Casting of Polyacrylamide/Poly (vinyl alcohol) Reinforced by Carbon Nano-Wire for Using into Proton Exchange Membrane Fuel Cell

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AMMA irradiations pose very important role for finally cross-linking, poly (vinyl alcohol) (PVA) and acrylamide (AAm) with different ratios. The addition of 0.3% carbon nano-wire into the cast mixture was well mixed prior to being exposure to the irradiation doses (kGy). It was found that; 10 kGy was the optimum dose in order to attain accepted physical properties for the fuel cell’s application. Phosphorylation of the casted membranes was performed to improve their specifications for the desired application. Characterizations of the casted membranes were carried out using Fourier transformer infra-red (FTIR), thermal gravimetric analysis (TGA) and scanning electron microscopy (SEM) for their investigations chemically, thermally and morphologically respectively. Studying the properties of the membranes was performed using ion exchange capacity (IEC), water uptake and tensile strength for testifying their availability into the proton exchange membrane fuel cell (PEMFC). Proton conductivity was measured (maximum value was $8.1 \times 10^{-2}$ S/cm) and free volumes sizes were evaluated using positron annihilation lifetime spectroscopy (PALS). The casted membrane confirmed their capability for using into PEMFCs by measuring the fuel cell performance and durability (up to 500 hours) comparing to compressed Nafion® 1110.

Keywords: Casting, Gamma irradiation, Acrylamide, Poly (vinyl alcohol), Carbon nano-wire, Fuel cell.
the PAAm gel by the formation of inter-polymer complex [19]. PVA has been chemically modified by crosslink co-polymerization of acrylamide (AM) in an aqueous solution of PVA and finally crosslinking the co-polymer to produce a full interpenetrating network (FIPN) membrane [21].

Carbon, a monolayer of sp² hybridized carbon atoms organized in honeycomb lattice, is a semi-metal (zero gap semi-conductor) [22]. Due to their chemical, electrical, optical and mechanical properties, they can be employed in an exceptionally wide application range, including sensing, smart textiles, catalysis, battery technology or composite reinforcement [23-25]. Finite termination of carbon into the membrane results in opening a band gap in carbon nano-wire structures, that is inversely proportional to the width [26].

In this work casting of phosphorylated PAAm and similar for PVA with different ratios and 0.3% carbon nano-wire was carried out. Characterization of the membranes using FTIR, TGA, and SEM. Some parameters were measured such as IEC, water uptake, proton conductivity, membrane thickness and tensile strength. The membrane proved their applicability into the PEMFC and their durability up to 500 hours compared to Nafion 1110.

Experimental

Materials

Acrylamide, carbon nano-wire phosphonic acid and hypophosphorous were kindly provided by Aldrich, England. PVA (Mw. 14000) were provided by Merck (Germany). All chemicals were of analytical grade and used without further purification.

Membrane synthesis

PVA was dissolved in deionized (DI) water by stirring at 90 °C to obtain a 5% (w/w) solution. Then, H₃PO₄: PVA mass ratio 6:1 was prepared by mixing aqueous solutions of 0.5 M H₃PO₄ and 5% PVA. After vigorous stirring of the mixtures at room temperature for 30 minutes, the solutions were cast into molds (polystyrene petri dishes) [27, 28].

AAm was dissolved by bi-distilled water for a ½ hour at 70 °C. It was placed in sealed glass ampoules with 5% diluted phosphonic acid. The phosphonic acid was diluted with co-solvents namely methylene chloride and methanol with a 1:1 volume ratio then pouring them with ratio 10, 20 and 30 % into the petri dishes. Adding 0.3% carbon nano-wire into the cast mixture was performed. The cast components were well mixed using magnetic stirrer for ½ hour. Membrane thickness was adjusted to 50 +/- 2 µm and measured by a digital micrometer (Flower China). Different gamma irradiation (Co-60) doses (5, 10, 15 and 20 kGy) were used to ensure finally crosslinking and complete phosphorylation prior to solvent evaporation (24 hours at room temperature). The prepared membranes were stored in a dry atmosphere after carefully detaching from the petri dishes [29]. Before characterization, the membranes were washed in deionized water to get rid of un-reacted phosphonic acids until neutrality was reached. The prepared membranes were stored in a dry atmosphere [3].

Fig. 1. The suggested structure of phosphorylated PVA [30].

