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Elucidating the reversible and irreversible self-assembly mechanisms of low-complexity aromatic-rich kinked peptides and steric zipper peptides†

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Many RNA-binding proteins such as fused-in sarcoma (FUS) can self-assemble into reversible liquid droplets and fibrils through the self-association of their low-complexity (LC) domains. Recent experiments have revealed that SYG-rich segments in the FUS LC domains play critical roles in the reversible self-assembly behaviors of FUS. These FUS LC segments alone can self-assemble into reversible kinked fibrils, which are markedly different from the canonical irreversible steric zipper β -sheet fibrils. However, the molecular determinants underlying the reversible and irreversible self-assembly are poorly understood. Herein we conducted extensive all-atom and coarse-grained molecular dynamics simulations of four representative hexapeptides: two low-complexity aromatic-rich kinked peptides from the amyotrophic lateral sclerosis-related FUS protein, FUS_{37–42} (SYSGYS) and FUS_{54–59} (SYSSYG); and two steric zipper peptides from Alzheimer's-associated A β and Tau proteins, A β _{16–21} (KLVFFA) and Tau_{306–311} (VQIVYK). We dissected their reversible and irreversible self-assembly dynamics, predicted their phase separation behaviors, and elucidated the underpinning molecular interactions. Our simulations showed that alternating stickers (Tyr) and spacers (Gly and Ser) in FUS_{37–42} and FUS_{54–59} facilitate the formation of highly dynamic coil-rich oligomers and lead to reversible self-assembly, while consecutive hydrophobic residues of LVFF in A β _{16–21} and IVY in Tau_{306–311} act as hydrophobic patches, favoring the formation of stable β -sheet-rich oligomers and driving the irreversible self-assembly. Intriguingly, we found that FUS_{37–42} and FUS_{54–59} peptides, possessing the same amino acid composition and the same number of sticker and spacer residues, display differential self-assembly propensities. This finding suggests that the self-assembly behaviors of FUS peptides are fine-tuned by the site-specific patterning of spacer residues (Ser and Gly). This study provides significant mechanistic insights into reversible and irreversible peptide self-assembly, which would be helpful for understanding the molecular mechanisms underlying the formation of biological liquid condensates and pathological solid amyloid fibrils.

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Introduction

Fibrillary deposits are the hallmark of many neurodegenerative disorders,^{1,2} such as Alzheimer's disease associated with the fibrillary aggregates of A β and Tau.^{3–5} Amyloid fibrils (also known as steric zipper β -sheet fibrils) share cross- β architecture, with extended β -strands stacking along the fibril axis into a β -sheet and the side chains of mating sheets interdigitating with each other.^{6–8} Attributed to their highly ordered intermolecular backbone hydrogen bonds (H-bonds) and complementary dry zipper interface, steric zipper β -sheet fibrils are stable and present strong resistance to dissociation under harsh conditions, such as being in the presence of chemical denaturants⁹ or at an elevated temperature.¹⁰

In addition to the highly stable cross- β fibrils (irreversible fibrils), in recent years, labile kinked β -sheet fibrils with

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thermal-responsive reversibility (reversible fibrils) have been reported to be formed by many RNA binding proteins, such as fused-in sarcoma (FUS),^{11–13} TAR DNA-binding protein 43 (TDP-43)^{14,15} and heterogeneous nuclear ribonucleoprotein A1 (hnRNP1)^{13,16} and A2 (hnRNP2).¹⁷ These proteins are capable of undergoing liquid–liquid phase separation (LLPS) to form liquid biomolecular condensates.^{18,19} On the one hand, LLPS is identified as the formation mechanism of the membrane-less organelles (MLOs, composed of proteins and RNAs)^{20–25} and is involved in many biological cellular functions.^{26–28} On the other hand, LLPS plays a critical role in the formation of toxic aggregates in diseases such as amyotrophic lateral sclerosis (ALS)^{29–33} and frontotemporal lobar degeneration (FTLD).^{34–37} Under abnormal conditions, such as mutations or continuous external stimuli, the droplets formed *via* LLPS can turn into solid fibrillary aggregates, a process which is called liquid-to-solid phase transition (LSPT).^{30,37} The amyloid fibrils are a common end-stage product of LSPT.⁷ Recently, the structure of the reversible fibril formed by the fibril core of the FUS low complexity (FUS LC) domain has been determined by solid state nuclear magnetic resonance (ssNMR).¹¹ This fibril core consists of residues 35–95 of FUS (FUS_{35–95}), a 61-residue segment. The FUS_{35–95} fibril, unlike the steric zipper β -sheet fibril, is labile for the lack of intermolecular side chain hydrophobic interactions (or tight interdigitation).³⁸ Apart from through maturation or aging (the time-dependent coarsening and loss of dynamicity) of a liquid-like droplet by RNA binding proteins such as FUS, pathogenic fibrils can also be formed by proteins such as A β directly through irreversible aggregation without LLPS, which is termed liquid–solid phase separation (LSPS).³⁹

In addition to full-length proteins and their LC domains, short peptides are able to phase separate, forming liquid condensates^{40–43} and reversible or irreversible fibrils.^{12,13,15}

Recent crystal structure studies of short peptides have provided different structural properties of the two types of fibrils.³⁹ Unlike the irreversible fibrils, the structures of reversible fibrils are organized in kinked strands perpendicular to the fibril spine, or extended β -strands with a hydrous sheet interface, or a stacking pattern with Asp residues from individual β -strands aligned along the fibril spine.^{7,12,13,16} For example, fibrils of FUS_{37–42} (PDB ID: 5XSG) comprise kinked strands without extended β sheets (Fig. S1A†), and fibrils of FUS_{54–59} (PDB ID: 5XRR) consist of pairs of β -sheets with a hydrophilic interface (Fig. S1B†).¹² In contrast, the irreversible fibril structures of A β _{16–21} (PDB ID: 3OW9)⁴⁴ and Tau_{306–311} (PDB ID: 2ON9)⁴⁵ (Fig. S1C ad S1D†) are composed of stacking β -strands and have large hydrophobic interfaces between β -sheets and surface complementarity. Eisenberg's group also solved the fibril structures of several segments from FUS, hnRNP1 and nup98 proteins,¹³ named LARKS (low-complexity aromatic-rich kinked segment). LARKS forms reversible hydrogel, rather than irreversible fibrils.¹⁵

Reversible fibrils are easily dissociated by heating or adding detergent.^{12,13,16} In the thermostability experiments performed by Liu's group,¹² it was found that fibrils formed from two tandem (S/G)Y(S/G) motifs of FUS, namely ₃₇SYSGYS₄₂ (FUS_{37–42}) and ₅₄SYSSYG₅₉ (FUS_{54–59}) at 4 °C dissolved upon heating to room temperature or higher. When cooled down back to 4 °C, both FUS_{37–42} and FUS_{54–59} fibrils were formed again.¹² In contrast, the amyloid fibrils (termed irreversible fibrils) of A β _{16–21} (Ace-₁₆KLFFFA₂₁) and Tau_{306–311} (Ace-₃₀₆VQIVYK₃₁₁) remained intact during the heating and cooling processes, displaying high thermostability.¹²

In spite of the structural characteristics and thermal stabilities of reversible and irreversible fibrils being well studied experimentally, the dynamic properties and phase behaviors of those reversible/irreversible self-assembling peptides as well as their underlying molecular mechanisms remain largely elusive. In this study, we investigated the reversible/irreversible self-assembly process and predicted the LLPS/LSPS of four hexapeptides, namely FUS_{37–42}, FUS_{54–59}, A β _{16–21} and Tau_{306–311}, using both all-atom molecular dynamic (AA-MD) and coarse-grained MD (CG-MD) simulations combined with explicit water solvents. We dissected the main differences in the dynamic and structural properties of reversible and irreversible peptide self-assembly, characterized the phase behaviors, and elucidated the underlying molecular interactions. Our computational work provides atomic insights into the reversible and irreversible self-assembly of short fibril-forming peptides, which may be helpful for the in-depth mechanistic understanding of LLPS and LSPS of proteins.



