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Supramolecular Self-Assembly to Control Structural and Biological Properties of Multicomponent Hydrogels

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Abstract

Self-assembled nanofibers are ubiquitous in nature and serve as inspiration for the design of supramolecular hydrogels. A multicomponent approach offers the possibility of enhancing tunability and functionality of this class of materials. We report on the synergistic multicomponent self-assembly involving a peptide amphiphile (PA) and a 1,3:2,4-dibenzylidene-D-sorbitol (DBS) gelator to generate hydrogels with tunable nanoscale morphology, improved stiffness, enhanced self-healing, and stability to enzymatic degradation. Using induced circular dichroism of Thioflavin T (ThT), electron microscopy, small-angle neutron scattering (SANS), and molecular dynamics approaches we confirm that the PA undergoes self-sorting while the DBS-gelator acts as an additive modifier for the PA nanofibers. The supramolecular interactions between the PA and DBS gelators result in improved bulk properties and cytocompatibility of the two-component hydrogels as compared to the single component systems. The tunable mechanical properties, self-healing ability, resistance to proteolysis, and biocompatibility of the hydrogels suggest future opportunities for the hydrogels as scaffolds for tissue engineering and drug delivery vehicles.

Introduction

Multicomponent supramolecular self-assembly is ubiquitous in natural processes leading to the formation of highly ordered and complex architectures.¹ The functionality of many biomacromolecules such as cytoskeleton actin and actin-binding proteins found in living organisms depend on the co-assembly of multiple building blocks into one-dimensional nanostructures.² These supramolecular assemblies result from cooperative and synergistic non-covalent interactions, which provide adaptive, dynamic, and responsive properties.

Multicomponent self-assembly is currently the focus of fundamental research aimed at dissecting the code embedded within natural supramolecular architectures³⁻⁵ to engineer innovative materials based on predictive molecular interactions.⁶⁻⁸ Such design strategies have been developed using well-defined molecular building blocks including β -sheet fibrillizing peptides, peptide amphiphiles (PAs), engineered polypeptides, and low-molecular-weight species.⁸⁻¹⁰ However, many of these artificial nanostructures are homotypic, consisting of one class of building block and thereby limiting the level of structural and functional complexity, diversity, and tunability. In contrast, multicomponent self-assembly offers the possibility to generate a wider range of more complex structures, enhance modularity, and provide spatio-temporal control of self-assembly.^{2,11} This approach has been used to harness synergistic properties as a result of using two different interacting molecular building blocks such as peptide-peptide,¹²⁻¹⁴ protein-peptide,¹⁵⁻¹⁷ PA-polysaccharide,¹⁸ protein-protein,^{19,20} and protein/peptide-DNA.²¹ The structures and properties emerging from these systems are opening new opportunities for the rational design of more complex and functional materials.⁶

Multicomponent self-assembly is a versatile strategy to design supramolecular hydrogels with molecular complexity, enhanced interfacial areas between nanofibers, tunable mechanical properties, *in situ* creation of new materials, and diverse morphology.²² These properties make such hydrogels suitable for a wide range of applications in drug delivery,²³ tissue engineering,²⁴ nanoreactor design, and optoelectronic materials.²⁵ Specifically, multicomponent gels can have advantages compared to individual component systems. For example, multicomponent gels can exhibit optoelectronic properties that cannot be accessed using the single building blocks.²⁶ There are also examples where cellular behaviours can be improved using multicomponent hydrogels.^{27,28} It is also possible to prepare systems with unusual behaviour such as delayed gelation³, or two-stage rheological control.^{29,30} Moreover, multicomponent self-assembly breaks the molecular homogeneity imposed by traditional single-building block hydrogels.

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5 PAs represent a fascinating class of self-assembling building blocks for designing bioactive
6 hydrogels.³¹⁻³⁴ These amphiphilic molecules consist of at least one lipid chain attached to a
7 peptide backbone and are able to self-assemble into high-aspect ratio cylindrical nanofibers
8 defined by the hydrophobic collapse of the alkyl region and electrostatic interactions between
9 the charged head groups.³⁵ PAs have been used in supramolecular co-assembling systems with
10 oppositely charged and structurally distinct molecules to develop sac-like structures,¹⁸ dynamic
11 tubular materials,¹⁶ and hydrogels.¹⁷ However, combination of PAs and other gelators with
12 distinct molecular structures exhibiting similar charge has not been explored. Therefore,
13 supramolecular self-assembly of a well-known PA molecule with another molecule with a
14 well-characterized self-assembly mechanism is of considerable interest in designing new breed
15 of multicomponent hydrogels.
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26 1,3(*R*):2,4(*S*)-Dibenzylidene-D-sorbitol (DBS) is a well-known sugar-based low-molecular-
27 weight gelator (LMWG) with a high capacity to self-assemble into fibrillar networks in organic
28 solvents and in water.³⁶ The effectiveness of DBS as a gelator is due to intermolecular hydrogen
29 bond interactions between the sugar units combined with π - π stacking and solvophobic
30 interactions between the aromatic rings, which characterize its “butterfly-like” wing
31 structure.³⁷ In the past century, DBS has been widely used in industrial applications as a
32 cosmetic additive, in dental composites, and as a polymer nucleation/clarification agent.³⁷
33 Recently, Smith and colleagues reported the development of DBS derivatives including DBS-
34 COOH³⁸ and DBS-CONHNH₂³⁹ that are capable of self-assembly in water into self-supported
35 hydrogels by pH-switching and heat-cool transitions, respectively. Using molecular dynamics
36 simulations, Knani and Alperstein determined that the carboxylic acid and acylhydrazone
37 groups on these molecules play an important role in modifying both their solubility and
38 capacity to self-assemble into fibrillar networks in aqueous systems.⁴⁰ Hydrogels based on
39 DBS-COOH and DBS-CONHNH₂ have potential applications in environmental clean-up,^{39,41}
40 drug encapsulation and delivery,⁴² electro-catalysis,⁴³ cell culture,⁴⁴ and as nanoreactors.⁴⁵
41 Furthermore, these gelators have been hybridized with polymers to produce robust^{38,46} and
42 multidomain^{30,47} hydrogels, and more recently DBS-COOH/DBS-CONHNH₂ co-assembled
43 hydrogels⁴⁸ but their combination with LMWGs of other types to design multicomponent
44 hydrogels has not been investigated.
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3 The combination of LMWGs from different classes remains relatively rare and offers
4 considerable potential for the emergence of new functions, which can be exploited to develop
5 materials with innovative properties. In this study, we report on the supramolecular design of
6 multicomponent hydrogels based on a well-known PA molecule and DBS-COOH using a slow
7 acidification protocol to fabricate hydrogels with emergent properties. The resulting hydrogels
8 exhibit tunable stiffness, self-healing capacity, enhanced stability to enzymatic degradation,
9 and tunable bioactivity. To the best of our knowledge, this is the first example of these two
10 LMWGs being combined.
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20 **Materials and Methods**

21 **Preparation of hydrogels**

22 Both **PA-E3**³² and **DBS-COOH**³⁸ were synthesized as previously reported. For single-
23 component hydrogels, **PA-E3** and **DBS-COOH** solutions (10 mM) were individually prepared
24 in phosphate buffer saline (pH 7.4, 200 μ L) and aliquots of NaOH (aq) (0.5 M, 10 μ L) were
25 added to dissolve the gelators (pH = 9). The solutions were then transferred to vials containing
26 glucono- δ -lactone (GdL) (4 mg), followed by thorough shaking to dissolve the GdL granules.
27 The vials were left overnight for gelation to occur. The multicomponent hydrogels were
28 prepared by mixing aliquot solutions of **PA-E3** and **DBS-COOH** at various molar ratios
29 expressed as percentages (100/0, 80/20, 50/50, 20/80 and 0/100) such that the final total
30 concentration of the mixtures was 10 mM in each case. The solutions were then transferred to
31 vials containing GdL, shaken and left overnight for gelation to occur.
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43 **Thioflavin T (ThT) fluorescence assay**

44 Aliquots of solutions (0.4 mM, 20 μ L) of **PA-E3**, **DBS-COOH** and **PA-E3/DBS-COOH**
45 mixtures were added to GdL (0.2 mg). Aliquots of ThT (0.4 mM, 20 μ L) were then added to
46 the gelator solutions. Each sample was mixed by pipetting up and down three times, loaded
47 into a 20 μ L sample flow cell, sealed with paraffin wax and immediately transferred onto an
48 inverted Laser Scanning Confocal Microscope (LSCM) (Leica Laser Scanning Confocal TCS
49 SP2) with 63 oil immersion objective, using excitation and emission wavelengths of 458 nm
50 and 468 nm, respectively. Images of the self-assembled nanofibers were acquired after
51 incubating the sols for 6 h. The flow cells were kept humid in petri dishes to prevent
52 evaporation. The hydrogels prepared using an identical protocol but without the ThT were used
53 as control and did not exhibit any fluorescent signal.
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Circular dichroism measurement for ThT interactions

Circular dichroism (CD) was measured with a ChirascanTM circular dichroism spectrometer (Applied Photophysics Limited, UK) using a quartz cell with 1 mm path length and the following parameters: data pitch – 0.5 nm, scanning mode – continuous, scanning speed – 100 nm/min, bandwidth – 2 nm and accumulation – 5. All CD data are presented as ellipticity and recorded in millidegree (mdeg). Samples for CD measurements were prepared by adding GdL (0.4 mg/mL) to **PA-E3**, **DBS-COOH** and equimolar **PA-E3/DBS-COOH** solutions (200 μ M). Then an aliquot of ThT (200 μ M) was added to the gelator solutions and the mixture was loaded into a quartz cell. CD spectra were obtained at 2 min interval from 190 to 500 nm at speed of 50 nm/min.

Transmission electron microscopy (TEM) characterization

Solutions (1 mM, 40 μ L) of **PA-E3**, **DBS-COOH** and **PA-E3/DBS-COOH** mixtures were added to GdL (0.2 mg), shaken thoroughly and incubated for 10 h. Samples were then mounted on holey carbon-coated copper grids that were pre-plasma treated (Agar Scientific, Stansted, UK). The grids were immersed in ultrapure water for 30 s to remove excess and unadsorbed samples. The grids were then immersed in a solution of uranylacetate (2%) for 30 s and excess uranylacetate solution was removed using filter paper. Grids were allowed to dry in a desiccator for 24 h at room temperature. Images were acquired on a JEOL 1230 transmission electron microscope fitted with Morada CCD camera and operated at an acceleration voltage of 80 kV.