Ion exchange capacity (IEC) and degree of substitution

IEC values of the casted polymers were measured using a typical titration method. The dried membranes in the protonic form were immersed into 25.00 mL of 3 M NaCl solution for 24 h. A large excess of Na⁺ in the solution ensured nearly complete ions have exchanged. Then, 10.00 mL of the 3 M NaCl solution containing the released H⁺ was titrated against 0.05 M NaOH solution using phenolphthalein as indicator. The IEC was calculated using the following equation:

$$I EC_{exp} = \frac{0.05 \times V_{NaOH} \times n}{W_{dry}} \text{ (m equiv./g)} \ldots (1)$$

where $V_{NaOH}$ (mL) is 0.05 M NaOH solution’s volume used for titration. $n$ is the factor corresponding to the ratio of the amount of NaCl solution taken to immerse the polymer (25.00 mL)
to the amount used for titration (10.00 mL), which is 2.5. \( W_{\text{dry}} \) (g) is the dry weight of the polymer electrolyte membrane in the protonic form [31].

**Water uptake and proton conductivity**

The prepared membranes were immersed in DI for different times (hours) intervals till attaining constant weight. The water uptake was calculated as the following:

\[
\text{Water uptake} = \frac{(W_{\text{wet}} - W_{\text{dry}})}{W_{\text{dry}}} \times 100
\]  

where \( W_{\text{wet}} \) and \( W_{\text{dry}} \) are the membrane weights in the wet and dry states, respectively. Proton conductivity was determined by impedance spectroscopy measurement using a LCR meter, Herioki, Japan at 20 Hz to 1 MHz frequency range. The samples were hydrated in water at 25 °C overnight before the measurements. The high-frequency intercept on the real axis of the Nyquist plot was used to calculate the membranes’ proton conductivity. The ionic conductivity (\( \sigma \)) was calculated according to the following equation:

\[
\sigma(\Omega^{-1}\text{cm}^{-1}) = \frac{L}{RA}
\]

where \( L \) is the thickness of the membrane sample, \( A \) or \( (\pi r^2) \) is the sample surface area and \( R \) is the resistance [32].

**Positron annihilation lifetime spectroscopy**

The PALS of the membranes were determined by detecting the prompt Y-ray (1.28 MeV) from the nuclear decay that accompanies the emission of a positron from the \(^{22}\text{Na} \) radioisotope and the subsequent annihilation Y-rays (0.511 MeV). A conventional fast-fast coincidence circuit of PAL spectrometer with a time resolution 240 Ps was used to record all PAL spectra. The PAL spectra containing \( 1.5 \times 10^6 \) counts were analyzed into three components (\( \tau_1, \tau_2, \) and \( \tau_3 \)) and their intensities (I1, I2, and I3) using PALS fit program [33].

**Fuel cell performance**

The (MEAs) were fabricated by hot-pressing the anode and the cathode to the membrane at 200 °C and 1500 psi for 90 seconds. The Pt loading in the anode and the cathode were 0.3 and 0.5 mg cm\(^{-2} \), respectively. The active area of MEAs was \( 5 \times 5 \text{ cm}^2 \). The MEA was placed in a single cell using stainless steel as the end plates and stainless mesh as the current collectors. To check the self-humidifying effect of the Nafton-1110 and cast (PVA and PAAm with 0.3 carbon nano-wire) membrane, the single cell was kept running with dry \( \text{H}_2 \) and \( \text{O}_2 \). Pure hydrogen and oxygen were used as a fuel and an oxidant with a flow of 0.2 L/min. regulated by a mass flow controller (Brooks). The MEAs were evaluated with a commercial fuel cell test system (Compu-cell, GT, Electrochem) running a single cell test. The cell voltages and power densities were recorded at different current densities after a stable cell performance was maintained [34]. The membranes were treated by boiling in 3% \( \text{H}_2\text{O}_2 \) and 0.5 M \( \text{H}_2\text{SO}_4 \) in deionized water to completely remove all impurities before the experiment was carried out [35].

**Equipment**

- The cast membrane (10/90/0.3%) functional groups were studied using a Mattson 1000, Pye-Unicam, England.
- The cast membrane (10/90/0.3%) surface investigation and magnification were carried out by SEM on a JEOL-JSM-5400 (Japan). The magnification was 30 kV X1000.
- The casted membrane’s (10/90/0.3%) thermal stability was analysed with a Shimadzu TGA-50, Japan. The heating rate was 10 °C/ minute in a nitrogen gas atmosphere.

