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Regular Article

Cell guidance on peptide micropatterned silk fibroin scaffolds

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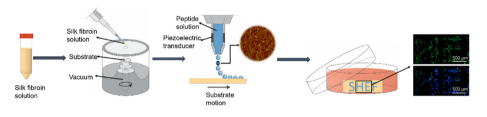
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ABSTRACT

Guiding neuronal cell growth is desirable for neural tissue engineering but is very challenging. In this work, a self-assembling ultra-short surfactant-like peptide I₃K which possesses positively charged lysine head groups, and hydrophobic isoleucine tails, was chosen to investigate its potential for guiding neuronal cell growth. The peptides were able to self-assemble into nanofibrous structures and interact strongly with silk fibroin (SF) scaffolds, providing a niche for neural cell attachment and proliferation. SF is an excellent biomaterial for tissue engineering. However neuronal cells, such as rat PC12 cells, showed poor attachment on pure regenerated SF (RSF) scaffold surfaces. Patterning of I₃K peptide nanofibers on RSF surfaces significantly improved cellular attachment, cellular density, as well as morphology of PC12 cells. The live / dead assay confirmed that RSF and I₃K have negligible cytotoxicity against PC12 cells. Atomic force microscopy (AFM) was used to image the topography and neurite formation of PC12 cells, where results revealed that self-assembled I₃K nanofibers can support the formation of PC12 cell neurites. Immunolabelling also demonstrated that coating of I₃K nanofibers onto the RSF surfaces not only increased the percentage of cells bearing neurites but also increased the average maximum neurite length. Therefore, the peptide I₃K could be used as an alternative to poly-L-lysine for cell culture and tissue engineering applications. As micro-patterning of neural cells to guide neurite growth is important for developing nerve tissue engineering scaffolds, inkjet printing was used to pattern self-assembled I₃K peptide nanofibers on RSF surfaces for directional control of PC12 cell growth. The results demonstrated that inkjet-printed peptide micro-patterns can effectively guide the cell alignment and organization on RSF scaffold surfaces, providing great potential for nerve regeneration applications.

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1. Introduction

The human nervous system is composed of the central nervous system (CNS) and the peripheral nervous system (PNS), which can be easily impaired by injuries, such as trauma and car accidents, as well as diseases, including Alzheimer's disease, Parkinson's disease, strokes and brain tumours.[1–3] Regeneration of both damaged CNS and PNS is challenging in tissue engineering.[3–5] It is therefore vitally important to develop well-defined functional scaffolds for nerve tissue regeneration to help guide neural cell attachment, alignment, spreading and proliferation.[6–8]

In addition, alignment and interconnection of neuronal cells in vitro allows the mimicking of neuronal architectures in vivo and help in understanding the underlying mechanisms needed to promote and accelerate regeneration of damaged neural tissue. Micro-patterning technology, which has already attracted significant attention, can enable the geometric control of neuronal cell alignment.[9–12] Lithography, including ultraviolet lithography (UVL), soft lithography (SL) and electron-beam lithography (EBL), is a traditional technology for micro-patterning proteins onto substrates.[13,14] Compared to UVL and EBL, SL is a convenient technique, [15] which has been widely used to micro-pattern neuronal cells.[9,10,12] For example, micro-patterned polydimethylsiloxane (PDMS) has been shown to enhance the attachment, alignment, spreading, proliferation, neurite formation and elongation of neuronal cells.[12] However, SL needs to be operated in a highstandard clean room, and samples can be easily contaminated during fabrication. [16,17] Inkjet printing, on the other hand, is a costeffective and flexible micro-patterning technique which is capable of patterning complex geometries at high precision.[13,14] Moreover, as inkjet printing is a non-contact technique, crosscontamination of the final product is significantly reduced. Therefore, it was used as a micro-patterning technique to pattern selfassembled peptide nanofibers on regenerated silk fibroin (RSF) surface to guide the growth of neuronal PC12 cells in this study.

Silk fibroin (SF), extracted from Bombyx mori (B. mori), has received significant attention due to its biocompatibility, tuneable biodegradability, low immunogenicity and excellent mechanical properties.[18,19] RSF possesses tuneable rheological properties and can be used to fabricate different types of scaffolds, such as hydrogels, [20] films [21] and sponges. [22] These scaffolds have been successfully applied in a variety of tissue engineering applications, such as skin, [23] vascular [24] and musculoskeletal tissue engineering to match the different properties of autologous tissues. [25] However, B. mori silk lacks the cell adhesive components. such as arginine-glycine-aspartic acid (RGD) sequence, which promotes cell attachment. [26-28] Therefore, without the addition of cell adhesive molecules, such as poly-L-lysine (PLL), and extracellular matrix (ECM) components, silk scaffolds normally have poor cell attachment, [29,30] which is particularly significant for neuronal cells.[31]

Cell adhesive molecules have been successfully applied as scaffolding materials in nerve tissue engineering.[32] During the last two decades, peptide sequences such as RGD, YIGSR and IKVAV have been used to promote neuronal cell attachment, proliferation, and neurite outgrowth.[33,34] Self-assembled peptide nanofibers (such as RADA16, EAK16) are novel biomaterials that can be fabricated through bottom-up approach and have the potential to be used as scaffold materials for tissue engineering.[35,36] Surfactant-like peptide AC-I₃K-NH₂ (I₃K) has an acetyl group on its N terminal and its C terminal was blocked by an amine group. Three hydrophobic isoleucine (IIe or 1) and one hydrophilic lysine residue (Lys or K) causes the peptide to possess the surfactant feature and promote the self-assembly of I₃K into long and uniform nanofibers in aqueous solutions with the K residues on the outside of the nanofibers.[37,38] Positively charged PLL has been shown to promote neuronal cell attachment.[39] Therefore, it is anticipated that the self-assembled I_3K peptide nanofibers also have great potential as a cell adhesive matrix for nerve tissue regeneration.

