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Specific phosphorylation patterns control the interplay between aggregation and condensation of Tau-R4 peptides†

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Specific phosphorylation patterns regulate the activity of proteins and play a central role in protein self-assembly. In Tau, such patterns drive the formation of disease-related condensates and aggregates. Understanding their functional impact is essential for studying Tauopathies such as Alzheimer's Disease. Here we show how specific phosphorylation patterns regulate Tau self-assembly and control the interplay between its aggregation and condensation, using a peptide-based approach that allows systematic analysis of libraries of specific phosphorylation patterns at the domain level and is complementary to the current protein-level methods. We applied our methodology to study the effect of specific phosphorylations on the aggregation and condensation of the R4 domain of Tau that is pivotal for its self-assembly, forming the β -helix motif that is common to various Tau patient fibrils. Using advanced phosphopeptide synthesis methods developed in our labs, we generated a library of multi-phosphorylated peptides derived from Tau R4. We found that phosphorylation at Ser341 promotes aggregation, while Ser352 enhances condensation. Phosphorylation at Ser356 inhibits both processes. The source of these different outcomes is the distinct microenvironments around each phosphorylated site. Our results provide a residue-level resolution of how the decision between Tau condensation and aggregation is being made. This was possible by using our peptide-based approach, which is complementary to protein-level method and enables efficient identification of active phosphorylation patterns. These can later be studied at the protein level.

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Introduction

Phosphorylation is one of the most abundant posttranslational modifications (PTM) in eukaryotes.^{1,2} Phosphorylation has a key role in regulating protein-protein interactions (PPI) and self-assembly of monomeric protein to oligomers.^{3–5} The most prevalent phosphorylation sites occur on serine and threonine, and most phosphoproteins are phosphorylated at multiple sites.^{4,6}

Proteins can oligomerize to different types of assemblies, ranging from coacervates formed *via* liquid-liquid phase separation (LLPS) to aggregates of different types such as amorphous, native-like, or amyloid fibrils.⁷ In cells, coacervates function as membrane-less organelles, which have several possible functions such as bioreactors that catalyze protein interactions and cellular processes.^{8–10} Aggregation, however, is frequently associated with pathological pathways that lead to disease.¹¹

Different phosphorylation patterns can regulate protein activity and dictate which pathway the protein will undergo. They regulate PPI and may control the oligomerization type of proteins, selecting between aggregation and condensation.^{3,12} The existence of multiple phosphorylation sites in proteins means that there are many possible phosphorylation patterns, each with a unique effect on oligomerization. The ideal way for studying the effect of each such pattern is by obtaining a library of multi-phosphorylated derivatives of the protein of interest, each with a unique and specific phosphorylation pattern. Such a library would cover all relevant patterns. However, obtaining such a diverse multi-phosphorylated protein library in sufficient quantities for extensive biophysi-

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cal, biochemical and structural studies is difficult. The methods for obtaining recombinant phosphorylated proteins for *in vitro* biophysical, biochemical and structural studies include kinase-mediated phosphorylation, the use of “phosphomimetic” amino acids and semisynthetic methods. These methods are complementary, each having its pros and cons, raising the need for using more than one method to obtain the full picture. There is thus a need for additional methods to be added to the existing toolbox.

Replacing phosphorylated amino acids with other negatively charged residues is very popular.^{13,14} This common substitution of phosphoserine by negatively charged glutamate or aspartate (commonly referred to as phosphomimic residues) allows for performing large scale studies at the protein level.¹⁵ This is an easy and fast method to make libraries. However, using this method results in a significant compromise as there is a crucial difference between a phosphate group and a carboxylic group. Replacing a phosphoserine or phosphothreonine by glutamate or aspartate retains only one negative charge at the specific site but overlooks the unique chemical and physical properties of the phosphate group.¹³ A phosphate has two negative charges at physiological pH, while glutamate has only one. Moreover, the pK_a of phosphate is 5.58, while the pK_a of glutamate is 4.25.¹⁶ The phosphate also occupies much larger volume that is derived from its tetrahedral geometry compared to the planar carboxylate group. These factors govern the distinctive interplay between the phosphorylated residue and its surroundings, dictating the role of the phosphate at each specific microenvironment within the protein. Comparative studies showed that Glu substitution of the phosphorylated Ser did not necessarily result in the same protein structure or function.^{17,18} It was demonstrated for α -synuclein that while phosphorylation of Ser129 inhibited fibrillation, mutation to Asp or Glu did not reproduce this activity.¹⁹ This difference is prevalent in the study of condensation, as it is a highly charge-dependent process.²⁰ *In vitro* phosphorylation of recombinant proteins by kinases although widely used, introduces constraints, as it usually results in a mixture of proteins with different phosphorylation patterns that are difficult to isolate and analyze, limiting our understanding of their precise functional roles.²¹ Enzymatic phosphorylation by different kinases is not totally selective.²¹ A prior knowledge of each specific kinase is needed for achieving the desired specific phosphorylation is required.¹⁸ The above limits the access to many possible phosphorylation patterns that can be of high importance. To obtain the specific phosphorylation pattern with high selectivity and high yields, chemical peptide synthesis should rather be used.

Semisynthetic strategies, which include the chemical synthesis of a phosphorylated domain followed by its ligation to a recombinant protein (such as native chemical ligation), have the advantage of introducing the actual phosphorylation pattern into the protein. However, these approaches are time consuming, low yielding processes that provide only small quantities of the desired proteins and are completely impractical for the preparation of libraries with phosphorylation

patterns.^{22,23} Thus, while excellent for studying a specific pattern at the protein level, these approaches are not suitable for identifying which pattern is the active one since it is practically impossible to produce libraries and systematically study them using the mentioned semisynthetic methods. Most importantly, protein synthesis depends on the availability of the corresponding multiphosphopeptides which is still a significant problem. Since the peptides are later incorporated into the proteins using NCL, the successful synthesis of the peptides is crucial for the success of the NCL later and is another layer in the process. If the peptides are difficult to make, the proteins could not be made either. The synthesis of proteins hence depends on the development of synthetic methods to prepare multi-PTM peptides and demonstrate their relevance in studies such as the one presented here and as described below.

The synthesis of long, multiphosphorylated peptides remains challenging due to the steric bulk and lability of protected phospho-serine and phospho-threonine residues under standard Fmoc conditions. These factors limit coupling efficiency and overall yield. Despite these hurdles, we have established robust synthetic method that enables the efficient production of phosphorylated peptide libraries derived from aggregation-prone domains.^{24–27} The method is based on a combination of high temperature and fast mixing for accelerating each coupling step while minimizing the formation of side products and maximizing the efficiency.²⁵ Using this method, we demonstrated the synthesis of peptides containing up to eight phosphorylation sites. The above strategy is ideal for synthesizing libraries of multiphosphorylated peptides. These libraries can then be used for systematic studies of the effect of specific phosphorylation patterns on the biological activity of proteins. Here we applied our method for studying the aggregation processes of the Tau protein.^{28,29}

The Tau protein forms amyloid fibrils in neurons as part of the pathology of neurodegenerative diseases (ND). *in vivo*, fibrils of Tau are hyperphosphorylated,³⁰ an event that is linked to aggregation and condensate formation by the protein.^{21,31–34} However, the role of the specific phosphorylation patterns in controlling the initial molecular events leading to amyloid aggregates or condensates is mostly unclear. Tau is a highly disordered protein that is composed of an N-terminal domain, a proline-rich domain, microtubule binding domain (RD) with five homologous repeats (R1–R4, R') and a C-terminal part (Fig. 1A). In Tauopathies such as Alzheimer's disease (AD) and chronic traumatic encephalopathy (CTE), Tau undergoes aggregation into amyloid fibrils *via* its RD. Fibrils from AD and CTE patients share a hairpin with a hydrophobic core that is formed by the fourth repeat (R4) of Tau-RD (β -helix; Fig. 1B). In fibrils, R4 has three beta-sheets, interconnected by disordered linkers. In addition, Tau-RD is essential for the condensation of Tau.^{34,35} Although it is known that phosphorylation of the full-length Tau initiates aggregation and condensation,^{21,36} there is almost no information about the effect of specific phosphorylations, mostly due to the difficulty in obtaining Tau derivatives with specific



