Abstract: Here, we report the synthesis and thermophysical properties of seven primarily aromatic, imidazolium-based polyamide ionenes. The effects of varied para-, meta-, and ortho-connectivity, and spacing of ionic and amide functional groups, on structural and thermophysical properties were analyzed. Suitable, robust derivatives were cast into thin films, neat, or with stoichiometric equivalents of the ionic liquid (IL) 1-benzy-3-methylimidazolium bistriflimide ([Bnmim][Tf$_2$N]), and the gas transport properties of these membranes were measured. Pure gas permeabilities and permselectivities for N$_2$, CH$_4$, and CO$_2$ are reported. Consistent para-connectivity in the backbone was shown to yield the highest CO$_2$ permeability and suitability for casting as a very thin, flexible film. Derivatives containing terephthalamide segments exhibited the highest CO$_2$/CH$_4$ and CO$_2$/N$_2$ selectivities, yet CO$_2$ permeability decreased with further deviation from consistent para-linkages.

Keywords: aromatic polyamides; ionic liquids; ionenes; membranes; gas separations

1. Introduction

Gas separation membranes consisting of polymeric materials have drawn significant research attention, as the technology and materials are scalable and performance competes with more energy-intensive conventional techniques including pressure swing adsorption, cryogenic distillation, and gas–liquid absorption [1–4]. Materials considered to possess state-of-the-art gas separation performance feature certain functional and structural moieties, which may impart higher free volume distribution or promote flexibility and thermal stability. Polyimides [5–10], polymers of intrinsic microporosity (PIMs) [11,12], thermally-rearranged polymers (TR) [13–16], polyamides [17–19], and ion-containing polymers [20,21] are classes of polymeric materials which have been prevalent in this research area. Polyamides are another desirable material class as they feature H-bonding, which often produce self-assembled or well-ordered materials due to chain alignment [22–24].

Polyamide membranes and related composites are versatile, and perform well at various separation scales. Amide functionality in polymers has been shown to improve the mechanical strength, chemical and thermophysical stability due to the H-bonding and the induced ordering of polymer chains. Certain polyamides exhibit interesting surface behaviors, rendering a platform amenable to post-polymerization modifications. These surface interactions have been studied to show suitability for grafting or deposition upon other frameworks [25–29], or harnessed as antifouling and antimicrobial coatings [30–32]. Polyamide films have been explored for nanofiltration, reverse/forward osmosis, desalination [33,34], gas separations [17,35–39], and wastewater treatment [40]. The structuring and chain packing of the polyamide matrix is also impacted by the preparation technique, as films have
been processed by solution casting [17,35], layer-by-layer fabrication [41,42], 3D-printing [43,44], and interfacial polymerization [34,45,46] methods. Tailoring polymer structure and morphology via rational molecular design is important to advancing progress in membrane-based separations, as well as improving mechanical and thermal properties.

Recently, we have synthesized polyimide ionene (PI-ionene) and polyamide ionene (PA-ionene) materials for gas separation membranes, with the combination of functional groups and ionic components yielding flexible yet robust thin films via solvent-casting or melt-pressing. These ionenes incorporate imidazolium cations within the main backbone paired with “free” (i.e., untethered) fluorinated anions, spacing functional groups or high free volume elements [6,8,47]. Ionenes are an form of ionic polymers that is typically synthesized via condensation reactions [48]. The chain entanglement and organization in these PA-ionenes are dictated by ionic, H-bonding, π-cation and π–π stacking intermolecular interactions [49]. The contributions of these noncovalent interactions influence self-assembly and structuring of the matrix, which can be further tuned by adding “free” ionic liquids (ILs) to the PA-ionenes.

This work presents the synthesis of five new highly aromatic amide-linked bis(imidazole) monomers, which were subsequently polymerized via Menshutkin reactions yielding a set of imidazolium PA-ionene isomers with systematically varied aromatic connectivity/regiochemistry. Select derivatives were impregnated with IL, which acts as a non-volatile plasticizer, in order to probe the impact of these ionic interactions on nanostructuring and chain alignment as well as membrane fabrication. The effects of alternating connectivity between functional and charged features on chain packing and structuring are discussed, and the performance and gas transport properties of several solvent-cast films are analyzed.

2. Materials and Methods

2.1. Materials

Terephthaloyl chloride (TC, >99%), α, α’-Dichloro-p-xylene (pDCX, >98%), α, α’-Dichloro-m-xylene (mDCX, >96%), and α, α’-Dichloro-o-xylene (oDCX, >97%) were purchased from TCI (Portland, OR, USA). Isophthaloyl chloride (IC, 98%) and potassium carbonate (K₂CO₃, 99% anhydrous) were purchased from BeanTown Chemical (Hudson, NH, USA). Lithium bistriflimide (LiTf₂N) was purchased from 3M (Minneapolis, MN, USA). Acetonitrile (CH₃CN, ACS Grade) and N-methylpyrrolidone (NMP, ACS Grade) were purchased from VWR (Atlanta, GA, USA).

