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## **PAPER**

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Bacterial cellulose (BC) nanofibers secreted by Komagataeibacter xylinus 10245 were applied alone or in combination with chitosan to prepare highly aligned and porous scaffolds through a combined liquid nitrogen-initiated ice "templating" and freeze-drying process. Their morphology and physical properties were controlled by adjusting the concentration of chitosan over a range of 1, 1.5, and 2% (wt%) and analyzed by Scanning Electron Microscopy (SEM), Brunauer-Emmett-Teller (BET), and X-ray diffraction methods. The SEM images confirmed a distribution of fibrils vectorially aligned in the freezing axis direction, while chitosan contributed to the development of a dense network, superior mechanical properties, and biomedical relevance of the final scaffolds. It was found that as the chitosan concentration increased, the crystallinity index decreased from 89% to 79% likely because of strong intermolecular bonding. However, the scaffolds containing chitosan demonstrated excellent shape recovery and structural stability after compressive tests and may act as excellent scaffolds for potential cartilage tissue engineering applications.

An environmentally benign approach to achieving

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vectorial alignment and high microporosity in

bacterial cellulose/chitosan scaffolds†

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## Introduction

Although cellulosic materials are ubiquitous in nature, in general, isolation of pure cellulose from terrestrial or aquatic biospheres requires expenditure of significant energy to remove contaminants such as lignin, heteropolysaccharides, and related biomolecules. Bacterial cellulose (BC), however, is a cellulosic material with the same chemical structure as plantor animal-based cellulose, but it does not contain lignin in contrast to plant cellulose and hence is much purer. It is known that various genera of bacteria such as Komagataeibacter within the appropriate environment can produce highly porous and fibrous BC networks. The biomaterial is characterized by high water sorption, gel-forming ability, biocompatibility, high modulus of elasticity, and extremely high purity. It has found applications in pulp & paper products, audio components (e.g.,

speaker diaphragms), conductive materials, and soft tissue engineering.1-3 It possesses unique morphological and mechanical properties as evidenced by nanoscale fibrillar structure and high tensile strength, respectively; most importantly, it displays biocompatibility from analogous chemical motifs to the extracellular matrix (ECM) component of naturally occurring tissues, i.e., collagen, that make it a desirable building block for soft tissue engineering.3-8

Chitosan, like cellulose, is an abundant and ubiquitous polysaccharide widely known for biomedical applications. It is a linear polysaccharide composed of glucosamine and N-acetyl glucosamine residues linked in a β-(1-4) regular structural pattern. It can acquire a cationic nature because of the -NH<sub>2</sub> groups on the chitosan backbone to modulate electrostatic interactions with anionic glycosaminoglycans (GAGs), proteoglycans, and sundry negatively charged molecules. The electrostatic property is of great importance because most growth factors are attached to GAG to help chitosan-based scaffolds to maintain the growth factors secreted by cells.9 Chitosan has secured significant interest for as a tissue engineered scaffold because in addition to its charge interactions with GAGs, it aids in wound healing, drug delivery, and displays antibacterial activity.

Engineered polymer-based tissue engineered scaffolds require several properties for successful application. Among them, the scaffold material must provide a platform for cell seeding, growth, and proliferation. In fact, human cells require a complex selection of pores, ridges, and fibrous materials

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within the extracellular matrix. Therefore, factors that enhance cell attachment, migration, and nutrient delivery to such cells include fiber size, porosity (pore size & structure), the void volume, and the architecture of the system. Scaffold porosity is among the most important parameters for cell migration and ultimately for vascularization. A porous scaffold will enable facile diffusion of nutrients, oxygen, and waste within implanted tissue. Finally, a high specific surface area will facilitate strong cell attachment. He phenotypic cell expression, however, can be altered if the fiber diameter does not coincide with the pre-requisite size of the cells within the ECM. Furthermore, uniform fiber directionality is an essential parameter in engineering scaffolds because it influences not only cell phenotypic expression, attachment, and proliferation, but final scaffold mechanics.

Different techniques have been adopted to mimic natural tissue properties such as pore size and fiber directionality. Among them are gas foaming, 16 solvent casting/particulate leaching, 17 phase separation, 18 fiber mesh generation, 19,20 and electrospinning. 11 In the gas-foaming technique, the scaffolds have closed cellular structures, whereas in solvent casting/particulate leaching, the morphology of the resulting scaffold cannot be controlled because chemical modifications occur during dissolution and removal of the solvent that alter final material properties. 11 Although electrospinning and fiber mesh techniques facilitate fiber orientation, the final scaffolds typically lack both acceptable structural stability and mechanical strength. 12

The freeze-drying method has been used to produce porous scaffolds because of its simplicity, low cost, and ease of use. It requires sublimation under specific temperature and pressure profiles over a defined time, but does not require chemical reactions. Thus, properties such as pore structure, size, and networking can be modulated and fine-tuned.<sup>11</sup> In addition, the method is considered highly functional because it allows production of porous materials with vectorially-guided morphologies. In fact, aligned scaffold porosity is critical to successful tissue engineering to ensure directional growth of cells.<sup>23</sup> Various functional polymer-based scaffolds possessing tailored microstructures and physical properties have been prepared by freeze-drying.<sup>24</sup>

