Supramolecular nano-fibrous peptide/polymer hydrogels for the multiplexing of bioactive signals

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ABSTRACT

The ability to provide multiple functions within a single scaffold biomaterial is a major goal in tissue engineering. Self-assembling peptide-based hydrogels are gaining significant attention as three-dimensional biomaterials because they provide a network of nanofibers similar to the native extracellular matrix while allowing the presentation of multiple biochemical cues for cell signalling. Herein, we combine a positively charged peptide amphiphile (PA) and the negatively charged synthetic polymer poly(sodium 4-styrenesulfonate) (PSS) to fabricate hybrid hydrogels through supramolecular self-assembly. PSS/PA hydrogels show rather high mechanical stiffness while being stable in buffered environment. The sulfonate functionality in PSS promotes hydrogel mineralization which can be controlled if undertaken in standard osteogenic medium. Loading proteins with different charges in the hydrogels reveals their ability to retain and sustain their release and indicates their potential for the controlled delivery of growth factors. Human mesenchymal stem cells encapsulated in PSS/PA hydrogels remain viable. The biomimetic nanofibrous structure of the hydrogels, together with multiplexing of bioactive signals, can provide a suitable environment for stem cell differentiation.

Key words: Supramolecular hydrogels, nanofibers, bioactive signals, biomimetic mineralization, protein delivery, 3D stem cell culture
**Introduction**

The extracellular matrix (ECM) of tissues serves multiple roles to ensure tissue homeostasis, from providing anchorage sites for cell adhesion and migration to regulating the availability of soluble growth factors (GFs) for cell proliferation and differentiation. Mimicking the multifunction features of the native ECM has been a major goal when designing biomaterials for tissue engineering and regenerative medicine. In the recapitulation of the ECM, several methods have been pursued, from macromolecular crowding and self-assembly to electrospinning and patterning. For example, to replicate the ECM biophysical properties, the typical nanofibrillar collagen network, electrospinning\(^1\) and self-assembly\(^2\) have been used. However, electrospinning uses polymer solutions dissolved in non-aqueous solvents and an electric field to produce nanofibers. These harsh conditions unable the incorporation of cells during nanofiber processing. In contrast, self-assembly can occur at physiological conditions generating nanofibers with diameters as small as 10 nm and smaller than those obtained by electrospinning. In addition, self-assembled peptide nanofibers can be designed to contain sequences derived from ECM proteins\(^3,4\) further extending the ability of mimicking ECM biochemical properties. The ability of the ECM to regulate the activity of GFs results from the supramolecular binding of GFs to insoluble ECM components, notably to heparan sulfate proteoglycans via electrostatic interactions.\(^5\) Several creative approaches have been reported to emulate the GF binding to sulfated glycosaminoglycans (GAGs) in native ECMs, including sulfation of polysaccharides\(^6,7\); incorporation of heparin through covalent bonds\(^8\) or supramolecular interactions using specific\(^9\) or non-specific\(^10,11\) peptide sequences and in polyelectrolyte complexes\(^12,13\); synthesis of heparin-like molecules\(^14\) (e.g. sulfonated peptides). From these studies, it becomes obvious that self-assembling peptides can provide much of the ECM functionality (cell adhesion motifs such as
RGD, sulfonated groups, nanofibrillar structure). Peptides can also be combined with (bio)polymers to self-assemble into multicomponent hydrogels, either through the formation of covalent linkages or non-covalent interactions.\textsuperscript{17}

Herein, we used self-assembly to prepare multi-functional hydrogels through electrostatic interactions between a positively charged peptide amphiphile (PA) and a negatively charged synthetic polymer bearing sulfonate functionality, poly(sodium 4-styrenesulfonate) (PSS) (Figure 1-A). Previous studies showed that sulfonate groups can promote the nucleation of calcium ions\textsuperscript{18,19} and we have thus exploited PSS-induced hydroxyapatite mineralization in this hydrogel system. The negative sulfonate groups in PSS can also provide binding sites for basic proteins, similar to sulfated glycosaminoglycans. The dual functionality of PSS has been utilized in combination with PAs of opposite charge to form self-assembled nanofibrous hydrogels. We show that these hydrogels are able to mineralize in cell culture conditions, control the release of positively charged proteins and support the viability and spreading of stem cells.

**Figure 1.** Self-assembling nano-fibrous hydrogels for the multiplexing of bioactive signals. (A) Self-assembly of peptide amphiphile (PA) and PSS forming a self-supporting hydrogel; (B) Calcium phosphate mineral nucleation on the hydrogel’s nanofibrous structure; Hydrogel can be used as a scaffold for protein delivery (C) and 3D cell encapsulation (D).
Experimental Section

Materials and methods

**Materials.** Fmoc-protected amino acids, Fmoc-Ala-OH, Fmoc-Lys(Boc)-OH, Fmoc-Val-OH, Fmoc-Arg(Pbf)-OH, Fmoc-Gly-OH, Fmoc-Asp(OtBu)-OH, Fmoc-Ser(Tbu)-OH and 4-methylbenzhydrylamine (MBHA) rink amide resin (100-200 mesh) were purchased from Novabiochem (Merck, UK). Triisopropylsilane (TIS) and piperidine were obtained from Alfa Aesar (UK). Poly(sodium 4-styrenesulfonate) with two different molecular weights (70 kDa and 1 MDa) and palmitic acid (C_{16}H_{32}O_{2}) were obtained from Sigma-Aldrich, UK. Fluorescent-tagged bovine serum albumin (BSA-FITC) was acquired from Sigma-Aldrich (UK) and lysozyme labelled with Rhodamine B and FITC (lysozyme-FITC) were obtained from Nanocs (US). Cell culture medium (DMEM), fetal bovine serum (FBS) and penicillin-streptomycin (Pen/Strep) were purchased from Gibco (Thermo Fisher scientific, UK). Other chemicals were purchased from Sigma-Aldrich (UK) unless otherwise stated.