Atomic force microscopy (AFM) characterization

Solutions of **PA-E3** (1 mM, 40 μ L, pH 11), **DBS-COOH** (1 mM, 40 μ L, pH 11) and **PA-E3/DBS-COOH** mixtures were mixed with GdL (0.4 mg) and the mixtures were dropped onto freshly cleaved mica surface so that they self-assemble on the substrate. The samples were air-dried at room temperature for 24 h. AFM characterization of nanofibers was performed on an Asylum Research MFP-3D atomic force microscope (Santa Barbara, CA, USA) without treatment. Ultrasharp silicon nitride tips (NSC15 noncontact silicon cantilevers, MikroMasch, Spain) were used. Typical scan settings involved the use of an applied piezo deflection voltage of 0.6 – 0.7 V at a scan rate of 0.7 Hz. All images were processed using Igor Pro software.

Small-angle neutron scattering (SANS) characterization

The required concentration (10 mM) of **PA-E3**, **DBS-COOH** and various mixtures of **PA-E3** and **DBS-COOH** were prepared in D₂O (400 μL). The suspensions were dissolved by adding aliquots of NaOD (0.5 M, 10 μL) and the solutions were mixed with GdL (5 mg). After a thorough shaking until GdL granules fully dissolved, the mixtures were transferred into 1 mm path length UV spectrophotometer grade quartz cuvettes (Hellma). Gelation of the solutions was allowed to proceed overnight. Small-angle neutron scattering (SANS) measurements were performed on the fixed-geometry, time-of-flight LOQ diffractometer (ISIS Neutron and Muon Source, Oxfordshire, UK). A white beam of radiation with neutron wavelengths spanning 2.2 to 10 Å was enabled access to Q [$Q = 4\pi\sin(\theta/2)/\lambda$] range of 0.004 to 0.4 Å⁻¹ with a fixed-sample detector distance of 4.1 m. The cuvettes were mounted in aluminium holders. The time taken for each measurement was approximately 30 min. All scattering data were normalized for the sample transmission, the background was corrected using a quartz cell filled with D₂O and the linearity and efficiency of the detector response was corrected using the instrument-specific software. Data were fitted using the appropriate models between 0.00485 <math>Q > 0.49132 Å⁻¹ for consistency. The scattering length density (SLD) for D₂O was set to 6.3 x 10⁻⁶ Å⁻²; SLD for scattering structures set to 2.197 x 10⁻⁶ Å⁻² for **DBS-COOH** only and 1.31 x 10⁻⁶ Å⁻² for **PA-E3** only. SLDs for mixtures of **PA-E3** and **DBS-COOH** were based on compositional average – these values estimated using the NIST calculator with a (assumed) density of 1.3. Data for the **PA-E3** hydrogel fit flexible elliptical cylinder model while a power law was used to fit the data for **DBS-COOH** hydrogels which suggests that the structures are large and outside the effective range of the SANS. For the **PA-E3/DBS-COOH** admixtures, combination of the flexible elliptical cylinder model with a power law was used to fully fit the data. Details of the parameters for all the fittings are presented in **Table S1**.

Rheological kinetics

Rheological measurements were performed using a Discovery Hybrid Rheometer, Rheo-DHR3 (TA Instruments). All data were collected at 25 °C. For each sample, immediately following the addition of GdL (4 mg/mL) to solutions of the gelators (10 mM, pH 9), 100 μL of the mixture was added to the center of the bottom plate and the upper geometry (parallel top plate with 20 mm diameter) was lowered to a gap of 50 μm. A time-sweep was performed where the storage and loss moduli were monitored for 3 hours at a constant frequency of 1 Hz and 0.5% strain at 25 °C. Light mineral oil was applied to the perimeter of the parallel plate to prevent evaporation of water over the course of the experiment.

Characterization of nanofiber growth by TEM

Solutions (0.5 mM, 100 μ L) of PA-E3, DBS-COOH and PA-E3/DBS-COOH mixtures were added to GdL (0.2 mg/mL) and shaken thoroughly. An aliquot (50 μ L) of each sample was added onto a flat sheet of parafilm. Holey carbon-coated copper grids (Agar Scientific, Stansted, UK) that were pre-plasma treated were carefully placed on top of each drop. The samples were kept in a temperature and humidity chamber to prevent evaporation. The grids were taken off at various time intervals, excess solution was removed using filter paper and the grids were then immersed in a solution of uranyl acetate (2%) for 30 s and excess uranyl acetate solution removed using filter paper. Grids were allowed to dry in a desiccator for 24 h at room temperature. Images were acquired on a JEOL 1230 transmission electron microscope fitted with Morada CCD camera and operated at an acceleration voltage of 80 kV.

Molecular dynamic simulations

The simulation was conducted using Material Studio 8.0 software (by Biovia). The dynamic atomistic simulation was performed according to the following steps:

Step 1: Building cubic cells

Five simulation cubic boxes (about 20 Å edge) were constructed using Amorphous cell module for each of the following: DBS-COOH, PA-E3 and their mixtures at various ratios: 1:5, 1:1 and 5:1 (DBS-COOH: PA-E3).

Step 2: Molecular Dynamics simulation

Dynamic simulation was performed at 300 K. The cells were subjected to 100,000 dynamic steps of 1 fs each at constant moles number, pressure and temperature (NPT ensemble) to determine their density. This stage was followed by a constant moles number, volume and temperature (NVT ensemble) refinement stage of 100,000 dynamic steps and a data collection stage of additional 400,000 NVT steps. All MD simulations were conducted using Forcite module with COMPASSII force field. The electrostatic term was considered using Ewald and the van der Waals term using atom-based summation methods with an accuracy of 10^{-3} kcal/mol. The repulsive cut-off for Van der Waals term was chosen as 12.5 Å. For NPT molecular dynamic simulations, Nose thermostat and Berendsen barostat were chosen.

Step 3: Analysis

The resulting dynamic trajectories were analysed using Forcite module analysis tools. The following properties were calculated:

- (i) ***Cohesive Energy Density (CED) and Solubility parameter***

Cohesive energy is the energy required to break the interactions between molecules. Generally, it is measured as the heat of vaporization of a liquid. The cohesive energy density (CED) corresponds to the cohesive energy per unit volume. The solubility parameter is the square root of the CED, and is a measure of the ability of materials to dissolve each other.

(ii) Enthalpy of mixing

CED values can be used to calculate the enthalpy of mixing (per unit volume) using the following equation:

$$\Delta H_{\text{mix}} = \Phi_a E_{\text{coha}} + \Phi_b E_{\text{cohb}} - E_{\text{cohab}} \quad (5)$$

Where: E_{coh} is CED of constituent **a**, **b** or the blend (**ab**); Φ_a and Φ_b are the volume fractions of the two components in the blended system.

The enthalpy of mixing is the released or taken-up heat upon mixing of two substances. Generally, a negative Gibbs free energy of mixing is a necessary condition for substances to form a miscible phase. The entropy change during mixing may be negligible when polymers are involved and miscibility can be determined from the enthalpy change of mixing instead of Gibbs free energy. An exothermic enthalpy of mixing is indicative of a miscible blend.

(iii) Radial distribution function (RDF)

Radial distribution function (also referred to as Pair correlation function) gives a measure of the probability that, given the presence of an atom at the origin of an arbitrary reference frame, there will be an atom with its center located in a spherical shell of infinitesimal thickness at a distance r from the reference atom. RDF may serve as a tool to estimate intermolecular interactions like hydrogen bonding.

Amplitude sweeps, critical strain, and self-recovery of hydrogels

A parallel plate geometry (parallel top plate with 8 mm diameter) was used with 150 μm gap distance to perform frequency and amplitude sweeps. **PA-E3** (10 mM), **DBS-COOH** (10 mM) and various stoichiometric mixtures of **PA-E3/DBS-COOH** hydrogels were prepared by adding GdL (10 mg/mL) to the gelator solutions. The hydrogels were left to form overnight before the measurements. Frequency scans were performed from 1 to 10 Hz under a strain of 0.5%. At this strain, the frequency sweeps were performed within the linear viscoelastic region, where the storage modulus (G') and loss modulus (G'') are independent of the strain amplitude. Critical strain values were determined from the point where G' values start to decrease with strain. The recovery properties of the hydrogels in response to applied shear forces were investigated using the following procedure; 0.1% (100 s), 100% (200 s), 0.1% (200 s), 100%

(200 s), (400 s) with the applied shear force, expressed in terms of strain (%) and duration (s) in parentheses. For the strain recovery, 10 mM of **PA-E3** was mixed with 10 mM of **DBS-COOH** to prepare **PA-E3/DBS-COOH** hydrogels.

Proteolytic stability of the hydrogels

Hydrogels of **PA-E3** (10 mM), **DBS-COOH** (10 mM), **PA-E3** (5 mM)/**DBS-COOH** (5 mM) were prepared as previously described above. The hydrogels were immersed in PBS for 1 hour, then incubated at 37 °C for 5 days in proteinase K solutions (5 mg/mL in PBS). Hydrogels were then prepared for scanning electron microscopy (SEM) by freeze-drying. SEM micrographs of the xerogels were acquired on Inspect F50 (FEI Comp, the Netherlands) after sputter-coating with gold (10 nm thick). Also, mass spectra of the mixtures of 0.1 mM **PA-E3**/proteinase K (5 mg/mL), **PA-E3/DBS-COOH**/proteinase K (5 mg/mL) were acquired on Liquid Chromatography-Mass Spectrometer, comprising a 1100 Series LC and SL Ion Trap MSD (Agilent Technologies, UK). Weight loss measurements were performed by incubating each hydrogels in 500 mL of proteinase K solution (5 mg/mL) or phosphate buffer saline (PBS 1x) at 37 °C at various time intervals. The solutions with or without the enzyme were replaced with the freshly prepared solutions after each weight measurement. Then, the weights of hydrogels were recorded to determine the weight loss compared to the initial weights of hydrogel. We computed the weight loss using equation $(W_f - W_i)/W_i \times 100$, where W_i and W_f represent initial and final hydrogel weights, respectively.

Cellular behaviours on PA-E3, DBS-COOH, and PA-E3/DBS-COOH hydrogels

PA-E3, **DBS-COOH**, and **PA-E3/DBS-COOH** hydrogels were prepared as previously discussed above. The hydrogels were cross-linked with CaCl_2 (50 mM) and washed with PBS until the hydrogels became neutral to pH strips (pH 7). Human adipose derived stem cells (hADSC, 5000 cells/mL) were cultured on the hydrogels and incubated for 4 days at 37 °C and 5% CO_2 . Using a Live/Dead assay kit (Life Technologies, UK), the assay was performed by incubating the hydrogels with calcein AM (10 mM) and ethidium homodimer-1 (1 mM) for 30 min. Imaging was performed on an inverted confocal laser scanning microscope (CLSM, Leica, Germany).

