Fungal chitosan as membranous material modified by atmospheric plasma

Quitosana fúngica como material membranoso modificado por plasma atmosférico

El quitosano fúngico como material membranoso modificado por plasma atmosférico

Abstract

Objective: This study produced a fungal chitosan membrane extracted from *Rhizopus stolonifer*, as well as its modification using dielectric barrier discharge plasma (DBD), aiming to improve the physicochemical characteristics of the membrane, optimizing its use in the medical research field. Method: The obtained chitosan was physically and chemically characterized (Molecular Weight, Fourier Transform Infrared, X-ray Diffraction), later were produced fungal chitosan membranes and DBD plasma was applied. The membranes were characterized before and after plasma application using the tests contact angle, swelling and atomic force microscopy (medium roughness) analyzes. Results: A fungal chitosan with a yield of 16.73 mg/g, and an apparent molecular weight of 4 kDa was obtained, being considered of low molecular weight and high degree of deacetylation (84%). It was possible to obtain the membrane and after application of DBD plasma, the contact angle dropped from 77.5° to 30.9°, making it more hydrophilic. Conclusion: Thus, the efficiency of the technique for increasing the hydrophilicity of the fungal chitosan membrane without the additive of chemical reagents during the process was confirmed and the membrane formed is a promising alternative can be used in different ways in the medical area.

Keywords: Biopolymers; *Rhizopus stolonifer*; Fungal chitosan; Dielectric barrier discharge; DBD plasma.

Resumo

Objetivo: Este estudo produziu uma membrana fúngica de quitosana extraída de *Rhizopus stolonifer*, bem como sua modificação utilizando plasma de descarga de barreira dielétrica (DBD), visando melhorar as características físico-químicas da membrana, otimizando seu uso na área de pesquisa médica. Método: O quitosano obtido foi caracterizado física e quimicamente (Peso Molecular, Transformada de Fourier no Infravermelho, Difração de Raios X), posteriormente foram produzidas membranas de quitosana fúngica e aplicado plasma DBD. As membranas foram caracterizadas antes e após a aplicação do plasma por meio dos testes de ângulo de contato, intumescimento e microscopia de força atômica (rugosidade média). Resultados: Foi obtida uma quitosana fúngica com rendimento de 16,73 mg/g, e peso molecular aparente de 4 kDa, sendo considerada de baixo peso molecular e alto grau de desacetilação (84%). Foi possível obter a membrana e após a aplicação do plasma DBD, o ângulo de contato caiu de 77,5° para 30,9°, tornando-o mais hidrofílico. Conclusão: Assim, foi confirmada a eficiência da técnica para aumentar a hidrofílicidade da membrana fúngica de quitosana sem o aditivo de reagentes químicos durante o processo e a membrana formada é uma alternativa promissora podendo ser utilizada de diferentes formas na área médica.

Palavras-chave: Biopolímeros; *Rhizopus stolonifer*; Quitosana fúngica; Descarga de barreira dielétrica; Plasma DBD.

Resumen

Objetivo: Este estudio produjo una membrana fúngica de quitosano extraída de *Rhizopus stolonifer*, así como su modificación mediante plasma de descarga de barrera dieléctrica (DBD), con el objetivo de mejorar las características físico-químicas de la membrana, optimizando su uso en el campo de la investigación médica. Método: El quitosano obtenido se caracterizó física y químicamente (Peso Molecular, Transformación de Fourier en infrarrojos, Difracción de Rayos X), posteriormente se produjeron membranas de quitosano fúngico y se aplicó plasma DBD. Las
membranas se caracterizaron antes y después de la aplicación del plasma mediante ensayos de ángulo de contacto, hinchamiento y microscopía de fuerza atómica (rugosidad media). Resultados: Se obtuvo un quitosano fúngico con un rendimiento de 16,73 mg / g y un peso molecular aparente de 4 kDa, considerándose de bajo peso molecular y alto grado de desacetilación (84%). Se pudo obtener la membrana y luego de aplicar el plasma DBD, el ángulo de contacto bajó de 77.5 ° a 30.9 °, haciéndola más hidrofílica. Conclusión: Así, se confirmó la eficacia de la técnica para incrementar la hidrofobicidad de la membrana fúngica de quitosano sin la adición de reactivos químicos durante el proceso y la membrana formada es una alternativa prometedora que puede ser utilizada de diferentes formas en el campo médico.

Palabras clave: Biopolímeros; Rhizopus stolonifer; Quitosano fúngico; Descarga de barrera dieléctrica; Plasma DBD.

1. Introducción

El quitosano es un copolímero formado a partir de la desacetilación de algunos residuos de glicopyranosa de chitosan (Annu et al., 2017). Tiene 2-acetamido-2-deox-D-glycopyranose y 2-amino-2-deox-D-glycopyranose unidades. Sin embargo, esta molécula se considera quitosano cuando más del 50% de los residuos de polímeros son 2-amino-2-deox-D-glycopyranose (Anwar et al., 2017). Cuando químicamente obtenido desde el chitín presente en la carapaza de crustáceos es conocido como un quitosano animal (Polymar, 2020). Además, el quitosano también puede ser obtenido extrayéndolo directamente desde la pared celular de hongos, especialmente aquellos que pertenecen a la clase de Zygomycetes (Paiva et al., 2017). El quitosano animal es el más comercialmente utilizado debido a su rendimiento que es aproximadamente 2 a 3 veces más alto que el del quitosano fúngico. Sin embargo, su purificación y procesamiento requiere un gran volumen de reactivos químicos, lo que aumenta los costos de producción y disminuye el rendimiento (Bento et al., 2011). Además, el quitosano animal depende de los factores estacionales de la región donde se cría el crustáceo, lo que dificulta la estandarización y el desarrollo industrial (Ghormade et al., 2017). Cuando se refiere a uso en alimentos y medicamentos, se sugiere otro problema: la potencialidad alérgica de algunos proteínas como tropomiosina y arginina que se acumulan en el quitosano cuando no se purifica correctamente (Stamford et al., 2007; Pascal et al., 2015; Faber et al., 2017).