**Peptide synthesis and purification.** Peptide amphiphiles (C_{16}V_{3}A_{3}K_{3} and C_{16}V_{3}A_{3}K_{3}G_{3}RGDS) were synthesized in an automated peptide synthesizer (Liberty Blue, CEM, UK) using standard 9-fluorenylmethoxycarbonyl (Fmoc)-based solid phase synthesis protocol. Peptides were grown on the solid support, MBHA rink amide resin, using (4:4:4) equivalents of Fmoc-protected amino acids, 1-hydroxybenzotriazole hydrate (HOBt, Carbosynth Ltd. UK) and N,N'-diisopropylcarbodiimide (DIC, Thermo Fisher, UK) for the coupling reactions. Fmoc group was deprotected using 20% (v/v) piperidine in dimethylformamide (DMF, VWR, UK). Before cleavage from the resin, the peptide N-terminal was capped with a 16-carbon alkyl chain (C_{16}, palmitic acid) using a similar coupling protocol. The removal of the side-chains protecting
groups and cleavage of the peptides from the resin was performed with a mixture of trifluoroacetic acid (TFA, Thermo Fisher, UK)/ triisopropylsilane (TIS)/ water (95%/2.5%/2.5% (v/v)) for 3 hours at room temperature. The peptide solution was collected and reduced down to a viscous solution in a rotary evaporator. Cold diethyl ether was used to precipitate the peptide. After centrifugation, the supernatant was discarded and the solid precipitate allowed to dry overnight. The peptide mass was confirmed by electrospray ionization mass spectrometry (ESI-MS) using a single quadrupole mass detector (SQ Detector 2, Waters, UK). A Waters 2545 Binary Gradient high-performance liquid chromatography (HPLC) system with a preparative reverse-phase C18 column (X-Bridge prep OBD, 5 μm, 30×150 mm column, Waters, UK) was used for purification of the crude peptides using a water/acetonitrile (0.1% TFA) gradient. Fraction collection was made based on the peptide mass and acetonitrile removed by rotary evaporation followed by lyophilisation to obtain the peptide in powder form. Pure peptide amphiphile was treated with hydrochloric acid (10 mM) in three cycles of lyophilization in order to remove the TFA counter ions. Finally, the peptides were dialyzed against ultrapure water using 500 MWCO dialysis membrane (Spectrum Labs, The Netherlands) and lyophilized. Purity of the peptide was confirmed using analytical reverse-phase HPLC (Alliance HPLC system coupled with 2489 UV/Vis detector, Waters, UK). Peptide sample (1 mg ml⁻¹) was injected and run through a C18 column (XBridge analytic 3.5 μm, 4.6 x 150 mm, Waters, UK) using a gradient of water/acetonitrile (0.1% TFA) at 1 ml/min.

**Zeta potential.** PA and PSS (70 kDa and 1 MDa) solutions were analysed in a Nano-ZS Zetasizer (Malvern Instruments) for measuring their zeta potential. Samples were prepared at 0.05 wt% in water (pH 7) and PBS (pH 7.4).
**Preparation of PSS/PA hydrogels.** PA and PSS solutions were prepared by dissolving the powders in phosphate buffer saline (PBS) to obtain the desired concentration. Hydrogels were produced in a 96 well plate by casting 50 µL of a 2% (wt%) PSS (70 kDa or 1 MDa) solution followed by 50 µL of 2% (wt%) PA solution. Hydrogels were formed after incubation at 37 °C for 30 minutes.

**Oscillatory Rheology.** All rheological experiments were performed in a Discovery Hybrid Rheometer (DHR-3, TA Instruments, USA) fitted with a 8 mm diameter plate geometry (advanced peltier plate, APP). Strain and frequency sweeps were performed on the various hydrogels (n≥3) at 1 Hz frequency and 0.1% strain, respectively. Measurements were carried out at 25 °C with a loading gap height set to 0.5 mm.

**Stability of hydrogels.** The stability of the PSS/PA hydrogels was studied by incubating them in aqueous conditions (water and PBS buffer pH 7.4) at 37 °C. The absorbance of the styrene ring at 260 nm was used to monitor the release of PSS in the medium over the time (28-days period). PSS concentration was calculated from a calibration curve obtained from PSS standards. Three repeats were performed for each hydrogel condition and time.

**Scanning electron microscopy (SEM).** The microstructure of the hydrogel surface and cross-section was examined by scanning electron microscopy (SEM). Samples for SEM observation were prepared by incubating the hydrogels in 2% glutaraldehyde in PBS for 1 hour at 4 °C. After three cycles of washing with phosphate buffer saline (PBS), hydrogels were progressively dehydrated using graded ethanol concentrations. Ethanol removal was performed using a critical point dryer (EMS 850, Electron Microscopy Sciences, USA). Samples were first coated with a gold layer (5-30 nm) using an Emitech SC7620 sputter coater (Quorum Technologies, UK) then
imaged in a field emission gun scanning electron microscope (FEG SEM, Inspect F50, FEI, Netherlands).

