ABSTRACT

Chitosan was modified by maleic anhydride, and it was then functionalized using heterogeneous and blending method to obtain the membrane. The results of the reaction between chitosan with maleic anhydride were signed by the new peak appears around 1475 cm\(^{-1}\) which attributed to C=C bending of the alkene. The new peak also appears at 1590 cm\(^{-1}\) which attributed to N-H bending of the amide. Chitosan-maleic anhydride membranes show microstructure of chitosan membrane with high porous density and rigidity while chitosan-maleic anhydride membranes have clusters. In addition, the thermal tenacity of membranes reached 500 °C. Modified membrane by heterogeneous and blending method have higher water uptake, ion exchange capacity, and proton conductivity than chitosan membrane. Moreover, the blending method is much more effective than the heterogeneous method that can be exhibited from ion exchange capacity and proton conductivity values of 1.08–6.38 meq g\(^{-1}\) and 1x10\(^{-3}\)–1x10\(^{-2}\) S cm\(^{-1}\), respectively. The results imply that modification of chitosan membrane with the addition of maleic anhydride using heterogeneous and blending method can be applied to proton exchange membrane.

Keywords: chitosan; membrane; maleic anhydride; heterogeneous; blending method

ABSTRAK

Kitosan dimodifikasi dengan anhidrida maleat yang selanjutnya difungsionalisasi menggunakan metode heterogen dan metode pencampuran polimer (blend) untuk menghasilkan membran. Hasil reaksi antara kitosan dan maleat anhidrida ditandai dengan munculnya puncak baru sekitar 1475 cm\(^{-1}\) yang merupakan vibrasi tekuk C=C alkena. Puncak baru juga muncul pada 1590 cm\(^{-1}\) yang merupakan vibrasi teku N-H amida. Membran kitosan-maleat anhidrida memperlihatkan mikrostruktur membrane kitosan dengan kerapatan pori dan kekakuan yang tinggi sedangkan memrantai kitosan-maleat anhidrida memiliki cluster-cluster. Sementara itu, ketahanan termal membran mencapai 500 °C. Modifikasi secara heterogen dan poli-blend kitosan dengan maleat anhidrida memiliki daya serap air, kapasitas penukar ion (KPI), dan konduktivitas proton yang lebih tinggi dibandingkan membran kitosan (10\(^{-4}\) S cm\(^{-1}\)). Bagaimanapun, metode poliblend lebih efektif daripada metode heterogen yang dapat dilihat dari nilai KPI dan konduktivitas proton yakni 1,08–6,38 meq g\(^{-1}\) dan 1x10\(^{-3}\)–1x10\(^{-2}\) S cm\(^{-1}\), 0,92–2,27 meq g\(^{-1}\) dan 1,53x10\(^{-4}\)–3,04x10\(^{-3}\) S cm\(^{-1}\). Hasil menyimpulkan bahwa modifikasi membran kitosan dengan penambahan anhidrida maleat menggunakan metode heterogen dan poliblend dapat diaplikasikan sebagai membran penukar proton.

Kata Kunci: kitosan; membran; anhidrida maleat; heterogen; metode pencampuran polimer

INTRODUCTION

Many attempts have been made to search and find the way to use energy efficiently as well as to conserve the environment. A fuel cell that can be used as a substitute for rechargeable batteries in portable applications is only Proton Exchange Membrane Fuel Cell (PEMFC). It has a simple system design and easy storage of high energy density fuel [1]. The membrane that used in PEMFC has the negative fixed charge which is usually called proton exchange membrane. The performance of this membrane can be increased by modifying the structure of polymer [2].

Some polymer with ionic charge or commonly known as polyelectrolyte can be applied as a proton exchange membrane. During this time the polymer used as a proton exchange membrane for fuel cell applications is sulfonated polytetrafluoroethylene or known as Nafion. It has a \(\text{SO}_3\text{H}\) group as a proton-exchange group. Nafion has been shown to have a

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good ability as a proton exchange membrane, but this membrane comprises tetrafluoroethylene that makes it very expensive and dangerous to the environment. Therefore, the development of new alternative proton exchange membranes that will provide improved character, environmentally friendly, and low production cost for fuel cell application is highly required [3].

