A novel antibacterial cellulose based biomaterial for hernia mesh applications†

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A novel bacterial cellulose/collagen–hydroxypropyltrimethyl ammonium chloride chitosan composite (BCC–H), as an ideal artificial hernia mesh, was synthesized by combining solution impregnation with the EDC/NHS chemical crosslinking method. Scanning electron microscopy was used to examine the morphology of the bacterial cellulose/collagen (BCC) and BCC–H composites. Infrared spectroscopy demonstrated that collagen and hydroxypropyltrimethyl ammonium chloride chitosan (HACC) were incorporated into the bacterial cellulose network successfully. Mechanical testing of the materials showed that the addition of HACC made the tensile strength decrease while the elongation at break was increased. Antibacterial assessment of the BCC–H composites revealed distinct growth inhibition against Escherichia coli and Staphylococcus aureus. In vitro cell experiments confirmed that the BCC–H composites had low cytotoxicity and good biocompatibility.

Introduction

A hernia, especially an external abdominal hernia, as a common disease of general surgery, causes damage to the digestive and reproductive system of patients and even threatens their life if severe. Nowadays, operation has been proven to be the only hernia treatment, and includes traditional inguinal herniorrhaphy, open tension-free herniorrhaphy and laparoscopic hernia treatment, and includes traditional inguinal herniorrhaphy. In contrast with the others, laparoscopic herniorrhaphy is a clinical operation since it is considered to be a minor surgery, involves a rapid recovery, presents less postoperative complications and has low recurrence. However, the artificial hernia mesh used is another determinant of the success of hernia surgery. Currently, artificial hernia meshes are based on polypropylene, presenting postoperative complications and has low recurrence. Therefore, one of the main objectives in the current research on hernias is to develop a light, anti-infective, anti-adhesive, biocompatible and bioactive mesh.

Bacterial cellulose (BC) is a polysaccharide composed of glucose molecules as common substrates linked by β-(1-4)-glycosidic bonds. BC has drawn lots of attention in the biomedical field due to its good mechanical properties, good stability, favorable biocompatibility and low toxicity. These types of materials have already been used for several applications, such as wound dressing, artificial skin for severe burns, artificial blood vessels and tissue engineering scaffolds. However, the non-antibacterial ability of BC confines its further application as an implantable biomaterial.

Hydroxypropyltrimethyl ammonium chloride chitosan (HACC) is a derivative of chitosan, which is known to be a natural antibacterial agent. Furthermore, HACC has better antibacterial properties than chitosan due to its positive charge. Meanwhile, the introduction of a quaternary ammonium group into chitosan can enhance the hydratability, resulting in better moisture absorption and moisture retention. However, the positive charge on the surface of HACC has an intense electrostatic interaction with negatively charged cells, causing cytotoxicity to a certain degree. Therefore, to address this issue, collagen, as the main protein of animals and which is widely used in the biomedical field, is introduced to improve the biocompatibility and bioactivity of the composites.

In this study, we tried to fabricate a novel antibacterial cellulose based biomaterial for hernia mesh applications. An EDC/NHS chemical crosslinking method is used to prepare the BCC composite. A solution impregnation method is used to incorporate HACC into BCC to enhance its antibacterial properties. Scanning electron microscopy and infrared spectroscopy were used to examine the morphology of the BCC composites and to ensure the successful incorporation of collagen and HACC into the bacterial cellulose network. Mechanical testing of the materials was also performed to ensure that the mechanical properties of the hernia mesh are suitable. Furthermore, antibacterial assessment and an in vitro cell
The chemical structures of the BCC-H composites were characterized using Fourier transform-infrared spectroscopy (FTIR, Nicolet, USA) in the range between 3800 and 800 cm\(^{-1}\) at room temperature. In order to do so, the materials were lyophilized and ground into powder and then combined with KBr to prepare a disk for observation.

