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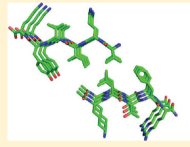


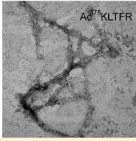
Secondary Nucleating Sequences Affect Kinetics and Thermodynamics of Tau Aggregation

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ABSTRACT: Tau protein was scanned for highly amyloidogenic sequences in amphiphilic motifs $(X)_n Z$, $Z(X)_n Z$ $(n \ge 2)$, or $(XZ)_n$ $(n \ge 2)$, where X is a hydrophobic residue and Z is a charged or polar residue. N-Acetyl peptides homologous to these sequences were used to study aggregation. Transmission electron microscopy (TEM) showed seven peptides, in addition to well-known primary nucleating sequences Ac²⁷⁵VQIINK (AcPHF6*) and Ac³⁰⁶VQIVYK (AcPHF6), formed fibers, tubes, ribbons, or rolled sheets. Of the peptides shown by TEM to form amyloid, Ac10VME, AcPHF6*, Ac375KLTFR, and Ac393VYK were found to enhance the fraction of β -structure of AcPHF6





formed at equilibrium, and Ac375KLTFR was found to inhibit AcPHF6 and AcPHF6* aggregation kinetics in a dose-dependent manner, consistent with its participation in a hybrid steric zipper model. Single site mutants were generated which transformed predicted amyloidogenic sequences in tau into non-amyloidogenic ones. A M11K mutant had fewer filaments and showed a decrease in aggregation kinetics and an increased lag time compared to wild-type tau, while a F378K mutant showed significantly more filaments. Our results infer that sequences throughout tau, in addition to PHF6 and PHF6*, can seed amyloid formation or affect aggregation kinetics or thermodynamics.

au is a microtubule-associated protein that regulates microtubule stability, neurite growth, and other microtubuledependent functions. The human protein exists as six isoforms, consisting of an acidic N-terminal containing up to two 29amino acid inserts and a basic microtubule binding region (MTBR) containing either three or four tandem 31 (or 32)-amino acid pseudorepeats (R1-R4) near the C-terminal. 1-4 Normal phosphorylation of tau by proline-directed kinases (GSK-3 β , MAPK, CDK5) and by non-proline-directed kinases (PKA, MARK, SADK, and Src family of tyrosine kinases) may affect tau's affinity for microtubules and regulate the dynamics of microtubules and establish neuronal polarity, axonal outgrowth, and axonal transport in mature neurons. 5,6 "Hyperphosphorylation" of tau and cleavage of tau by cell proteases (thrombinlike proteases, cathepsins, caspases, and calpains) appear to lead to the aggregation of tau into dimers, oligomers, and paired helical filaments (PHFs) which make up the neurofibrillary tangles (NFTs) in patients with Alzheimer's disease (AD) and other tauopathies.^{7–9} Recent evidence suggests that, while smaller aggregates of tau occur as early events during the course of the disease and are responsible for neurotoxicity, larger filaments and filament bundles are neuroprotective. 10-1

While early work showed recombinant tau and tau isolated from microtubule preparations to be largely unstructured or "natively unfolded", aggregated forms of the protein displayed physical characteristics of amyloid, including a cross- β X-ray diffraction pattern. 14-18 Nucleating sequences or "hot spots" in the protein sequence, ²⁷⁵VQIINK (PHF6*) in R2 and ³⁰⁶VQI-VYK (PHF6) in R3, were identified, and it has been shown that one or more of these sequences is essential for filament formation. 19-22 Previous work from our laboratory has shown that an N- and C-terminal blocked peptide AcPHF6 aggregates into filaments displaying a cross- β -X-ray pattern and, in the presence of smaller peptides, produces twisted filaments with morphology similar to PHFs. 23 More recent X-ray work on the PHF6 peptide has shown it to pack in a "steric zipper" with antiparallel layers of parallel, in-register aligned peptides, 24-26 while EPR and NMR of tau protein have shown that PHF6 or PHF6 and PHF6* interact intermolecularly in a parallel or antiparallel fashion in protein aggregates. 12,27-29 Furthermore, NMR and FRET studies have shown that while the backbone of the protein monomer appears to be in rapid exchange with an ensemble of conformations, the average conformation is one in which the C-terminal is folded over the MTBR and the Nterminal is folded over the C-terminal in a "paper clip" conformation. 11,12,30-34 It has been hypothesized that phosphorylation of tau at specific sites opens the paper clip conformation, exposing the MTBR and catalyzing aggregation of the protein or exposing a portion of the N-terminal which inhibits anterograde fast axonal transport, resulting in neurotoxicity. 30,35

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Tau lacking the PHF6 still has the ability to form aggregates, implying that PHF6* or other sequences in tau can nucleate tau polymerization or stabilize oligomeric structures. 20-22,36 A Cterminal tail peptide that included the PHF6-like sequence ³⁹²IVYK³⁹⁵ but lacked either PHF6 or PHF6* was found to form straight filaments containing a high degree of β -sheet conformation.³⁷ While deletion of the C-terminal tail at Lys321 enhances aggregation, a similar deletion at Asp314 renders tau incapable of aggregation, suggesting that ³¹⁴DLSKVTS is essential for PHF formation. A recent NMR study on full length tau has shown that several other sequences including ³³⁶QVEVKSEKLD, ⁸⁶GKQAAAQ, and ²²⁴KKVAVVR are in the β -sheet conformation. ³¹ Surprisingly, deletion of 336QVEVKSE from the MTBR of R4 or the entire R4 sequence enhanced polymerization, suggesting that sequences within R4 have an inhibitory effect. 21,38,39