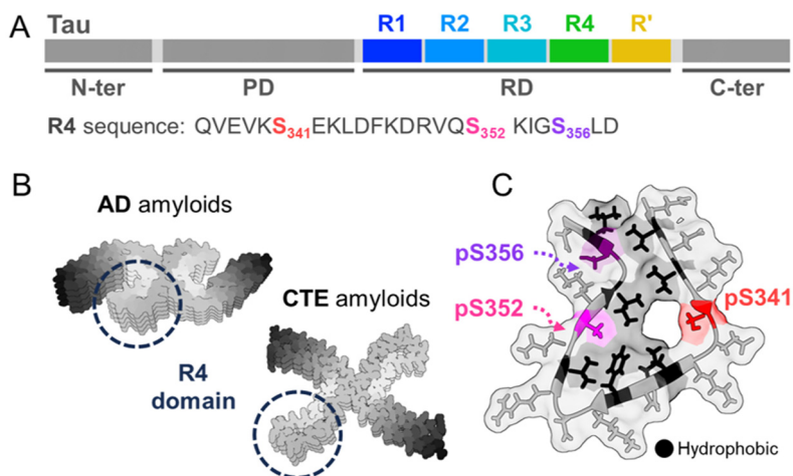


Fig. 1 The Tau R4 domain (residues 336–358). Tau 336–358 is derived from the R4 repeat and forms a hairpin structure within native Tau fibrils. (A) Domain structure of Tau: the N-terminal part (N-ter), the proline-rich domain (PD), the repeat domain (RD) with its five repeats R1–R4, R' (highlighted), and the C-terminus part (C-ter). The sequence of Tau R4 (serine 341 in red, 352 in magenta and 356 in purple). (B) Cryo-EM structures of Tau filaments isolated from patients of different Tauopathies: AD amyloid filament (PDB ID: 5O3T),³⁸ CTE amyloid filaments (PDB: 6NWQ),³⁹ Tau R4 that adopts a hairpin structure is framed. (C) The hairpin structure that Tau R4 forms in AD PHF patients. Serine 341 labeled in red, 352 in magenta and 356 in purple. hydrophobic residues in black.

phosphorylation patterns. For example, single phosphorylations at Ser293 in R2 or Ser305 in R3 promote amyloid fibril formation *in vitro*.³⁷ However, the effect of specific phosphorylation patterns on oligomerization of R4 is yet unknown. R4 has three sites that were found to be phosphorylated in ND: Ser341, Ser352 and S356. Each of these serine residues is located at a different chemical environment. Ser341 is located between two charged residues, Lys340 and Glu342, within the disordered linker. Ser352 is positioned next to Gln351 and Lys353 and is located within a loop structure, while Ser 356 is located between two aliphatic residues, Gly355 and Leu357 at the end of the loop of R4. The side chains of Ser341 and Ser352 are facing toward the core of the hairpin, while Ser356 is facing out (Fig. 1C).

Due to the challenges associated with synthesizing full-length Tau protein, studies of Tau self-assembly often rely on the use of Tau-derived fragments. These fragments vary in length, from short hexapeptides such as those derived from key amyloidogenic motifs in the R3 and R2 domains⁴⁰ to larger constructs of ~100 residues, including the First Intermediate Amyloid (FIA) fragment⁴¹ and well-established aggregation models such as K18 and K19.⁴² Peptides are vastly used to explore molecular mechanisms of canonical domains that undergo phase separation.^{43,44} Peptides have several advantages in bridging the gap between the authentic yet inaccessible multiphosphorylated proteins to the much-needed biophysical and biochemical input. First, synthesis of peptides can provide each specific pattern at high purity, specificity and reliability. Second, synthetic processes guarantee that the peptides can be obtained reproducibly and at a sufficient quantity. Third, peptides can be labeled by different chemical tags to enable analysis *via* diverse biochemical assays. Most importantly: they provide an easy and accessible route to libraries.

Here we harnessed our novel approaches for multiphosphorylated peptide synthesis^{12,24,45} to study systematically how specific phosphorylation pattern affect aggregation and condensation of the Tau 336–358 (Tau R4 domain). We designed and synthesized a library of peptides representing the R4 domain and containing all the possible phosphorylation patterns. Then, we determined whether each peptide undergoes condensation or aggregation by fluorescence and electron microscopy. We found that the fully phosphorylated peptide can condensate or aggregate depending on the conditions, while the non-phosphorylated peptide cannot oligomerize at all. Phosphorylation of Ser341 promoted aggregation, while phosphorylation of Ser352 promoted condensation. Our research emphasizes the importance of a methodological study that explores the effects of phosphorylation patterns on oligomerization using phosphorylated peptide libraries, as a complementary method for the other protein-level methods that are currently in use.

Results

Synthesis of a phosphorylated peptide library derived from Tau R4

To study the effect of phosphorylation on the condensation and aggregation of R4, we synthesized a library of peptides with patterns including all combinations of phosphorylations in Ser341, Ser352, and Ser356. The peptides were synthesized using our newly developed Accelerated MPP synthesis,²⁴ which enabled us to synthesize the library in a short time and in sufficient amounts (see Table 1 for peptide sequences and ESI Fig. 1 and 2† for the analytical data of the peptides). The synthesis was performed with a constant overhead stirring of 1200



Table 1 Library of Tau R4 and its derived phosphorylated peptides

Peptide	Name	Sequence	Net charge
Tau336–358	0P	Q VEVKSEK <u>LD</u> FDKDRVQ SK IGSLD	0
Tau336–358(pS341)	1Pa	Q VEVK p SEK <u>LD</u> FDKDRVQ SK IGSLD	–2
Tau336–358(pS352)	1Pb	Q VEVKSEK <u>LD</u> FDKDRVQp SK IGSLD	–2
Tau336–358(pS356)	1Pc	Q VEVKSEK <u>LD</u> FDKDRVQ SK IGpSLD	–2
Tau336–358(pS341,352)	2Pa	Q VEVK p SEK <u>LD</u> FDKDRVQp SK IGSLD	–4
Tau336–358(pS341,356)	2Pb	Q VEVK p SEK <u>LD</u> FDKDRVQ SK IGpSLD	–4
Tau336–358(pS352,356)	2Pc	Q VEVKSEK <u>LD</u> FDKDRVQp SK IGpSLD	–4
Tau336–358(pS341,352,356)	3P	Q VEVK p SEK <u>LD</u> FDKDRVQp SK IGpSLD	–6

Hydrophobic residues are highlighted in yellow, positively charged residues in blue, and negatively charged residues in red. Serines are shown in bold black, with phosphorylated serines underlined in red. All peptides synthesized both with and without fluorescein labeling. Labeled peptides comprising herein a prefix of FL before their name.

rpm, at 90 °C. The assembly of each peptide was completed in less than 35 min. Sixteen fluorescent and non-fluorescent peptides were obtained in over 95% purity (ESI Fig. 1 and 2†) and multi-mg scale in less than two weeks of synthesis. The peptides were then studied for their aggregation and condensation using biophysical analyses as described below.

Full phosphorylation of Tau R4 promotes its aggregation and condensation

To test if the phosphorylation of Tau R4 affects its aggregation and condensation, we first compared the fully phosphorylated (**3P**) and the non-phosphorylated (**0P**) peptides. We performed turbidity assays as a quick screening method to test first if the peptides undergo any type of high-order oligomerization. This may indicate possible condensation or aggregation of the peptides. Mixtures of 5% Fluorescein labeled (FL) **FL-0P** and **FL-3P** with 95% non-labeled peptides were tested at NaCl concentrations of 0, 150, 500 and 1000 mM, and peptide concentrations of 50, 300 and 1000 μM. Testing the dependence of the peptide oligomerization on the ionic strength was performed to examine whether the process is charge-dependent. This is important since the phosphorylation adds extra negative charges to the peptides. **3P** showed high turbidity in 1 mM peptide at all NaCl concentrations. It also showed high turbidity values in 1 M NaCl at all peptide concentrations. In contrast, **0P** showed low turbidity at all tested conditions except 1 M NaCl (Fig. 2A).