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University in December 2017. Her research focuses on understanding the molecular mechanism of the assembly of peptides into ordered nanostructures or amyloid fibrils and the regulation of pathological fibrillization by endogenous proteins with molecular dynamics simulations.

Models and methods

System setup

We selected four representative short peptides as our study models, two from the ALS-related FUS protein, FUS_{37–42} (₃₇SYSGYS₄₂) and FUS_{54–59} (₅₄SYSSYG₅₉),¹² and another two from the Alzheimer's-associated A β /Tau proteins, A β _{16–21}

(Ace-₁₆KLFFFA₂₁)⁴⁴ and Tau_{306–311} (Ace-₃₀₆VQIVYK₃₁₁).⁴⁵ We simulated a total of 12 systems for the four peptides: four all-atom fibril systems, four all-atom oligomer systems, and four coarse-grained oligomer systems. All the systems together with the simulation setups are provided in Table 1.

All-atom fibril systems. The initial fibril structures of the four hexapeptides used in our study consisted of 12 peptide chains (Fig. S1†). They were taken from the fibril structures of FUS_{37–42} (PDB ID: 5XSG) and FUS_{54–59} (PDB ID: 5XRR) solved by Luo *et al.*,¹² and Aβ_{16–21} (PDB ID: 3OW9) and Tau_{306–311} (PDB ID: 2ON9) fibril structures determined by the Eisenberg group.^{44,45} Aβ_{16–21} and Tau_{306–311} fibrils were acetylated at the N-terminal to keep peptides electrostatically neutral, consistent with capped peptides used for the thermostability measurements by Liu's group.¹² These peptide fibrils were placed in the center of a box with a side length of ~7.0 nm.

All-atom oligomer systems. The initial state of each oligomer system consisted of six monomers with random coil conformations, with each monomer randomly placed in the simulation box with dimensions of ~8.5 × 8.5 × 8.5 nm³. Peptide monomers were first constructed using PyMOL⁴⁶ in extended coil states, then simulated under 410 K for 50 ns.

Coarse-grained oligomer systems. In the initial state of each system, 200 monomers were randomly displaced in an aqueous simulation box containing about 43 000 water beads and 527 Na⁺ and 527 Cl[−] ions (~0.15 M NaCl). The simulation box has a size of ~18 × 18 × 18 nm³.

AA-MD simulations

AA-MD simulations were performed using the GROMACS-2016.4 software package⁴⁷ in combination with the AMBER99SB-ILDN force field⁴⁸ and TIP3P water model. Na⁺ and Cl[−] ions at a physiological concentration of 0.15 M were added to all the simulation boxes. We used the Verlet cutoff scheme for neighbor searching and the particle mesh Ewald (PME) method^{49,50} for electrostatic interaction calculations. A real-space cutoff of 1.4 nm was used for electrostatic interaction. The cutoff for van der Waals interaction was also set as 1.4 nm. Periodic boundary conditions were applied for all

simulations. Before the production run of the MD simulations, all the systems were sequentially equilibrated under an NVT (310 K) ensemble for 100 ps, and then under an NPT (310 K and 1 bar) ensemble for 100 ps.⁵¹ After that, each simulation was run for 500 ns, with an integration time step of 2 fs. Velocity rescale⁵² and the Parrinello-Rahman⁵³ methods were used for temperature and pressure coupling, respectively. To obtain statistically significant results, two independent simulations at 310 K were conducted for each fibril system, and five independent simulations at 310 K were performed for each of the oligomer systems. To examine the fluidity of the peptides at a higher temperature, another 500 ns simulation was conducted for each of the four oligomer systems at 340 K, starting from the final state of the MD simulation at 310 K.

CG-MD simulation

All CG-MD simulations were performed using the GROMACS-2018.3 software package. The peptides were described using the MARTINI coarse-grained force field (version 2.2)^{54,55} with modifications described as follows. Electrostatic interactions were calculated using the reaction field method⁵⁶ with a real-space cut-off of 1.4 nm in the CG-MD simulations. The other simulation setups were the same as those used in the AA-MD simulations. Three independent 6 μs simulations were conducted for each system to ensure that each of them reached equilibrium. The previous literature has reported the overestimation of inter-residue interaction strength using the MARTINI forcefield,^{57–59} especially for Ser and Thr residues. Several groups have utilized a scaling factor α to adjust the pair-interaction well depth, which is aimed at reproducing experimental observations.^{57,58} As FUS_{37–42} and FUS_{54–59} peptides are enriched in Ser residues, rescaling the pair interaction is crucial for accurately characterizing their assembly capabilities. However, the selection of the α value is system-dependent, and experimental data for the condensates formed by these two FUS segments are not available. In a recent work, the Hummer group has shown that FUS-LCD condensation occurs above a critical α value of 0.6, and the experimental densities

Table 1 A summary of all the systems and simulation setups

Systems ^a			Number of chains	Number of atoms/beads	Temperature	Simulation time
All-atom	Fibrils	FUS _{37–42}	12	33 575	310 K	500 ns × 2
		FUS _{54–59}		33 584		
		Aβ _{16–21}		33 653		
		Tau _{306–311}		33 620		
	Oligomers	FUS _{37–42}	6	60 134	310 K	310 K: 500 ns × 5
		FUS _{54–59}		60 137		
		Aβ _{16–21}		60 161	340 K	340 K: 500 ns × 1
		Tau _{306–311}		60 203		
Coarse-grained	Condensates	FUS _{37–42}	200	46 854	310 K	6 μs × 3
		FUS _{54–59}		46 854		
		Aβ _{16–21}		47 054		
		Tau _{306–311}		46 854		

^a FUS_{37–42}: ₃₇SYSGYS₄₂; FUS_{54–59}: ₅₄SYSSYG₅₉; Aβ_{16–21}: Ace-₁₆KLFFFA₂₁; Tau_{306–311}: Ace-₃₀₆VQIVYK₃₁₁.