In a previous study amyloidogenic propensities of individual amino acid residues were extrapolated from measured properties of a family of AcPHF6 peptides (VQIVXK, where X is the substituted residue) using a principal component analysis approach. 40 Using individual amino acid propensities, amyloidogenic propensities of sequences within tau conforming to structural motifs, $(X)_n Z$, $Z(X)_n Z$, or $(XZ)_n$ $(n \ge 2)$, where X is a hydrophobic residue and Z is a charged or polar amino acid, were estimated. The goal of the present study was to determine if sequences within tau predicted to have high amyloidogenic propensities could, in fact, seed filaments or interact with other known nucleating sequences within the protein (PHF6 and PHF6*). Surprisingly, we found that peptides with these amyloidogenic sequences were able to form all different morphological expressions of amyloid including filaments, tubes, ribbons, and rolled sheets. Furthermore, CD difference spectra and kinetics measurements showed that some of these peptides were able to interact with PHF6 or PHF6* peptides in a manner which either enhanced or inhibited aggregation. We found that point mutations in tau which severely reduce the amyloid propensities of these sequences affect the kinetics of aggregation or the number of filaments formed at equilibrium.

MATERIALS AND METHODS

Materials. Fmoc-amino acids were purchased from EMD Chemicals (Gibbstown, NJ). High molecular weight heparin (HMWH; porcine mucosa; 18 kDa average MW; 6–30 kDa range) was obtained from Sigma-Aldrich Chemical (St. Louis, MO) and was used without further purification. All other chemicals and buffers were of reagent grade.

Amyloidogenic Propensities of Sequences in Tau Protein. Tau protein was scanned for sequences which satisfied the amphiphilic motifs $(X)_n Z$, $Z(X)_n Z$, or $(XZ)_n$ $(n \ge 2)$, where X is a hydrophobic and Z is a polar or charged amino acid residue. Amyloidogenic propensities of sequences satisfying these motifs were evaluated using the sum of the absolute values of their constituent residue PC1 scores.

Peptide Synthesis. *N*-Acetyl peptide amides corresponding to potentially nucleating sequences selected in tau were prepared according to previously published procedures and purified by reverse-phase HPLC using a water—acetonitrile gradient. ^{23,40,41} Purity was checked using ESI-MS or MALDITOF MS.

Far-UV CD Spectroscopy. Stock solutions of the peptides were prepared ($\sim 1-5$ mg/mL) in deionized water. Final peptide concentrations were determined by absorbance using published

molar extinction coefficients for tyrosine $[\varepsilon_{276} \text{ of } 1390 \text{ M}^{-1} \text{ cm}^{-1}]$ or for the peptide bond $[\varepsilon_{192} \text{ of } 7110 \text{ M}^{-1} \text{ cm}^{-1}].^{42,43}$ For CD measurements, peptide stocks were diluted to 100 μ M in 5 mM MOPS/0.15 M NaCl, pH 7.2 (MOPS-Cl buffer) containing 0.1 mg/mL of heparin. Samples were transferred to a 0.1 cm path length quartz cell and bath sonicated for 1 min. Far-UV CD spectra were acquired an AVIV 202 CD spectrophotometer, following a 1 h incubation at room temperature. ^{23,40,44} CD measurements of peptide binary mixtures were prepared in a similar manner with each of the peptides present at 100 μ M. In order to calculate mean residue molar ellipticity, the total number of residues was set to the sum of the total number of residues in the peptides included in the mixture.

TEM. Samples for TEM were prepared from peptide stock solutions in MOPS-Cl buffer containing 0.3 mg/mL heparin. Following incubation overnight at room temperature, samples were loaded onto carbon-coated Formvar copper grids (200 mesh) and stained with 2% w/w uranyl acetate. A JEOL 1200 EX scope interfaced to a digital camera was used to visualize samples. Distance measurements were made using Adobe Photoshop software (Adobe, Inc., San Jose, CA).

Kinetics of Tau Peptide Aggregation. Polymerization kinetics were measured by monitoring the increase in Thioflavin S (ThS) fluorescence at 490 nm (440 nm excitation) in a manner similar to that used to follow aggregation in other amyloid-forming peptides and proteins.^{40,41,45–47} For each peptide, a 1 mg/mL sample stock solution was filtered through a Millipore Ultrafree-MC 100 000 NMWL filter unit (Billerica, MA) by centrifuging for 10 min at 14500 rpm, using an Eppendorf MiniSpin plus benchtop centrifuge (Westbury, NY). Following filtration, the concentrations of the peptides were determined by UV absorbance. Stock solutions were then diluted in 200 uL wells of a 96-well clear bottom ELISA plate such that the final concentrations were 100 μ M peptide, 0.3 mg/mL heparin, 100 μ M ThS, in MOPS-Cl buffer, or in 5 mM sodium phosphate containing 0.15 M NaCl, pH 3.0 (phosphate-Cl buffer). Fluorescence data were collected in triplicate in kinetic mode, using a Molecular Devices SPECTRA Max Gemini XPS spectrofluorimeter (Sunnyvale, CA), with measurements taken at 15 s intervals. Mixing between readings was accomplished using a 1/8 in. PTFE grade 2 polished mixing bead (Orange Products, Allentown, PA). In order to minimize time before the first data were collected, peptide solutions were added just before the commencement of the readings. A blank containing only buffer, heparin, and ThS was recorded concurrently with the samples. Baseline corrected kinetic data were fit to Gompetz growth curves according to

$$y = e^{-k't} A e^{-e^{(t-t_1)/b}}$$
 (1)

where y is defined as the fluorescence signal at time t, t_i corresponds to the inflection point of the curve or the time of maximum growth rate, A is the maximum fluorescence observed for a given sample, and $b=1/k_{\rm app}$, where $k_{\rm app}$ is the rate constant of aggregation, in units of s⁻¹. The pregrowth curve exponential, ${\rm e}^{-k't}$, was inserted into the traditional Gompetz growth curve equation to take into account the slow decrease in fluorescent intensity resulting from ThS bleaching, quenching, and gravitational precipitation of the amyloid—ThS complex. Fach experiment showing peptide aggregation was run three times, and the error in fitted parameters was determined from the standard deviation of the fitted parameters. Kinetic data for

mixtures of some peptides were fit to a sum of growth curve functions according to

