consists of 441 residues and contains several domains, including the proline rich flanking regions P1 and P2 (residues 151–244) and four repeats R1–R4 (residues 244–369) (Figure 1). All four repeats constitute the

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Figure 1. Domain organizations of the tau isoform htau40 consists of 441 residues and contains several important domains: four pseudorepeats R1-R4 and two proline-rich flanking regions P1 and P2. The colors in the primary sequence of the pseudorepeats R2, R3, and R4 highlight the secondary structure of the repeats:  $\beta$ -structure (red), turn structure (blue), and disordered structure (green).

microtubule-binding domain, and residues in repeats R2–R4 possess a high propensity to adopt a  $\beta$ -structure.<sup>24–27</sup> Figure 1 illustrates the secondary structure of the three repeats indicating the propensity of  $\beta$ -structure. AFM, FT-IR, CD, and X-ray fiber diffraction experiments have shown that fragments of the  $\beta$ -structure in these repeats can stably interact with each other and with themselves and can aggregate into fibrils.<sup>28-30</sup> Petterson and co-workers<sup>31</sup> suggested that the hexapeptides motifs <sup>275</sup>VQIINK<sup>280</sup> and <sup>306</sup>VQIVYK<sup>311</sup> in repeats R2 and R3, respectively, act as mediators of intermolecular interactions between tau monomers which were stabilized by the anionic factor heparin. EPR spectra and EM measurements also indicated that fragments in the R2 repeat of tau have a parallel in-register arrangement of  $\beta$ -strands.<sup>32</sup> Monte Carlo simulations<sup>33</sup> of the <sup>306</sup>VQIVYK<sup>311</sup> in repeat R3 have shown that the tau molecule not only promotes the formation of stable oligomers but also is capable of stabilizing these structures because such fragments are characteristically rich in  $\beta$ -sheet structures. The simulations<sup>33</sup> have shown that the <sup>306</sup>VQIVYK<sup>311</sup> fragment in repeat R3 has a mostly parallel arrangement of  $\beta$ -strands.

According to the experimental studies above, tau aggregation occurs via interactions between  $\beta$ -strand fragments in the tau protein. Furthermore, they indicate that tau interacts with  $A\beta$ . Since all amyloidogenic peptides, including the tau protein, polymerize into fibrils, have very similar properties and share similar sheet intermediates, it is conceivable that they may interact with these  $\beta$ -pleated sheet domains via a similar fibril formation mechanism; however, how and which interactions between amyloidogenic peptides and tau protein could be involved in fibril formation are not completely understood. Here we test interactions between tau and A $\beta$  by constructing A $\beta$ -tau oligomer complexes: we combine fragments of tau R2, R3, and R4 repeats with A $\beta_{1-42}$  and A $\beta_{17-42}$  oligomers to figure out the preferred tau repeat $-A\beta$  interactions and organization. Our study revealed a stable A $\beta$ -tau oligomer complex where the fragment in tau protein that interacts with A $\beta$  to form a stable association derives from repeat R2, while those fragments from repeats R3 and R4 which interact with A $\beta$  exhibit less stable oligomeric conformations. Nonetheless, amyloids are inherently polymorphic,<sup>34,35</sup> and  $A\beta$ -tau complexes are also expected to present a rugged landscape, e.g., through variants of turn shapes and sizes and shifts in residue registration in  $\beta$ -strands via sidechain orientations. Understanding the range of structural features of the aggregates is important for effective drug design to reduce aggregate formation.

# MATERIALS AND METHODS

Molecular Dynamics (MD) Simulations Protocol. MD simulations of solvated tau and tau $-A\beta$  oligomers were performed in the NPT ensemble using the NAMD program<sup>36</sup> with the CHARMM27 force field.<sup>37,38</sup> The oligomers were energy mini-mized and explicitly solvated in a TIP3P water box<sup>39,40</sup> with a minimum distance of 15 Å from any edge of the box to any A $\beta$ atom. Any water molecule within 2.5 Å of the A $\beta$  was removed. Counterions Na<sup>+</sup> were added at random locations to neutralize the A $\beta$  charge. The Langevin piston method<sup>36,41,42</sup> with a decay period of 100 fs and a damping time of 50 fs was used to maintain a constant pressure of 1 atm. The temperature 330 K was controlled by Langevin thermostat with a damping coefficient of 10  $ps^{-1.36}$  The short-range van der Waals (VDW) interactions were calculated using the switching function, with a twin range cutoff of 10.0 and 12.0 Å. Long-range electrostatic interactions were calculated using the particle mesh Ewald method with a cutoff of 12.0 Å.43,44 The equations of motion were integrated using the leapfrog integrator with a step of 1 fs.