In this study, RSF/I₃K peptide scaffolds were fabricated to guide neuronal cell attachment. The coating and patterning of peptide nanofibers were achieved through spin coating and inkjet printing, respectively. The glass/silicon wafer substrates were coated with a layer of negatively charged RSF[40] before the coating, or printing, of peptide nanofiber solutions. The cationic peptide nanofibers adhered onto the RSF surfaces through charge-charge interactions. Rat pheochromocytoma (PC12) cells were cultured onto RSF/I₃K scaffolds to investigate the effect of the I₃K peptide nanofibers on cell attachment, proliferation and viability. Atomic force microscopy (AFM) was used to further analyse cell morphology, height and footprint on the RSF/I₃K scaffold surfaces. [41] The results indicated that I₃K peptide nanofibers promoted cell attachment, proliferation and neurite outgrowth of PC12 cells. Immunolabelling also demonstrated that coating of I₃K nanofibers onto RSF surfaces, not only increased the percentage of cells bearing neurites, but also increased the average maximum neurite length. Cells attached along the inkjet-printed peptide nanofiber patterns, demonstrating that inkjet printing is a promising technique to pattern scaffolds for geometrical guidance of neuronal cell growth as well as investigation of neurite development and formation *in vitro*.[42]

2. Experimental section

2.1. Materials

The peptide Ac-I₃K-NH₂ (purity > 98%, w/w) was purchased from GL Biochem Ltd. (Shanghai, China). *B. mori* silkworm cocoons were supplied by Biological Science Research Centre, Southwest University, China. PC12 Adh (CRL-1721.1) cell line was obtained from the American Type Culture Collection (ATCC). Silicon wafers were purchased from Compact Technology Ltd, UK. Unless otherwise specified, chemicals and regents (analytical grades) were purchased from Sigma Aldrich, UK.

2.2. Preparation of regenerated silk fibroin

B. mori silkworm cocoons were cut into small pieces (~1 cm²) and degummed in 0.02 M Na₂CO₃ solution at 100 °C for 1.5 h under stirring. Degummed silk was rinsed three times with deionized water (DI water) to ensure the removal of sericin. After which the degummed silk fibres were dried for 2 days in a drying oven at 60 °C and dissolved under stirring in Ajisawa's reagent (CaCl₂/ ethanol/deionized water = 1:2:8 M ratio) at 80 °C for 1.5 h. The resulting viscous solution was dialyzed against DI water until a conductivity below 10 μ S of the dialysis fluid was reached. The resulting RSF solution was then centrifuged for 10 min at 10,000 rpm to remove any particulates. The RSF concentration was determined by weighing dried RSF peptide residues on microscope slides. Stock RSF solutions of 5 mg/mL and 40 mg/mL were made by diluting with DI water and stored at 4 °C prior to use.

2.3. Preparation of RSF / peptide samples

I₃K peptides were dissolved in 20 mM HEPES buffer (pH 6.0) at 5 mg/mL and incubated for 7 days under ambient conditions for self-assembly. The sample was then diluted with 20 mM HEPES buffer (pH 6.0) to 4, 3, 2, 1 mg/mL prior to use. RSF/peptide bilayer scaffolds were made by spin coating (Laurell Technologies Corporation, USA) onto 1 cm² microscope cover glasses or silicon wafers. The first layer of RSF (30 μ L, 8,000 rpm, 25 s) was coated followed by fixing using 95% wt/vol ethanol (20 μ L, 4,000 rpm, 25 s), to convert the RSF layer from soluble random coil structure (silk I) to insoluble β -sheet structure (silk II).[43] The second layer, i.e. the positively charged peptide (30 μ L, 8,000 rpm, 25 s) was coated and adhered onto the negatively charged RSF substrate via charge interaction. The solution concentration ratios of RSF/peptide were 5:0; 40:0; 40:1; 40:2; 40:3; 40:4; 40:5 and 0:5, respectively.

2.4. Atomic force microscopy

AFM measurement (Bruker Dimension Icon, Bruker Corporation, USA) was performed in tapping model with SCANASYST-AIR probes at room temperature. To image peptide nanostructures, peptide solution was dropped onto freshly cleaved mica and dried under gentle air flow. AFM was also used to characterise the topography of the RSF/I₃K scaffolds and attached cells. PC12 neuronal cells were fixed with 3.7% paraformaldehyde (PFA) in phosphate buffered saline (PBS) for 45 min at room temperature. The PBS solution was then removed carefully with a pipette and the samples were washed with DI water gently to avoid crystallisation of PBS buffer salts. The samples were then left to dry at room temperature for 1 min prior to AFM characterisation. Images were analysed by NanoScope Analysis software (Version 1.5).

2.5. Inkjet printing

All glass slide surfaces were cleaned with 5% Decon90 solution and rinsed with plenty of DI water before printing: 1 layer of RSF solution (40 mg/mL) was coated via spin coating on the glass slides (30 μ L per cm², 8000 rpm, 25 s) followed by 95% wt/vol ethanol solution (20 μ L, 4000 rpm, 25 s) via spin coating. A Jetlab 4xL (MicroFab Inc., Texas, US) equipped with a piezoelectric drop-ondemand (DoD) jetting device (60 μ m nozzle diameter) was used for the printing of the I₃K peptide nanofiber ink. The actuation voltage and frequency used were 90 V and 300 Hz, respectively. The distance between the nozzle tip and the substrate was approximately 10 mm. To investigate the effect of printed I₃K peptide nanofiber pattern on the growth of PC12 cells, 1 layer of the pattern "SHEF" was printed.