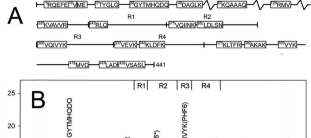
$$y = Ae^{-e^{(t-t')/b}} + A'e^{-e^{(t-t'')/b'}}$$
 (2)

where $b' = 1/k'_{app}$. **2N4R Tau Experiments.** Site-directed tau mutants M11K, F378K, and Y394K were generated from 2N4R tau (pT7Chtau40 (a gift from Dr. Jeff Kuret)) using the Stratagene QuikChange Site-Directed Mutagenesis kit following manufacturer's protocols. Primers for site-directed mutagenesis were synthesized by Integrated DNA Technologies. The 6x histagged proteins were purified using Ni²⁺ charged chelating sepharose fast flow chromatography, followed by Superdex 200 gel filtration chromatography as previously described. 49 2 µM of wild type or mutants of 2N4R tau (M11K, F378K, and Y394K) were incubated with 75 μ M arachidonic acid in polymerization buffer (100 mM NaCl, 10 mM Hepes pH 7.64, 0.1 mM EDTA, 5 mM DTT) for 240 min at room temperature. Right angle laser light scattering (LLS) readings were taken at various time points, and images were analyzed by Adobe Photoshop for histogram intensity of the light scattering band. 50 Data were fit to the Gompertz growth curve as previously described. A (final extent of polymerization), $k_{\rm app}$ (proportional to the rate of polymerization), and lag (time until significant polymerization), which equals $t_i - b$, were determined as the mean \pm SEM of three trials.⁵¹ After 240 min, reactions were diluted 1:10 and fixed with 10% glutaraldehyde. TEM grids were prepared, and filaments were stained with 2% uranyl acetate. Images were collected at 3600× using a Technai F20 XT field emission TEM (FEI). Quantitation of 5 TEM images for filament number, length, and mass per field was done using Image Pro Plus 6 with a minimum length cutoff of 15 nm.

RESULTS

Amyloidogenic Sequences in Tau Protein. In a previous study we used a family of tau-related peptides, AcVQIVXK, to define an amyloidogenic propensity of the amino acid residue X based on a group of experimental observables typically used to define amyloid. 40 We found that peptide sequences with an amphiphilic motif (X), Z, where X is a hydrophobic residue and Z is a charged or polar amino acid, have a high propensity to act as nucleating sequences. Others have shown that amphiphilic sequences in which hydrophobic and polar residues alternate above and below the extended peptide chain $((XZ)_n (n \ge 2))$ or have a $Z(X)_n Z$ motif also have a high tendency to nucleate amyloid formation. 24,52-55 We scanned tau protein for these amyloid-forming motifs and calculated their amyloidogenic propensities using their constituent residue amyloidogenic propensity scores. Using this approach, we found 20 high propensity sequences (Figure 1): 8 sequences (5RQEFEV, 10VME, ¹⁷TYGLG, ²⁸GYTMHQDQ, ⁴⁰DAGLK, ⁸⁷KQAAAQ, ¹²⁶RMV, ²²⁵KVAVVR) in the N-terminal region, 6 in the microtubule binding region (242RLQ in R1; 275VQIINK (PHF6*) and ²⁸²LDLSN in R2; ³⁰⁶VQIVYK (PHF6) in R3; ³³⁷VEVK and ³⁴³KLDFK in R4), and 6 in the C-terminal region (³⁷⁵KLTFR, ³⁸²AKAK, ³⁹³VYK, ⁴¹⁹MVD, ⁴²⁸LAD, ⁴³²VSASL). Of these, PHF6* and PHF6 in the MTBR have been shown experimentally to act as nucleating sequences in tau-related peptides or full length tau protein.19

TEM of Predicted Amyloid Forming Sequences in **Tau.** We prepared all 20 predicted amyloidogenic sequences as their N-acetyl peptide amides, since N- and C-terminal charges



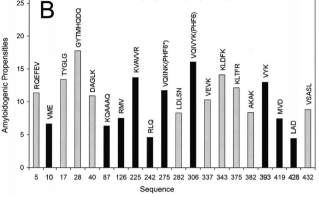


Figure 1. (A) Schematic of full-length tau showing the 20 sequences which fit the motifs $(X)_n Z$ or $Z(X)_n Z$ $(n \ge 2)$ (black) or $(XZ)_n$ $(n \ge 2)$ (gray), where X is a hydrophobic residue and Z is a charged or polar amino acid. The model assumes that these sequences have a high propensity to act as nucleating sequences in amyloid formation. (B) Amyloidogenic propensities for sequences calculated by taking the sum of the absolute values of constituent amino acid propensities.⁴

have been shown to have a profound effect on the ability of a peptide to form amyloid.⁵⁶ We studied the behavior of these peptides in the presence of HMW heparin, a known polyanion inducer of tau protein aggregation. ^{19,29,45,46,57–60} Using TEM, we observed fibers with diameters between 3 and 7 nm for Ac5RQEFEV, Ac40DAGLK, AcPHF6*, AcPHF6, Ac343KLDFK, Ac³⁷⁵KLTFR, Ac³⁹³VYK, and Ac⁴¹⁹MVD following overnight incubation in MOPS-Cl buffer with HMW heparin (MOPS-Cl-HMWH; Figure 2). We also observed tubes, ribbons, and rolled sheets for Ac10VME, AcPHF6*, and Ac375KLTFR. TEM data for these peptides can be found in Table 1.