2.2. Characterization

¹H-NMR and ¹³C-NMR spectra were obtained using 360 MHz or 500 MHz Bruker Avance instruments (Madison, WI, USA). The thermal stabilities of these PA-ionenes were evaluated by thermogravimetric analysis (TGA) at a heating rate of 10 °C-min⁻¹ under a N₂ atmosphere (Seiko TG/DTA 7300). The glass transition temperature (Tg) of each PA-ionene was observed by DSC (TA Instruments, DSC Q200) (New Castle, DE, USA) from 30–300 °C with a scan rate of 10 °C-min⁻¹ under N₂. The wide-angle X-ray diffraction (WAXD) patterns of the materials were measured using a Bruker D8 Discover diffractometer by employing a scanning rate of 4° min⁻¹ in a 2θ range from 5 to 70° with a Co Kα₁ X-ray source (λ = 0.17886 nm). The d-spacing values were calculated using Bragg’s law (d = λ/2sinθ) and Diffrac-EVA software (Bruker). Number average molecular weight (Mₙ) values of the PA-ionenes were determined via MALDI-TOF MS (Bruker Ultraflex).
2.3. Synthesis of Bis-Imidazole Monomers Containing Amide Functionality

2.3.1. Synthesis of 4-(1H-imidazol-1-yl)aniline “I4A”, 3-(1H-imidazol-1-yl)aniline “I3A”, and 2-(1H-imidazol-1-yl)aniline “I2A”

The synthesis of 4-(1H-imidazol-1-yl)aniline (I4A) starting from imidazole and 4-fluoronitrobenzene followed by reduction in EtOH with H2 over Pd/C was reported in our previous work [47]. 1H NMR (500 MHz, DMSO-d6) δ 7.95 (s, 1H), 7.48 (s, 1H), 7.22 (d, 2H), 7.01 (s, 1H), 6.64 (d, J = 8.8, 2H), 5.26 (s, 2H). The synthesis of 3-(1H-imidazol-1-yl)aniline “I3A” was synthesized via a similar procedure and was outlined in our previous work [6]. 1H NMR (360 MHz, DMSO-d6) δ 8.08 (s, 1H), 7.57 (t, J = 1.3 Hz, 1H), 7.13 (t, J = 1.3 Hz, 1H), 7.08 (s, 2H), 6.73 (t, 1H), 6.72–6.67 (m, 1H), 6.60–6.53 (m, 1H), 5.41 (s, 2H). The synthesis of 2-(1H-imidazol-1-yl)aniline “I2A” was synthesized via a similar procedure and was outlined in our previous work [47]. 1H NMR (360 MHz, DMSO-d6) δ 7.74 (t, J = 1.1 Hz, 1H), 7.30 (t, J = 1.3 Hz, 1H), 7.14 (ddd, J = 8.1, 7.3, 1.6 Hz, 1H), 7.10 (t, J = 1.1 Hz, 1H), 7.03 (dd, J = 7.8, 1.6 Hz, 1H), 6.86 (dd, J = 8.1, 1.4 Hz, 1H), 6.64 (td, J = 7.5, 1.4 Hz, 1H), 5.06 (s, 2H).

2.3.2. Synthesis of Bis-Imidazole Amide Monomers

Five bis-imidazole monomers were synthesized, via condensation of 2 eq. of I4A, I3A, or I2A with 1 eq. TC or IC to form the amide linkage, according to Scheme 1.

TC (6.96 g, 34.3 mmol), I4A (12.0 g, 75.4 mmol), and K2CO3 (20.8 g, 150.7 mmol) were added to a 500 mL round-bottom-flask. CH3CN (150 mL) was added, and the flask was equipped with a stir bar and reflux condenser. The reaction was heated to 85 °C for 24 h. The mixture was allowed to cool to...
RT, and then was poured into 350 mL of deionized (DI) water. The precipitate was filtered and washed with 2 × 50 mL of deionized water, then dried in a vacuum oven at 100 °C for 24 h to yield the product “TC I4A” as a white powder (10.2 g, 67%). 1H NMR (360 MHz, DMSO-d6) δ 10.60 (s, 2H), 8.26 (s, 2H), 8.16 (s, 4H), 7.97 (d, J = 8.8 Hz, 4H), 7.76 (s, 2H), 7.69 (d, J = 8.8 Hz, 4H), 7.13 (s, 2H).