**Results and Discussion**

**Theoretical approach**

This work aims to achieve a cheap membrane with highly electrical conductivity, environmentally friendly and having high stability. Phosphorylating the poly (vinyl alcohol) and similar for acrylamide were carried out. Adding phosphorylated PV A to phosphorylated PAAm with carbon nano-wire and mixing well was performed prior to being irradiated. Gamma irradiation has a potential role for complete phosphorylate the PAAm and PV A and finally cast membranes crosslinking. The carbon nano-wire has a serious role for membrane reinforcement, gasses passage control, and electrical conductivity improvement. This work has been illustrated in Fig. 2.

**Membrane characterization**

The evidence of co-polymerization and network formation by characterizing the casted membrane (PVA / PAAm with 0.3 carbon nanowire).

In the FTIR spectrum shown in Fig. 3, the peak at 3400 cm\(^{-1} \) is due to the O-H stretching.
vibration band. Some broadness in OH, due to intermolecular hydrogen bonding between the hydroxyl groups, participated along with the chains of the co-polymer network structure. Phosphorylates display bands due to the PO stretching vibration mainly in the 900-1200 cm\(^{-1}\) range, especially the (POH) at approximately 925 cm\(^{-1}\) and PO\(_2^\text{2-}\) at approximately 970 cm\(^{-1}\) vibrations [36]. The absorption band at 3433 cm\(^{-1}\) is attributed to the vibration corresponding to the NH groups from the crosslinking, while the asymmetric valence vibration from 2926 cm\(^{-1}\) is assigned to the CH\(_2\) groups included in the macromolecular chains and crosslinking [37]. In addition, increasing the intensity of the stretching vibration band of the C-C group at 1450 cm\(^{-1}\) provides evidence of a crosslinking co-polymerization reaction [38]. Increase phosphorylated PVA at the cast membranes, increase the peaks intensity and express their importance role for the PEMFC applications.

**Surface Morphology study**

Scanning electron microscope of phosphorylated PVA/ phosphorylated PAAm cast membranes with 0.3% carbon nanowire was illustrated in Fig.4.

Figure 4(a) showed obvious differentiated two polymers shared into the cast membranes. The small cube-like structure with bright, whitish
color and bigger matrix with dark grayish color can be distinguished. Figure 4(b) showed less distinguishable polymers’ compartments of the casted membranes. Bigger non-geometrical structural were obviously seen. Large pores and grooves have been irregularly distributed onto the prepared membrane’s surface. Figure 4(c) illustrated nearly non-distinguishable cast membrane’s compartments. The membrane has wavy, smooth surface with small pores and channels. This illustration showed the optimum compatibility of the three kinds of the prepared membranes. These observations recommend it to be successfully applied into PEMFC. There is no dots aggregation onto the polymer matrix which ensured well distribution (homogenous distribution) of carbon nanowire into the casted membranes.

**Thermal gravimetric analysis**

The adding 0.3 carbon nanowire into the 90% PVA and 10% PAAm cast membrane improved their thermal stability as seen in Fig. 5 and Table 1. The thermogram could be divided into 5 divisions.

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**Fig. 4.** Scanning electron microscope of phosphorylated poly (vinyl alcohol)/phosphorylated (poly acrylamide) and 0.3% carbon nano-wire at 10 kGy gamma irradiation dose. a) Phosphorylated PVA: Phosphorylated PAAm (70:30%), b) Phosphorylated PVA: Phosphorylated PAAm (80:20%), c) Phosphorylated PVA: Phosphorylated PAAm (90:10% by weight.

Fig. 5. Thermogram of 90% phosphorylated poly (vinyl alcohol): 10% phosphorylated poly (acrylamide) with 0.3% carbonnano-wire at 10 kGy gamma irradiation dose.

