Effect of water and alkali on purification bacterial cellulose membrane from Kombucha

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Abstract
Bacterial cellulose membrane (BCM) is a biomaterial synthesized by bacteria of the genus *Gluconocetobacter hansenii* with a higher degree of purity than plant cellulose. The commonly used raw material for manipulating bacterial cellulose is kombucha, a beverage consumed by a vast population around the world that promises health benefits. The beverage is composed of tea species *Camellia sinenses* and a carbon source, refined sucrose, and a starter culture of bacteria and yeast with 10% fermented tea (starter tea) to activate the fermentative process. The Kombucha’s bacterial cellulose membranes (KBCM) are formed over 7 to 10 days on the surface of the fermented product and have the appearance of a gelatinous membrane, this being the by-product of interest. In this work, the objective was to obtain a membrane composed of cellulose via Kombucha and purify it to obtain crystalline cellulose. The purification was performed with distilled water and 0.5M NaOH sodium hydroxide solution to remove residues from the fermentation, successfully removing sugars and bacteria. At the end of the experiments, a lighter film was obtained with coloration close to white, and comparative analyses were performed to verify the structural chemical composition, crystallinity, and morphology of the samples by techniques FTIR, DRX, and SEM, respectively. Then, once the biomaterial was purified, the range of applications expanded to several products to meet the biomedical area, sustainable packaging, and even the fashion industry.

Keywords: Bacterial cellulose; Kombucha; Purification; Crystallinity; Water; Alkali.
1. Introduction

The interest in developing new biomaterials has expanded over time with analyses and studies in various industrial sectors because they have a shallow environmental impact. An example of a biopolymer to be used as a biomaterial is bacterial cellulose, whose format is purer than the cellulose provided by plants (Costa & Biz, 2017).

Cellulose is a polysaccharide extracted from plant cells of plants such as eucalyptus, cotton, and trees (Wang et al., 2019). Besides non-woody plants such as hemp, jute, and sisal leaves (Alila et al., 2013). The problem for implementing the cellulose as a biotechnological material is the submission to drastic chemical treatments to remove lignin and hemicellulose, substances that are not of industrial interest. Moreover, a promising biotechnological material is a bacterial cellulose, which is in high demand for its unique properties, such as good crystallinity, high tensile strength, moldability, high degree of polymerization, and being a hundred times thinner than the cellulose fibers obtained from plants (Vandamme et al., 1998).

To be synthetically produced, it requires raw materials of high production cost (Vazquez et al., 2013). However, it was observed that BCM could be synthesized by good bacteria and come from some industrial residues such as sugarcane bagasse, soy molasses, coconut cream, fruits, and others (Dima et al., 2017).

There is a work that reports the effectiveness of obtaining bacterial cellulose by the bacterium *Gluconacetobacter xylinus* through rotten fruit; Jozala et al., (2015) reported in their study that it is essential to reduce waste and deploy these by-products to extract another bioproduct of interest in this case BC for different applications. Lin et al., 2014 demonstrated in their work that beer yeast residuals are a good nutrient source for *Gluconacetobacter Hansenii* for BC production, and the final properties and microstructures were as good as the conventional chemical method.

It is possible to obtain this biopolymer cheaply and easily through the ancient drink kombucha, consumed by many people worldwide (Aditiawati et al., 2021). This beverage consists of fermenting the tea of the *Camellia Sinenses* species along with an initial inoculum of bacteria, with the main bacteria being acetic: *Acetobacter xylinum*, *A. xylinoides*, *A. aceti*, *A. pausterianus* and *Bacterium gluconicum and yeasts, such as Candida sp.*, *Kloeckera sp.*, *Schizosaccharomyces pombe*, *S. ludwigii*, *S. cerevisiae*, *Torulaspora sp.*, *Zygosaccharomyces bailii*, and *Pichia* species along with refined sucrose used as a carbon source (Ugale, 2021). During fermentation of the liquid, a thick film forms on the surface of the beverage. This beverage, besides containing bacteria, yeast, and sucrose, contains cellulose (Goh, 2012). Therefore, it is of great interest to separate this material for study in biomedical applications, tissue engineering, orthopedics for bone regeneration, dental implants, vascular...
grafts, and adhesives to heal wounds, among others being studied for applications (Amarasekara et al., 2020). In addition to the biomedical area, the Kombucha’s bacterial cellulose membranes (KBCM) can be applied in the textile industry, for the production of new sustainable leather-like fabrics (Domskiene et al., 2019) and in the biodegradable packaging industry (Tapias et al., 1947). However, purification of this kombucha membrane is required before any application to improve the biomaterial of interest.

Thus, in this work, the main goal was to submit the Kombucha’s bacterial cellulose membranes (KBCM) to purification with an alkaline solution of 0.5M sodium hydroxide to remove the fermentative residues that are nucleic acids, sugars, and bacteria, leaving only the cellulose fibers in evidence, and increase the material crystallinity to extend the range of applications.

