Preparation and characterization of an alkaline anion exchange membrane from chlorinated poly(propylene) aminated with branched poly(ethyleneimine)

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Preparation and characterization of an alkaline anion exchange membrane from chlorinated poly(propylene) aminated with branched poly(ethyleneimine)

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1. Introduction

Anion exchange polymers have applications as ion exchange resins and as membranes for water purification [1], Li−air batteries, and in polymer exchange membrane (PEM) fuel cells [2]. PEM Fuel cells show promise as alternatives to current energy conversion devices in transportation and stationary applications as well as for use in portable devices. There is currently a large amount of interest in using anion exchange membranes (AEMs) as the electrolyte in alkali fuel cells, because of the increased power density obtained with a solid electrolyte. AEM fuel cells (AEMFCs) have advantages over proton exchange membrane fuel cells, with the potential for direct use of methanol or more complex fuels, and the potential use of non-precious metal catalysts [2]. Commercial AEMs are available from companies such as Tokuyama [3], Solvay Plastics [4], Dupont, and Fumatech [5], although many of these materials are optimized for non-fuel cell applications such as dialysis. Commercial membranes with acceptable mechanical properties, long-term alkaline stability and higher anionic conductivity at lower RH for fuel cell applications are still needed [6,7].

In base oxidative attack of the polymer by hydroxide is less important than degradation of the cationic functionalities [2]. It is, therefore, possible to design polymer systems with methylene chain backbones. A simple approach would be to simply react an aminated polymer with a halogenated polymer to form amiated materials that could then be readily quaternized to form a potentially inexpensive AEM. In previous work, Hong et al. have synthesized similar materials by amination of chlorinated polypropylene (CPP) with ethylenediamine (EDA) [8] and amination of PPC with low molecular weight linear PEI [9,10]. It was shown that these materials exhibit good chloride ionic conductivities in water after soaking in NaCl with a maximum of 0.008 and 0.013 S cm−1 for the quaternized EDA/PPC and the quaternized PEI/PPC material respectively. To extend this group of materials, we selected a branched high molecular weight polyethyleneimine (PEI) as a starting material due to the high concentration of quaternizable amine groups throughout the polymer. Polyethyleneimine (PEI) has most commonly been investigated in biological applications where it is often crosslinked and quaternized to form stable cationic polymers. Resulting products have been shown to be

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effective in gene delivery systems and for use as antimicrobial metal surface coatings [11,12]. Its quaternized form has anti-bacterial properties [13,14].

One issue of hydroxide exchange membranes is that hydroxide reacts rapidly with the ambient CO₂ in the air, forming a mixture of carbonate and bicarbonate counter-ions [15]. The presence of bi(carbonate) ions in place of hydroxide ions is detrimental to the conductivity in the membrane due to inherently lower ion mobility. A reaction of a CO₂ molecule with a hydroxide anion in solution can produce a bicarbonate ion as shown in Eq. (1) and it may also produce a carbonate ion in the presence of two hydroxide ions as shown in Eq. (2). When CO₂ reacts with hydroxide ions located as counter anions in a polymeric membrane, the ratio of bicarbonates and carbonates can be determined by titration or FT-IR.

\[ \text{CO}_2 + 2\text{OH}^- \rightarrow \text{CO}_3^{2-} + \text{H}_2\text{O} \]  
(1)

\[ \text{CO}_2 + \text{OH}^- \rightarrow \text{HCO}_3^- \]  
(2)

Suzuki et al. have observed these reactions of hydroxide with atmospheric CO₂ within the first minutes of exposure of hydroxide form Tokuyama AEMs to air [16].

Although our synthesized membrane had only a moderate IEC of 0.74(0.02) mequiv. mol⁻¹, unique water uptake and transport were observed through Dynamic Vapor Sorption (DVS) and pulsed-field gradient nuclear magnetic resonance (PFG-NMR) spectroscopy. This paper discusses the transport of anions in the context of beginning to understand the morphology of the material. Quantitative analysis of the formation of carbonates when hydroxide counter-ions in the membrane react with atmospheric CO₂ is included.

2. Experimental

2.1. Materials

Branched polyethyleneimine (PEI) (solution (Mₖd ~ 2000 by LS, 50 wt% water), chlorinated polypropylene (CPP) (Mₖd ~ 100,000), trimethylamine (≥99.0%) and iodoethane (99%) were used as received (Sigma–Aldrich). All solvents were reagent grade.

