Mechanically robust cationic cellulose nanofibril 3D scaffolds with tuneable biomimetic porosity for cell culture†

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3D foam scaffolds were produced in a “bottom-up” approach from lyophilised cationic cellulose nanofibril (CCNF) dispersions and emulsions (CCNF degree of substitution 23.0 ± 0.9%), using a directional freezing/lyophilisation approach, producing internal architectures ranging from aligned smooth walled micro channels, mimicking vascularised tissue, to pumice-like wall textures, reminiscent of porous bone. The open, highly porous architecture of these biomimetic scaffolds included mesopores within the walls of the channels. A combination of SEM and NMR cryoporometry and relaxometry was used to determine the porosity at different length scales: CCNF foams with aligned channels had an average macropore (channel) size of 35 ± 9 μm and a mesopore (wall) diameter of 26 ± 2 nm, while CCNF foams produced from directional freezing and lyophilisation of Pickering emulsions had mesoporous walls (5 ± 3 μm) in addition to channels (54 ± 20 μm). Glyoxal crosslinking both enhanced robustness and stiffness, giving Young’s moduli of 0.45 to 50.75 MPa for CCNF foams with degrees of crosslinking from 0 to 3.04 mol%. Porosity and channels are critical scaffold design elements for transport of nutrients and waste products, as well as O2/CO2 exchange. The viability of MG-63 cells was enhanced on crosslinked, mechanically stiff scaffolds, indicating that these exquisitely structured, yet robust, foams could provide biomaterial scaffolds suitable for industrial applications requiring 3D cell culturing.

Introduction

Current demand for donor organs and tissues for transplantation vastly surpasses availability. For example, more than 100,000 US patients wait on the organ donor list in 2018 and, on average, 22 will die per day. The goal of tissue engineering is to develop new cell constructs that can be implanted into a patient to restore function to the damaged organs. This process requires a biocompatible scaffold to support the cells, often by mimicking the extracellular matrix (ECM), as these proliferate and grow into tissues. Traditionally, adherent cells are cultured on a two-dimensional (2D) scaffold in vitro. Cells growing on 2D scaffolds tend to only be attached to the substrate at their periphery, which forces the cells into a monolayer culture, rather than promoting layering as would be found in vivo, as well as limiting the size of the cell population produced. Once the cell layer has reached 100% confluence (all available surface is covered), cell viability can decrease, and cell death can occur as mass transfer of nutrients and oxygen diffusion is limited to depths of 100–200 μm. To retain cell viability beyond this depth, a vascularised network is required.

Furthermore, if primary keratinocytes are grown to 100% confluence, the phenomenon of “terminal differentiation” can arise: cells receiving insufficient nutrients die and those that remain alive become senescent, limiting the size of the cell construct that can be grown on a 2D scaffold.

On the other hand, 3D scaffolds require internal porosity to mimic the vascular morphology of native tissue, facilitating transfer of nutrients and gases and allowing larger tissue fragments to be grown. Ideally, the scaffolds should be resorbed and replaced, over time, by the newly regenerated tissue at the site of implant. A 3D arrangement of cells enables more complex cell–cell and cell–ECM interactions as cells are surrounded by ECM or scaffold, due to the greater surface area available for adhesion. Thus, cells cultured in a 3D scaffold...
more accurately mimic the response and behaviour of cells in vivo, beneficial for applications in both cell culturing for tissue engineering and as model tissues for drug development.10

Interconnecting porous networks can be used to promote cell growth, migration and mass transfer of nutrients.11 Generally, a larger pore size is considered to be beneficial for improving both cell migration and nutrient flow, however, this decreases the specific surface area of the scaffold, in turn reducing the matrix ligand density available for cell binding. Instead of migrating into the bulk of the material, cells tend to cluster around the edges of the scaffold.12 O’Brien et al., demonstrated that increasing pore size from 96 to 151 μm increased nutrient permeability through scaffolds,12 however, a pore size of between 20 and 120 μm was required for optimal balance between nutrient flow and ligand binding density.11 Therefore, a compromise must be reached between the two factors, pore size and specific surface area, of a 3D scaffold.13

Decellularised tissue has been investigated as a “top-down” method of obtaining the complex tissue structure and ECM composition needed for tissue regeneration, whilst retaining the microvasculature (<10 μm in diameter) that cannot readily be fabricated by current techniques such as 3D printing.14 Human donor or animal tissues can be used, but need to be sterilised and decellularised by enzymatic or detergent methods to remove the native cells and proteins. The remaining material is the ECM, which can vary in composition and structural properties, depending on the source.8 It is important that the decellularising process does not disrupt the ECM or adversely affect the biological activity or mechanical integrity of the remaining structure.15 The performance of the tissue can degrade with age and human tissue in particular, is in short supply.16 Decellularised scaffolds based on plant tissue have been reported,17 but, despite much of the vascularised structure being in place, scaffold shapes are limited to the shape of the plant tissue from which it was derived and further shape modulation is challenging.