**Hydrogels mineralization.** The hydrogels were incubated in stimulated body fluid (SBF) solution for 3, 7 and 14 days at 37 °C. The SBF solution contains ion concentrations similar to those of the human blood plasma (Na\(^+\) 142.0, K\(^+\) 5.0, Ca\(^{2+}\) 2.5, Mg\(^{2+}\) 1.5, Cl\(^-\) 147.8, HCO\(_3^-\) 4.2, HPO\(_4^{2-}\) 1.0, SO\(_4^{2-}\) 0.5 mM).\(^{20,21}\) SBF solution was replenished twice a week. After each immersion time, the hydrogels were removed from SBF, washed with distilled water and prepared for SEM observation, as described in prior section. Hydrogels were also tested for their mineralization ability in culture medium with and without osteoinducer agents (10 mM β-glycerophosphate, 50 μg ml\(^{-1}\) 2-Pho2-Phospho-L-ascorbic acid trisodium salt and 10 nM dexamethasone) for 7, 14 and 21 days. Samples were also washed with PBS before sample preparation for SEM. To determine the composition of the mineral formed, hydrogels were analysed by SEM (FEG SEM, Inspect F50, FEI, The Netherlands) equipped with energy dispersive electron X-ray (EDX) spectroscopy (Oxford Instruments, UK). For the EDX analysis, carbon coating (carbon thread evaporation) was used and the samples were analysed at 10.0 kV voltage. The atomic percentage of minerals were determined using INCA software (Oxford Instruments, UK).

**FTIR spectroscopy.** In order to confirm the formation of calcium phosphate minerals on the hydrogels, mineralized hydrogels were burnt at 700 °C to obtain the powder containing the inorganic component. Then, the IR spectra of mineral powder and dried non-mineralized hydrogels were analysed using Brukler Tensor 27 FTIR spectrometer using an attenuated total reflection (ATR) diamond crystal accessory (Bruker, Germany). The absorbance spectrum was measured in the range of 2000–400 cm\(^{-1}\) by averaging 32 individual scans per sample at a
resolution of 4 cm⁻¹.

**Protein loading and release.** Hydrogels were loaded with two labelled model proteins having positive (lysozyme-FITC) and negative (BSA-FITC) charges to examine their release profile over time in physiological conditions. The proteins were loaded in the hydrogels at final concentration of 0.05 mg·ml⁻¹ by first dissolving them in PSS (BSA-FITC) or PA (lysozyme-FITC) solutions followed by the formation of the gel as described before. Then the loaded hydrogels were incubated in 1 mL of PBS (pH 7.4) at 37 °C and aliquots (1 mL) were collected at predetermined time intervals and replaced with fresh PBS (1 mL). Collected samples were kept frozen until experiment completion and then lyophilized. Release experiments were done in triplicate and the protein concentration was determined by dissolving the lyophilized samples in 200 µl of PBS and then measuring the fluorescence intensity at excitation/emission wavelengths of 485/520 nm for both BSA-FITC and lysozyme-FITC and calculating the corresponding concentration from a calibration curve constructed for each labelled protein. The amount of protein loaded in each gel was determined by extracting the protein from the gels through physical/mechanical breakdown of the gels (pipetting and sonication) followed by repeated centrifugation and washings of the gel pellet until no fluorescence was detected in the supernatant. The percentage of protein released at each time point was calculated based on the protein amount loaded in the gel and cumulative release plotted over time. The distribution of proteins in the hydrogels was examined by confocal microscopy (Leica TCS SP2) using 488 nm and 544 nm laser beams for BSA-FITC, lysozyme-FITC and lysozyme-Rhodamine, respectively.

**Lysozyme activity determination.** For determining the activity of released lysozyme from the PSS/PA gels, unlabelled enzyme (from hen egg white, Sigma-Aldrich) was loaded in the 1 MDa PSS/PA gels and release studies were conducted as described above. Aliquots were collected and
kept frozen until further analysis. Later, samples were first lyophilized and dissolved in 100 μl of reaction buffer. Enzyme activity was measured using the EnzChek_Lysozyme Assay Kit (Thermo Fisher Scientific) following the instructions provided by the supplier. The initial activity of the enzyme loaded in the gel was determined for calculating the cumulative lysozyme activity released over time. A control (free enzyme in solution incubated at 37 °C for different periods of time) was performed in triplicate to estimate the effect of incubation time and temperature on enzyme deactivation.

**Cell culture and maintenance.** hMSCs (passage number 6, PromoCell GmbH, Germany) were used to assess the ability of PSS/PA gels to support cell viability. Cells were first cultured in 75 cm² flask at density $3 \times 10^3$ cells cm$^{-2}$ and maintained in culture medium (DMEM, low glucose, GlutaMAX™ supplement, Gibco, Thermo Fisher, UK) supplemented with 10% FBS, 1% Pen/Strep and 0.1% bFGF (FGF-Basic (AA 10-155) recombinant human protein, Thermo Fisher Scientific, UK). Flasks were incubated at 37°C, 5% CO₂. Medium was refreshed twice a week and cells were trypsinized when reached 80% confluency.