2.2. Synthesis

In preliminary studies, branched PEI solution (50 wt% water) was quaternized with a 10-fold excess of iodoethane, under flowing nitrogen at 50 °C using a condenser to minimize evaporation. The final solution was dried in vacuo to produce a highly hydroscopic white powder.

2.2.1. Quaternized PEI-polypropylene (PP-PEI)

The synthetic procedure is outlined in Fig. 1. CPP (5 g) was dissolved in toluene (50 ml) at 60 °C and added to PEI solution (10 g, 50 wt% water) at 60 °C. The amination reaction was performed at 60 °C for 48 h. The hydrochloric acid (HCl) produced during the reaction was neutralized with excess trimethylamine over 24 h. Five fold excess, based on the amine groups in the starting PEI, iodoethane was added to quaternize all the amine groups on the polymer at 60 °C for 48 h. The remaining solvent and excess trimethylamine reactants were then removed by evaporation, and the resulting polymer was washed with dionized (DI) water to extract any unreacted PEI. The resulting material was soaked in toluene, swelling considerably, and spread onto a Teflon casting block to form a membrane. The toluene was removed in vacuo at 40 °C. The resulting membrane was soaked in a 1 M NaCl solution, and again washed with DI water to guarantee that the sample was in a chloride counter-ion form for further characterization. The final
product will have quaternary ammonium groups with an unknown mix of ethyl groups and poly(propylene) chains attached to each nitrogen.

2.3. Characterization

2.3.1. Spectroscopy

FTIR was performed using a Thermo-Nicolet 2100 FTIR spectrometer in attenuated total reflectance mode. $^{1}$H, and $^{13}$C magic angle spin (MAS) NMR were performed on a Bruker Avance III 400 Spectrometer with a 50 kHz B1 field. A 4 mm MAS BB/BB probe was used. $^{1}$H MAS NMR was taken at 400 MHz, with spin rates of 12 kHz with a 6 s recycle delay. $^{13}$C MAS NMR was recorded at 100 MHz, with a spin rate 8 kHz, a 6 s recycle delay and a 1 ms contact time.

2.3.2. Ion exchange capacity

Ion exchange capacity (IEC) was measured using the Mohr titration method to determine the chloride concentration in a given sample of membrane. The sample was first dried in a vacuum oven at 40 °C overnight and weighed to determine a starting dry mass. Three samples were tested.

2.3.3. Morphology

Small and wide angle X-ray scattering (SAXS and WAXS) experiments were performed at the Advanced Photon Source at Argonne National Lab on beamline 12 ID-B. A Platus 2M SAXS detector was used to collect scattering data with an exposure time of 1 second. The X-ray beam had a wavelength of 1 Å and power of 12 keV. The intensity (I) is a radial integration of the 2D scattering pattern with respect to the scattering vector (q).

Temperature and humidity were controlled during SAXS experiments within a custom built sample oven [17]. Typical experiments studied three membrane samples and one empty window so a background spectrum of the scattering through just the Kaption™ windows and nitrogen environment could be obtained for each experimental condition. The humidity of the sample environment was controlled by mixing heated streams of saturated and dry nitrogen and measured independently with humidity probe (Vaisala). Sample holders were inserted into an oven environment of 60 °C and 95% relative humidity and allowed to equilibrate for 15 min. The humidity was stepped to 75% RH, 50% RH, 25% RH, dry and back to 95%RH with 17 min steps. Spectra were taken during the last 2 min of each humidity condition.

2.3.4. Imaging

Environmental electron scanning microscope (ESEM) images were acquired with an FEI Quanta 600i ESEM and IR microscopy was acquired using a Thermo IN-10 FTIR microscope.

2.3.5. Diffusion coefficient experiment

The self-diffusion coefficient of water through the membrane was measured by pulsed-field gradient nuclear magnetic resonance (PFG-NMR) spectroscopy with a stimulated echo pulse sequence. Pulse gradient NMR experiments are analyzed using the Stejskal-Tanner-Equation given in Eq. (3).

\[
\frac{S}{S_0} = e^{-\gamma^2 G^2 (\Delta - \delta/3) D}
\]