The solvated systems were energy minimized for 2000 conjugated gradient steps, where the hydrogen bonding distance between the  $\beta$ -sheets in A $\beta_{17-42}$  is fixed in the range 2.2–2.5 Å. The counterions and water molecules were allowed to move. The hydrogen atoms were constrained to the equilibrium bond using the SHAKE algorithm.<sup>45</sup> The minimized solvated systems were energy minimized for additional 5000 conjugate gradient steps and 20 000 heating steps at 250 K, with all atoms allowed to move. Then, the systems were heated from 250 to 300 K for 300 ps and equilibrated at 330 K for 300 ps. All simulations ran for 30 ns. These conditions are applied to all cases.

Generalized Born Method with Molecular Volume (GBMV). To obtain the relative structural stability of tau,  $A\beta$ , and tau– $A\beta$  oligomers, the oligomer trajectories of the last 5 ns were first extracted from the explicit MD simulation excluding water molecules. The solvation energies of all systems were calculated using the generalized Born method with molecular volume (GBMV).<sup>46,47</sup> In the GBMV calculations, the dielectric constant of water was set to 80. The hydrophobic solvent-accessible surface area (SASA) term factor was set to 0.005 92 kcal/(mol Å<sup>2</sup>). Each conformer is minimized using 1000 cycles, and the conformational energy is evaluated by grid-based GBMV. The minimization does not change the conformations of each conformer; it only relaxes the local geometries due to thermal fluctuation which occurred during the MD simulations.

Analysis Details. We examined the structural stability of the studied oligomers by following the changes in the number of the hydrogen bonds between  $\beta$ -strands with the hydrogen bond cutoff set to 2.5 Å and by monitoring the change in the intersheet distance (C $\alpha$  backbone—backbone distance) in the core domain of all of the oligomers. In all models we constructed, the core domain is between residue 22 and residue 35, as previously done by Miller et al.<sup>34</sup>

We further investigated the averaged water molecules around each side-chain C $\beta$  carbon within 4 Å, for the tau oligomers from repeat R2 with and without interactions with  $A\beta_{17-42}$  and, similarly,  $A\beta_{1-42}$ .

# RESULTS

Construction of the Conformational Ensemble of A $\beta$  with **Tau Repeats.** Interactions between the  $A\beta$  amyloid and the tau protein are more likely to appear in regions of residual  $\beta$ -structure in the tau protein that have the potential to serve as seeds for aggregation. Mutations in tau repeats were shown to lead to a strong tendency for aggregation which correlates with their high propensity of  $\beta$ -structure.<sup>48</sup> Our constructed A $\beta$  oligomers, tau oligomers, and A $\beta$ -tau oligomeric complexes are based on the NMR model of Lührs et al.<sup>49</sup> for the A $\beta_{17-42}$ structure and on recent experimental data<sup>25,50</sup> that provide some details of the secondary structure of the tau protein, such as  $\beta$ -structure and the position of the U-turn in the sequence. We also constructed A $\beta$ -tau oligomeric complexes using the NMR model of Tycko and co-workers;<sup>51</sup> however, our simulations revealed unstable complexes (Figure S1). To match tau with  $A\beta_{17-42}$  oligomers, we extracted tau fragments that have the ability to serve as seeds for aggregation. The highest propensity of  $\beta$ -structure content in the tau protein is in repeats R2, R3, and R4 (Figure 1). These repeats are known to be involved in the abnormal aggregation of tau.<sup>25,50</sup> Moreover, the high propensity of  $\beta$ -structures in these repeats is also known to be essential for paired helical filaments formation.<sup>28,48</sup> In addition to the  $\beta$ -structure in the repeats, we also consider other similar structural properties between  $A\beta$  and each of the R2, R3, and R4 repeats. More specifically, we consider the tendency to form stable interactions between A $\beta$  and tau repeats in both the  $\beta$ -structure domain and the U-turn domain, e.g., hydrophobic interactions, electrostatic interactions, polar interactions, and salt bridge that stabilize the turn. Consequently, the tau repeats consist of two  $\beta$ -strands connected by U-turn, similarly to A $\beta$ peptides.

