Aromatic Stacking Facilitated Self-Assembly of Ultrashort Ionic Complementary Peptide Sequence: β-Sheet Nanofibers with Remarkable Gelation and Interfacial Properties


ABSTRACT: Understanding peptide self-assembly mechanisms and stability of the formed assemblies is crucial for the development of functional nanomaterials. Herein, we have adopted a rational design approach to demonstrate how a minimal structural modification to a nonassembling ultrashort ionic self-complementary tetrapeptide FEFK (Phe4) remarkably enhanced the stability of self-assembly into β-sheet nanofibers and induced hydrogelation. This was achieved by replacing flexible phenylalanine residue (F) by the rigid phenylglycine (Phg), resulting in a constrained analogue PhgEPhePhgK (Phg4), which positioned aromatic rings in an orientation favorable for aromatic stacking. Phg4 self-assembly into stable β-sheet ladders was facilitated by π-staking of aromatic side chains alongside hydrogen bonding between backbone amides along the nanofiber axis. The contribution of these noncovalent interactions in stabilizing self-assembly was predicted by in silico modeling using molecular dynamics simulations and semiempirical quantum mechanics calculations. In aqueous medium, Phg4 β-sheet nanofibers entangled at a critical gelation concentration ≥20 mg/mL forming a network of nanofibrous hydrogels. Phg4 also demonstrated a unique surface activity in the presence of immiscible oils and was superior to commercial emulsifiers in stabilizing oil-in-water (O/W) emulsions. This was attributed to interfacial adsorption of amphiphilic nanofibrils forming nanoemulsified microspheres. To our knowledge, Phg4 is the shortest ionic self-complementary peptide rationally designed to self-assemble into stable β-sheet nanofibers capable of gelation and emulsification. Our results suggest that ultrashort ionic-complementary constrained peptides or UICPs have significant potential for the development of cost-effective, sustainable, and multifunctional soft bionanomaterials.

INTRODUCTION

Molecular self-assembly has been exploited in nature for the engineering of complex higher macromolecular structures of the proteome. Inspired by nature, the bottom-up design of de novo self-assembling peptides has been extensively investigated in the last three decades to innovate a wide variety of stable nanostructures for biomedical applications.1−5 Zhang and co-workers were the first to propose the self-assembling ionic self-complementary peptide sequence pattern, with alternation of hydrophobic (A) and hydrophilic charged (B) amino acids: (ABAB)n, where n is the number of the pattern repeats.6−9 This design was inspired by the unusual sequence pattern EAK16 identified in a segment of the Z-DNA binding protein Zuotin, which was purified from the nuclear extract of the baker’s yeast Saccharomyces cerevisiae.6

Peptides based on this sequence pattern were shown to self-assemble into stable β-sheet nanofibers, with typically ~8−16 residues (n = 2−4) required to maintain the thermodynamic stability of the assembled nanostructures.6,7−9,11 Interestingly, the tetrapeptide FEFK (Phe4), which is the shortest reported sequence with the ABAB pattern (n = 1), failed to self-assemble into stable β-sheet structures, implicating the importance of the sequence length for the stability of the formed assemblies (Figure 1A).8,9 This could possibly be attributed to the fact that longer sequences provide more backbone amide bonds readily available for intermolecular hydrogen bonding between peptide chains, stabilizing the assembled β-sheet structures.

Herein, we investigated whether the structural modification of Phe4 to the more constrained phenylglycine based analogue Phg4 (Figure 1B) would lead to self-assembly into stable amphiphilic β-sheet fibers in aqueous medium and formation of a hydrogel. Our hypothesis was that the introduction of the rigid phenylglycine (Phg) residue would result in a more
interactions of the aromatic rings projecting from the hydrophobic face. In monophasic medium, the packed form is more abundant forming the amphiphilic single ladder will adsorb on the oil thermodynamically stable packed \( \beta \)-rings alongside the intermolecular hydrogen bonding between backbone amides. The single ladder form is in equilibrium with the more by gray color and the hydrophilic by violet.

Figure 1. Molecular structures of (A) Phe4 (FEFK) and (B) Phg4 (PhgEPghK) peptides. (F, phenylalanine; E, glutamic acid; K, lysine; Phg, phenylglycine). (C) Schematic illustration of Phg4 self-assembly into the amphiphilic \( \beta \)-sheet single ladder form, though \( \pi \)-stacking of the aromatic rings alongside the intermolecular hydrogen bonding between backbone amides. The single ladder form is in equilibrium with the more thermodynamically stable packed \( \beta \)-sheet ladders in aqueous medium, where the latter is formed by lateral growth through hydrophobic interactions of the aromatic rings projecting from the hydrophobic face. In monophasic medium, the packed form is more abundant forming nanofibers, which above critical gelation concentration (CGC) entangle into a nanofibrous network forming hydrogels. In oil–water biphasic media, the amphiphilic single ladder will adsorb on the oil–water interface forming oil-in-water (O/W) emulgel (EMG). The hydrophobic face is denoted by gray color and the hydrophilic by violet.