2.6. Culture of PC12 neuronal cells on scaffolds

PC12 neuronal cells were cultured in Dulbecco's Modified Eagle Medium (DMEM) with high glucose and supplemented with 10% foetal calf serum (FCS), 1% penicillin/streptomycin,1% glutamine and 0.5% fungizone in an incubator at 37 °C under 5% (v/v) CO₂. The medium was replaced every 3 days. RSF/I₃K scaffolds were sterilized under ultraviolet light for 30 min before being washed in PBS for three times, and then placed in 12 well plates under metal rings (to secure the samples). Confluent cells were detached with 0.25% (w/v) trypsin-EDTA (ethylenediaminetetraacetic acid) and then seeded onto scaffolds' surface at 10,000 cells/cm² through the holes of metal rings. These samples were cultured in DMEM medium containing 10% FCS for 6 days.

2.7. Cell adhesion assay

Following incubation, neuronal cells were fixed in 3.7% PFA for 45 min at room temperature followed by washing twice with PBS and incubated for a further 45 min with 0.1% Triton X-100. Finally, the cells were washed twice with PBS and stained with FITC-phalloidin to visualize actin filaments and 4,6-diamidino-2-phenylindole dihydrochloride (DAPI) to visualize the nuclei. The samples were then imaged using a fluorescence microscope (Nikon Eclipse LV100).

2.8. Resazurin assay

Metabolic activity of PC12 cells was assessed after 24, 72, and 144 h in culture. Culture medium was removed, and samples were cultured in a 100 μ M resazurin salt in PBS, and assay dependent culture media for 4 h at 37 °C and 5% CO₂. Triplicates of 100 μ L, of reduced formazan product, were then transferred to a black 96 well plate and the fluorescence was read in a FLx800 fluorescence plate reader (Biotek Instruments Inc.) at 540/635 nm. Background fluorescence readings were measured and subtracted from results.

2.9. Live and dead assay

A live / dead assay was carried out by exchanging the medium with serum-free medium containing 0.001% (v/v) Syto-9TM (Invitrogen) and 0.0015% (v/v) propidium iodide (PI) and then incubated for 30 min in an incubator at 37 °C under 5% (v/v) CO₂. The samples were then imaged using an upright Zeiss LSM 510 confocal microscope. An argon ion laser was used to visualise live cells stained with Syto-9 TM (λ_{ex} = 494 nm / λ_{em} = 515 nm) and a helium-neon laser for dead cells stained with PI (λ_{ex} = 536 nm / λ_{em} = 617 nm). ImageJ software (National Institutes of Health, USA) was used to count the number of live and dead cells for several images of 2500 µm² sample areas randomly and averaged. Microscope images were converted to grayscale 8-bit images and then converted to a binary image via selecting the best threshold to generate a high contrast image, cell number was then counted via the 'analyse particles' algorithm in ImageJ.

2.10. Immunostaining of the neurites assay

Neuronal cell differentiation, on samples, was assessed by measuring the lengths of neurites extending from cells. PC12 cells were washed with PBS before cells were fixed with 3.7% (v/v) PFA for 20 min at room temperature. Following a PBS wash, cells were permeabilized with 0.1% Triton X-100 for 20 min, at room temperature and unreactive binding sites were blocked with 3% bovine serum albumin (BSA) in PBS for 30 min. PC12 cells were incubated with a mouse anti- β III-tubulin (neurite marker) antibody (1:250 dilution from Promega, Chilworth, United Kingdom) diluted in 1% BSA in PBS and incubated at 4 °C for 24 h. After a PBS wash, PC12 cells were labelled with Texas Red-conjugated anti-mouse IgG antibody (1:200 dilution in 1% BSA from Vector Labs, Burlingame, USA) in 1% BSA, for 90 min at room temperature. Samples were imaged with an upright Zeiss LSM 510 confocal microscope, using a helium-neon laser (543 nm) for Texas Red excitation (λ_{ex} = 589 nm / λ_{em} = 615 nm). Images were analysed, and neurites were measured using the ruler tool on ImageJ software.

2.11. Statistical analysis

GraphPad Prism V.6 software was used to analyse data quantitatively. One-way or two-way analysis of variance (ANOVA) with multiple comparisons was used for all multiple group experiments, and equality of variances was confirmed by Tukey's multiple comparisons test. P values < 0.05 were deemed significant. Values in graphs are presented as mean ± one standard deviation.

3. Results and discussion

3.1. Attachment of PC12 neuronal cells on SF films

RSF scaffolds have been extensively used in tissue engineering applications.[23] However, pure RSF lacks cell recognized molecules, and therefore, it normally has poor cell attachment.[26–28]

To investigate the attachment of PC12 neuronal cells on the RSF scaffold surfaces, RSF solutions, at different concentrations, were spin coated onto clean cover glasses or silicon wafers. AFM images (Fig. 1A) demonstrated different surface topographies, and consequently cell attachment. The RSF coated glass surfaces (both 5 and 40 mg/mL) showed smooth coated layers (Fig. 1A (a-b)) with roughness's at the nanoscale $(0.78 \pm 0.02 \text{ nm} \text{ and } 0.93 \pm 0.01 \text{ nm})$. The RSF scaffolds investigated here showed poor performance in cell attachment and spreading using PC12 neuronal cells (Fig. 1A (d-e)). A significant decrease in cell density on the surfaces (Fig. 1B) was observed using RSF coated glasses both at 5 mg/mL (9600 ± 700 cells/cm²) and 40 mg/mL (900 ± 600 cells/cm²) as substrates compared to clean glass (21900 \pm 1600 cells/cm²), which demonstrates that PC12 cells have a low binding efficiency to RSF coated substrates. These results indicate that RSF coatings, in particular at high concentrations, can be used as cell-repellent surfaces for PC12 cells. When combined with cell adhesive moieties, surfaces could be used to pattern PC12 cells, to guild the growth of the cells.