(3)

where $S$ and $S_0$ represent the signal amplitude and the signal amplitude at zero gradient, $\gamma$ is the gyro magnetic ratio, $G$ is the gradient strength, $\delta$ is the gradient pulse length, $\Delta$ is the diffusion time, and $D$ is the apparent diffusion coefficient. Membrane samples were wound into a cylinder and suspended above saturated potassium chloride salt solution in a 5 mm NMR tube to generate an environment of 84% relative humidity at 30 °C. A pulse gradient stimulated echo sequence was performed using a Bruker AVANCEIII NMR spectrometer and 400 MHz wide bore Magnex Magnet. Proton diffusion measurements were made using a 5 mm Bruker single-axis DIFF60L Z-diffusion probe. The 90 degree pulse length was of the order of 5.0 μs. Typical parameters at 25 °C were $G = 0$–128 G cm$^{-1}$, incremented in 16 steps, $\delta = 1$ ms, $\Delta = 6$–700 ms. The Bruker TopSpin software was used for data acquisition and analysis. Multiple diffusion experiments were performed varying the time between pulses ($\Delta$) between 6 and 500 ms with a constant gyromagnetic ratio ($\gamma$) of 4258 Hz G$^{-1}$ and pulse length (δ) of 1 ms. The maximum gradient strength for each experiment was chosen to produce a full decay of the signal intensity over the length of the experiment. The decay of the signal intensity was plotted against gradient strength for each experiment and was fit to Eq. (1) to determine the diffusion coefficient.

2.3.6. Water uptake

Water uptake experiments were conducted using a dynamic vapor sorption (DVS) apparatus, DVS Advantage (Surface Measurement Systems Ltd). The DVS Advantage was used to measure the mass of water absorbed by the membrane in various humidity environments at 60 °C. The sequence of humidity conditions starts with 4h of dry nitrogen flow to ensure an accurate initial dry mass. A series of humidity steps follow, where humidified nitrogen is controlled to 20%, 40%, 60%, 80% and 95% relative humidity (RH) for 2 h each. The process is then reversed with 2h at each humidity back down to dry conditions.

Water uptake experiments were also conducted under environmental conditions mimicking the conditions of the small-angle X-ray scattering experiments. In these experiments humidity steps lasted only 17 min and cycled through 95% RH, 75% RH, 50% RH, 25% RH, dry and back to 95% RH nitrogen gas flow.

2.3.7. Ionic conductivity

In-plane conductivity was measured by electrochemical impedance spectroscopy (EIS) using a Bio-Logic VMP3 potentiosat. Data were collected and analyzed using EC Laboratories software. The membrane was held in a four-electrode test cell, with platinum electrodes. Chloride ion conductivity measurements were made while the sample was in a TestEquity H1000 oven that controls temperature and RH. At each RH studied, the temperature was varied from 50 to 90 °C by steps of 10 °C. The relative humidity is measured with a Vaisala humidity sensor. Feedback of this information controls saturated and dry air flow rates as necessary to maintain desired relative humidity throughout the experiment. Samples were allowed to equilibrate to each temperature set-point for 35 min before data was collected.

2.3.8. Counter-ion reaction in air

Infrared spectroscopy was used to monitor the reaction of hydroxide ions with atmospheric CO$_2$ to form (bi)carbonate species in the membrane. Quaternized membrane with a chloride counterion was suspended in chloroform and spin coated onto a glass slide. The membrane coated slide was left to dry in atmospheric conditions of temperature and pressure for about a week and a thin membrane (ca. 5 μm) was peeled from the glass slide. A few drops of water on the slide eased sample removal. The chloride counterions were exchanged to hydroxide ions by immersing the sample in 1 M NaOH solution for 1 day. The membrane in the hydroxide form was rinsed thoroughly with high purity nitrogen gassed DI water and patted with a Kimwipe to absorb excess water. The sample was transferred as quickly as possible to the ATR stage in order to maximize the amount of the (bi)carbonate reaction observed. ATR infrared spectra were collected using a Nexus 470 FT-IR spectroscopy unit. The ATR was equipped with a diamond ATR crystal and a MCT/A detector. 256 scans were averaged to obtain a
single beam sample spectrum and 10 scans were averaged to obtain a single beam background spectrum. FTIR spectra also were collected for a standard 1 M solution of potassium carbonate and potassium bicarbonate using FTIR spectroscopy. The range of wavenumbers observed for carbonates and bicarbonates were taken as benchmark position to identify the carbonates and bicarbonates peaks during dynamic spectra collection.