**Experimental section**

**Materials**

Bacterial peptone, bacto-agar and yeast extract were purchased from Guangdong Huankai Microbial Technology Limited Company (Guangzhou, China). MgSO\(_4\) and CaCl\(_2\) with purities higher than 99.9% were purchased from Tianjin Chemical Factory (Tianjin, China). 1-(3-Dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (EDC-HCl) and N-hydroxysuccinimide (NHS) with purities higher than 99.9% were provided by GL Biochem Ltd. (Shanghai, China). 2-Hydroxypropyltrimethyl ammonium chloride chitosan (HACC, Medical Grade) was provided by Zhejiang Aoxing-biology Ltd. (Taizhou, China). Acetic acid, ethyl alcohol (absolute), sodium hydroxide, sodium dodecyl sulfate and glucose (AR) were provided by Sinopharm Chemical Reagent Co., Ltd. (Shanghai, China). A dialysis bag (MWCO8000-1.44w) was provided by Guangzhou Qiyun biotechnology Ltd. (Guangzhou, China). Collagen (beef tendon, Type 1) was provided by Guangzhou Huameikanglian Biotechnology Ltd. (Guangzhou, China). *Acetobacter xylinus* was purchased from ATCC (USA).

**Preparation and purification of BC**

*Acetobacter xylinus* was inoculated on sterilized medium, which consisted of fermented coconut juice, 1% yeast extract powder, 0.6% bacto-peptone, 0.02% MgSO\(_4\), 0.01% CaCl\(_2\) and 2% glucose. Subsequently, the bacterium was cultured statically to get a bacterial cellulose membrane. Thereafter, the BC membrane was purified by sequential washing steps of tap water, distilled water, 0.1 M NaOH, 2% sodium dodecyl sulfate and distilled water, in that order. The purified BC was cut into 3 cm \(\times\) 4 cm slices and stored in deionized water at a low temperature to prevent drying.

**Preparation of BCC and BCC-H composites**

The prepared BC pieces were immersed in a 0.25% (w/w) collagen solution and gently rocked for 24 h in a shaking table. Then, the BC pieces were crosslinked in a 0.1% EDC/NHS solution (EDC : NHS = 2 : 1) solution for 24 h at 4°C. The pieces were afterwards washed with distilled water, and a 24 hour dialysis followed. The water was changed every 12 h. The prepared BCC composites were dried at 37°C for 24 h. Subsequently, 0.5%, 0.25% and 1.0% (w/v) HACC solutions were prepared. The prepared BCC composites were immersed in the 0.5%, 0.25% and 1.0% HACC solutions for 12 h, respectively. Finally, the prepared BCC-H materials were dried at 37°C and named as BCC-0.25H, BCC-0.5H and BCC-1.0H, respectively.

**Infrared analysis of the BCC-H composites**

The chemical structures of the BCC-H composites were characterized using Fourier transform-infrared spectroscopy (FTIR, RSC Advances, 2017, 7, 11601–11607)

**Preparation of BCC-H composites**

The prepared BCC-H materials were characterized using scanning electron microscopy (SEM, EVO18, Germany). Prior to the SEM observation, the materials were lyophilized and coated with an ultrathin layer of gold by an ion sputter.

**Water absorptivity characterization**

The equilibrated water absorptivity of the BCC-H materials was measured. First, the weight \((M_0)\) of the BCC and BCC-H materials treated under different conditions was measured. The specimens were then immersed in a phosphate-buffered saline solution (PBS, pH = 7.4) at 37°C. After determined time points, the specimens were removed and their weight \((M_t)\) was measured. The water content ratio (W%) was calculated according to the following equation:

\[ W\% = \frac{(M_0 - M_t)}{M_0} \times 100\% \]

**Morphology of the BCC-H composites**

The morphologies of the prepared BCC-H materials were characterized using scanning electron microscopy (SEM, EVO18, Germany). Prior to the SEM observation, the materials were lyophilized and coated with an ultrathin layer of gold by an ion sputter.

