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Biomimetic composite scaffold from an *in situ* hydroxyapatite coating on cellulose nanocrystals

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A novel nanocomposite scaffold was developed by homogeneous deposition of hydroxyapatite (HAP) on a cellulose nanocrystals (CNCs) matrix suspended in a simulated body fluid (SBF). By adjusting the pH of the SBF, the HAP content in the nanocomposite could be controlled between 15 wt% and 47 wt%. Physical and chemical characteristics of the nanocomposites were analyzed by SEM, FTIR, XRD, SAED, and TEM, which confirmed the successful incorporation of HAP onto the CNCs. The nanocomposites were then freeze-casted into porous scaffolds by different solidification technologies (*i.e.*, directional freezing (DF), plunging in liquid N₂ (PL) or in a -20 °C freezer (FZ)) followed by lyophilization. Compression testing of the HAP/CNCs foams indicated that DF caused significant improvement in mechanical properties due to the specific orientation and anisotropic porous structure compared to conventional freezing methods such as PL and FZ. Moreover, the scaffold with high HAP content exhibited improved mechanical and thermal properties, which holds potential for application in bone tissue engineering.

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1. Introduction

Materials mimicking the natural composite of bone in both chemical and structural aspects have been studied for decades due to its extraordinary mechanical properties and biological functionality.¹ Inspired by the intrinsic relationship between hydroxyapatite (HAP) and human bone, different types of 3D scaffolds have been developed in order to remedy the drawbacks of currently applied bone substitutes.^{2,3} Human bone is a hierarchically organized composite made up of collagen fibrils and apatite crystals (hydroxyapatite).⁴ The primary role of HAP in human bone is its osteoconductivity in which osteoprogenitor cells can adhere, migrate and finally form new bone.⁵ HAP in bone is precipitated between the oriented collagen and thus forms mineralized collagen fibers with a hierarchical structure.⁶ This type of structure gives the bone great mechanical properties.⁷ Many polymeric materials, like polyvinyl alcohol (PVA), poly(lactic acid) (PLA) and chitosan have been combined with

HAP to enhance their osteoconductivity and hemocompatibility.⁸⁻¹⁰ However, these approaches have not yet fully addressed all the requirements needed for successful bone scaffolds due to their inherent drawbacks like poor mechanical properties and lack of flexibility,¹¹⁻¹³ and research is ongoing in several fields, including on the use of several alternative materials such as nanocellulose.¹⁴

Cellulose nanocrystals (CNCs), a strong and promising natural material derived from native cellulose, have attracted tremendous interest due to their excellent physical and chemical properties and hold much potential for conversion into bio-based materials that are rigid, biocompatible, non-immunogenic and renewable.¹⁵ These rod-like crystalline cellulose structures are typically 5–20 nm in diameter and several hundred nanometers in length.¹⁶ It has been reported that the CNCs have an average Young's modulus around 130 GPa, even stronger than steel.¹⁷ In addition, CNCs are highly hydrophilic with a multitude of hydroxyl groups available, which can offer binding sites for bone cells and make it superior to other materials such as collagen and silk fibroin.¹⁸ Because of these excellent properties, CNCs have been successfully applied in many fields, such as drug delivery systems, tissue substitutes, hydrogels, conductive films, filtration membranes, and in composites as a reinforcement agent to enhance their physical properties.^{19,20} Although CNCs are not capable of being degraded in the human body, it is highly biocompatible which will not cause any inflammatory response after implanting.²¹ The use of CNCs associated with HAP as a template for developing bone substitute is a field of growing

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interest. Kumar *et al.* (2014) reported a polyvinyl alcohol/HAP/CNCs composite, showing enhanced mechanical property as with the incorporation of CNCs.²² Through a sol-gel method, Herdicia-Lluberes *et al.* (2015) fabricated a CNCs/HAP pellet composite which could promote the proliferation of the osteoblast cells.²³ Moreover, Saska *et al.* (2011) conducted *in vivo* test with a novel bacteria cellulose/HAP membrane, and results showed that the membrane could accelerate the new bone formation.²⁴ These results suggest a promising perspective for using this composite in bone tissue engineering.

However, almost all the studies concerning CNCs/HAP composite were prepared by just blending HAP and CNCs, resulting in a heterogeneous mixture with relatively poor mechanical properties. During the preparation of nanocellulose by sulfuric acid hydrolysis, the surface of such CNCs was activated by the introduction of sulfonic groups. It was hypothesized that ionic groups could be induced to precipitate on CNCs by electrostatic interactions if such activated CNCs was put into a solution containing various ionic groups. Thus, a novel *in situ* HAP coating on individual CNCs fiber by immersing into a simulated body fluid (SBF) containing calcium ions and phosphate groups was developed herein this study. The content of HAP particles was controlled by adjusting the pH of the solution. Scanning electron microscopy (SEM), transmission electron microscopy (TEM), X-ray diffraction (XRD), selected area electron diffraction (SAED) and Fourier transform infrared spectroscopy (FTIR) were used to confirm the existence of such HAP coating. Moreover, the HAP coated CNCs nanocomposite was then freeze-casted into porous scaffolds by different freezing technologies, in which the directional freezing formed an orientated structure and was anticipated to improve the mechanical property.