constrained sequence with the aromatic rings directly attached to the peptide backbone readily positioned for intermolecular \( \pi \)-staking (Figure 1B). This will help stabilize the assembled peptide chains alongside hydrogen bonds formation between the chains backbone amides along the fiber axis, forming a stable amphiphilic \( \beta \)-sheet ladder (Figure 1C).\(^{14,15}\) A single ladder will in turn interact with a second one through lateral \( \pi \)-interactions of the aromatic rings, which become packed between the dimer ladders shielding the hydrophobic faces from the surrounding aqueous medium (Figure 1C). Further lateral growth of the packed fibers can happen through the possible electrostatic interaction between the exposed charged E and K residues side chains, forming thicker fiber bundles (Figure 1C). To rationalize the observed effectiveness of Phg4 self-assembly at the atomic level and confirm the role of the aforementioned noncovalent interactions in stabilizing self-assembly, we have performed in silico modeling on the packed dimer ladder in the form of Prime minimization and side chain predictions, molecular dynamics (MD) simulations, and semiempirical quantum mechanics (SQM) calculations.

We have also investigated the interfacial activity of the unpacked amphiphilic \( \beta \)-sheet ladder, which is in thermodynamic equilibrium with the packed form, for stabilizing oil-in-water (O/W) emulgels (EMGs) in biphasic media (Figure 1C).\(^{16–19}\) EMGs stability was tested versus a number of commercial emulsi bers under a range of environmental and storage conditions. For these investigations, a variety of techniques were used for the molecular, mesoscopic, macroscopic, and mechanical characterizations of both hydrogels and EMGs; mainly including Fourier transform infrared spectroscopy (FTIR), scanning electron microscopy (SEM), transmission electron microscopy (TEM), atomic force microscopy (AFM), small-angle X-ray scattering (SAXS), and oscillatory shear rheology.

We have demonstrated the importance of understanding the mechanisms of peptide self-assembly and stability of the assembled structures at a molecular level, which are of paramount importance for the rational engineering of peptide based nanomaterials.

MATERIALS AND METHODS

Materials. The Phe4 (FEFK) and Phg4 (PhgEPghK) peptides were purchased from Cambridge Reagents Ltd. with \( \geq \)95% purity, as estimated by HPLC (Figure S1). Melissa oil was acquired from Essential Oils Direct Ltd. Sodium dodecyl sulfate (SDS), sodium chloride, potassium phosphate monobasic, and potassium thiocyanate were purchased from Acros Organics with a purity of \( \geq \)99%. Potassium chloride was procured from Alfa Aesar. Polysorbate 80 (Tween 80; 99%), poly(ethylene glycol)-block-poly(propylene glycol)-block-poly(ethylene glycol)/poloxomer (Pluronic F-68; 99%), and alkyltrimethylammonium bromide (cetrimide; \( \geq \)95%) were obtained from Sigma-Aldrich. HPLC grade water and chloroform were supplied by Thermo Fisher.

Computational Details. Initial models of single chain Phg4 and an antiparallel amphiphilic packed dimer \( \beta \)-sheet ladder model composed of 16 Phg4 peptide chains (\( \times \times \)) were built using Maestro and Protein Preparation Wizard,\(^{20}\) with the protonation states assigned on the basis of PROPKA\(^{21}\) at pH 7.0. To design these initial models, the solved structure of an amyloid forming peptide KLVFFA from amyloid beta (PDB code 3OW9) was used as a scaffold for editing to Phg4. Different alignments and conformations of the phenyl side chains were manually explored in terms of the
potential association between the two sheets in the packed dimer, considering favorable π-stacking distances and nonpolar C–C, C–H, and H–H van der Waals distances that are in the attraction region.30–34 Prime v4.5 minimizations and side chain predictions were performed on the starting models.35 For the latter, rotamer libraries were used for random side chain adjustments together with minimizations, until the side chain rotamers were converged. The OPLS force field was used and water solvation effects accounted for using the implicit variable-dielectric generalized Born model (VSBG), which incorporates residue dependent effects.36,37 Single chain Phg4 and the packed dimer model of Phg4 then underwent Desmond v4.7–7,25,27 100 ns MD simulations in the NPT ensemble again with the OPLS force field but water solvation effects now explicitly accounted for using the TIP3P model. Three independent MD simulations were performed for the packed dimer using different seeds for assignment of the initial velocities and the trajectory postprocessed packed-dimer energies, again using Prime. Full details of the simulations and postprocessing are given in the Supporting Information. Following the dynamics, single chain and peptide aggregation geometries from the trajectories (three simulations combined for the packed dimer) were clustered into the recommended number of clusters (20 and 31 for single chain and packed dimer, respectively) based on atomic RMSDs (heavy atoms) and the affinity propagation method.35 1.4