TEM is perhaps the most sensitive measure of the ability of a peptide to form amyloid since the hydrophobic nature of amyloid structures are attracted to the hydrophobic surface of the carbon-coated TEM grid. Previously, we observed amyloid fibers formed from peptides containing proline, a known β -sheet breaking amino acid residue. However, it was later determined by other measurements that the majority of the peptide existed in a random coil conformation. ⁴⁰ Hence, the TEM experiment is biased toward the observation of amyloid-like structures, even when only a relatively small fraction of the sample forms amyloid. Equally important are those sequences that were not observed to form amyloid structures by TEM. These sequences can be eliminated as nucleating sequences and are less likely to affect the kinetics of amyloid formation by their interaction with other nucleating sequences in the protein.

CD of Predicted Amyloid Forming Sequences in Tau. We used CD spectroscopy to study the conformation of the predicted amyloid forming sequences in tau. In water, the CD of AcPHF6 (Figure 3A) and AcPHF6* (not shown) displayed a pattern characteristic of random coil peptides with near zero ellipticity at higher wavelength and a single negative band present near 195 nm. Under these conditions these nucleating peptides possess a very low fraction of characteristic of aggregating species having a significant fraction of β -sheet structure. In

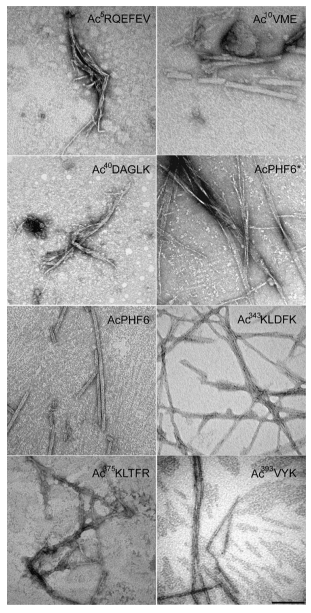


Figure 2. TEMs of high propensity peptide sequences in MOPS-Cl-HMWH buffer. Scale bar represents 100 nm.

MOPS-Cl-HMWH, the CD spectra of AcPHF6* and AcPHF6 showed classical spectra of extended β -sheet, with a negative nπ* band at 217 nm and a positive ππ* band near 198 nm. ^{23,61} With the exception of Ac³⁹³VYK and Ac¹⁰VME, other peptides studied showed patterns characteristic of random coil peptides in MOPS-Cl-HMWH. The tripeptides Ac³⁹³VYK and Ac¹⁰VME displayed a CD spectra which appeared to be a combination of spectra arising from β -sheet and random coil structures, having a negative shoulder between 215 and 220 nm and a strong negative band below 198 nm.

We used CD to study the interaction between peptides in MOPS-Cl-HMWH. Figure 3B shows the CD of AcPHF6, Ac¹⁰VME, a 1:1 mixture of the two peptides, and the CD that would be calculated from the spectra of the peptides by themselves, in the absence of any interaction. As expected, there is a negative 215 nm band for the calculated spectra for AcPHF6 and Ac¹⁰VME (dashed line), with an intensity that is halfway between that for the spectra of the two peptides alone. In contrast, the observed

Table 1. TEM Results for Tau-Related Peptides^a

peptide	amyloid structures	fiber/tube diam (nm)
Ac ⁵ RQEFEV	fiber bundles	2.7 (±0.4)
Ac ¹⁰ VME	tubes	20.3 (±1.5)
Ac ¹⁷ TYGLG	n.o.	
Ac ²⁸ GYTMHQDQ	n.o.	
Ac ⁴⁰ DAGLK	fiber bundles	$4.0 \ (\pm 0.7)$
Ac ¹²⁶ RMV	n.o.	
Ac ²²⁵ KVAVVR	n.o.	
Ac ²⁴² RLQ	n.o.	
AcPHF6*	rolled sheets, fibers, ribbons	5.5 (±1.3)
Ac ²⁸² LDLSN	n.o.	
AcPHF6	fibers	6.5 (±1.2)
Ac ³³⁷ VEVK	n.o.	
Ac ³⁴³ KLDFK	fibers	$3.3 (\pm 0.4)$
Ac ³⁷⁵ KLTFR	rolled sheets, fiber bundles	$4.0 \ (\pm 0.9)$
Ac ³⁸² AKAK	n.o.	
Ac ³⁹³ VYK	fibers	$5.3 (\pm 0.4)$
Ac ⁴¹⁹ MVD	fibers, spherical aggregates	6.8 (± 0.4)
Ac ⁴²⁸ LAD	n.o.	
Ac ⁴³² VSASL	n.o.	

"Results labeled as "n.o." indicate no amyloid structures were observed.

CD spectrum for a 1:1 mixture of the two peptides (solid line) has a negative 215 nm band with an intensity significantly more negative than the calculated spectrum, suggesting that $Ac^{10}VME$ interacts with AcPHF6 to shift the equilibrium toward the aggregated form. It is also noteworthy that the observed spectrum of the 1:1 mixture shares an isodichroic point at 205 nm with the spectra of the two isolated peptides, suggesting that only two forms of the peptides, monomeric and aggregated, are present. A plausible model for this type of interaction is one in which $Ac^{10}VME$ participates in the same β -sheet structure characteristic of the AcPHF6 amyloidogenic state. In contrast, the spectrum of a AcPHF6:AcPHF6* mixture (Figure 3C) does not appear to share an isodichroic point with spectra of the two pure peptides, suggesting a type of interaction involving something more complex than a simple two-state model.

We define a positive interaction as one in which the observed CD of a 1:1 mixture is more negative above (higher wavelength) the isodichroic point and more positive below this point. In contrast, a negative interaction implies that a smaller fraction of the sample exists in an aggregated state when the two peptides are mixed. A summary of the results is presented in Figure 3D. The greatest positive interactions with AcPHF6 result from peptides Ac¹⁰VME, AcPHF6*, Ac³⁷⁵KLTFR, and Ac³⁹³VYK while the greatest positive interactions with AcPHF6* result from AcPHF6 and from C-terminal sequences Ac³⁷⁵KLTFR, Ac³⁹³VYK, Ac⁴¹⁹MVD, Ac⁴²⁸LAD, and Ac⁴³²VSASL. It should be noted that interactions evaluated in this manner are a reflection of pairwise interactions at thermodynamic equilibrium and do not necessarily reflect how these interactions will affect aggregation kinetics.