2. Experimental section

2.1. Materials

CNCs purchased from the University of Maine were prepared by sulfuric acid hydrolysis of wood pulp, possessing a dimension of 5–20 nm in width and 150–200 nm in length. The obtained product was an aqueous slurry (12.1 wt%) and the sulfur content of the CNCs was 0.95 wt%. All the other chemicals of analytical grade were purchased from Fisher Scientific (Waltham, MA, USA) and used as received without further purification unless otherwise noted.

2.2. Fabrication of CNCs/HAP composite

A simplified SBF was prepared based on a previous report by Ishikawa *et al.* (2015).⁴ NaCl, CaCl₂·2H₂O, Na₂HPO₄·12H₂O, and HCl were dissolved in DI water so as to set the final Na⁺, Ca²⁺, Cl⁻ and HPO₄²⁻ concentrations to 141.0 mM, 2.5 mM, 165.0 mM and 1.0 mM, respectively. The CNCs aqueous suspension (1.044 g) was added to the prepared SBF (200.0 mL) in a flask with a calcium ion (in SBF solution) and sulfonic group (in CNCs) molar ratio of 13 : 1. The suspension was then stirred until no bulk CNCs particles were apparent and then transferred to an ultrasonic water bath for 30 min to evenly

disperse CNCs. The mixture was then adjusted to different pH values with tris(hydroxymethyl)aminomethane solution (50 g L⁻¹). Next, the suspension was put in an orbital shaker at 37 °C and 150 rpm for 1 h. This process of ultrasound and subsequent mild stirring was repeated three times. The solid contents in suspensions were then collected by centrifugation at 1705g for 10 min and washed with anhydrous ethanol three times to remove the residual salts. Finally, the CNCs/HAP composites were dried for two days in a conventional oven at 45 °C.

2.3. Preparation of porous CNCs/HAP scaffolds

The porous scaffolds were prepared by freeze-casting 5% (w/w) CNCs or CNCs/HAP suspensions solidified by directional freezing (DF), plunging in liquid N₂ (PL, -196 °C) and putting in a -20 °C freezer (FZ). This was accomplished by adding dry CNCs and CNCs/HAP sample in DI water under magnetic stirring for 2 h to obtain a homogeneous suspension with 5 wt% consistency. Then, ~3.5 mL of the solutions were transferred into 10 mL glass vials and frozen by FZ (-20 °C for 24 h), PL (-196 °C for 10 min) or DF (-196 °C for 10 min). In the case of DF, only the bottom of the vials contacted liquid N₂ and vertical heat transfer was prevented by insulation of the vials with polystyrene foam. Upon solidification, the samples were immediately put in a freeze drier (-80 °C and 12 Pa) for 48 h to obtain the porous scaffolds.

2.4. Characterization of nanocomposites

The surface morphology of the CNCs/HAP composites and the longitudinal side view of the porous scaffolds were observed employing a SEM (Zeiss EVO MA15, Germany) at an accelerating voltage of 5 kV. Before SEM analysis, the samples were put in a 70 °C oven to remove the moisture and then sputter coated with gold. The dimensions of CNCs or CNCs/HAP fibers were observed by TEM. Before the TEM analysis, samples were placed in DI water to obtain a 0.02% (w/v) solution. A drop of the diluted solution was then deposited on a carbon-coated copper grid to improve the dispersion of the CNCs. The residual water was removed with a filter paper from the periphery of the copper grid and dried in the air for 1 min. The specimens were observed using a Zeiss TEM (Zeiss Libra 200 MC, Germany) with an acceleration voltage of 200 kV. The lengths and widths of the CNCs and CNCs/HAP composite were analyzed with ImageJ software, and around 100 fibers were selected for characterization.²⁵ SAED patterns were obtained with the same TEM based on the selected areas in the TEM images. FTIR analysis was accomplished using a Nicolet 6700 spectrometer (Thermo Scientific, MA, USA). Spectral data was acquired in the transmittance mode with 16 scans at 4 cm⁻¹ resolution over the range of 4000–500 cm⁻¹. The XRD patterns were recorded using a D8 advanced instrument (Bruker, Germany) with Cu K α radiation at a voltage of 40 kV and a current of 30 mA. The scanning speed was 2° min⁻¹ within the 2 θ angular range of 5–60°.

Thermogravimetric analysis (TGA) was carried out using a Q5000 instrument (TA Instruments, New Castle, DE, USA). About 5–10 mg of the nanocomposites were analyzed from RT to 700 °C at a heating rate of 10 °C min⁻¹ under high-purity



nitrogen. The HAP content in the nanocomposites was also determined by TGA following a published methodology except for that technical air (79% N₂ and 21% O₂) was used to facilitate complete combustion of organic compounds in the samples.²⁶ The HAP contents were then calculated based on the initial and residual weights of the samples.