2. Methodology

2.1 Obtaining Kombucha’s bacterial cellulose membranes (KBCM)

The methodology was based on the work of Jayabalan et al., (2014). For obtaining the bacterial cellulose membrane, it is necessary to prepare a sweet tea infusion with 1 liter of water together with 100g of white sugar and 7 grams of black tea. The tea was left to infuse for 15 minutes. Then, the tea was cooled to room temperature and was placed in a glass container with a starter culture of bacteria and yeast with 100ml of already fermented tea to activate the fermentative process. The glass was covered with paper towels and held in place with a rubber band to avoid contamination for 10 days. Finally, a gelatinous film was obtained on the air-liquid surface and removed for further purification with distilled water and sodium hydroxide for comparison. The flow chart for obtaining the cellulosic membrane is shown in Figure 1.

![Flowchart representative of obtaining the cellulosic membrane.](source: Authors)

2.2 Purification of bacterial cellulose membranes from kombucha

Two comparative purification methods were used in the purification step: the first with distilled water and the second with sodium hydroxide NaOH. Both purifications used water bath equipment. These different ways of purifying the bacterial cellulose were used to compare results and to establish the most efficient method to obtain crystals.

2.2.1 Purification of Kombucha’s bacterial cellulose membranes (KBCM) with distilled water

From purification stage, about 400mL of distilled water was added to the KBCM and left in a water bath for 1 hour at 80°C. The process was repeated four times until a white coloration was obtained. Subsequently, this membrane was separated and placed in a petri dish to dry in an oven at 50°C for 18 hours.
2.2.2 Purification of Kombucha’s bacterial cellulose membranes (KBCM) with NaOH

The methodology of purification was based on the work of (Rangaswamy et al., 2015) and collaborators. An alkaline solution of 0.5M NaOH was prepared for purification of the kombucha bacterial cellulose membrane. Around 75mL of NaOH was added to a beaker with the KBCM. It was left in a water bath at 80°C for 15 minutes. Subsequently, to neutralize the pH, 400mL of distilled water was added to the KBCM, again in a water bath at 80°C for one hour. This purification step was repeated four times until neutralization of the pH. The steps are represented in Figure 2. Afterward, this membrane was separated and placed in a petri dish to dry in an oven at 50°C for 18 hours.

Figure 2. Alkaline purification steps.

Source: Authors

2.3 Characterization

2.3.1 Fourier Transform Infrared Absorption Spectroscopy (FTIR)

The main functional groups present in the membranes were characterized via FTIR. It was also used for comparison of the chemical composition between the KBCM in natura and purified with distilled water and NaOH. The characterization was performed in the Espectro 400 GX FT-IR equipment, in the Central Multiuser Laboratories in transmittance mode, with a range between 4000 and 500 cm⁻¹.

2.3.2 X-Ray Diffraction (XRD)

To analyze the material's crystallinity, it was necessary to characterize by X-ray diffractometer (XRD). The analysis was performed in the film mode, with a divergence slit of 0.3 mm, incidence angle theta = 1.5°, and angle 2 theta ranging between 10.0° and 80.0°, with speed 2°/min and step of 0.02°. The X-ray beam generated by the copper target with a nickel filter allows only X-rays with wavelength λ =1,5413 Å to pass through with a wide range of applications in materials engineering. The XRD technique was also used to estimate the average crystallite size using the Scherrer equation (Maiti et al., 2013).

\[ D = \frac{k\lambda}{\beta\cos\theta} \]  

Equation 1

Where: k is an empirical proportionality constant, which varies with the geometric shape of the crystallite. To measure the peak width at half-height, K varies between 0.84 and 0.89, depending on the geometry. If the geometry of the crystallite is
not known, the spherical shape is adopted, and it is worth 0.90. \( \lambda \) is the wavelength measured at the center of the peak obtained via XRD, \( \beta \) is the peak width measured at the average height, and \( \theta \) is the Bragg reflection angle measured in radians.

2.3.3 Scanning Electron Microscopy (SEM)

To analyze the morphology of the KBCM, the dry sample was coated with a gold film via sputtering. The morphology of the fibers was analyzed via SEM and was performed in the Central Multiuser Laboratory (CLM in a Carl Zeiss EVO MA 10 Scanning Electron Microscope coupled with an integrated EDX system.

3. Results and Discussion

3.1 KBCM Depigmentation

Figure 3 contains four photographs of the KBCM depigmentation process in water and four photographs of the process with NaOH. It can be comparatively observed in Figures 3(d) and 3(h) that there was discoloration in both the step with water and the step with NaOH.

![Figure 3. Depigmentation of the bacterial cellulose membrane](image)

Source: Authors

It is possible to observe in Figures 4b and 4d, the evolution of the depigmentation of the kombucha bacterial cellulose membrane; from a yellowish-brown coloration, it started to have a lighter coloration close to white; through this visualization, it is possible to observe the coloration variation between KBCM purified only with water and KBCM purified with 0.5M NaOH. The article by Laavanya et al., (2021) reports a relevance in the treatment through visual analysis. Moreover, they also visualized this depigmentation through the alkaline treatment in their research.
Figure 4. a) KBCM before purification; b) KBCM after purification; c) KBCM before purification and dried; d) KBCM after purification and dried.