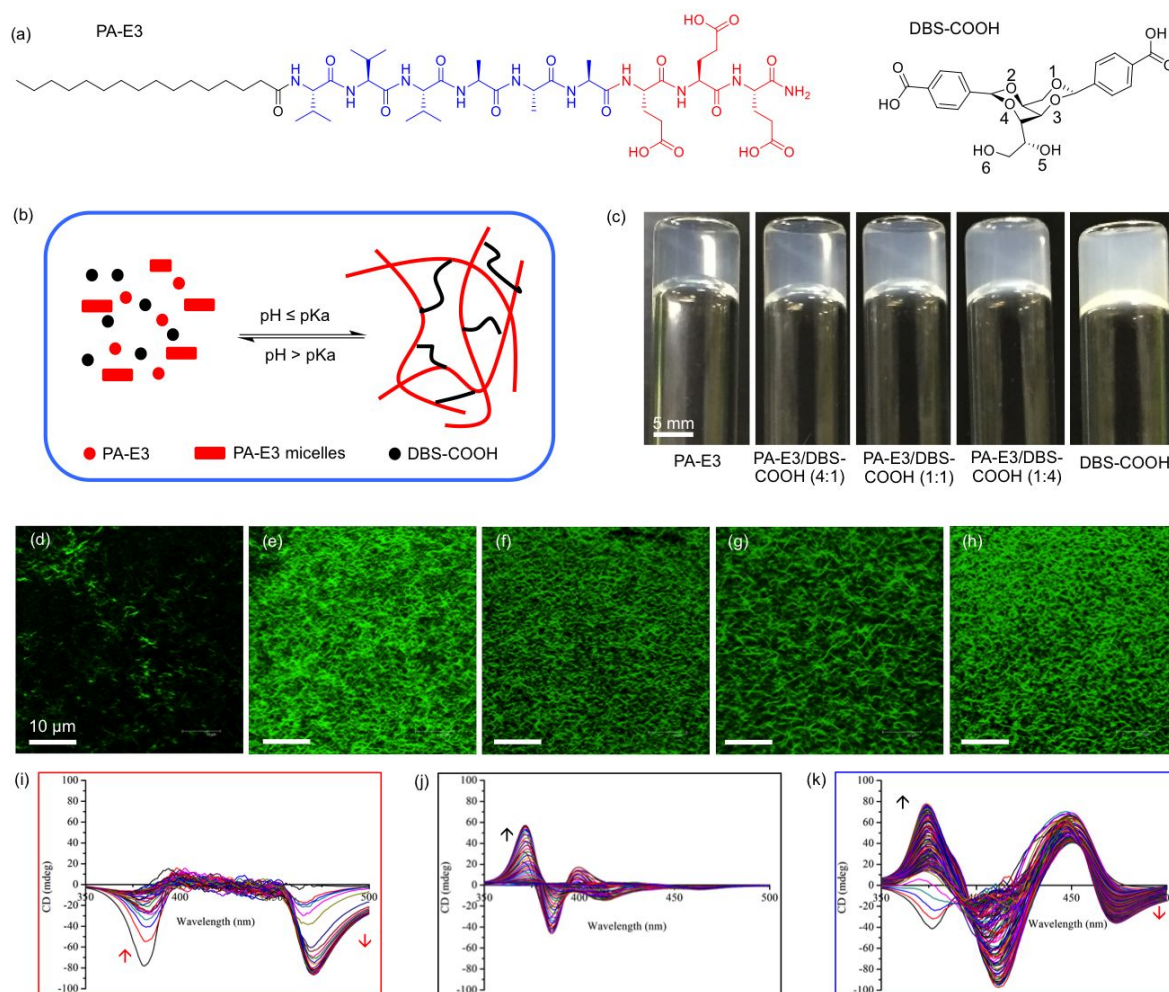


Figure 1. Molecular information and sol-gel phase transition. (a) Molecular structure of **PA-E3** and **DBS-COOH**. (b) Schematic representation of supramolecular self-sorting of **PA-E3** and **DBS-COOH** by pH-switching. (c) Photographs of hydrogels of **PA-E3**, **DBS-COOH** and **PA-E3/DBS-COOH** mixtures. Laser scanning confocal micrographs of nanofibers of (d) **PA-E3**, (e) **PA-E3/DBS-COOH** (4:1), (f) **PA-E3/DBS-COOH** (1:1), (g) **PA-E3/DBS-COOH** (1:4), and (h) **DBS-COOH**. Induced circular dichroism spectra of ThT during the self-assembly of (i) **PA-E3**, (j) **DBS-COOH** and (k) **PA-E3/DBS-COOH** (1:1).

Results and Discussion

Self-assembly and formation of hydrogels

The PA used in this study is the negatively charged peptide $\text{CH}_3-(\text{CH}_2)_{14}-\text{CONH-VVVAEEEE}-\text{CONH}_2$ (**PA-E3**), palmitoylated at the N-terminus (Fig. 1a). This PA is known to assemble into micrometre-long nanofibers by calcium ion coordination.^{31,49} Conversely, **DBS-COOH** (Fig. 1a) has been shown to assemble into nanofibers that are microns in length as a result of pH switching.³⁸ Aqueous solutions (10 mM) of gelator **DBS-COOH** and gelator **PA-E3** were prepared and the aliquots of each were mixed to provide a series of solutions with various stoichiometric ratios of **PA-E3** to **DBS-COOH** (100/0, 80/20, 50/50, 20/80, and 0/100 %) such that a constant total concentration (10 mM) was maintained in all mixtures. Stable

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3 hydrogels were formed by slow acidification of the basic solutions by adding GdL (10 mg/mL).
4 GdL undergoes a base-catalysed hydrolysis to gluconic acid as previously described. The slow
5 hydrolysis allows significantly more controlled and homogeneous gels to be formed as
6 compared to the use of mineral acids.⁵⁰ Consequently, the gradual decrease in pH below the
7 pK_a of the terminal carboxylic acid ($pK_a \sim 4 - 5$) of **DBS-COOH** and **PA-E3**^{30,51} allows the
8 slow protonation of the carboxylate groups (Fig. 1b). This protonation directs the gradual self-
9 assembly of both **PA-E3** and **DBS-COOH** over a period of 4 h, leading to kinetically
10 controlled formation of homogenous self-supported hydrogels (Fig. 1c).
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19 *Characterization of assembly*

20 Hydrogels were formed for all tested materials including **PA-E3**, **DBS-COOH**, and their
21 combinations. The hydrogels of **PA-E3** alone were optically transparent while those of **DBS-**
22 **COOH** were translucent. Moreover, there was an increasing degree of translucency with
23 increasing concentration of **DBS-COOH** in the two-component hydrogels, suggesting an
24 increasing **DBS-COOH** nanofiber density in the materials. At the nanoscale, fluorescent
25 microscopy using ThT was used to further confirm self-assembly of **PA-E3**, **DBS-COOH**, and
26 their mixtures. As revealed by LSCM, nanofibers of individually assembled **PA-E3** (200 μ M)
27 and **DBS-COOH** (200 μ M) containing an equimolar concentration of ThT (200 μ M)
28 fluoresced when excited at 488 nm (Fig. 1d and 1h). Similarly, the nanofibers of **PA-E3/DBS-**
29 **COOH** mixtures (200 μ M) also exhibited strong fluorescence (Fig. 1e-g) when excited at the
30 same wavelength, which results from the integration of ThT within the hydrogel
31 nanostructures. However, the fluorescence is more intense on **DBS-COOH** (Fig. 1h) and **PA-**
32 **E3/DBS-COOH** (Fig. 1e,f,g) nanofibers than on the **PA-E3** nanofibers (Fig 1d). We observed
33 no fluorescence from the DBS-gelator solution itself (Supplementary Figure 2). It is well
34 established that ThT is a molecular rotor and its fluorescence is viscosity dependent.^{52,53} No
35 fluorescence was observed pre-gelation, as the ThT is not incorporated into viscous
36 environments. However, upon gelation, hydrophobic rigid fibers are formed and, incorporation
37 of ThT into these, results in an increase in fluorescence.⁵² Therefore, our systems support the
38 idea that the reason for the characteristic increase in the ThT fluorescence intensity following
39 its incorporation into the **DBS-COOH** and the two-component (**PA-E3/DBS-COOH**)
40 nanofibers than the **PA-E3** alone nanofibers is due to the rigidity of the nanofibers, which
41 prevents the rotation of the benzothiazole ring relative to the aminobenzene ring in the excited
42 state.⁵³ Previous studies have demonstrated that molecular gelator nanofibers are able to
43 sequester cationic dyes such as methylene blue (MB) from aqueous system.⁴¹ We have also
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3 demonstrated this with **DBS-COOH** using MB (data not shown). Therefore, we reasoned that
4 **DBS-COOH** nanofibers sequester the cationic ThT dye in a similar fashion, which can also
5 explain the observed increased fluorescence intensity.
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10 We also confirmed by circular dichroism (CD) spectroscopy that the achiral ThT molecule
11 acquired a chiral signature following its interaction with the self-assembled **PA-E3**, **DBS-**
12 **COOH**, and **PA-E3/DBS-COOH** nanofibers in a differential and time-dependent manner
13 (Supplementary Figure S3). Such chirality transfer is believed to result from a “sergeants-and-
14 soldiers” type of interaction⁵⁴ between ThT and the self-assembled nanofibers. Upon
15 incorporation into the **PA-E3** nanofibers during self-assembly, ThT exhibited a negative band
16 at 385 nm, which disappeared within 30 min of self-assembly while another negative band
17 emerged at 470 nm as the self-assembly progressed (Fig. 1i, Supplementary Figure 3a). On the
18 other hand, ThT exhibited a bisignate Compton effect with positive and negative bands at 370
19 and 385 nm, respectively (Fig. 1j, Supplementary Figure 3b). Interestingly, the CD spectrum
20 of ThT upon incorporation into the **PA-E3/DBS-COOH** nanofibers combines the distinct
21 chiral signatures of both **PA-E3** and **DBS-COOH** as well as the emergence of a new band at
22 450 nm (Fig. 1k, Supplementary Figure 3c). This result suggests that ThT is able to
23 discriminate between different chiral environments in the multicomponent hydrogels. Also,
24 with this approach, we could infer that the two building blocks constituting our
25 multicomponent hydrogels exhibited self-sorting self-assembly.
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39 *Characterization of nanostructures*