The porosity of different samples was determined according to a modified displacement method in a previous report.²⁷ In brief, the volume and weight of the cylindrical foams were measured as V_0 and W_0 , respectively. The volume was calculated based on the diameter and height of the cylinder determined by a digital Vernier caliper. Then the specimen was immersed into anhydrous ethanol for 48 h until saturation. The weight of the saturated foam was measured as W_1 . Finally, the porosity of the samples was obtained as the formula of $P = (W_1 - W_0)/(\rho \times V_0)$, where ρ is the density of the anhydrous ethanol (0.789 g mL⁻¹). Measurements were made on three samples of each foam, and the results represent an average value. Compression strength of different nanocomposite foams was determined with an INSTRON 5567 universal test machine (Canton, MA, USA) using a load cell of 500 N. The cylindrical foams were subjected to a constant compression rate of 0.1 mm s⁻¹ until 80% strain and the compression modulus was calculated from the initial linear region (5%) of the stress-strain curve. Samples were stored in a desiccator before analysis.

3. Results and discussion

3.1. Synthesis of CNCs/HAP nanocomposites

Simulated body fluid (SBF) is a solution containing inorganic ions similar to those in human blood plasma which includes Ca²⁺, Na⁺, PO₄³⁻ and Cl⁻ as its main ions. SBF has been widely used to evaluate the bioactivity of different materials by forming the apatite layer on its surface.¹⁸ Unlike conventional bone substitutes such as titanium and stainless steel alloys, HAP crystals formed in SBF are nanosized and tend to grow and elongate in the *c* axis (vertical orientation), the same as the orientation in real bones.²⁵ The original SBF solution developed by Kokubo (1991) is composed of various inorganic ions such as Na⁺, K⁺, Mg²⁺, Ca²⁺, Cl⁻, HCO₃⁻, HPO₄²⁻ and SO₄²⁻.²⁸ However, a recent study illustrated the presence of Mg²⁺, HCO₃⁻, SO₄²⁻ had only a slight effect on the formation and deposition of the HAP crystals.²⁹ Thus, a simplified SBF containing Ca²⁺, Na⁺, Cl⁻, and HPO₄²⁻ was adopted in this study and this is consistent with the literatures.^{4,30,31} The existence of a large amount of Na⁺ and Cl⁻ is necessary to equalize the ion strength. The SBF is a metastable solution, and no HAP crystals were observed under the mild pH condition if no high-affinity substrate was introduced.²⁹ In this study, CNCs were employed as a substrate to facilitate the deposition of HAP. To investigate the HAP coating, different pH values of 6.0 (acid condition), 7.4 (body condition) and 8.5 (basic condition) were chosen and the morphologies of CNCs after HAP coating were analyzed with SEM (as shown in Fig. 1). SEM images clearly indicated the successful synthesis of the CNCs/HAP composite. Specifically, the surface of CNCs was flat and neat (Fig. 1a), whereas after immersion in SBF at pH 6.0, a small amount of grain-like HAP particles could be seen

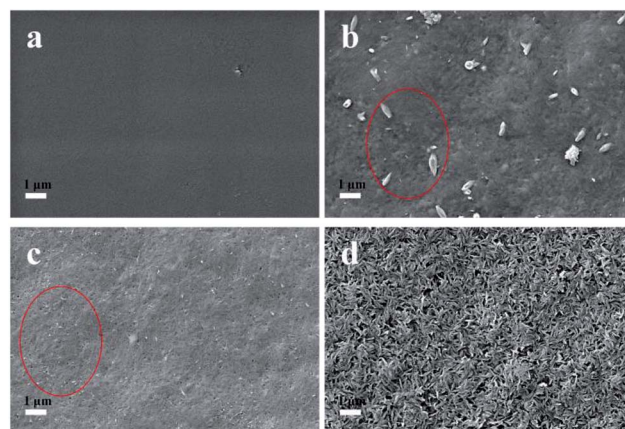


Fig. 1 The SEM images of CNCs before and after hydroxyapatite (HAP) coating. (a) Pure CNCs; (b) at SBF of pH 6.0; (c) at SBF pH 7.4; (d) at SBF pH 8.5.

embedded into the CNCs matrix (Fig. 1b). The deposition of HAP is primarily induced by the anionic functional groups (*i.e.*, sulfonic and hydroxyl groups) on the surface of CNCs that can initiate the nucleation of HAP and promote the coating of HAP on CNCs matrix.⁴ Increasing the pH to 7.4, the particles became smaller compared to the ones accomplished at pH 6.0. At a higher pH value of 8.5, small HAP particles aggregated due to the strong electrostatic interactions with CNCs and became predominant in the SEM image (Fig. 1d). It is known that HAP is soluble in an aqueous solution under acidic conditions. On the other hand, HAP tends to aggregate at high pH values. In our study, at the pH values of 6.0 and 7.4, only a few HAP deposits on CNCs were observed. Further increasing the pH to 8.5 significantly promoted the growth of the CNCs/HAP composites.