Source: Authors

3.2 Fourier transform infrared spectroscopy (FTIR)

The Fourier Transform Infrared Absorption Spectroscopy was aimed at presenting the molecular structure of which the cellulose membrane is composed in natura, blue line, purified with distilled water, red line, and purified with NaOH, black line. We observe bands at 3259, 3291, and 3342 cm\(^{-1}\) representing an intramolecular hydrogen stretching vibration, which are hydroxyl groups related to the water molecule. A decrease of these water groups on the membrane is noticed through these bands as the purification proceeds. The FTIR spectrum also shows at 2899, 2918, and 2886 cm\(^{-1}\) less pronounced absorption bands, which indicates the stretching of C-H bonds, and a much smaller peak of these compounds can be seen when the membrane was purified with NaOH. In the bands 1652 and 1633 cm\(^{-1}\), it shows absorption of the C=O group. The bands at 1145, 1151, and 1106 cm\(^{-1}\), indicate the presence of the NH group. According to Li et al., (2019) and collaborators the peaks representing proteins are 1062.6-1059.1 cm\(^{-1}\); however, in this work, these peaks for the membrane proteins were 926, 1010, and 1016 cm\(^{-1}\). The results of the spectra of pure KBCM, KBCM purified with water, and KBCM purified with NaOH are shown in Figure 5. The amide and protein groups were reduced, probably due to the temperature and pH action of the NaOH treatment.
3.3 X-Ray Diffraction (XRD)

Crystallized cellulose is found in four polymorphic phases, but two are widely studied: type I - cellulose (native) and type II - cellulose (regenerated). Native cellulose is the most common and can be divided into Iα and Iβ. KBCM has typical characteristics of type Iα cellulose, as observed in the diffractogram of Figure 6, the prominent peaks at 14.5, 16.6, and 22.6, typical values of type I crystalline cellulose with structures (-1 0 1), (1 0 1), and (0 0 2), according to previous studies by Karina et al., (2012). There was an increase in the peaks when the membrane was subjected to alkaline treatment compared to the low peaks of pure KBCM; therefore, the membrane showed an increase in crystallinity. The X-ray diffractometers of Pure and NaOH-purified KBCM are shown in Figure 6.
Table 1 contains the data and the estimated average crystallite size for the pure KBCM samples, purified with distilled water + heating and purified with 0.5molar NaOH solution + heating. Note that the crystalline planes that intensified the most after the purification process were (0 0 2) and (1 0 1), with the (0 0 2) plane showing the most remarkable change in diameter, indicating greater efficiency in the purification process with NaOH.

Table 1. Data and estimated average crystallite size for pure and purified KBCM samples with water and NaOH

| Samples      | K   | L(μm) | Kα(μm) | β (FWHM)* | θ   | θcosθ | D=Kλ (θcosθ)-1 (nm) | Crystalline planes |
|--------------|-----|-------|--------|-----------|-----|-------|---------------------|-------------------|
| KBCM-H2O     | 0.90| 15.41 | 13.87  | 3.20      | 14.60 | 0.97  | 4.48                | (-1 0 1)          |
| KBCM-NaOH    | 0.90| 15.41 | 13.87  | 2.58      | 14.61 | 0.97  | 5.56                | (0 0 2)           |
| KBCM-pur2    | 0.90| 15.41 | 13.87  | 2.04      | 22.63 | 0.92  | 7.37                | (0 0 2)           |
| KBCM-pur3    | 0.90| 15.41 | 13.87  | 2.18      | 22.51 | 0.92  | 6.89                | (0 0 2)           |

* Full width at half maximum of the peak (FWHM). Source: Authors.

3.4 Scanning Electron Microscopy

The morphology of the cellulosic membrane was analyzed by the SEM technique and is represented in Figure 7. One can observe in Figure 7(a) the morphology of KBCM in natura, (b) Morphology of KBCM purified with water (c), and Morphology of KBCM purified with NaOH (d). The presence of bacteria of the genus Gluconacetobacter hansenii, which produces the cellulose fibrils, can be observed. After alkaline purification, no more bacteria could be seen, as they were removed from the membrane. In Figures 7c and 7d, only a tangle of fibrils randomly disordered with respect to directions was noticed. It
was reported in the work of Souza and Recouvreux, (2016) the randomness of cellulosic fibrils produced by the bacterium *Gluconacetobacter hansenii*.

**Figure 7.** KBCM morphology a) and b) Morphology of KBCM purified with water c) and d) Morphology of KBCM purified with NaOH.

4. Conclusion

It was possible to obtain bacterial cellulose membrane by fermenting the millennial beverage kombucha, made from sweetened tea with sucrose and a symbiotic culture of bacteria and yeast. This membrane was separated for treatment with distilled water and weak alkaline treatment of 0.5M, where it was possible to dry these membranes and characterize them by FTIR, XRD, and SEM techniques to analyze the molecular composition, crystallinity, and morphology of the fibrils. The results obtained were satisfactory, decreasing the water band of the biomaterial, increasing the crystallinity, and removing impurities in the membrane, such as bacteria. With the purification performed, it will be possible to further study the material for use as a product, such as in biomedicine and tissue engineering, as a biomaterial reinforcement.

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