Permeation dynamics of dimethyl methylphosphonate through polyelectrolyte composite membranes by in-situ Raman spectroscopy

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1. Introduction

Protective materials against chemical warfare agents (CWAs), such as nuclear, biological and chemical (NBC) protective clothing, have been evolving from impermeable butyl rubber-based materials to permeable polyurethane foams mixed with activated carbon [1]. One of the challenges in designing protective wearable gear is the ability of the material to quickly permeate sweat and dissipate heat while blocking penetration of CWAs through the protective material. Earlier generations of NBC protective gear such as butyl rubber-based gloves, masks and boots, are excellent at blocking hazardous chemicals, since they are impermeable, but are not able to dissipate heat or sweat from the body causing serious heat strain [2]. As protective clothing technology evolved through time more protective features were included within textiles of military uniforms. An example of such technological development for chemical-biological (CB) protection includes the Joint Service Lightweight integrated Suit Technology (JSLIST), a multilayered air-permeable adsorptive textile, as it provides chemical protection. In addition, the properties of PEMs can be tailored by loading functional metal-oxide nanoparticles (MONP) using the in-situ growth mechanism [8]. Growing MONP within PEM substrates allows for easy scalability for mass production as membranes go through very simple steps in order to incorporate inorganic catalysts. The incorporation of MONP within PEM substrates provide enhanced chemical stability [9], along, as shown below, with the ability to hinder the permeation of harmful substances, thus providing chemical protection. In addition, the proposed alternative protective materials could have the potential to have self-decontaminating properties as CWAs could be decomposed by the

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ABSTRACT

Reliable measurement of permeability of membranes with respect of different chemical compounds is of paramount importance for the design of novel materials for selective separations, protective barriers against chemical warfare agents and toxic industrial chemicals. An original in-situ Raman spectroscopy experimental setup is devised to measure the dynamics of permeation of toxins with example of dimethyl methylphosphonate (DMMP), a nerve agent surrogate, across polyelectrolyte composite membranes. Efficiency of the proposed method is demonstrated on two types of commercial membranes, Nafion 117 and Nexar MD9200, modified with metal oxide nanoparticles. It was found that loading membranes with ZnO nanoparticles significantly reduces agent permeation, enhancing its protective capabilities against hazardous substances. The proposed methodology can be adopted and applied for characterization of permeability of other types of CWAs, their simulants, and other chemicals through polymeric membranes of different origin.

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nano-crystalline metal-oxides opening an opportunity towards recyclable garments.

In this study, we report a novel experimental setup to measure via in-situ Raman spectroscopy the membrane protective capabilities against harmful substances. As a practical example, we study the permeation of DMMP, a chemical warfare agent surrogate (CWS) for nerve agents, through composite MONP-PEM membranes. In-situ and in-operando Raman spectroscopy has become a powerful technique to study dynamic experiments, particularly those that require high sensitivity towards chemical and physical changes, such as changes in restructuring of metal-oxides during a chemical reaction [10,11], observing electrochemical changes in metal-oxides for charge storage applications [12], and monitoring reaction products during biomass conversion [13], as well as electrocatalytic water splitting reactions [14]. Moreover, in-situ Raman spectroscopy has been useful for monitoring liquid-phase kinetic phenomena such as functionalization of organic compounds for the production of urea-derivatives [15], measuring concentrations of CO2 within water at high pressures [16], and real-time measurement of aqueous corrosion of borosilicate glass [17]. In-situ Raman spectroscopy measurements for permeation through membrane barriers have also been utilized for electrolytes [18,19].

Two sets of PEMs were used to assess their capabilities as protective materials, Nafion and Nexar. Nafion membranes, produced by DuPont, are part of a larger family of perfluorosulfonated ionomers (PFIS) that have been used in several applications such as proton exchange membrane fuel cells (PEMFCs) [20–22], super-acid catalysis [23,24], selective drying and humidification of gases [25,26], and chlor-alkali electrolysis [27,28]. Nafion membranes consist of a sulfonated tetrafluoroethylene (PTFE)-backbone that incorporate grafted perfluorovinyl ether groups terminated with sulfonate groups onto its PTFE backbone. Nafion shows great chemical stability to a vast amount of chemicals including very corrosive compounds such as hydrogen fluoride and hydrogen chloride gases. In addition, they present excellent mechanical and thermal stability thus making them ideal candidates for high temperature reactions in fuel cells [29,30] and particularly advantageous for chemical protection. Nafion 117 has an IEC of 0.91 meq/g, which yields in average polymer networks with 1100 equivalent weight (EW).