La utilización de hongos para la producción de quitosano tiene algunos beneficios sobre el proceso tradicional. (1) Los hongos se pueden cultivar con simples y Economicamente viables sustratos, como residuos industriales y alimentos, reduciendo los costos de producción; (2) El quitosano fúngico puede ser producido durante el año sin restricciones cuando se usa un entorno controlado, garantizando un proceso continuo de producción (3) Tiene bajos contenidos de calcio y otros iones, lo que reduce el riesgo de acumulación de minerales en el cuerpo (Arcidiacono et al., 1989; Synowiecki & Ali-Khatteeb, 2003; Queiroz et al., 2015; Batista et al., 2018).

En los últimos años, algunas investigaciones han mostrado el potencial de quitosano fúngico, como antifungal, antibacterial, anticancer and antiparasitic action (Paiva et al., 2014; Souza Neto et al., 2017; Almutairi et al., 2020; Batista et al., 2020; Sathiyaseelan et al., 2020). Por estas razones, el quitosano fúngico ya se está produciendo y comercializando por compañías en diferentes países, como Bélgica, Canadá y Estados Unidos. Todo para aplicaciones médicas o farmacéuticas (Kitozyme, 2020; Mycodev, 2020).

Estudios han demostrado que el grado de acetilación y peso molecular del quitosano tiene un directo influencia en la absorción de líquidos, el mayor grado de acetilación (entre 80-90%) y molecular weight, el mayor el rate de water absorption (Schipper et al., 1996; Pankaj, 2015; Chumwangwapee et al., 2016). Esto se explica por la predominancia de amino groups en el quitosano que da este biopolímero una alta polaridad (Assis et al., 2002).

Debido a interferencias de características fisicoquímicas, para mantener las posibilidades de uso de biomateriales, modificaciones son requeridas. Técnicas de modificación biomaterial convencionales son tratamientos solventes, soluciones base acídica, abrasión mecánica y activación química, favoreciendo la parametrización de polisacáridos, oscilaciones de cadenas y producción de desperdicio tóxico.

Para resolver los problemas de técnicas convencionales, el uso de plasma de atmósfera fría, un gas ionizado débil operando a condiciones de atmósfera normal, se ha desarrollado. Under estas condiciones, solo una fracción pequeña de átomos que cargan y moleculas de gas colisionan con electrones de alta energía, resultando en un aumento de la excitación, ionización, y
dissociation, while the plasma remains "cold" (Klampl et al., 2012). Among the chemical clusters identified in atmospheric plasma are molecules (OH, NO, CN, O3), atomic radicals (H, O, N), and other species of active groups: such as N2+, which has properties that the gas alone doesn’t have, such as magnetism and conductivity (Machala et al., 2007).

Treating materials with atmospheric plasma give to the materials characteristics for their use in different areas. Among these characteristics are: (1) ability of its action to be restricted to the surface of materials without affecting their mechanical properties; (2) introduction of reactive radicals and their conversion into functional groups; (3) do not require high cost vacuum or compression systems; (4) be environmentally friendly as its use generates less environmentally toxic compared to chemical methods of material modification; (5) antimicrobial effect of Escherichia coli and Bacillus subtilis endospores, Staphylococcus aureus and Clostridium difficile cells (Hegemann et al., 2003; Hong et al., 2009; Chen et al., 2010; Salem et al., 2011; Sasmazel, 2011; Galvin et al., 2013; Shahidi et al., 2015; Ren et al., 2017).

One of the most versatile types of atmospheric plasma is dielectric barrier discharge (DBD) plasma, which consists of a discharge occurring in a space between two metal electrodes, where at least one dielectric are inserted between these electrodes. By adding a potential difference between the electrodes, the electric charges begin to accumulate on the dielectric surface. At some point, the charge accumulation on the surface reaches its limit, breaking the dielectric rigidity of the gas, causing micro-discharge, which is what we call plasma. This process is repeated every half cycle of the applied voltage pulse between the metal electrodes (Bogaerts et al., 2002).

DBD has been shown to be an effective method for producing homogeneous unbalanced plasma at atmospheric pressure with simple operation system. Surface modification of materials such as grafting, surface cleaning and hydrophilic polymer modification, powders and seed surface modification is widely used (Morent et al., 2011; Molina et al., 2014). In a view of the diversity of applications of fungal chitosan and the advantages of the treatment of biopolymers by dielectric barrier discharge plasma. The present work aims to treat fungal chitosan membranes with atmospheric plasma in order to observe how the membrane hydrophilicity and liquid absorption, will behave after the treatment. Thus, we intend Aiming to potentiate the use of fungal chitosan, leaving it in optimal conditions for the application in different medical areas.

2. Methodology

2.1 Fungal Collection, Cultivation and Chitosan Extraction

Soil samples were collected at 10 different points of the ESEC Seridó reserve (Seridó Ecological Station), under the license number 36672-1 SISBIO / ICMBio, in the county of Serra Negra do Norte / RN / Brazil, at the geographic coordinates 06° 35’ e 06° 40’ South, and 37° 20’ and 37° 39’ West. After monospore growth, the fungi were isolated and identified as Rhizopus stolonifer (Vital & Zilli 2010).