To enhance cell attachment, composite RSF scaffolds have been widely used to facilitate the cell affinity.[25] RSF scaffolds are normally combined with cell recognized molecules such as collagen, gelatin and PLL to enhance cell attachment. For example, gelatin was cross-linked with RSF scaffolds for the repair of cartilage injury in vitro and in vivo.[44] The scaffolds not only provided a mechanical protection before neocartilage formation, but also a suitable 3D microenvironment for BMSC (endogenic bone marrow stem cells) proliferation, differentiation, and ECM production. RSF scaffolds have also been modified with RGD sequences through the side groups of aspartic and glutamic acids to improve the cell adhesion.[25] While the previous methods mostly involved chemical modification of the SF material through covalently bonding, here we use the electrostatic interaction of self-assembled cationic peptide nanofibers to enhanced the cell attachment onto the SF scaffolds.

3.2. Characterization of RSF/I₃K scaffolds

The peptide AC-I₃K-NH₂ (in short I₃K) was able to self-assemble into long and uniform nanofibers after incubation as previously reported.[37,38] According to previous studies[37], the selfassembly process of I₃K is a dynamic process via non-covalent interactions. Upon complete dissolvement of I₃K molecules, small I₃K fragments form interdigitated bilayers with hydrophobic iso-

leucine residues kept in the interior, and charged lysine located on the surface of bilayers. Small I₃K fragments then assemble into short stacks through hydrophobic interactions and hydrogen bonding. Subsequently, based on molecular chirality and surface curving, these stacks tend to grow into twisted fibres. Further growth of which leads to the formation of long and uniform nanofibers. The width of the formed nanofibers was around 50 nm (Figure S1) while the length of the nanofibers can reach up to 10 µm. Persistence length is a characteristic length scale that has been used to determine the conformation of a uniform chain length.[45] Cox et al.[46] recently measured the contour length (the distance between two ends of I₃K fibres) by AFM and stochastic reconstruction microscopy and then used this value to calculate persistence length of I₃K fibres. Their results indicated that self-assembled I₃K fibres have an average contour length of around 6 µm and persistence length of 10.1 ± 1.2 µm.

AFM was further used to characterise how concentrations of I₃K affect the surface topography of the I₃K coated RSF scaffolds. As shown in Fig. 2(a), only a few self-assembled nanofibers were observed on the 1 mg/mL I₃K coated RSF scaffolds. By increasing the concentration of I₃K, the number of nanofibers increased gradually forming a near full coverage at 3 mg/mL (Fig. 2(b-c)). Further increasing the concentration of I₃K resulted in more stacked selfassembled nanofibers, which can potentially lead to cell detachment during cell culture. No significant difference of surface topography between the multi-material RSF/I₃K scaffold (at concentration ratio 40:5) and I₃K-only scaffold (5 mg/mL coated glass) could be detected. However, I₃K-only scaffolds assembled on blank glass could easily be washed off when immersed in the aqueous solutions due to low adhesion properties to the glass surface (Figures S2 and S3), hence making it ineffective for cell culture applications. Therefore, the multi-material combination of I_3K and RSF generates a structurally stable scaffold that can easily withstand normal cell culture procedures and thus provides an excellent way to generate peptide-based scaffolds for cell culture applications. The strong adhesion between RSF (negatively charged) and I₃K (positively charged) is a result of the strong electrostatic charge-charge interactions.[37.38]

3.3. Neuronal cell attachment and morphology on RSF/I₃K scaffolds

To investigate PC12 neuronal cell attachment and viability on different RSF/I_3K scaffold surfaces, a live / dead assay was carried

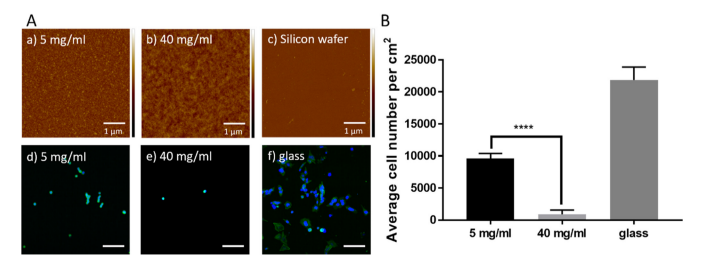


Fig. 1. A, AFM topographical images of RSF coated scaffolds on Si-wafers at (**a**) 5 mg/mL; (**b**) 40 mg/mL and (**c**) bare silicon wafer control, Z scale height = 30 nm. Fluorescence images of PC12 cells attached to (**d**) 5 mg/mL and (**e**) 40 mg/mL RSF coated surfaces and (**f**) glass control. (Blue: DAPI staining for nucleus. Green: FITC-phalloidin staining for F-actin.) Scale bar = 100 μ m. **B**, Average cell numbers (per cm²) attached to the RSF coated surfaces at RSF concentrations of 0 (glass control), 5 and 40 mg/mL. n \geq 3, **** p < 0.0001. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

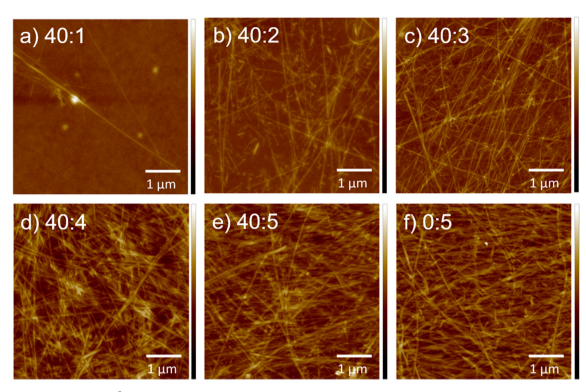


Fig. 2. AFM topographical images (25 μm²) showing 1-layer RSF/l₃K scaffolds coated at different concentration ratios: (**a**) 40:1, (**b**) 40:2, (**c**) 40:3, (**d**) 40:4, (**e**) 40:5, (**f**) 0:5. The Z scale (height) for all images is 120 nm.