Freeze-drying is an approach for removing water from BC, 8,25-27 but until this report, no work has shown vectorially-guided final morphologies by using liquid nitrogen. In the past, homogenous and well-structured foams of cellulose have been prepared by dissolving cellulose in solvents to develop gels, while vectorial alignment in cellulose nanofibers was obtained by dispersing fibers in water under high-pressure homogenization. However, the latter techniques tend to result in a loss of desirable physical and chemical qualities.<sup>28,29</sup>

The current study has developed a porous BC scaffold by using a green approach that consists of an environmentally benign liquid nitrogen-initiated ice-templated freezing methodology coupled with freeze-drying. Chitosan was added to the system to impart relevant biological properties; the systems

were evaluated in terms of their morphological, chemical, and physical characteristics.

## Experimental section

#### **Materials**

Komagataeibacter xylinus (ATCC 10245) was purchased from American Type Culture Collection. Yeast extract, bacto-peptone (becton, Dickinson and Company), di-sodium phosphate, and citric acid were used for preparing the appropriate culture media. All the reagents were purchased from Sigma unless otherwise stated. Chitosan was obtained from Vanson Company. Methanol (99%) was used to wash the chitosan samples. Hydrochloric acid solution, 0.1 N (N/10) (Certified) and sodium hydroxide solution 0.1 N (N/10) (Certified), both obtained from Fisher Chemical, were used in acid-base titrations to determine the degree of deacetylation (% DD) found to be 90%. Sodium acetate anhydrous (Fused Crystals/Certified ACS) was obtained from Fisher Chemical. All reagents and polymers were used as received unless stated otherwise.

#### Methods

Cell culture and preparation of bacterial cellulose film. Komagataeibacter xylinus (ATCC 10245) was grown on a media consisting of 2.5% p-mannitol, 0.5% yeast extract, and 0.3% bacto-peptone dissolved in deionized water at initial pH 6.5. All the cells were pre-cultured in a test tube for 1 day at 110 rpm at 28 °C to increase the population of bacteria for the main inoculation. The suspension passed through 16 layers of gauze as inoculums for future cultivations. The BC film formed in the test tube was then inoculated into a 500 ml Erlenmeyer flask containing 100 ml of culture broth. Static cultivations were carried out at 28 °C over 14 days.

The BC film after the cultivation period was sterilized with washing with a 1 N NaOH solution at 80  $^{\circ}$ C for 90 min to dissolve remaining bacteria. The samples were rinsed thoroughly with deionized water. The purified BC was dried to constant weight at 80  $^{\circ}$ C and then weighed to ascertain the production of BC. Finally, the purified wet cellulose film was stored in distilled water at 4  $^{\circ}$ C for future experiments.

Characterization and deacetylation of chitosan. The obtained chitosan having a degree of deacetylation of 70% was further deacetylated by an alkaline treatment using 50 wt% NaOH per a reported procedure. Briefly, 100 g of chitosan was suspended in 1.1 l of 50% aqueous NaOH solution and stirred at 110 °C for two hours. Nitrogen gas was continuously sparged into the mixture to avoid degradation of sample. The mixture was cooled down to 80 °C and then rinsed in water until pH 7 after which it was air-dried for two days. Subsequently, it was washed with methanol to remove impurities. Lastly, the sample was subjected to rinsing with water until neutral pH and air-dried over two days.

The degree of deacetylation was measured by acid-base titration. A detailed explanation is provided elsewhere.<sup>31,32</sup> Briefly, a fixed amount of chitosan was placed in a 250 ml beaker and dissolved by adding 10 ml of 0.1 N hydrochloric

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acid. Next, 90 ml of distilled water was added to the beaker in addition to a few drops of phenolphthalein as an indicator. The conductivity meter (Orion conductivity cell, model 013030) was immersed into the solution while the solution was kept at constant stirring. The solution was then titrated with 0.1 N NaOH with by a 10 ml burette. The readings were noted after

NaOH with by a 10 ml burette. The readings were noted after adding the NaOH solution and plotted as volume of NaOH solution *versus* conductivity.

The molecular weight of chitosan was determined by viscosity measurements using a Ubbelohde viscometer. A series of dilute solutions with increasing chitosan concentrations were prepared by dissolving chitosan in a solution of 0.2 M CH<sub>3</sub>COOH/0.1 M CH<sub>3</sub>COONa. A solution of 10 ml was then placed into a Cannon–Ubbelohde semi-micro viscometer (size 75 J-134) and the instrument was immersed into a constant water bath at 30 °C. It was kept for 20 min to equilibrate before the readings were recorded. The viscosity average molecular weight ( $M_v$ ) was calculated from the measured intrinsic viscosity [ $\eta$ ] and found to be 8.2 × 10<sup>5</sup> by use of the Mark–Houwink–Sakurada equation, [ $\eta$ ] =  $kM^a$  where k = 1.64 × 10<sup>-30</sup> × (% DD) and a = -1.02 × 10<sup>-2</sup> × (% DD) + 1.82.<sup>32</sup>