To test if the high turbidity of **3P** resulted from aggregation or condensation, we analyzed the oligomers using fluorescence microscopy. For **3P**, we observed condensates formation in 50 μM and 300 μM of peptide at all NaCl concentrations and aggregation of the peptide at 1000 μM of peptide in all NaCl concentrations. For **0P** we did not observe aggregation nor condensation of peptide except at very high salt concentration (Fig. 2B and C). This also confirmed that the fluorescein labeling itself did not promote condensation or aggregation.

An important feature of self-assembly processes is their dynamics, meaning the ability of the peptides to diffuse in and out of the condensates. This defines the rate of condensate maturation which is the conversion of condensates into a more viscous gel-like phase.⁴⁶ To evaluate the dynamics of the

condensate formation by **FL-3P** and test whether they have similar properties to the condensates of the recombinant Tau,⁴⁶ we used fluorescence recovery after photobleaching (FRAP). This method allows monitoring the diffusion of molecules such as peptides in and out of the condensates. Higher recovery means a higher diffusion rate and less viscous condensates. We bleached the entire **3P** condensate areas 15 and 40 minutes after they were formed. ~80% of the condensates that were bleached after 15 min recovered after 30 seconds. More mature condensates that were bleached 40 min after formation showed only partial recovery of ~35%. In both cases, a plateau was reached at approximately 40 seconds (Fig. 2D). These results indicate that after 15 minutes the condensates are liquid-like, which allows fast diffusion of **3P** in and out of the condensates. The partial recovery of the mature condensates can indicate a solidification process that led to a more gel-like character (Fig. 2D and E).

TEM and SEM images of **3P** taken after 10 days (Fig. 3A and B) showed that it formed spherical aggregates of 10–60 nm. These units together formed higher-order oligomeric structures with a flower-like morphology. Under the same conditions, **0P** did not aggregate even after this long incubation period (Fig. 3A and B).

Specific phosphorylation patterns control Tau-R4 condensation and aggregation

Controlling the condensation and aggregation of Tau R4, we characterized the ability of the mono and di-phosphorylated peptides to form condensates at NaCl concentrations of 0, 150, 300 and 1000 mM (Fig. 4A and B) using fluorescence microscopy. Mixtures of 5% fluorescein-labeled and 95% of non-labeled peptides were prepared. **1Pa**, **1Pb** and **2Pa** formed condensates, while **1Pc**, **2Pb** and **2Pc** did not form condensates. Like **3P**, **1Pb** and **2Pa** formed condensates in 0, 150, 300 and 1000 mM NaCl. In contrast, **1Pa** formed condensates only at lower salt concentrations of 0 and 150 mM. **2Pb** is the only peptide that underwent condensation only at 1000 mM NaCl. These findings suggest that: (1) phosphorylation is required for condensation of Tau R4; (2) phosphorylations of Ser341 and Ser352 facilitate condensation; (3) phosphorylation of Ser356 inhibits condensation. A closer investigation of the size



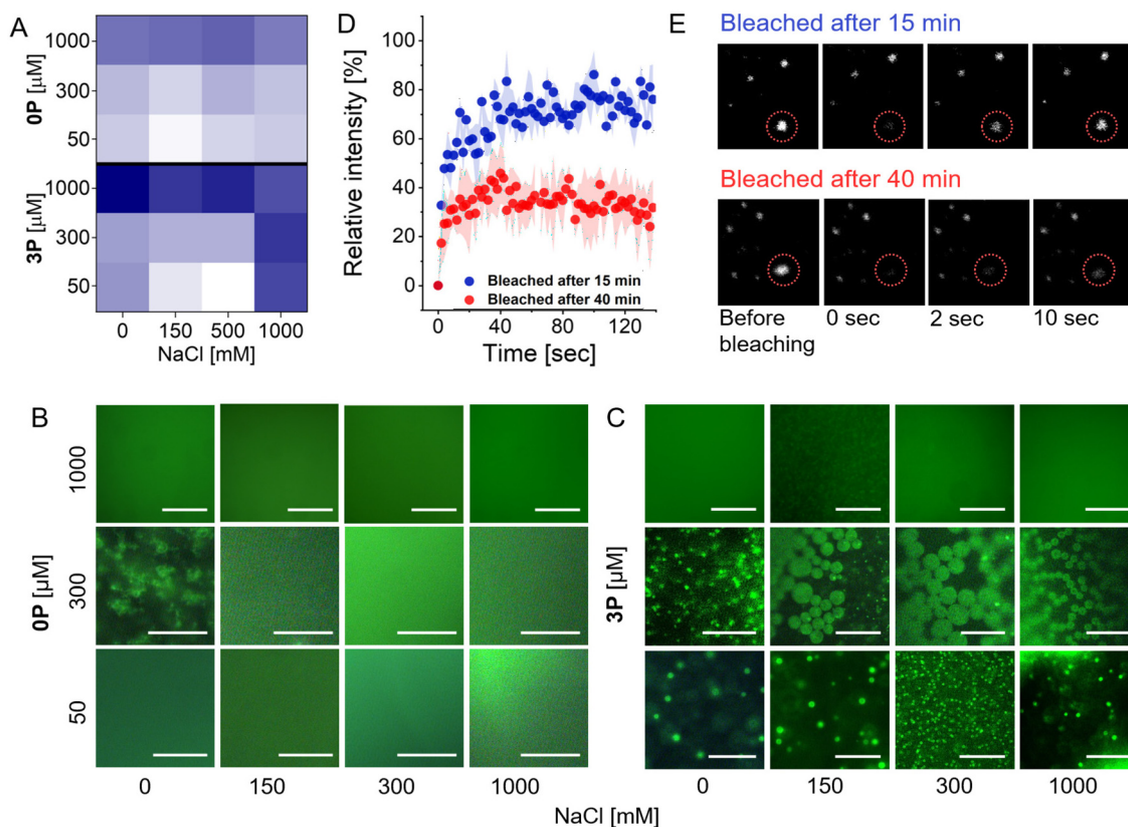


Fig. 2 (A) Turbidity assay for initial evaluation of possible oligomerization of **OP** and **3P**. Turbidity was measured in 0, 150, 300 and 1000 mM NaCl concentrations. Darker colors indicate higher optical density that correlates with high order oligomerization (B and C) fluorescence microscopy images of **3P** and **OP** condensates formed in 0, 150, 300 and 1000 mM NaCl (Y axis) and 50 and 300 μM peptide (X axis) (D) FRAP of **3P** mixed with 5% of FL-**3P** after incubation of 15 min (blue) and 40 min (red). Shown is the time dependence of the relative fluorescence intensity (E) recovered **3P** bleached condensates (circled), bleached after 15 min incubation on the left and after 40 min incubation on the right, before and during the recovery process.

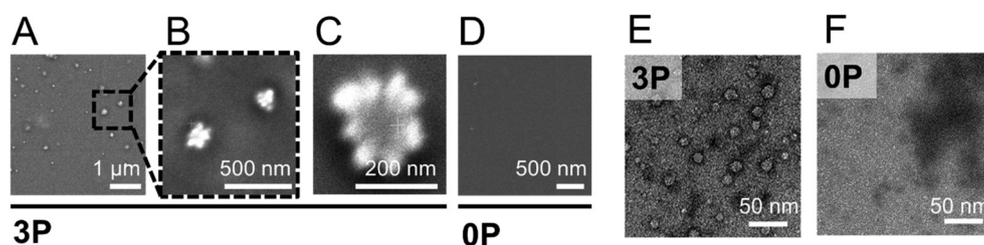


Fig. 3 (A–D) Characterization of **3P** aggregates using SEM microscopy. Image of **3P** (A), followed by zooming in on the flower-like structure (B) and an enlarged image of flower-like aggregates (C). **OP** did not aggregate (D). (E and F) Characterization of **3P** aggregates using TEM microscopy. **3P** showed spherical aggregates (E). **OP** did not aggregate (F).

of the condensates showed that while **1Pa** had an average diameter of ~ 3.5 μm, **1Pb** formed condensates with diameters that ranged between 4 to 12 μm. The largest condensates of **1Pb** were in the same range as the mean diameter of **2Pa** ~ 12 μm, and **3P** ~ 12.3 μm. Condensates of **1Pa**, **2Pa** and **3P** were uniform in their size, with relatively small standard deviations, while **1Pb** showed a large range of condensate size (Fig. 4C). The intensity of the fluorescence in each condensate was examined since it is an indication of peptide

concentration. We showed that while the peptide concentration of **1Pb**, **2Pa** and **3P** was homogeneous across the condensate droplets. The edge of **1Pa** condensates had higher fluorescence intensity (Fig. 4D) suggesting that peptides accumulated on the surface of the condensate, which can stop the growth of the condensates to give smaller entities.