Chitosan, extracted from shrimp waste, both as dissolved polymer and as cast membrane, has attracted significant interest in many potential applications as a cationic biopolymer and bioactive compound [4]. Chitosan has pointed out that cationic polyelectrolyte such as chitosan has unique character due to the presence of both amine and hydroxyl groups [5-6]. Therefore, it makes chitosan membrane an excellent material to be further developed. In the previous work of our group, chitosan was extracted and characterized from shrimp shell that obtained chitosan with a high degree of deacetylation (DD) [7].

Herein, we extend this work by reporting on maleic anhydride-modified chitosan membrane for proton exchange membrane. As known, maleic anhydride has carboxyl groups that increase the hydrophilic properties of the chitosan membrane. It means that ion exchange capacity increases if the carboxyl groups exist in the membrane. Chitosan was modified by maleic anhydride using both homogeneous and heterogeneous blending methods to improve the ion exchange capacity (IEC) and proton conductivity of proton exchange membrane.

EXPERIMENTAL SECTION

Materials

Chitosan with a deacetylation degree of 81% was obtained from previous research team by Ahmad [8]. Analytical grade reagents use in this paper were acetic acid, N,N-dimethyl formamide (DMF), maleic anhydrous, sodium hydroxide, sulfuric acid, acetone, copper(II) sulfate (CuSO₄) were purchased from the Emsure Merck, Jakarta. De-ionized water was used in all experiments.

Instrumentation

Fourier transform infrared spectroscopy (FTIR) (Shimadzu 8400) (4,000–500 cm⁻¹, resolution 4 cm⁻¹), differential thermal analysis-thermogravimetric analysis (TGA) Analyzer (Mettler Toledo) (operation condition: 30–550 °C), Scanning Electron Microscopy (SEM) (Jeol JSM-6360), and Impedance Analyzer (Agilent™ E4980A).

Procedure

Homogeneous poly-blend method

Chitosan membrane solution of maleic anhydride (CSMA) (the first polymer). Chitosans solids weighed as much as 1 g then dissolved in 50 mL of 2% solution of acetic acid. It was stirred until all chitosans soluble (homogeneous). A solution of maleic anhydride was added to mole ratio as chitosan. A solution of maleic anhydride was prepared by dissolving 0.549 g of solid maleic anhydride in 10 mL of DMF. The solution mixture was stirred for 24 h to form a solution CSMA₁. The same procedure was used for the preparation of the solution dope CSMA₂ that had a mole ratio of maleic anhydride with chitosan of 2, 3 and 4 with a mass of chitosan and maleic anhydride. The dope solution was denoted as CSMA₁, CSMA₂, CSMA₃, and CSMA₄.

Membrane solution of chitosan (CS) (the second polymer). Chitosan solution of 2% made by dissolving 1 g of solid chitosan in 50 mL of 2% solution of acetic acid that result is called CS₁ solution. The same procedure was conducted for the solution of chitosan 3, 4 and 5% by weight of each is 1.5; 2; 2.5 g, respectively. The results were denoted as CS₁, CS₂, CS₃, and CS₄.

The casting of the poly-blend membrane of CSMA-CS. Chitosan solution (CS) was added to chitosan-maleic anhydride (CSMA) with various mole ratio (Table 1.). Subsequently, the solutions were stirred for 3 h until homogenous (dope solution). The dope solutions were cast onto petri-dish and dried at room temperature for 72 h. Next, 1 N sodium hydroxide solution was added to the dry membranes in the petri-dish several times. The membranes were washed several times with deionized water to neutralize the membranes. As proton exchange application, membranes were soaked onto sulfuric acid for 24 h and neutralized by water.

Heterogeneous method

Fabrication of chitosan membrane. Chitosan was weighed as much as 2 g and then dissolved in 100 mL of acetic acid of 2% (v/v). It was then stirred for 24 h. Subsequently, chitosan solution was filtered and cast onto petri-dish. It was dried at room temperature for 72 h. Next, 1 N sodium hydroxide solution was added to the dry membranes in the petri-dish several times. To neutralized, the membranes were washed several times with deionized water.