**Mechanical properties**

The mechanical properties were determined using a dynamic biomechanical testing machine (Bose ELF3200, USA). In brief, the specimens were cut into dumbbell-shaped sticks with sizes of 35 mm \(\times\) 5 mm. The sticks were then stretched at a cross-head speed of 100 mm min\(^{-1}\) to reach a constant strain rate. Five parallel samples were used for each group. The tensile strength \((\delta)\) and elongation at break were calculated according to the following equations:

\[ \text{Tensile strength} \quad \delta = \frac{F}{(a \times b)} \]

\[ \text{Elongation at break} = \frac{\Delta L}{L} \times 100\% \]

where \(F\) represents the breaking load, \(a\) is the width of the stick, \(b\) is the thickness of the stick, \(\Delta L\) is the elongation of the stick when it fractures, and \(L\) is the original length of the stick.

**Antibacterial assessment**

The antibacterial properties of BC, BCC and BCC-H were analyzed against *Escherichia coli* as a Gram-negative bacterium and *Staphylococcus aureus* as a Gram-positive bacterium, using the colony forming count method.\(^{29-31}\) *E. coli* and *S. aureus* were provided by the Guangzhou industrial microbiology testing center (Guangzhou, China). Both types of bacteria were cultivated in a nutrient broth. The composites were placed in 24-well culture plates and inoculated with 750 \(\mu\)L of bacterial suspension \((1.0 \times 10^8\) cfu mL\(^{-1}\)) at 37°C. After being cultured for 3 h, the bacteria were washed and the solutions were diluted to 50 \(\mu\)L. Then, 10 \(\mu\)L of the diluted solution was taken and placed on agar plates. Colonies were counted visually after incubating for 24 h at 37°C. The antibacterial rate was calculated using the following equation:

\[ A = \frac{(C_0 - C_t)}{C_0} \times 100\% \]

where \(C_t\)
represents the bacteria number of the experiment group and \( C_0 \) represents the bacteria number of the control group.

Cell proliferation and morphology

The cell proliferation was assessed using the widely used Cell Counting Kit CCK-8 (CCK-8, Dojindo Laboratories, Japan). In brief, mouse-derived fibroblast cells (L929, Cell Bank of the Chinese Academy of Sciences, Shanghai, China) were propagated in 1640 Medium (PAA, Germany) supplemented with 10% fetal bovine serum (FBS, Life Technologies, Gibco, USA). The BCC composites were placed in 48-well plates and sterilized using Co\(_{60}\)-gamma radiation. After pretreatment with the culture medium for 12 h, the cells were seeded at a density of \( 2.0\times10^4 \) cells per well and cultured for 1 d, 3 d and 5 d. At the determined time points, the CCK-8 working solution \((1640 : \text{CCK-8} = 10 : 1)\) was added to each sample, which was then incubated at 37 °C for 1 h. Subsequently, the supernatant medium was extracted and the absorbance of 450 nm light was measured with a microplate reader (Thermo 3001, USA) \((n = 5)\).

A laser scanning confocal microscope (CLSM, Leica SP8, Germany) was used to observe the cell viability and morphology of the L929 cells on the composites. Specifically, the obtained BCC and BCC–H membranes were placed in 48-well plates after having been sterilized using Co\(_{60}\)-gamma radiation. L929 cells were seeded on the membranes at a density of \( 2.0\times10^4 \) cells per well and inoculated at 37 °C in an incubator with 5% CO\(_2\). After being cultured for 1 d, the samples were washed in PBS and then a living/dead staining solution was added for 20 min at room temperature. Thereafter, the cells were washed again with PBS. Finally, the cell viability and morphology of the L929 cells on the surface of the materials was observed using the laser scanning confocal microscope.

Statistical analysis

A one-way analysis of variance (ANOVA), followed by Tukey’s test for comparison of means, was used to assess the level of significance, by employing the SPSS 19.0 statistics software. The results were expressed as the mean ± standard error and *\( p < 0.05 \) or **\( p < 0.01 \) was designated as statistically significant.

