Surface-Modified Bacterial Cellulose with Mercaptosilane as a Multifunctional Platform

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Abstract: Cellulose synthesized by bacteria has unique properties such as high water retention capacity, biocompatibility, biodegradability and flexibility. Nevertheless, modification of this biomaterial is required in order to obtain multifunctional materials, which may be applied in several high-value added products, as catalytic and cell culture platforms. The surface of bacterial cellulose (BC) can be modified by several approaches, namely: (i) physical treatment by plasma, (ii) adsorption of molecules onto BC surface, and (iii) chemical modification. In this sense, the aim of this study was to modify the BC surface by silanization reaction at room temperature using a mixture of ethanol and water, using two different protocols. Thus, BC membranes synthesized by Komagataeibacter xylinus were modified by adding the thiol (SH) functional group with (3-mercaptopropyl) trimethoxysilane under mild conditions. The produced materials were analyzed by elemental analysis, ATR–FTIR, TGA and SEM, and the successful modification was proven by elemental analysis and SEM.

Keywords: Bacterial cellulose; Surface Modification; Silanization; Mercaptosilane.

Introduction

Cellulose is the most abundant biopolymer on planet and it can be produced by green plants, fungi and bacteria. Cellulose consists of β-D-glycopyranose units linearly arranged by β-(1→4) glycosidic bonds. Plant-derived cellulose is usually associated with several components as hemicellulose, lignin and pectin, while bacterial cellulose (BC) is obtained chemically pure. Besides, BC presents nanometer-sized fibers once it is excreted by a microorganism, therefore BC is also known as bacterial nanocellulose

The strain used for the BC production was Komagataeibacter xylinus. The bacteria were cultured under static conditions in Hestrin–Schramm (HS) media, composed of D-glucose, yeast extract, peptone, disodium hydrogen phosphate, citric acid, and distilled water. Dried BC membranes of surface area and thickness of 1.5 cm² and 0.05 mm, respectively, were treated by silanization reaction with MPTMS by two different methods. A mixture of ethanol and water 10:1 (v/v) was used as solvent.

For Method A (sample BC–SH–A), it was used 0.538 mmolL⁻¹ of MPTMS. The system was kept at stirring for four hours at room temperature. Then the excess of solution was discarded and the membranes were washed three times with acetone and dried at room temperature for three days.

For Method B (sample BC–SH–B), the membranes were put in contact with the solvent for 5 minutes. Then NH₄OH was added to medium followed by the addition of siloxane — the final medium concentration was 0.163 mmolL⁻¹ of MPTMS, three times lower than that used in the first method. The system was kept at stirring for 12 hours at room temperature. Then, as the process done for the previous method, the excess of solution was discarded and the membranes were washed three with acetone and dried at room temperature for three days.

Experimental Procedures

BC Production and Surface Modification

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Characterization

The membranes were characterized by elemental analysis, Attenuated Total Reflectance Fourier Transform Infrared (ATR–FTIR), Scanning Electron Microscopy (SEM) and Thermogravimetric Analysis (TGA).

Elemental Analysis

Hydrogen, sulfur, nitrogen and carbon contents were determined by dry combustion in a Perkin Elmer elemental analyzer, model 2400 series II.

ATR–FTIR

The transmittance spectra were obtained in the infrared region by attenuated total reflectance (ATR–IR) in a Cary 630 Agilent spectrometer in the range of 4,000–650 cm⁻¹.

SEM

The morphological characterization was performed using a Joel JSM 7500F Field Emission Scanning Electron Microscope. The samples were covered with a thin layer of conductive carbon. The images were taken at 10,000 times magnification.

TGA

Thermogravimetric analyses were performed on a TA Instruments SDT Q600 thermal analyzer under the following conditions: heating rate of 10 °C/min with a synthetic air flow of 100 mL/min, from 30 °C to 600 °C. Alumina pan was used as reference.