The representative structures from each group were then considered in SQM PM7 optimizations using a restricted Hartree–Fock (RHF) wave function and molecular mechanics correction to CONH bonds (MM05x setting). All optimizations included the COSMO implicit solvation model (conductor-like screening model) with a dielectric constant of 78.4 for water.30,31 The Localized Molecular Orbitals method available in MOPAC in the form of the linear scaling SCF MOZYME algorithm was used to speed up calculations.32,33 Models with the lowest heats of formation were considered for further analyses. The SQM calculations were performed using MOPAC2016.34

Preparation of Hydrogels. Peptide powder was dissolved in HPLC water under vortex at 2500 rpm for 1 min. To trigger self-assembly and gelation the pH was then adjusted to ∼4.5–5 by stepwise addition of 0.5 M NaOH solution, while agitation for 30–60 s in between additions. After pH adjustment, the final volume was corrected using HPLC water to obtain the required concentration. Hydrogels were left in fridge at 4 °C overnight to equilibrate and used the following day.

Preparation of Emulsions and Emulgels. The oil phase (either chloroform or Melissa oil) was added dropwise to the peptide aqueous phase of pH values preadjusted separately to ∼2, 4.5–5, or 8.6 (by stepwise addition of 0.5 M NaOH solution as detailed under preparation of hydrogel), with mixing for 30 s in between additions at 2500 rpm using vortex. Different volumes of the oil phase were added to obtain W/O volume ratios of 2:8, 7:3, 6:4, 5:5, 4:6, 3:7, and 8:2.

Stability Testing of Emulsions and Emulgels. The stability of the emulgels (EMGs) with Melissa oil phase (EMG-M) was compared with the traditional surfactants; SDS, poloxamer (Pluronic F-68), cetrimide, and Tween 80 all used at 50 mM and the emulgels (EMGs) with Melissa oil phase (EMG-M) formulation was carefully removed using a spatula, and the sample was washed using 10 μL of water and finally negatively stained with 10 μL of a solution of 4% (w/w) uranyl acetate. After 4 days, the samples were analyzed after being completely dried.
RESULTS AND DISCUSSIONS

Self-Assembly and Gelation Properties. A phenylglycine-based ionic self-complementary tetrapeptide sequence (Phg4) was designed as a constrained analogue of the previously investigated phenylalanine-based tetrapeptide (Phe4) (Figure 1A,B).\(^\text{8,9}\) Guilbaud et al. showed that the Phe4 sequence neither self-assembled into \(\beta\)-sheet structures nor formed a self-supported hydrogel, as would normally be expected with ionic self-complementary sequences based on Zhang designs.\(^\text{8,9}\) Self-assembly and gelation were only expected with ionic self-complementary sequences based on \(\beta\)-sheet nanostructures.\(^\text{8,9}\)

Here, the secondary structure of both Phe4 and Phg4 peptides was studied at different concentrations using ATR-FTIR at pH 4.5–5 (Figure 2A,B). As predicted from Guilbaud et al. previous findings,\(^\text{8,9}\) the Phe4 did not show the tendency to self-assemble into \(\beta\)-sheet structures; instead, unstructured random conformation at all the investigated concentrations was observed, where all spectra showed a strong broad peak at 1645 and 1640 cm\(^{-1}\) (Figure 2A).\(^\text{35,36}\) Similarly, Phg4 spectra of 5 and 10 mg/mL solution samples showed an unstructured conformation. However, 20, 30, and 40 mg/mL concentrations showed prominent peaks at 1624 and 1650 cm\(^{-1}\) as well as 1524 cm\(^{-1}\) amide II band; corresponding to the formation of an extended \(\beta\)-sheet conformation (Figure 2B). Strikingly, Phg4 peptide showed to form self-supported cloudy hydrogel at a critical gelation concentration (CGC) ~ 20 mg/mL (Figure 2C, top panel), which is consistent with the FTIR spectra (Figure 2B). On the other hand, Phe4 failed to form self-supported materials at the same concentration range (5–40 mg/mL, Figure 2C, bottom panel), as it lacked structure and did not self-organize into \(\beta\)-sheet structures (Figure 2A).

The concentration dependency of Phg4 for the sol–gel transition and mechanical properties at pH 4.5–5 were investigated using shear rheometry. The shear moduli (\(G'\)) significantly increased from ~40 Pa at 10 mg/mL to ~1350 Pa at 20 mg/mL and ~4500 Pa at 30 and 40 mg/mL (Figures 2D and S2A,B), showing the formation of self-supported viscoelastic hydrogels with CGC ~ 20 mg/mL, confirming results from the inverted vial test (Figure 2C, top panel), molecular conformations (Figure 2B), and phase diagram (Figure 2E). Materials of \(G'\) ≥ 1500 Pa or slightly below were thought to be weak gels, which are self-supportive and do not flow by effect of gravity in the inverted vial test, but can be forced to flow if the tube is strongly tapped. Gels of \(G'\) > 1500 Pa are strong gels and would not flow with tapping in the