3.2. Structure characterization

3.2.1. FTIR and XRD analysis of the nanocomposites.

During the HAP coating process, the CNCs and HAP were combined by hydrogen bonds between sulfonic and hydroxyl groups or by the formation of ionic bonds (including [HO⁻-Ca²⁺ and [HSO₃⁻-Ca²⁺]) through the hydroxyl and sulfonic groups and calcium ions in HAP and CNCs.²² To analyze the structure changes during the HAP coating process, FTIR was employed, and results are shown in Fig. 2a. The FTIR spectrum of CNCs exhibited an adsorption at 1637 cm⁻¹, which is due to the O-H bending of the water adsorbed by cellulose.³² Typical peaks of CNCs can be observed at 1428 cm⁻¹ (CH₂ scissoring motion), 1369 cm⁻¹ (C-H bending), 1337 cm⁻¹ (O-H in-plane bending), 1316 cm⁻¹ (CH₂ wagging vibration), 1159 cm⁻¹ (C-C ring stretching) and 1054 cm⁻¹ (C-O-C stretching vibration).^{4,22} The band at 895 cm⁻¹ was assigned to the cellulosic β -glycosidic linkages.³³ Furthermore, the FTIR spectral data of the samples after the HAP coating showed all the typical signals of CNCs, indicating the existence of CNCs in the nanocomposite. Moreover, signals associated with functional groups in HAP emerged in the CNCs/HAP nanocomposites. The signal at 1020 cm⁻¹ was due to the asymmetric stretching of the P-O

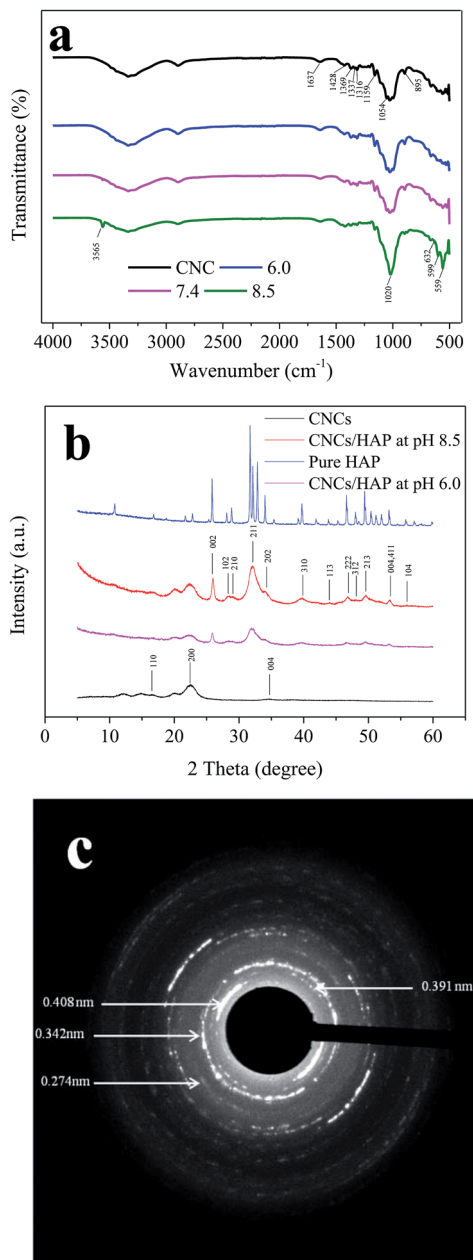


Fig. 2 FTIR (a), XRD (b) and SAED (c) analysis of the CNCs and HAP coated CNCs.

bonds in the phosphate group.³⁴ The signals at 599 cm^{-1} and 559 cm^{-1} were assigned to be the degenerated bending of the O-P-O bonds.³⁵ Also, signals at 3570 cm^{-1} and 632 cm^{-1} which represent the stretching and bending modes of hydroxyl groups in HAP could be also observed.³⁶ These results revealed that the layer coated on CNCs is actual HAP rather than normal calcium phosphate. The HAP was further confirmed by measuring the Ca/P ratio of the sample with an energy dispersive X-ray diffraction (EDX) attached to the SEM, which is a widely used technology to analyze the elements in HAP.^{18,34} EDX results showed that ratio of Ca and P in the nanocomposite was 1.76 which is consistent with the values in other reports and close to

the value of 1.67 in natural human bone, indicating the coated layer was HAP.^{18,37,38} This higher Ca/P ratio is likely to be caused by the generation of some other calcium contained salts like calcium carbonate and calcium hydroxide during the depositing process.