of both the dilute and dense phases were accurately replicated at $\alpha = 0.65$.⁵⁸ In spite of a much shorter sequence length, FUS_{37–42} and FUS_{54–59} have been identified as two reversible amyloid cores, which mediate the dynamic assembly of FUS LCD.¹² We thus infer that $\alpha = 0.65$ may also be suitable for studying the self-assembly process of FUS_{37–42} and FUS_{54–59}. To test this hypothesis, we selected six α values (0.2, 0.4, 0.5, 0.6, 0.65, and 0.8) and performed short (600 ns) simulations to compare their performance in characterizing the assembly capability of the peptides. The scaling was performed following the method of Stark *et al.*⁵⁵ by down-scaling the van der Waals parameters between Ser pseudo-atoms and the backbone beads of all residues, $\epsilon_\alpha = \epsilon_0 + \alpha(\epsilon_{\text{original}} - \epsilon_0)$. A value of $\alpha = 0$ corresponds to a repulsion-dominated interaction in the MARTINI model ($\epsilon_0 = 2 \text{ kJ mol}^{-1}$), and a value of $\alpha = 1$ recovers the full interaction in the MARTINI force field ($\epsilon_1 = \epsilon_{\text{original}}$). The results suggest that FUS_{37–42} and FUS_{54–59} presented increasing assembly capability with increasing scaling factor values (Fig. S2†). They displayed weak assembly capabilities at low α values (<0.6), especially FUS_{37–42}, while they presented higher assembly ability than A β _{16–21} and Tau_{306–311} at a high α value (0.8). At a moderate α value of 0.65, FUS_{37–42} and FUS_{54–59} possessed a certain degree of assembly capability, which was lower than that of A β _{16–21} and Tau_{306–311} in good agreement with previous experimental observations.¹² We thus selected 0.65 as the scaling factor for performing the coarse-grained simulations. To offer a quantitative comparison with all-atom simulations, we conducted coarse-grained simulations for each of the four peptides starting from six randomly dispersed peptide chains. At a scaling factor of $\alpha = 0.65$, the collapse degree and SASA fluctuation values predicted by coarse-grained simulations were of the same order of magnitude as those predicted by the all-atom simulations (Fig. S3†). Moreover, both simulations predicted that FUS_{37–42}/FUS_{54–59} possessed lower assembly capability and higher liquidity than A β _{16–21}/Tau_{306–311}.

Analysis methods

Data analysis was performed using the tools implemented in the GROMACS software package and our in-house developed codes. For data analysis of the AA-MD simulations, an atomic contact was defined as being when two aliphatic carbon atoms were within 0.54 nm of one another or when any other two heavy atoms came within 0.46 nm.^{60–62} When two monomers were located with a minimum distance between heavy atoms of less than 0.54 nm, they were regarded as belonging to one cluster. The intra- and inter-chain dihedral correlations were evaluated following previous literature.⁶³ The correlation coefficients between two dihedrals (x and y) were calculated utilizing the method put forward by Jammalamadaka and Sengupta,⁶⁴ using the following equation:

$$\rho_{xy}^{\text{circular}} = \frac{\sum_{i=1}^N \sin(x_i - \bar{x}) \cdot \cos(y_i - \bar{y})}{\sqrt{\sum_{i=1}^N \sin^2(x_i - \bar{x}) \cdot \sum_{i=1}^N \sin^2(y_i - \bar{y})}}$$

A hydrogen bond (H-bond) was considered to be formed on the basis of two conditions:^{65–67} (1) the distance between the H-bond donor (D) and the acceptor (A) is less than 0.35 nm, and (2) the angle of D–H...A is larger than 150°. The secondary structure was analyzed using the DSSP tool.⁶⁸ The end-to-end distance of a monomer refers to the distance between the first and the last C α atoms of the monomer. Two aromatic residues were in the π – π stacking state when the distance between the centroids of two aromatic rings was less than 0.7 nm.⁶⁹ Three different types of stacking patterns were defined according to the angles of the two aromatic rings: 0°–30° for parallel, 30°–60° for herringbone, and 60°–90° for T-shape. The statistical results of the AA-MD simulations were all based on the data from 300 to 500 ns.

In the data analysis of CG-MD simulations, two monomers were considered to have molecular contact if their minimum distance was less than 0.6 nm. In accordance with our recent study, the aggregation propensity of each peptide was characterized using the collapse degree and the clustering degree.⁴³ The collapse degree was defined as the ratio of SASA of all hexapeptide molecules in the initial state to their SASA in the final configuration.⁷⁰ The clustering degree was defined by molecules in clusters divided by the total number of molecules in the system.⁴³ The fluidity of an aggregate was characterized using the fluctuation of SASA, fluctuation of the clustering degree and the exchange rate of the interactions.⁴³ The statistical results of the CG-MD simulations were all based on data from 3 to 6 μ s. All the structure representations were drawn using the VMD program⁷¹ or PyMOL.⁴⁶

Results and discussion

Fibrils formed by the four hexapeptides display different thermoresponsive stabilities

We first examined the structural stability of the preformed fibrils of the FUS_{37–42}, FUS_{54–59}, A β _{16–21} and Tau_{306–311} peptides. Two independent 500 ns simulations were carried out at 310 K for each fibril system and the results are shown in Fig. 1. It is noted that the structures of the FUS_{37–42}/FUS_{54–59} fibril and the A β _{16–21}/Tau_{306–311} fibril were solved at 4 °C and 18 °C,^{12,44,45} which were lower than our simulation temperature (310 K). As seen in Fig. 1a and b, the C α -root-mean-square-deviation (C α -RMSD) and the radius of gyration (R_g) values of both FUS_{37–42} and FUS_{54–59} far exceeded those of A β _{16–21} and Tau_{306–311} and fluctuated considerably, indicating that the fibril structures of FUS_{37–42} and FUS_{54–59} were much less stable than those of the A β _{16–21} and Tau_{306–311} fibril structures. The fibril structures of FUS_{37–42} and FUS_{54–59} were completely disrupted in the final state. In contrast, both the C α -RMSD and R_g curves of A β _{16–21} and Tau_{306–311} maintained a plateau at small values (Fig. 1a and b), although the fibril structures of A β _{16–21} and Tau_{306–311} were twisted and the β -sheet contents decreased a bit (Fig. 1d). Then, the side chain dynamics of the four systems were analyzed by calculating the rate of side chain dihedral angle (χ_1) transition between the