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41 Transmission electron microscopy (TEM) revealed that the **PA-E3** solution (0.1 mM, pH 10)
42 on its own has pre-formed micellar structures that are ~200 nm long. Such pre-gelation
43 assembly of PAs into cylindrical micellar nanostructures has been previously reported to take
44 place as a result of the hydrophobic collapse of the alkyl moieties in an aqueous environment.⁵⁵⁻
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48 ⁵⁷ On the other hand, while such structures were not observed in the solution of pure **DBS-**
49 **COOH** (Supplementary Figure S4), they were observed in the equimolar mixture of **PA-E3**
50 and **DBS-COOH**. Upon gelation triggered by GdL hydrolysis, the **PA-E3** solution further
51 assembled into typical PA nanofibers that are microns in length and ~7 nm in diameter (Fig
52 2ai, top panel), while the nanofibers of **DBS-COOH** (0.1 mM) on its own self-assembled into
53 bundles of ribbons that are several microns in length and ~80 nm in width as previously
54 reported⁴⁶ (Fig 2av). Atomic force microscopy (AFM) was used to confirm these structures
55 (Fig. 2bi, middle panel) (Fig. 2bv, middle panel). Interestingly, nanofibers of the **PA-E3/DBS-**
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COOH hydrogels (0.1 mM) appeared to have different geometries compared to those of the single-component hydrogel systems and be more entangled (Fig 2a ii-iv top panel). Furthermore, the density of this nanofiber network seemed to increase with increasing DBS-COOH concentration. Such entanglement is possibly due to the formation of interconnected self-sorting networks.⁵⁸ The nanofiber bundling phenomenon is more pronounced with the PA-E3/DBS-COOH hydrogels than in the hydrogels of DBS-COOH alone (Fig. 2b ii-iv, middle panel). It is important to mention that the difference in appearance of the nanofiber images acquired with the two techniques (TEM and AFM) is probably due to different sample preparation procedures. In the case of AFM, the solutions were allowed to self-assemble on the substrates, while with TEM, images were acquired on pre-formed partial hydrogels.

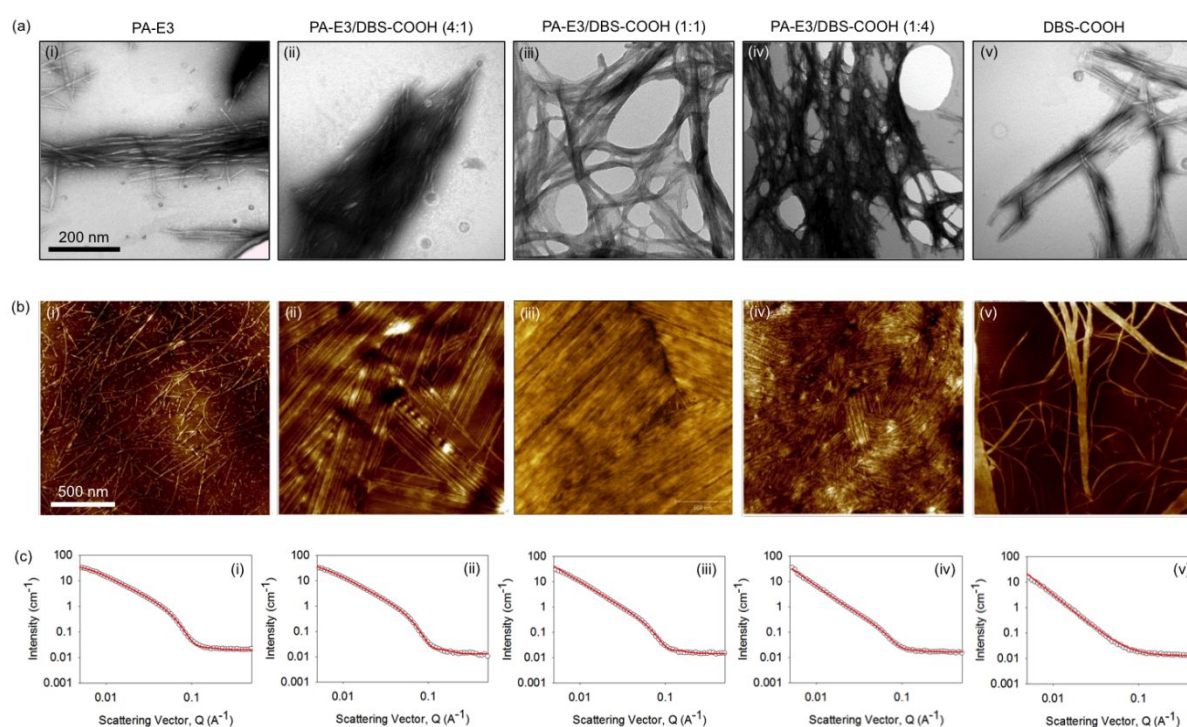


Figure 2. Nanostructure characterization. Nanostructures of nanofibers of (i) PA-E3, (ii) PA-E3/DBS-COOH (4:1), (iii) PA-E3/DBS-COOH (1:1), (iv) PA-E3/DBS-COOH (1:4) and (v) DBS-COOH at 0.1 mM concentration gelator concentration revealed by (a) TEM and (b) AFM. (c) SANS data for gels of (i) PA-E3, (ii) PA-E3/DBS-COOH (4:1), (iii) PA-E3/DBS-COOH (1:1), (iv) PA-E3/DBS-COOH (1:4) and (v) DBS-COOH at 1 mM gelator concentration. In all cases, the fits to the data are shown as the red line through the open symbols.

To further characterize the internal structuring of the hydrogels without drying and staining effects that accompany imaging techniques, synchrotron small-angle neutron scattering

(SANS) measurements were performed on the hydrogels. SANS data for the hydrogels of **PA-E3** (10 mM) fitted best to a flexible elliptical cylinder model after the initial fixing of the background and setting the length to 2000 Å (an arbitrarily value greater than the resolution of the technique) (Fig. 2k, lower panel). Other parameters to fit the data for **PA-E3** include a radius of 26.9 ± 0.1 Å, an axis ratio of 1.64 ± 0.02 , and a Kuhn length of 134.0 ± 2.0 Å. On the other hand, data for the hydrogels of **DBS-COOH** fitted to a power law only with a power-law exponent of 2.72 ± 0.01 , which implies that the structures are relatively large and outside the effective range of the SANS (Fig. 2o). In order to fully fit the SANS data for the two-component hydrogels (10 mM), we needed to combine the flexible elliptical cylinder model with a power law (Fig. 2l,m,n) as previously demonstrated.⁵⁹ With this approach, we obtained a radius of 27.5 ± 0.3 Å, an axis ratio of 1.53 ± 0.03 Å, and a Kuhn length of 111.6 ± 2.1 Å for the 4:1 molar ratio **PA-E3/DBS-COOH** hydrogel; a radius of 28.4 ± 0.7 Å, an axis ratio of 1.36 ± 0.06 , and a Kuhn length of 96.3 ± 3.2 Å for the 1:1 molar ratio **PA-E3/DBS-COOH** hydrogel; and a radius of 27.9 ± 1.1 Å, an axis ratio of 1.52 ± 0.10 , and a Kuhn length of 114.4 ± 9.21 Å for the 1:4 molar ratio **PA-E3/DBS-COOH** hydrogel. These results indicate that the nanoscale flexible elliptical cylinders are similar across the data set in terms of radius, axis ratio, and Kuhn length. The small differences in the fitted parameters can be attributed to the minimal perturbation effect of **DBS-COOH** on the **PA-E3** fibre formation during the self-assembly process⁶⁰, suggesting that **PA-E3** fibre assembly still occurs in the presence of **DBS-COOH**. The exponents for the power law are 2.47 ± 0.05 , 2.76 ± 0.04 , and 2.95 ± 0.07 for the hydrogels with 4:1, 1:1, and 1:4 molar ratio of **PA-E3/DBS-COOH**, respectively. For convenient comparison, the data for the fittings are summarized in Supplementary Table S1. These power law exponents suggest that there is an increasing nanofiber network formation with increasing concentration of **DBS-COOH**, in agreement with the LSCM (Fig 1e,f,g), TEM (Fig. 2b,c,d, top panel), and AFM (Fig. 2g,h,i, middle panel) images. However, it is important to mention that the radius of the nanofibers is such that the scattering at mid to high Q range is dominated by the flexible elliptical model as opposed to the power law. We confirmed this dominant flexible elliptical model by plotting the intensity at Q of 0.06151 against % **PA-E3** in the mixture to obtain a straight line fit as opposed to a non-linear fit expected if **DBS-COOH** is entrained within the nanofibers of **PA-E3** (Supplementary Figure S5). Linking this all together, we propose that self-sorting dominates, leading to fibres that consist of **PA-E3** only and others that consist of **DBS-COOH** only. Since the **PA-E3** assembles first, it will do so in the presence of unassembled **DBS-COOH**, and hence the assembly of **PA-E3** may be affected

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3 by presence of the additive. The **DBS-COOH** then assembles to give the self-sorted system,
4 which is known to play an important role in determining bulk properties of hydrogels.⁶¹
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8 **Kinetics of self-assembly**