out as shown in Figure S4. RSF-only scaffolds showed very low cell adhesion due to a lack of cell recognizable groups such as RGD. Furthermore, they are negatively charged, which has been known to have negative effects on cell attachment.[47,48] Therefore, by adding positively charged I₃K peptide nanofibers onto the negatively charged RSF substrates, the positively charged lysine residues deposited on the scaffold surface promoted PC12 cell binding. [49] PC12 cells showed poor attachment on I₃K-only and RSF/I₃K scaffolds at low I₃K concentrations (<3 mg/mL) compared with RSF/I₃K scaffolds with high I₃K concentrations (3-5 mg/mL). This is most likely because I₃K nanofibers are easily washed off without RSF base during cell culture as shown in the AFM images in Figures S2 and S3, and the RSF/I₃K scaffolds with low concentrations of I₃K provide insufficient anchoring points for cell attachment. A significant difference in cell density was observed between RSF-only and RSF/I₃K (3–5 mg/mL) scaffolds, indicating the promotion of PC12 neuronal cell attachment on RSF/I₃K scaffolds through the addition of positively charged I₃K nanofibers. However, with an I₃K concentration of 5 mg/mL, the number of cells decreased slightly, where no cell attachment in some areas was observed. This was possibly due to some detachment of stacked I₃K nanofibers occurring at this and higher concentrations, as previously discussed in section 3.2. The stacked I₃K nanofibers promoted excessive cell attachment and cells grew quickly becoming over confluent during the incubation time, thus detaching from the scaffold surface. Additionally, the live / dead assay indicated that RSF/I₃K scaffolds have excellent biocompatibility. The results indicated that I₃K is a promising candidate which can be used as a functional scaffold material similar to other peptides (such as PLL[49] or gelatin[4]) previously reported for tissue engineering.

The investigation of cell morphology attached on different ratios of RSF/I₃K scaffolds was carried out via AFM (Fig. **3A and S5**). As previously noted, cells spread out well on I₃K-only coated surfaces but showed patchy attachment due to the peptide being

washed off during cell culture. Cells showed an excellent spreading on I₃K-only coated surfaces (Fig. 3**A**(**a**)) whereas cells barely spread out their terminals on RSF-only coated surfaces (Fig. 3**A**(**b**)). Cells started to attach on RSF/I₃K coated surfaces at concentration ratio 40:1 (Fig. 3**A**(**c**)). By increasing the concentration of I₃K from 2 to 4 mg/mL deposited onto RSF (40 mg/mL) coated surfaces, cellular spreading improved drastically (Fig. 3**A**(**d**-**g**)), indicating a peptide concentration dependence of cell spreading. Additionally, FITCphalloidin and DAPI was used to stain actin filaments and cell nuclei respectively to further characterize the morphology of cells adhered on RSF/I₃K scaffolds (**Figures S6**). Cells on RSF/I₃K coated substrates also showed excellent spreading and flattening with this being enhanced as the concentration of coated I₃K increased. These results are consistent with previous AFM results shown in Fig. 3**A**.

The first step of the cell adhesion process is the cell-polymer interactions, which is essential for cell communication, regulation, tissue development and maintenance. [50,51] Cell-polymer interactions can be divided into three types, i.e. non-adhesion, passive adhesion and active adhesion. [50] The interactions between cells and RSF surfaces corresponds to passive adhesion, which means cells attach easily but can also easily detach from surfaces.[52] Cells attached on RSF/I₃K scaffolds on the other hand undergo active adhesion, in which cells spontaneously adhere onto the surface and the adhesion is tight, therefore, it is difficult for cells to detach.[53] Additionally, the positively charged lysine residues in I₃K activates cell changing morphology and causes spreading for attachment-dependent phenotypes. Furthermore, there are three phases that can describe the process of cell adhesion onto cellactive polymer surfaces. Cells adhere onto the RSF-only scaffolds via complex physicochemical interactions including Van-der-Waals, coulombic and hydrophobic forces, known as Phase I cell attachment.[54] The action of the cells starting to spread and become flattened on the RSF/I₃K scaffold surface (Fig. 3A(c-d)), due to integrin binding, is known as Phase II. Full spreading and

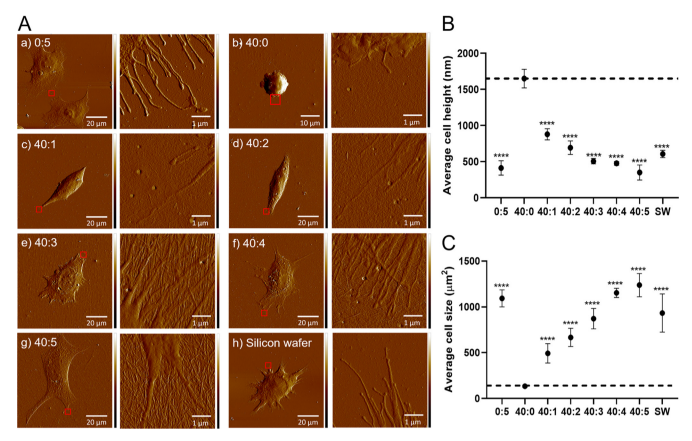


Fig. 3. A, AFM Peak Force Error images of PC12 neuronal cells attached on a series of RSF and I_3K coated surfaces. In addition, enlarged AFM images of red box areas are shown in right of each image. The concentration ratios of RSF and I_3K were: (**a**) 0:5, (**b**) 40:0, (**c**) 40:1, (**d**) 40:2, (**e**) 40:3, (**f**) 40:4, (**g**) 40:5 and (**h**) silicon wafer control. The force setpoint constant is 30 nN for images of cells and 5 nN for enlarged areas. **B** and **C** indicate average cell heights and sizes in all scenarios. ($n \ge 3$; ^{***} p < 0.0001; ^{***} p < 0.001; and ^{*}p < 0.05). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

formation of focal adhesions for the cytoskeleton of PC12 neuronal cells on the RSF/I₃K (Fig. **3A(e-g**)) is regarded as Phase III. Therefore, RSF/I₃K scaffolds, where I₃K can be considered a celladhesive polymer, are able to activate the cell adhesion process inducing spreading and flattening. Increasing the concentration of I₃K resulted in improved and expedited cell adhesive properties.