**Cell encapsulation and viability in the hydrogels.** Cells were encapsulated in 1 MDa PSS/PA and 1 MDa PSS/PA-RGDS (10% PA-RGDS) hydrogels using $200 \times 10^3$ cells per 100 μl hydrogel by mixing half of the cells with the PSS solution and the other half with PA solution. 1 MDa PSS/PA-RGDS hydrogels were prepared as before where the PA solution contained 10% of PA-RGDS and 90% of standard PA at peptide final concentration of 2% (wt.%). After self-assembly (20 minutes), hydrogels with cells were incubated in culture medium (DMEM supplemented with 10% FBS and 1% Pen/Strep) at 37°C, 5% CO₂ for 1 and 3 days. Viability of the cells encapsulated in the hydrogels was analysed using LIVE/DEAD® Viability/Cytotoxicity Kit for mammalian cells (Invitrogen, UK). Briefly, each hydrogel sample was treated with 4 μM calcein
AM and 2 μM ethidium homodimer-1 and incubated for 15-20 minutes at room temperature. Hydrogels were imaged using confocal microscopy (Leica TCS SP2) and images were analysed in Image J for counting the number of live and dead cells. Three hydrogel replicates were used and compared to cell culture as monolayers (control). Statistical differences in viability data were obtained with GraphPad Prism 7 software (USA) using a two-way analysis of variance (ANOVA) with a Bonferroni’s multiple comparison test (*p < 0.05; **p < 0.01; ***p < 0.001; NS: not significant).

**Scanning electron microscopy of cell-encapsulated hydrogels.** The interaction of MSCs with hydrogel’s nanofibers was analysed by SEM. MSCs encapsulated in the hydrogels were prepared for SEM as above. The samples were coated with gold layer before observation.
Results and discussion

**Mechanical properties and stability of self-assembled PSS/PA hydrogels.** Peptide amphiphiles (PAs) designed by Stupp’s laboratory are known to self-assemble into cylindrical nanofibers maintained by strong β-sheet secondary structures. These PAs are charged molecules that upon mixing with electrolytes or polymers of opposite charge are able to form hydrogels, membranes or capsules. The self-assembly of PAs with high molecular weight (HMw) polysaccharides, such as hyaluronic acid (HA), alginate and \(\lambda\)-carrageenan results in the formation of solid membranes at liquid-liquid interface either in the form of a sac or planar membrane. However, the use of HMw synthetic polymers has not been fully exploited.

Following a similar approach, we combined the positively charged PA (Figure 1A), containing an alkyl chain (C\(_{16}\)), a β-sheet forming segment (V\(_3\)A\(_3\)) and three lysine residues (K\(_3\)) with negatively charged poly(sodium 4-styrenesulfonate) (PSS). Contrary to the HA/PA self-assembling systems, or with other polysaccharides, mixing PA with PSS of different Mw (70 kDa and 1 MDa) resulted in the formation of self-supporting opaque hydrogels (Figure 1A). It was shown previously that polyelectrolytes with different charge density interact differently with positively charged PAs affecting the micro- and nanostructure of the resulting membrane. PSS is a strong polyelectrolyte showing a zeta potential (ζ) of \(-49.62 \pm 2.55\) mV in water at pH 7 (Figure S2). Viscosity measurements of PSS solution (1 wt%, 1 MDa) revealed perfect Newtonian behaviour with constant viscosity in a wide shear range and low viscosity close to water (0.003 Pa·s) at 37 °C and shear rate of 70 s\(^{-1}\). In contrast, HA solutions of same concentration and similar molecular weight (1.5 MDa) showed higher viscosity (0.682 Pa·s) and shear-thinning behaviour. Because fluid flow is highly dependent on the viscosity, one should expect rapid mixing of PSS and PA solutions leading to PA self-assembly into a random network.
of nanofibers and formation of gel, without the creation of a diffusion barrier as seen in the HA/PA system. The low viscosity of PSS solutions (5 wt%, 126.7 kDa) was also observed in other studies with SAXS patterns showing characteristic behaviour of salt-free polyelectrolyte solutions and correlation lengths (calculated from $q_{\text{max}}$) between 8 and 70 nm depending on the PSS concentration (from 8.7 to 0.6 wt%, respectively).\textsuperscript{26} While the viscosity of polymer solutions depends mainly on the polymer molecular weight and concentration, intra- and inter-hydrogen bonding formed between polar hydroxyl groups also play a role in determining the final viscosity, and the extent of hydrogen bonding decreases with increasing temperature and addition of electrolytes into the solution. This is the case in HA molecules, where carboxylate, acetamido and hydroxyl groups enable the formation of intra hydrogen bonds and results in increased viscosity at neutral pH.\textsuperscript{27}

Hydrogels were formed by mixing the PA and PSS solutions at the same volume ratio. However, we have first optimized the hydrogel formulation by testing different PA and PSS concentrations followed by rheological analysis. The rheology data showed changes in the storage modulus with peptide and polymer concentration (Figure S3). However, after incubation in culture medium (DMEM) for 7 days, hydrogels showed different behaviours. Formulations with higher concentrations of PSS made the DMEM environment acidic (as detected by colour change of the medium) and the hydrogels with lower concentration of PSS were unstable. Therefore, formulations of similar concentration of peptide and PSS were used for subsequent experiments. PA solution of 2 wt% was combined with PSS solution of the same concentration and allowed to self-assemble for 10 minutes. PSS of different Mw (1 MDa and 70 kDa) were used to investigate the effect of the polymer length on the assembly and mechanical properties of the hydrogel.