Petri dishes with BDA synthetic medium (Potato Dextrose Agar) and monospore growth were incubated at 28° C for 72 hours. The spores were collected and stored in 15 mL of sterile distilled water (standard solution). About 105 spores/mL of the standard solution were added in 400 mL of YPD (Yeast Extract 10 g; Peptone 20 g; Dextrose 20 g per liter) and incubated at 28° C/ 96 hours in static mode (Stamford et al., 2007). Biomass was filtered, lyophilized, and chitosan was extracted according Hu et al. (1999).

2.2 Fungal chitosan Characterization

Size exclusion chromatography was applied to determine the apparent molecular weight of chitosan. 7.8 x 300 mm Ultrahydrogel 500 and 250 columns (Waters Corp., Milford, Massachusetts, USA), connected in series, were coupled to an Accela® refractive index detector HPLC (Thermo Scientific, Waltham, Massachusetts, USA). The eluent was filtered (0.22 μm membrane) in ultrapure water with 0.1 M NaNO3, with a flow rate of 0.6 mL / min at 30 °C. A set of dextran standards (4,
6, 10, 40, 72.1, 147 and 270 kDa) was used to construct the standard curve and to determine the apparent molecular weight of the biopolymer (Galinari et al., 2017).

Fourier Transform Infrared Spectroscopy (FTIR) were performed with a Shimadzu spectrophotometer model IR Prestige-21, with attenuated total reflectance accessory (from English, ATR) coupled with the following analysis conditions: Region 4000 - 600 cm\(^{-1}\); Resolution: 4 cm\(^{-1}\); Accumulations: 20; Mode: transmittance. For the checking the degree of deacetylation of the samples, formula (1) was applied (Brugnerotto, 2001).

\[
DA = \frac{[(A1320/A1420) – 0.3822]}{0.03133} \quad (1)
\]

\[DD = 100 - DA\]

X-ray diffraction measurements were performed based on the technique of Signini & Campana Filho (2001) with modifications. It was used a SHIMADZU X-ray diffractometer (Model XRC-6000) with copper tube (\(\lambda = 1.54 \text{ Å}\)). The voltage and current used were of the order of 40 kV and 40 mA, respectively. These measurements were performed at a range of 3-50 °C with a scan rate of 1° / minute in 0.02° steps (Signini & Campana Filho, 2001).

### 2.3 Biomembrane Production and Application of the Dielectric Barrier Discharge Plasma (DBD)

The powder was dissolved in 2% acetic acid with constant stirring for 24 hours. After this period, the chitosan solution were filtered twice, the first time with a nylon filter and the second with a 0.45 mm filter (Millipore, Billerica, MA, USA) (Macedo et al., 2012).

The application of plasma to fungal chitosan membranes was performed according to the methodology proposed by Napartovich et al. (2001) with modifications. Plasma was generated using 1 L / min flow Helium gas and applied as describes in Table 1. The dielectric barrier discharge plasma generation system can be seen in Paiva (2017).

#### Table 1: Parameters of dielectric barrier plasma (DBD) applications and the relationships between wettability and average roughness for the different treatments analyzed.

| Treatments | Voltage(kV)/Time(min)* | Wettability  | Average roughness (nm) |
|------------|------------------------|--------------|------------------------|
| 1          | 15/10                  | 69.5º ± 0.51\(^a\) | 624.76 ± 7.37\(^b\) |
| 2          | 15/5                   | 39.5º ± 0.57\(^b\) | 1346.05 ± 5.84\(^b\) |
| 3          | 15/1                   | 30.9º ± 0.57\(^b\) | 2118.78 ± 2.31\(^*\) |
| 4          | 30/10                  | 64.0º ± 0.50\(^*\) | 732.82 ± 4.59\(^b\) |
| 5          | 30/5                   | 58.5º ± 0.80\(^b\) | 1023.57 ± 2.72\(^b\) |
| 6          | 30/1                   | 57.3º ± 0.56\(^b\) | 847.25 ± 2.61\(^b\) |
| 7          | No treatment           | 77.5º ± 0.82\(^*\) | 668.30 ± 3.43\(^b\) |

\(^a\) All tests were at the frequency of 500 Hz; \(^ab\) Different letters denote statistically different data.

Source: Authors.

### 2.4 Characterization of the Chitosan Membrane

For the analysis of the wettability, the sessile drop method was chosen, where a drop was deposited on the surface of
the sample. The samples were placed on the flat base and then a drop of 10 μl of distilled water was deposited on the surface of the membranes. The researchers monitored the process in real time, and saved it on a computer using the Surftens 4.5 software. After, the images were saved for the desired instants: 0, 10, 20, 30, 40, 50 and 60 seconds and analyzed by the same software. The contact angle values presented in this study reflected the average of the measurements of three images. Five measurements were taken for each image (Alves Jr. et al., 2016).

The degree of swelling or absorption of the fungal chitosan membranes was evaluated by immersing them in PBS solution (Phosphate Buffered Saline pH ~ 7.4). The test was performed in triplicate for each assay. After a 24-hour immersion period the samples were removed from the PBS solution and weighed in analytical balance (excess liquid was removed with paper towels). Each membrane was weighed 4 times to reduce the risk of weighing errors.