Complementing the “top-down” approaches to the creation of porous 3D scaffolds, there are also a range of “bottom-up” methodologies to produce complex structures for cell culture, including: electrospinning polymer solutions to form 3D nanofiber meshes,18–20 3D printing of “bioinks” to fabricate complex scaffold architectures21 and the lyophilisation of dispersions or solutions to produce open porous aerogels or foams.22 Additive manufacturing approaches can reportedly give access to structures with minimum feature lengths of <100 μm,21 while reverse templating can be used to achieve porosities of 50–85% and interconnecting pores of 50 μm in size.23,24

There are several other “bottom-up” approaches to fabricate scaffolds suitable for applications in tissue engineering. For example, foam templating by generation of bubbles in solutions of polysaccharides containing surfactants,25 followed by freeze drying and crosslinking with a carbodiimide cross-linker yields a porous structure with pore diameters of 206–250 μm.26 Foam templating routes have also been applied to poly(vinyl alcohol) solutions27 and dextran–methacrylate solutions.28 Scaffolds so produced present highly ordered pore arrays with uniform and tuneable pore sizes.29 As pores are interconnected, mass transfer of nutrients to cells within such scaffolds is enhanced.

Freeze drying, or lyophilisation, is a popular method to introduce porosity into the scaffold due to its low cost, simplicity and versatility.24 In this process, ice crystals act as a porogen, leaving a porous structure post sublimation under vacuum.30 The rate of cooling during the freezing step can affect the size and distribution of ice crystals formed and thus the size and form of the pores within in the resulting foam.31 An advantage of using this technique is that water is commonly the solvent; hence impurities, such as surfactants, are not added during the lyophilisation stage, making it especially beneficial for biological applications. The introduction of emulsions, nanoparticles and dilute polymer solutions into the material can further modulate macro- and mesopore formation.30 Ice crystals grown in a unidirectional fashion leave behind channels of aligned pores, mimicking the vascularity of native tissue32 and can be applied to a range of scaffold materials, including: hydroxyapatite,33 silk fibroin,14 gelatin,35 dextran,36 and, more recently, chitosan–alginate blends,37 cellulose–chitosan blends,38 and cellulose solutions.39 The resultant scaffolds have the potential to expand the use of these materials from 2D films to 3D porous scaffolds and many authors suggest that the channels can promote vascularisation in vivo, although corroborating cell studies are only reported for some materials.

Scaffolds fabricated from cellulose, using both “top-down” and “bottom-up” approaches, have been tested in a range of cell culture applications.40–43 Cellulose offers many beneficial attributes to tissue engineering, such as biocompatibility, versatile chemical and physical properties, and ease of processing. It is also a cost effective and sustainable material, which makes it suitable for industrial applications.44–48 However, previous reports of directionally frozen/freeze dried cellulose hydrogels for scaffolds have largely focussed on the materials characterisation aspects – few reports of the growth of cells on these 3D scaffolds have appeared.38,39

Previously, we have demonstrated that cellulose surfaces grafted with cationic moieties allow for attachment of MG-63 cells in the absence of matrix ligands.49 Here we use similarly modified nanofibrillar materials to develop novel porous 3D scaffolds that we suggest offer many advantages over “top-down” scaffolds such as those derived from decellularised plant tissue. In particular the use of a scalable method offering ease of manufacture, tunable porosity and mechanical properties, to prepare materials that can be stored without special requirements, yet which mimic vascularised tissue, may provide supports beneficial for various cell culture applications.

Materials and methods

Cellulose powder (C8002), sodium hydroxide pellets (≥98%), glycidyl trimethylammonium chloride (GTMAC, ≥90%), DMSO (≥99%), 1-ethyl-3-methylimidazolium acetate (97%),
cyclohexane (99.5%), absolute ethanol (≥99%) and N,N-dimethylacetamide (DMAc) were purchased from Sigma-Aldrich and used as received (except DMAc, which was dried over 3 Å molecular sieves prior to use). For crosslinking modifications, glyoxal 40 w/w% aqueous solution was purchased from Alfa Aesar and made up to required concentrations with deionised (DI) water. Mobile phases for HPLC were prepared from H₂SO₄ (99.99%) and sodium phosphate monobasic monohydrate, sodium phosphate dibasic and sodium azide powders purchased from Sigma Aldrich.