Preparation of BC and BC-chitosan composites. A wet BC film was placed on a glass rod for ten minutes to remove excess water.<sup>33</sup> The BC film was placed inside a polypropylene tube (dimensions: 10 mm in diameter and 40 mm in length) that was filled up to the 20 mm mark with deionized water.<sup>34</sup> The polypropylene tube was placed inside an insulating Styrofoam container so only the bottom surface of the polypropylene tube was exposed.<sup>35</sup> The Styrofoam container that housed the tube was allowed to float on liquid nitrogen (-196 °C) for 15 minutes (Fig. 1). During the process, the BC film inside the tube was instantly vectorially aligned from the bottom to the top *via* a ice crystal-templated growth. The solidified BC was then lyophilized at -40 °C for 2 days to remove ice crystals by sublimation.

To fabricate a composite scaffold of chitosan, a wet BC film was placed between two filter papers to remove excess water. Then the BC film was immersed in 1% chitosan solution (previously dissolved in a 1% aqueous acetic acid solution) at  $50\,^{\circ}\text{C}$  for 24 hours. Afterwards, the BC film was withdrawn from

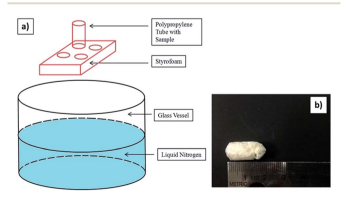


Fig. 1 Schematic procedure of preparing porous sample: (a) the sample is placed into a polypropylene tube which can contact with the liquid nitrogen in the glass vessel to induce a bottom-to-top freezing event instantly. (b) A digital photograph of the final ice-templated/freeze-dried sample.

the chitosan solution beaker and excess chitosan solution was removed by dabbing with filter paper. Afterwards, the BC film that was immersed in the chitosan solution was put in polypropylene tube and vectorially aligned frozen in a similar manner to above (*vide infra*). Thus, BC–Ch-*x* was prepared where BC is bacterial cellulose, Ch is chitosan, and *x* represents the wt% of chitosan solution. The BC and BC-Ch-*x* scaffolds after freeze-drying were removed and stored in silica gel desiccators for further characterization.

#### Characterization

Scanning Electron Microscopy (SEM). Oven-dried and freezedried samples of pure bacterial cellulose and BC-chitosan scaffolds were analyzed by SEM (Field Emission Inc, FEI Phenom). The samples were mounted on a metal stub and coated with a layer of gold approximately 100 Å thick via a gold sputtering machine to reduce charge interruptions. Samples were viewed at magnification between 5000-20 000 times original sizes. Those images were used to evaluate fiber diameter and consistency. Revolution software and ImageJ software were used to process the SEM images to determine the surface morphology and the cross-sectional image of BC and composite scaffolds. Fifty readings were measured from two different SEM images and the readings were averaged to determine the mean fiber diameter and fiber distribution. The porosity of the scaffolds was analyzed based on a literature method using ImageJ software. The SEM images were first converted into gray scale images and then the threshold value was adjusted to observe the first layer of the porous structure and measure the number of filled versus unfilled pixels.

**Absorption properties.** The absorption properties were determined by placing the cylindrical dried sample of 8 mm diameter  $\times$  6 mm height in phosphate buffer saline (PBS) for 24 hours at 37 °C to simulate body fluid. The cylindrical wet samples were taken out and the excess surface fluid was removed by placing the samples on filter paper before weighing.

The percent adsorption was calculated using following equation:

Adsorption(%) = 
$$\frac{W_t - W_0}{W_0} \times 100\%$$
 (1)

where  $W_0$  and  $W_t$  are the weights of the sample before and after the absorption test, respectively. All the results are the mean values from four samples.

**Mechanical properties.** Compression moduli of the scaffolds were tested on an Instron Model 5544 at a cross-head speed of 2 mm min $^{-1}$  at room temperature and collected using the Bluehill<sup>TM</sup> version 1.00 software. The testing machine was equipped with a 100 N load cell. Samples with an 8 mm diameter  $\times$  6 mm height dimension were tested in both the dry and wet states. In the wet state, the scaffolds were immersed in PBS and incubated for 24 hours at 37  $^{\circ}$ C to simulate *in vivo* conditions. Before the compression test, excess surface fluid was removed by blotting the scaffold with filter paper. Four samples were used to obtain mean values.

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Fourier Transform Infrared Spectroscopy (FTIR). FTIR spectra of samples of pure BC and BC-chitosan were obtained using a Nicolet 510P FTIR spectrometer. Scans were completed between 4000-400 cm<sup>-1</sup> with 16 convolutions at a resolution of 4 cm<sup>-1</sup>. A total 64 scans were completed for each sample. The baselines of all the samples were corrected and samples were analyzed using Omnic software.