Modification of chitosan membranes. The various mole ratio of chitosan toward to maleic anhydride solutions (1:1, 1:2, 1:3 and 1:4) was poured to chitosan membranes in an Erlenmeyer. Subsequently, it was
shaken for 24 h. The modified-chitosan membranes were washed with deionized water then with acetone to remove unreacted maleic anhydride. The membranes were then dried at room temperature. The results were denoted as CS-MA1, CS-MA2, CS-MA3, and CS-MA4.

**Water uptake measurements**

The water uptake of the membrane was determined by measuring the weight difference of membrane before and after immersion in distilled water. Samples of the membrane were dried at room temperature for 24 h and then weighted (M_{dry}). The membrane then was immersed in distilled water at room temperature for 24 h. After removing residual water from the surface of the membrane, the membrane was then weighted (M_{wet}). The water calculated based on Eq. 1.

\[
\text{Water uptake} (\%) = \frac{M_{wet} - M_{dry}}{M_{dry}} \times 100\% \tag{1}
\]

where \(M_{wet}\) and \(M_{dry}\) are the weights of the membrane in the wet and dry states, respectively.

**Ion exchange capacity (IEC)**

The membranes were cut to size of 3 cm\(^2\), then soaked in 50 mL of 0.01 M NaOH for 12 h at room temperature. A total of 10 mL of NaOH solution used for soaking the membrane was titrated by 0.01 M HCl with phenolphthalein indicator. In addition, 10 mL of 0.01 M NaOH solution as a blank was also titrated with a solution of HCl 0.01 M. Ion exchange capacity calculated based on Eq. 2.

\[
\text{IEC} = \frac{v_{blanko} - v_{sample}}{m} \times [\text{base}] \times f \tag{2}
\]

where \(m\) is the weight of membrane and \(f\) is a correction factor.

**Proton conductivity**

The membranes were cut to size of 12 cm\(^2\). The proton conductivity of fully hydrated membranes was measured as a function of temperature. Measurements were carried out in two point-probe conductivity cells using two platinum wire electrodes as the working electrode. The electric resistance data were obtained using impedance analyzer (Agilent\textsuperscript{TM} E4980A) and were recorded between 20 Hz–2 MHz at working voltage amplitude of 1 V. The impedance spectra were fitted on ZView 2 software by Scribner Associates Inc. for curve fitting procedure. The proton conductivity values were obtained according to Eq. 3.

\[
\sigma = \frac{\Delta \text{S}}{R \times A \times L} \tag{3}
\]

where \(\Delta \text{S}\) is a gap of electrodes, \(A\) is wide of electrodes, \(L\) is the thickness of membrane and \(R\) is membrane resistance derived from the low intersection of the high-frequency semicircle on a complex impedance plane with Re (2) axis.

**RESULT AND DISCUSSION**

**Fabrication of CSMA, CSMA-CS and CS-MA Membranes**

Though active amino groups and hydroxyl groups are beneficial to grafting reaction, poor water-solubility prevents chitosan from becoming a perfect grafting biomaterial. In general, chitosan is dissolved in acetic acid solution [9-10]. The usage of acetic acid as a solvent may make amino groups of chitosan protonated with no possibility to donate electrons, which makes against side group attachment to amino groups. In this study, both of blending and heterogeneous methods were using inversion phase method with evaporation at room temperature.

The CSMA-CS and CS-MA membranes are obtained from amidation reaction between chitosan, and maleic anhydride in a DMF solvent. The free electrons at the nitrogen atom of amines interact with C carbonyl of maleic anhydride. Consequently, maleic anhydride undergoes ring opening, and C carbonyl of maleic anhydride binds to the amine group as shown in Fig. 1.

**Characterization of CSMA, CSMA-CS and CS-MA Membranes**

The maleic anhydride modified-chitosan was characterized using FTIR, SEM, and TGA. Water uptake, ion exchange capacity, and proton conductivity were measured as well.

**FTIR analysis**

FTIR technique was applied to analyze the structural changes in maleic anhydride-modified chitosan directly. Fig. 1 shows FTIR spectra of maleic anhydride-modified chitosan for CSMA, CSMA-CS, and CS-MA membranes.