For cell studies, Dulbecco’s Modified Eagle Medium (DMEM, GlutaMAX™), non-essential amino acids, sodium pyruvate, trypsin (0.05%) and trypan blue (0.4%) were purchased from Gibco and stored at 4 °C. Foetal bovine serum (FBS, non-USA origin), MG-63 human osteosarcoma cells, and methylthiazolyldiphenyltetrazolium bromide (MTT) powder were purchased from Sigma-Aldrich. Phosphate buffer solution (PBS, 0.1 M) and the mixture stirred for 1 h at room temperature, resulting in a 4 wt% CC solution. For comparison, a solution of unmodified cellulose (UC) was prepared using the same procedure.

CC and UC films were formed by casting the solutions described above onto a clean glass sheet using an Elcometer 3700 reservoir and an Elcometer 4340 Automatic film applicator, with a gap of 0.8 mm between the blade and glass surface. The films were regenerated by immersion in an ethanol anti-solvent bath for 24 h. Residual EMImOAc and DMSO were removed by Soxhlet extraction with ethanol overnight. The films were washed twice with copious amounts of DI water to remove excess EtOH before being stored in 20 wt% MeOH solution to inhibit bacterial growth.

**Scaffold characterisation**

**NMR cryoporometry.** Using a procedure adapted from Johns et al., foam and film samples were hydrated in PBS overnight, excess PBS was removed, the samples placed in individual NMR tubes and sealed using damp absorbent paper to maintain humidity. The ¹H NMR signal was recorded on a 400 MHz Bruker Avance spectrometer equipped with a 5 mm BBO probe, running with the boil-off from liquid nitrogen as cooling gas, and a BVT3200 temperature control unit with a precision of ±0.1 K. Actual versus recorded temperatures had previously been verified using methanol. A simple spin echo sequence was used, with an echo time of 2.2 ms, to ensure minimal suppression of signal from liquid water and complete suppression of signals from both cellulose materials and frozen water. Measurements were performed by decreasing the temperature to 218 K in order to completely freeze the sample, followed by stepwise temperature increase in 5 K increments up to 258 K, then in 1 K increments to 267 K, 0.2 K increments to 271 K, and finally to bulk melting temperature using a temperature step of 0.1 K. At each increment signals were recorded after establishment of thermal equilibrium, achieved by a waiting time of 20 min (Fig. S8–S13, ESI†).
The melting point depression, $\Delta T$, is related to the pore radius, $r$, via bulk properties of the probe liquid, $P$, as described by the Gibbs-Thomson equation:

$$\Delta T \propto P(r - s)$$  

where $P$ is 25 nm for water and $s$ represents the thickness of a pre-molten liquid-like layer on the surface of the substrate, here assumed constant over the temperature range at two monolayers thick, i.e. 0.6 nm.\(^{56,57}\)

**NMR $T_2$ relaxation.** NMR Carr-Purcell Meiboom-Gill (CPMG) experiments were performed using a Bruker Minispec MQ-20 spectrometer operating with a magnetic field of 0.5 T ($^1$H Larmor frequency of 20 MHz); 50,000 echoes were acquired with echo time of 70 $\mu$s and recycle delay of 15 s. The CPMG decay curves were processed to obtain the $T_2$ distribution using a non-negative least square procedure known as a numerical Inverse Laplace Transform, ILT.\(^{58,59}\) The obtained $T_2$ distributions were deconvoluted using log-gaussian functions to provide the contribution of each component in the pore structure. Fluid bound in a pore interacts with the pore surface, restricting the molecular mobility of the fluid. Such restriction is reflected by a decrease of the transverse relaxation time, $T_2$, and can be quantified, in the fast diffusion regime, by the relation:\(^{60,61}\)

$$1/T_2 = \rho(S/V) = 2\rho/r,$$  

where $\rho$ is the surface relaxivity, which depends on the particular porous media and is usually unknown, and $S$ and $V$ are the pore surface area and volume, respectively. In a realistic scenario, the distribution of pore sizes and differences in the fluid mobility within the pores result in a multieponential decay of the CPMG signal, i.e., a distribution of $T_2$ times.