**TABLE 1. Thermogram results characterization.**

<table>
<thead>
<tr>
<th>Thermograms divisions</th>
<th>Temperature range °C</th>
<th>Weight loss %</th>
<th>46% Grafting yield</th>
</tr>
</thead>
<tbody>
<tr>
<td>Part I</td>
<td>(Ambient) - 98</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Part II</td>
<td>98-180</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>Part III</td>
<td>180-360</td>
<td>20</td>
<td></td>
</tr>
<tr>
<td>Part IV</td>
<td>360-460</td>
<td>30</td>
<td></td>
</tr>
<tr>
<td>Part V</td>
<td>460-560</td>
<td>46</td>
<td></td>
</tr>
</tbody>
</table>

The first part described the membrane stability upon raising the temperature (the membrane working temperature) at which, no weight loss have been noticed. Hence, this drying step, namely the evaporation of moisture where the temperature range was under 100 °C [39]. Second part showed increasing the weight loss to 4% by raising the temperature range up to 180 °C. It displayed tiny mass loss, which could be due to the loss of light volatiles [40]. Third part illustrated decrease 20% of the original weight that was described as active pyrolysis region, and most of the mass loss associated with the degradation of organics and in sequence [41]. The fourth part showed 10% loss weight as raising up the temperature to 460 °C. The fifth part illustrated the weight decrease 16% by raising the temperature to 560 °C. Hence, the shoulder was not obvious in the sample, decomposition process described as, like tails. In the final stage, the residues were carbonized into stable and complex organic [42].

Membranes characterizations for using in the fuel cell

Characterizations of the phosphorylated PVA/phosphorylated PAAm with different ratios and 0.3% carbon nano-wire to investigate their availability into PEMFC were performed and tabulated in Table 2.
Some parameters such as water uptake, IEC, proton conductivity, free volume sizes and tensile strength as a function of phosphorylated PVA ratio into the casted membranes (as a predominant factor to attain the optimum compatibility) were studied. Water uptake and IEC increased as phosphorylated PVA ratio increased. It was due to increase the phosphorylation yields, hydroxyl one and the more membrane compatibility. Proton conductivity increased with increasing the phosphorylated PVA ratio while at a high ratio (90%) their value became constant. The significantly high proton conductivities of the casted multilayer membranes were due to high water content retention [28].

PALS is an important method for studying sub-nanometer size holes and for the determination of their size distribution and free-volume fractions [43]. The very important point of interest is the size of the nano-holes, which is strongly correlated with the o-Ps (ortho positron spectroscopy) lifetime. Since it had no information about nano-hole shape, it is assumed that the nano-holes in the prepared membranes were spherical. According to the classical free-volume theory, the segmental motion of polymers is considered to incessantly generate and dissipate free-volume nano-holes [43]. The penetrant molecules in polymers can diffuse when they find neighboring holes large enough to move. It has been recognized that $D$ (molecules diffusion) of penetrants in polymers is affected by $rV$ (gas constant) and other parameters revealed that $D$ in polymers strongly depended on the individual nano-hole size, $V$, measured by PALS. It can be proposed the following empirical relationship between $D$ and $V$ is as follows:

$$D = A rT \exp\left(-\frac{a}{V}\right) \quad (4)$$

where $A$ and $B$ denote the constants related to the types of gasses and polymers [44].

Phosphorylated PVA up to 90% revealed an increase in membranes free volumes. The porous membranes had strong ability to enhance $\text{H}^+$ diffusion and so increasing the ion exchange capacity, water uptake, and proton conductivity. Increasing pores numbers and their volumes onto the membranes enhanced $\text{H}^+$ exchanging capacity and so increasing proton conductivity. The uniformly dispersed molecules, (especially for 90% phosphorylated PVA casted membranes) could form bridges among functional groups (a hydroxyl, $–\text{PO}_2$ and an amine) clusters and shorten the distance for proton hopping under high humidity [28]. It can be strengthened by adding carbon nano-wire. The incorporation of carbon nano-particles into the prepared membranes materials contributed to a better filler interaction due to the fact that it facilitated the gap between the membranes components structures. It leads to an infinite conductive path development that aided in increasing electrical conductivity and better formation of the conducting network [45-47].

The conductivity via membranes is depending mainly on an ion exchanging process. The electrons motion is not the main process as the carbon nanowire low concentration which seemed as isolated islands.