X-ray diffractometry. X-ray diffraction (XRD) patterns of BC and BC-composites were recorded using a Siemens type F X-ray diffractometer equipped with Ni-filtered Cu Ka radiation at a wavelength of 1.54 Å. The X-ray generator voltage and current were 40 kV and 20 mA, respectively. The diffractograms were recorded at  $2\theta$  angular scanning angles of 5-40° with a 0.05 step size. The X-ray data were used without baseline correction. The crystallinity indices (CI) were obtained from X-ray diffraction data:

$$CI = \frac{I_{(at \ 200)} - I_{(amp)}}{I_{(at \ 200)}} \times 100$$
 (2)

where  $I_{(at 200)}$  is the intensity at (2 0 0) peak at 22.93° and  $I_{(amp)}$  is the minimum peak between (1 1 0) and (2 0 0).

Brunauer-Emmett-Teller (BET) surface area. The surface areas of pure and BC-chitosan scaffolds were measured using Autosorb-1C instrument equipped with the ASWin software package (Quantachrome Instruments, FL, USA). The samples were degassed at room temperature (25 °C) for 5 hours to remove moisture to ensure precision and accuracy in all of the experiments. Nitrogen adsorption/desorption isotherms were performed on every sample. The experiments analyzed surface area (m<sup>2</sup> g<sup>-1</sup>) by N<sub>2</sub> adsorption at 77 K using multipoint Brunauer-Emmett-Teller theory.

## Results and discussion

### Preparation and characterization of bacterial cellulose nanofibers

The original random morphology of BC film was aligned by a vectorially aligned freeze-drying method to assert a level of biomimetic control of scaffold architecture for tissue

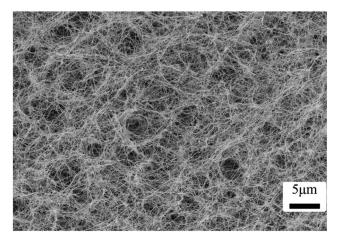


Fig. 2 SEM image of BC produced by Komagataeibacter xylinus.

engineering. Because BC hydrogels upon mechanical loading cannot recover their original shape, chitosan was added as a reinforcing polymer within the BC network in addition to providing useful biological properties. SEM was used to observe the BC film after purification and drying. Fig. 2 shows the SEM image of the macroscopic sponge-like material that at the microscopic level displays an interconnected nonwoven nanofibrous network. The average diameter of these nanofibers is 64  $\pm$  15 nm.

The freeze-drying process can freeze the water in the cellulose network and dry it (remove solvent) below the glass transition or melting temperature of water (sublimation). The removed frozen solvent consequently leaves behind void spaces that characterize the resultant porous structure.34 Even when various amounts of chitosan were added and freeze-dried, all the morphological analyses by SEM revealed a highly porous structure (Fig. 3).

Fig. 3a-d provide cross sections of pure BC and BC-chitosan scaffolds. Pure BC (Fig. 3a) did not possess pores that were only circular; rather they were circular and ovaloid within an interconnected network. Pores were created in the cross-sectional area perpendicular to the freezing direction. The channel-like porosity was created in the sample in the cross section parallel to the freezing direction, a phenomenon shown in Fig. 3b. The channel pores are connected with short nanofibrils in which the red arrow indicates the direction of (aligned) porosity in the sample. It should be noted that this alignment was not present in the original bacterial cellulose nanofibrous network (Fig. 2). The freeze-drying method requires that the bottom surface first contacts the liquid nitrogen to induce nanofibers to stretch and display random morphology because the ice crystals nucleate and propagate from underneath. Thin film formation was observed on the outer scaffold surface that was in contact with the inner wall of the polystyrene/glass tube. The film shows nonwoven nanofibrous morphology; however, it does not display any pore formation, a result that has been documented in other reports.9,35

The SEM analyses were done to determine the effect of chitosan concentration on the morphological changes inside the scaffold. Fig. 3c shows the interconnected pores within a threedimensional network in which the structure of the pure BC nanofibrils was lost due to the incorporation of chitosan. The inner section of BC-Ch-1% demonstrates an alignment of the porosity. The channel like formation can be attributed to a vectorially aligned freezing of the sample as the porosity developed from the bottom to the top of the sample. BC with 1 wt% chitosan (Fig. 3a) was even more porous than the other two samples that had chitosan concentrations of 1.5 and 2 wt% (Fig. 3e and g). They displayed an interconnected porous network within a three-dimensional network similar to the pure BC scaffold. The distance between two channels was largely due to chitosan penetration inside the BC pores to form a film on the individual surfaces. It was found, not unlike what has been previously reported, that the surfaces became more compact when chitosan concentration increased.<sup>27</sup> Mean pore diameter and porosity of BC and composite scaffolds are listed in Fig. 4a and b.

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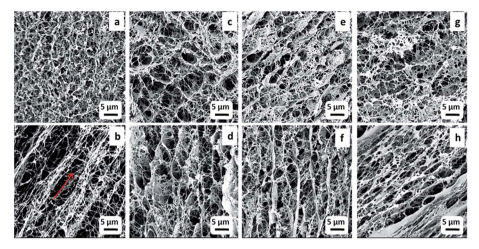


Fig. 3 SEM photographs of freeze-dried pure BC and BC-Ch composites possessing different levels of chitosan (a, b: pure BC; c, d: BC-Ch-1%; e, f: BC-Ch-1.5%; g, h: BC-Ch-2%). Top row is the cross section and bottom row is the inner wall. (all images are at 5000× magnification.)