Recent EM and AFM images and dimensions of the tau fibrils<sup>52</sup> revealed that only the R2, R3, and R4 tau repeats can



**Figure 2.** Initial structural models: (A)  $A\beta_{17-42}$  tetramer interacts with tau tetramer and (B) tau octamer. The tau protein fragment of each monomer is taken from the repeat R2. Blue arrows illustrate hydrophobic interactions between  $A\beta_{17-42}$  and repeat R2.

fit. It is likely that the other tau protein domains that do not have a preponderance for  $\beta$ -structure "hang out" and are not part of the regular fibrillar organization. In constructing the fibril, we assume that the conventional turn shape in  $A\beta$  oligomers, which is also observed in other amyloids,<sup>53,54</sup> can act as seed for fragments of tau repeat oligomers and examine whether the simulated tau and  $A\beta$ —tau oligomers can retain the  $\beta$ -strandturn- $\beta$ -strand shape. Our assumption is based on the fact that amyloids from different species or with different sequences (such as in prion) have close structural similarity.<sup>55</sup> We further assume that the secondary structures of octamers of the tau repeats R2, R3, and R4 studied here are similar to those of  $A\beta$  oligomers, as illustrated by experimental studies<sup>25,50</sup> and detailed below.

To construct A $\beta$ -tau oligomeric complexes, we used tetramers of repeats R2, R3, and R4 and associated each tetramer with A $\beta_{17-42}$  tetramers. We first test ensemble of alignments between  $A\beta_{17-42}$  and sequences in tau repeats which maximize the overlap between the hydrophobic residues. For example, hydrophobic residues in the <sup>275</sup>VQIINK<sup>280</sup> motif in repeat R2 can match the hydrophobic residues <sup>17</sup>LVFFAE<sup>22</sup> in  $A\beta$  (Figure 2), and the U-turn and C-terminal regions of A $\beta$  can then superimpose on the continuous repeat sequence in tau. Similarly, the fitting of the hydrophobic residues in <sup>306</sup>VQIVY<sup>310</sup> motif in repeat R3 matches the  $A\beta$  <sup>17</sup>LVFFA<sup>21</sup> segment (Figure 3) and the R4 repeat <sup>337</sup>VEVK<sup>310</sup> motif matches the  $A\beta$  <sup>17</sup>LVFF<sup>20</sup> segment (Figure 4). The final alignments are indicated in Figures 2, 3, and 4 for the R2, R3, and R4 repeats, respectively. As can be seen for repeat R2, four hydrophobic interactions between tau and A $\beta$  appear in one  $\beta$ -strand and two hydrophobic interactions appear in the second. For repeat R3, five hydrophobic interactions between tau and A $\beta$  appear in the first strand and two hydrophobic interactions in the second. Finally, for repeat R4, three hydrophobic interactions between tau and A $\beta$ 



**Figure 3.** Initial structural models: (A)  $A\beta_{17-42}$  tetramer interacts with tau tetramer and (B) tau octamer. The tau protein fragment of each monomer is taken from repeat R3. Blue arrows illustrate hydrophobic interactions between  $A\beta_{17-42}$  and repeat R3.



**Figure 4.** Initial structural models: (A)  $A\beta_{17-42}$  tetramer interacts with tau tetramer and (B) tau octamer. The tau fragment of each monomer is taken from repeat R4. Blue arrows illustrate hydrophobic interactions between  $A\beta_{17-42}$  and repeat R4.

appear in one strand and two hydrophobic interactions appear in the second.