The crystalline structures of the pure CNCs, CNCs/HAP composites synthesized at pH 6.0 and 8.5, and pure HAP were analyzed by XRD (Fig. 2b). The characteristic peaks of cellulose at 16.7° , 22.5° , and 34.7° were assigned to the typical planes (110), (200) and (004) of cellulose, respectively.³⁹ These diffraction peaks can also be found in numerous nanocellulose composites.^{18,22} After coating, the samples exhibited typical characteristic peaks of HAP at $\theta = 26.0^\circ$ (002), 28.4° (102), 29.0° (210), 32.1° (211), 34.3° (202), 39.7° (310), 44.0° (113), 46.7° (222), 48.2° (312), 49.7° (213) and 56.1° (104), respectively. The peaks at 53.4° indicated planes (004/411) presented in the HAP.⁴⁰ All of these characteristic peaks of our synthesized HAP could be found in the pure HAP (Fig. 2b). In addition, it was found that the CNCs/HAP composites obtained at different pH values almost have the same crystalline structure, indicating that the pH has very limited effect on the structure of generated composites. The crystalline structure of the coating was further analyzed with SAED (Fig. 2c). The diffraction pattern exhibited several strong and broad diffuse rings, indicating the strong crystalline structure of the HAP. From the pattern, planes of (200), (002) and (111) could be clearly seen with $d_{200} = 0.408\text{ nm}$, $d_{002} = 0.342\text{ nm}$ and $d_{111} = 0.391\text{ nm}$, respectively. In addition, an interplanar spacing of 0.274 nm could be found in the diffraction pattern which could be assigned to the planes of (211), (112) or (300) in the HAP.^{41,42} The crystalline results of XRD and SAED supported the FTIR result and confirmed the sample was composed of CNCs and HAP.

3.2.2. TEM images of the CNCs before and after HAP coating. To confirm the HAP is coated on the CNCs, we further adjusted the SBF pH to 8.5 without adding any CNCs substrates. However, no HAP precipitation was obtained in that condition, indicating the coating process was initiated by the negatively charged CNCs. The mechanism of the HAP deposition has been previously reported and involves HAP nucleation and subsequent growth.¹⁸ When the CNCs were added into the SBF, Ca^{2+} and PO_4^{3-} could bind to the surface of the CNCs by ionic bonds and hydrogen bonds, and this process formed a stable nucleus. After that, the HAP nucleus grew and aggregated as more and more Ca^{2+} and PO_4^{3-} attached to the CNCs and finally generated a HAP layer coated on the CNCs. This process is predisposed to occur under basic conditions. The images of the CNCs fibers before and after the HAP coating were measured by TEM and results are shown in Fig. 3. The dimensions of the original CNCs were determined to be 9.5 nm in width and 85.3 nm in length, which increased to 42.8 nm and 161.6 nm , respectively, after coating most of the CNCs with fine HAP particles. This result indicated the coated HAP layer have a size around $16.7\text{--}38.2\text{ nm}$ (calculated by the increase of CNCs' width and length).

The synthesis process was optimized by adjusting the pH of the SBF from 5.0 to 11.0. Thermogravimetric analysis (TGA) method was used to analyze the content of HAP in different



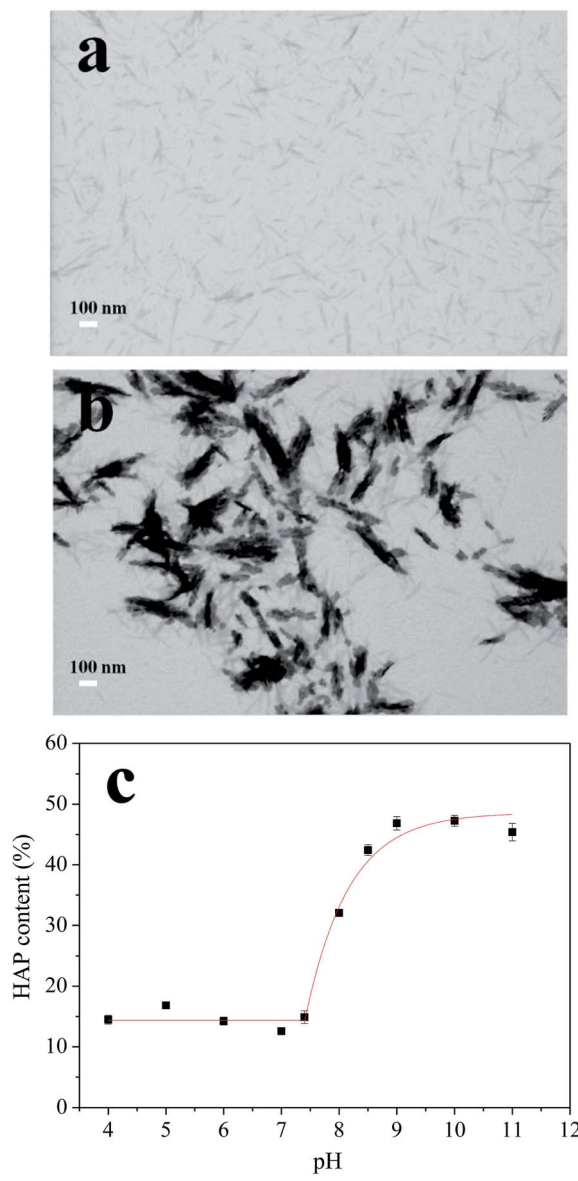


Fig. 3 Micrographs of CNCs before (a) and after (b) HAP coating, scale bars 100 nm. The relationship between the HAP content and the pH values (c).