As the concentration of chitosan increased to 1.5%, the distance between two channels became shorter because of chitosan aggregation inside the channels (Fig. 3f). The channels displayed a specific degree of alignment due to increased penetration of chitosan clearly witnessed at the 2 wt% loading mark (Fig. 3h) that induced layer formation on the BC film.

However, the channels were still interconnected by cellulose nanofibrils.

From the cross-sectional SEM images, mean pore diameters (Fig. 4a), and porosities (Fig. 4b), it appeared that the pores at 1 wt% chitosan were larger compared to pure BC. The mean pore diameter and porosity in the pure scaffold was 2  $\pm$  1  $\mu m$  and 75

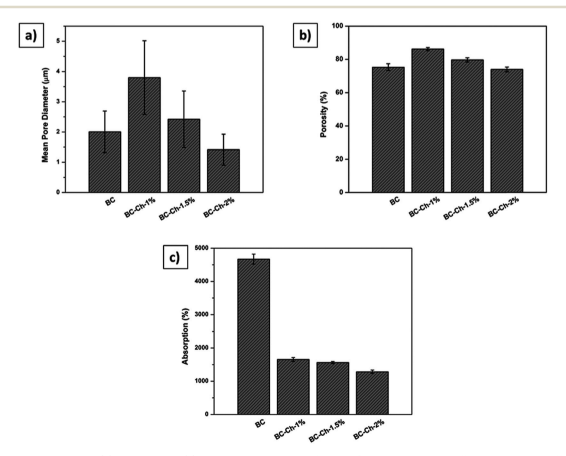


Fig. 4 (a) Mean pore diameter, (b) porosity, and (c) absorption properties of pure BC and composite scaffolds possessing different weight percentages of chitosan (1%, 1.5%, and 2%).

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 $\pm$  2%, respectively, whereas, for BC–Ch-1%, it was 4  $\pm$  2  $\mu m$  and  $87 \pm 1\%$ . The increase in pore diameter and porosity was due to increased concentration as well as the viscosity of the overall sample that facilitated the BC film to rearrange themselves into highly oriented pores and channels. Water acted as a solvent during BC scaffold preparation. Many small size ice crystals tended to form during the freezing process to induce small pores and reduce the porosity in the final sample. The overall porous structure was reduced as the chitosan concentration increased from 1 to 2 wt% as shown in Fig. 3c, e, g and 4a, b. The pore diameter and the porosity of BC-Ch-1.5% was 2.5  $\pm$  1  $\mu$ m and 80  $\pm$  1%, while for BC–Ch-2%, it was 1.4  $\pm$  1  $\mu m$  and 74  $\pm$ 2%, respectively. This may be because increasing chitosan concentration interfered with ice crystal nucleation during the freezing step. Water is sequestered, while the "spectator" chitosan macromolecules prevented the water molecules from rearranging and forming larger ice crystals during a rapid freezing process. Another reason for the random and reduced pore distribution might be due to the viscosity of the chitosan solution. As the concentration increased, the viscosity of the solution also increased. Thus, during the freezing step, a higher viscosity sample prevented water molecules from rearranging and thus hindered ice crystal nucleation and gave rise to random crystal shapes.35 Therefore, as the concentration of chitosan increased, the distance between two juxtaposing channels decreased as did the pore size. The 2 wt% chitosan sample had a random distribution of channels with relatively small pore size. However, all the scaffolds retained a porous structure. This was observed in a previous effort.35 The current result indicates that adjusting chitosan concentration may control the nano-structure of the BC/chitosan scaffold. The reduced pore size may also be beneficial to retain synthesized matrix molecules.36 The porous structure and the interconnection of fibrils are clearly also critical for cellular attachment and proliferation, thus making it an attractive scaffold for tissue engineering.

#### **Absorption properties**

The absorption properties of pure BC and associated composite scaffolds were tested by immersion in PBS at 37 °C for 24 hours. The results showed that pure BC has the highest absorption percentage because of the abundance of hydrophilic functionalities. Additionally, because the scaffold dimensions are fixed before adsorption, the scaffold tended to swell, accommodating as much as water inside the porous network as possible and thereby increasing its volume. During absorption, the hydroxyl

groups attracted water because of hydrogen bonding forces. When chitosan (1%, 1.5%, and 2%) was introduced into the scaffold, water sorption reduced as shown in Fig. 4c. The water penetration was limited due to bond formation between chitosan -NH2 groups and the hydroxyl groups of BC; thus, the original porous structure was retained because chitosan limited the scaffold swelling given that it has a tendency to retain its morphology and hence the original porous structures/ dimensions. The results were similar to former reports that demonstrated that intermolecular hydrogen bonds affected the interaction between water and polymer chains directly.37,38 Though the absorption properties decreased with inclusion of chitosan, this may have been offset by the favorable mechanical and biological activities of chitosan. Very fortuitously, chitosan may act as a structural stabilizer for optimizing tissue engineering applications.

#### Mechanical properties

Compressive properties were tested in both the dry and wet states. The wet studies were specifically executed as close to *in vivo* conditions as possible. Table 1 shows the compressive moduli of pure BC and composite scaffolds.