Considering the  $\beta$ -structures and the turn regions in repeats R2, R3, and R4 in tau, <sup>50</sup> we constructed three parallel octamers of

Table 1. Conformational Energy of the Constructed Oligomers

oligomer model	conformational energy <sup>a</sup> (kcal/mol)	Figure	
$A\beta_{17-42}$	-2244.66 (103.01)	Fig. 6 and Fig. 1B in Ref. 34	
A $\beta_{\rm 17-42}-{\rm tau}~{\rm R2}$	-4238.12 (99.15)	Fig. 2A	
A $\beta_{\rm 17-42}-{\rm tau}~{\rm R3}$	-2757.63 (104.40)	Fig. 3A	
A $eta_{17-42}-$ tau R4	-5319.35 (116.07)	Fig. 4A	
Tau R2	-6206.32 (127.23)	Fig. 2B	
Tau R3	-3387.86 (106.66)	Fig. 3B	
Tau R4	-8580.03(128.77)	Fig. 4B	
<sup><i>a</i></sup> Conformational energies were computed using the GBMV calcula- tions. The standard deviation values are presented in parentheses.			

26 continuous residues from each repeat which form organizations similar to those obtained previously by Miller et al.<sup>34</sup> for  $A\beta_{17-42}$  octamers: the first octamer consists of a fragment in repeat R2, residues V275–V300 (Figure 2), the second of a fragment in R3, residues V306–K331 (Figure 3), and the third octamer consists of a fragment in R4, residues V337–H362 (Figure 4).

We further constructed the full-length  $A\beta_{1-42}$  dodecamer by applying the Lührs model<sup>49</sup> (PDB code ID: 2BEG) for  $A\beta_{17-42}$ . Then, for each monomer we linked residues 1–16 in a manner similar to our recent construction of the tubular  $A\beta_{1-42}$  fibrils.<sup>56</sup> For the tau dodecamer, only the R2 repeat was used; however, we extended the sequence in the N-terminal for the structure shown in Figure 2A, i.e., each monomer in the tau oligomer consists of residues K259–V300. Finally, we constructed the  $A\beta_{1-42}$ –tau dodecamer complex by interacting hexamers of  $A\beta_{1-42}$  with hexamers of the tau fragment K259–V300. Figure 5 illustrates the tau,  $A\beta_{1-42}$  and  $A\beta_{1-42}$ –tau oligomers.

Reaction Coordinates for the Formation of  $A\beta$ -Tau Oligomeric Complexes from  $A\beta$  and Tau Oligomers Demonstrate That the R2 Repeat in Tau Is More Likely to Interact with  $A\beta$ . To investigate the stability of each soluble  $A\beta$ -tau oligomeric complex for each of the fragments in repeats R2, R3, and R4 interacting with  $A\beta$ , the conformational energies were computed for  $A\beta$  oligomers, tau oligomers, and the  $A\beta$ -tau oligomeric complexes (Table 1). The conformational energies for each oligomer are based on the energy computed with the generalized Born method with molecular volume (GBMV).<sup>46,47</sup> We estimate the relative stability of each  $A\beta$ -tau oligomeric complex by comparing its energy with the two separate components,  $A\beta$  and tau oligomers, as illustrated by the following chemical "reaction":

$$(\mathbf{A}\beta)_{2n} + (\mathrm{Tau})_{2n} \leftrightarrow 2[(\mathbf{A}\beta)_n \cdot (\mathrm{Tau})_n] \tag{1}$$

where *n* indicates the number of monomers within an oligomer. Figure 6 demonstrates the reaction coordinates for the three repeats. As seen from Figure 6, the  $A\beta$ -tau R2 repeat oligomeric complex is more stable energetically than its two separate components,  $A\beta$  and tau oligomers. One can view the reaction coordinates for R2 as an exothermic reaction, where the product ( $A\beta$ -tau oligomeric complex) is more stable than the reactants ( $A\beta$  and tau oligomers). However, for the repeats R3 and R4, the picture is the opposite: the two separate  $A\beta$  and tau oligomer components are more stable than the  $A\beta$ -tau oligomeric complex. Hence, these relative stabilities calculations indicate that repeat R2 in the tau protein is more likely to have the ability to interact with  $A\beta$  to form a stable oligomeric complex.

Tau Repeat R2 Oligomers Structurally Stabilize the  $A\beta$  Oligomers. While parallel  $A\beta$  oligomers are stable in experiment



**Figure 5.** Initial structural models: (A)  $A\beta_{1-42}$  tetramer interacts with tau tetramer and (B) tau octamer. The tau protein fragment of each monomer is taken from repeat R2.