Rheology measurements were carried out first by subjecting the hydrogels to an amplitude sweep
in a shear strain ranging from 0.01% to 10% at constant frequency of 1 Hz. In all tested hydrogels, the storage modulus exceeded the loss modulus, which is a typical behaviour of hydrogel systems and both types of hydrogel (1 MDa PSS/PA and 70 kDa PSS/PA) lost their linearity over strain of 1% (Figure 2-A,B). Following the linear viscoelastic region (LVER), strain of 0.1% was used in frequency sweep measurements.

Figure 2. Rheological characterization of PSS/PA hydrogels measured at 25 °C. Amplitude (A, B) and frequency (C, D) sweep graphs of 1 MDa PSS/PA (A, C) and 70 kDa PSS/PA (B, D); circles refer to storage modulus (G’) and triangles refer to loss modulus (G’’).

To measure the stiffness of the hydrogels, frequency sweeps were conducted at 0.1% strain over a range of frequencies. The storage modulus for both 1 MDa PSS/PA and 70 kDa PSS/PA
hydrogels is dominant over the loss modulus indicating an elastic rather viscous material.\textsuperscript{28,29} Moreover, the tan(\(\delta\)) (\(G''/G'\)) variation with frequency showed values below 1 (\(\tan(\delta) < 1\)), indicative of a gel system with \(G'\) always exceeding the \(G''\).\textsuperscript{29} \(\tan(\delta)\) is constant across a frequency range 1-10 Hz (\textbf{Figure S4}). A frequency of 20 Hz was reported for optimal osteoblast proliferation\textsuperscript{30}. Both 1 MDa PSS/PA and 70 kDa PSS/PA hydrogels show frequency-independent behavior at frequency range 1-10 Hz and they are considered as stiff hydrogels (30-40 kPa)\textsuperscript{31,32} (\textbf{Figure 2-C,D}). Different variations of PA hydrogels, formed in the presence of electrolytes, have a stiffness in the range of 20 Pa - 83 kPa depending on their molecular structure and concentrations\textsuperscript{28,33,34}. To increase the stiffness of self-assembling peptide gels, several approaches have been attempted by combining with either peptides or polymers of opposite charge).\textsuperscript{17,31} For example, different types of self-assembling peptides have been combined with sulfated polysaccharides, including heparin\textsuperscript{10,12,35}, chondroitin sulfate\textsuperscript{36} and fucoidan\textsuperscript{37}, to obtain gels with increased mechanical strength compared to gels crosslinked with multivalent ions. However, the storage modulus obtained did not exceed 50 kPa, whereas the mechanical properties of PSS/PA hydrogels can be in the range of 200 Pa – 200 kPa (\textbf{Figure S3}).

To investigate the stability of the PSS/PA hydrogels in aqueous environments of different ionic strength, they were incubated either in water or phosphate buffered saline (PBS) over a period of 28 days. The stability was assessed by monitoring the release of PSS into the medium. 1 MDa PSS forms more stable hydrogels compared to the lower Mw PSS. As seen in \textbf{Figure 3-A}, the 70 kDa PSS/PA hydrogels dis-integrated by day 7 in PBS. Moreover, the amount of PSS released to the medium was minimal for both hydrogels over 28 days incubation period (\textbf{Figure 3-B}).

The differences in stiffness and stability of the hydrogels can be explained through the PSS Mw (chain length) and the number of sulfonate groups available for interacting with the PA. The
higher stability of 1 MDa PSS/PA gels can be due to higher number of sulfonate groups (also confirmed by mapping analysis of the hydrogels (Figure S5) available for interaction with PA fibers leading to a robust intertwined network. However, in the lower Mw PSS/PA gels (70 kDa PSS/PA), the shorter chain length (hence less sulfonate groups) resulted in a less entangled network leading to decreased stability.

Figure 3. Stability of the hydrogels in water and PBS at 37 °C. (A) Comparison between 1 MDa PSS/PA hydrogel with 70 kDa PSS/PA (with disintegration by day 7 in PBS). (B) Graphs showing the PSS release from 1 MDa PSS/PA (red bars) and 70 kDa PSS/PA (green bars) hydrogels when incubated in water or PBS over 28 days.