Kinetics of Tau Peptide Aggregation. The kinetics of aggregation of tau peptides was measured in MOPS-Cl-HMWH by following the time dependence of ThS fluorescence at 490 nm and fitting the data to Gompertz growth curves. 20,40,41,45,47,48 Under these conditions only AcPHF6 and AcPHF6* displayed measurable aggregation ($k_{\rm app}=(23.3\pm1.4)\times10^{-3}~{\rm s}^{-1}$ and $(9.2\pm0.7)\times10^{-3}~{\rm s}^{-1}$, respectively). We also measured aggregation of these two peptides in the presence of the other

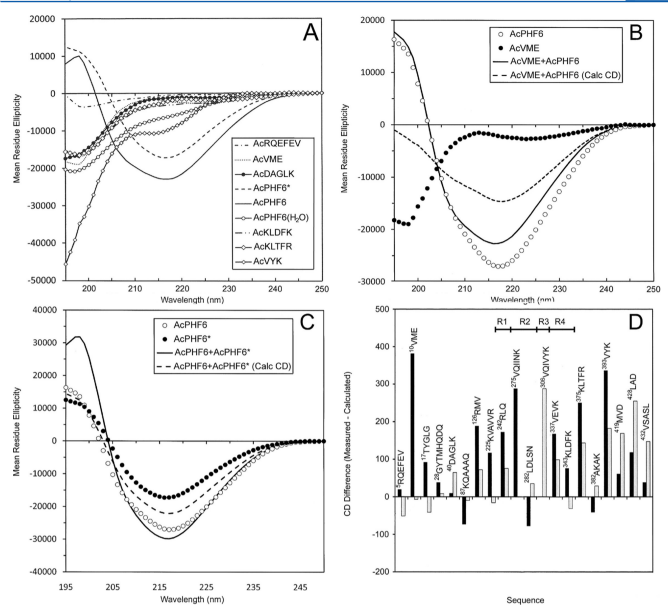


Figure 3. (A) Far-UV CD spectra of 100 μ M N-acetylated peptide amides in water or MOP-Cl-HMWH buffer (pH 7.2). (B) CD of AcPHF6, Ac¹⁰VME (MOPS-Cl-HMWH), the calculated mean residue ellipticity of a 1:1 stoichiometric mixture, and the observed ellipticity of the mixture. (C) as in (B) for AcPHF6 and AcPHF6*. (D) Difference between calculated and observed ellipticity for the peptides. A positive interaction is one in which the observed CD of a 1:1 mixture is more negative above (higher wavelength) the isodichroic point and more positive below this point. In contrast, a negative interaction infers that a smaller fraction of the sample exists in an aggregated state when the two peptides are mixed. Black and gray shading refer to interactions with AcPHF6 and AcPHF6*, respectively.

predicted amyloidogenic peptides by mixing peptides at a 1:1 stoichiometric ratio. Only Ac²⁷⁵KLTFR had any significant effect on the aggregation ($p \le 0.05$), slowing the aggregation kinetics of both AcPHF6 and AcPHF6* ((19.4 \pm 0.5) \times 10⁻³ $\rm s^{-1}$ and $(3.9 \pm 0.6) \times 10^{-3} \rm s^{-1})$. The effect of Ac²⁷⁵KLTFR on the aggregation of AcPHF6* is shown in Figure 4A as a function of increasing Ac²⁷⁵KLTFR:AcPHF6* stoichiometry. At less than a 1:1 stoichiometric ratio Ac²⁷⁵KLTFR acts as an aggregation inhibitor, affecting both the $k_{\rm app}$ and the lag time, while at a 2:1 Ac²⁷⁵KLTFR:AcPHF6* stoichiometric ratio there are two discernible aggregation events. Plots of $\log(k_{
m app})$ and lag time obtained from fitting vs Ac²⁷⁵KLTFR:AcPHF6* stoichiometry for the first aggregation event showed a linear relationship (Figure 5; $r^2 \ge 0.97$), suggesting that both nucleation and elongation are affected in a nucleation-elongation mechanism.

We found it surprising that Ac^{275} KLTFR was an inhibitor of AcPHF6 and AcPHF6* aggregation but that Ac^{343} KLDFK, a peptide whose sequence is homologous, was not. We hypothesized that it may be the charge—charge repulsion between aspartic acid side chains that led to its inability to fit into the AcPHF6 or AcPHF6* steric-zipper structure. To test this hypothesis, we measured the aggregation of AcPHF6* at different stoichiometric ratios of Ac^{343} KLDFK:AcPHF6* in phosphate buffer at pH 3.0, where the aspartic acid carboxyl is uncharged (Figure 4B). Under these conditions Ac^{343} KLDFK inhibits the aggregation of AcPHF6* in a dose-dependent manner, similar to the inhibition by Ac^{275} KLTFR at higher pH. A plot of $log(k_{app})$ vs Ac^{343} KLDFK:AcPHF6* stoichiometry was linear (Figure 5A), but we did not observe a linear dependence in lag time with increasing Ac^{343} KLDFK concentration.