nanocomposites and results were shown in Fig. 3c. When the pH value of the SBF increased from 4.0 to 7.4, the HAP contents in the nanocomposite were nearly constant (~15%). Further adjusting the SBF system to basic conditions, the HAP content increased exponentially by increasing pH values, with the highest content of ~47% obtained at pH 9.0. The deposition of HAP revealed a high degree of HAP supersaturation in the basic condition. Furthermore, when pH was increased beyond 9.0 to 10.0 and 11.0, HAP contents did not change that much which indicated that all of the Ca^{2+} and PO_4^{3-} ions have deposited on CNCs at pH 9.0. These results suggested that the significant HAP coating process started at pH 7.4 and saturated at pH 9.0. Therefore, pH values of 7.4 and 8.5, which were in this deposition pH range, were chosen for further experiments.

3.3. Freeze-casting of the nanocomposite for porous scaffolds

3.3.1. Morphology of freeze-casted foams by different freezing technologies. The porous structure of biomaterial plays an important role in tissue engineering in regards to promoting the desired cellular response and development of the new tissue.⁴³ Pore size, shape, and volume of biomaterials can affect strength and ability of the regenerated tissues, and their uptake of nutrients and release of waste products.⁴⁴ Freeze-casting has long been regarded as a versatile and promising technology to fabricate the porous scaffolds.⁴⁵ This process involves freezing of water into ice crystals followed by sublimation of ice under reduced pressure. The resulting product is a porous structure material in which the pores are the replica of the ice.⁴⁶ Thus, the freezing technology has a significant influence on the pore formation in the final foams.

In this study, nanocomposites prepared by HAP coating on CNCs were freeze-casted into porous foams by different freezing technologies including conventional freezing and directional freezing in liquid N_2 (DF). The morphology of the freeze-casted foams is shown in Fig. 4. The samples prepared by directional freeze-casting had oriented and continuous channels (Fig. 4a–c), compared to samples frozen by PL. During directional freezing, only the bottom of the vials was in contact with liquid N_2 , thus ice crystals were induced to grow vertically, parallel to the imposed temperature gradient.⁴⁷ Moreover, it can be found that the size of channels in the pure CNC scaffold was around 534.3 μm , while it decreased to 217.0 μm (at pH 7.4) and 227.1 μm (at pH 8.5) with the introduction of HAP. In addition, in the samples with only CNCs, pores in the channels were irregular and randomly distributed (Fig. 4a). With the addition of HAP, a more uniform and homogeneous pore structure was observed (Fig. 4b and c). This phenomenon was possibly caused by various adhesive forces (such as ionic bonds and hydrogen bonds) between CNCs and HAP that benefitted construction of a stable, uniform and oriented porous architecture.²² However, when the nanocomposites were solidified by plunging into the liquid N_2 , the ice crystals could nucleate at any places, and this resulted in a structure with randomly orientated pores (Fig. 4d–f).

3.3.2. Thermal properties. The thermal stability and degradation properties of the scaffolds were measured by TGA under a N_2 atmosphere and results are shown in Fig. 5. The curves of pure CNCs and low HAP content foams almost overlapped and included three weight loss regions. The first weight loss occurred near 105 °C, corresponding to the initial moisture loss. The main degradation appeared in the second region from 239 °C to 325 °C, with weight loss of 64.9% and 62.0% for pure CNCs foam and foam prepared at pH 7.4, respectively. This significant weight loss was primarily due to dehydration, depolymerization, and carbonization of the nanocomposites.⁴⁸ The last weight loss region between 325 °C and 450 °C is due to the further breakdown of the char into lower molecular weight gas compounds. Of significance, the thermal property of the nanocomposite was significantly improved with the incorporation of a large amount of HAP (sample prepared at pH 8.5). The main degradation temperature region increased to 269–381 °C