The degree of swelling was evaluated as the mass percentage gain of the wet samples and was calculated by:

Equation 2

\[
\text{Loss of Mass (\%) } = \left( \frac{\text{inicial mass} - \text{final mass}}{\text{inicial mass}} \right) \times 100
\]

To analyze the roughness and topography of the chitosan membrane samples, a Shimadzu Corp. atomic force microscope, model SPM-9600 (Japan) was used. The equipment was used in dynamic or intermittent mode at a scan rate of 1 Hz. Random areas of 10 μm x 10 μm were scanned and analyzed by SPM Manager software Version 3.4 (Japan). The parameters of arithmetic average roughness (Ra) were analyzed. The analysis of the topography was done through the 3D images obtained in the AFM (Macedo et al., 2012).

2.5 Statistical analysis

For the statistical analysis of the contact angles and mean roughness, the ANOVA test was used to measure the variability between the values and then the Tukey test was used to evaluate if there was a significant difference between the means. The researchers performed all tests in triplicate and analyzed by the PAST program.

3. Results and Discussion

3.1 Characterization of fungal chitosan

The edaphic samples from Seridó Ecological Station was identified as R. stolonifer and it generated a yielded of chitosan of 16.73 mg/g by dry biomass. The chitosan produced was characterized with molecular weight of 4.12 kDa, degree of deacetylation of 84%. The molecular weight obtained by a High-performance liquid chromatography and stable in solution, was characterized. Indicating whether it is a polymer with a low molecular weight and a low colloid formation capacity.

These results contrast with those found by Cardoso et al. (2012) and Berger et al. (2018) within a respective yield of 29.3 mg/g and 49.31 mg/g of chitosan for R. arrhizus. Although they are all from the same class of Zygomycetes, they have specific genetic differences and the carbon and nitrogen source used in the culture medium may also have influenced the production of fungal chitosan.

3.1.1 Fourier Transform Infrared and X-ray

The Fourier transform infrared characterize the chitosan as for the degree of deacetylation (DD) and it is considered one of the most important chemical characteristics that can influence solubility, chemical reactivity and biodegradability. Fungal chitosan with high degree of deacetylation has a high-density positive charge due to the free amine groups, making it unique for several biological applications (Berger et al., 2014; Abdel Gaward et al., 2017). Freier et al. (2005) demonstrated
how this degree favors a low degradation of the biopolymer, as well as the cell adhesion when close to a 100%. Huang et al. (2004) showed that the greater the degree of deacetylation, the greater the absorption of nanoparticles by fibroblasts in cell culture.

The DD of 84% estimated by the FT-IR in this study characterizes the polymer as chitosan, according to the EUCHIS (European Chitin Society) (Figure 1A). Indicating that this biopolymer has few protonated groups in its chain and has an amine group in the carbon 2 in more than 60% of its residues (Zhang et al., 2017). These results are similar to those obtained by Gharieb et al. (2015) who produced chitosan from the fungi species Mucor rouxxi, Rhizopus sp., Cunninghamella elegans, with DD of 80.3%, 81.5%, 80.3%, respectively. However, compared to the degree of deacetylation of others fungi chitosan, the results contrast with the ones found by Mondala et al. (2015) and Zimoch-Korzycka et al. (2016) who obtained a degree of deacetylation of 60% and 72%, respectively, using Mucor rouxxi, and Aspergillus niger. These contrasts indicate that the degree of deacetylation varies with the type of fungus and the culture medium, and this variation can be about 30%. It is important to note that there are no studies to characterize the fungus Rhizopus stolonifer, so the comparisons are between others same fungi class, Zygomycetes.

The X-ray diffraction in the sample extracted from Rhizopus stolonifer, demonstrated the presence of a peak around 20° indicating the crystalline region of the material, and a 9° peak indicating the amorphous region of chitosan (Figure 1B).

The results of diffraction Raio-X corroborate with studies founded which confirm these peaks as characteristic regions of chitosan, be it fungal or animal. Associated with the FT-IR result, it is indicated that the extracted polymer is effectively chitosan (Berger et al., 2014; Berger et al., 2018; Zhang et al., 2017).

**Figure 1.** Structural analysis of the fungal chitosan molecule by using the Fourier transform infrared spectroscopy technique (A) and X-ray diffraction technique (B). (A) Peaks ~3378.25 cm⁻¹ (O-H stretch), ~2922.00 cm⁻¹ (C-H stretch), ~1644.94 (carbonyl group C = O (amide I)), ~1417.14 (amide I and II) and ~1030.49 cm⁻¹ (saccharide structure). (B) Peak around 20° indicating the crystalline region and a 9° peak indicating the amorphous region of fungal chitosan.

3.2 Production and characterization of the fungal chitosan membrane

The chitosan membrane was produced according to the proposed methodology and afterwards it was divided into pieces measuring 2x2 cm to carry out the characterizations before and after treatment by DBD plasma (Figure 2).
The treatments 2 and 3 (described in Table 1) obtained a significant lower contact angle than the untreated sample, indicating that atmospheric plasma increased wettability in those specific conditions (Figure 3).

Statistical analysis of the data indicated a significant difference ($p < 0.05$) between the contact angles of treatments 2 and 3, in comparison to the others, showing that the treatments had the better effect. When performing the Tukey test, the difference between the contact angles was obtained on the 6 different treatments.

The measurement of the contact angle of a liquid with the surface of a sample has its importance in the correlation of solubility of that sample in that liquid. Based on this idea, the use of DBD plasma in chitosan membranes favor the incorporate
polar functional groups on these surface, which can alter its absorption capacity and interfere in the contact angle.