3. Results and discussion

3.1. Synthesis

Initial attempts to synthesize an AEM from the branched high MW PEI and iodoethane resulted in a hygroscopic material that was obviously impractical due to its solubility in water. When the dried product was exposed to ambient conditions (20 °C, 25% RH), it absorbed enough water to completely dissolve the material. Since a membrane could not be formed from the simple quaternized PEI we reasoned that animating with another polymer followed by quaternization should result in a water stable film. The reaction scheme in Fig. 1 was pursued to form a water insoluble material. To form a film, the product was treated with toluene to form a viscous paste and spread over a Teflon block. After the toluene evaporated, a yellow opaque visually defect free membranes was formed, 20 μm in thickness. The film was stable to water dissolution and was stored until use in water. The membrane ion exchange capacity, as measured by Mohr titration was found to be 0.74(0.04) mequiv. mol⁻¹.

3.2. Characterization

3.2.1. Reaction confirmation

The FTIR (Fig. 2) shows the starting CPP and the resultant quaternized PP-PEI polymer. The reaction is confirmed with the disappearance of the C-Cl stretch at 550 cm⁻¹ and the appearance of a broad peak that grows in at approximately 3450 cm⁻¹ corresponding to the N-C bonds in the ammonium groups and OH stretches of the water absorbed by the sample [9]. The spectra do not show all the same changes as those reported for the linear PEI/CPP materials and it appears that much of the polypropylene starter remains in the polymer.

Fig. 3 shows the ¹H MAS-NMR spectrum (a) and the ¹³C MAS-NMR spectrum (b). Because of the strong proton homonuclear dipolar coupling, the proton spectrum is very broad. The 12 kHz spin rate was insufficient to remove the dipolar couplings in the ¹H spectrum. The starting materials show well defined peaks for the methylene protons. The peak at δ = 100 ppm of the ¹³C spectrum confirms that the chloride in the starting CPP has fully reacted. The complex peak at ca. 40 ppm is assigned to the methylene carbons in the polymer backbone and the peak at 70 ppm to the methylene carbons adjacent to the quaternary ammonium cations.

3.2.2. Morphology

Wide-angle X-ray scattering (WAXS), not shown, had one peak corresponding to an amorphous polymer structure. The small angle X-ray scattering (SAXS) results are shown in Fig. 4. The scattering pattern had a constant Porod slope between 3.7 and 4.1 in the q-range of 0.006–0.04 Å⁻¹. In the dried sample this corresponds to a smooth spherical surface on the length scale of 16–105 nm. The reduction the porod slope after humidification is associated with a rougher or less spherical surface. The fact that there are no shoulders or peaks in the scattering pattern shows that there is no ordered structure on the length-scale associated with the q-range.
tested [18]. The material does not appear to phase separate or swell in the size range studied by SAXS.

Relative scattering intensity in organic polymer systems is generally correlated with water content. However, the scattering patterns in Fig. 4 show that the humidified samples have lower intensity relative to the dry sample. This could be attributed to water moving into structure with length scales too large to be observed in the q-range studied here.

3.2.3. Imaging

An ESEM image of the film from the top down, Fig. 5a shows that the material has an heterogeneous appearance with some 1 μm spherical clusters. This is very similar to the morphology observed by Hong et al. for the materials formed with the linear PEI [9]. When the same material is imaged with a visible microscope, not shown, further heterogeneity is observed. More intriguing is the apparent network of channels that appear to be have a width of <0.5 μm and extend across the entire imaged field. These features can be differentiated in the IR. An IR spectrum of this film is shown in Fig. 5b, for reference, In Fig. 5c we show the IR contour plot for the features corresponding to 3600 cm⁻¹ (the OH stretch of water). It indicates that water forms steaks across the film with a width of 150 μm. When we image the film at 2900 cm⁻¹ (Fig. 5d) corresponding to the methylene chains of the backbone of the polymer we see features again of a width of 150 μm, but more widespread, suggesting that the polymer may segregate into larger hydrophobic and hydrophilic domains.

3.2.4. Diffusion experiment

It is to be expected that anion transport will be mediated by water in this material and so we investigated the self-diffusion of water in these materials by PFG-NMR. Restricted diffusion was probed by a series of NMR experiments with varying times between pulses. The diffusion coefficient for each experiment was plotted vs. Δ as shown in Fig. 6. By linear regression of the low range data to zero Δ, D₀ was calculated to be $1.77 \times 10^{-3}$ cm² s⁻¹.
close to the value $2.5 \times 10^{-5}$ cm$^2$ s$^{-1}$, for free water. The maximum measurable diffusion time was $\Delta = 500$ ms here the diffusion coefficient appears to be approaching the infinite diffusion coefficient and so a simple extrapolation was used to obtain this value, $D_\infty = 3.72 \times 10^{-6}$ cm$^2$ s$^{-1}$, although the diffusion coefficient may continue decreasing slightly beyond this diffusion time. The continued decrease of the diffusion coefficient even at long diffusion times may be a result of transport observed in the much larger feature observed in the ESEM and IR imaging or due to the low resolution of the instrument at very large $\Delta$. Nevertheless it appears that there is considerable restriction of the diffusion in the material indicating tortuosity in the anion transport.