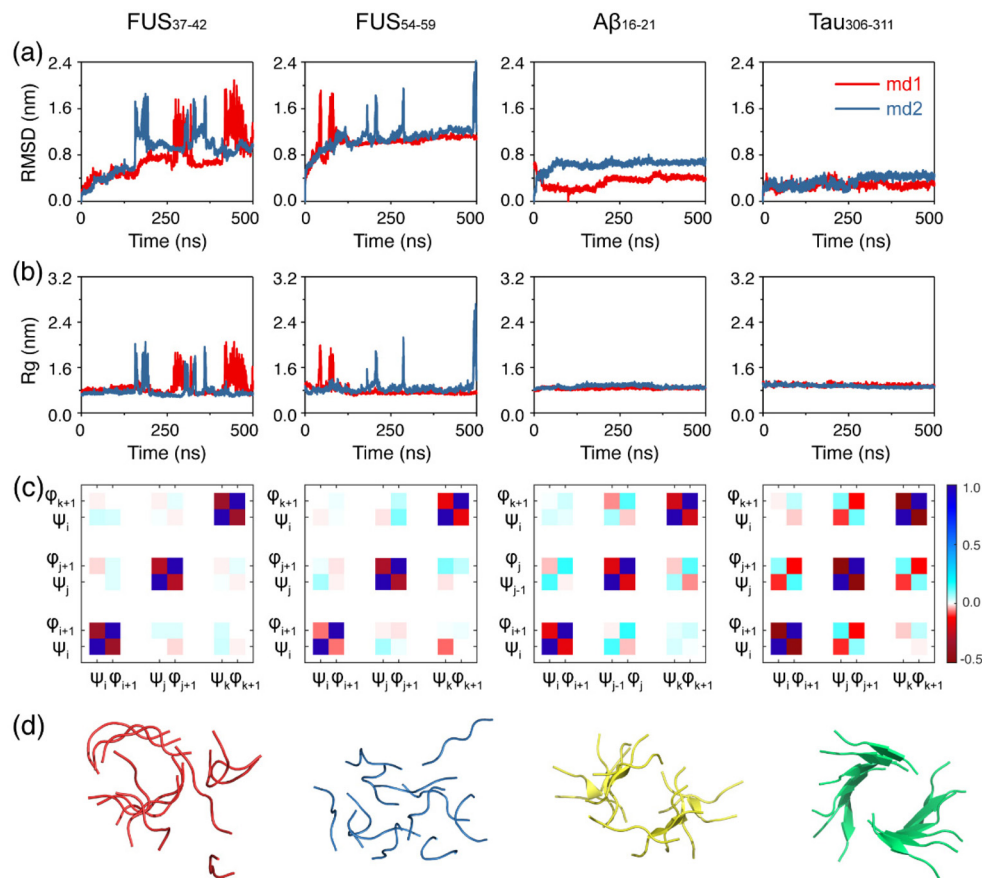


Fig. 1 Thermostability of the preformed FUS₃₇₋₄₂, FUS₅₄₋₅₉, Aβ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ fibrils. Time evolution of (a) RMSD and (b) R_g values of the FUS₃₇₋₄₂, FUS₅₄₋₅₉, Aβ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ fibrils. (c) The dihedral angle correlation coefficients within and between the peptide chains in the four fibril systems. (d) Snapshots of the final states of the four fibril systems. Peptides were drawn using PyMOL.

$C^{\gamma-exo}$ and $C^{\gamma-endo}$ conformations.⁷² It can be seen from Fig. S4† that FUS₃₇₋₄₂ and FUS₅₄₋₅₉ had much higher χ_1 transition rates (>2.0) than Aβ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ (<1.2), suggesting a much higher degree of side chain dynamics of the LARKS peptides. This high degree of dynamics likely played a role in the low stability of the FUS₃₇₋₄₂ and FUS₅₄₋₅₉ fibrils. The previous literature has revealed that correlated backbone motions are one of the fundamental properties of the β -sheet.⁶³ To investigate the backbone correlation of the four fibrils, we calculated the circular cross-correlation coefficients of main chain dihedrals within and between the fibril chains (Fig. 1c). The definitions of the dihedral angles for parallel and antiparallel β -strands are given in Fig. S5.† Strong local anticorrelations are observed between the φ and ψ values of each peptide chain in all of the four systems. This anticorrelation is consistent with the previous literature and indicates the rigidity of the peptide plane, which couples the motion of the dihedral angles while preserving the structure of the strands. Strong inter-chain correlations are also observed between the adjacent chains in the Aβ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ systems, indicating the stability of their fibrils. Interestingly, in the Aβ₁₆₋₂₁ system φ - φ and ψ - ψ dihedral pairs are correlated while the φ - ψ pair is anticorrelated, and *vice versa* in the Tau₃₀₆₋₃₁₁ system. This is

due to the antiparallel and parallel nature of the Aβ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ β -sheets. In contrast, the inter-chain correlations of the FUS₃₇₋₄₂ and FUS₅₄₋₅₉ systems are much weaker, further demonstrating the instability of their fibrils. The computationally observed lower stability of the LARK segments FUS₃₇₋₄₂/FUS₅₄₋₅₉ than that of the Aβ₁₆₋₂₁/Tau₃₀₆₋₃₁₁ peptides is consistent with previous experimental observations,¹² demonstrating the accuracy and suitability of the force field and the methodology that we used.

The FUS₃₇₋₄₂/FUS₅₄₋₅₉ peptides form highly dynamic oligomers, whereas the Aβ₁₆₋₂₁/Tau₃₀₆₋₃₁₁ peptides assemble into stable oligomers

After the structural stability examination of the fibril structures, we investigated the self-assembly processes of the four hexapeptides using AA-MD simulations. It is shown in Fig. 2a that the numbers of clusters of FUS₃₇₋₄₂ and FUS₅₄₋₅₉ vary during the oligomerization process. The monomers of FUS₃₇₋₄₂ and FUS₅₄₋₅₉ gathered together first, then were separated, and came together again, indicating that their self-assembly processes are highly dynamic, and their aggregates are labile and easy to dissolve. In the Aβ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ systems, in marked contrast, most monomers self-assembled

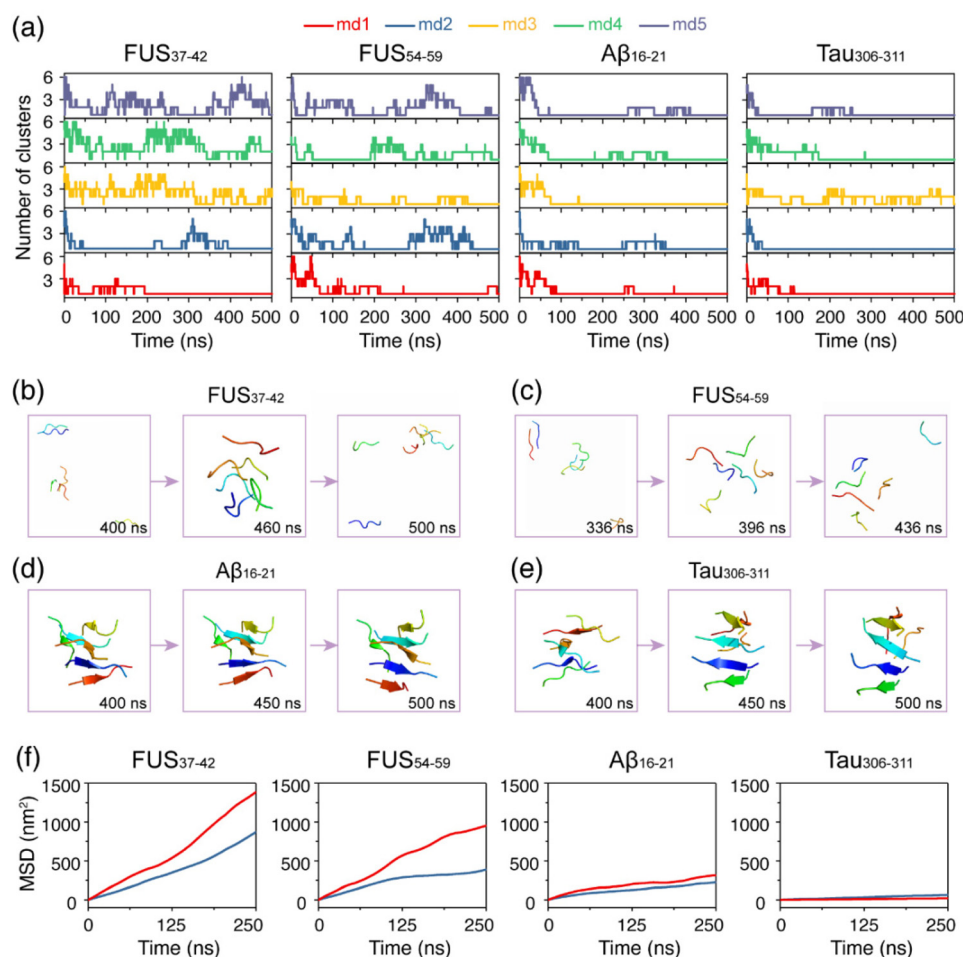


Fig. 2 The self-assembly properties of the FUS₃₇₋₄₂, FUS₅₄₋₅₉, Aβ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ hexapeptides into oligomers. (a) Time evolution of the number of clusters formed by each peptide at 310 K. (b–e) Snapshots of the representative states in one out of the five simulations for each system. Peptides were drawn in cartoon form using PyMOL with different chains in different colors. (f) Mean square displacement (MSD) of the four systems at 310 and 340 K. The MD simulations at 340 K started from the final states obtained at 310 K.

into stable aggregates, reflecting the irreversibility of their self-assembly processes. Additionally, side chain dihedral transition rates were also examined. Much higher transition rates were observed for FUS₃₇₋₄₂ and FUS₅₄₋₅₉ than for the Aβ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ oligomers, indicative of the higher side chain dynamics of the LARKS peptides (Fig. S4†). The different self-assembly dynamics were also demonstrated in the representative states of the four systems shown in Fig. 2b–e.