Small Angle X-ray Scattering (SAXS). SAXS experiments were performed on beamline I22 at the Diamond Light Source (DLS) facility in Didcot, U.K. The energy of the beam was 12.4 keV corresponding to the X-ray wavelength of 0.1 nm. Quartz capillaries (1.5 mm outer diameter, 0.01 mm wall thickness) were supplied from the Capillary Tube Supplies Ltd. A total of 20 mg/mL of Phg4 hydrogel was introduced to capillaries via syringe. Acquisition time was 1 s, and the area pixel array detector used to collect SAXS data transferred vector range of 0.086 (nm\(^{-1}\))\(^{-}\)(4\(\pi\)\(\lambda\)/2), where \(\theta\) is the scattering angle and \(\lambda\) is the wavelength of incident photons. Calibration of the momentum transfer was performed using silver behenate powder. ddH\(_2\)O was used as background and subtracted from all measurements, while the subtraction mask was created using glassy carbon. Data were reduced using the processing tools at DawnDiamond software suite. The 2D scattering photon patterns were integrated and the detector was fixed to 2.211 m, resulting in a momentum transfer vector range of 0.086 (nm\(^{-1}\)) < \(q\) < 7.76 (nm\(^{-1}\)).

Strikingly, Phg4 peptide showed to form self-supported cloudy hydrogel at a critical gelation concentration (CGC) ~ 20 mg/mL (Figure 2C, top panel), which is consistent with the FTIR spectra (Figure 2B). The concentration dependency of Phg4 for the sol–gel transition and mechanical properties at pH 4.5–5 were investigated using shear rheometry. The shear moduli (\(G'\)) significantly increased from ~40 Pa at 10 mg/mL to ~1350 Pa at 20 mg/mL and ~4500 Pa at 30 and 40 mg/mL (Figures 2D and S2A,B), showing the formation of self-supported viscoelastic hydrogels with CGC ~ 20 mg/mL, confirming results from the inverted vial test (Figure 2C, top panel), molecular conformations (Figure 2B), and phase diagram (Figure 2E). Materials of \(G'\) ≥ 1500 Pa or slightly below were thought to be weak gels, which are self-supportive and do not flow by effect of gravity in the inverted vial test, but can be forced to flow if the tube is strongly tapped. Gels of \(G'\) > 1500 Pa are strong gels and would not flow with tapping in the

Figure 2. ATR-FTIR spectra for the two peptides: (A) Phe4 and (B) Phg4 at a series of concentrations. (C) Inverted vial test of Phg4 (top panel) and Phe4 (bottom panel) peptides at pH 5. (D) Shear modulus (\(G'\); y-axis set to log 10) of Phg4 hydrogels at 6 rad s\(^{-1}\) obtained from the frequency sweep experiments performed at 0.2% strain. (E) Phase diagram of Phg4 peptide as a function of pH and concentration. Three distinctive states were observed: strong gel, weak gel, and solution. (F) ATR-FTIR spectra for Phg4 (30 mg/mL) showing phase transition from unstructured sol form at pH 4.5 and back to unstructured sol form at pH 8.6.
inverted vial test (Figure 2E). In contrast, the shear moduli of Phg4 at the same concentrations and pH did not show the formation of stable viscoelastic hydrogel, as observed from the very low shear moduli ($G'$) values compared to Phg4 (data not shown).

Electrostatic interactions play a key role in the self-assembly and gelation properties of β-sheet forming peptides.37 With this in mind, we have investigated self-assembly and sol–gel phase transitions as a function of Phg4 peptide charge. To probe the charge status experimentally for comparison with theoretical calculations, peptide solution (30 mg/mL) was titrated using 0.5 M NaOH. The experimental titration curve indicates that the net peptide charge is neutral within pH range of 4.5–8 when compared to the theoretical charge curve (Figure S2C). Phase diagram of Phg4 hydrogel formation followed its symmetrical charge profile (Figures 2E and S2C) with strong hydrogels being formed at pH in the range ~4.5–8.5, shadowing the charge profile of Phg4 peptides of neutral net charge. This agrees with Caplan and co-workers results, which suggested that a reduction in electrostatic intermolecular repulsion is required to enable stable molecular assembly of ionic self-complementary peptides.37

Remarkably, ATR-FTIR showed the structural switch of Phg4 from unstructured at pH 2.3 to β-sheet structure at pH 4.5 and back again to unstructured at pH 8.6, correlating well with the sol–gel–sol phase transitions and charge status, confirming that the gel formation is a result of the Phg4 self-assembly into β-sheet order (Figures 2E,F and S2C). Furthermore, fluorescence spectroscopy using Thioflavine T (ThT), which has strong binding affinity to β-sheet-like fibrils,38 has corroborated the conformational switch of Phg4 from unstructured at pH 2.3 to β-sheet order, as indicated by the enhancement of ThT fluorescence intensity at pH 4.5, which was insignificant in the case of Phg4, indicating the unique tendency of Phg4 to self-assemble (Figure S2D).