SEM was used to characterize the aggregates formed by the phosphorylated peptides after 10 days of incubation (Fig. 5A).



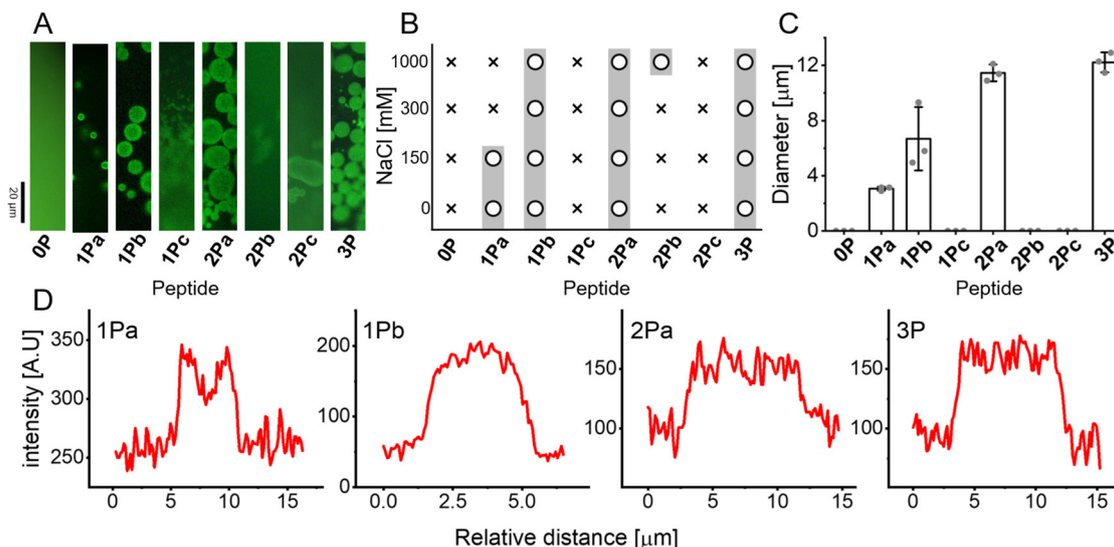


Fig. 4 (A) Fluorescence microscopy of the peptide library. Only **1Pa**, **1Pb**, **2Pa** and **3P** made condensates. (B) The dependence of peptides condensates on ionic strength. 'O' with grey background: condensates formation; 'X': No condensates formation. X axis is the peptide library, Y axis is NaCl concentration in mM. (C) The maximum diameter size of the condensates of the different peptides. (D) Cross-section analysis of condensates of the peptides that underwent condensation. X axis is the position within the cross-section of the condensate in μm . The 0 μm position was defined as the starting point of the measurement. Y axis is the intensity of each position. **1Pa** has higher fluorescence intensity around the condensate edges while in the other peptides, the intensity is uniform.

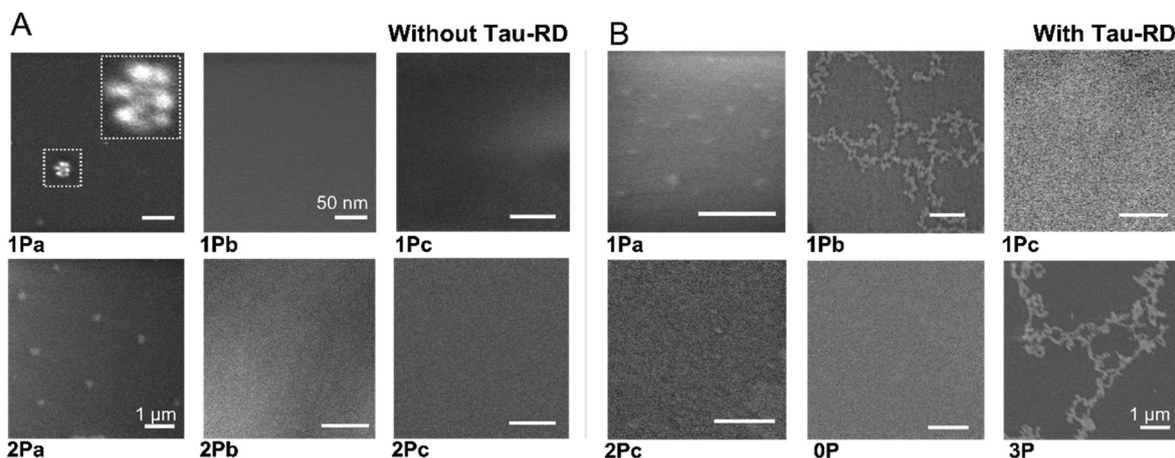


Fig. 5 (A) SEM of mono- and di-phosphorylated Tau336–358 peptides without Tau-RD (B) SEM of **0P**, **1Pa**, **1Pb**, **2Pc** and **3P** Tau336–358 peptides with Tau-RD. Scale bar = 500 nm.

2Pa, **1Pb**, **1Pc**, **2Pb** and **2Pc** did not aggregate (ESI Fig. S2†). Like **3P**, **1Pa** formed spherical aggregates with flower-like morphology. These results suggest that phosphorylation of Ser341 contributes to the aggregation behavior of Tau R4, while phosphorylation of Ser352 and Ser356 did not contribute to the phenomena.

The effect of specific R4 phosphorylations and their combinations on the Tau-RD protein (244–372) aggregation

Since it is technically impossible to study the effect of phosphorylation patterns at the protein level *via* a library of phosphoproteins, we decided to study the effect of phosphorylation

patterns on Tau oligomerization by testing the effect of the phosphorylated Tau R4 peptide library on aggregation the full R domain of Tau (Tau-RD, residues 244–372), which is responsible for its amyloid aggregation. *In vitro* Tau forms amyloids *via* its MTBR only at the presence of an inducer such as heparin, which is highly negatively charged. Here, we tested if the phosphorylated peptides have a similar effect, due to their negative charges. When Tau-RD was incubated for 10 days with **3P**, spherical aggregates with a diameter of 20–60 nm were formed, interconnected by larger amorphous aggregates. This suggests that aggregation is indeed dependent on the interactions between **3P** and Tau-RD.



To test the effects of the specific phosphorylation patterns at the protein level, **0P**, **1Pa**, **1Pb**, **1Pc** and **2Pc** were incubated with the recombinant Tau-RD for 10 days at 37 °C and the samples were tested using SEM (Fig. 5B). **3P** and **1Pb** induced Tau-RD aggregation, resulting in aggregates with string-like morphology. **1Pa**, **1Pc**, and **2Pc** incubation with Tau-RD did not result in oligomer formation. Tau-RD also did not aggregate by itself (ESI Fig. 3†).