Further AFM analysis revealed the average cell height (Fig. 3B) on RSF-only scaffolds was 1650 ± 110 nm. In the case of I₃K-only scaffolds, an average cell height of 410 ± 80 nm was measured. The average height of cells on RSF/I₃K peptide scaffolds decreased with increasing I_3K concentrations (1 to 5 mg/mL) from 880 ± 60 nm to 350 ± 90 nm. For the blank silicon wafer, the average cell height was 600 ± 40 nm, which was similar to (but slight lower than) that of the RSF/I₃K (ratio 40:2) samples. The average size of attached cells (Fig. 3C) on RSF-only scaffolds was found to be 140 \pm 10 μ m². A gradual size increase (up to 9-fold at 5 mg/ mL I₃K) was observed as I₃K concentration was increased. Both average cell height and size on the silicon wafer substrates were similar to the RSF/I₃K scaffold at a concentration ratio of 40:3. The results indicated that RSF/I₃K scaffolds with I₃K concentrations above 3 mg/mL are ideal for PC12 cell adhesion and spreading, outperforming the cell behaviour on RSF-only scaffolds. The red boxes in Fig. 3A also indicate areas of interest that were enlarged to further investigate the cell morphology. The AFM images show neurites of PC12 cells adhering to the RSF-only or RSF/I₃K coated surfaces (Fig. 3A(b-g)). In contrast to I₃K-only scaffolds, Fig. 3A (h) shows the neurites of PC12 cells adhering on exposed bare silicon wafer substrate. The results were similar to those reported by Gupta et al.[55] who indicated that neural cells can adhere and differentiate on chitosan-based scaffolds. We conclude that I₃K can promote PC12 cell attachment, spreading and neurite formation.

3.4. Differences between RSF/I₃K scaffolds and collagen scaffolds on the function of PC12 cells

Collagen is one of the basic components of the ECM that can provide a natural environment for cell growth, and proliferation, and is widely used in nerve tissue repair.[56,57] In addition, collagen has been proven to possess a good adherence and proliferation ability for PC12 cells.[58] Therefore, the following experiments, including resazurin assay, live / dead assay and immunostaining of neurites, were carried out comparing the differences in PC12 cell functions on RSF/I₃K scaffolds and collagen scaffolds. Note that, three types of RSF/I₃K scaffolds were chosen, where the RSF scaffold concentration was 40 mg/mL and RSF/I₃K scaffold concentration ratios were 40:3 and 40:4 (represented as RSF/I₃K 40:3 and RSF/I₃K 40:4).

The metabolic activity of PC12 cells on sample surfaces (Type I collagen; RSF; RSF/I₃K 40:3 and RSF/I₃K 40:4) was determined after 24, 72 and 144 h in culture using a resazurin assay, and control groups performed on bare glass and TCP (tissue culture plastic) substrates (Fig. 4). Metabolic activity was observed to increase gradually between 24 and 144 h on all surfaces. Cells adhered on RSF coated surfaces showed the lowest metabolic activity amongst all test surfaces at 24 and 144 h, while for RSF/I₃K scaffolds, the metabolic activity increased more, indicating I₃K can promote PC12 cell proliferation. RSF/I₃K scaffolds at a ratio of 40:4 showed the highest metabolic activity which surpassed that of Type I collagen scaffolds (between 24 and 144 h), indicating a difference in proliferation on the surfaces. For control groups, cells on glass had a lower metabolic activity at all time points in contrast to TCP. The highest cell metabolic activity was observed on TCP samples compared to test surfaces at 24 h. However, the increase in

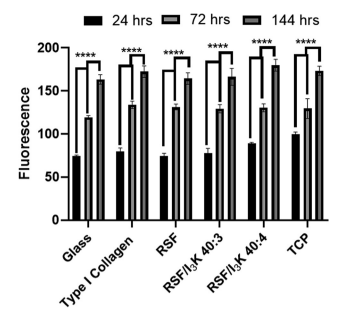


Fig. 4. Metabolic activity of PC12 cells adhered on different surfaces assessed using resazurin assay after 24-, 72-, and 144-hours culture. TCP represents tissue culture plastic. ($n \ge 3$; ***** p < 0.0001).

metabolic activity observed on the TCP surface between 72 and 144 h was similar to that of the other surfaces. This indicates excellent cell proliferation on the coated surfaces.