Nanostructure and mineralization of the PSS/PA hydrogels. Yoreo et al. showed that a PSS matrix was able to induce the biomimetic nucleation of calcium carbonate minerals. Negatively charged groups in biomacromolecules, including carboxylic, phosphate and sulfate groups, play
a crucial role in the nucleation of calcium-based minerals.\textsuperscript{18,38,39} In a similar way, we hypothesize that the presence of sulfonate groups in PSS/PA hydrogels could induce the nucleation of calcium phosphate mineral as precursor of hydroxyapatite. In order to test this hypothesis, hydrogels were first incubated in a simulated body fluid (SBF) for a period of 14 days.\textsuperscript{20,21} After 14 days incubation, samples were analysed by SEM and EDX, which showed the formation of calcium phosphate minerals over the hydrogel surface (\textbf{Figure S6}). However, SBF is a highly saturated medium, which does not resemble a controlled biomimetic mineralization. Thus, the hydrogels were incubated in culture medium over a period of 21 days (3-weeks, minimum time period for osteogenic differentiation).\textsuperscript{40} Samples were incubated in four different culture media: DMEM without FBS (DMEM-FBS), DMEM with 10\% FBS (DMEM+FBS), osteogenic differentiation medium without FBS (ODM-FBS) and osteogenic differentiation medium with 10\% FBS (ODM+FBS) at 37 °C and 5\% CO\textsubscript{2} to replicate culture conditions. Mineral nucleation was detected by day 7 covering the hydrogels surface (\textbf{Figure 4}), but traces of calcium phosphate mineral were also observed in the hydrogel interior (\textbf{Figure S7}, SEM images of hydrogel cross-section). Minerals are spherical in morphology and deposited alongside the fibres; though it is also possible to get high concentrated areas of minerals (70 kDa PSS/PA, D21). The ratio of Ca/P changed with incubation time and is also different in the two hydrogel systems. In 1 MDa PSS/PA hydrogels, the Ca/P is close to hydroxyapatite (Ca\textsubscript{10}(PO\textsubscript{4})\textsubscript{6}(OH)\textsubscript{2}, 1.67) by day 7, whereas in 70 kDa PSS/PA it takes up to 21 days to get a Ca/P of 1.63 (\textbf{Figure 4}). PEDOT/PSS scaffolds without cells and incubated in proliferating medium (7 days) first, followed by 21 days in osteogenic medium, showed calcium deposition only after 28 days with no significant calcium accumulation by day 7.\textsuperscript{19} Calcium-phosphate nodules were seen and detected in 1 MDa PSS/PA gels after 7 days (\textbf{Figure 4}). The nanofiber structure of the PSS/PA
gels may provide a more biomimetic template for hydroxyapatite mineralization.

Figure 4. SEM images of mineralized hydrogels (1 MDa PSS/PA and 70 kDa PSS/PA) incubated in DMEM-FBS medium at different incubation periods.

The effect of serum and osteogenic inducers in the medium on the mineralization of the gels was also investigated and Figure 5 summarises the data from these studies. Osteogenic medium contains β-glycerophosphate, an organic phosphate that can be cleaved by alkaline phosphatase (ALP) enzyme secreted by osteoblast cells, into free phosphate ions necessary for hydroxyapatite mineralization. The time-dependent formation of PO$_3^{2-}$ by ALP is expected to prevent rapid and uncontrolled mineral precipitation in solution. Once phosphates are cleaved, they can be incorporated in the calcium-nucleated PSS/PA nanofibers. The Ca/P ratios of the mineral formed on 1 MDa PSS/PA hydrogels incubated in basal medium with and without FBS (DMEM-FBS
and DMEM+FBS) is close to tetra calcium phosphate (TTCP, Ca$_4$(PO$_4$)$_2$O) with higher calcium content compared to the osteogenic media (ODM-FBS and ODM+FBS), which are calcium-deficient hydroxyapatite. This indicates that minerals with different resorbability can be formed on these gels depending on the mineralizing medium. Furthermore, these studies were performed without cells, which are expected to play a role in the mineralization process, by secreting ALP, while here the release of phosphate depends on the ALP present in FBS.

**Figure 5.** Mineralization of hydrogels in four different media (DMEM-FBS, DMEM+FBS, ODM-FBS and ODM+FBS) after 21 days of incubation shown by SEM micrographs and EDX spectra; (A) 1 MDa PSS/PA, (B) 70 kDa PSS/PA.

The storage modulus of the gels after mineralization in the different media tended to increase,
but differences were not statistically significant for most of the samples tested (Figure S8). To further characterize the minerals formed on the hydrogels, samples were heated up to 700 °C to completely burn the organic phase (peptide and polymer components). The remaining inorganic component (mineral, ~ 4.5%, Figure S9) was analysed by FTIR (Figure S10). Several bands corresponding to amines in the peptide’s structure can be assigned in the spectrum of the non-mineralized hydrogel before heating, including amide I (1635–1670 cm⁻¹), amide II (1545 cm⁻¹) and amide III (1400, 1445 cm⁻¹). Moreover, the characteristic bands of SO₃⁻ are visible with a slight shift from 1187 to 1162 cm⁻¹ and from 1043 to 1033 cm⁻¹ corresponding to the asymmetric and symmetric stretching vibrations, respectively. The styrene ring of PSS overlapped with amide I band of PA hence giving an intense peak at 1613 cm⁻¹. In the heated samples, the bands corresponding to the peptide disappeared and the bands associated with minerals are more visible, such as HPO₄²⁻ (1090–1100 cm⁻¹), ν₃ PO₄³⁻ (1045, 603, and 570 cm⁻¹), and ν₁ PO₄³⁻ (960 cm⁻¹). However, the heating of samples above 600 °C can convert amorphous calcium phosphate into poorly crystallized apatite.

Based on the mechanical properties and mineralization capacity of the PSS/PA hydrogels, 1 MDa PSS/PA hydrogels were selected for further studies considering their greater stability and stiffness and faster mineralization capability (7 days).