To assess the pore structure of the cellulose foams, DMAc (HPLC grade) was used as a molecular probe, due to its weak interaction with the cellulose structure. DMAc does not significantly modify the cellulose pore structure, while water, often used as molecular probe for relaxometry experiments, alters the pore distributions in biomass samples.\(^{62}\) Lyophilised modified cellulose foams and films were dried under reduced pressure (640 mmHg) for 24 h at 80 °C. Samples were soaked in DMAc and kept in a desiccator under a 600 mmHg vacuum for 24 h, following which excess DMAc was removed by centrifuge filtration at 600 g for 1 min (Corning Costar Spin-X, 0.45 $\mu$m, nylon membrane filter). All measurements were carried out in duplicate and the CPMG decay of the mean used in the ILT procedure (Fig. S14–S16, ESI†).

**Scanning electron microscopy.** Field Emission Scanning Electron Microscopy (FESEM) was used to characterise the internal morphology and porosity of the different CCNF foams. A JEOL FESEM 6301F was used to image the surface of the foams as well as the cross section and internal structure of the lyophilised CCNF foams at ultrahigh resolution. To prepare the samples for surface imaging, the foams were carefully adhered to a metal stub with double sided carbon tape. To obtain a cross section, the dried foams were frozen in liquid nitrogen, fractured using a very sharp blade and attached to the stubs, prior to vacuum drying for 24 h. All samples were sputter coated with a 20 nm layer of chromium in an argon environment prior to imaging. A high sensitivity backscattered electron detector was used for computational imaging. The samples were imaged at an acceleration voltage of 5 kV at magnifications between 50× and 25 000×.

**Mechanical properties.**

**Foams.** The Young’s moduli of the CCNF foams were determined using an Instron 3343 electromechanical test machine. CCNF (23.0 ± 0.9% DS) crosslinked with 0, 2.5, 5, and 10 wt% glyoxal solutions were tested. The dry foams were placed between steel plates and a 1000 N load cell was used to deliver a compressive load at a rate of 1 mm min$^{-1}$ to the foams until deformation or failure occurred. Four samples were tested per crosslinked foam and an average reported. The Young’s modulus of the foams was calculated using the equation:

$$E = \frac{F}{A} \Delta L/L,$$

where $F$ is the compressive load, $A$ is the sample area, $\Delta L$ is the degree of sample compression and $L$ is the original sample height.

**Films.** The bulk elastic moduli of the regenerated cellulose scaffolds were determined using a Dynamic Materials Analysier (DMA1 STAR® System, Mettler Toledo). The samples used were films regenerated from 4 wt% cellulose and CCNF (DS = 23.0 ± 0.9%) solutions, with a range of crosslinking in both sets (DXL 0 to 3%). Both hydrated “never-dried” and dried films were cut into strips $\geq$1.50 cm in length by 0.50 cm width and the thickness recorded using a steel digital vernier micrometer calliper. The film strips were gripped between titanium tension clamp sample holders and a preload force of 1 N applied to the sample. An offset of 10 $\mu$m was set at a frequency of 1 Hz and the elastic moduli were determined over 5 min. Five samples were tested for each film and an average reported.

**Cell studies.**

**Cell culture.** MG-63 cells were cultured in DMEM supplemented with 10% FBS, 1 v/v% non-essential amino acids, 1 v/v% sodium pyruvate and 1 v/v% penicillin streptomycin at 37 °C in a humidified incubator with 5% CO$_2$. When MG-63 cultures reached 80% confluence (3–4 days) the cells were passaged and reseeded in a T75 flask at a density of 5000 cells per cm$^2$ with fresh media.

**Cell viability.** Lyophilised foams, placed in 48 well tissue culture plates and sterilised using a Hoyer™ UVC 500 crosslinker at 254 nm for 15 min prior to being rehydrated and soaked in three 0.5 mL aliquots of PBS, to remove any unreacted crosslinker. MG-63 cells were used to assess the viability of the CCNF foams as cell scaffolds. The scaffolds were seeded with a cell density of 10 000 cells cm$^{-2}$ (corresponding to a total cell number of 9500 cells per sample) in growth media and placed in an incubator at 37 °C in 5% CO$_2$ to proliferate for a given time. Tissue culture plastic was used as a control “scaffold” for the MTT assay and material controls were performed for each sample. $N=8$ replicates per sample were measured.
A MTT stock solution of 50 mg mL\(^{-1}\) in PBS was prepared. After appropriate incubation time (1, 4 and 7 days), the growth medium was replaced with 250 μL of 5 mg mL\(^{-1}\) MTT in growth medium and incubated for a further 4 h. The culture medium was gently removed from the well plate and 200 μL of DMSO added to dissolve the formazan crystals generated in this assay. Well plates were placed on a plate shaker and gently agitated for 10 min. The DMSO-formazan crystal suspension was transferred to a 96 well plate and the absorption intensity recorded at 570 nm and 690 nm, using a Biotek Synergy HT plate reader. To determine the activity the following equation was used:

\[
\text{MTT activity} = \frac{(S_{570\text{nm}})}{(MC_{570\text{nm}})}
\]

where \(S\) is the absorption of the sample and \(MC\) is the material control, i.e., absorption due to the sample material sans cells at the given wavelengths.

**Cell visualisation.** First, cells were grown on CCNF and CCNF crosslinked (XL) scaffolds for 24 h in growth medium. A fixative solution was prepared by adding 2 mL glutaraldehyde (GDA at 25 wt%) to 10 mL double strength cell culture medium (sans serum) and diluted to 20 mL with DI H\(_2\)O. This gave a fixative solution of 2.5 wt% GDA in normal strength culture medium.

The culture medium was very slowly removed from the scaffold keeping the pipette at the edge of the scaffold away from the cells. The samples were gently washed in 2 × 1 mL fresh medium (normal strength culture medium without serum) again keeping the pipette at edge of the scaffold. The wash medium was removed as above, replaced with the fixative solution and left for 2 h. The samples were rinsed three times with 1 mL wash medium. The cells were post-fixed in aqueous 1 wt% osmium tetroxide for 1 h in a fume hood at ambient temperature, then washed in 1 mL DI H\(_2\)O three times at 15 min intervals.

The cells were dehydrated by sequential washing with solutions of acetone in H\(_2\)O gradually increasing in acetone content to 100% dry acetone. Specifically solutions of 50, 70, 90, 95% and 100% acetone in H\(_2\)O were applied to the cells (on scaffold) for 15 min, repeated three times for each concentration. Acetone was replaced with 1:1 dry acetone:HMDS solution for 15 min, followed by three sequential treatments with 100% HMDS for 30 min each. Excess HMDS was removed and the samples allowed to air dry in a fume hood for 2 h prior to mounting onto SEM stubs.

**FBS absorption.** A phosphate buffer solution (mobile phase) was prepared by adding a solution of 13.8 g sodium phosphate monobasic monohydrate in 500 mL DI H\(_2\)O to 14.2 g sodium phosphate dibasic in 500 mL DI H\(_2\)O until the pH was 6.8. The slightly acidic solution was made up to 1 L with DI H\(_2\)O, 0.2 g of sodium azide added and the solution degassed. A solution of 10 wt% FBS in PBS was prepared, 250 μL of the FBS solution was added to each sample and the plate stored in an incubator for 24 h. The amount of FBS absorbed onto the scaffold surface was determined using an Agilent Technologies 1260 infinity HPLC. A range of FBS solutions with concentrations in the range 0.1 to 30 wt% were prepared and a calibration curve constructed. HPLC analysis: size exclusion column (TSKgel G4000 PWXl, 7.8 mm, 10 × 30 mm, 50 °C), mobile phase phosphate buffer (0.7 L min\(^{-1}\)), and UV detector. The amount of FBS present was calculated by subtracting the concentration present in the solution from the initial concentration of 10 wt%.

**Results and discussion**

Application of directional freezing and lyophilisation to aqueous dispersions of CCNF and CCNF stabilised oil-in-water Pickering emulsions, led to preparation of 3D cationic cellulose scaffolds in the form of foams with exquisite tuneable structure. Unidirectional ice crystal formation resulted in channels in the tens of micron range, mimicking vascular structures, and preservation of spherical pores arising from emulsions, yielded pumice-like wall textures, reminiscent of porous bone (Fig. 1).

Crosslinking, previously shown to enhance cell spreading\(^{63}\) (indicative of cell viability and scaffold compatibility), was used to render the delicate structures robust enough for handling.
and repeated immersion in cell growth media, as would be required for use in tissue engineering applications.

To illustrate their utility as tissue culture scaffolds, the attachment and viability of MG-63 cells was assessed using MTT assays and SEM analysis to discern how well cells became integrated into the bulk of the putative scaffold material and whether or not these cells began to proliferate.