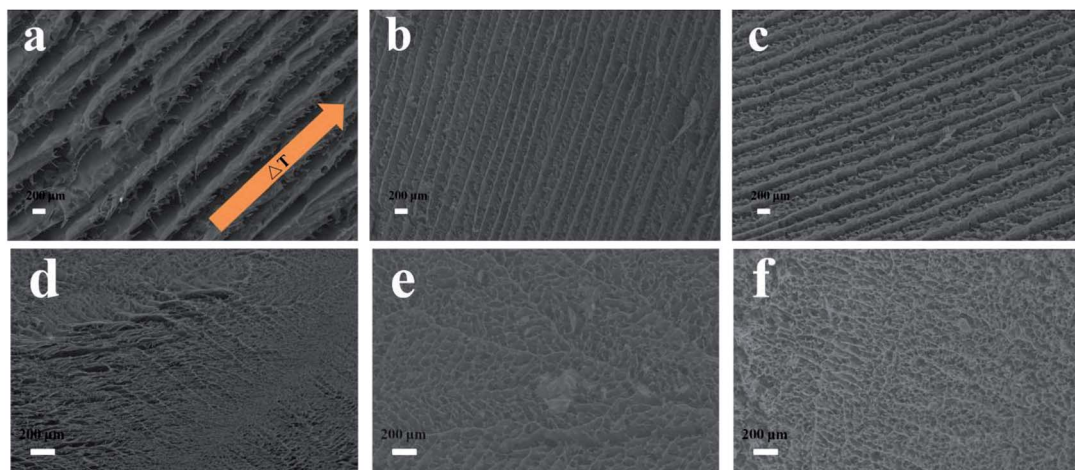


Fig. 4 SEM images of the longitudinal side views of the freeze-casted porous foams. (a–c) Directionally frozen; (d–f) frozen by plunging into liquid N₂. (a and d) Pure CNCs; (b and e) prepared at pH 7.4; (c and f) prepared at pH 8.5.

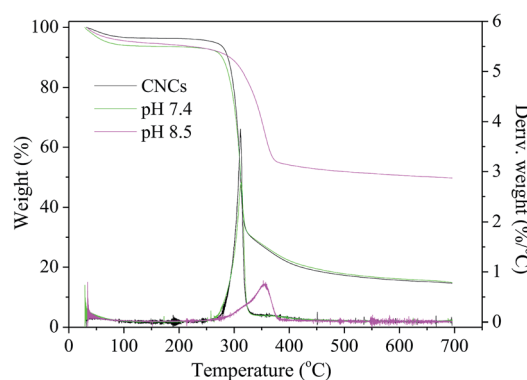


Fig. 5 TGA-DTG curves of the CNCs/HAP scaffolds.

with a weight loss of 54.9%. In addition, it can be seen from the DTG curve that the maximum weight loss decreased from 3.9% (at 311 °C) to 2.8% (at 313 °C) and 0.8% (at 357 °C) per °C, respectively, as the HAP content in the nanocomposites was increased, indicating the introduction of HAP is beneficial to the thermal property. This improvement was mainly attributed to the strong ionic and hydrogen bonds between CNCs and HAP and thus improved interfacial bonding in the mixture.²

3.3.3. Porosity, density and mechanical property of the scaffolds. The porosity of the samples was determined by the ethanol displacement method and the results are shown in Table 1. The results demonstrated that the porosity of the scaffold was significantly related to the solidifying methods employed (*i.e.* freezing temperature: FZ > DF > PL). The samples acquired by FZ freezing showed a higher porosity than freezing by PL and DF. For example, the porosity of pure CNCs foam increased from 83.2% (DF) to 85.6% (PL) and further reached 91.2% with solidification by FZ. The same findings have also been reported by other researchers, indicating the freezing methodology significantly affected pore formation and the FZ method at a higher freezing temperature contributed to pore formation and resulted in higher porosity.⁴⁹ In addition,

porosity tended to decrease with increasing HAP content in the nanocomposite. The porosity of the pure CNCs foam and the CNCs/HAP prepared at pH 7.4 were almost the same irrespective of the solidification method. However, porosity dropped by ~10% by further increasing the HAP content to 42.4% (prepared at pH 8.5). This could be attributed to the increasing HAP content which could have occupied more interstitial space in the frozen samples, resulting in lowered porosity after ice sublimation.⁹ High porosity structure is beneficial to tissue growth. Generally, scaffolds with a porosity higher than 70% could significantly facilitate nutrient delivery, cell adhesion, and attachment.²⁷ The porosity of the scaffolds from this study is consistent with other HAP contained material, showing high porosities that have met the requirement for bone substitute scaffold materials.

As to density, it also varied with the solidification method (Table 1). DF caused the highest density for all the three samples. For example, for CNCs/HAP nanocomposite prepared at pH 7.4, density increased from 46.1 to 48.1 and 50.8 mg cm⁻³ with freezing methods of FZ, PL, and DF, respectively. This finding is consistent with the porosity results, as increased porosity results in decreased density. According to Xu *et al.* (2014), density variation of samples was caused by the volume shrinkage during the freeze-casting process, which indicated that DF had a higher volume shrinkage compared to other two technologies, thus leading to a higher density. Concerning the effect of HAP content on density, a slight increase was observed with increasing HAP content which could be explained by the fact that the HAP particle is heavier than CNCs.⁴⁷

Scaffolds for bone substitution also need to be mechanically strong to provide enough structural stability to support cell adhesion. A recent study reported that a 20 times improvement in mechanical strength was obtained with the HAP coated CNCs, compared to the normal CNCs/HAP mixture.⁵⁰ The compression results of the synthesized CNCs/HAP scaffolds from different solidification technologies are summarized in Table 1. The mechanical properties of the scaffolds were also significantly related to the freezing technologies (freezing