In silico modeling was exploited to help understand at the atomic level the observed propensity of Phg4 toward self-assembly, modeled as antiparallel β-sheet ladders. Prime minimization/side chain predictions led to input models for single chain and packed dimer ladder Phg4 MD simulations. A 2 × 8 chain Phg4 dimer model was considered a good compromise between sheet length and accuracy, allowing us to postprocess MD trajectory structures using theoretically more rigorous SQM PM7 calculations that better account for potential π-stacking interactions in the aggregated assembly. The packed dimer model maintained a stable aggregation structure for the full 100 ns simulation (three independent simulations). The RMSD (system heavy atoms) was seen to stabilize after ~10–20 ns (Figure S3) and the last 80 ns of these simulations used for further analysis. In the models, hydrophobic phenyls from both sheets were intercalated limiting exposure to solvent, and the top-to-bottom aligned charged side chains of hydrophilic K and E residues from consecutive chains formed a stable network of favorable ionic and hydrogen bond interactions throughout. The average number of interchain backbone hydrogen bonds over the three different packed dimer simulations was 2.7, 2.8, and 2.8 (overall average of 2.8). Each chain had stronger interchain hydrogen bonding on one side of the chain (averages of 3.5, 3.5, and 3.5 for the individual simulations; average when combined of 3.5) compared to the other (individual averages 1.7, 2.0 and 2.0; average when combined of 1.9). Generally at least two interchain hydrogen bonds were present throughout. We calculated an association energy for the propensity of single chains to aggregate into the models as $\Delta E_{\text{assoc}}$ (eq S1, Supporting Information) using the last 80 ns of each simulation with values of $-540.3$, $-539.9$, and $-550.9$ kcal mol$^{-1}$ obtained or an average ± standard deviation of $-543.7$ ± 6.2 kcal mol$^{-1}$ and indicative of significant stabilization. A video presentation of the simulation with $\Delta E_{\text{assoc}} = -550.9$ kcal mol$^{-1}$ is included as Supporting Information. Despite the relatively good consistency of results, we highlight that other models of aggregation are possible than these explored here and there are many possibilities (degrees of freedom) in terms of optimal π-stacking for the aggregated dimer. The PM7 optimization of the 31 trajectory cluster representatives
allowed us to more closely analyze the nature of the phenyl interactions in the aggregated dimer, with the predicted most stable model based on the heats of formation shown in Figure 3A. The side view (Figure 3A) demonstrates the formation of a hydrophobic inner core with phenyl rings from both sheets, and the network of hydrogen bonding involving K and E side chains. The interchain backbone hydrogen bonds can also be seen but are more clearly visible in Figure 3B. Stacking and sandwich intercalation of the hydrophobic phenyl side chains from the two sheets is evident when viewed from the top (Figure 3C) and includes T-stacking and parallel-displaced π-stacking interactions (Figure 3B). Antiparallel chain arrangement allows the phenyl ring side chains of nonterminal Phg residues (PhgE, PhgK) from each chain to align forming a hydrophobic central core favorable for aggregation. Predicted stabilization (from PM7/COSMO heat of formations) of the 16-chain Phg4 dimer model structure compared to free single peptide chains was considerable (−524.1 kcal mol⁻¹; relatively similar in magnitude to the ΔE_assoc values from the MD simulations), in further support of experiment and the packed aggregation of Phg4 chains in aqueous media. The stabilization values from PM7 optimizations can in fact be lower for other optimized frames from the MD simulations, with only MD cluster representatives used due to the computational expense involved.

On the mesoscopic scale, the ability of Phg4 peptide to form extended β-sheet nanofibrillar structures was investigated using SEM, AFM, and TEM. SEM showed a dense network of nanofibrillar structures, which is typical for hydrogel formation (Figure 4A). The AFM micrograph clearly showed slightly twisted ribbon-like morphology, which stack in a hierarchical manner to form β-sheet fibrils (Figure 4B). TEM showed relatively straight long entangled nanofibers with Y-shaped branching, as well as lateral associations along the fibers length forming thicker bundles with sizes varied from ~7−11 nm (Figure 4C). Unlike Phg4, Phe4 samples did not show any nanofibrillar structures when examined with the three microscopy techniques (data not shown).

SAXS was further used to investigate the formation and structure of the Phg4 nanofibers (Figure 4D–F). The ln[I_q(q)] versus log q representation indicated no fiber formation for 5 mg/mL, for which scattering similar to water level background was observed (Figure 4D). For 10 and 20 mg/mL samples, q values of ~−1.77 and 1.29 were obtained, respectively (Figure 4D). Typical ~q⁻¹ behavior indicating rods and ~q⁻² indicating discs/flat objects were also plotted. As previously observed from FTIR measurements (Figure 2B), 10 mg/mL did not exhibit a structured β-sheet peak. Likewise, SAXS confirmed that at this concentration Phg4 did not assemble into a rod-like shape but random aggregates (q ~ −1.77), whereas at 20 mg/mL rod-like objects closely resembling fibers were observed with q value 1.29; closer to q⁻¹ behavior. The slight deviation from the q ~ −1 behavior of 20 mg/mL, is most likely coming from the tendency of Phg4 peptide to form flattened fibers and fiber bundles through interfiber lateral growth, as observed from AFM (Figure 4B).