For these reasons, phosphorylated PVA and carbon nanowire contained into the casted membranes raised the proton conductivity values and confirmed their utility for the fuel cell application. The casted membranes (90% phosphorylated PVA/10% phosphorylated PAAm) tensile strength showed more value compared to Nafion-1110. It was found that; the

<table>
<thead>
<tr>
<th>No.</th>
<th>Kinds of membranes</th>
<th>Thickness (mm)</th>
<th>Water uptake (%)</th>
<th>Ion Exchange Capacity (mmolg$^{-1}$)</th>
<th>Proton conductivity $\times 10^2$ (Scm$^{-1}$)</th>
<th>Free volumes sizeÅ</th>
<th>Tensile at break Strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-</td>
<td>70% PVA casted membrane</td>
<td>50</td>
<td>43</td>
<td>1.17</td>
<td>7.9</td>
<td>9.053</td>
<td>21</td>
</tr>
<tr>
<td>2-</td>
<td>80% PVA casted membranes</td>
<td>50</td>
<td>58</td>
<td>1.27</td>
<td>8.7</td>
<td>9.471</td>
<td>28</td>
</tr>
<tr>
<td>3-</td>
<td>90% PVA casted membrane</td>
<td>50</td>
<td>66</td>
<td>1.46</td>
<td>9.6</td>
<td>9.094</td>
<td>39</td>
</tr>
<tr>
<td>5-</td>
<td>Nafion-1110</td>
<td>52</td>
<td>35</td>
<td>0.91</td>
<td>6.1</td>
<td>8.371</td>
<td>35</td>
</tr>
</tbody>
</table>

TABLE 2. Casted membranes characterization compared to Nafion-1110.
tensile strength at less than 90% phosphorylated PVA cast membranes decreased which may be due to less compatibility of the polymers shared into the casted membranes. These results appreciated using 90% phosphorylated PVA cast membranes into the PEMFC.

Fuel cell performance

The PEMFC performance was carried out using Nafion 1110 and 90:10% phosphorylated PVA/phosphorylated AAm with 0.3% carbon nano-wire cast membrane.

Figure 6 & 7 showed the polarization and performance curves for the PEMFC at 100% RH and 80 °C with H2 as a fuel and O2 as an oxidant under ambient pressure. From the data, it can be seen that prepared membrane showed better performance than Nafion 1110. Higher open-circuit voltages (OCVs) observed for the understudied membranes indicating that there are no problems related to gas permeability and no significant electronic conductivity which would otherwise adversely affected the OCV. The trends observed in polarization plot were in akin to the proton conductivity and IEC measurements for the membranes. In the prepared membranes, the proton conductivity is account for the transfer of protons through hydrogen bonding with water-filled ion pores. The high density of –PO2 groups of the cast membrane act as a solid acid proton conducting medium which provides potential proton conducting sites, and facilitates achieving higher proton conductivity [48]. However, for Nafion 1110 membrane, lower in proton conduction and performance is observed. A large fuel cell performance at high current density region is clearly seen for the prepared membrane and Nafion1110. From the foregoing, it is obvious that the understudied membranes’ performance is closely related to the issue of water management. The limited availability of water at the anode may be due to an electro-osmotic drag of water from the anode to the cathode and insufficient water back-diffusion from the cathode to the anode cause the MEA to dehydrate. Membrane dehydration resulted in an increase in the ohmic resistance of the cell, leading to decreasing in cell performance. By contrast, if the membranes hold more generated water and help the membranes sufficiently wet which maintains the proton conductivity value and enhances the power density. At high load current densities, water generation rates at the cathode are high enough to substantially affect the effective area of the gas-diffusion layers inside the electrode and cause flooding in the understudied membranes. However, high water uptake nature especially of the prepared membrane resulted in water is back diffused towards the anode electrode and maintains reasonable hydration level across the membrane and mitigates the cathode flooding [49].

The highest open circuit voltages were 0.90 V for Nafion 1110 and cast membrane at 80 °C. The carbon nano-wire favor the mass transport (minimize gas crossover through the plane of the membrane) and reduce power losses when using the casted membranes. These facts increase effectively the membrane durability. At low current densities (up to 100 mA cm-2), when the fuel cell is a kinetic controlled both membranes showed nearly the same power densities, at a higher current densities the performance of the casted membrane became higher than the Nafion 1110 membrane [50].

Fig. 6. Cell performance (current densities against voltage) of 90% phosphorylated poly (vinyl alcohol)/phosphorylated poly (acrylamide) with 0.3% carbon nanowire casted membrane and Nafion-1110 at 80 °C temperature and 100% RH.
Fuel cell durability

The durability of the casted membrane (10 % phosphorylated PAAm with phosphorylated PVA and 0.3% carbon nanowire) and Nafion-1110 was performed and illustrated in Fig. 8.