9 *Micelles act as nuclei for faster assembly on PA containing systems*

10 Understanding the kinetics of self-assembly is not only important for gaining insight into the
11 molecular mechanism, but also for the design and modulation of properties of multicomponent
12 hydrogel systems. In this study, we decided to use dynamic oscillatory rheology to monitor the
13 self-assembly kinetics of **PA-E3**, **DBS-COOH**, and the equimolar **PA-E3/DBS-COOH**
14 mixture. According to the time-sweep experiment within the linear viscoelastic region (LVR)
15 of the hydrogels, **PA-E3** displayed the expected rapid self-assembly (Fig. 3a) whereas **DBS-**
16 **COOH** exhibited a delay of about 20 min prior to self-assembly (Fig. 3b). The **PA-E3** system
17 exhibits a two stage increase in G' and G'' . By analogy to other work,⁶² this can be associated
18 with interactions between initially formed fibers and a decrease of charge in the nanofibers.
19 We stress, however, that given the difficulty to deconvolute the relative effects provided by
20 fiber bundling, cross-linking, and reduction of charges in the fibres, interpretation of the
21 rheology data must be taken with care. Interestingly, with the inclusion of **DBS-COOH**, the
22 equimolar mixture of **PA-E3/DBS-COOH** exhibited a slightly shorter delay of about 10 min
23 prior to self-assembly upon the addition of GdL (Fig. 3c). This suggests that the two-
24 component system displayed a different self-assembly mechanism to pure **DBS-COOH**. It is
25 well known that **PA-E3** tends to form cylindrical micellar nanostructures in water due to
26 desolvation of the apolar palmitoyl group⁶³ and electrostatic repulsion between the negatively
27 charged headgroups.⁶⁴ These structures are expected to provide nucleation sites and thereby
28 reduce the entropic cost of PA self-assembly both in the single and two-component systems.
29 This hypothesis is based on a non-classical nucleation theory, in which nucleation and growth
30 is much faster in a heterogeneous system comprising nuclei than the homogenous counterpart
31 having no preformed nuclei.^{65,66} Therefore, as expected, unlike the **DBS-COOH** system on its
32 own, both **PA-E3** and the equimolar mixture of **PA-E3/DBS-COOH** underwent faster self-
33 assembly.
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55 The self-assembly of **PA-E3**, **DBS-COOH**, and the equimolar **PA-E3/DBS-COOH** mixture
56 into a solid-like gel, as revealed by G' , reached a plateau within 115, 92, and 63 min,
57 respectively (Fig. 3a,b,c), implying that the gels were supersaturated at the plateau region.
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Beyond the gelation point, $\tan \delta$ also decreases with time for the self-assembly of **PA-E3**, **DBS-COOH**, and **PA-E3/DBS-COOH** mixture and then levels off, representing the completion of self-assembly (Supplementary Figure S6). The observed time difference for the full assembly of **PA-E3**, **DBS-COOH**, and **PA-E3/DBS-COOH** equimolar mixtures can be attributed to their distinct self-assembly events. Bearing in mind that **PA-E3** already assumed a pre-gelation nanostructure, there is a high tendency for **PA-E3** to undergo an assembly whereby there is a gradual increase in the number and length of the nanostructures until an insoluble sample-spanning network of nanofibers is formed.⁶⁷ Given the sigmoidal rheographs of **PA-E3** and that of the **PA-E3/DBS-COOH** mixture, we suggest that the self-assembly mechanism of **PA-E3** is consistent with such isodesmic assembly with three set lag phases, initial rearrangement/bundling of the cylindrical micelles, elongation, and supersaturation (Supplementary Figure S1). On the other hand, **DBS-COOH** alone exhibits a different cooperative self-assembly (nucleation-growth) mechanism via an initial formation and growth of nuclei, above which a critical point gelation occurs.⁶⁸ The initiation of nucleation by **DBS-COOH** results in a delayed emergence of the assembly for about 20 min following a prolonged lag phase (Supplementary Figure 1). However, once assembly is initiated, it then rapidly proceeds in a cooperative manner. Similar to the assemblies of **PA-E3**, **DBS-COOH** assembly has three sets of lag phases but with a different initial event – nucleation, elongation, and supersaturation. Interestingly, the **PA-E3/DBS-COOH** assembly shows the fastest establishment (~ 63 min) of an overall gel network. This system (**PA-E3/DBS-COOH**) combines the initial nuclei of the **PA-E3** system, reducing the lag phase, with the cooperative network assembly of **DBS-COOH**, hence optimising both the initial nucleation step and the assembly of the full sample-spanning network (Supplementary Figure 1). This is a clear example of the dual benefits of this two-component assembly system.

The Avrami exponent, n , for the temporal changes in the structure of **PA-E3**, **DBS-COOH** and the equimolar mixture of **PA-E3/DBS-COOH** as a result of gradual pH-switching that drives self-assembly is 2.0, 2.2 and 2.7, respectively (Supplementary Figure S7 – S9). With these values, we deduced that **DBS-COOH** and **PA-E3** self-assembled into a 2D nanostructure while the Avrami exponent obtained for the equimolar **PA-E3/DBS-COOH** mixture suggests the formation of a 3D nanostructure or networks. It is worth noting that these Avrami exponents for **DBS-COOH** are higher than those previously determined by NMR and CD methods.³⁸ This reflects the fact that determining Avrami exponents by rheology will indicate the formation of a sample spanning network in two/three dimensions, whereas the other methods report directly

on one-dimensionally nanofiber assembly. These results indicate a greater degree of three-dimensionality in the presence of **DBS-COOH**, which is in agreement with the earlier discussion about the ability of this gelator to enhance nanofiber aggregation and sample-spanning nanofiber network formation.

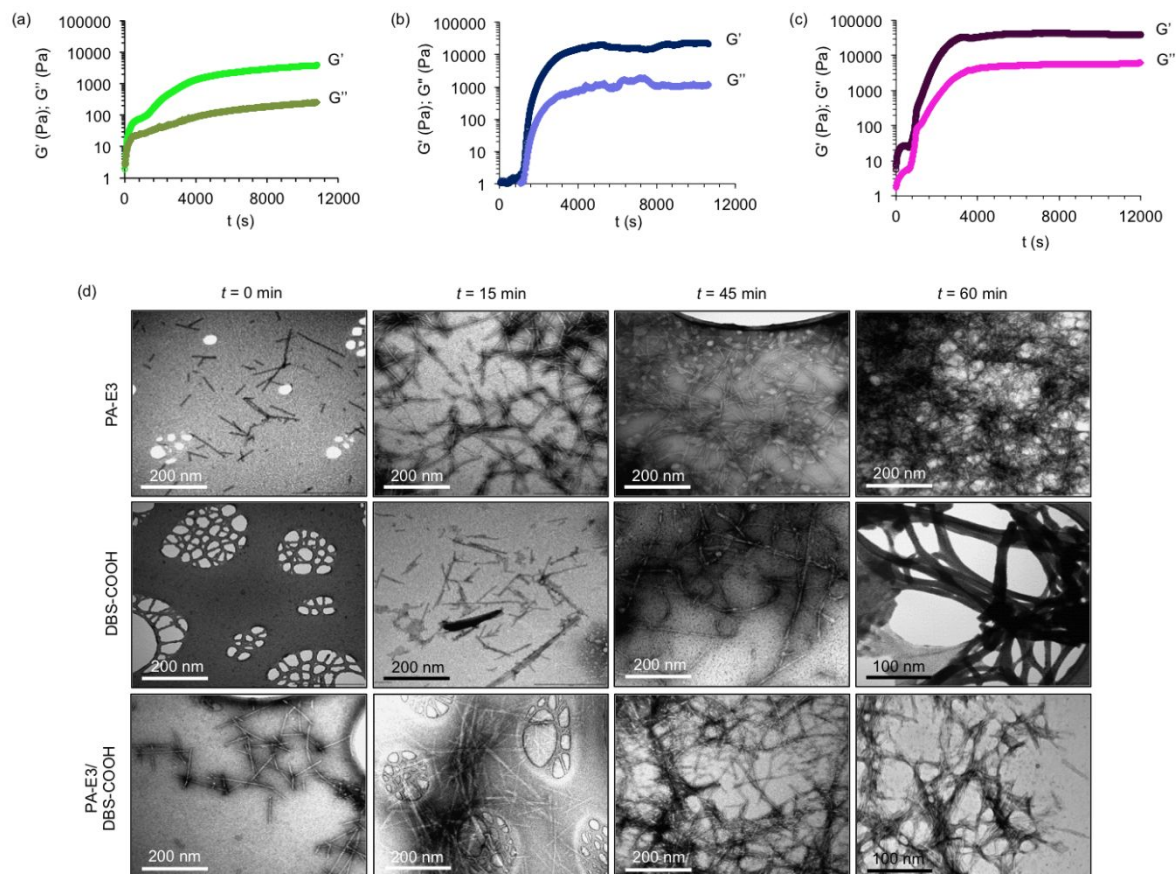


Figure 3. Kinetics of self-assembly into nanofibers. Evolution of nanofibers is monitored during the sol-hydrogels phase transition by recording storage (G') and loss (G'') modulus values as a function of time for 10 mM of (a) **PA-E3**, (b) **DBS-COOH** and (c) equimolar mixture of **PA-E3/DBS-COOH** at 25 °C and self-assembly is initiated by the addition of GdL (10 mg/ml); frequency = 10 Hz, strain = 0.5%. (d) TEM images showing self-assembly of **PA-E3**, **DBS-COOH** and **PA-E3/DBS-COOH** into nanofibers at various time intervals.

Remarkably, the slow self-assembly kinetics that result from using low gelator concentrations (0.01 mM, pH 9) and the *in situ* protonation resulting from GdL hydrolysis facilitated real-time observations of the growth of the nanofibers by TEM. As shown in Fig. 3d, at the onset of self-assembly ($t = 0$), **PA-E3** already exists in a pre-assembled micellar form while **DBS-COOH** exists in a free and monomeric form at time $t = 0$ min. There are also micellar nanostructures with similar dimensions in the equimolar solutions of **PA-E3/DBS-COOH**. Interestingly,

within 15 min of adding GdL, **DBS-COOH** self-assembled into ~200 nm long nanofibers, while the **PA-E3** and **PA-E3/DBS-COOH** micelles grew into ~600 nm long nanofibers. Further into the incubation period (45 and 60 min), the nanofibers grew into entangled, interpenetrated, and long nanofiber networks in all cases. **DBS-COOH** and **PA-E3/DBS-COOH** nanofibers are clearly more bundled than **PA-E3** and **DBS-COOH** alone – in agreement with the kinetic observations from the rheological study described above. The hierarchical nanofiber growth further confirms the nucleation – elongation – supersaturation mechanism we proposed for the stepwise increase in the G' values observed in Fig. 3a,b,c and Fig. 4, which also follows the kinetic model of step-growth of β -amyloid peptide nanofibers.⁶⁹ It is important to mention that the differences between nanofibers of **PA-E3**, **DBS-COOH**, and **PA-E3/DBS-COOH** are not sufficiently significant given the need to air-dry the samples.