### 3.2.5. Water uptake

Water uptake (%) by mass was high as compared to representative AEMs investigated to date [19]. Water uptake reached its high of 70.3 wt% when the sample was at equilibrium with a 95% RH environment, nearly doubling when the relative humidity increased from 80% to 95%. The IEC and water uptake values were used to calculate lambda (Fig. 7). The highest water uptake converted to lambda is 53 water molecules per quaternary ammonium cation. The high lambda values are due to the extreme hydrophilicity of the quaternized PEI which is immobilized within the crosslinked polymer. Interestingly the membranes does not appear to swell dimensionally and so the water is absorbed by the free volume of the quaternized PEI moieties in the material.

### 3.2.6. Ionic conductivity

As expected, the ionic conductivity is highest under the highest temperature and humidity conditions tested (90°C and 95% RH). The highest chloride ion conductivity observed was 0.293 mS cm$^{-1}$, as shown in Fig. 8. This result is lower than for comparable AEMs. Interestingly, the chloride anion conductivity at 95%RH does not show Arrhenius behavior. When the film is tested at the drier 95%RH condition the conductivity decreases off at the higher temperatures. These samples were tested from low to high temperature and so this may be an indication of either cation instability or a significant loss of water. We were unable to measure hydroxide conductivity for these materials under RH cycling conditions as they did not survive the rigorous hydroxide exchange treatment used to obtain and mount films in the fixture in pure hydroxide form.

### 3.2.7. Counter-ion exchange in air

Despite the issues with attempting to make rigorous hydroxide conductivity measurements we were able to demonstrate that they reacted with ambient CO$_2$. Time resolved FTIR of the hydroxide film placed on an ATR crystal and exposed to ambient CO$_2$ is shown in Figs. 9 and 10. The growth of peaks at 1460 cm$^{-1}$ and 1380 cm$^{-1}$ assigned to the increasing carbonate and bicarbonate
counter-ions respectively within a hydroxide form membrane exposed to air (Figs. 9 and 10). These peaks were assigned based on comparisons to spectra of standard solutions of potassium carbonate and potassium bicarbonate, although slight differences were observed due to interactions with functional groups in the membrane. A maximum intensity for the two species is reached after 7 min, which presumably corresponds to the consumption of all of the hydroxide. After this time the ratio of carbonate to bicarbonate is maintained as the film slowly reaches equilibrium. A steady-state mixture is reached after 20 min of exposure to atmospheric CO₂ concentrations. These integrated areas are not corrected for the relative absorbance of the two different anions and so it is not implied that the concentration is greatest at 7 min simply that the initial ratio of anions produced eventually relaxes to an equilibrium value.

4. Conclusions

A new randomly crosslinked alkaline exchange membrane was produced through the amination of chlorinated polypropylene (CPP) with polyethyleneimine (PEI) and quaternization with iodoethane. The new material was characterized by FTIR and MAS-NMR. Environmentally controlled small/ wide angle X-ray analysis showed relatively featureless amorphous morphology through a full range of humidity environments over the 16–105 nm length scale. Larger scale ordering over >1 μm was observed in ESEM and IR microscope images implying that the water swelling occurs in larger features. However these features swell internally as little dimensional swelling of the material is observed.

Water self diffusion studies indicate rapid short-range diffusion, but heavily restricted diffusion over longer length scales. Temperature and humidity effects on the in-plane chloride ion conductivity were measured using electrochemical impedance spectroscopy (EIS). The highest chloride conductivity observed was 0.29 mS/cm at 90 °C and 95% relative humidity. Infrared spectroscopy was used to monitor the rate of counter-ion exchange from hydroxide to a mixture of carbonate and bicarbonate when exposed to air, which occurred in a time frame of minutes. Unfortunately these materials are too unstable to hydroxide for use in hydroxide exchange membrane fuel cells, but show interesting properties, such as high water uptake with little dimensional swelling, that may make them suitable for use in low temperature carbonate exchange membrane electrochemical systems.

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