Live / dead assay results (Fig. 5) indicated a low rate of cell mortality on all surfaces after 6 days of culture. As can be seen from Fig. 5**B**, there was a slightly lower proportion of live cells on type I collagen (96.7% \pm 0.3%) compared to RSF (100%), RSF/I₃K 40:3 (99.3% \pm 0.1%) and RSF/I₃K 40:4 (98.4% \pm 0.2%) scaffolds. Although collagen is a well-known biocompatible material [59], RSF/I₃K showed better cell growth and proliferation overall. The population of cells adhered onto uncoated RSF surfaces was poor (Fig. 5C), only 34 ± 28 cells were observed, which is much lower than the adherence onto RSF/I₃K 40:3 (180 \pm 50) and RSF/I₃K 40:4 (290 \pm 50). Please note that, I₃K coated on RSF surfaces, resulted in a significant difference in observed percentage of live cells. That is due to RSF surfaces having a poor cell attachment, resulting in cells easily detaching from the RSF surfaces during cell culture, and only a few cells remaining. The results demonstrated that I₃K coated on RSF scaffolds can increase neuronal cell proliferation and attachment. Furthermore, the population of cells increased with increasing I₃K concentration, which is consistent with the results reported in section 3.3. According to Wiatrak et al. [58]. PC12 Adh cells show good attachment on plastic surfaces. As can be seen in Fig. 5C (results obtained from much larger areas than showed in Fig. 5A), there is no significant difference between RSF/I₃K 40:4 and TCP (note that cells on TCP showed patches with some areas having more cells (e.g., Fig. 5A(f)) and some areas having less). Therefore, RSF/I₃K 40:4 scaffold also promotes good PC12 cell attachment. However, there was still a significant higher cell population (650 \pm 30) on type I collagen than RSF/I₃K 40:4 scaffolds, indicating that peptides with multiple amino acids / functional groups are required to increase cell densities. The main amino acids in the collagen peptide are glycine, proline and alanine, [59] which can be used as building blocks to further design a modified self-assembled peptide based on I₃K to improve the performance in nerve tissue engineering.[60]

PC12 neuronal cells were labelled for β III-tubulin, a specific neurite formation marker (Fig. 6A). Short neurite outgrowth was observed for cells adhered to uncoated RSF surfaces and on glass substrates. However, at an I₃K concentration of 3 mg/mL coated onto RSF surfaces, neurite formations were observed, but slightly shorter than those on cells grown on type I collagen. With increas-

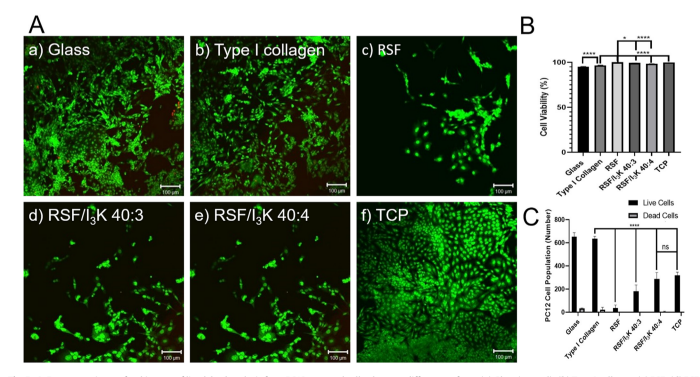


Fig. 5. A, Representative confocal images of live / dead analysis from PC12 neuronal cell culture on different surfaces, (**a**) Glass (control); (**b**) Type I collagen; (**c**) RSF; (**d**) RSF/ I₃K 40:3; (**e**) RSF/I₃K 40:4 and (**f**) TCP (control). The live cells (green) were stained by Syto-9TM and dead cells (red) were stained by propidium iodide, scale bar = 100 μ m. **B**, Percentage of cell viability. **C**, The population of live and dead cells. TCP represents tissue culture plastic. (n \ge 3; ^{****} p < 0.0001). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

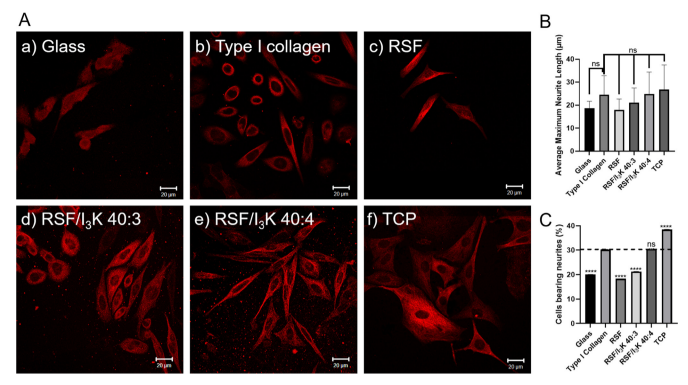


Fig. 6. A, confocal images of PC12 neuronal cells adhered onto different surfaces, including (**a**) Glass (control); (**b**) Type I collagen; (**c**) RSF; (**d**) RSF/I₃K 40:3; (**e**) RSF/I₃K 40:4 and (**f**) TCP (control). Neurites (red) were stained by anti- β III-tubulin, scale bar = 20 μ m. **B** and **C** indicate average neurite lengths and the percentage of cells bearing neurites in all scenarios. TCP represents tissue culture plastic. (n \ge 3; ^{****} p < 0.0001). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

ing concentration of I_3K to 4 mg/mL, longer neurite formation was observed, which indicated that the concentration of I_3K directly affects the length of neurites (Fig. **6B**), which is consistent with the previous AFM images (Fig. 3). The measurement of average maximum neurite lengths per neuronal cell revealed no significant difference between cells grown on type I collagen ($25 \pm 8 \mu m$) and RSF/I₃K 40:4 surfaces (Fig. **6B**). Experiments were terminated after 6 days, due to long neurite lengths being physically impaired at high cell density.[61] Comparison to glass and RSF surfaces, type I collagen and RSF/I₃K 40:4 surfaces had significantly higher percentage of cells bearing neurites (Fig. **6C**). However, no significant difference was observed between type I collagen and RSF/I₃K 40:4 surfaces, suggesting that I₃K at a concentration of 4 mg/mL possesses similar neuronal cell differentiation to type I collagen.