NEXAR® MD9200 membranes, produced by Kraton, are sulfonated penta-block copolymers comprised of styrene and ethylene-co-propylene blocks, where the central styrene block is sulfonated. They have been used in several applications such as alcohol dehydration [31,32], water purification [33], separation of solvents [34], nanofiltration [35], fuel cells [36], and for chemical protection [37]. These pentablock copolymers have been known to possess high water-vapor transmission rates, excellent mechanical and thermal stability [38] making them ideal for high performance clothing. Nexar has an ion exchange capacity (IEC) of 2.0 meq/g, which yields in average polymer networks with 500 equivalent weight (EW). The in-situ growth of MONP within PEM was performed as reported in our previous publication [8].

2. Experimental

2.1. Materials

Nafion-117 ionomer membranes with an equivalent weight of 1100 g/SO3H were purchased from Ion Power, Inc. NEXAR® MD9200 solutions (11 wt% in Cyclohexane and Heptane) were obtained through Kraton. Salt reagents zinc nitrate hexahydrate (98%), and sodium hydroxide (> 97%) were purchased from Sigma-Aldrich.

2.2. Solvent casting of Nexar membranes

Nexar films were solvent-casted from NEXAR® MD9200 solutions (11 wt% in Cyclohexane and Heptane) using a mixture of 100 mL of tetrahydrofuran (THF) and 30 mL of NEXAR® MD9200 solution. The polymer solution was thoroughly mixed for 6 h at room conditions. Afterwards, 25 mL of the solution was poured into 50 mL Teflon evaporation dishes and the dishes were covered with P2 filter paper (to help slow down the evaporation rate of solvent) and were left overnight at room conditions. After 3 days, Nexar films were slightly brown and about 500 μm in thickness and were qualitatively robust and flexible.

2.3. In-situ growth of ZnO within poly electrolyte membranes

Nexar and Nafion membranes were cut into 0.5 inch by 0.5 inch pieces and were immersed in HCl (1 M) at room conditions for 1 h to remove any impurities and then immersed in deionized (DI) water to remove any excess HCl from the previous step. Once washed, these membranes were labeled as “PEM-H+” where PEM represent either Nafion or Nexar. PEM-H+ membranes were immersed in 200 mL of 0.05 M Zn(NO3)2 solutions at room conditions for 1 h to allow for Zn2+ exchange and these samples were labeled “PEM-Zn2+”. Afterwards, PEM-Zn2+ membranes were rinsed in DI water and immersed in 200 mL of 0.5 M NaOH at 60 °C for 1. This step allows for Zn2+ to hydrolyze and form Zn(OH)2. Membranes were rinsed again in DI water and blotted dry before putting the membranes in an oven at 100 °C for 1 h. These samples were labeled “PEM-ZnO”.

2.4. Physical characterization

X-Ray Diffraction (XRD). X-ray diffractograms were obtained using a Phillips XPert diffractometer (Bragg-Brentano geometry) with a CuKα anode (1.5405 Å). The instrument was operated at 45 kV and 40 mA in a continuous scanning mode at a 0.02°/step acquisition rate with a dwell time of 2 s/step from 20 angles over the range of 10°–90° on a 20 scale using a 0.3 mm fixed receiving slit. Transmission Electron Microscopy (TEM). Transmission electron micrographs were obtained by a JEOL 1200EX electron microscope with an accelerating voltage of 80 kV and 2-s sample exposure time. Samples were supported on lacy carbon type-A 300 mesh copper grids.

2.5. Spectroscopic characterization

Raman spectra were acquired by using a Horiba LabRAM HR Evolution spectrometer with spectral resolution. The incident beam, which was a laser of 532 nm, and 80 mW initial power, was directed via a 90° macro lens of 40 mm focal length in the middle of the 2 mm vial that used to measure DMMP diffusion. The scattered light was collected through backscattering geometry and analyzed with an 1800gr/mm grating. Collection of the scattered light was achieved with an air-cooled open electrode 1024x256 pixels CCD at the temperature of −75 °C. The acquisition time of each spectrum was 60–120s, depending on the signal to noise (S/N) ratio and the number of accumulations varied in the range of 6–12. The spectral slit was fixed at 100 μm.