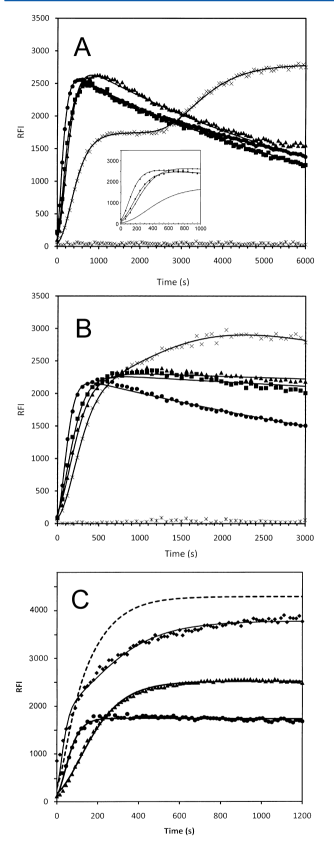


Figure 4. Aggregation kinetics of peptides and peptide mixtures in MOPS-Cl-HMWH followed by fluorescence of ThS at 490 nm (440 nm excitation). AcPHF6* was 100 μ M. (A) AcPHF6* (filled circles), AcPHF6*:Ac³⁷⁵KLTFR at 2:1 (filled squares), 1:1 (filled triangles), and 1:2 stoichiometry (crosses). Solid lines represent best fits to the data, assuming a single Gompetz growth curve model for pure AcPHF6* and AcPHF6*:Ac³⁷⁵KLTFR mixtures at less than 1:1

Figure 4. continued

stoichiometry. The AcPHF6*:Ac³⁷⁵KLTFR mixture at 1:2 stoichiometry was modeled using a sum of Gompetz growth curves (eq 2). (B) as in (A) with AcPHF6* and AcPHF6*:Ac³⁴³KLDFK mixtures in phosphate-Cl buffer at pH 3.0. (C) Aggregation kinetics of AcPHF6 (filled circles), AcPHF6* (filled triangles), and a 1:1 stoichiometric mixture of AcPHF6:AcPHF6*. Solid lines represent best fits to a single Gompetz growth curve, in the case of pure peptides, or a sum of two Gompetz growth curves, in the case of the mixture. The dashed line represents a sum of fitted curves for AcPHF6 and AcPHF6*, assuming no interaction between the peptides.

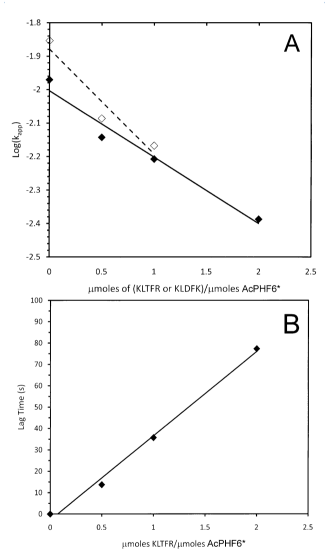


Figure 5. Plots of parameters obtained from fitting kinetic data for AcPHF6* and AcPHF6*:Ac 375 KLTFR and AcPHF6*:Ac 343 KLDFK mixtures. (A) Log($k_{\rm app}$) for first aggregation event of AcPHF6* and AcPHF6*:Ac 375 KLTFR mixtures at pH 7.2 (filled diamonds) and for AcPHF6* and AcPHF6*:Ac 343 KLDFK mixtures (exempting AcPHF6*:Ac 343 KLDFK at 1:2 stoichiometry) at pH 3.0 (open diamonds). (B) Lag time for first aggregation event of AcPHF6* and AcPHF6*:Ac 375 KLTFR mixtures. Data could be fit with an $r^2 > 0.97$.

Figure 4C shows the effect that mixing AcPHF6* with AcPHF6 at 1:1 stoichiometry has on aggregation kinetics. Using the same fitted kinetic parameters that we obtained from the kinetics of AcPHF6 and AcPHF6* aggregation, we simulated the aggregation

of the AcPHF6:AcPHF6* mixture from a sum of growth curves (dashed line). While the curve comes close to the observed data at short times, it deviates from the data at longer time points. The data could be fit to a sum of growth curves by varying the kinetic parameters for both AcPHF6 and AcPHF6* (solid line), and we found the best fit could be obtained with a significantly faster $k_{\rm app}$ for AcPHF6* ((46 ± 3) × 10⁻³ s⁻¹) and a significantly slower $k_{\rm app}$ for AcPHF6* ((5.6 ± 0.6) × 10⁻³ s⁻¹), again suggesting a complex kinetic interaction between these two peptides.

Aggregation Kinetics of 2N4R Tau Mutants. While the CD interaction experiments and kinetic studies on predicted amyloidogenic tau peptides suggest that interactions between sequences may effect tau aggregation rate or the overall fraction of tau participating in an aggregated state, they do not address whether such interactions take place intra- or intermolecularly or what additional constraints the full protein may impose. To address this issue, we prepared three single site mutants of tau (M11K, F378K, and Y394K) that were predicted to decrease the amyloidogenic propensities of ¹⁰VME, ²⁷⁵KLTFR, and ³⁹³VYK by substituting a positively charged lysine for a hydrophobic residue.⁴⁰ Kinetics of polymerization for 2N4R lysine mutants were assayed by right angle laser light scattering against wildtype tau (Figure 6A). M11K showed a significantly longer lag time than WT (12.69 \pm 3.41 min compared to 7.20 \pm 0.61 min; $p \le 0.05$) while F378K and Y394K showed no significant difference (7.41 \pm 0.07 and 7.22 \pm 0.42 min respectively). The $k_{\rm app}$ for M11K decreased significantly (0.09 \pm 0.03; $p \leq$ 0.05), while the k_{app} for F378K (0.17 \pm 0.05) and Y394K (0.15 \pm 0.007) were slightly larger than k_{app} for WT (0.13 \pm 0.02). M11K, F378K, and Y394K showed a trend for slightly higher maximum light scattering values (126.63 \pm 18.92, 110.50 \pm 13.55, and 113.87 \pm 15.16, respectively) than WT (95.40 \pm 5.22). TEM images, however, showed no difference in the morphology of filaments (Figure 6B-E). Quantitation of filaments revealed M11K had significantly fewer filaments, while F378K had significantly more filaments ($p \le 0.0001$) and Y394K had approximately the same number as compared to WT (Figure 6F). Filament lengths were only increased in M11K as compared to WT, with F378K and Y394K having the same average length of filaments per field (Figure 6G). The mass of filaments per field (length × number) was similar to the changes seen for the number of filaments per field (Figure 6H).