Results and Discussion

In terms of chemical reaction of silanization, water has the function of induce siloxane hydrolysis to silanol (Figure 1, Step 1), while ethanol favors the materials drying process. The interaction between hydroxyls favors the condensation reaction (Figure 1, Step 2). Thus, in this type of modification the condensation reaction occurs between silanol groups (Si–OH) as well as between the silanol and the BC hydroxyls.(10). The condensation step can also be induced by employing strong or weak bases such as NaOH(7) and NH₄OH(9).

The efficiency of BC surface modification by the applied methods was verified by elemental analysis of pristine BC, used as reference, compared to the treated samples (Table 1).

Modifying the material with MPTMS necessarily implies increasing sulfur (S) content once this siloxane carries the thiol functional group. BC–SH–A has a sulfur percentage very close to the native (reference) while BC–SH–B has a sulfur percentage almost 4 times higher compared to the native. This indicates that only method B was indeed efficient in modifying the BC surface through the covalent Si–O–Si bond formed between the silane moiety and the polymeric matrix.

FTIR–ATR spectrum (Figure 2) of native BC and BC treated by Methods A and B have the same profile. In all of them, there is a weak band at 900 cm⁻¹, attributed to vibrational modes of the C–O–Cβ (1→4) glycosidic bonds between glucose units, a band at 1,054 cm⁻¹ attributed to the C–OH stretch, and a band at 1,163 cm⁻¹, attributed to asymmetric deformation of the glucose pyranose ring C–O–C. The 3,340 cm⁻¹ band is attributed to the O–H stretch of water adsorbed to the cellulose surface(11). This analysis was not sensitive attest the surface functionalization even for the BC treated by Method B, once it was not possible to observe the band corresponding to the deformation of the Si–O–Cellulose group in the region between 1,100–1200 cm⁻¹(8).

In the SEM images obtained from the sample surfaces (Figure 3), it is observed that the native BC has a three–dimensional nanometric structure with randomly arranged nanofibers. It can be seen that the surface of the sample submitted to Method A did not change (there was no modification), unlike the surface of the sample submitted to Method B, where a distinct structure material deposited on the nanofibers is observed. These results corroborate with the data obtained by elemental analysis.

According to the thermogravimetric curves (Figure 4), it can be seen

Figure 1 – Silanization reaction with MPTMS.
Table 1 – Elemental composition of samples.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Hydrogen (%)</th>
<th>Nitrogen (%)</th>
<th>Sulfur (%)</th>
<th>Carbon (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Native BC</td>
<td>5.75</td>
<td>0.87</td>
<td>1.04</td>
<td>42.45</td>
</tr>
<tr>
<td>BC-SH-A</td>
<td>5.95</td>
<td>0.94</td>
<td>1.37</td>
<td>42.23</td>
</tr>
<tr>
<td>BC-SH-B</td>
<td>6.26</td>
<td>0.91</td>
<td>3.3</td>
<td>40.16</td>
</tr>
</tbody>
</table>

Figure 2 – ATR–FTIR spectrum of Native BC, BC–SH–A, and BC–SH–B.
Figure 3 – Surface image by SEM of (A) Native BC (B) BC–SH–A and (C) BC–SH–B.
that the membranes treated by both Methods show mass losses with more pronounced inflection points in relation to native BC. In addition, the percentage of surface water loss is lower for treated membranes: about 3% for treated samples and 5% for BC. As well as the FTIR analysis, the TGA was not sensitive to verify the BC functionalization once it was not observed mass increase by 600°C due to silicon oxide formation.

Figure 4 – TGA and DTG curves of (A) Native BC (B) BC–SH–A and (C) BC–SH–B.
Conclusions
Among the silanization methods employed in this study for BC surface modification, only the one whose condensation step was induced by employing NH$_4$OH was indeed effective. Elemental analysis and SEM analysis confirmed the efficiency of this particular surface functionalization using MPTMS.

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