3.5. Micropatterning PC12 cells on RSF scaffolds via inkjet printing of peptide nanofibers

The micro-patterning of complex biomaterial structures plays an essential role in guiding cell adhesion, migration, differentiation and proliferation.[62,63] Inkjet printing can be used as an effective tool to micro-pattern complex structures of biomaterials onto a vast variety of bio-substrates including protein scaffolds[64]. Therefore, it has been deployed here to print I₃K (3 mg/mL) peptide nanofibers as the letters "SHEF" onto RSF (40 mg/mL) coated substrates. Cell culture studies revealed cells grew almost exclusively along the printed I₃K letters as shown in Fig. 7. The results were consistent with those reported by Poudel et al.[11] who used photolithography to pattern collagen type I on cell-repellent surfaces and demonstrated neural cell growth along the patterns. It was noticed that PC12 cells prefer to grow on the edge of the letter lines rather than their central areas. This is attributed to the so-called 'coffee ring effect' resulting from the inkjet printing possess, thus resulting in more I_3K nanofibers accumulating on the edge of the letters. [65,66] It is possible to reduce this effect by the addition of additives to alter the surface tension and spreading of the I_3K ink during the printing process, which might be deemed beneficial in the future. In nerve tissue engineering, the alignment of cells is important in axonal regeneration and direction [42]. Therefore, we have shown here the micro-patterning of PC12 cells via inkjet printing of the self-assembled I_3K peptide nanofibers onto RSF substrates may provide an excellent approach to enable the analysis of axonal development *in vitro*.[64,67]

4. Conclusions

The surfactant-like ultrashort peptide I₃K is able to selfassemble into nanofibrillar structures with a hydrophobic isoleucine tailed fibre core, and positively charged lysine residues located outside the fibres. [46,68] The self-assembled I₃K peptide nanofibers have been successfully used as templates for the fabrication of silica nanotubes.[37] In this study, I₃K peptide nanofibers were used as a cell-attractive agent to modify RSF scaffold surfaces to encourage neuronal cell (PC12 cell) attachment and growth. Commonly, positively charged PLL has been used for facilitating cell attachment and proliferation[69,70]. However, PLL can be cytotoxic, especially due to its high-molecular-weight.[71] Therefore, the structure of PLL should be modified via incorporation of segments that can reduce toxicity.^[72] In addition, the I₃K peptide used here, is more cost-effective than PLL due to its short sequence. Overall, we speculate that the positively charged I₃K peptide nanofibers could be used as an alternative to PLL for cell culture and tissue engineering applications.

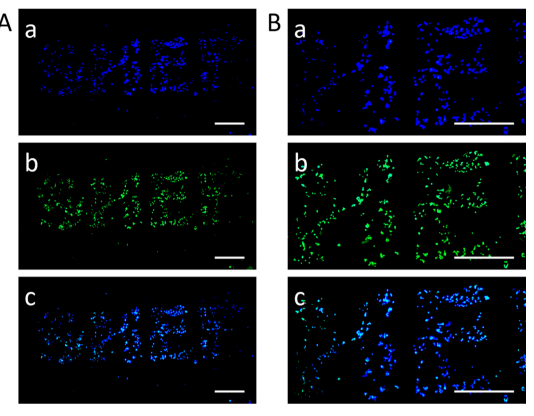


Fig. 7. A, Inkjet-printing of 1 layer of "SHEF" letters onto RSF coated glass surfaces (40 mg/mL) using I₃K peptide (3 mg/mL) as the ink, with PC12 neuronal cells growing along the printed letters. **B**, Enlarged letters ("HE"). (**a**, DAPI staining for nucleus; **b**, FITC-phalloidin staining for F-actin; **c**, merged images of a & b), scale bar = 500 μm.

It was found in our study that RSF-only coated surfaces had poor PC12 cell attachment, due to a lack of cell binding functional groups. However, this could be counteracted by introducing I_3K peptides nanofibers onto the scaffolds during the fabrication process. The peptide nanofibers naturally immobilise onto the RSF scaffold surfaces via charge-charge interactions, where the RSFonly scaffold surface is negatively charged and the I_3K peptide nanofibers are positively charged. The results showed that the presence of I_3K peptides promoted PC12 cell binding efficiency as the positively charged lysine residues facilitate cell attachment, which is equivalent to PLL[39]

Two methods (spin coating and inkjet printing) were applied to prepare scaffolds, with a series of different concentration ratios of RSF and I₃K onto glass and silicon wafer as substrates. Cells grown on the prepared scaffolds showed variable attachment, proliferation, and morphology including the formation of neurites. Additionally, the live and dead assay demonstrated that both RSF and I₃K demonstrated negligible cytotoxicity toward PC12 neuronal cells, indicating that both materials are promising scaffold materials for neural tissue engineering. In addition, the RSF/I₃K (ratio 40:4) produced scaffolds that optimally supported cell adhesion indicating excellent biocompatibility and differentiation. This also demonstrates that the ultra-short peptide I₃K could be used as an alternative to PLL for cell culture and tissue engineering applications.

Inkjet printing has been shown to be an excellent micropatterning method for the guidance of cell attachment.[73] The charge-charge interactions between positively charged I_3K peptide nanofibers and negatively charged RSF coated surfaces facilitated the robustness of the scaffold system during fabrication and cell culture work. Thus, enabling excellent cell growth along the printed patterns. This patterning method is a strength of inkjet printing offering a promising approach for analysing and understanding fundamental cellular functions such as neurite development and cell-cell interaction *in vitro*. [74,75]

CRediT authorship contribution statement

Weizhen Sun: Conceptualization, Methodology, Validation, Formal analysis, Investigation, Data curation, Writing - original draft, Visualization. Caroline S. Taylor: Methodology, Validation, Formal analysis, Investigation, Data curation, Writing - original draft. Yi Zhang: Methodology, Validation, Formal analysis, Investigation, Data curation, Writing - original draft. David A. Gregory: Methodology, Writing - review & editing. Mhd Anas Tomeh: Writing review & editing. John W. Haycock: Writing - review & editing. Patrick J. Smith: Resources. Feng Wang: Resources. Qingyou Xia: Resources. Xiubo Zhao: Conceptualization, Methodology, Writing - review & editing, Visualization, Supervision, Project administration, Funding acquisition.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary material

Supplementary data to this article can be found online at https://doi.org/10.1016/j.jcis.2021.06.086.

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