Figure 4. (A) SEM image of 10× diluted and freeze-dried Phg4 hydrogel at 30 mg/mL. (B) AFM micrographs of Phg4 hydrogel at 30 mg/mL diluted to 0.5 mg/mL. (C) TEM micrographs of 10× diluted Phg4 hydrogel at 40 mg/mL, with arrows pointing to fiber bundles of ~7−11 nm size. (D) ln[I_q(q)] vs log q representation of SAXS of 5, 10, and 20 mg/mL of Phg4, q⁻¹ and q⁻² behavior are also presented. (E) Kratky plot (q² I_q(q) vs q representation) of 10 and 20 mg/mL Phg4. The curve shapes are marked by arrows corresponding to folded (assembled) and partially folded states. (F) SAXS scattering pattern of a 20 mg/mL Phg4 at low q plotted in a ln[q²IN(q)] vs q² representation.
and TEM (Figure 4C), respectively. In fact, Kratky plots \( q^2 I(q) \) vs \( q \) for those two concentrations showed a clear transition between partially folded structures at 10 mg/mL to fully folded (assembled) structures at 20 mg/mL (Figure 4E), respectively. It has been previously shown that, for rod objects (i.e., approximately fibers), for \( q R \sigma < 1.3 \) \((R \sigma: \text{cross-section radius of gyration})\), the scattering intensity can be written as \(^4\) \(^2\)

\[
\ln I(q) \propto -\frac{R^2}{2} q^2
\]

If the scattering observed is of the form described by the above eq 1, then at low \( q \), a linear behavior should be obtained in a \( \ln [I(q)] \) versus \( q^2 \) representation, which was the case with Phg4 (Figure 4F). The cross-section radius of gyration, \( R \sigma \), of the fibers can be estimated from the slope of the linear section. If the fibers can be modeled by an infinitely long cylinder, the \( R \sigma \) is related to the diameter of the fiber, \( d \), through the following equation:

\[
R \sigma \approx \sqrt{\frac{d^2}{8}}
\]

By fitting the linear region, the nanofiber diameter size can be estimated as 9.7 ± 0.4 nm, which is in good agreement with the values obtained from TEM (Figure 4C). In contrary, Phe4 did not exhibit any fibril-like scattering patterns (data not shown).

The above-discussed results clearly showed the unique tendency of the ultrashort constrained Phg4 sequence to adopt stable \( \beta \)-sheet conformation in a concentration-dependent and pH-responsive fashion, surpassing its Phe4 counterpart that failed to self-assemble into \( \beta \)-sheet structure. Interestingly, the seemingly minimal change in the chemical structure had large influence on the molecular stability of self-assembly for these tetrapeptide sequences (Figure 1A,B). This is attributed to the aromatic rings orientation in Phg4, which are directly attached to the peptide sequence backbone, making them readily positioned for \( \pi \)-stacking stabilizing the self-assembled structure alongside the hydrogen bonds formed between backbone amides and the salt bridges between K and E residue side chains; all together forming a very stable network of noncovalent interactions from top to bottom along the fiber axis (Figure 3).

**Self-Assembly and Interfacial Properties.** Both Phe4 and Phg4 tetrapeptides are ionic self-complementary sequences with alternating hydrophobic and hydrophilic residue pattern (Figure 1A,B). The amphiphilic-like design suggests that both peptides can exhibit interfacial activity. To test this, the peptide aqueous phase (20 mg/mL) was mixed with chloroform as the immiscible oil phase in 1:1 volume ratio at a range of pH values. Strikingly, only Phg4 managed to form stable emulsion, but not Phe4 at pH 5 (Figure 5A). Also, both peptides showed immediate phase separation after mixing at pH 2 and 8.5 (data not shown). This implies that the surface activity is due to the adsorption of Phg4 amphiphilic nanofibers at the oil–water interface, which are formed at pH 5, rather than the monomeric nonassembled peptide chains at pH 2 or 8.5 (Figure 5A). Phe4 failed to stabilize emulsions at all pH ranges, as it does not self-assemble into amphiphilic \( \beta \)-sheet nanofibers. Fluorescence microscopy of Phg4 emulsion revealed the formation of oil-in-water (O/W) macroemulsion

![Figure 5](https://dx.doi.org/10.1021/acs.biomac.0c00366)
Figure 6. (A) Series of Phg4 Melissa oil mixtures with a range of peptide concentrations (10−60 mg/mL) at 1:1 W:O volume ratio. (B) EMG-M (30 mg/mL) mixed at a range of different W:O volume ratios (2.8−8:2) tested for phase separation over 10 weeks at room temperature. (C) Shear moduli (G') of EMG-M7:3 at a range of peptide concentrations (20−60 mg/mL), performed at 0.2% strain, obtained from the frequency sweep experiments performed at 0.2% strain. (D) Inverted vial test of EMG-M7:3 (20 mg/mL) at pH S. (E) ATR-FTIR spectra of EMG-M7:3 (30 mg/mL) showing phase transition from an unstructured immiscible form at pH 2 to a β-sheet structured EMG form at pH 5. (F) Emulsion stability profiles of EMG-M7:3 vs poloxomer (Pluronic F-68), cetrimide (alkyltrimethylammonium bromide), sodium dodecyl sulfate (SDS), and tween 80 at 50 mM emulsifier concentration after incubation for 3 h at 60 °C (top panel) and after incubation with 100 mM potassium phosphate monobasic for 1 week (bottom panel). (G) Left: SEM image of EMG-M (30 mg/mL), which has been 10X diluted and vacuum-dried, showing two oil droplets stabilized by a peptide fibrous network at the interface. Right: illustration of the O/W EMG-M showing the Melissa oil droplets in yellow entrapped within the peptide nanofibrous network suspended in the continuous aqueous hydrogel phase in blue.