Figure 6. Relative energies of the two separate components A $\beta$  and tau and of A $\beta$ -tau complex for repeats R2, R3, and R4.

and simulations, experimental structural data of soluble  $A\beta$ -tau oligomeric complexes are unavailable. Herein, we test models for parallel  $A\beta$ -tau oligomeric complexes and parallel tau oligomers. We investigate their conformational stability by analysis of the hydrophobic association among the  $\beta$  strands.

Previously, Miller et al.<sup>34</sup> demonstrated that for the  $A\beta_{17-42}$  parallel octamers based on the Lührs model >70% of the hydrogen bonds are retained during the 30 ns simulations, and the  $\beta$ -sheet distances across the U-turn are in the 12–13 Å range. Here, for the  $A\beta_{17-42}$ -tau R2 oligomeric complex, >70% of the hydrogen bonds are also retained during the simulations, and assisted by the backbone hydrogen bonds, the association is slightly better among the  $\beta$ -strands compared with the  $A\beta_{17-42}$ 

parallel octamer (Figure 7A). We also examined three different  $\beta$ -sheet distances across the U-turn for the  $A\beta_{17-42}$ -tau oligomeric complexes: between two  $\beta$ -sheets of  $A\beta$ , between two  $\beta$ -sheets of the tau repeat, and between a monomer of  $A\beta$  and a nearby monomer of a tau repeat (Figure 8). All three  $\beta$ -sheet distances across the U-turn for  $A\beta_{17-42}$ -tau R2 oligomeric complex are in the  $\sim 12-13$  Å range, indicating that the tau R2 oligomers stabilize the  $A\beta_{17-42}$ -tau R4 oligomeric complexes demonstrate different values, in the range of 12-15 and 10-20 Å, respectively. Proline residues are disfavored in  $\beta$ -structures. In repeat R3, Pro312 may explain the relatively large  $\beta$ -sheet distances across the U-turn for  $A\beta_{17-42}$ -tau R3 oligomers.



**Figure 7.** Fraction of the number of hydrogen bonds (in percentage) between all  $\beta$ -strands compared with the initial oligomer structures of A $\beta$ -tau for repeats R2, R3, and R4 (A) and of tau oligomers for repeats R2, R3, and R4 (B).



**Figure 8.** C $\alpha$  backbone–backbone distance of A $\beta$ –tau for repeats (A) R2, (B) R3, and (C) R4. Three different distances were examined: averaged distance between two A $\beta$  peptides (blue), between A $\beta$  peptide and tau monomer (red), and the averaged distance between two monomers of tau (green).

The large  $\beta$ -sheet distances across the U-turn for  $A\beta_{17-42}$ -tau R4 are due to the few hydrophobic interactions between the two  $\beta$ -strands (three hydrophobic interactions between the  $A\beta$  peptide and tau repeat in one strand and only two appear in the second). Although there are similarities in the structural properties in the turn region for  $A\beta_{17-42}$ -tau for both repeats R3 and R4, the interactions between the  $\beta$ -strands are probably more important for structural stability than the turn region. Although the  $A\beta_{17-42}$ -tau R4 oligomeric complexes present only slightly less than 70% of their hydrogen bonds for the last 10 ns of the simulations (Figure 7A), the differences among the  $\beta$ -sheet distances suggest that the repeats R3 and R4 are less preferred to interact with  $A\beta$  (Figure 9B,C).

Interestingly, in tau oligomers consisting of repeats R3 and R4, the percentage of hydrogen bonds that are retained during the simulations are less than 60% for R3 and less than 50% for R4, while in repeat R2, >70% of the hydrogen bonds are retained. Hence, the tau oligomer with the R2 repeat has a better association among the  $\beta$ -strands assisted by backbone hydrogen



Figure 9. C $\alpha$  backbone-backbone distance of tau oligomers for repeats R2, R3, and R4.



bonds (Figure 7B), indicating that repeat R2 has a high propensity of  $\beta$ -structures that may interact with  $A\beta$  to form a stable oligomer. We note that the percentages of hydrogen bonds in  $A\beta$ -tau oligomeric complexes for all repeats and in particular for R3 and R4 are higher than in tau oligomers that seem to have similar structural properties. The reason for this difference could lie in the electrostatic interactions between the  $\beta$ -strands: for  $A\beta$ -tau R2, two ( $A\beta$  Glu22-tau Lys280;  $A\beta$  Asp23-tau K281); for  $A\beta$ -tau R3, one ( $A\beta$  Glu22-tau K311); and for  $A\beta$ -tau R4, one interaction ( $A\beta$  Asp23-tau K343). Yet, the  $\beta$ -sheet distances expand to 20–21 Å for tau R2 oligomers; therefore, the R2 repeat interacts more favorably with  $A\beta$  than with itself.