To further explore the thermostability of the oligomers formed at 310 K, we performed a 500 ns MD simulation for each system at a higher temperature of 340 K starting from the final state obtained at 310 K. Fig. 2f shows that the slope of the mean square displacement (MSD) curves of FUS₃₇₋₄₂ and FUS₅₄₋₅₉ are both very large at 340 K and there is a big gap between the curves at 310 K and those at 340 K, suggesting strong temperature dependence of their fluidity. In sharp contrast, the MSD values of Aβ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ are quite similar at both temperatures, suggesting poor temperature dependence of their fluidity. Moreover, Aβ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁

have much smaller MSD values than FUS₃₇₋₄₂ and FUS₅₄₋₅₉, indicating their weaker mobility at both temperatures. Overall, the LARK segments FUS₃₇₋₄₂ and FUS₅₄₋₅₉ self-assemble into dynamic aggregates with high mobility and low thermal stability, while Aβ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ self-assemble into more stable aggregates.

Peptide chains in the FUS₃₇₋₄₂ and FUS₅₄₋₅₉ oligomers tend to be kinked with low β-sheet contents, while those in the Aβ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ oligomers are extended with high β-sheet contents

We further analyzed the structural features of the oligomers formed by each of the four peptides. Fig. 3a and b shows the probability density function (PDF) of the end-to-end distance of all the peptide chains in each system. It can be seen that the end-to-end distance distribution curves of the FUS₃₇₋₄₂ and FUS₅₄₋₅₉ peptides have two wide peaks, located at around 0.5 nm and 1.4 nm (Fig. 3a), respectively, corresponding to the collapsed and extended single-chain conformations (illustrated

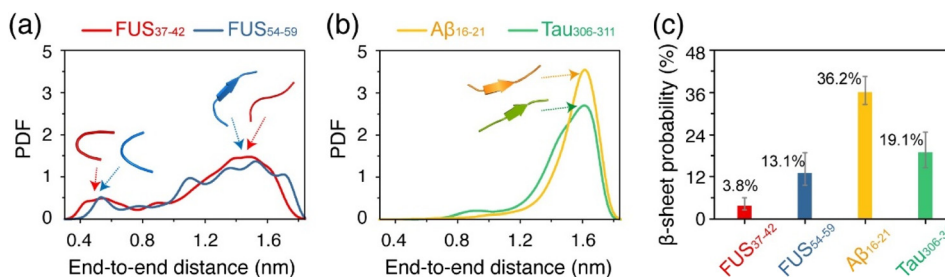


Fig. 3 The structural properties of the oligomers formed by each of the four different peptides. Probability density function (PDF) of the distance between the first and the last C α atoms of all single chains in (a) the FUS₃₇₋₄₂ and FUS₅₄₋₅₉ systems and (b) the Aβ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ systems. Arrows in (a) and (b) show the extended/kinked and extended single chain conformations. (c) β-sheet probability of the four systems. Data are from 300 to 500 ns. The error bars were calculated using a bootstrap procedure of 50 resamplings, with a confidence interval of 10%.

by the representative snapshots in the dashed boxes). In contrast, the curves of the Aβ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ peptides have only one peak at around 1.6 nm (Fig. 3b). The results suggest that the peptide chains in the FUS₃₇₋₄₂/FUS₅₄₋₅₉ oligomers adopted both extended and collapsed conformations, whereas those in the Aβ₁₆₋₂₁/Tau₃₀₆₋₃₁₁ oligomers had mostly extended conformations. Moreover, the FUS₃₇₋₄₂/FUS₅₄₋₅₉ chains (with an end-to-end distance of 1.4 nm) were less extended than the Aβ₁₆₋₂₁/Tau₃₀₆₋₃₁₁ chains (1.6 nm), indicative of the kinked conformations of FUS₃₇₋₄₂/FUS₅₄₋₅₉ peptides. Collapsed peptides were likely to form more intra-molecular interactions, while extended peptides tended to form more inter-molecular interactions. Therefore, the peak at 0.5/1.4 nm of the distance distribution curves of the FUS₃₇₋₄₂ and FUS₅₄₋₅₉ systems signified that the collapsed/kinked peptides were less likely to form stable inter-molecular interactions, which may disfavor the formation of stable cross-β structures.

The β-sheet probability of the peptides in the four systems was calculated and is shown in Fig. 3c. The four systems with β-sheet contents from high to low are: Aβ₁₆₋₂₁ (36.2%), Tau₃₀₆₋₃₁₁ (19.1%), FUS₅₄₋₅₉ (13.1%) and FUS₃₇₋₄₂ (3.8%). Intriguingly, the FUS₃₇₋₄₂ (SYSYGS) oligomers had much lower β-sheet content than the FUS₅₄₋₅₉ (SYSSYG) oligomers, although the two peptides had the same amino acid composition (see below for a more detailed discussion). These results demonstrate that both Aβ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ peptides had a higher propensity to form β-sheet structures than the FUS₃₇₋₄₂ and FUS₅₄₋₅₉ peptides. Considering that β-sheets form the spines of stable amyloid fibrils, the different β-sheet preferences of the FUS₃₇₋₄₂/FUS₅₄₋₅₉ and Aβ₁₆₋₂₁/Tau₃₀₆₋₃₁₁ peptides may lead to their reversible and irreversible self-assembly.

The alternating stickers and spacers in the FUS₃₇₋₄₂/FUS₅₄₋₅₉ peptides result in the formation of reversible aggregates, while hydrophobic patches in the Aβ₁₆₋₂₁/Tau₃₀₆₋₃₁₁ peptides are responsible for their irreversible self-assembly

After examining the reversible and irreversible self-assembly properties of the two types of peptides, we further explored the underlying driving forces by calculating the number of inter-molecular contacts and H-bonds (Fig. 4). There were similar numbers of contacts between the main chains (MC-MC)

among the four systems, while the contacts between the side chains (SC-SC) were considerably different, *i.e.* much higher in the Aβ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ systems than in the FUS₃₇₋₄₂ and FUS₅₄₋₅₉ systems (Fig. 4a). These results indicate that the SC-SC contacts in the four systems were the main interactions driving the reversible and irreversible self-assembly of the FUS₃₇₋₄₂/FUS₅₄₋₅₉ and Aβ₁₆₋₂₁/Tau₃₀₆₋₃₁₁ peptides. In addition, more MC-MC H-bonds were formed in the FUS₅₄₋₅₉, Aβ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ systems than in the FUS₃₇₋₄₂ system (Fig. 4b), consistent with the results of the formation of more β-sheets in the former three systems. Fig. 4b also shows that the SC-SC H-bonds only accounted for a minor part of the forces driving the peptide self-assembly in all the systems. Moreover, the numbers of H-bonds formed were different in the FUS₃₇₋₄₂ and FUS₅₄₋₅₉ oligomers. Although having the same amino acid compositions and containing the tandem SYS and SYG/GYS motifs, FUS₅₄₋₅₉ (SYSSYG) with Ser in the middle of the sequence had a greater ability to form intermolecular H-bonds than FUS₃₇₋₄₂ (SYSYGS) with Gly in the middle of the sequence, indicating that the Ser residue at middle positions had more significant influences on peptide self-assembly than the Gly residue and that the site-specific patterning of the spacer residues (Ser and Gly) can subtly tune the self-assembly behavior.