~50−200 μm with the continuous aqueous phase stained with fluorescein isothiocyanate (FITC) green fluorescence (Figure 5A).

AFM of the dried sample revealed the formation of Phg4 microspheres surrounded with nanofibers (Figure 5B) similar to those forming hydrogels (Figure 4B), implying the formation of emulsions (EMGs). This suggests that the amphiphilic nanofibrils formed of unpacked β-sheet single ladders are adsorbed at the surface of the oil phase (chloroform) with the Phg aromatic rings oriented toward chloroform and the hydrophilic E and K residues facing the aqueous phase, by this following the “oriented wedge theory” first suggested by Harkins and Keith in the early 1900s (Figure 1C).44,45 Based on the AFM micrograph, we also hypothesize that chloroform evaporation can lead to the formation of nanofibrilized hollow microspheres entangled within the nanofibrous network of the surrounding aqueous medium. Evidently, SEM micrograph of vacuum-dried emulsion clearly showed the formation of nanofibrilized porous microspheres within the nanofibrous network of the continuous phase, while the air-dried sample showed the formation of solid peptide microspheres (Figure 5C). It is worth mentioning here that artifacts could happen from sample preparation, where it was observed that the nanofibrous network surrounding microspheres was slightly cracked from vacuum drying (Figure 5C, left micrograph). In general, Phg4 microspheres showed similar morphology in SEM to the Fmoc-YL that was previously reported by Ulijn and co-workers to stabilize chloroform-in-water emulsions.16 as well as the surfactant-like decapeptide A_9R that was reported by Hamley and co-workers to stabilize water-in-1-bromohexadecane emulsions.19 The TEM micrograph of diluted emulsion manifested short β-sheet fiber bundles adopting spherical arrangement with bundle diameter size of ~9 nm (Figure 5D), which is within the size range of the hydrogel forming fibers (Figure 4C,F).

Phg4 Peptide Emulgels (EMG). We have then changed the oil phase from chloroform to the pharmaceutically relevant and more viscous oil phase, Melissa oil, which is extracted from the leaves of Melissa officinalis species of the Lamiaceae family. It has been reported that Melissa oil possesses antimicrobial,46 antifungal,47 antiviral,48 and antioxidant49 activities. The Phg4-based O/W emulgels with Melissa as the oil phase (EMG-M) has been formulated by simple mixing of oil droplets in the peptide aqueous phase at self-assembly pH S. First, the effect of Phg4 peptide concentration on the formulation stability was studied at 1:1 W:O ratio. No emulsion formation was observed at 10 mg/mL, but at 20 mg/mL emulsions were obtained with minimal phase separation observed at concentrations ≥30 mg/mL (Figure 6A). This correlates with the FTIR spectra, where Phg4 was unstructured at 10 mg/mL and acquired the β-sheet structure at 20 mg/mL, where self-assembly into the emulsifying amphiphilic β-sheet ladders occurs (Figure 5A). The effect of the W:O ratio on the formulation stability was evaluated at 30 mg/mL peptide concentration. After 10 weeks of storage, we noticed that increasing the water volume above 50% yielded the most stable formulations (Figure 6B). Indeed, FTIR spectroscopy confirmed these visual observations with strong β-sheet peak at ~1624 cm⁻¹ for W:O ratios 6:4, 7:3,
Shear rheology confirmed the formation of viscoelastic hydrogel-like system for W:O 7:3 samples at a range of peptide concentrations (20–60 mg/mL) with shear moduli ~20 kPa (Figures 6C and S5A). The high stiffness of EMG-M, as indicated from $G'$ values, compared to hydrogels of $G'$ hundreds Pa (Figures 2D and S2B) could be attributed to network structure changes associated with the formation of nanofibrillated microspheres; which could have enhanced the mechanical strength of the nanofibrous network of the continuous aqueous phase. The EMG-M stiffness was not significantly affected by the W:O ratio (Figure S5C). Also, EMG-M formulation showed to be not only shear-thinning, but also pseudoplastic thixotropic, where instantaneous, but partial recovery from shear strain was observed from the controlled strain profile of EMG-M$_{7.3}$ at 40 mg/mL Phg4 concentration (Figure S5D). This implicates the injectability of these peptide emulgels (Figure S5D), where fast recovery kinetics is a desirable parameter to avoid uncontrolled release of cargos post injection; which is a useful material property for drug and cell delivery applications. Inverted vial test revealed a CGC for a self-supported emulsion-gel or “emulgel” (EMG) system of ~20 mg/mL for a 7:3 W:O ratio (Figure 6D), similar to that observed for the hydrogel formulations (Figure 2C, top panel).