DISCUSSION

Whether or not proteins form amyloid depends on the presence of nucleating sequences or "hot spots", short sequences of amino acids that have a high propensity (HP) to form parallel or antiparallel β -sheet structure between protein monomers. ^{62–65} However, the presence of these nucleating sequences is not a sufficient condition for oligomerization, particularly for most proteins where HP sequences occur on the interior of the protein. ^{63,66} Perturbation of the protein away from its native conformation, either by partial denaturation or by the presence of mutations, controls the exposure of these sequences to the solvent and thus controls their ability to interact intermolecularly with other monomers and form toxic oligomers or filaments. ^{62,64} In "natively unfolded" proteins or peptides, HP sequences may inhibit the formation of amyloid, either by controlling the folding of the backbone or by forming intramolecular β -structure. ^{62,67,68} Tau is a loosely folded flexible protein with two nucleating sequences, PHF6* and PHF6 at the beginning of R2 and R3 in the MTBR. ^{19,31,69} The PHF core is formed by only a portion of

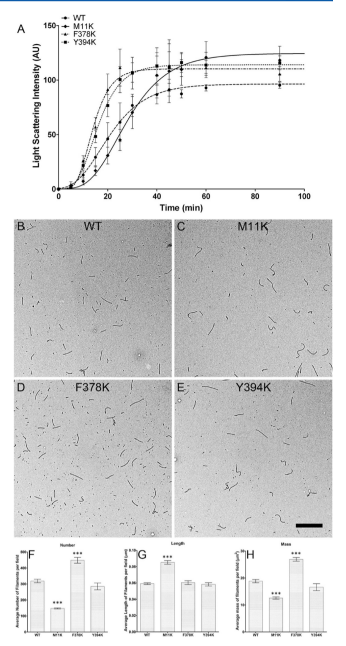


Figure 6. 2N4R lysine mutants show variations from WT tau. Mutant forms (M11K, F378K, and Y394K (diamond, triangle, and square, respectively)) and WT (circle) 2N4R tau were assayed by LLS over the course of 240 min (first 100 min shown) (A). Data were fit to the Gompertz growth curve (M11K, F378K, Y394K, and WT (solid, long/short dash, short dash, and long dashed lines, respectively)). Representative TEM images of WT (B), M11K (C), F378K (D), and Y394K (E) were collected at 3600× magnification. Images were quantitated using Image Pro Plus for filament number (F), length (G), and mass (H). Data in (F–H) are averages of 5 fields \pm SEM ($p \le 0.0001$ for bars marked with asterisks). Scale bar represents 1 μ m.

tau near the C-terminal, including the MTBR, while N-terminal forms the outer "fuzzy coat" of PHF. 14,70–72 Shorter tau related peptides containing PHF6 and PHF6* aggregate much more rapidly than full length tau, suggesting that other segments of the full length protein assume a regulatory role by interacting intramolecularly with the MTBR region. 23,40,73 Other segments of tau distant from the nucleating PHF6 and PHF6* regions, such as the N-terminal inserts, tend to promote aggregation. 33,38

Given the evidence suggesting that regions of tau remote from the MTBR are instrumental in regulating polymerization, we were curious to see if (1) HP sequences, other than PHF6 and PHF6*, could nucleate filament formation and (2) HP sequences could interact with the known nucleating sequences PHF6 or PHF6* to increase the fraction of β -sheet structure present or to control aggregation kinetics. We scanned tau for HP motifs and quantitated their propensities using an algorithm previously described by us. 40 Other algorithms exist for determining sequence amyloidogenic propensity based on sequence or on their interaction energy with other nucleating sequences. 58,74-77 While they are able to find PHF6 and PHF6* in tau, no other potential HP regions have been reported. Our model flagged 20 HP sequences, including 87KQAAAQ, ²²⁵KVAVVR, and ³³⁷VEVK, in addition to PHF6 and PHF6*. These sequences have previously been shown to populate β sheet structure by NMR studies of full length tau protein.³¹ In summary, our simple method of finding potential HP regions of tau is capable of locating sequences that other more complex algorithms cannot, although it may be that our method is so sensitive that it flags sequences which neither form amyloid nor interact with other known nucleating sequences.

We used TEM to experimentally evaluate the amyloid forming propensity of N-acetylated peptide amides with sequences identical to the 20 HP sequences in tau flagged by our algorithm. Filaments and tubes were observed for N-terminal peptides $Ac^{5}RQEFEV^{10}$ and $Ac^{10}VME$. In the past, a $\Delta 2-18$ mutant has been shown to aggregate less effectively than wild-type tau, suggesting that residues in the N-terminal help seed aggregation in the presence of arachidonic acid inducer.³³ Tau with frontotemporal dementia mutation R5L aggregates faster than wild-type tau and forms 4.5 times more filaments, again emphasizing the role this segment plays in oligomerization.³ We observed rolled sheets, fibers, and ribbons for AcPHF6* and filaments for AcPHF6 and Ac343KLDFK. These sequences occur in the MTBR and have previously been shown by NMR to populate β -sheet structure. We observed no amyloid structures for Ac87KQAAAQ, Ac225KVAVVR, or Ac337VEVK, sequences which were observed to populate β -structure in the NMR of full length tau. 31 We observed rolled sheets and fiber bundles for Ac³⁷⁵KLTFR, a sequence in tau which lies just downstream of R4, and has been shown by NMR to be in an inflexible region of PHFs.⁶⁹ We have previously shown that Ac³⁹³VYK forms filaments, and mixtures of AcPHF6 and Ac393VYK form twisted filaments with identical morphology to bona fide PHFs.²³ The ³⁹²IVYK sequence likely is the nucleation sequence for the amyloid formed from the 383-337 C-tail 8 peptide, previously shown to possess a high degree of β -sheet structure and form filaments in the absence of polyanion inducer.³⁷ We also observed fibers and spherical aggregates for the peptide Ac⁴¹⁹MVD. Caspase-3 mediated cleavage after Asp421 can be detected in mild cognitive impairment and AD but not in nondemented control brains. It is tempting to speculate that C-terminal truncation at this site may allow the 419MVD sequence sufficient solvent accessibility to help seed oligomerization.