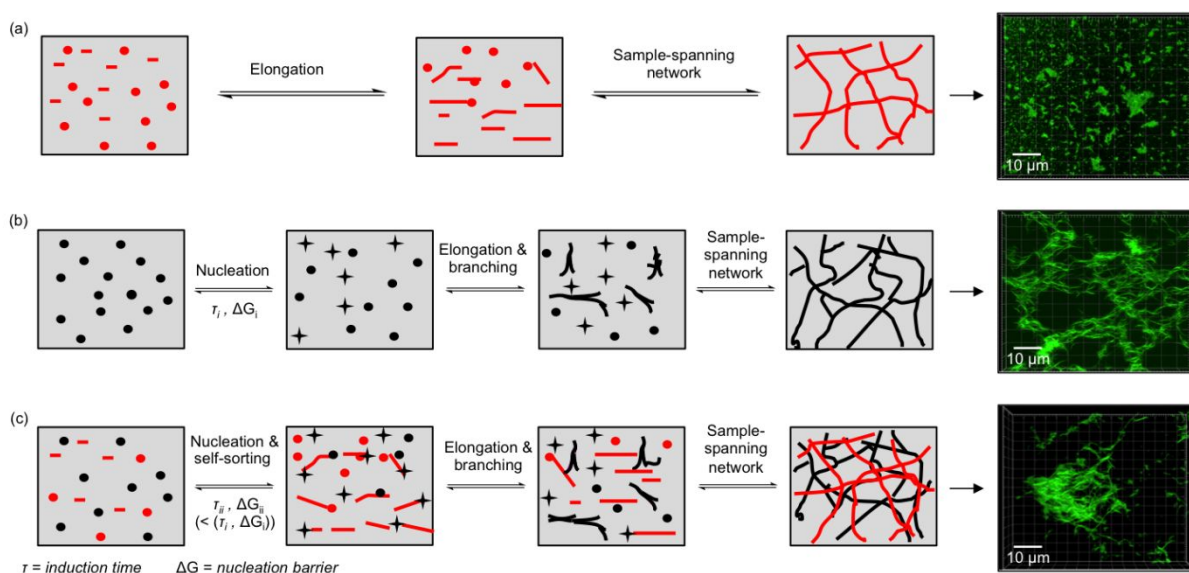


Figure 4. Mechanism of self-assembly. Proposed mechanisms of self-assembly of (a) **PA-E3**, (b) **DBS-COOH** and (c) **PA-E3/DBS-COOH** when triggered with GdL in aqueous systems. The green fluorescent images represent ThT fluorescent micrographs of self-assembled nanofibers.

All-atom molecular dynamics (MD) simulations of self-assembly

It has been established that the structures, self-assembling driving forces, surface charge, and hydrophobicity of building blocks in a multicomponent self-assembling system are key to determining whether they will co-assemble or self-sort.⁷⁰⁻⁷² Therefore, molecular dynamics simulations were performed using Material Studio 8.0 software from Biovia Software Inc (San Diego, California)⁷³ in order to understand how the structural differences, driving forces, and hydrophilicity of **DBS-COOH** and **PA-E3** dictate the type of internal nanostructuring in these

two-component hydrogels. According to solubility parameter calculations, the theoretical solubility parameters of **DBS-COOH** and **PA-E3** are $24.36 \text{ (J/cm}^3)^{1/2}$ and $23.13 \text{ (J/cm}^3)^{1/2}$, respectively, which implies that **DBS-COOH** is slightly more polar than **PA-E3**. Upon mixing the two components, the cohesive energy density (CED) decreases as the ratio of **PA-E3** increases (Supplementary Figure S10). Moreover, the enthalpy of mixing (ΔH_{mix}) values of **DBS-COOH** and **PA-E3** were negative and became more negative with decreasing amount of **PA-E3**, indicating mixing of the two components (Supplementary Figure S11).

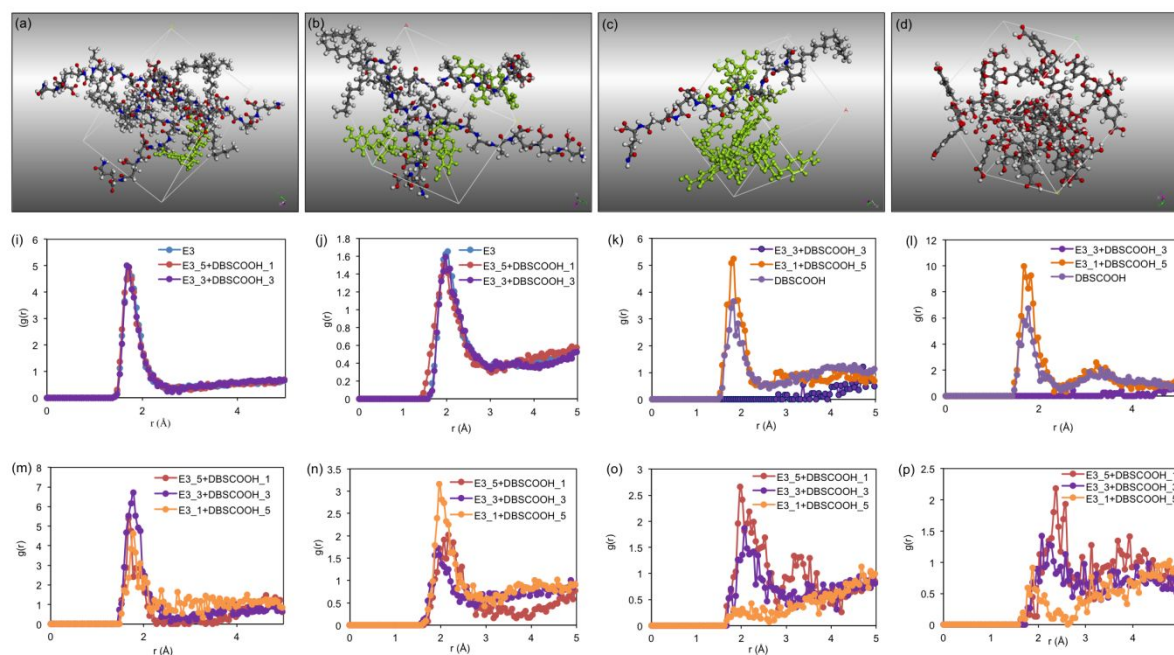


Figure 5. Molecular dynamics simulation. Periodic cubic cell containing (a) 1 molecules of **DBS-COOH** (green) and 5 molecules of **PA-E3**, (b) 3 molecules of **DBS-COOH** (green) and 3 molecules of **PA-E3**, (c) 5 molecules of **DBS-COOH** (green) and 1 molecule of **PA-E3**, (d) 12 molecules of purely **DBS-COOH** after 500 ps dynamic simulation. Intermolecular interaction intensities between (e) H-carboxyl and O-carbonyl of **PA-E3** as pure component and in the presence of **DBS-COOH** at various molar ratios, (f) H-amide and O-carbonyl of **PA-E3** as pure component and in the presence of **DBS-COOH** at various molar ratios, (g) O-carbonyl of **DBS-COOH** and H-O6 of **DBS-COOH** as pure component and in the presence of **PA-E3** at various molar ratios, (h) O6 of **DBS-COOH** and H-O6 of **DBS-COOH** as pure component and in the presence of **PA-E3** at various molar ratios, (i) O-carbonyl of **DBS-COOH** and H-carboxyl of **PA-E3** at various molar ratios, (j) O-carbonyl of **DBS-COOH** and H-amide of **PA-E3** at various molar ratio, (k) O6 of **DBS-COOH** and H-amide of **PA-E3**, and (l) O5 of **DBS-COOH** and H-amide of **PA-E3**.

Molecules of **PA-E3** and **DBS-COOH** were constructed and optimized using DMOL3 quantum mechanical module. Self-assembly of **PA-E3**, **DBS-COOH** and mixtures of both were studied by dynamic simulation using Forcite module with COMPASS II force field. The results after 500 ps dynamic steps (Fig. 5b-e) indicate that introduction of some molecules of **DBS-COOH** into the simulation box of **PA-E3** does not disturb the **PA-E3** self-assembly, suggesting self-sorting of **PA-E3**. However, **DBS-COOH** molecules did not self-sort and instead bind to the surface of the **PA-E3** molecules through multiple hydrogen bonding interactions. At a higher **DBS-COOH/PA-E3** ratio, the bound **DBS-COOH** molecules within close proximity self-assemble, thereby facilitating interfacial interaction with the **PA-E3** (Fig. 5d). This interaction might explain the tendency of the two-component **PA-E3/DBS-COOH** system to form bundled nanofibers with improved mechanical properties. Radial distribution function (RDF) was used to estimate the intermolecular (hydrogen bond) interaction distance between molecules of the individual components (**PA-E3** and **DBS-COOH**) as well as their mixtures. It was found that the most significant intermolecular interactions that underpin **PA-E3** self-assembly are formed between H-carboxyl and O-carbonyl groups (interaction distance of 1.7 Å and intensity of 5 (Fig. 5e)). On the other hand, the dominant intermolecular interactions between **DBS-COOH** molecules are O-carbonyl/H-O6 and O6/H-O6 with interaction distances of 1.8 Å and 1.7 Å, respectively (Supplementary Table 2). Interestingly, when **PA-E3** molecules self-assemble in the presence of **DBS-COOH**, the intensity of the interaction distance between H-carboxyl and O-carbonyl groups as well as H-mide and O-carbonyl of **PA-E3** remain constant (Fig. 5e,f), suggesting that the presence of **DBS-COOH** does not affect self-assembly of **PA-E3**. However, by measuring the interaction distance between O-carbonyl/H-O6 as well as O6/H-O6 of **DBS-COOH** at 1:1 (**DBS-COOH:PA-E3**) molar ratio, we observed no interaction formed between **DBS-COOH** molecules (Fig. 5g,h). Thus, indicating that **DBS-COOH** molecules preferentially interact with the self-assembled **PA-E3** structure at this molar ratio. When the molar ratio of **DBS-COOH** increases to 5:1 (**DBS-COOH:PA-E3**), a significant interaction is formed between **DBS-COOH** molecules (Fig. 5g,h). In all cases, intermolecular hydrogen bond (H-bond) interactions are formed between **PA-E3** and **DBS-COOH** by several groups but the most significant interaction is formed between O-carbonyl of **DBS-COOH** and H-carboxyl of **PA-E3** with an interaction distance of 1.7 Å (Supplementary Table S2). The interactions that the H-Amide of **PA-E3** form with various groups of **DBS-COOH** are less significant and in all cases the interaction intensity decreases with the decrease of **PA-E3** concentration (Fig. 5i,j,k,l). **DBS-COOH** does not affect the interactions between **PA-E3** as revealed by RDF results, suggesting that **PA-E3** undergoes

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3 self-sorting. However, the presence of **PA-E3** causes **DBS-COOH** to alter its self-assembly
4 fashion, which might corroborate our findings with SANS and dynamic time sweep rheology.
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8 In summary, the interactions mediating the self-assembly can be discussed on the basis of the
9 energetics of molecular interactions. The interactions that formed the self-assembled structures
10 between **PA-E3** and **DBS-COOH** will be a balance between all possible energies. The **PA-E3**
11 self-assembly is driven by the hydrophobic collapse of the hydrocarbon tail while the polar
12 heads are displayed on the periphery of the nanofibers. Therefore, the energy associated with
13 the assembly of **PA-E3** is too large for **DBS-COOH** to fully disrupt, but then **DBS-COOH**
14 can form some adventitious interactions with parts (polar head group) of **PA-E3** that are not
15 responsible for self-assembly in order to provide some favourable energetics. In this way, small
16 amounts of **DBS-COOH** interact with the assembling **PA-E3** but without disrupting it. As the
17 concentration of DBS increases, self-self interactions become more important for **DBS-COOH**
18 because there are more potential **DBS-COOH** binding partners, resulting in interactions
19 between **DBS-COOH** molecules becoming more energetically significant.
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30 **Viscoelastic and self-recovery properties of hydrogels**