Results and discussion

Infrared analysis of the BCC–H composites

In this study, the bacterial cellulose/collagen–hydroxypropyltrimethyl ammonium chloride chitosan composite (BCC–H) was synthesized by combining solution impregnation with the EDC/NHS chemical crosslinking method. In order to identify if HACC was introduced into the BCC material successfully, FTIR analysis was conducted to characterize the functional group changes of the BCC composites before and after recombination with HACC. The results (Fig. 1) showed that a new absorption peak was present for the BCC–H composites. Besides, the intensity of the absorption peak was enhanced when the HACC content increased, which demonstrated that HACC was incorporated into the BCC material successfully.

Surface morphology of the BCC–H composites

The surface morphologies and structures of the BCC and BCC–H composites were also investigated using SEM to further validate the successful incorporation of HACC. As shown in Fig. 2, the BCC composites clearly showed a three-dimensional fibrous network structure. Compared to the BCC composites, the surface and section morphology of the BCC–H composites was different due to the immersion in the HACC solution. As can be seen in Fig. 2a–d, there was no difference between the surface morphologies of BCC and BCC–0.25H. However, when the HACC content increased, the surface morphology of the BCC–H composites showed a significant change and the BCC–1.0H composite was covered with a layer of HACC particles, which were distributed uniformly on the surface of the bacterial cellulose (Fig. 2d). Furthermore, the cross-section morphology showed that the interlayer distance also changed when the amount of HACC that penetrated into BCC increased (Fig. 2e–g). It was noted that the BCC–H composites showed a more fluffy layered structure compared to BCC when the HACC content increased, which may be facilitated by adhesion and grown into the materials. Therefore, from the SEM results, we can conclude that HACC was incorporated into the BCC composites successfully and this resulted in the changes in the structure of the BCC composites.

Water absorptivity

It has been the consensus that a good implanted biomaterial should not expand in volume excessively after absorbing water, so that it possesses stable physical and chemical properties. Besides, the shrinking of materials is an important factor resulting in the recurrence of hernias. In this study, the water absorptivity of the BCC–H composites was determined in PBS at 37 °C and the water absorption curves of the BCC–H composites are shown in Fig. 3. The results showed that a balanced water absorption was reached quickly after immersing BCC–H in PBS.
for 30 min, which demonstrated that the BCC–H composites that we fabricated possessed a good water absorptivity ability. Furthermore, compared with the BCC composites, the equilibrium moisture content of the BCC–H composites increased when the HACC content increased (Fig. 3b). For instance, the equilibrium moisture content of BCC–1.0H improved by almost 50% compared with BCC. The possible reasons may be that the hydration ability of the molecules was strengthened because of the introduction of quaternary ammonium salt groups into chitosan. In addition, we also see that the volumes of BCC–0.25H and BCC–0.5H in the water equilibrium state presented no significant difference, which contributes to their use as implantable materials. Nevertheless, for BCC–1.0H, the volume expansion was obvious, which means that it may not be suitable as a good implantable biomaterial.

Mechanical test

Elastic deformation is one of the major parameters of artificial hernia meshes. It is reported that the natural abdominal wall possesses good elasticity and about 38% elastic deformation under 32 N cm⁻¹ tension. Therefore, similar elastic deformability should be required of an artificial mesh for hernia repair. Here, we studied the mechanical properties using a dynamic biomechanical testing machine and the results are shown in Fig. 4. Although when compared with BC, the elongation at break of BCC was significantly increased, it still needed to be further enhanced for use as an artificial mesh. As is shown in Fig. 4a, the elongation at break of the BCC composites greatly increased after immersing them in HACC solution. With increasing the content of HACC solution, the elongation at break of the BCC–H composites was also enhanced. However, contrary to the elongation at break, the corresponding tensile strength of the BCC–H composites showed the opposite change (Fig. 4b). The results showed that the tensile strength of the BCC–H composites decreased as the HACC content increased. As shown in Fig. 4, the elongation at break of BCC–0.5H (close to 38%) and tensile strength (25 MPa) can match the clinical requirements.