The analysis was performed on chitosan-maleic anhydride membranes to determine the success of the process of adding maleic anhydride to the chitosan membrane to form of chitosan-maleic anhydride membranes. The addition of maleic anhydride to the chitosan membrane will generate amide functional group on chitosan membrane so that it will change the pattern of the infrared spectra and will bring up the absorption peaks at wave numbers that identify amide groups in chitosan.

The characteristic band at 1558 and 1311 cm\(^{-1}\) are attributed to N-H and C-N groups (NHCOCH\(_3\), Chitosan's amide group.
Fig 1. The proposed reaction between chitosan and maleic anhydride

Fig 2. FTIR spectra of functional of chitosan with maleic anhydride for (a). (1. CSMA$_1$-CS$_1$, 2. CSMA$_1$-CS$_2$, 3. CSMA$_2$-CS$_3$, 4. CSMA$_3$-CS$_3$, 5. CSMA$_4$) polyblend membranes and (b). (1. CS, 2. CS-MA 1:1, 3. CS-MA 1:2, 4. CS-MA 1:3, and 5. CS-MA 1:4) heterogeneous membranes

amide II and III). These increase in absorptions indicate that the maleic anhydride is reacted with chitosan membrane for both blend and heterogeneous methods. The increasing of maleic anhydride which reacted with chitosan membrane lead to the increasing of amide groups contained in chitosan-maleic membranes. The reaction between chitosan and maleic anhydride is supported by the absorption at the peak of 1600–1560, 1260–1255 and 1430–1420 cm$^{-1}$ which indicated a ring-opening reaction of maleic anhydride. In addition, there is also absorption at around 1475 cm$^{-1}$ which attributed to C=C alkenes. Moreover, the absorption band at 1716 cm$^{-1}$ due to the C=O stretching vibration of ester bonds were detected in spectra (a) and (b) (Fig. 2). The intensity of the peak increased with the increase of maleic anhydride amount [11]. Results from the FTIR
the analysis confirmed that the chitosan-maleic membranes were successfully obtained through the reaction of chitosan with maleic anhydride via ring-opening reaction and also ionic crosslinking.

**Morphology analysis (SEM)**

SEM micrograph was utilized to further characterize physical structures of chitosan, CSMA-CS, and CS-MA membranes. Fig. 3 shows that chitosan exhibited a flaky nature with relatively rough and small surface, which can be attributed to strong interactions between chitosan molecules. The surface of the poly-blend membrane (CSMA-CS) and heterogeneous membrane (CS-MA) are non-porous. In addition, all membranes are homogeneous enough and had no significance for phase dissent.

**TGA studies**

The thermal degradation behavior of membranes was examined following the weight loss as a function of temperature. The TGA curves for CSMA-CS and CS-MA membranes are shown in Fig. 4. The TGA curve of CSMA-CS blend membrane (Fig. 4a) exhibits two endothermic transitions at 120 and 400 °C. The transition at 120 °C is attributed to the decomposition of a water molecule with weight loss about 12%. The temperature ranging between 140–290 °C is attributed to the decomposition of amide bond and maleic groups with weight loss about 4.4%. The loss of weight drastically occur at 300–330 °C and then slowed down at a temperature of 540 °C with a mass remaining about 54% that attributed to the decomposition of chitosan polymer [12-13].

The TGA curve of CS-MA (Fig. 4b) shows three weight loss stage in the different temperature range. The decomposition of water molecules occurs at a range between 64.41–107 °C. The temperature at 216–295 °C is attributed to the decomposition of chitosan and maleic moiety. The decomposition of chitosan polymer also occurs at a range between 300–330 °C then significantly decreased to 55.5% of membrane weight at 540 °C [12-13].