**“Bottom-up” fabrication of vascularised 3D scaffolds**

3D scaffolds, with variable form, internal architecture and robustness, were constructed using a “bottom-up” approach from “cationic cellulose”, CC, bearing quaternary ammonium groups derived from glycidyl trimethylammonium chloride (GTMAC) grafting, either as CC nanofibrils, or dissolved and regenerated CC films.

To induce directional freezing, low weight% CCNF dispersions (weak hydrogels) were placed, in containers, on metal blocks immersed in liquid nitrogen. This resulted in exclusion of the CCNF into “walls” surrounding oriented ice crystals, which, upon lyophilisation, yielded aligned channels, reflecting the size of the ice crystals formed. Cyclohexane in water CCNF stabilised Pickering emulsions treated similarly resulted in both channel-like porosity and walls bearing much smaller pores, 3–5 μm in diameter, due to oil droplet templation of the hydrogel. These materials are designated “CCNF PE” henceforth.

Although the foams contained both aligned micro channels reflecting the longitudinal ice crystal growth, and pores, reflecting the emulsion structure, these were very delicate and easily compressed, or even dispersed. Even gentle agitation in cell culture medium caused the scaffolds to break apart, rendering them useless as scaffolds in cell culture applications. Hence, the foams were crosslinked to enhance robustness, especially these as scaffolds in cell culture applications. Hence, the foams were crosslinked to enhance robustness, especially when hydrated, and to retain the complex internal architecture.

To reinforce these delicate porous materials, glyoxal, a crosslinker, was added to the CCNF dispersions and later reacted by heating, in order to fix the complex internal structure, through *in situ* glyoxalation – these materials are designated as “CCNF XL”. In common with most reactive crosslinkers, unreacted glyoxal is cytotoxic, but once reacted to form acetal and hemiacetal linkages with cellulose, does not inhibit cell viability.

This modification served to support the delicate structures, maintaining the integrity of the vascularised or porous internal architecture, but it has also been shown previously that crosslinking further increases the spreading of attached MG-63 cells on cationised cellulose scaffolds. Furthermore, modification of the mechanical properties of the scaffold can act as a stimulus to up-regulate physiological processes and signalling pathways within the cell cycle, thus promoting cell growth.

Thus, these materials could serve as very flexible scaffolds, allowing external shape/size, internal architecture, rigidity and robustness to be varied while also offering opportunities to modulate cell response, as described later.

**Scaffold porosity**

To determine the effect of lyophilisation and glyoxalisation on macroporosity, the CCNF, CCNF XL, and CCNF PE scaffolds were characterised using SEM and compared to regenerated lyophilised films (Fig. 2 and Fig. S17–S19, ESI†). Structures accessible ranged from unidirectional, vascularised materials with smooth walls (Fig. 2a), foamed walls (Fig. 2c) to more open pore structure materials (Fig. 2b), compared with regenerated cationic cellulose films (Fig. 2d). (For comparison, an SEM image and pore size distribution for unmodified cellulose films is included in the ESI† (Fig. S19).) To determine the average pore size diameter SEM images were analysed using ImageJ software yielding the following average pore sizes: CCNF = 35 ± 9 μm, CCNF XL = 60 ± 20 μm, CCNF PE = 54 ± 20 μm and 5 ± 3 μm, and CC film = 20 ± 10 μm. The presence of 5 wt% glyoxal in the CCNF XL hydrogel appeared to reduce the alignment of ice-crystals formed during the freezing process, while emulsion templating introduced smaller pores, ~5 μm in

![Fig. 2](image-url) SEM images and pore size distributions of lyophilised foams (a) 2 wt% CCNF hydrogel, CCNF; (b) 2 wt% CCNF hydrogel + 5 wt% glyoxal, CCNF XL; (c) 1 wt% CCNF Pickering emulsion templated hydrogel, CCNF PE; and (d) regenerated 4 wt% cationic cellulose film, CC.
diameter, into the walls of the lyophilised hydrogel structure. Film structures, while porous, appear to show less connectivity between pores and a dense “skin” layer.