Specific phosphorylation patterns control Tau-R4 condensation and aggregation

Charge and steric hindrance alone cannot explain the different effects of the three phosphorylations on the oligomerization of Tau R4. We used the PEP-FOLD3 predictor to examine the conformations and secondary structures of Tau R4.⁴⁷ The most probable 29 models that were calculated and generated (ESI Fig. 4†) shared the same closed hairpin-like conformation as was found in AD amyloids. All these models contain a combination of disordered and structured domains (Fig. 1C). A notable difference is that according to the models, the Tau R4

fragment has a helical domain whereas the same region in the full-length protein has a beta-sheet structure (Fig. 7A and B). Indeed, peptides usually adopt helical rather than a β -sheet structures since the stabilizing partner strands are missing. In all the calculated models, Ser341 is in the disordered region of the peptide, from Q356 to Phe346. Ser352 is in the helical domain that includes 7 residues from Lys347 to Lys353, and Ser356 is located in the IDR tail from Ile354 to Asp358 (Fig. 6A), similar to their locations in the AD amyloid structure. The CD spectrum of **0P**, had a minimum at \sim 200 nm and \sim 225 nm, corresponding to disordered and helical conformations (Fig. 6B). **0P** and **3P** had similar CD spectra suggesting they obtain the same secondary structure (Fig. 6B). Using the PEPstrMOD server,⁴⁸ we generated a *de novo* structure prediction of Tau R4 β -helix motif phosphorylated at all three p-sites. The results show that the phosphorylations do not affect the secondary structure (Fig. 6C). Thus, we added the phosphate groups to the backbone of **0P** using a PyTM plugin (version 1.8.4.0).⁴⁹ Using PyMol measurement wizard,⁵⁰ we found that Ser341 and Ser356 are more solvent-exposed than Ser352

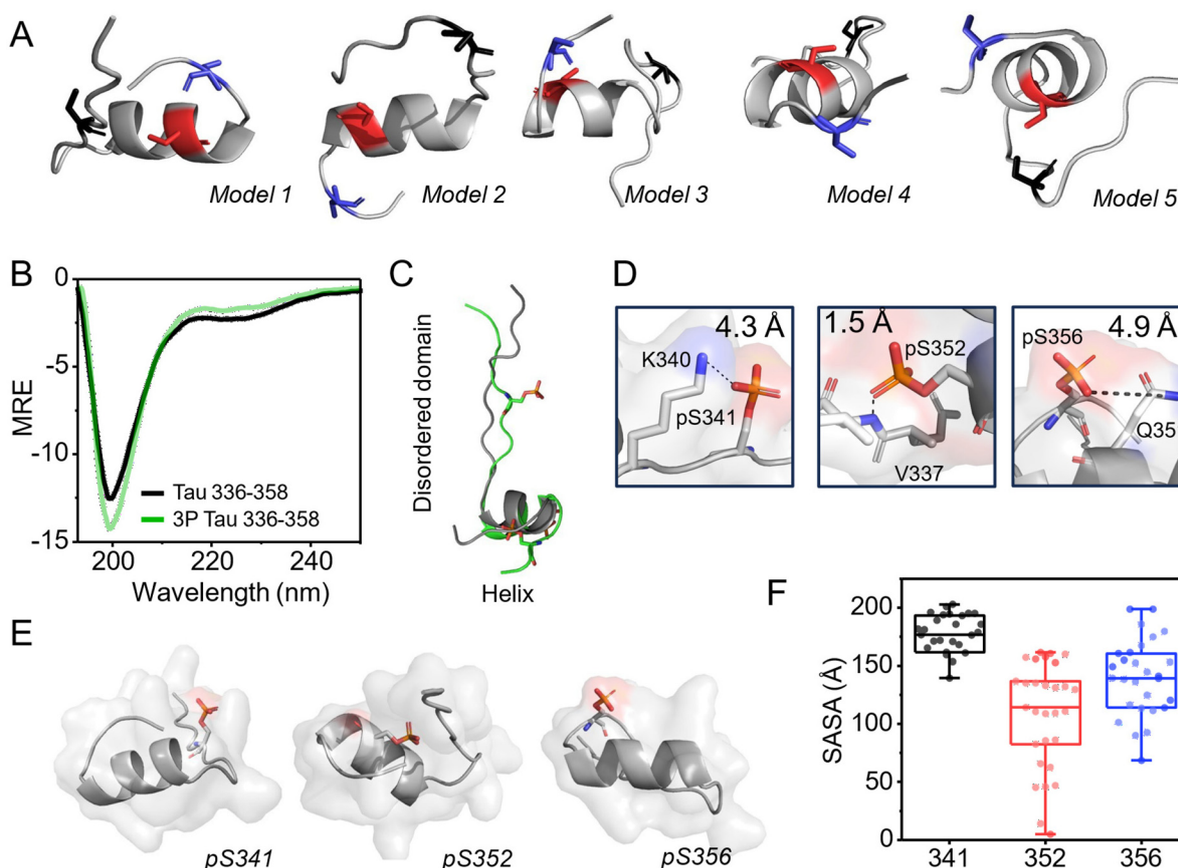


Fig. 6 (A) The five most probable models calculated using PEP-FOLD3⁴⁷ show that Tau R4 has an IDR tail, a helical domain, and a disordered region. In all models Ser341 (black) is located in the disordered region, Ser352 (red) in the helical domain and Ser356 (blue) in the IDR tail. (B) CD spectra of **0P** (black) and **3P** (green) indicate the same secondary structures, comprising of helical and disordered domains. (C) models of **0P** (black) and **3P** (green) as calculated by PEP-FOLD3 and PEPstrMOD.⁴⁸ Phosphate groups are shown in orange. (D) Zoom in on the chemical environment and interactions of each phosphorylation site (Ser341 and 356 shown in model 1, Ser352 in model 2). (E) The position of each phosphorylation site in the Tau peptide, as shown by Pymol for model 1. (F) SASA of each phosphorylation site. Boxplots represent the median value and interquartile (25–75% percentiles) of SASA. Each dot represents one of the 29 models that were examined.



(Fig. 6D). Calculated distances between the p-site and their nearest entity showed that: (1) pSer352 may interact with the amine group on the backbone of Val337 *via* hydrogen bond of 1.5 Å, (2) pSer341 had a 4.3 Å distance from the most adjacent positive charged Lys351, and (3) pSer356 had a 4.7 Å distance from the adjacent amide of the side chain of Gln351 (Fig. 6E). Altogether, pSer 352 is the least exposed of all three. pSer341 and pSer356 are more exposed, and in contrast to pSer352, do not have significant interactions with their surroundings.

Using the Pymol software we calculated the solvent-accessible surface area (SASA)⁵¹ for pSer341, pSer352 and pSer356 in the most probable 29 models of the peptide. The phosphate groups were added by PyTM.⁴⁹ to the models that were generated by the predictor. The average SASA of pSer352 was $108 \pm 45 \text{ \AA}^2$ while pSer341 had an average SASA of $169 \pm 36 \text{ \AA}^2$. pSer356 had an average SASA of $137 \pm 36 \text{ \AA}^2$.

Discussion

In the current study, we combined phosphorylated peptide libraries with an ensemble of biophysical methods to elucidate the specific effects of phosphorylation patterns on the different association processes of Tau R4. We found that phosphorylation at specific sites of the β -helix motif dictate whether the peptides can assemble to form oligomers at all, and which type of oligomerization process they undergo or induce (condensation or amyloid aggregation). Our results highlight the importance of Ser352 as a major phosphorylation site for inducing Tau oligomerization. This result could be obtained only with the help of our synthetic methods enabling the synthesis of phosphorylated peptide libraries.

The role of the phosphorylation pattern in switching between condensation and aggregation of Tau R4

Our results show that the Tau R4 peptide (**OP**) does not form any type of high-order oligomers by itself and does not induce oligomerization of Tau-RD. Phosphorylation of one or more serine residues within peptides derived from this R4 domain results in significant changes in oligomerization processes. The tri-phosphorylated peptide **3P** formed condensates or aggregates, depending on the time and ionic strength, and induced aggregation of the Tau-RD. This indicates that phosphorylation controls both the condensation and aggregation of this domain and the specific phosphorylation is critical for the decision which of the processes will take place. These phenomena are unique, as recombinant non-phosphorylated Tau monomers do not tend to aggregate even within days.⁵²

The contributions of each phosphorylation within these patterns are not necessarily additive. Phosphorylation of Ser352 activated condensation of R4 as shown by the ability of the **1Pb** peptide to undergo condensation, while phosphorylation of Ser341 activated aggregation of R4 as shown by the aggregation of the **1Pa** peptide. Phosphorylation of Ser356 prevented any oligomerization of R4 as reflected by the inability of **1Pc** to form condensates or aggregates. The effects of combi-

nations of two phosphorylations were synergistic. **2Pa**, which contained phosphorylations at residues Ser341 and Ser352, generated condensates with similar size and propensity to **1Pb** containing a single phosphorylation at Ser352. We conclude that Ser352 is more dominant in promoting condensation than Ser341. Combining phosphorylation of Ser356 with phosphorylation of Ser341 (**2Pb**) or with phosphorylation of Ser352 (**2Pc**) resulted in no condensates, like the effect of Ser356 phosphorylation alone (**1Pc**) (Fig. 7C). We conclude that Ser356 phosphorylation prevents condensate formation even in the presence of the oligomerization-activating phosphorylation at Ser341 or Ser352. However, when the 3 phosphorylations were combined (**3P**), we observed condensate formation despite the phosphorylation at position 356. This shows that a combination of both Ser341 and Ser352 phosphorylations overcame the inhibitory effect of phosphorylating Ser 356. Using a peptide library with all possible combinations of phosphorylations makes it possible to elucidate the role of each phosphorylation in controlling the oligomerization processes of Tau. Moreover, it enables demonstrating the synergism between the different phosphorylation sites. The library thus provides a detailed mechanistic insight about how the combinations of the phosphorylations contribute to the oligomerization processes.