**Protein loading and release.** Binding of GFs to the ECM takes place through different forms. One of the most prominent form is through electrostatic interactions with sulfated GAGs. Most GFs, such as basal fibroblast growth factor (bFGF, also known as fibroblast growth factor-2, FGF-2) and BMPs (bone morphogenetic proteins) are basic proteins having net positive charges. Considering the role of sulfate groups in GF binding, PSS/PA hydrogels could also
serve as depots to store basic proteins and control their release for cell activation.

Two model proteins of different charges and sizes were chosen to study the ability of the PSS/PA gels for protein retention and release. Bovine serum albumin (BSA) is negatively charged at physiological pH (pI = 5.4) and has a size of $M_w$: 67 kDa\textsuperscript{46}. Based on its negative charge, the protein was incorporated in the hydrogel by mixing it first with PSS solution (to avoid interaction with positive PA) and then assembled with PA. As positively charged protein, lysozyme (pI = 11.4) with a size of 14 kDa\textsuperscript{46} was used and first mixed with PA before self-assembly with PSS. Because self-assembly (gelation) occurs within 5 minutes at physiological conditions, the proteins can be entrapped inside the hydrogel without compromising their structure. To confirm the efficient incorporation of the proteins in the hydrogels, both proteins were labelled with fluorescent dyes to allow their visualization and distribution in the hydrogel by confocal microscopy and monitoring their release by fluorescence spectroscopy.

The possibility of loading multiple proteins in the PSS/PA gels was first investigated. Proteins were labelled with different fluorophores and their distribution in the hydrogel visualized by confocal microscopy (Figure 6). Green fluorescence (BSA-FITC) is observed in the form of nanofibers, indicating preferential interaction with PA, while the red fluorescence (lysozyme-rhodamine) appears mostly as clumps (potential interaction with PSS). EDX mapping of sulfur (Figure S5) shows a uniform distribution of sulfonic groups on the hydrogel.
To study the protein release, BSA and lysozyme were labelled with same fluorophore to eliminate the potential effect of fluorophore on the release pattern. In the first 24 hours of incubation, PSS/PA hydrogels released only 5% of BSA reaching 11% after 30 days (Figure 7-A,B). On the contrary, the amount of lysozyme released was higher, with 38% released after 30 days. The diffusion of proteins through hydrogels can be controlled through the physical (mesh size) and/or the chemical (affinity) characteristics of the hydrogel. Protein (e.g. BSA) release from hydrogels typically occurs within 1-3 weeks depending on the type of hydrogel and crosslinking density, even from sulfated HA hydrogels, indicating low protein retention. Both components of the PSS/PA gel are charged (Figure S2) providing binding sites for proteins through electrostatic interactions. Lysozyme is smaller in size compared to BSA and SEM images of PSS/PA hydrogels (Figure S5) show a very dense network of nanofibers. Considering similar affinity of the proteins for the gel components, release is expected to be controlled mainly by diffusion. The higher amount of lysozyme released compared to BSA is confirmed by the confocal microscopy images of the gels taken after the release experiment (Figure S11-A).
showing less fluorescence in the lysozyme-FITC loaded gels after 30 days. The slow release of BSA is further confirmed by the strong fluorescence exhibited by the gels loaded with BSA-FITC after the release study and indicates strong interactions with gel components, probably with PA nanofibers. The effect of electrostatic interactions between proteins and self-assembling peptide hydrogels of high charge density was shown to modulate their release patterns. Minimal protein release (7% over 28 days) was observed when unlabelled lactoferrin (77 kDa, pI = 8.4-9.0, positively charged at pH 7.4) was incorporated into peptide gels bearing a negative charge, while negatively charged or neutral proteins were completely released over 5 days. Appel and co-workers reported extended BSA release up to 160 days from neutral supramolecular polymer gels, compared to lysozyme (40 days). The hydrogels were formed through host-guest interactions of poly(vinyl alcohol) modified with viologen (first guest) and the macrocyclic host cucurbit[8]uril and hydroxyethyl cellulose functionalized with a naphthyl moiety (second guest). For this hydrogel system, protein release is mainly controlled by the size of the protein and hydrogel concentration that determines the mesh size. With respect to GFs, incorporation and controlled release of recombinant FGF-2 (Mw = 17.2 kDa, pI = 9.7) has been described using heparin-containing carriers. For example, self-assembling heparin/PA (2/3 wt.%) gels showed the ability to retard the release of FGF-2 from 100% in control gels (without heparin) to 57.1% after 10 days and this delayed release has been reported for other self-assembling heparin/peptide and heparin-containing hydrogels. Considering that both lysozyme and FGF-2 are basic proteins with similar Mw, one can assume similar release of FGF-2 from PSS/PA gels. Protein release results reveal the potential PSS/PA hydrogels for the continuous delivery of GFs or other proteins (e.g. antibodies) for considerable periods of time (long-acting release) to achieve robust therapies.
Figure 7. Cumulative release (mean ± SD) of BSA-FITC and lysozyme-FITC loaded in 1 MDa PSS/PA hydrogels at early time points (A, 24 h) and for extended periods of time (B, 30 days). Cumulative lysozyme activity release (mean ± SD) from the hydrogels within the first 24 h (C) and over a month (D).