Antibacterial studies

At present, artificial hernia meshes in clinical applications are always associated with infections. In this study, we try to develop a novel antibacterial hernia mesh by the addition of
HACC into BCC composites. In order to verify its ability, the antibacterial efficacy of BC, BCC and BCC–H against *E. coli* and *S. aureus* was measured. As is shown in Fig. 5, the BCC–H composites showed a better antibacterial effect than the BC and BCC composites, while the BC and BCC composites have no antibacterial ability (Fig. 5a–c). Moreover, with increasing HACC content, the BCC–H composites have increased antibacterial efficacy against *E. coli* and *S. aureus* (Fig. 5d). Contrary to the BCC–H composites, the increasing number of bacterial colonies for the BCC composites may result from the collagen on the surface of the material, which helped the bacteria to adhere to the material’s surface. To characterize this observation in a quantitative manner, the number of bacterial colonies was calculated for the BC, BCC and BCC–H composites. The results (Fig. 5b and c) showed that the bacteriostatic rates of BCC–1.0H and BCC–0.5H reached up to 99% and 88%, respectively. Besides, we also compared the antibacterial efficacy against *E. coli* and *S. aureus* among the BCC–H composites (Fig. 5d). It was obvious that the antibacterial efficacy of the BCC–H composites was enhanced when the content of HACC
increased. In addition, we can also see that the antibacterial effect of BCC–0.25H against *E. coli* and *S. aureus* was different. Previous works showed that the antibacterial effect of BCC–H composites against *E. coli* is better than that against *S. aureus*, which may be related to the different cell wall structures of Gram-negative bacteria and Gram-positive bacteria. The cell wall of Gram positive bacteria possesses a thicker peptidoglycan layer and can prevent more antibacterial agent from entering the cytoplasm to destroy the cellular structure.\(^{35}\)

**Cytotoxicity and biocompatibility**

Cytotoxicity and biocompatibility are the important parameters for artificial hernia meshes to be applied in clinic.\(^{44}\) To characterize the cytotoxicity and biocompatibility, L929 cell proliferation on BCC and BCC–H composites was assessed via CCK-8 assay after 1, 3, and 5 days of culturing. As shown in Fig. 6a, the L929 cells show normal cell proliferation on all samples. On the 1st day, there was no difference between the cell densities on the BCC and BCC–H composites. With prolonged culture time (the 3rd day), although the cell density on the BCC–H composites decreased gradually with increased content of HACC, there was no difference among them. On the 5th day, BCC–1.0H still presented the lowest cell proliferation rate. Although the rate of cell proliferation of BCC–0.5H was still less than that of BCC, there were no significant differences among BCC–0.5H and BCC. Besides, after culturing the L929 cells on BCC and BCC–H for 1 d, live/dead staining was also carried out to ensure cell viability and adhesion. As is shown in Fig. 6b, the L929 cells possessed good cell adhesion and growth on the BCC and BCC–H composites. In addition, there were few dead cells after culturing for 1 d, which demonstrated that the BCC and BCC–H composites showed no obvious cytotoxicity. With the culture time extended (the 5th day), BCC–1.0H and BCC–0.5H showed low cytotoxicity, which may result from the slow releasing of HACC from the BCC–H composites. These data indicate that, with the increase of HACC content in the BCC–H composites, the proliferation and cell viability of the L929 cells were not affected greatly, which demonstrated that the BCC–H composites possess good cell proliferation and biocompatibility.

**Conclusions**

In summary, we successfully fabricated novel BCC–H composites with differing HACC content by incorporating collagen and quaternary ammonium chitosan with BC, and explored their potential for hernia mesh application. A series of property characterizations were performed in detail. Our results showed that the BCC–H composites, especially BCC–0.5H, possess appropriate mechanical properties, and good antibacterial efficacy and biocompatibility, which prove them to be excellent candidates for artificial meshes in hernia surgery.

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