31 Given the increasing hydrogel translucency that is observed with growing concentration of
32 **DBS-COOH** in the **PA-E3/DBS-COOH** hydrogels (Fig. 1b) as a result of higher nanofiber
33 density, we used dynamic oscillatory rheometry to quantify its effect on the bulk properties
34 (stiffness) of the hydrogels. Based on amplitude-sweep measurements, hydrogels for **PA-E3**
35 (10 mM) and **DBS-COOH** (10 mM) exhibited storage moduli (G') of ~ 9.5 kPa and ~ 27 kPa,
36 respectively (Fig. 6a). Interestingly, G' values increased linearly with increasing percentage of
37 **DBS-COOH** in the **PA-E3/DBS-COOH** hydrogels (Fig. 6a and Supplementary Figure S12),
38 indicating a synergistic effect. Also, from the amplitude-sweep measurements, the critical
39 strain (i.e. the strain at which the hydrogels breaks) for **DBS-COOH** and **PA-E3** were 0.8
40 (Supplementary Figures S13) and 2.7% (Supplementary Figures S14), respectively. The
41 critical strain values for the two-component hydrogels range between 1 and 2%
42 (Supplementary Figures S15-18) and slightly decrease as the amount of **PA-E3** in the hydrogels
43 decreases, suggesting that **PA-E3** is able to promote formation of resilient hydrogels. Previous
44 studies have shown that π - π stacking between DBS gelators confers enhanced rigidity on their
45 self-assembled structures.⁴⁰ This rigid structure formation suggests a reason for higher G'
46 values (also lower values of $\tan \delta$) of **DBS-COOH** hydrogels and the **PA-E3/DBS-COOH**
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than the non-aromatic **PA-E3** hydrogels (Supplementary Figure S13). It is important to note that the total concentration of the gelators remain fixed (10 mM) in all multicomponent hydrogels, so differences cannot be assigned to total gelator loading.

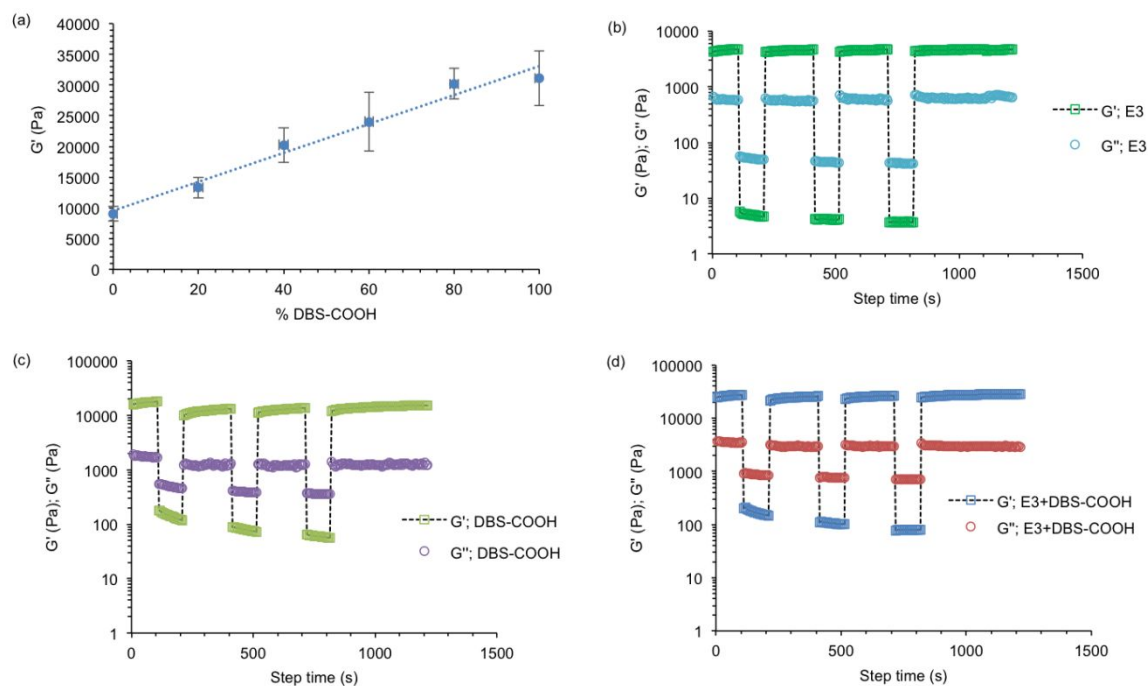


Figure 6. Mechanics and self-healable properties. (a) Plots of G' against percentage of **DBS-COOH** in the hydrogels. (b), (c) and (d) show the self-recovery of **PA-E3**, **DBS-COOH** and equimolar concentration of **PA-E3** and **DBS-COOH**, respectively. In all cases, the hydrogels were subjected to a constant frequency of 10 Hz and strain of 0.1% (100 s), 100% (100 s), 0.1% (200 s), 100% (100 s), 0.1% (200 s), 100% (100 s) and 0.1% (400 s).

Self-assembly through non-covalent bonds such as π - π stacking, hydrogen bonding, and host-guest interactions holds great potential for constructing self-healable hydrogels due to their intrinsic reversibility.⁷⁴⁻⁷⁶ For example, Stevens and co-workers have harnessed the ability of non-covalent β -sheet cross-links to fabricate self-healable hybrid hydrogels owing to their ability to reassemble and recover fully after being strained to failure.⁷⁷ Giving the structural dissimilarity between the aromatic sugar gelator (**DBS-COOH**) and the β -sheet forming PA gelator (**PA-E3**), we wanted to dissect the impact of the interplay of the π - π stacking and hydrogen bonding that **DBS-COOH** provides and the non-covalent β -sheet cross-links that **PA-E3** offers to the self-healing ability of the **PA-E3/DBS-COOH** hydrogel.

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3 To test this hypothesis, the thixotropic properties of the hydrogels after high shear loads were
4 measured in dynamic time sweep experiment under strain amplitudes within (0.1%) and
5 beyond (100%) the linear viscoelastic region (LVR) of the hydrogels (Supplementary Figure
6 S12 for the LVR). Under high shear load, the hydrogels undergo internal breakage as indicated
7 by the significant decrease and inversion of G' and G'' values. This signifies that the liquid-like
8 behaviour of the hydrogels dominates under high strain amplitude, which indicates that the
9 hydrogels have been broken. After 3 cycles of strain amplitude, hydrogels of **PA-E3** displayed
10 full (100%) recovery (Fig. 6b) while **DBS-COOH** hydrogels only recovered 62% (Fig. 6c) and
11 hydrogels with a 1:1 molar ratio of **PA-E3/DBS-COOH** exhibited 98% recovery (Fig. 6d).
12 The two-component gel, therefore, exhibited enhanced stiffness relative to individual building
13 blocks as a result of the **DBS-COOH** aromatic structure while maintaining a high level of
14 recovery compared to **DBS-COOH** hydrogels due to the **PA-E3** non-covalent β -sheet cross-
15 links. This result demonstrates how the two distinct gelators cooperate in the multicomponent
16 hydrogels. Also, these data reveal how interactions of self-assembling building blocks at the
17 nanoscale and the spatio-temporal arrangement of the nanofibers can translate into materials
18 bulk properties at the macroscale level.^{78,79}

31 32 **Proteolytic stability and effects on cell morphology**

33 To further demonstrate the synergistic material properties that can emerge from the
34 multicomponent **PA-E3/DBS-COOH** hydrogels, we assessed the material's resistance to
35 proteolysis. PAs have been widely used *in vivo* as therapeutic agents in regenerative medicine.
36 However, the peptide bonds with L-chirality are known to quickly degrade after systemic
37 delivery.⁸⁰ Instability of alkylated L-peptides to proteolysis has also been reported.⁸¹ Therefore,
38 D-enantiomers with a better resistance to proteolysis have previously been used as a substitute
39 or a protective corona for the L-enantiomers by co-assembly.⁸²⁻⁸⁴ Covalent attachment of
40 polymers to the peptides surface is another approach that has previously been explored.⁸⁰ For
41 example, Stupp and co-workers employed a supramolecular co-assembly strategy to develop
42 proteolytically stable PA nanofibers consisting of an anti-tumour PA in combination with its
43 PEGylated analogue.⁸⁵ The use of a non-peptidic low-molecular weight gelator as a protective
44 corona for L-peptides against premature proteolysis is another approach that has not been
45 explored in the field of biomaterials development. Our self-assembled system bearing well-
46 defined structures of a lipidated L-peptide and acetalized D-sugar would provide an alternative
47 approach to extend the functionality of PAs.