Only AcPHF6 and AcPHF6* showed measurable aggregation kinetics as well as a high fraction of β -structure at equilibrium determined by CD. Of the peptides shown by TEM to form amyloid, Ac¹⁰VME, AcPHF6*, Ac³⁷⁵KLTFR, and Ac³⁹³VYK were found to enhance the fraction of β -structure of AcPHF6 formed at equilibrium, and Ac³⁷⁵KLTFR was found to affect AcPHF6 and AcPHF6* aggregation kinetics. Others have shown that interactions between PHF6 and PHF6* exist in tau or tau-

related peptides. 19-22,27,28 Ac375KLTFR slowed the aggregation kinetics of AcPHF6 by roughly 15% but slowed the $k_{\rm app}$ for AcPHF6* aggregation by 60% at 1:1 stoichiometry. We obtained linear plots of $\log(k_{\rm app})$ or lag time for the first aggregation event versus the Ac³⁷⁵KLTFR/AcPHF6* ratio, suggesting that Ac375KLTFR acts as a competitive inhibitor of AcPHF6* aggregation. Given the sequence homology between Ac³⁷⁵KLTFR and Ac³⁴³KLDFK, we found it surprising that Ac343KLDFK was not also an inhibitor of AcPHF6* aggregation. However, we found that Ac343KLDFK was an inhibitor at pH 3.0 where the carboxyl of the aspartic acid side chain is protonated. Apparently, its limited ability to inhibit AcPHF6* aggregation arises from unfavorable interaction between the charged carboxylate side chains in the peptide at neutral pH. Both Ac343KLDFK and Ac375KLTFR satisfy a model based on the steric zipper where they occur in alternating layers with AcPHF6*. A hypothetic model shows antiparallel layers of parallel in-register peptide β -structure (Figure 7). ^{24,25}

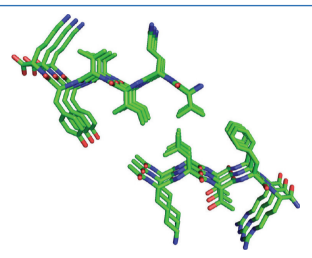


Figure 7. Hypothetical model for AcPHF6 and Ac³⁷⁵KLTFR in hybrid steric zipper. At 1:1 stoichiometry antiparallel layers of parallel inregister chains of AcPHF6 and Ac³⁷⁵KLTFR continue to form amyloid, albeit at a reduced rate of aggregation.

Although data from studies with HP peptides suggest that the sequences ¹⁰VME, ³⁷⁵KLTFR, and ³⁹³VYK have the potential of interacting with PHF6 and PHF6* to enhance the fraction of amyloid present or inhibit the rate of aggregation, it is difficult to predict how these interactions will influence tau aggregation without having some idea of solvent accessibility. Intramolecular interactions which enhance amyloid formation might be expected to inhibit oligomerization while similar intermolecular interactions would be expected to enhance it. In the past, we and others have shown that two consecutive C-terminal positively charged residues transform an amyloidogenic sequence into a non-amyloidogenic one. 40,41,55 In the case of the M11K mutant, fewer filaments were observed, and there was an increase in lag time and a decrease in the rate of aggregation compared to wild-type tau, suggesting that the N-terminal sequence ¹⁰VME enhances arachidonic acid induced nucleation and elongation. In contrast, the F378K mutant showed an increased rate of aggregation and more filaments compared to wild-type tau. Because of the proximity of the ³⁷⁵KLTFR sequence to PHF6 and PHF6*, it is tempting to speculate that the region just downstream of MTBR interacts intramolecularly with these nucleating sequences, protecting them from interacting intermolecularly.

The Y394K mutant produced about the same number of filaments and had a similar rate of aggregation as wild-type tau. This result is consistent with NMR evidence which shows that 392 IVYK does not form β -structure in full length tau. 31,37,59

In conclusion, we have shown that HP sequences in tau other than PHF6 and PHF6* can nucleate amyloid formation and can help regulate the kinetics of amyloid formation. That being said, amyloid nucleation is a complex event in tau, governed by the folding of the peptide backbone, phosphorylation of amino acid side chains, environmental conditions (reducing or non-reducing, salt concentration, pH), the nature of polyanionic or long chain fatty acid inducers, and other HP sequences remote from the primary nucleating site. These secondary HP sequences in tau or other proteins may have evolved in order to protect biological systems from protein oligomerization and toxicity.

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ABBREVIATIONS

PHF6*, ²⁷⁵VQIINK; PHF6, ³⁰⁶VQIVYK; PHF, paired helical filament; MTBR, microtubule binding region; R1–R4, 32 amino acid pseudorepeats 1–4 in the MTBR; DTT, dithioerythritol; MOPS, 3-[N-morpholino] propanesulfonic acid; HMWH, hight molecular weight heparin; ThS, thioflavin S; EDTA, ethylenediaminetetracetic acid; SEM, standard error of the mean; HP, high propensity; CD, circular dichroism; TEM, transmission electron microscopy; MALDI-TOF MS, matrix-assisted laser desorption/ionization mass spectrometry; ESI, electrospray ionization; FRET, fluorescent resonant energy transfer; PC1 scores, principal component scores; LLS, right angle laser light scattering.

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