The results of TGA analysis indicate that chitosan reacted with maleic anhydride for both blend and heterogeneous methods. However, the addition of maleic anhydride to the chitosan membrane increasing the thermal resistance of the membrane and reduce the decomposition of chitosan. Chitosan is polymer...
Table 1. Water uptake, Ion Exchange Capacity (IEC) and Proton Conductivity of CSMA-CS and CS-MA membranes

<table>
<thead>
<tr>
<th>No</th>
<th>Sample</th>
<th>Mole ratio</th>
<th>Water uptake (%)</th>
<th>Ion exchange capacity (meq g⁻¹)</th>
<th>Proton conductivity (x 10⁻³ S cm⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>CSMA₁-CS₁</td>
<td>1:1:1</td>
<td>106</td>
<td>4.2</td>
<td>6.2</td>
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<tr>
<td>2</td>
<td>CSMA₁-CS₂</td>
<td>1:1:1.5</td>
<td>150</td>
<td>3.8</td>
<td>1.8</td>
</tr>
<tr>
<td>3</td>
<td>CSMA₁-CS₃</td>
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<td>165</td>
<td>2.1</td>
<td>5.0</td>
</tr>
<tr>
<td>4</td>
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<td>1:1:2.5</td>
<td>155</td>
<td>1.8</td>
<td>3.7</td>
</tr>
<tr>
<td>5</td>
<td>CSMA₂-CS₁</td>
<td>1:2:1</td>
<td>157</td>
<td>6.5</td>
<td>2.9</td>
</tr>
<tr>
<td>6</td>
<td>CSMA₂-CS₂</td>
<td>1:2:1.5</td>
<td>110</td>
<td>2.8</td>
<td>3.5</td>
</tr>
<tr>
<td>7</td>
<td>CSMA₂-CS₃</td>
<td>1:2:2</td>
<td>152</td>
<td>3.6</td>
<td>7.3</td>
</tr>
<tr>
<td>8</td>
<td>CSMA₂-CS₄</td>
<td>1:2:2.5</td>
<td>118</td>
<td>3.7</td>
<td>6.3</td>
</tr>
<tr>
<td>9</td>
<td>CSMA₃-CS₃</td>
<td>1:3:2</td>
<td>102</td>
<td>2.2</td>
<td>6.8</td>
</tr>
<tr>
<td>10</td>
<td>CSMA₄-CS₄</td>
<td>1:4:2.5</td>
<td>166</td>
<td>2.0</td>
<td>3.5</td>
</tr>
<tr>
<td>11</td>
<td>CS-MA₁</td>
<td>1:1</td>
<td>63</td>
<td>0.92</td>
<td>1.0</td>
</tr>
<tr>
<td>12</td>
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<td>1.69</td>
<td>1.1</td>
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<tr>
<td>13</td>
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<td>70</td>
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<td>1.7</td>
</tr>
<tr>
<td>14</td>
<td>CS-MA₄</td>
<td>1:4</td>
<td>70</td>
<td>2.27</td>
<td>3.5</td>
</tr>
</tbody>
</table>

Fig 5. The IEC and proton conductivity curve of CSMA-CS and CS-MA membranes

compound that composed of glucosamine monomer units that need a high temperature for its decomposition.

**Water uptake**

Water uptake has an important influence on membrane performance such as proton transport. Water uptake is used to determine the amount of water absorbed by the membrane. Several factors that determine water absorption capability of a membrane are hydrophilic group number and structure of the polymer chain [14]. The homogenous and heterogeneous poly-blends membrane of chitosan-maleic contains hydrophilic groups such as an amine (N-H), hydroxyl (O-H) and carbonyl (C=O). Basically, the present of maleic anhydride makes the structure of polymer opened. It leads the water molecule to diffuse into the membrane more easily. Water uptake of both membranes is shown in Table 1.

Reducing crystallinity of the membrane causes the hydrophilicity of membrane increase and produce greater water uptake. It affects the performance of the membrane in applications as ion exchange membrane. The higher the water uptake leads to higher in proton conductivity due to much more water molecule in the membrane as proton transfer media [15].

**Ion Exchange Capacity (IEC)**

Ion exchange capacity (IEC) reveals the number of groups randomly distributed in the polymer chain that capable to exchange the protons in solution. Poly-blend membranes are made, the group is expected to support the activity of proton exchange is an amine and hydroxyl groups on the chitosan and the carboxyl group of maleic.

Carboxylate groups are a polar group. Adding the carboxylic group into the polymer chain of chitosan can make chitosan more polar. The increased polarity on chitosan membrane due to increasing the hydrophilic properties of the chitosan membrane. The value of ion exchange capacity increase with the increasing concentration of maleic anhydride that was reacted with chitosan. This relation indicates that the
hydrophilic properties are directly proportional to the ion exchange capacity. This is due to the chitosan-maleic anhydride membrane ionic groups that support the activity of ion exchange are carboxylic groups (-COOH) [16].