Our results revealed that phosphorylation of Ser352 is highly important for Tau aggregation. Phosphorylated Ser352 can be found in a type of Tau filament, named PHF, which was isolated from brains of AD patients.⁵³ However, it could not be phosphorylated *in vitro* by kinases combination including PKA, GSK, PKA + GSK, P12, P20, PKA + GSK3 β and PKA + SAPK4.^{21,53–55} This suggests that phosphorylation of this site *in vitro* is challenging, explaining why this site was not studied extensively in previous research. Working at the peptide level, using our advanced phosphorylated peptide synthesis methods, allowed us to pinpoint Ser352 as crucial site for Tau aggregation.

The effect of the phosphorylated Tau R4 peptides on the aggregation of Tau-RD protein

Studying biological systems in their native forms is ideal, but creating a relevant library of phosphoproteins is currently not feasible. To overcome this challenge, we combined synthetic phosphopeptide libraries with recombinant Tau-RD to investigate the effect of phosphorylation on aggregation at the protein level. We focused on the Tau-RD domain because it plays a critical role in amyloid formation in the brains of Alzheimer's disease (AD) and chronic traumatic encephalopathy (CTE) patients. While inducers such as heparin and poly-RNA have been widely used to study Tau aggregation and condensation by simulating certain aspects of the physiological environment,⁵⁶ they can introduce non-native interactions, potentially distorting the interpretation of aggregation mechanisms. Our study shows that using 100 μM of peptides with specific phosphorylation patterns alone can induce Tau aggregation and condensation without the need for exogenous inducers. This approach provides native cellular conditions, ensur-



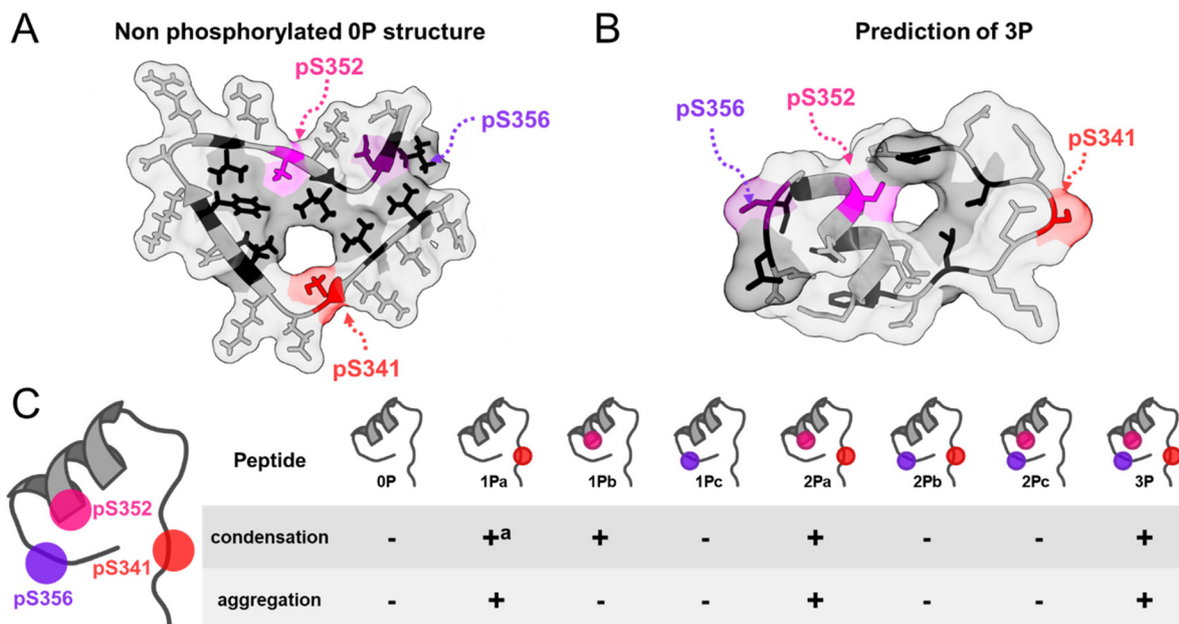


Fig. 7 (A) Structure of Tau336–358 (0P) as found in amyloids isolated from AD patients (11). (B) Predicted secondary structure and conformation of tri-phosphorylated Tau336–358 (3P) as calculated by PEP-FOLD3 (37). Color code for A and B: hydrophobic groups are black. Serine 341 is labeled in red, 352 in magenta and 356 in purple. (C) Summary of the effects of the different phosphorylation patterns on Tau oligomerization. For condensation, (+) represents condensation in all tested conditions; (+^a) represents condensation that occurred at certain ionic strength conditions but not all conditions tested; (–) represents peptides that did not undergo condensation. For aggregation, (+) represents the formation of aggregates; (–) represents peptides that did not undergo aggregation.

ing that the observed effects are driven solely by the intrinsic properties of the phosphorylated peptides, and most importantly, in the specific phosphorylation pattern.

The micro-environment of the phosphorylation sites is responsible for controlling Tau R4 oligomerization

The computational part represents a monomeric, non-aggregated peptide, which is why its structure does not align with the beta-sheet structure found in full-length Tau amyloids (Fig. 6). This difference in structure highlights the effect of the microenvironment on peptide conformation, that is crucial for understanding aggregation processes. SASA calculations of the 29 models of Tau R4 β -helix motif show that pSer352 is significantly less exposed to the solvent compared with pSer341 and pSer356. In the calculated conformations, pSer352 is buried and forms hydrogen bonds with the backbone of Val337 (e.g. Fig. 6D model 2). A closer examination of model 1 shows that pSer341 and pSer356 are located at the peripheral parts of the peptide with the closest interactions being at distances larger than 4 Å (Fig. 6D). This suggests that these phosphate groups do not bind any backbone or side chain residues within the peptide (Fig. 6E). Ser341 and Ser356 still differ in their micro-environments. Ser341 is in the IDR of Tau R4, while Ser356 is located within the IDR tail, adjacent to the structured domain of the peptide. The residues surrounding Ser341 are the charged Lys340 and Glu342, while the residues surrounding Ser356 are the small Gly355 and the hydrophobic Leu357.

The microenvironments described above for each phosphorylation site explain the observed effects of the different phosphorylations on the oligomerization of the Tau R4 peptide. Phosphorylation of Ser352 resulted in phosphoserine which is less accessible to water and salt. Thus, the peptides including this phosphorylation, **1Pb**, **2Pa**, **2Pc** and **3P**, are less soluble and more available to form interactions with additional peptides or proteins (such as Tau-RD) resulting in condensation or aggregation. Phosphorylation of Ser356, as in **1Pc**, did not show any type of oligomerization. This can be explained by a possible interaction of the phosphate group with water molecules, which leads to better solubility of the entire peptide. Our suggested model correlates with the experimental findings regarding the activating and inhibiting phosphorylation.