Zhang et al. (2012) demonstrated, through mass spectrometry, that when using DBD plasma formed by helium gas there is a promotion of the formation of a larger number of reactive groups that act modifying the surface and leaving the surface layer of the membrane more hydrophilic. Among these reactive groups are: N+, O+, OH+, H2O+, H3O+, N2+, N2H+, NO+, O2+, N2O+, NO2+, O-, OH-, H3O-, O2-, NO2-, O-(H2O)2, CO3-, HCO3-, NO3-.

In addition to helium gas experiments some authors are using atmospheric air at room temperature. Theapsak et al. (2012) and Dorraki et al. (2015) managed to promote the formation of polar reactives on the surface of membranes containing oxide polyethylene and animal chitosan. These authors demonstrated that a reduction of 50.52% and 40.57%, respectively, in the contact angle for solubilization in water when under conditions 15 kV / 10 sec / 350 Hz and 14 kV / 6 min / 6 Hz, respectively. Thus, they indicate the availability of using DBD plasma in the preparation of membranes containing chitosan for medical and food use among others. In our study, the better treatment the contact angle going from 77.5° to 30.9°, a reduction of 39.87% (15 kV / 1 min / 500 Hz treatment 3, Table 1) demonstrates the economic viability of using this technique to also increase the hydrophilicity of a fungal chitosan membrane.

An increase in the wettability of a membrane favors the hypothesis of its great importance for a drug delivery application. Such an application requires a material that readily absorbs the drug to be applied, in other words, to a more hydrophilic material.

To demonstrate the effects of plasma on the surface of the fungal chitosan membrane, an atomic force microscopy was performed on the membranes. The average roughness of the samples was measured and this analysis being the average of the peaks found on the surface of the treatment.

The untreated membrane had an average roughness of 668.61 nm, with topography with peaks concentrated in a single region (Figure 4A). The treatment 3 (Figure 4B) resulted in a higher average roughness value of the fungal chitosan membrane, reaching a value of 2118.78 nm, showing an increase in roughness after treatment with DBD plasma.

The increase of the average roughness indicates a surface modification of the membranous material, which is also evidenced by the topographic modification visible in the images, related to the wettability. The results demonstrate that the higher the average roughness, the greater the wettability of the membrane, in other words, the more hydrophilic the membrane will be (Table 1).

Our data contrast with the work of Marques et al. (2016) which obtained a mean roughness of 4.0 nm of membrane produced from animal chitosan with a degree of deacetylation of 88%. The studies of Cleymand et al. (2016) which obtained a mean roughness of 9.32 nm in the animal chitosan membrane with 85.9% deacetylation associated with nanoliposomes; and with Tamburaci & Tihminlioglu (2017), obtained a result of 2.38 nm for roughness of the composite of chitosan animal and diatomite. These differences can be explained by the origin of chitosan, degree of deacetylation, molecular weight and mode of obtaining the membrane. Since, according to its characteristics, the polymer can behave in different ways. A high degree of deacetylation (above 70%) influences permeation and absorption, thus increasing the average roughness (Pankaj, 2015). Macedo et al. (2012) obtained chitosan membranes with high roughness when treated with a low pressure plasma with different atmospheres. The results were 4800, 5911 and 5062 nm for the atmospheres of (O2, N2, H2), respectively. When nitrogen plasma was used, the surface of the membrane became more uneven, demonstrating that there was more superficial modification of the membrane.
4. Conclusion

According to the results, it was possible to obtain a good quality chitosan from the fungi *Rhizopus stolonifer*, as well as its transformation into membranous material. In addition, the use of the DBD plasma technique at room temperature with atmospheric air for the addition of polar chemical reagents resulted in an increase in the average of the roughness and wettability of the formed membrane. Thus, the efficiency of the technique to increase the hydrophilicity of the chitosan fungal membrane without the addition of chemical reagents during the process has been confirmed and the formed membrane is a promising alternative that can be used in different ways in the medical field.

For additional analysis, we suggested *in vitro* tests with the membranous to confirm its antimicrobial potential, cellular regeneration and cellular adhesion as described by literature for different membranous of chitosan (Braga et al., 2019; Rosendo et al., 2020).

Credit authorship contribution statement

The work was carried out jointly in association with authors from different universities and areas, with the following division of tasks: Conceptualization – Wesley de Souza Paiva; Data curation – Wesley de Souza Paiva; Formal analysis – Wesley de Souza Paiva, Francisco Ernesto de Souza Neto and Erika de Souza Paiva; Funding acquisition – Anabelle Camarotti de Lima Batista; Investigation – Wesley de Souza Paiva; Methodology - Anabelle Camarotti de Lima Batista, Wesley de Souza Paiva and Francisco Ernesto de Souza Neto; Project administration - Anabelle Camarotti de Lima Batista; Resources - Anabelle Camarotti de Lima Batista; Software - Anabelle Camarotti de Lima Batista, Wesley de Souza Paiva and Francisco Ernesto de Souza Neto; Supervision - Anabelle Camarotti de Lima Batista; Validation - Anabelle Camarotti de Lima Batista, Wesley de Souza Paiva, Francisco Ernesto de Souza Neto and Erika de Souza Paiva; Writing – original draft - Wesley de Souza Paiva; Writing – review & editing - Anabelle Camarotti de Lima Batista, Wesley de Souza Paiva, Francisco Ernesto de Souza Neto and Erika de Souza Paiva.
Acknowledgments

To Coordination for the Improvement of Higher Education Personnel (CAPES – Brazil) for the scholarships; to Federal Rural University of Semiárido (UFERSA), Mossoró, Rio Grande do Norte; and to Federal University of Paraíba (UFPB), Bananeiras, Paraíba, for being of great importance for the execution of the research.