Solvation of Backbone Residues in Tau and A $\beta$ -tau Oligomeric Complexes Indicate the Stability of the Com**plexes.** We further examine the stability of the A $\beta$ -tau R2 oligomeric complexes by investigating the backbone solvation. To this aim, we compared the backbone solvation of tau oligomers from the R2 repeat with the tau oligomers from the R2 repeat that are complexed with A $\beta$  oligomers. Figure 10 illustrates the backbone solvation values for each residue in both tau $-A\beta_{17-42}$  and tau $-A\beta_{1-42}$  and the corresponding tau oligomers. For both tau oligomers, with 26 residues in the R2 repeat and the tau-A $\beta_{17-42}$  complex, the backbone solvation values for all residues are similar, except for residue 37. The solvation is related also to the character of residue 37 in the sequences for the oligomers: in the A $\beta$  sequence, it is G37 while in the tau it is D37, indicating that small residues are more solvated. Thus, overall, tau and tau-A $\beta_{17-42}$  oligomers illustrate similar stabilities. This result is in agreement with the hydrogen bond analysis for these oligomers (Figure 7): both oligomers have relatively high percentage of hydrogen bonds during the simulations.

For the full-length tau $-A\beta_{1-42}$  oligomers, the backbone solvation illustrates smaller values compared with the tau oligomers (Figure 7), indicating that the tau $-A\beta_{1-42}$  oligomers have better  $\beta$ -sheet interactions and thus higher stability than the tau oligomers.

## DISCUSSION

The aggregation of tau into NFTs, and A $\beta$  into amlyoids, are processes long known to be related to AD progression. More recent experimental studies led to the proposition that A $\beta$  may directly or indirectly interact with tau to accelerate NFTs formation.<sup>4-8,10-14</sup> Yet, the mechanism and the related interactions through which tau and A $\beta$  self-assemble together have been controversial. Herein, we investigate the interactions between tau oligomers and  $A\beta$  oligomers at the molecular level to figure out the preferred organizational state. Using all-atom MD simulations in explicit solvent, we investigated the parallel oligomeric architectures of three different repeats in the tau protein-R2, R3, and R4—and the interactions of these parallel tau architectures with A $\beta_{17-42}$  oligomers. While the absence of experimental examination of the interactions of purified tau variants with A $\beta$ oligomers limits our work to the prediction of the interactions through computational tools, our models are based on experimental data: the oligomers were based on the experimental model by Lührs et al.<sup>49</sup> as previously studied by Miller et al.<sup>34</sup> for  $A\beta_{17-42}$ ; and the tau oligomer models are also based on experimental data which illustrate high propensity of  $\beta$ -structure in tau repeats R2, R3, and R4 pointing to likely intermolecular  $\beta$ -sheet interactions. In addition to the  $\beta$ -structure of the peptide <sup>306</sup>VQIVYK<sup>311</sup> in repeat R3,<sup>28</sup> other studies by NMR spectroscopy,<sup>57,58</sup> CD measurements,<sup>59</sup> and FTIR spectroscopy<sup>60</sup> suggest  $\alpha$ -helical structures in repeat R3. Herein, we did not test these  $\alpha$ -helical structures, and further investigations will need to be performed.

In our proposed  $A\beta$ -tau models, we consider two main factors: hydrophobicity and flexibility of these domains. The hydrophobicity factor plays a key role in forming hydrophobic core and enhancing the interactions between  $A\beta$  oligomers and tau oligomers. Recently, hydrophobicity and flexibility were illustrated to be the primary factors for oligomer assembly.<sup>61</sup> A recent study<sup>62</sup> speculated that  $A\beta$  oligomers may be responsible for accumulation of the transactive response DNA-binding protein 43 (TDP-43). Further studies are needed to elucidate the nature of the  $A\beta$ -TDP-43 interactions.