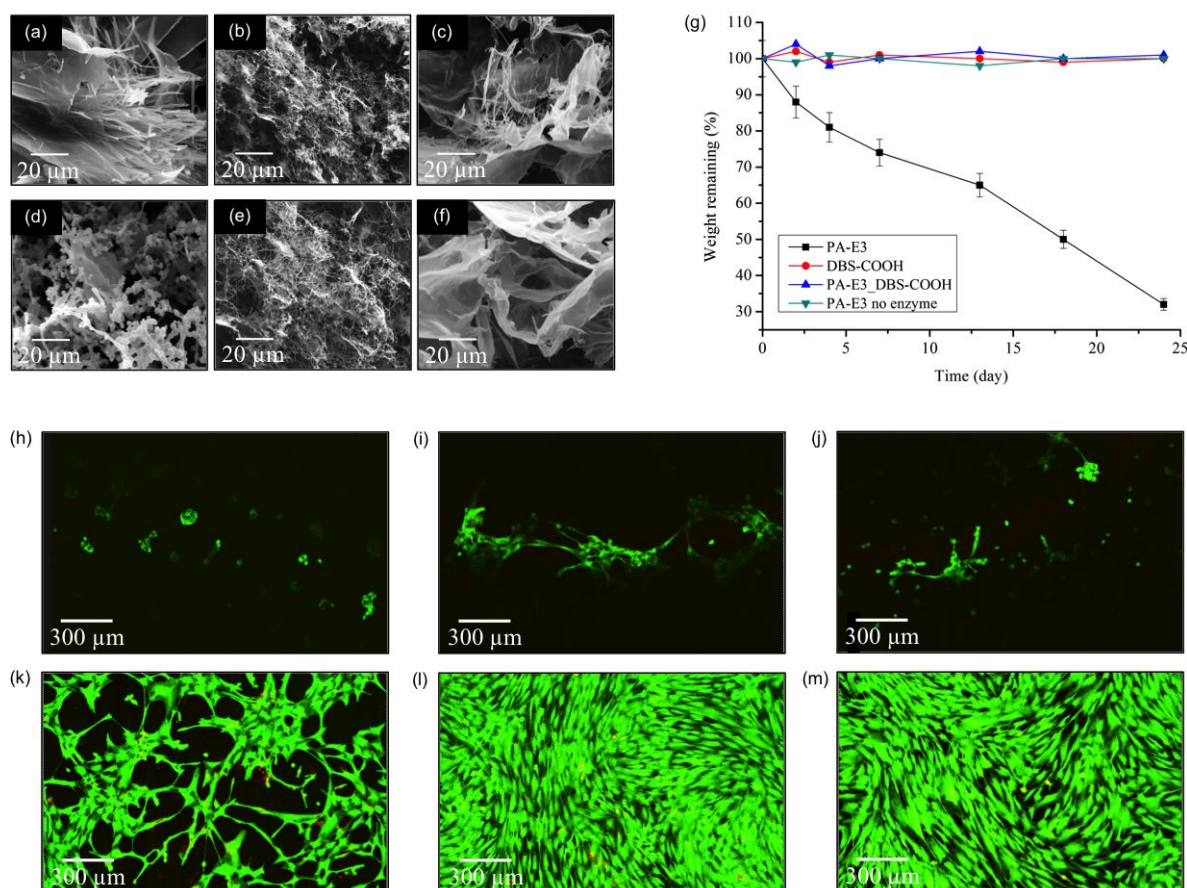


Figure 7. Biological functionalities. Stability of nanofibers of (a) **DBS-COOH**, (b) **PA-E3** and (c) 1:1 molar ratio of **DBS-COOH** and **PA-E3** submerged in PBS solution. (d) **DBS-COOH**, (e) **PA-E3** and (f) 1:1 molar ratio of **DBS-COOH** and **PA-E3** after incubation with proteinase K for 5 days, as revealed by SEM. (g) Degradation profile of **PA-E3**, **DBS-COOH** and **PA-E3/DBS-COOH** hydrogels after incubation with proteinase K at 37 °C for 22 days. **PA-E3** hydrogels were also incubated in PBS under similar conditions. LSCM images of human adipose-derived stem cells seeded on hydrogels of (h) **PA-E3**, (i) **PA-E3/DBS-COOH** (4:1), (j) **PA-E3/DBS-COOH** (1:1), (k) **PA-E3/DBS-COOH** (1:4), (l) **DBS-COOH**, and (m) cells pated on TCP for 4 days. Scale bar: 300 μm.

To investigate proteolytic stability, **PA-E3/DBS-COOH** and control hydrogels were interacted with aqueous solution of proteinase K (50 mg/mL). Digestions were assessed using scanning electron microscopy (SEM) to characterize the morphology of **PA-E3**, **DBS-COOH** and **PA-E3/DBS-COOH** hydrogels. Unlike the nanostructure of **PA-E3** hydrogels submerged in PBS (Fig. 7a), Fig. 7d shows that the **PA-E3** hydrogel nanostructures were digested and transformed into aggregates of nanoparticles after incubating with proteinase K for 5 days. This structural transformation is reminiscent of a report by Hamley and co-workers using chymotrypsin to

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3 induce nanofiber-nanoparticle transformation in a PA solution.⁸¹ As expected, **DBS-COOH**
4 hydrogel nanostructures (Fig. 7b) remained intact after treatment with proteinase K (Fig. 7e),
5 suggesting that **DBS-COOH** lacks the specific motifs for enzymatic cleavage. Indeed,
6 equimolar mixtures of **PA-E3/DBS-COOH** hydrogels were resistant to enzymatic digestions
7 as the nanostructures remained intact (Fig. 7f) when compared with nanostructure of the
8 hydrogels that were not treated with proteinase K (Fig. 7c), suggesting that **DBS-COOH** acted
9 as a protective corona around **PA-E3** nanofibers in the two-component hydrogels, making it
10 impossible for the enzyme to access peptide bonds in **PA-E3**. To further elucidate the
11 enzymatic degradation of the **PA-E3** by proteinase K, liquid chromatography-mass
12 spectrometry (LC-MS) was employed. As shown in Supplementary Figure S19a,aii, an LC-
13 MS spectrum for the **PA-E3/proteinase K** mixture reveals multiple peaks with m/z values of
14 1177.80 Da, 1155.70 Da, 1078.70 Da, 847.5 Da, 846.5 Da, 817.7 Da, 520.2 Da and 418.3 Da.
15 The sharp peaks with m/z values 1177.80 Da and 1155.70 Da correspond to $[M+Na]^+$ and
16 $[M+H]^+$, respectively, where M is the calculated mass of **PA-E3**. The remaining peaks are the
17 masses of the **PA-E3** fragments following proteolysis. Also, the ultraviolet-visible (UV)
18 chromatogram displayed two broad bands after digestion (Supplementary Figure S19c).
19 Interestingly, only two peaks with m/z values 1177.80 Da and 1155.70 Da which correspond
20 to $[M+Na]^+$ and $[M+H]^+$, respectively were observed on the spectrum of **PA-E3/DBS-**
21 **COOH/proteinase K** mixture (Supplementary Figure S19b), suggesting that **DBS-COOH**
22 indeed acted as a protective corona for **PA-E3** in the multicomponent hydrogels, limiting the
23 proteolysis of **PA-E3**. Impacts of this morphological transformation on the macroscopic
24 hydrogels were also assessed by a weight loss measurement approach. After incubating the
25 **DBS-COOH** and **DBS-COOH/PA-E3** hydrogels in enzyme solution as well as **PA-E3**
26 hydrogels in PBS for 22 day, the weight ratio remained constant. In contrast, the weight ratio
27 of **PA-E3** hydrogels in enzyme solution drastically reduced within 22 days of incubation (Fig.
28 7g). These results further support our argument that the macroscopic properties of
29 multicomponent hydrogels is dependent of the nanoscale composition.

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51 In order to investigate the response of cells to the multicomponent hydrogels, we cultured
52 human adipose derived stem cells (ADSCs) on the hydrogels prepared with various
53 stoichiometric ratios of **PA-E3** and **DBS-COOH**. After 4 days of incubation, the cells were
54 stained with live/dead dyes. Cells stained with the green dye (calcein AM) were alive while
55 those stained with the red dye (ethidium homodimer) were dead. As shown in Fig. 7h, the cells
56 on the **PA-E3** hydrogels assumed a round morphology after 4 days in culture. In contrast,
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3 hydrogels of **DBS-COOH** were fully covered with live cells. Also, the cells exhibited an
4 elongated morphology (cell spreading) (Fig. 7l) similar to cells cultured on tissue culture plastic
5 (TCP) (Fig. 7m). For the multicomponent hydrogels with 4:1 and 1:1 ratios of **PA-E3/DBS-**
6 **COOH**, there was a mixture of both round and elongated cells. However, the cells seeded on
7 hydrogels with 1:4 ratio of **PA-E3** and **DBS-COOH** formed a network of elongated
8 morphology (Fig. 7k). Evidently, cells were more viable on **DBS-COOH** hydrogels than **PA-**
9 **E3** hydrogels. The multicomponent hydrogels are intermediate between the two – clearly **DBS-**
10 **COOH** can impart some of cell compatibility to these hybrid materials. The tunable
11 morphology of the cells also show how molecular composition and mechanical properties of
12 supramolecular hydrogels can potentially be used to influence cell growth, and potentially in
13 future, decide cell phenotypes. We hope to characterize cell differentiation in detail in the
14 future.

27 Conclusion

28 We report on the synergistic properties arising from the self-sorting of a PA and a DBS-based
29 gelator by slow acidification. Compared to the individual components, the resulting
30 multicomponent hydrogels exhibited a series of properties including tunable nanoscale
31 morphology, improved stiffness (endowed by DBS gelator), enhanced self-healing (endowed
32 by the PA), stability to enzymatic degradation (provided by the DBS gelator), and cell
33 compatibility (provided by the DBS gelator). Induced circular dichroism of ThT, SANS and
34 molecular dynamics simulations demonstrated self-sorting with the **DBS-COOH** initially
35 acting as an additive adsorbed on the surface of the **PA-E3** nanofibers through multiple
36 hydrogen bonding interactions. The adsorbed sugar molecules within close proximity then
37 interact with each other, thereby facilitating interfacial interactions between nanofibers. We
38 reason that these interfacial interactions impart the nanofiber network with enhanced stiffness
39 as well as resistance to enzymatic degradation. With regards to the assembly mechanism, the
40 **PA-E3** exhibits different assembly kinetics than **DBS-COOH**, with the **PA-E3** benefitting
41 from the presence of multiple cylindrical micellar nuclei, avoiding the lag-phase inherent in
42 the **DBS-COOH**. However, the assembly of the DBS gelator is more cooperative and rapid
43 once nucleation has occurred. Interestingly, the multicomponent system benefits from both
44 advantages of the PA and DBS systems, and is the fastest to establish a full-sample-spanning
45 gel network. Finally, human adipose-derived stem cells could be cultured on the DBS gelator
46 and the two-component hydrogels with sufficiently high concentration of **DBS-COOH**,

opening new opportunities for developing complex hydrogels for regenerative medicine. In summary, we have developed and characterized a **PA-E3/DBS-COOH** hybrid system that expands the scope and capacity of both **PA-E3** and **DBS-COOH** with each of the two LMWG systems endowing the resulting hybrid gel with its own unique capabilities.

Supporting information

The Supporting Information is available free of charge on the ACS Publications website at DOI: <https://eur01.safelinks.protection.outlook.com/?url=http%3A%2F%2Fpubs.acs.org&data=02%7C01%7Ca.mata%40qmul.ac.uk%7Cf1b67868864547098c4408d725963039%7C569df091b01340e386eebd9cb9e25814%7C0%7C1%7C637019199296442776&sdata=6Dq5Ix6jNx%2Fy04aVYJBbE9XtZfkTyoFWW2wQ8X26Rqk%3D&reserved=0>.

Overview of multicomponent self-assembly; ThT fluorescent microscopy for sol samples; full range CD spectra for ThT-gelator interactions; TEM images of gelator sols; SANS fitting parameters; tan delta values for gelator self-assembly kinetics; Avrami equations and plots; molecular dynamics computation; amplitude sweeps and critical strain for hydrogels; UV-TIC spectra for hydrogel digestions.

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TOC