Conflicts of Interest

The authors declare no conflict of interest.

References

Almutairi, F. M., El Rabey, H. A., Tayel, A. A., Alalawy, A. I., Al-Duaib, M. A., Sakran, M. I. & Zidan, N. S. (2020). Augmented anticancer activity of curcumin loaded fungal chitosan nanoparticles. *Int J Biol Macromol*, 155, 861-867.

Alves Jr, C., Vitoriano, J. O., Silva, D. L. S., Farias, M. L. & Dantas, N. B. L. (2016). Water uptake mechanism and germination of Erythrina velutina seeds treated with atmospheric plasma. *Sci Rep*, 6, 1-7.

Annu, S. A., Ahmed, S. & Ikram, S. (2017). Chitin and chitosan: history, composition and properties. In Chitosan: derivatives, composites and applications, Ed. Ahmed, S., Ikram, S., Beverly, M. A.: Scrivener Publishing, Wiley.

Anwar, M., Anggраeni, A. S. & Al Amin, M. H. (2017). Comparison of green method for chitin deacetylation Cite as: AIP Conference Proceedings, 1823, 020071-1.

Assis, O. B. G., Vieira, D. C., Vasques, R. A. & Campana-Filho, S. P. (2002). Formed-in-place chitosan-carboxymethyl cellulose supported microfiltration membranes for water purification. in: Proceedings of the 4th ISNAPOL (Natural Polymers and Composites IV), 2002, 341.

Arcidiacono, S., Lombardi, S. J. & Kaplan, D. L. (1989). Fermentation, processing and enzyme characterization for chitosan biosynthesis by Mucor rouxii. In: Sjak-Braek, G., Anthonse, T., Sandford, P. Chitin and chitosan: sources, chemistry, biochemistry, physical properties and applications. Elsevier, 1989. 835.

Batista, A. C. L., Souza Neto, F. E. & Paiva, W. S. (2018). Review of fungal chitosan: past, present and perspectives in Brazil. *Polímeros*, 28, 275-283.

Batista, A. C. L., Melo, T. B. L., Paiva, W. S., Souza, F. S., & Campos-Takaki, G. M. (2020). Economic microbiological conversion of agroindustrial wastes to fungi chitosan. *An Acad Bras Ciênc*, 92, 1-13.

Bento, A. R., Stamford, T. L. M., Stamford, T. C. M., Andrade, S. A. C. & Souza, E. L. (2011) Sensory evaluation and inhibition of Listeria monocytogenes in bovine pâté added of chitosan from Mucor rouxii. *Leb-WissenTech*, 44, 588-591.

Berger, L. R. R., Stamford, T. C. M., Stamford-Arnaud, T. M., Alcantara, S. R., Silva, A. C., Silva, A. M., Nascimento, A. E. & Campos-Takaki, G. M. (2014). Green conversion of agroindustrial wastes into chitin and chitosan by Rhizopus arrhizus and Cunninghamella elegans strains. *Int J Mol Sci*, 15, 9082-9102.

Berger, L. R. R., Stamford, T. C. M., Oliveira, K. A. R., Pessoa, A. M. P., Lima, M. A. B., Pintado, M. M. E., Câmara, M. P. S., Franco, L. O., Magnani, M. & Souza, E. L. (2018). Chitosan produced from mucorales fungi using agroindustrial by-products and its efficacy to inhibit colletotrichum species. *Int J Biol Macromol*, 108, 635–641.

Bogaerts, A., Neyts, E., Gibjels & Mullen, R. J. V. (2002). Gas discharge plasmas and their applications. *Spectrochim Acta Part B at Spectros*, 57, 609-658.

Braga, L. A. S., Flauzino Junior, A., González, M. E. L., & Queiroz, A. A. A. de. (2019). Membranas termossensíveis baseadas em redes poliméricas semi-interpenetrantes de Quitosano e Poli(N-isopropilacrilamida). *Res Soc Dev*, 8, e3883748.

Brugnerotto, J., Lizardi, J., Goycoolea, F. M., Argüelles-Monal, W., Desbiériès, J., & Rinaudo, M. (2001). An infrared investigation in relation with chitin and chitosan characterization. *Polymmer*, 42, 3569–3580.

Cardoso, A., Lins, C. I., Santos, E. R., Silva, M. C. & Campos-Takaki, G. M. (2012). Microbial enhance of chitosan production by Rhizopus arrhizus using agroindustrial substrates. *Molecules*, 17, 4904-4914.

Chen, C., Ogino, A., Wang, X. & Nagatsu, M. (2010). Plasma treatment of multiwall carbon nanotubes for dispersion improvement in water, *Appl Phys Lett*, 96, 131504-131504-2.

Chumwungwaeep, S., Chingsungnoen, A. & Siric, S. A. (2016). A plasma modified cellulose-chitosan porous membrane allows efficient DNA binding and provides antibacterial properties: A step towards developing a new DNA collecting card. *Forensic Sci Int Genet*, 25, 19-25.

Cleymand, F., Zhang, H., Dostert, G., Menu, P., Arab-Tehrany, E., Velot, E. & Mano, J. F. (2016). Membranes combining chitosan and natural-origin nanopolysomes for tissue engineering. *RSC*, 6, 83626–83637.

Dorraki, N., Safa, N. N., Jahanfar, M., Ghomi, H. & Ranaei-Siadat, S. (2015). Surface modification of chitosan/PEO nanofibers by air dielectric barrier discharge plasma for acetylcholinesterase immobilization. *Appl Surf Sci*, 349, 940-947.