The mechanical strength of pure BC was highest,  $9.51\pm0.23$  MPa, among all the scaffolds in the dry state. This is due to the reduced pore size and porosity of the scaffold. During the initial loading, the fibers had better chances of aligning themselves between the channels of the nanofibrils. This rearrangement might be the reason for the higher modulus of pure BC. The addition of chitosan tends to decrease the compressive modulus, but the values were approximately similar for BC containing 1% and 1.5% chitosan. The addition of chitosan increased the structural integrity and formed networks within the BC nanofibrils, except for the scaffold containing 2% chitosan. This reduction in dry state might be due to the coating of individual BC nanofibrils that upon drying become brittle and thus under normal loading display lower compressive strengths.

The compressive properties of the scaffold have mixed results in the wet state. For pure BC, the modulus increased 3.6× compared to its dry state. This is because the BC has a strong affinity for water that tends to fill the empty pores thereby swelling and increasing the volumetric dimensions of the scaffold and directly affecting the mechanical properties. In addition, the scaffold undergoes many physical and chemical changes during water absorption that are highest for pure BC. The water content helps the nanofibers slide over each other such that upon mechanical loading can rearrange. Though the

**Table 1** BET surface area, crystallinity index, and compressive modulus of pure BC and composite scaffolds with different weight-percentages of chitosan in both dry and wet states

Sample name	Surface area $(m^2 g^{-1})$	Crystallinity index (%)	Dry compressive modulus (MPa)	Wet compressive modulus (MPa)
BC	201	89.45	$9.51\pm0.23$	$37.32\pm1.07$
BC-Ch-1%	29.5	83.57	8.04 $\pm$ 0.47	$0.58\pm0.17$
BC-Ch-1.5%	22.06	83.94	$9.13 \pm 0.40$	$0.82\pm0.15$
BC-Ch-2%	19.85	79.46	$4.74\pm1.00$	$\textbf{9.64} \pm \textbf{0.46}$

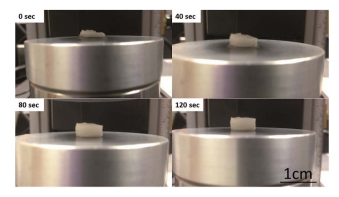


Fig. 5 Photographs of the BC–Ch-1% scaffold (original dimensions: 8 mm diameter  $\times$  6 mm height) that show a compressed form (at 0 s) that undergoes shape recovery after a two-minute interval to virtually its original dimensions.

pure BC possesses high mechanical strength, the scaffold can recover its original shape even after 2 hours at room temperature.<sup>39</sup> The compressive moduli for BC-Ch-1% (0.58  $\pm$  0.17) and BC-Ch-1.5% (0.82  $\pm$  0.15) were found to be much lower than in their dry state. Because the BC-Ch-composite display reduced absorption and chitosan shows a tendency to retain its original shape, the scaffold was lubricated between the chitosan and BC nanofibers to reduce internal friction, ultimately reducing the compressive strength.40 The wet compressive modulus of the BC-Ch-2% was found to be higher (9.64  $\pm$  0.46) than in its dry state (4.74  $\pm$  1.00), despite its lower absorption properties. This might be due to the chitosan having a high degree of deacetylation (90%), which means it has more amino groups. Moreover, the chitosan had an entanglement concentration of 1.85 wt% (it tends to form a charged complex in acidic solution) that surpassing causes an increase in intermolecular hydrogen bonding and restriction of individual polymer chains.

In addition, the lower pore size facilitates a viscous chitosan charged complex inside the porous network upon uniform compressive loading, ultimately increasing its compressive strength.<sup>31</sup> All the wet scaffolds showed higher compressive moduli at the same order of magnitude as native human

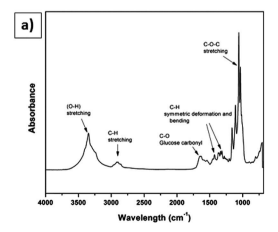
cartilage (0.677  $\pm$  0.198).<sup>41</sup> All the scaffolds containing chitosan showed 100% recovery after two minutes of removing the load (Fig. 5). This type of scaffold is useful for waste removal during compression and diffusion and exchange of nutrients during the recovery of scaffolds given that natural articular cartilage lacks the necessary blood vessels for such functions. Additionally, previous studies confirmed these potential applications: the porous scaffolds and three-dimensional scaffolds having high compressive strength could be used for meniscus regeneration, culturing primary human fibroblasts, periosteal cells, osteoblasts, and other cellular tissues.  $^{42-45}$ 

#### Surface area measurements

From inspection of the SEM images, pure BC scaffold has a more open structure than the BC-chitosan scaffold. The BET technique was used to obtain the surface area to give a total surface area for pure BC of 201 m<sup>2</sup> g<sup>-1</sup> (Table 1). However, the total surface area drastically reduced with incorporation of chitosan. The total surface areas were 29.5 m<sup>2</sup> g<sup>-1</sup> for BC-Ch-1%, 22.06 m<sup>2</sup> g<sup>-1</sup> for BC-Ch-1.5%, and 19.85 m<sup>2</sup> g<sup>-1</sup> for BC-Ch-2%. Solution facilitates chitosan penetration inside the BC interconnected fibrils to block empty pores. Moreover, a higher content of chitosan decreased pore size as clearly seen from SEM images in Fig. 3. The fibrils having a close arrangement became dense and compact ultimately leading to a lower surface area. Another reason for the reduction in surface area was due to a strong interaction of chitosan molecules with the BC chains to form a dense network of fibrils and ultimately close the empty spaces among the BC nanofibrils.46