The stability profile of EMG-M$_{7.3}$ (30 mg/mL) formulation was studied under various physical, chemical, and environmental conditions. The EMG-M stability was indeed affected by pH, where the colloidal system was most stable at the self-assembly pH 5 due to the formation of the emulsifier amphiphilic β-sheet ladders (Figure 6E). Phase separation was observed at pH 2 (Figure 6E) and 8.5 (data not shown), as the peptide is unstructured at these pH values (Figures 2F and 6E). The correlation of structural and phase transitions suggests that the amphiphilic nanofibers are the effective emulsifier and not the nonassembled individual peptide chains. Over long storage (up to 5 weeks), EMG-M showed to be more stable, compared to a variety of commercial surfactants belonging to different classes of surface active agents and commonly used in pharmaceutical and cosmetic products such as the cationic surfactant cetrimide (alkyltrimethylammonium bromide), the anionic sodium dodecyl sulfate (SDS), and the nonionic tween 80 that all showed phase separation when used at the same concentration (50 mM; Figure S6A). However, only the nonionic surfactant poloxomer (Pluronic F-68) showed similar stabilization as EMG-M, as both formed a self-supported hydrogel that helped stabilize suspended oil droplets through the highly viscous continuous phase (Figure S6A). EMG-M was the only formulation that showed thermal stability when heated for 3 h at 60 °C, while phase separation occurred with all other emulsifiers, including poloxomer (Figure 6F, top panel). The salting-out effect was studied as well, using a variety of salts (phosphate, chloride and thiocyanate), and in all cases, only EMG-M and poloxomer emulsions were shown to be stable for one week (Figure 6F, bottom panel, and Figure S6B). Uljin and co-workers have also reported the ability of the β-sheet forming Fmoc-capped short aromatic sequences to stabilize chloroform-in-water emulsions for 24 h better than SDS when incubated with phosphate, chloride, and thiocyanate salts.16

SEM of diluted EMG-M showed peptide nanofibrils adsorbed at the oil–water interface and surrounding the oil droplets (Figure 6G, left panel). This confirms the stabilization of the EMG system through formation of amphiphilic nanofibers that not only adsorb at the oil–water interface, but also form a fibrous network that helps stabilize the system by entrapping the suspended oil droplets within the highly viscous hydrogel continuous phase (Figure 6G, right panel).

Our results, therefore, have demonstrated the considerable potential of using Phg4 amphiphilic β-sheet nanofibers capable of gelation and emulsification. This was achieved by the rational design of a constrained tetrapeptide analogue of the previously reported nonassembling Phg4, exploiting the properties of a more rigid phenylglycine residue. We hypothesized the atomistic interactions leading to stability of self-assembled fibrils through hybrid molecular dynamics simulations and SQM calculations, which highlighted the strong potential for favorable aggregation through π-stacking interactions involving the aromatic Phg side chains from different sheets. These fibrils showed both the ability to form hydrogels in monophasic aqueous medium and interfacial activity in biphasic media forming stable emulgels. Our results highlight the importance of complementary interfacial noncovalent forces alongside backbone amide hydrogen bonding to stabilize the molecular self-assembly of short peptides as exemplified by favorable aromatic stacking in this study. Ultrashort ionic-complementary constrained peptides or UICPs have significant potential for the development of cost-effective, sustainable, and multifunctional soft nanomaterials based on robust nanoassemblies possessing both hydrogelation and emulsification properties: beneficial attributes for the development of a wide range of materials for biomedical, pharmaceutical, cosmetic, and food applications.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.biomac.0c00366.

Molecular dynamics details, HPLC chromatograms of used peptides, additional oscillatory rheology data for hydrogels and emulgels (strain sweep, frequency sweep, and strain recovery), theoretical net charge and titration curve of Phg4 peptide, ThT fluorescence for the two peptides, additional ATR-FTIR for emigel EMG-M (different peptide concentrations and W:O ratios), and emulsion stability profiles for EMG-M compared to commercial emulsifiers (PDF)

Molecular dynamics simulation video for Phg4 packed dimer (MP4)

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https://dx.doi.org/10.1021/acs.biomac.0c00366

Biomacromolecules 2020, 21, 2676–2680
assistance with the AFM experiments, Xinyi Zhu (UoM) for training R.P. on TEM sample preparation, and Diamond Light Source for the Beam Time Award (SM17102) to M.A.E. and J.K.W. and the staff on Beamline I22 for their support with the SAXS experiments, in particular, Andrew J. Smith. Authors would also like to thank Nadia Moinuddin (UCLan) for acquiring preliminary data on chloroform-in-water emulsions funded by UCLan Undergraduate Research Internship Scheme (URIS).

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