We note that there is an increase in the average contact number for the Aβ₁₆₋₂₁ systems starting from ~300 ns. To examine whether the contact number further rises with the increase of simulation time, we extended all five Aβ₁₆₋₂₁ oligomer simulations for an additional 150 ns and calculated the time evolution of their main chain and side chain contact numbers (Fig. S6†). For some simulations, the contact numbers remained mostly constant after 500 ns (such as the main chain contact for MD3 and the side chain contact for MD5). For most simulations, the contact numbers fluctuated around their equilibrium values. Importantly, trends of continued growth were not observed after 500 ns, suggesting that the 500 ns duration was sufficient to reach a convergence for these contact numbers.

The intermolecular MC-MC and SC-SC residue-wise contacts and H-bonds were also analyzed (Fig. 4c, d and S7, S8†). As seen from Fig. 4c, only Tyr residues, separated by Ser and Gly, had strong interactions with each other in the FUS₃₇₋₄₂ and FUS₅₄₋₅₉ systems, suggesting that Tyr residues are crucial

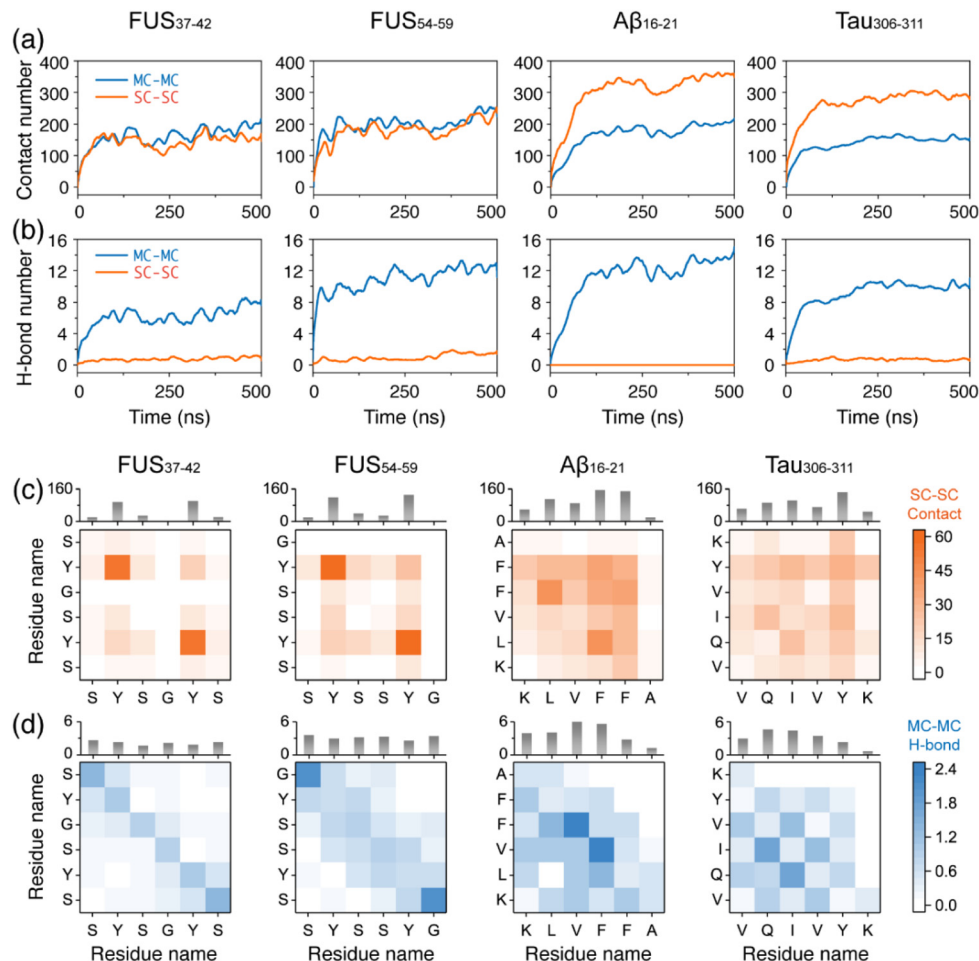


Fig. 4 The driving forces underlying the reversible self-assembly of the FUS₃₇₋₄₂/FUS₅₄₋₅₉ peptides and irreversible self-assembly of the A β ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ peptides. Time evolution of (a) main chain–main chain (MC–MC) and side chain–side chain (SC–SC) contacts, as well as (b) MC–MC and SC–SC H-bonds in the FUS₃₇₋₄₂, FUS₅₄₋₅₉, A β ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ systems. Inter-molecular residue-wise (c) SC–SC contacts and (d) MC–MC H-bonds of the four systems. The grey bar charts show the cumulative contact numbers between each residue and other residues.

stickers of FUS₃₇₋₄₂ and FUS₅₄₋₅₉. The important role of Tyr and repeated motifs containing Tyr in contributing to protein LLPS has been reported in many studies.⁷³⁻⁷⁵ The amphipathic side chain of Tyr allowed it to form a hydrogen bond *via* its phenol group, as well as π - π interaction through the aromatic ring, the latter of which is important for the assembly of various amyloid peptides into β -rich fibrils,^{76,77} and has emerged as a major driver of LLPS.⁷⁸⁻⁸⁰ The intermolecular π - π stacking interactions were calculated between aromatic rings in the four systems. It is shown in Fig. S9† that more π - π stacking interactions were formed in the FUS₃₇₋₄₂ and FUS₅₄₋₅₉ systems than in the A β ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ systems, although the number of aromatic rings was the same in FUS₃₇₋₄₂, FUS₅₄₋₅₉ (containing two Tyr residues) and A β ₁₆₋₂₁ (containing two Phe residues). The distribution patterns of the π stacking angle were alike in the four all-atom oligomer systems, mainly preferring perpendicular stacking, except for FUS₅₄₋₅₉ with an equal preference for T-shape stacking. Although with strong SC–SC interactions *via* π - π stacking, H-bonding and hydrophobic interactions between Tyr residues, the occurrence of

the polar spacers of Ser and Gly impeded the formation of successive strong interactions in the FUS₃₇₋₄₂ and FUS₅₄₋₅₉ systems, but enabled transient intermolecular interactions favorable for LLPS. The sticker-spacer architecture is ubiquitous in LLPS-prone proteins⁸¹ and the sticker-sticker, spacer-spacer and sticker-spacer interactions interplay⁸² during their LLPS process. FUS₃₇₋₄₂ and FUS₅₄₋₅₉ peptides possess alternating stickers (Tyr) and spacers (Ser and Gly), and the transient interactions among stickers and spacers may facilitate the reversible self-assembly. In contrast, the central consecutive hydrophobic residues LVFF in A β ₁₆₋₂₁ and IVY in Tau₃₀₆₋₃₁₁, acting as hydrophobic patches, have higher contact numbers with all residues through side chain contacts. The hydrophobic patch provides persistent interactions, contributing to peptide irreversible self-assembly. Additionally, the correlations between side chain and main chain dynamics were examined by calculating the Pearson correlation coefficients between main chain dihedrals ϕ/ψ and side chain dihedral χ_1 . Our calculations suggest a low main chain–side chain correlation, with correlation coefficients <0.3 (Fig. S10 and S11†). These

results further demonstrate that main chain and side chain play different roles in the assembly process.