3. Results and discussion

3.1. Characterization of MONP-PEM composites

The samples were examined by x-ray diffraction (XRD) to confirm the presence of ZnO within the membranes. In Fig. 1(a), one can observe that Nafion and Nexar do have the characteristic peaks of hexagonal lattice ZnO as they match the supersposed peaks of reference bulk ZnO. Once the presence of ZnO was confirmed by XRD, we apply transmission electron microscopy techniques in an effort to image and quantify the size of the ZnO nanoclusters within the Nexar substrates. Fig. 1(b,d) presents the ZnO nano-aggregates within PEM substrates. The ZnO nano-aggregate sizes range from 8 to 35 nm in Nafion-ZnO samples, Fig. 1(c). In the Nexar-ZnO samples, larger nanoggregates with a wider distribution are observed with ZnO aggregate size
distributions that range from 10 to 120 nm, Fig. 1(e).
Although both PEM, Nafion and Nexar, were subjected to the same conditions for the in-situ growth of ZnO, there is a significant difference between the ZnO nanocrystallite size distributions. The polymers microstructure plays an important role during the nucleation and growth of ZnO nanocrystals. These membranes nano-segregate in such a way that the hydrophilic subphases, containing the sulfonate ions, coalesce and form interconnected channels that allow for water and ion transport upon hydration [39]. Since the nano-crystalline ZnO grow within the hydrophilic subphases of the membranes, the growth of such crystals may be limited by the confinement of the hydrophilic domains and the limited supply of Zn$^{2+}$ ions available, which allow for nano-sized crystals to grow without a capping agent. It has been demonstrated that by using organic solvents, growth of ZnO within membranes can be manipulated to favor the growth of particular crystalline planes within ZnO. It should be noted, since the studied Na case, the growth of ZnO nanoparticles decreases significantly with the ZnO incorporated membranes. It is evident that the size of the nanoparticles is larger than the hydrophilic subphases of the membranes. The growth of such nanoparticles, Nafion-H$^+$ and Nexar-H$^+$, and is the peak intensity is significantly less with the ZnO incorporated membranes. It is evident that the incorporation of nanoparticles does have a great influence over the permeation rate of DMMP across the PEM, as it is seen on both Nafion and Nexar cases. It is possible that the addition of ZnO–DMMP may hinder DMMP diffusion as ZnO serves as attractive centers from which DMMP may be strongly adsorbed to, as it has been reported previously, DMMP strongly adsors to metal-oxide surfaces [41]. The reduced Signal/noise ratio in some of our measurements is due to the combination of the low concentration as well as fast acquisition time. After performing a background subtraction and peak fitting for each spectra the AUC is calculated and later concentrations of permeated DMMP as a function of time are extracted from the calibration curve in Fig. 3(b).

When plotting the permeated DMMP concentration on the receiving compartment as a function of time, as in Fig. 6(a), it can be seen that Nafion-H$^+$ has an observable breakthrough time, $t_{b,na}$, defined as the time of the first observation of the non-zero concentration of DMMP in the receiving compartment, of 15 min when compared to Nafion-ZnO, which has a breakthrough time of 135 min, an order of magnitude longer. A similar increase of the breakthrough time upon addition of ZnO is found for Nexar samples, 45 and 145 min for Nexar-H$^+$ and Nexar-ZnO, respectively. In addition, the permeation rate of Nexar-H$^+$ is approximately 1.8, 2.3, and 3.8 times faster than Nafion-H$^+$, Nexar-ZnO, and Nafion-ZnO, respectively. These results show that the incorporation of ZnO nanoparticles has a large impact on both, breakthrough time and permeation rate in Nafion and Nexar samples, which demonstrates the enhancement in the protective capabilities that PEM-ZnO have against DMMP.

Once the calibration curve for DMMP in water was obtained, a series of spectra were collected for all permeation experiments against the different membranes studied, see Fig. 4.