High salt concentration inhibited condensate formation by **1Pa**, containing phosphorylation only at Ser341, but not by peptides containing phosphorylation at position 352. This correlates well with our model and can be explained by the higher exposure of Ser 341 to the environment and thus to the salt, which at high concentrations can inhibit the molecular interaction leading to condensation. The micro-environment of the peptides can also explain why **1Pa** formed smaller condensates than **1Pb**. The same principles also explain why **1Pa** condensates are smaller and the peptide is concentrated at the boundaries of the condensate. The buried pSer352 was not reactive enough to initiate aggregation on its own (it initiated condensation) but it did induce Tau-RD aggregation. pS356 on the



other hand is too exposed to allow participation of the peptide in aggregation events with or without Tau-RD.

In conclusion, by using phosphopeptide libraries, our study pinpoints the essential role of specific phosphorylation sites in driving Tau's self-assembly and the decision when condensation or aggregation will take place, further advancing the understanding of its role in neurodegenerative diseases.

Methods

Peptide synthesis, labeling and purification

The peptides were synthesized on a rink amid resin with a 0.48 mmol gr^{-1} loading after 30 min of swelling in DMF. All peptides were synthesized using the Stirring-based accelerated peptide synthesis approach.²⁴ A reactor containing a sintered glass filter, and a heating jacket connected to a circulating 90 °C water bath. The reactor was equipped with an overhead 5-fin turbine PTFE impeller. Solvents and reagents were inserted directly by a feeding line and drained by vacuum filtration. Each cycle started with Fmoc deprotection followed by a 5 mL DMF wash, continued with 1 min coupling and a second wash. Fmoc-Ser/(PO₃Bzl)-OH and its sequential residue were double coupled. 3 mL of coupling mixture containing 3 equiv. of protected amino acid, 2.9 equiv. of HATU, and 8 equiv. of DIEA was added. Fmoc deprotection was done by adding 3 mL of a 0.5% DBU solution in DMF for 10 s. Peptides were cleaved using a freshly prepared TFA cocktail, and were purified by Waters preparative HPLC, using a reverse-phase C18 column with a TDW/ACN gradient. The column was equilibrated with 5% ACN for 5 minutes, followed by a linear increase from 5% to 60% ACN over 30 minutes. After elution, the column was washed with 95% ACN for 12 minutes to remove residual hydrophobic components. Finally, the ACN concentration was reduced to 5% over 10 minutes to re-equilibrate the column for subsequent runs. All peptides were characterized by electron spray ionization mass spectrometry (ESI-MS) (ESI Fig. 1†). Peptide purity was tested by dissolving 80 μL of peptide in water injecting into analytical HPLC. Peptides were fluorescein labeled for FRAP and fluorescence microscopy experiments, with 5(6)-carboxy fluorescein at their N' termini. Fractions containing the desired phosphorylated peptides were lyophilized and the peptides were used without further purifications.

Turbidity assay

150 μL **0P** and **3P** (with 5% FL-peptide each) were tested at peptide concentrations of 50, 300 and 1000 μM . The solutions contained 25 mM phosphate buffer at pH of 7.4 and 10% PEG 400 (buffer A) with NaCl concentrations of 0, 150, 300 and 1000 mM. Triplicates were measured at $\lambda = 350$ in BioTek Synergy H1 plate reader.

Fluorescence microscopy

150 μL **0P** and **3P** (with 5% FL-peptide each) at peptide concentrations of 50, 300 and 1000 μM were also studied using fluo-

rescence microscopy. 10 μL of 50 μM peptide from each sample were applied onto a glass slip and visualized under Zeiss Axio scope A1 microscope with a green-fluorescent filter. Images were taken with an AxioCam ICc 3 camera, using Zen software.

Fluorescence recovery after photobleaching (FRAP) experiments

FRAP experiments were performed using an FV-1200 confocal microscope (Olympus, Japan) equipped with a sim scanner which enables bleaching while scanning. All experiments were done using a 60 \times /1.42 oil objective. Imaging was done using the 488 nm laser line and emission was collected 500–540 nm. 5% of **FL-3P** and 95% of **3P** were dissolved in 100 μL buffer A to final concentration of 25 mM phosphate buffer and 150 mM NaCl and 50 μM peptide. The measurement started by adding 10% (V/V) PEG-400 and samples were bleached after 15 and 45 min to evaluate the maturation of condensates. A specific region in the sample was exposed to 100% laser intensity for 120 ms. The interval between image scans was 2 s. Photobleaching correction and recovery were calculated using the OriginPro 2022 software. Final FRAP recovery curves are the average from measuring three different condensates.

CD of 0P and 3P

120 μM of **0P** and **3P**, in a 25 mM phosphate buffer at pH of 7.4 and 150 mM NaCl CD spectrum were determined using JASCO J-810 spectropolarimeter and Spectra manager software. The temperature was 25 °C and controlled by Peltier thermostat, in a 0.1 cm quartz cuvette for far-UV CD spectrometry, ranged from 190 to 260 nm.

Aggregation assay

To examine the aggregation properties of the peptides, 100 μM of peptides were incubated with and without 10 μM Tau-RD and 20 mM phosphate buffer at pH of 7.4 and 150 mM NaCl. 150 μL of the final solutions were incubated at a temperature of 37 °C for 10 days. The solution was then dried on silicon wafers and examined using SEM Apero 2S LoVac. Each experiment was repeated between three to nine times.

Conflicts of interest

The authors declare no conflict of interest.