Our study leads to a major conclusion: A $\beta$  oligomers interact better directly with fragments of the tau repeat R2 than repeats R3 and R4. Further experiments, such as surface plasmon resonance may be useful to investigate the affinities of the various tau repeats to A $\beta$  oligomers. The U-turn root-mean-square deviations (RMSDs) indicate that the oligomer of A $\beta_{17-42}$  that interacts with fragments of repeat R2 is slightly more stable than the A $\beta_{17-42}$  oligomer that interacts with fragments of repeats R3 or R4 (Figure S2). Figure S3 illustrates that the U-turn RMSDs of tau repeats R2 and R3 oligomers are similar to values observed previously<sup>34</sup> for A $\beta_{17-42}$ , and the RMSDs of oligomers of tau repeats R4 are slightly larger. Overall, the R2, R3, and R4 oligomers are structurally stable in the turn region; however, the  $\beta$ -sheet distances exhibit that the R2 oligomer cannot form a stable oligomer alone, while the tau repeats R3 and R4 oligomers can form stable structures (Figure 9). Nonetheless, the tau R2 oligomer is the more likely candidate to interact with  $A\beta_{17-42}$  as compared to R3 and R4 oligomers (Figure 8). We emphasize, however, that because amyloids including A $\beta$  are polymorphic,<sup>63</sup> A $\beta$ -tau complexes are also expected to be polymorphic, either in their gross organizations or on a minor scale. The polymorphic range can be broad, as was observed by cryo-EM, ssNMR, and modeling, with and without ion binding.<sup>34,35,56,64-66</sup> Two likely features that can lead to polymorphism is the U-turn shape, and the organization between the sequences of tau repeats with respect to the sequence of A $\beta$ . Additional arrangements of the A $\beta$  and tau repeats which relate to the shift of the A $\beta$  sequence along the tau repeats can expand the polymorphic range of the complexes. In addition, populations of the tau can be shifted: while the R2 oligomer cannot form a stable oligomer alone, it gets stabilized when interacting with A $\beta$  oligomer, thus shifting the tau ensemble. Recent EM and AFM measurements revealed highly polymorphic fibrils for tau<sup>52,67,68</sup> that differ significantly in fibril parameters such as cross section and twist. These measurements illustrate irregular fibril structures with no symmetry which most likely relate to the turn shapes in the tau repeats. Since the U-turn conformations in the tau repeats may differ as the sequence registration is shifted, the cross sections may be different, and consequently the number of protofilaments that wind around each other to form a mature fibril may also be different. In summary, the U-turn appears to play a key role in the amyloid polymorphic landscape, shifting the populations of the oligomers, fibrils, and gross-scale organization toward different states, as observed in the EM and AFM images and dimensions.

The hydrophobic regions in both R2 and R3 interact with the microtubules (MTs) and stabilize them. In AD, tau becomes excessively phosphorylated, loses its ability to bind to MTs, and aggregates into NFTs. One can argue that when tau loses its ability to bind to MTs, it interacts with  $A\beta$  to form a more stable oligomer, accelerating aggregation. We further suggest that when A $\beta$  interacts directly with tau repeat R2, the tau does not interact with the MTs, and consequently MTs degradation takes place, leading to toxicity. Mutations in tau in repeats R2, R3, and R4 showed no decrease in the binding of tau to the MTs.<sup>69</sup> However, tau mutations lead to faster aggregation into paired helical filaments (PHFs).<sup>69–74</sup> Some of the mutations may weaken tau's ability to bind to MTs75-77 and thus cause degradation of the MT and toxicity. A recent study<sup>78</sup> indicated that  $A\beta$  oligomers induce tau aggregation and formation of  $\beta$ -sheet rich neurotoxic tau oligomers; yet the mechanism for the toxicity is still controversial, and further investigations will need to be performed.

## ASSOCIATED CONTENT

**Supporting Information.** Figures S1–S3. This material is available free of charge via the Internet at http://pubs.acs.org.

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## ABBREVIATIONS

NFT, neurofibrillary tangles; MT, microtubule; AD, Alzheimer's disease; MD, molecular dynamics; R, tau repeat.

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