#### XRD analysis of prepared scaffolds

Fig. 6b shows the recorded X-ray diffractogram. The XRD of BC films were recorded in parallel with the X-ray beam. Three main peaks were observed in both spectra. The  $14.80^{\circ}$  was assigned to  $(1 - 1 \ 0)$ , the  $16.6^{\circ}$  was assigned to  $(1 \ 1 \ 0)$ , and the  $22.93^{\circ}$  was assigned to  $(2 \ 0 \ 0)$  crystallographic planes. These results confirm the presence of cellulose I in the BC films.  $^{26,27,33}$  The crystallinity index of BC was 89.45%. Thus, the data gleaned from the specific chemical peak position and the crystal



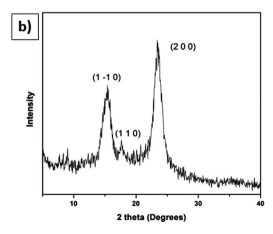


Fig. 6 Characterization of BC membrane used for making scaffolds: (a) FT-IR spectra, (b) XRD diffractogram.

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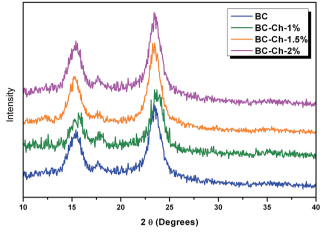


Fig. 7 XRD patterns of pure BC and at different weight-percentages of chitosan.

structure from the combined results of FTIR and XRD confirm the presence of cellulose I production by *Komagataeibacter xylinus*.

From Fig. 6b, the presence of cellulose I in the BC cultured from mannitol as a carbon source was confirmed. One method employed to measure the crystallinity of the cellulose I was peak height from X-ray diffraction. This technique measures the difference in intensity of the highest peak (2 0 0) and the lowest peak that is attributed to the amorphous region. Fig. 7 shows the XRD pattern and Table 1 shows the crystallinity indices of pure BC and composite scaffolds, respectively.

For the pure BC and composite scaffolds, three major peaks were found at 14.80° that are assignable to (1-10), 16.6° to (110), and 22.93° to (2 0 0) crystallographic planes. These findings confirm the presence of cellulose I in the BC film. The crystallinity of the composite scaffolds decreased with an increase in chitosan concentration. The crystallinity index of pure BC was 89.45% that reduced to 83.57% for BC-Ch-1%, 83.94% for BC-Ch-1.5%, and then further reduced to 79.46% for BC-Ch-2%. This indicates that the high crystallinity of the pure BC can be reduced by addition of chitosan. Similar results have been found in which the crystallinity index of cellulose I reduced with the addition of additives.26,27,33 This decrease was due to penetration of chitosan inside in the BC inter-fibrillar network. The chitosan molecules interact with the BC network through intermolecular hydrogen bonding to significantly alter the BC network by creating packing disruptions that lead to a reduction in the crystallinity.

#### FTIR analysis of prepared scaffolds

The FTIR results (Fig. 6a) confirmed the presence of stretching vibrations for O–H and C–H at 3347 cm $^{-1}$  and 2897 cm $^{-1}$ , respectively. A sharp and intense band appears at 1651 cm $^{-1}$  represents the glucose carbonyl (C=O) of cellulose. Two important bands appeared at 1427 cm $^{-1}$  and 1364 cm $^{-1}$  characteristic of the symmetric deformation and bending vibration of C–H, respectively. The sharp band at 1033 cm $^{-1}$  represents C–O–C stretching.  $^{27}$ 

Fig. 8 shows the IR spectra of pure BC and BC-chitosan composite scaffolds with different weight percentages of chitosan.

The BC scaffold results confirmed the presence of the stretching vibrations of O–H and C–H at 3346 cm<sup>-1</sup> and 2896 cm<sup>-1</sup>, respectively. An intense band appears at 1632 cm<sup>-1</sup> represents the glucose carbonyl stretch of cellulose. Two more bands at 1427 cm<sup>-1</sup> and 1370 cm<sup>-1</sup> showed the symmetric deformation and bending vibration of CH, respectively. The band over 1035 to 1033 cm<sup>-1</sup> represents the C–O–C stretching (Table 2).