Along with the macroporous network, visible in SEM images, mesopores occur within the walls of the freeze dried foams. NMR cryoporometry was used to probe the permeability of the foam walls to PBS (Fig. 3 and Table 1). PBS was used to reflect typical cell culture media. Importantly, this technique allows the pore structure of hydrogels to be analyzed in the hydrated materials, thus more accurately mimicking the microenvironment cells would experience in scaffolds. The smallest mesopore diameter, 20 ± 2 nm, was observed in CCNF XL foams, where the pore size was found to be very similar to that in the control regenerated CC films, 18 ± 2 nm. The wall pore modal diameter determined for CCNF foams was 25 ± 2 nm and was not significantly affected by the concentration of initial CCNF dispersion used. Crosslinking appeared to lead to a slight reduction in wall pore diameter to 20 ± 2 nm. In contrast the CCNF PE foam does not allow for penetration of PBS solution into the walls, as reflected by a modal diameter matching that determined for the bulk PBS solution control. It is likely that this reflects the densification of CCNF at the oil/water interface in the Pickering emulsions, yielding structures permeated by a plethora of pores arising from oil droplets, but with dense close-packed CCNF walls. These 3D scaffolds, formed from CCNF dispersions and emulsions, thus have porosity at three different scales: large pores or channels resulting from ice-crystal templation; smaller pores arising from templation by oil droplets in Pickering emulsions; and nanoscale pores reflecting packing of the CCNF within the walls of the foamed structures. As this differing porosity could be beneficial for both penetration of cells into the scaffold and mass transfer of nutrients, gases and waste products to and from cells during cell growth, we sought to unify the porosity measurements to allow comparison of potential scaffold materials and thus to inform future selection of the form of templates.

NMR T$_2$ relaxometry experiments were used to probe the accessibility of DMAc to three different scales on the materials, given by each component in the T$_2$ distribution. Such scales are related to pores of a few nanometers, for T$_2$ values of about 10$^{-1}$ s, mesopores of dozens of nanometers, for T$_2$ values ranging from 10$^{-2}$ to 10$^{-1}$ s and large pores of hundreds of nanometers up to one micrometer, for T$_2$ values above 10$^{-1}$ s. The T$_2$ distributions for all scaffolds are shown in the ESI† (Fig. S14–S16, ESI†).

The proportion of DMAc accessing pores on each length scale can be estimated from the respective relative areas of the log-gaussian obtained from deconvolution (Fig. 4). The greater the accessibility, the more pores are filled with the fluid at a particular length scale. Thus, CCNF XL has a greater accessibility of pores on the smallest scale in comparison to CCNF and CCNF PE, suggesting that crosslinking increases the number of accessible nanopores in the cell wall. This reflects the observation that crosslinked cellulose materials are less prone to hornification than uncrosslinked celluloses.

While the T$_2$ distributions do not give precise pore sizes due to the unknown surface relaxivity for each sample, if the

![Fig. 3](image)

**Table 1** Effect of CCNF surface modification and crosslinking on the mesopore modal diameter, determined by NMR cryoporometry, and macropore diameter, determined by analysis of SEM images.

| Scaffold type | Sample    | Mesopore diameter/nm | Macropore diameter/μm |
|--------------|-----------|-----------------------|------------------------|
| a            | CCNF 1 wt%| 25 ± 2                | 35 ± 9                 |
| a            | CCNF 2 wt%| 26 ± 2                |                        |
| a            | CCNF 4 wt%| 27 ± 2                |                        |
| a            | CCNF XL 2 wt% | 20 ± 2                | 60 ± 20                |
| b            | CCNF PE 1 wt% | 30 ± 2                | 5 ± 3†, 54 ± 20†       |
| c            | CC 4 wt%  | 18 ± 2                | 20 ± 10                |
| c            | UC 4 wt%  | 35 ± 2                | 0.6 ± 0.3              |
| Control      | PBS       | 32 ± 2                | n/a                    |

*a* Directionally frozen dispersions. *b* Directionally frozen Pickering emulsions. † Cast regenerated films. ‡ Crosslinker concentration in solution = 5 wt%. *f* Macropore diameter – emulsion droplets. *f* Macropore diameter – channels.
previously\textsuperscript{63}), glyoxalation was effected by sequential soaking in glyoxal solution and heating (as used previously\textsuperscript{64}, \textsuperscript{65}, \textsuperscript{66}). This enabled the foams to be handled without breaking during cell viability studies.

**Mechanical properties and robustness**

As mentioned previously, uncrosslinked CCNF foams were very delicate, becoming broken up and dispersing readily in DI H\textsubscript{2}O or buffer upon even very gentle agitation. Foams formed in moulds to create more complex 3D shapes, as may be required for cell studies, could be handled and survived the manipulations required for cell studies (e.g., aspiration of media), yet were not rendered too brittle to be of utility; CCNF foams with a crosslinker concentration of 2.5 wt% were used in cell culture studies (Fig. S20–S27, ESI\textsuperscript{[f]}).