The results showed that the 10% phosphorylated PAAm cast membranes slowed their rate of degradation compared to the compressed Nafion 1110 used. The polymer membrane degradation is likely to be the one mostly...
responsible for the failure of the PEMFC [51]. Mechanical and thermal membrane degradations could be mentioned [52], however they have a little effects on the fuel cell performances evolution during operation, except at the very end of the cell lifetime when its permeation current increases dramatically. It is a very sudden phenomenon which could be correlated also with the MEA initial fabrication defects and the bipolar plate mechanical constraints. Another critical degradation mechanism in PEMFC concerns is the catalyst layers and especially the active surface area loss during fuel cell operation. For instance, the irreversible losses can be due to carbon support corrosion, catalyst dissolution / re-deposition and catalyst layer micro-porous structure modification [53]. Conversely to the membrane degradation, these mechanisms affect directly the fuel cell performances with time but not directly a fuel cell failure. Degradation of Gas Diffusion Layers (GDL) and bipolar plates can also be observed but they seem to be a less.

A possible degradation mechanism was that; reactive species formed at the electrodes diffused into the polymer membrane and resulted in polymer chemical degradation. The OCV decay rates observed in this study are high, such values would be expected as membrane degradation due to the radical formation is enhanced under OCV operation and low relative humidity [54]. In the OCV durability test, the results are clear indications of the understudied membranes degradation decreased, which is most likely due to reduced release of harmful radicals formed at the anode. These radicals are believed to be captured by the active groups at the understudied membranes and become eliminated before they can attack the casted membrane material and cause the thinning and pinhole formation that are responsible for the increasing membrane permeability, and thereby gas crossover and OCV decay [55].

The reason for the fuel cell performance stability being improved with the casted membranes could be due to the final crosslinking restricting radicals’ diffusion into the membranes matrix [28].

**Acknowledgment**

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**Conclusion**

This work aims to synthesize low cost and efficient polymer membranes used in proton exchange membrane fuel cell (PEMFC). Acrylamide phosphorylation was performed and mixed with phosphorylated PVA to prepare the cast composite. 0.3 carbon nanowire was inserted into the cast mixture for membrane reinforcement, gas passage control and has a role for the excess of hydrogen proton sorption. The cast mixture was irradiated with different doses (10 and 20 K Gy) for finally crosslinking and ensure complete phosphorylation. Characterization of the membranes using FTIR, TGA, and SEM recommend their usage in PEMFC application. Measurements of some impacts such as water uptake, $IEC$, and hardness were carried out. Proton conductivity and porosity (evaluated by $PALS$) investigations confirmed the membranes utility into the fuel cell. The casted membrane at 90% phosphorylated PVA has high performance and durability compared to compressed Naftion1110 which recommend it to be successfully applied at the PEMFC.

**References**

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استخدام طريقة الصب للأكريلاميد/بولي فينيل الكحول و المدعم بكربون نانووير و ذلك لإستعماله في خلايا طاقة البروتون المستبدل

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تمثل اشعة جاما أهمية كبرى لتمام التشابك لكل من البولي فينيل الكحول و الاكريلاميد و ذلك بنسبة مختلفة و قد تم إضافة 3% كربون نانووير للخلوتيو و ذلك قبل التشعيع الجامعي و قد كان 200 كيلو جرام الكهف المائي للحصول على خواص فيزيائية مناسبة لتطبيق في خلايا الطاقة وقد تم تصنيف الأغشية باستخدام تقنيات الأشعة تحت الحمراء و التحليل الوزني الحراري و ميكروسكونتوب السمج الحراري و ذلك لفحص الأغشية كيميائيا و حراريا و مورفولوجيا على الترتيب كما تم دراسة سعة الاستبدال الأولي و التشرب بالماء و اختبار قوة الشد و سمكية الأغشية لكلاً من الأغشية في خلايا الطاقة وقد وصلت التوصيلية للبروتون الى 8.1$\times 10^{-2}$ S/cm كما تم قياس التفاعلية بتقنية كفاءة البروتون و قد اثبتت الأغشية قدرتها بالعمل في خلايا الطاقة لمدة تصل الى 500 ساعة و بدرجة مقارنة للغشاء التجاري نافيون 1110.


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