The role of water in the self-assembly process of these peptides was examined by calculating the number of hydrogen bonds (H-bonds) formed between the side chain of each residue and water molecules (Fig. S12†). The results suggest that the FUS_{37–42}/FUS_{54–59} peptides formed much more H-bonds with water through five out of six residues (Ser and Tyr). In contrast, only three residues (Gln, Tyr, and Lys) in the Tau_{306–311} peptides, and only one residue (Lys) in the A β _{16–21} peptide formed side chain H-bonds with water. These results are consistent with the lower assembly capability and the higher fluidity of FUS_{37–42}/FUS_{54–59} condensates than A β _{16–21}/Tau_{306–311} aggregates, and demonstrate the importance of water in the reversible/irreversible assembly process. A previous MD study⁸³ has reported that the interaction of water plays an important role in the fibrillization of the Gln-containing polar heptapeptide, GNNQQNY, from the prion domain of Sup35. Interestingly, among all residues in the four peptides,

the glutamine residue in the Tau_{306–311} peptide formed the highest number of H-bonds with water molecules, suggesting its role in mediating the assembly of Tau_{306–311}.

Taken together, the aromatic sticker residue (Tyr) in the LARK segments, FUS_{37–42} (SYSGYS) and FUS_{54–59} (SYSSYG), are separated by spacer residues (Ser and Gly). The alternating stickers and spacers cause FUS_{37–42} and FUS_{54–59} to self-assemble into reversible aggregates. In contrast, the side chain interactions between the hydrophobic patches cause irreversible self-assembly of the A β _{16–21} and Tau_{306–311} peptides. These two peptides self-assemble into aggregates with high β -sheets. Side chain hydrophobic interactions and main chain H-bonding interactions collectively stabilize the aggregates (Fig. 4c and d).

FUS_{37–42}, FUS_{54–59}, A β _{16–21} and Tau_{306–311} present distinct phase separation propensities

An increasing number of studies have demonstrated that LLPS is a general property of peptides and proteins,^{18,19,25} and even

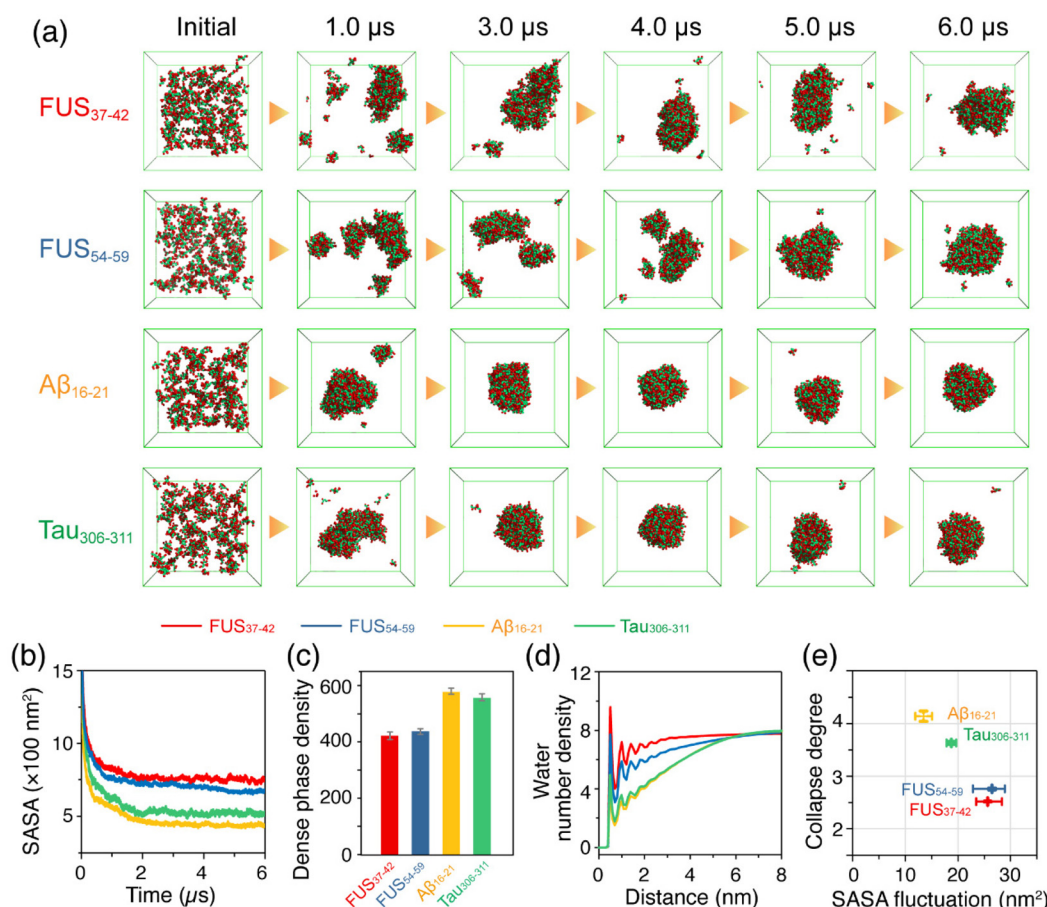


Fig. 5 Phase separation behaviors of the FUS_{37–42}, FUS_{54–59}, A β _{16–21} and Tau_{306–311} hexapeptides predicted by multiple microsecond CG-MD simulations. (a) Representative snapshots of each system at six time points in 6 μ s CG-MD simulations. (b) Time evolution of SASA of the FUS_{37–42}, FUS_{54–59}, A β _{16–21} and Tau_{306–311} peptides. Values were averaged over three independent 6 μ s MD simulations for each system. (c) Concentrations of the dense phase of the four systems. (d) Number density of water molecules as a function of distance from the center of the condensed phase in the four systems. (e) The collapse degree values and the fluctuation of SASA of the four systems. Detailed descriptions of the parameters used are listed in the Models and Methods section.