The ability of the PSS/PA gels to maintain the bioactivity of entrapped proteins was assessed by measuring the activity of lysozyme released from the gel (Figure 7-C, D). Increasing enzyme activity is observed as function of time, reaching 20% of initial activity after 30 days. Considering similar amounts of enzyme released (as measured for the lysozyme-FITC), a 50% loss in the enzyme activity is estimated (Figure S11-B). However, when lysozyme is incubated in buffer only, it loses 80% of its initial activity after 14 days, similar to what was found by Appel,51 and indicates the protective effect of PSS/PA gel against enzyme deactivation.

Cell Viability. Cytotoxicity of the hydrogels was assessed with human bone marrow-derived
mesenchymal stem cells (hMSCs). hMSCs were encapsulated in the hydrogels without or with the cell-adhesive sequence RGDS: PSS/PA hydrogel and PSS/PA-RGDS hydrogel (PSS/PA hydrogel containing 10% of RGDS-containing PA, Figure S1-D). Incorporation of 10% of PA-RGDS in the hydrogel formulation is aimed at promoting cell adhesion. Characterization of the mechanical properties of the resulting hydrogels did not show significant differences compared to plain gels (without 10% of PA-RGDS, Figure S12).

Figure 8. Viability of hMSCs encapsulated in 1 MDa PSS/PA hydrogels, with or without RGDS, for 1 and 3 days. (A) Confocal images of calcein-stained live cells and ethidium homodimer-stained dead cells; (B) Cell viability in the hydrogels compared to 2D culture. (C) Cell density in the hydrogels remained constant from day 1 to day 3. (No significant difference between the groups)

Figure 8-A illustrates the distribution and morphology of the cells inside the hydrogels, as well as the overall viability, with live cells stained green and dead stained red (hydrogels showed auto-fluorescent in the red channel, hence the red background in the images). The viability increased slightly in both hydrogels over 3 days of culture compared to 2D culture (Figure 8-B),
but statistical analysis showed no significant difference between the groups. This result confirmed that the gelation time (without DMEM) did not affect the cell viability and the hydrogel fabrication method is compatible for the encapsulation of cells (Figure 8-C). Cells appear round which is typically seen when encapsulated in hydrogels. The interaction of cells with the hydrogel nanofibers was also analysed by SEM (Figure 9). Cells are initially entrapped inside the dense nanofibrous structure of the gels, exhibiting round-shaped morphology even after 3 days of culture. However, in both PSS/PA and PSS/PA-RGDS gels, few cells were found with spread morphology. Similar findings were previously observed when culturing human dermal fibroblasts within self-assembled peptide capsules. By changing the nanofiber density, fibroblasts could recover their typical extended morphology. Matrix stiffness increases with matrix density and PSS/PA hydrogels with lower stiffness can be formed (Figure S3) and used to test the effect on cell morphology and differentiation. Taken together, the positive cell viability results and SEM analysis suggest that PSS/PA hydrogel can provide a biomimetic 3D environment for growing stem cells and studying their differentiation into different lineages.
Figure 9. SEM micrographs of 1 MDa PSS/PA hydrogels showing cells embedded in the dense nanofiber network at day 1 and their spreading at day 3.
Conclusions

Self-assembly of a known polyelectrolyte (PSS) with a PA of opposite charge in physiological conditions resulted in self-supporting stable hydrogels. The obtained hydrogels have nanofibrous structure and mechanical properties of stiff materials, which can be tuned by varying the concentration of individual components. The negative sulfonate groups of PSS provided nucleation points for calcium phosphate deposition. Depending on the mineralization medium used, different types of calcium phosphate mineral were formed. These hydrogels also showed the ability to retain and control the release of charged proteins, while preventing their rapid deactivation. Although preliminary, the positive results from cell culture studies suggest the potential of the PSS/PA hydrogels as 3D environments for cell differentiation either in vitro or in vivo.
Supporting Information Available

The following files are available free of charge.

Figure S1: Characterization of peptide amphiphiles used this study
Figure S2: Zeta (ζ) potential of PA and PSS solutions
Figure S3: Storage modulus (G′) PSS/PA hydrogels in function of their composition
Figure S4: Ratio of G′′ to G′ (tan(δ)) of PSS/PA hydrogels
Figure S5: SEM/EDX analysis of PSS/PA hydrogels
Figure S6: SEM/EDX of SBF-incubated hydrogels
Figure S7: SEM images of hydrogel cross-sections after incubation in DMEM-FBS
Figure S8: Storage modulus (G′) of PSS/PA hydrogels after mineralization
Figure S9: TGA analysis of PSS/PA hydrogels after incubation in mineralization medium
Figure S10: FTIR spectra of the hydrogels before and after burning
Figure S11: Confocal microscopy images of PSS/PA hydrogels loaded with labelled proteins before and after the release experiments; comparison between protein release and activity for lysozyme; decay of lysozyme activity over time at 37 °C when free in solution.
Figure S12: Mechanical characterization of 1 MDa PSS/PA-RGDS hydrogel

ACKNOWLEDGMENT

We thank Mr Lazar Kirov, an undergraduate student at the School of Engineering and Materials Science for measuring the viscosity of PSS and HA solutions. We thank the financial support from The Royal Society (Research Grant RSG\R1\180005).

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Supramolecular nano-fibrous peptide/polymer hydrogels for the multiplexing of bioactive signals

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