To avoid compromising the exquisite structures formed by sequential soaking in glyoxal solution and heating (as used previously\textsuperscript{63}), glyoxalation was effected \textit{in situ}: glyoxal solutions were added to the hydrogels prior to freezing and the curing step took place post lyophilisation on the now dehydrated foam. This simple procedure is attractive for manufacturing as it reduces the number of processing steps required in foam scaffold fabrication, making scaffold manufacture readily scalable. The dry foams thus produced had degrees of cross-linking from 1.18 to 3.04 mol\%, as determined by HPLC\textsuperscript{[f]} analysis (Fig. S6, ESI\textsuperscript{[f]}), comparable with glyoxalation conducted on cellulose films,\textsuperscript{63}

As the concentration of glyoxal crosslinker used increased from 2.5 to 10 wt\%, the crosslinked scaffolds became more robust with mechanical properties, such as Young’s modulus and compressive strength, increasing accordingly (Fig. 6a and b). Compressive strength, in particular, was enhanced, with the Young’s modulus, $E$, increasing from 0.1 MPa for uncrosslinked CCNF foams to 50.8 ± 8 MPa at 10 wt\% glyoxal (Fig. 6a). It was apparent from the compressive load versus compressive extension profiles that compression was occurring in two stages; initial compression at lower loading was assigned to the collapse of the large porous network and a second stage, at much higher loads, reflected the bulk of the material under compression (Fig. 6b). $E$ was determined from the stress/strain at 30\% deformation. Although the highest $E$ value occurred in the most crosslinked foams, these tended to fracture more quickly, reflecting a more brittle structure; hence foams up to 5 wt\% were used in cell culture studies (Fig. S20–S27, ESI\textsuperscript{[f]}).

Such foams represented a “sweet spot”: materials were robust enough to handle and survive the manipulations required for cell studies (e.g., aspiration of media), yet were not rendered too brittle to be of utility; CCNF foams with a moderate level of crosslinking (exposed to 2.5 and 5 wt\% glyoxal, yielding DXL of 1.18 and 3.04 mol\%) remained intact in PBS and only disintegrated if subjected to very vigorous agitation. This enabled the foams to be handled without breaking during cell viability studies.

**Cell response to 3D scaffolds**

We have previously investigated the attachment of cells on the surface of CC films,\textsuperscript{68} but here the presence of a “skin” on the surface of the regenerated CC films (Fig. S17, ESI\textsuperscript{[f]}) rendered the porous internal structure beneath inaccessible to cells, so that cell studies were confined to the foamed scaffolds produced by directional freezing, lyophilisation and crosslinking.

The cell viability was determined by assessing cell metabolism using an MTT assay, which showed that MG-63 cells were viable...
It has been reported previously that the mechanical properties of a cell scaffold affect the mechanosensitive cell response, although this can be dependent on the cell type. For example, fibroblasts, smooth muscle cells and epithelial cells spread more and develop larger focal adhesions and actin stress fibres than on compliant ones. Cells receive mechanical feedback from the substrate to which they adhere, even in the absence of externally applied forces, as they attempt to deform the substrate. As cells elongate on stiff scaffolds they experience a higher stress than on softer more compliant scaffolds, which promotes the assembly of the cytoskeleton into actin stress fibres and focal adhesions, and subsequently triggers signalling cascades that ultimately promote cell expansion.

Thus, we suggest that the cells adhered to the CCNF XL scaffolds due to the cationic surface charge, which promotes the initial phase of cell attachment, facilitating cell binding through electrostatic interactions and possibly aided by enhanced FBS sorption, particularly in the second phase of cell adhesion (Fig. 8). Once attached, the stiffer CCNF XL scaffolds provided the cells with a larger feedback force, promoting...
of cell waste products, beneficial for cell growth. Porosity was characterised at all length scales by a combination of SEM image analysis, NMR cryoporometry and T2 relaxation studies. These scaffolds can complement the library of 3D porous materials derived from decellularised plant tissue, yet have the advantage over the “top-down” scaffolds, which usually require several treatments prior to application and can be limited in structure and shape, whereas, a variety of bulk 3D shapes can be prepared through process moulding the frozen hydrogel prior to lyophilisation and crosslinking.

These “bottom-up” scaffolds derived from dispersed CCNF require minimal modification, use known and scalable chemistries, and can be easily processed through the use of moulding techniques to create the desired 3D constructs, with modulated vascularity and wall porosity.

Conflicts of interest

There are no conflicts to declare.

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