Full spectra from Fig. 4 was truncated within the wavenumber range of 615 and 825 cm$^{-1}$ to clearly observe the dynamics of the evolution of the DMMP peak located at 715 cm$^{-1}$ as a function of time as shown in Fig. 5. As it can be observed, the intensity of the peak at 715 cm$^{-1}$ after 1 h of permeation are most intense in the membranes without nanoparticles, Nafion-H$^+$ and Nexar-H$^+$, and is the peak intensity is significantly less with the ZnO incorporated membranes. It is evident that the incorporation of nanoparticles does have a great influence over the permeation rate of DMMP across the PEM, as it is seen on both Nafion and Nexar cases. It is possible that the additional ZnO–DMMP may hinder DMMP diffusion as ZnO serves as attractive centers from which DMMP may be strongly adsorbed to, as it has been reported previously, DMMP strongly adsors to metal-oxide surfaces [41]. The reduced Signal/noise ratio in some of our measurements is due to the combination of the low concentration as well as fast acquisition time. After performing a background subtraction and peak fitting for each spectra the AUC is calculated and later concentrations of permeated DMMP as a function of time are extracted from the calibration curve in Fig. 3(b).

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Fig. 1. (a) Wide angle x-ray diffraction spectra for Nafion-ZnO (black), Nafion-H+ (green), and hexagonal-lattice bulk ZnO reference (blue). (b) TEM micrographs of ZnO in Nafion, (c) ZnO aggregate size distribution in Nafion, (d) TEM micrographs of ZnO in Nexar, (e) ZnO aggregate size distribution in Nexar. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)
equilibrium).

Data shown in Fig. 6(a) could be fit to by using Equation (1) and transport properties could be extracted. To compare membrane permeabilities across multiple samples, one has to account for the difference in the membrane thickness, approximately 190 µm for Nafton and 500 µm for Nexar. Rearranging Equation (1) into the following form below in Equation (2),

$$\Delta C_B(t)V_BL = P(t - t_{0,\text{obs}})$$

provides a direct comparison between the sample permeabilities. Equation (2) presents the data shifted to the point, $t_{0,\text{obs}}$, of the first observation of a non-zero DMMP concentration in the receiving compartment and $\Delta C_B(t) = C_B(t) - C_B(t_{0,\text{obs}})$. The observed breakthrough times, $t_{0,\text{obs}}$, were obtained from Fig. 6(a). The permeation data in the shifted coordinates of Equation (2) is shown in Fig. 6(b). The slopes of the linear fits of permeation curves in Fig. 6(b) give the permeability constants, $P$. Linear fits of permeation curves in Fig. 6(b) had excellent agreement with an $R^2$ of 0.99, for the exception of Nexar-H+, which had an $R^2$ of 0.98.

Obtained transport parameters are given in Table 1. No reaction products were observed for DMMP within the time-frame and environmental conditions studied as DMMP is more stable than sarin given that it has a methyl group rather than the much more reactive fluorine that is present in sarin. However, the possibility of very small quantities of byproducts that fall below the detection limit of our Raman measurements cannot be excluded.

4. Conclusions

A novel in-situ Raman spectroscopy setup is built to measure agent permeation across protective barriers and applied to study DMMP permeation through ZnO loaded Nafton and Nexar polyelectrolyte membranes. It is shown that, incorporation of ZnO nanoparticles increases the breakthrough time and reduces the agent permeability, providing better protective capabilities of composite membranes. Of all samples studied, Nafton-ZnO seems to have the best protective capabilities against DMMP permeation. The proposed technique, which monitors the change of concentration in situ, as the permeation proceeds, has significant advantages compared to the existing methods for testing perm-selective membranes that are based on the measurements of permeants in the vapor phase that are transported to a detector by a flow of carrier gas. In-situ measurements provide high accuracy and require much shorter time due to a small sample cell. The proposed methodology can be adopted and applied for characterization of permeability of other types of CWAs, their simulants, and other chemicals through polymeric membranes of different origin.

Conflicts of interest

There are no conflicts to declare.

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Table 1

<table>
<thead>
<tr>
<th>SPF</th>
<th>DMMP Conc. (mol/L)</th>
<th>t_{obs} (min)</th>
</tr>
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<tr>
<td>Nafton-20</td>
<td>17.14</td>
<td>30</td>
</tr>
<tr>
<td>Nafton-30</td>
<td>4.72</td>
<td>135</td>
</tr>
<tr>
<td>Nafton-H+</td>
<td>14.20</td>
<td>45</td>
</tr>
<tr>
<td>Nafton-ZnO</td>
<td>6.27</td>
<td>45</td>
</tr>
</tbody>
</table>

*Measurements have an uncertainty of ± 15 min.

Willis and Blackwell from Kraton for providing the Nexar MD9200 samples.

References