Faber, M. A., Pascal, M., El Kharbouchi, O., Sabato, V., Hagendorens, M. M., Decuyper, I. I., Bridts, C. H. & Ebo, D. G. (2017). Shellfish allergens: tropomyosin and beyond. *Allergy*, 72, 842-848.
Freier, T., Koh, H. S., Kazazian, K. & Shoichet, M. S. (2005). Controlling cell adhesion and degradation of chitosan films by N-acetylation. Biomaterials, 26, 5872-5872.

Galimari, E., Sabry, D. A., Sassaki, G. L., Macedo, G. B., Passos, F. M. L., Mantovani, H. C. & Rocha, H. A. O. (2017). Chemical structure, antiproliferative and antioxidant activities of a cell wall α-d-mannan from yeast Kluyveromyces marxianus. Carbohydr Polym, 157, 1298-1305.

Galvin, S., Cahill, O., O'Connor, N., Cafolla, A. A., Daniels, S. & Humphreys, H. (2013). The antimicrobial effects of helium and helium-air plasma on Staphylococcus aureus and Clostridium difficile. Lett Appl Microbiol, 57, 83-90.

Gharib, M. M., El-Sabbagh, S. M., Shalaby, M. A. & Darwesh, O. M. (2015). Production of chitosan from different species of zygomycetes and its antimicrobial activity. Inter J Scien Eng Res, 6, 1-5.

Ghoramade, V., Pathan, E. K. & Deshpande, M. V. (2017). Can fungi compete with marine sources for chitosan production? Int J of Biol Macromol, 104, 1415-1421.

Hegemann, D. H., Brunner, H. & Oehr, C. (2003). Plasma treatment of polymers for surface and adhesion improvement. Nucl.Instrum. Methods Phys Res, B, 208, 281-286.

Hong, Y. F., Kang, J. G., Lee, H. Y., Uh, H. S., Moon, E. & Park, Y. H. (2009). Sterilization effect of atmospheric plasma on Escherichia coli and Bacillus subtilis endospores. Lett Appl Microbiol, 48, 33-37.

Hu, K. J., Yeung, K. W., Ho, K. P. & Hu, K. (1999). Rapid extraction of high-quality chitosan from mycelia of Absidia glauca. J Food Biochem, 23, 187-196.

Huang, M., Khor, E. & Lim, L. (2003). High-quality chitosan from mycelia of Absidia glauca. Lett Appl Microbiol, 37, 472-479.

Kitozyme. (2020). Vegetal Chitosan: Unique, patented biopolymer from fungal origin. https://www.kitozyme.com/en/ingredients/chitosan/.

Macedo, M. O. C., Macedo, H. R. A., Silva, G. C., Silva, M. A. M. & Alves Jr., C. (2012). Estudo comparativo da modificação superficial de membranas de quitosana tratadas por plasma de oxigênio, nitrógeno e hidrogênio. REMAP, 7, 95–103.

Machala, Z., Janda, M., Hensel, K., Jedlovský, I., Leštinská, L., Foltin, V., Martišovít, V. & Morovová, M. (2007). Emission spectroscopy of atmospheric pressure plasmas for bio-medical and environmental applications. J. Mol Spectrosc, 243, 194–201.

Marques, J. S., Chagas, J. A. O. D., Fonseca, J. L. C. & Pereira, M. R. (2016). Comparing homogeneous and heterogeneous routes for ionic crosslinking of chitosan membranes. React Funct Polym, 103, 156–161.

Molina, R., Jovancio, P., Vilchez, S., Tranzon, T. & Solans, C. (2014). In situ chitosan gelation initiated by atmospheric plasma treatment. Carbohydr Polym, 103, 472–479.

Mondala, A., Al-Mubarak, R., Atkinson, J., Shields, S., Young, B., Senger, Y. S. & Pekarovic, J. (2015). Direct Solid-State Fermentation of Soybean Processing Residues for the Production of Fungal Chitosan by Mucor rouxii. J Chem Eng, 3, 11-21.

Morent, R., Eyter, N., Desmet, T., Dubrueil & P., Leys, C. (2011). Plasma Surface Modification of Biodegradable Polymers: A Review. Plasma Process Polyom, 8, 171-190.

Mycodev. (2020). Production. http://mycodevgroup.com.

Napartovich, A. P. (2001). Overview of Atmospheric Pressure Discharges Producing Nonthermal Plasma. Plasm Polym, 6, 1-14.

Paiva, W. S. (2017). Quitosana fúngica na produção de biomaterial membranoso modificado por plasma de descarga em barreira dielétrica (DBD). Mossoró: Federal Rural University of Semiarid, Rio Grande do Norte, Brazil. https://sgaa.ufersa.edu.br/sgaa/public/programa/defesas.jsf?k=pt_BR&id=828 .

Paiva, W. S., Souza Neto, F. E. & Batista, A. C. L. (2014). Avaliação da atividade antibacteriana da quitosana fúngica. Persp Onl: Bio Salud, 13, 37-43.

Paiva, W. S., Souza Neto, F. E. & Batista, A. C. L. (2017). Characterization of Polymeric Biomaterial Chitosan Extracted from Rhizopus stolonifer. J Polym Mater, 34, 115-121.

Pankaj, S. K., Bueno-Ferrer, C., O’Neil, L., Tiwari, B. K., Bourke, P. & Cullen, P. J. (2015). Dielectric barrier discharge atmospheric air plasma treatment of high amylole corn starch films. LWT - Food Sci Technol, 63, 1076-1082.