The results of the ion exchange capacity of CSMA-CS and CS-MA membranes are shown in Table 1. It can be seen that increasing chitosan and maleic anhydride concentration did not obviously clear affect the IEC for CSMA-CS membranes (blend method). It is assumed that when chitosan reacts with maleic anhydride, the NH₂ group binds to maleic anhydride thereby reducing the active group acting as a proton exchange medium. The increasing of concentrations increase the number of hydrophilic groups but decrease homogeneity of the membrane by increasing the phase difference. It can be seen from the results of SEM where an increase in membrane phase through form aggregates. The existence of these aggregates reduces the capacity of the ion exchange membrane [17]. The aggregates are formed from the interaction of NH₂ groups between the chitosan molecules. Therefore, an increase in the concentration of chitosan reduces the IEC value due to reducing of NH₂ groups which can exchange the proton.

The CS-MA membranes (Fig. 5) show the increasing of the IEC value as increasing the maleic anhydride concentration in the membrane. The highest IEC of CS-MA membrane is obtained for CS-MA₄ with the IEC value of 2.27 m_eq.g⁻¹. Based on the data obtained, the chitosan membrane becomes more hydrophilic because the carboxylic group of the maleic anhydride successfully substituted chitosan into the polymer chain. The presence of the carboxylic group on chitosan membranes can act as ion exchangers. Therefore, chitosan membrane can be applied to an ion exchange membrane.

**Proton conductivity analysis**

The proton conductivity of CSMA-CS and CS-MA membranes was determined by using the complex impedance method. All impedances were carried out after hydration of the membranes. In the fuel cell system, electricity is generated from proton transfer involving functional groups that can redeem proton thus proton conductivity associated with IEC. In addition, the proton conductivity is also associated with the water uptake for proton transport medium. During measurement, the membrane is wet by the water supplied electric current so that the water molecules adsorbed in the membrane will be dissociated into H⁺ and OH⁻ ions. The H⁺ ions are then bound by a proton exchange group such as -NH₂, -OH and COOH for their ion pairs on the N and O atoms, which then migrate and move along the ion exchange groups as a result of the electrical current flowing along the membrane polymer chain. Proton conductivity measurement is the back and forth movement of protons moving in the opposite direction of the original motion [5].

Table 1 and Fig. 5 show that the chitosan modified-membranes have higher proton conductivity values than the chitosan membrane. The results of previous studies have shown that chitosan membrane with a high molecular weight and degree of deacetylation had the proton conductivity of approximately 10⁻⁴ S cm⁻¹. The existence of hydroxyl and amino groups on chitosan framework cause chitosan has high enough hydrophilicity. However, in normal circumstances, chitosan membrane has low electrical conductivity. Chitosan monomer has three atoms of hydrogen in its structure, but the hydrogen atoms are strongly bound to the framework of chitosan and could not be moved under an electric field so that the chitosan film could not be used as proton conductor [6]. However, if the chitosan is dissolved in acetic acid and then were cast as a membrane (thin film), the H⁺ or H₃O⁺ and CH₃COO⁻ ions on chitosan membrane system will spread in the framework of chitosan. These ions can be moved under the influence of an electric field. If the H⁺ or H₃O⁺ ions move more easily than CH₃COO⁻ ion, the chitosan films will be a proton conductor. Chitosan in acidic media can also act as a polyelectrolyte through the protonation of -NH₂ groups. The complex was then able to generate an electrical current in the membrane.

**CONCLUSION**

In conclusion, functionalization of chitosan with maleic anhydride hydride using both heterogeneous and blending method improved the ability of chitosan membrane as proton exchange membrane. It is supported by the results of analysis of ion exchange capacity, water uptake, and proton conductivity. The addition of maleic anhydride also improves the compatibility of the polymer matrix then increase the polymer matrix interactions. The synthesis of functionalized chitosan with maleic anhydride in this present study paves the way for being a promising candidate in the fuel cell membrane.

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