short peptides possess sufficient interaction sites for LLPS.^{42,43} In order to predict the phase separation propensity of the hexapeptides, we carried out CG-MD simulations on large space and time scales based on a modified MARTINI-2.2 force field with a down-scaling of the van der Waals parameters between Ser pseudo-atoms and the backbone beads of all the amino acid residues in the FUS₃₇₋₄₂ and FUS₅₄₋₅₉ systems.^{54,55} Three individual 6 μ s CG-MD simulations were carried out for each of the hexapeptide systems: FUS₃₇₋₄₂, FUS₅₄₋₅₉, A β ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁, starting from the initial states with 200 monomers randomly dispersed in the box. A detailed description of the scaling method is given in the Models and Methods section. The representative snapshots of the four hexapeptide systems in Fig. 5a show that FUS₃₇₋₄₂, FUS₅₄₋₅₉, Tau₃₀₆₋₃₁₁ and A β ₁₆₋₂₁ were all able to self-assemble into a dense phase with ascending phase separation capabilities qualitatively consistent with the self-assembly propensity in the AA-MD simulations of the oligomer systems. Specifically, the FUS₃₇₋₄₂ and FUS₅₄₋₅₉ condensates exhibited a loosely packed feature and were highly dynamic. In contrast, A β ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ condensates were more compact and remained relatively stable in shape as the simulation time increased. The FUS₃₇₋₄₂ and FUS₅₄₋₅₉ aggregates had a larger solvent accessible surface area (SASA) than the A β ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ aggregates (Fig. 5b), suggesting lower phase separation propensities. The concentrations of the FUS₃₇₋₄₂ and FUS₅₄₋₅₉ dense phases were lower than those of the A β ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ dense phases (Fig. 5c). These results indicate that the assemblies formed by the A β ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ peptides were probably hydrogel-like with a high degree of solvent exclusion, and FUS₃₇₋₄₂ and FUS₅₄₋₅₉ assemblies were more like liquid droplets containing water molecules (Fig. 5d, larger number density of water molecules).

Additionally, the cross-sections of the assemblies formed by the four peptides were provided to further demonstrate the presence of water molecules in the condensates. It can be seen from Fig. S13[†] that the FUS₃₄₋₄₂ and FUS₅₄₋₅₉ assemblies were embedded with water molecules. These water molecules may have contributed to their fluidities and also played a role in stabilizing the FUS₃₇₋₄₂/FUS₅₄₋₅₉ condensates. In contrast, the A β ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ assemblies were densely packed and solid-like. Therefore, we infer that the LARK segments FUS₃₇₋₄₂ and FUS₅₄₋₅₉ underwent LLPS, and the A β ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ peptides underwent LSPS. To more quantitatively characterize the different properties of the phase separations of the four peptides, we selected two order parameters, namely the collapse degree and the fluctuation of SASA (Fig. 5e), to manifest the phase separation propensities and the liquidity of the dense phase, respectively. It is shown that the aggregates formed by the A β ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ peptides have a larger collapse degree and smaller SASA fluctuation, while those formed by the FUS₃₇₋₄₂ and FUS₅₄₋₅₉ peptides have a smaller collapse degree and larger SASA fluctuation, indicative of the higher fluidity of the FUS₃₇₋₄₂/FUS₅₄₋₅₉ aggregates.

Collectively, our CG-MD simulations suggest that the LARK segments FUS₃₇₋₄₂ and FUS₅₄₋₅₉ have low phase separation propensities and their condensates are highly dynamic and loosely packed with high mobility. In contrast, the A β ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ peptides have high phase separation propensities and the aggregates are more like solids with low degrees of liquidity. Therefore, we postulate that FUS₃₇₋₄₂ and FUS₅₄₋₅₉ undergo liquid-liquid phase separation, while A β ₁₆₋₂₁ and Tau₃₀₆₋₃₁₁ peptides undergo liquid-solid phase separation.

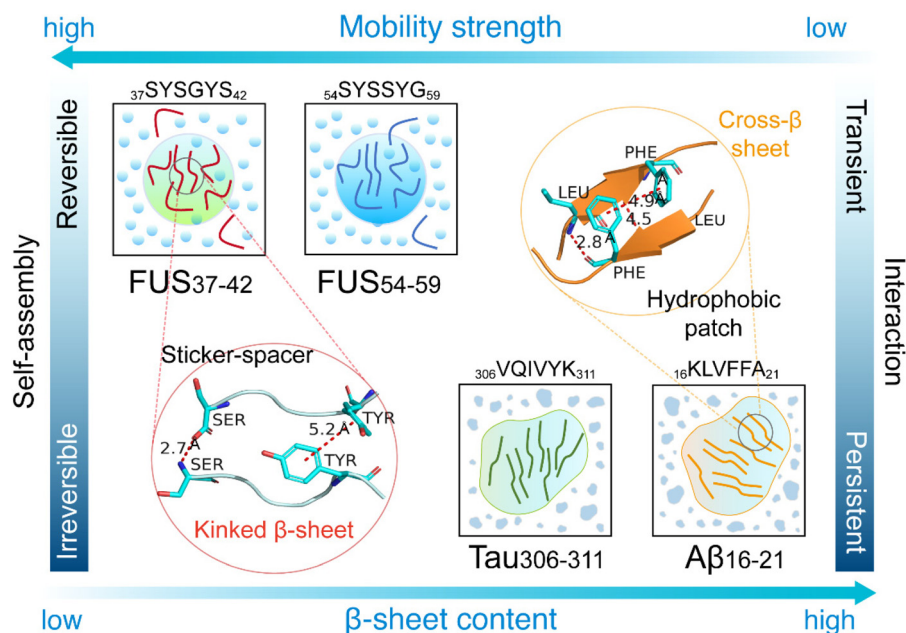


Fig. 6 Schematic diagram illustrating the reversible and irreversible self-assembly behaviors and mechanisms of the four hexapeptides.

Conclusions

In summary, we have investigated the reversible and irreversible self-assembly of four representative hexapeptides, FUS_{37–42}, FUS_{54–59}, A β _{16–21} and Tau_{306–311}, by performing AA-MD and CG-MD simulations. We characterized the self-assembly dynamics of the four hexapeptides and predicted their LLPS/LSPS propensities. Our simulations showed that the four hexapeptides self-assembled into aggregates with different physical properties. The LARKS peptides, FUS_{37–42} and FUS_{54–59}, formed coil-rich aggregates with high mobility and poor thermostability, while A β _{16–21} and Tau_{306–311} self-assembled into β -sheets with high thermostability. Interaction analyses indicated that the persistent interactions between the hydrophobic patches, namely LVFF in A β _{16–21} and IVY in Tau_{306–311}, acted as the main forces driving the irreversible self-assembly of A β _{16–21} and Tau_{306–311} and maintaining the stability of the irreversible aggregates. In contrast, the alternating sticker (Tyr) and spacers (Ser and Gly) in the FUS_{37–42} and FUS_{54–59} peptides provided transient interactions, which were strong enough to drive reversible self-assembly or LLPS and yet weak enough to inhibit irreversible aggregation or LSPS. Intriguingly, the FUS_{37–42} (SYSGYS) and FUS_{54–59} (SYSSYG) peptides had the same sequence composition and the same number of sticker and spacer residues, but exhibited different self-assembly properties, suggesting that the site-specific patterning of spacer residues (Ser and Gly) had a non-negligible effect on peptide self-assembly behaviors. Fig. 6 offers a visual illustration of the varying assembly behaviors and mechanisms of these hexapeptide aggregates. Our current work reveals the molecular mechanisms of sequence-dependent reversible and irreversible self-assembly and the phase separation behaviors of short fibril-forming peptides from amyloid proteins and LLPS-prone proteins, paving the way for further in-depth understanding of reversible LLPS and irreversible aggregation of disease-related proteins including FUS, TDP-43, and tau proteins.

Author contributions

G. W., C. G., and Z. L. conceived the project and constructed the systems. Z. L. performed the all-atom simulations and analyzed the simulation data. Y. T. performed the coarse-grained simulations. G. W. and Z. L. drafted the manuscript. C. G. and G. W. edited the manuscript. All authors analyzed the data, and reviewed and approved the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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