Pascal, M., Grishina, G., Yang, A. C., Sánchez-García, S., Lin, J., Towe1, D., Bañez, M. D., Sastre, J., Sampson, H. A. & Ayuso, R. (2015). Molecular Diagnosis of Shrimp Allergy: Efficiency of Several Allergens to Predict Clinical Reactivity. J Allergy Clin Immunol Pract, 3, 521-529.

Polymar. (2020). Nososos productos. http://www.polymar.com.br/.

Queiroz, M. F., Melo, K. R., Sabry, D. A., Sassaki, G. L. & Rocha, H. A. O. (2015). Does the Use of Chitosan Contribute to Oxalate Kidney Stone Formation? Mar drugs, 13, 141-158.

Ren, Y., Ding, Z., Wang, C., Zang, C., Zhang, Y. & Xu, L. (2017). Influence of DBD plasma pretreatment on the deposition of chitosan onto UHMWPE fiber surfaces for improvement of adhesion and dyeing properties. Appl Surf Sci, 396, 1571–1579.

Rosendo, R. A., Andrade, A. A., Figueiredo, A. B. M., Tavares, A. H. dos S., Castro, D. L. de S., Siqueira, R. R. de., Santos, A. dos., Medeiros, M. F. de., Penha, E. S. da., & Medeiros, L. A. D. M. de. (2020). Estruturas de quitosana utilizadas para regeneração óssea in vivo: uma revisão de literatura. Res Soc Dev, 9, e891974538.
Salem, T. S., Uhlmann, S., Nitschke, M., Calvimontes, A., Hund, R. & Simon, F. (2011). Modification of plasma pre-treated PET fabrics with poly-DADMAC and its surface activity towards acid dyes. Prog Org Coat, 72, 168–174.

Sasmazel, H. T. (2011). Novel hybrid scaffolds for the cultivation of osteoblast cells. Int J Biol Macromol, 49, 838-846.

Sathiyaseelan, A., Saravanakumar, K., Mariadoss, A. V. A. & Wang, M-H. (2020) Biocompatible fungal chitosan encapsulated phytogenic silver nanoparticles enhanced antidiabetic, antioxidant and antibacterial activity, Int J Biol Macromol, 15, 153-163.

Shahidi, S., Ghoranneviss, M. & Wiener, J. (2015). Improving synthetic and natural dyeability of polyester fabrics by dielectric barrier discharge, J Plast Film Sheeting, 31, 286–308.

Schipper, N. G. M., Varum, K. M. & Artursson, P. (1996). Chitosans as absorption enhancers of poorly absorbable drugs: Influence of molecular weight and degree of acetylation. Eur J Pharm Sci, 13, 1686-1692.

Signini, R. & Campana Filho, S. P. (2001). Características e propriedades de quitosanas purificadas nas formas neutra, acetato e cloridrato. Polímeros, 11, 58-64.

Silva, A. M., Stanford, T. C. M., Souza, P. M., Berger, L. R. R., Leite, M. V., Nascimento, A. E. & Campos-Takaki, G. M. (2015). Antifungal Activity of Microbiological Chitosan and Coating Treatment on Cherry Tomato (Solanum lycopersicum var. cerasiforme) to Post-Harvest Protection. Int J Curr Microbiol App Sci, 4, 228-240.

Souza Neto, F. E., Silva, H. C. A., Paiva, W. S., Torres, T. M., Rocha, A. C. P., Bezerra, A. C. D. S. & Batista, A. C. L. (2017). Quitosana fúngica sobre larvas de nematoides gastrintestinais de caprinos. Arq Inst Biol, 84, 1-5.

Stanford, T. C. M., Stanford, T. L. M., Stanford, N. P., Barros Neto, B. & Campos-Takaki, G. M. (2007). Growth of Cunninghamella elegans UCP 542 and production of chitin and chitosan using yam bean médium. Electron J Biotechnol, 10, 1-6.

Synowiecki, J. & Ali-Khateeb, N. A. A. Q. (2003). Production, properties and some new applications of chitin and its derivatives. Crit Rev Food Sci Nutrit, 43, 145-171.

Tayel, A. A. (2016). Microbial chitosan as a biopreservative for fish sausages. Int J Biol Macromol, 93, 41-46.

Theapak, S., Watthanaphanit, A., & Rujiravanit, R. (2012). Preparation of Chitosan-Coated Polyethylene Packaging Films by DBD Plasma Treatment. ACS Appl Mater Interfaces, 4, 2474-2482.

Tamburaci, S. & Tihminlioglu, F. (2017). Diatomite reinforced chitosan composite membrane as potential scaffold for guided bone regeneration. Mat Sci Eng: C, 80, 222-231.

Vital, M. J. S. & Zilli, J. E. (2010). Protocolo Básico de Coleta de Amostras de Solo para Caracterização da Diversidade Microbiana. Retrieved from: http://ppbio.inpa.gov.br/protocolos.

Zhang, Z., Xu, Z., Cheng, C., Wei, J., Lan, Y., Ni, G., Sun, Q., Qian, L., Zhang, H., Xia, W., Shen, J., Meng, Y. & Chu, P. K. (2017). Bactericidal Effects of Plasma Induced Reactive Species in Dielectric Barrier Gas–Liquid Discharge. Plasma Chem Plasma, 37, 415-431.

Zmoch-Korzycka, A., Śmieszek, A., Jarmoluk, A., Nowak, U., & Marycz, K. (2016). Potential Biomedical Application of Enzymatically Treated Alginate/Chitosan Hydrosols in Sponges—Biocompatible Scaffolds Inducing Chondrogenic Differentiation of Human Adipose Derived Multipotent Stromal Cells. Polymers, 8, 320-344.