Table 1 Porosity, density and compression results of the CNC/HAP scaffolds

Samples	Solidification technology	Porosity (%)	Density (mg cm ⁻³)	Stress at 50% strain (kPa)	Compressive modulus (kPa)
CNC	FZ (-20 °C)	91.2 ± 3.3	44.7 ± 1.2	17.7 ± 1.6	29.6 ± 1.2
	PL (-196 °C)	85.6 ± 2.1	46.4 ± 1.7	23.9 ± 1.1	42.4 ± 2.7
	DF (-196 °C)	83.2 ± 2.4	50.4 ± 0.9	37.0 ± 0.3	80.6 ± 1.4
CNC/HAP at pH 7.4	FZ (-20 °C)	90.3 ± 3.1	46.1 ± 2.5	13.6 ± 2.1	31.4 ± 1.2
	PL (-196 °C)	86.9 ± 1.3	48.1 ± 1.7	34.0 ± 1.0	72.8 ± 0.7
	DF (-196 °C)	82.6 ± 0.7	50.8 ± 2.2	43.1 ± 0.9	119.6 ± 2.7
CNC/HAP at pH 8.5	FZ (-20 °C)	81.0 ± 1.3	46.8 ± 1.6	23.2 ± 1.4	54.1 ± 1.4
	PL (-196 °C)	75.4 ± 2.2	47.4 ± 1.0	35.4 ± 1.2	100.8 ± 3.6
	DF (-196 °C)	73.6 ± 0.3	51.1 ± 1.8	61.7 ± 1.1	227.6 ± 2.7
CNC/HAP mixture	FZ (-20 °C)	83.2 ± 0.5	45.9 ± 0.9	20.6 ± 1.2	35.7 ± 2.6
	PL (-196 °C)	80.3 ± 1.5	48.2 ± 0.3	27.4 ± 0.7	56.9 ± 0.8
	DF (-196 °C)	78.2 ± 0.7	50.3 ± 1.2	40.3 ± 1.7	92.5 ± 2.3

temperature: FZ > DF > PL), including compressive stress and modulus, as shown in Table 1. For example, the compressive stress of CNCs scaffold increased from 17.7 kPa to 23.9 kPa and 37.0 kPa when FZ was replaced by PL and DF. The same trend can also be found in compressive modulus which increased from 29.6 kPa to 42.4 kPa and 80.6 kPa with different freezing methods. It could be concluded that both the freezing rate and freezing direction affected the mechanical properties. As observed, the fast freezing rate improved the mechanical properties (comparing PL to FZ). Moreover, the best compression results were obtained for the DF samples indicating that the oriented structure of the scaffold with strong anisotropy also improved the mechanical properties. During DF, ice crystals were induced to grow vertically, along with the direction of the imposed thermal gradient. The process formed an oriented and continuous channel architecture parallel to the direction of the compression force, this may explain why the DF resulted in the better mechanical performance than FZ and PL. On the other hand, with the coating of HAP, both the compression stress and compression modulus significantly increased, with highest values of 61.7 kPa and 227.6 kPa obtained with the nanocomposite at pH 8.5. This enhancement could be attributed to the improvement of stress transfer from the CNCs matrix to the strong HAP particles, and thus resulting in the superior mechanical properties. We further investigated the mechanical properties of the plain CNCs/HAP mixture (with the HAP content same to the CNCs/HAP composite prepared at pH 8.5), as shown in Table 1. The highest mechanical stress and modulus of the CNCs/HAP mixture were 40.3 kPa and 92.5 kPa (prepared by DF), respectively, which were much lower than our CNCs/HAP composite due to the heterogeneous dispersion of HAP in CNCs. This result indicated that our *in situ* HAP coated CNCs composite was superior to the CNCs/HAP mixture. Although the mechanical strength is still lower than a actual bone, we've made an improvement compared to some other bone scaffolds, including 8.13 times compression modulus of a chitosan-hyaluronic acid scaffold,⁵¹ 6 times of a 3-D HAP composite,⁵² and 2.0 times of a hydrogel/fiber scaffold,⁵³ which holds the potential for application as a substitute to the traditional metallic materials and bioceramics due to its biocompatibility. Further bone cell

related research will be done in the following experiment to test its perspective as a real bone substitute.

4. Conclusion

Our study illustrated a facile approach to fabricate a biomimetic scaffold by cultivating the CNCs in a simulated body fluid. With such methodology, the HAP was successfully coated on the CNCs, having a HAP layer size around 16.7–38.2 nm. The HAP content of the nanocomposite could be controlled by adjusting the pH of the solution, with the highest value of ~47% prepared at pH 9.0. Moreover, after being freeze-casted into solid scaffolds, these nanocomposites all showed high porosity (higher than 70%) and light weight (density around 50 mg cm⁻³). Compression results indicated the introduction of HAP increased the compressive properties of the freeze-casted scaffolds. On the other hand, the scaffolds solidified by DF have a better mechanical property than by PL and FZ due to the orientated structure it caused. These results demonstrated that the use of such *in situ* HAP coating followed by the directional freezing technology generated a biomimetic scaffold with high porosity and improved mechanical strength that can serve as a potential substitute for native bone.

Conflicts of interest

The authors declare no competing financial interests.

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