Data availability

The data supporting this article have been included as part of the ESI.†

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References

- Q. Zhong, X. Xiao, Y. Qiu, Z. Xu, C. Chen, B. Chong, X. Zhao, S. Hai, S. Li, Z. An and L. Dai, *MedComm*, 2023, **4**, e261.
- F. Ardito, M. Giuliani, D. Perrone, G. Troiano and L. Lo Muzio, *Int. J. Mol. Med.*, 2017, **40**, 271–280.
- S. G. Bressler, A. Mitrany, A. Wenger, I. Nathke and A. Friedler, *Int. J. Mol. Sci.*, 2023, **24**, 6478.
- T. Pawson and J. D. Scott, *Trends Biochem. Sci.*, 2005, **30**, 286–290.
- M. Samarasimhareddy, G. Mayer, M. Hurevich and A. Friedler, *Org. Biomol. Chem.*, 2020, **18**, 3405–3422.
- M. Mann, S.-E. Ong, M. Grønberg, H. Steen, O. N. Jensen and A. Pandey, *Trends Biotechnol.*, 2002, **20**, 261–268.
- T. Garfagnini, L. Ferrari, M. B. Koopman, F. A. Dekker, S. Halters, E. Van Kappel, G. Mayer, S. Bressler, M. M. Maurice, S. G. D. Rudiger and A. Friedler, *Chem. – Eur. J.*, 2024, **30**, e202400080.
- B. Wang, L. Zhang, T. Dai, Z. Qin, H. Lu, L. Zhang and F. Zhou, *Signal Transduction Targeted Ther.*, 2021, **6**, 290.
- T. P. Lopez-Palacios and J. L. Andersen, *Trends Cell Biol.*, 2023, **33**, 649–666.
- X. Tong, R. Tang, J. Xu, W. Wang, Y. Zhao, X. Yu and S. Shi, *Signal Transduction Targeted Ther.*, 2022, **7**, 211.
- A. Aguzzi and T. O'Connor, *Nat. Rev. Drug Discovery*, 2010, **9**, 237–248.
- M. Samarasimhareddy, D. Mayer, N. Metanis, D. Veprintsev, M. Hurevich and A. Friedler, *Org. Biomol. Chem.*, 2019, **17**, 9284–9290.
- A. Kozelekova, A. Naplavova, T. Brom, N. Gasparik, J. Simek, J. Houser and J. Hritz, *Front. Chem.*, 2022, **10**, 835733.
- P. E. Thorsness and D. E. Koshland, *J. Biol. Chem.*, 1987, **262**, 10422–10425.
- N. E. Mammeri, A. J. Dregni, P. Duan and M. Hong, *Proc. Natl. Acad. Sci. U. S. A.*, 2024, **121**, e2316175121.
- S. M. Pearlman, Z. Serber and J. E. Ferrell Jr., *Cell*, 2011, **147**, 934–946.
- N. M. Otto, W. G. McDowell, D. M. Dickey and L. R. Potter, *Mol. Pharmacol.*, 2017, **92**, 67–74.
- S. A. Johnson and T. Hunter, *Nat. Methods*, 2005, **2**, 17–25.
- K. E. Paleologou, A. W. Schmid, C. C. Rospigliosi, H.-Y. Kim, G. R. Lamberto, R. A. Fredenburg, P. T. Lansbury Jr., C. O. Fernandez, D. Eliezer, M. Zweckstetter and H. A. Lashuel, *J. Biol. Chem.*, 2008, **283**, 16895–16905.
- G. Tesei, T. K. Schulze, R. Crehuet and K. Lindorff-Larsen, *Proc. Natl. Acad. Sci. U. S. A.*, 2021, **118**, e2111696118.
- J. X. Meng, Y. Zhang, D. Saman, A. M. Haider, S. De, J. C. Sang, K. Brown, K. Jiang, J. Humphrey, L. Julian, E. Hidari, S. F. Lee, G. Balmus, R. A. Floto, C. E. Bryant, J. L. P. Benesch, Y. Ye and D. Klenerman, *Nat. Commun.*, 2022, **13**, 2692.
- D. Ellmer, M. Brehs, M. Haj-Yahya, H. A. Lashuel and C. F. Becker, *Angew. Chem.*, 2019, **131**, 1630–1634.
- Y. Li, J. Heng, D. Sun, B. Zhang, X. Zhang, Y. Zheng, W.-W. Shi, T.-Y. Wang, J.-Y. Li and X. Sun, *J. Am. Chem. Soc.*, 2021, **143**, 17566–17576.
- D. Grunhaus, E. R. Molina, R. Cohen, T. Stein, A. Friedler and M. Hurevich, *Org. Process Res. Dev.*, 2022, **26**, 2492–2497.
- J. N. Naoum, I. Alshanski, G. Mayer, P. Strauss and M. Hurevich, *Org. Process Res. Dev.*, 2022, **26**, 129–136.
- D. Ben Abba Amiel and M. Hurevich, *Eur. J. Org. Chem.*, 2022, 202200623.
- P. Strauss, F. Nuti, M. Quagliata, A. M. Papini and M. Hurevich, *Org. Biomol. Chem.*, 2023, **21**, 1674–1679.
- M. Samarasimhareddy, G. Mayer, M. Hurevich and A. Friedler, *Org. Biomol. Chem.*, 2020, **18**, 3405–3422.
- M. Samarasimhareddy, D. Mayer, N. Metanis, D. Veprintsev, M. Hurevich and A. Friedler, *Org. Biomol. Chem.*, 2019, **17**, 9284–9290.
- R. Rajmohan and P. H. Reddy, *J. Alzheimer's Dis.*, 2017, **57**, 975–999.
- D. Eisenberg and M. Jucker, *Cell*, 2012, **148**, 1188–1203.
- T. P. Knowles, M. Vendruscolo and C. M. Dobson, *Nat. Rev. Mol. Cell Biol.*, 2014, **15**, 384–396.
- S. Boyko and W. K. Surewicz, *Trends Cell Biol.*, 2022, **32**, 611–623.
- S. Ambadipudi, J. Biernat, D. Riedel, E. Mandelkow and M. Zweckstetter, *Nat. Commun.*, 2017, **8**, 275.
- N. M. Kanaan, C. Hamel, T. Grabinski and B. Combs, *Nat. Commun.*, 2020, **11**, 2809.
- S. Wegmann, B. Eftekharzadeh, K. Tepper, K. M. Zoltowska, R. E. Bennett, S. Dujardin, P. R. Laskowski, D. MacKenzie, T. Kamath, C. Commins, C. Vanderburg, A. D. Roe, Z. Fan, A. M. Molliex, A. Hernandez-Vega, D. Muller, A. A. Hyman, E. Mandelkow, J. P. Taylor and B. T. Hyman, *EMBO J.*, 2018, **37**, e98049.
- D. Ellmer, M. Brehs, M. Haj-Yahya, H. A. Lashuel and C. F. W. Becker, *Angew. Chem., Int. Ed.*, 2019, **58**, 1616–1620.
- A. W. P. Fitzpatrick, B. Falcon, S. He, A. G. Murzin, G. Murshudov, H. J. Garringer, R. A. Crowther, B. Ghetti, M. Goedert and S. H. W. Scheres, *Nature*, 2017, **547**, 185–190.



- 39 B. Falcon, J. Zivanov, W. Zhang, A. G. Murzin, H. J. Garringer, R. Vidal, R. A. Crowther, K. L. Newell, B. Ghetti, M. Goedert and S. H. W. Scheres, *Nature*, 2019, **568**, 420–423.
- 40 M. Von Bergen, P. Friedhoff, J. Biernat, J. Heberle, E. M. Mandelkow and E. Mandelkow, *Proc. Natl. Acad. Sci. U. S. A.*, 2000, **97**, 5129–5134.
- 41 S. Lovestam, D. Li, J. L. Wagstaff, A. Kotecha, D. Kimanius, S. H. McLaughlin, A. G. Murzin, S. M. V. Freund, M. Goedert and S. H. W. Scheres, *Nature*, 2024, **625**, 119–125.
- 42 M. Haj-Yahya, P. Gopinath, K. Rajasekhar, H. Mirbaha, M. I. Diamond and H. A. Lashuel, *Angew. Chem., Int. Ed.*, 2020, **59**, 4059–4067.
- 43 D. N. Edun, M. R. Flanagan and A. L. Serrano, *Chem. Sci.*, 2020, **12**, 2474–2479.
- 44 M. Abbas, W. P. Lipinski, K. K. Nakashima, W. T. S. Huck and E. Spruijt, *Nat. Chem.*, 2021, **13**, 1046–1054.
- 45 D. Grunhaus, A. Friedler and M. Hurevich, *Eur. J. Org. Chem.*, 2021, **2021**, 3737–3742.
- 46 S. Wegmann, B. Eftekharzadeh, K. Tepper, K. M. Zoltowska, R. E. Bennett, S. Dujardin, P. R. Laskowski, D. MacKenzie, T. Kamath, C. Commins, C. Vanderburg, A. D. Roe, Z. Fan, A. M. Molliex, A. Hernandez-Vega, D. Muller, A. A. Hyman, E. Mandelkow, J. P. Taylor and B. T. Hyman, *EMBO J.*, 2018, **37**, e98049–e98049.
- 47 A. Lamiabile, P. Thevenet, J. Rey, M. Vavrusa, P. Derreumaux and P. Tuffery, *Nucleic Acids Res.*, 2016, **44**, W449–W454.
- 48 S. Singh, H. Singh, A. Tuknait, K. Chaudhary, B. Singh, S. Kumaran and G. P. Raghava, *Biol. Direct*, 2015, **10**, 73.
- 49 A. Warnecke, T. Sandalova, A. Achour and R. A. Harris, *BMC Bioinf.*, 2014, **15**, 370.
- 50 *The PyMOL Molecular Graphics System*, Version 1.8, Schrödinger, LLC, New York, NY, USA, 2015.
- 51 S. Mitternacht, *F1000Research*, 2016, **5**, 189.
- 52 W. Zhang, B. Falcon, A. G. Murzin, J. Fan, R. A. Crowther, M. Goedert and S. H. Scheres, *eLife*, 2019, **8**, 43584.
- 53 C. X. Gong, F. Liu, I. Grundke-Iqbal and K. Iqbal, *J. Neural Transm.*, 2005, **112**, 813–838.
- 54 K. Tepper, J. Biernat, S. Kumar, S. Wegmann, T. Timm, S. Hubschmann, L. Redecke, E. M. Mandelkow, D. J. Muller and E. Mandelkow, *J. Biol. Chem.*, 2014, **289**, 34389–34407.
- 55 D. P. Hanger, H. L. Byers, S. Wray, K. Y. Leung, M. J. Saxton, A. Seereeram, C. H. Reynolds, M. A. Ward and B. H. Anderton, *J. Biol. Chem.*, 2007, **282**, 23645–23654.
- 56 P. Duan, A. J. Dregni, N. E. Mammeri and M. Hong, *Proc. Natl. Acad. Sci. U. S. A.*, 2023, **120**, e2310067120.