The spectral analysis of BC–Ch shows all the characteristic bands of BC. For the BC–chitosan scaffold, a broad band appears at 3245 cm<sup>-1</sup> (combination of O–H stretching and N–H stretching). C–O-stretching of BC–Ch decreased due to increased hydrogen bonding in the composite and more interactions between BC and Ch. The –NH<sub>2</sub> deformation vibration at 1555 cm<sup>-1</sup> indicates the presence of chitosan molecules inside BC. The C–N stretching peak at 1409 cm<sup>-1</sup> was due to the presence of primary amides. All the data indicate that the chitosan molecules interact with the BC through intermolecular hydrogen bonding networks and penetrate well within the BC fibrils.<sup>27</sup>

#### Mechanism of vectorially aligned freezing

The freezing step is the most critical step for creating a porous scaffold. Because BC contains water (solvent), these solvent molecules are converted in form to ice crystals during freezing. The randomly distributed web of BC nanofibers (Fig. 2) is vectorially aligned (in the direction of the freezing – from low to high temperature gradient) by freezing and subsequent freezedrying. The BC nanofibrillar network becomes disentangled during liquid nitrogen-induced freezing process by the ice-templating effect; in other words, the nanofibrils follow

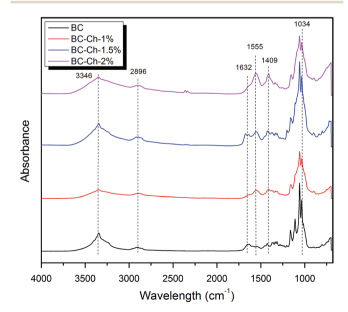


Fig. 8 FTIR spectra of pure BC and at different weight-percentages of chitosan.

Table 2 Spectral peak assignment of pure BC and BC-chitosan (1

wt%) scaffolds prepared by vectorially aligned freeze-drying

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Pure BC sample wave number (cm <sup>-1</sup> )	Functional group	BC-Ch sample wave number (cm <sup>-1</sup> )	
3346	O-H/N-H	3345	
2896	С-Н	2894	
1632	C-O	_	
_	$NH_2$	1555	
1427	C-H deformation	_	
1370	C-H bending	1373	
_	C–N stretching (primary amide)	1409	
1035	C-O-C	1034	

vectorial alignment (growth) of the ice nuclei by hydrogen bonding to the water as it freezes. The liquid nitrogen induced rapid formation of ice nuclei which consequently propagated in the direction of freezing.34 The ice-templating effect thus had a profound impact on the nanofibrillar alignment. The samples were vectorially aligned by the liquid nitrogen freezing of the water solvent. More specifically, the water-ice crystals grew parallel to the direction of freezing (i.e., from the low temperature to high temperature gradient). After drying, the scaffold displayed a highly vectorially aligned channel-like character that possessed a huge number of small voids (hence, its highly porous character). In sum, rapid cooling by liquid nitrogen of the water within the BC nanofibrils caused the formation of many small sized ice nuclei to result in a scaffold of dimensionally small pore size. Fig. 9 provides an illustration of the process that allows for ready visualization of how the wet BC film was frozen in one direction.

The ice crystals started to grow from the bottom to the top of the sample due to differences in the temperature gradient. The random nonwoven BC film fibrils were stretched and pulled by hydrogen bonding forces toward the ice crystals and consequently, they both aligned themselves parallel to the freezing direction. Interestingly, the chitosan macromolecules tended to

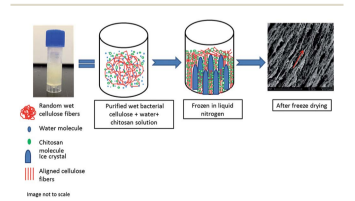


Fig. 9 Mechanism of orienting randomly distributed bacterial cellulose/composite scaffold by the vectorially aligned freeze-drying method. During freezing, ice crystals grow in freezing direction (vertically) that exclude the bacterial cellulose fibrils and chitosan molecules and aligns the nanofibrils in the spaces between the ice crystals (parallel to direction of freezing).

separate from the frozen solvent likely due to their milder hydrophobicity relative to the cellulose (see representation using green dots in Fig. 9). These chitosan macromolecules then aggregated and aligned themselves around the ice crystals and into the pores of the BC fibrils.

In general, the method is extremely simple, useful, and powerful for obtaining aligned porosity from a randomly nonwoven cellulose nanofibrillar network. The preparation of aligned porous materials with micrometre-sized pores is of particular importance for applications such as tissue engineering that require subsequent modification with entities such as biological cells, which can attach along the length of the channels of aligned porous scaffolds. Also, the synthetically aligned porous structures may also be used to mimic natural aligned materials such as bone.<sup>23</sup>

## Conclusions

Pure BC and BC-chitosan scaffolds were prepared by vectorially aligned freeze-drying. The incorporation of chitosan inside the BC network increased the interaction between BC nanofibrils and chitosan macromolecules that after freeze-drying displayed a vectorial alignment of fibrils along the freezing direction. Morphology considerations such as pore size and distance between channels could be altered by adjusting chitosan concentration; however, the crystallinity index tended to decrease with increasing chitosan concentration likely because of its well known strong intermolecular interactions with the cellulose backbone. However, chitosan penetration inside BC fibrils facilitated the formation of a dense network that allowed the scaffolds to maintain their structural integrity, display excellent shape recovery, and provide superior overall mechanical properties relative to pure BC after load removal, a finding that was very comparable to native human cartilage. These unique properties of the current prepared scaffolds may specifically be tailored for cartilage tissue regeneration.

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