IC (4.93 g, 24.3 mmol), I3A (8.5 g, 53.4 mmol), and K2CO3 (14.8 g, 106.8 mmol) were reacted in 120 mL of CH3CN via a similar procedure to yield the product “IC I3A” as a white powder (10.4 g, 95%). 1H NMR (500 MHz, DMSO-d6) δ 10.69 (s, 2H), 8.60 (s, 1H), 8.20 (d, J = 6.0 Hz, 4H), 8.08 (s, 2H), 7.81 (d, J = 8.1 Hz, 2H), 7.75 (t, J = 7.7 Hz, 1H), 7.68 (s, 2H), 7.53 (t, J = 8.1 Hz, 2H), 7.43–7.38 (m, 2H), 7.15 (s, 2H).

TC (4.93 g, 24.3 mmol), I3A (8.5 g, 53.4 mmol), and K2CO3 (14.8 g, 106.8 mmol) were reacted in 120 mL of CH3CN via a similar procedure to yield the product “TC I3A” as a white powder (10.3 g, 94%). 1H NMR (500 MHz, DMSO-d6) δ 10.63 (s, 2H), 8.20 (s, 2H), 8.15 (br, 2H), 8.08 (s, 2H), 7.80–7.77 (m, 2H), 7.69 (s, 2H), 7.57–7.53 (m, 2H), 7.43–7.39 (m, 2H), 7.15 (s, 2H).

IC (4.93 g, 24.3 mmol), I4A (8.5 g, 53.4 mmol), and K2CO3 (14.8 g, 106.8 mmol) were reacted in 120 mL of CH3CN via a similar procedure to yield the product “IC I4A” as a white powder (10.7 g, 97%). 1H NMR (500 MHz, DMSO-d6) δ 13.13 (br, 2H) 8.93–8.84 (m, 2H), 8.46 (t, J = 7.7 Hz, 2H), 7.96 (d, J = 8.9 Hz, 4H), 7.74 (m, 4H), 7.67 (m, 4H), 7.11 (s, 2H).

TC (6.96 g, 34.3 mmol), I2A (12.0 g, 75.4 mmol), and K2CO3 (20.8 g, 150.7 mmol) were reacted in 150 mL of CH3CN via a similar procedure to yield the product “TC I2A” as a brown powder (2.07 g, 42%). 1H NMR (360 MHz, DMSO-d6) δ 13.13 (br, 2H) 8.93–8.84 (m, 2H), 8.46 (t, J = 7.7 Hz, 2H), 8.17–7.95 (m, 6H), 7.85–7.75 (m, 6H), 7.75–7.69 (m, 2H).

2.4. Synthesis of PA-Ionene Isomers

Seven PA-ionenes (Figure 1) were formed via the Menshutkin reaction, from bis-imidazole amide monomers with pDCX, mDCX, or oDCX.

The PA-ionenes were formed from equimolar amounts (1:1) of the respective bis-imidazole monomers and DCX derivatives. The monomers were added with 200 mL of NMP and a stir bar to a 500 mL round-bottom heavy-walled pressure vessel (Ace Glass, Vineland, NJ, USA) and sealed with PTFE screw caps. The reaction was heated at 150 °C for 24 h. Following polymerization, the reaction mixture was poured into a flask containing 500 mL DI water containing 2.5 eq. of LiTf2N under vigorous mechanical stirring to promote ion exchange from the [Cl−] to the [Tf2N−] form. The ionenes precipitate as a coarse powder which was collected by vacuum filtration, washed with 2 × 100 mL DI water, then collected to dry in a vacuum oven for 24–48 h at 100 °C. For brevity, the gram/molar amounts of each monomer and the respective yields for each PA-ionene are outlined in Table 1.
Many peaks in the $^1$H-NMR spectra for the PA-ionene are broad, due to the polymeric character. 

Table 1. Reactant Amounts and Polymer Yields for all seven PA-ionenes.

| ID | PA-Ionene | Bis-Imidazole | Amount | Linkage | Amount | LiTf$_2$N | Product Yield | Yield |
|----|-----------|---------------|--------|---------|--------|-----------|---------------|-------|
| 1  | [TC I4A pX][Tf$_2$N] | TC I4A | 10.1 g | 22.6 mmol | pDCX | 3.95 g | 22.6 mmol | 16.2 g | 21.5 g | 86% |
| 2  | [IC I3A mX][Tf$_2$N] | IC I3A | 8.00 g | 17.8 mmol | mDCX | 2.50 g | 5.57 mmol | 12.8 g | 14.1 g | 71% |
| 3  | [TC I3A mX][Tf$_2$N] | TC I3A | 2.50 g | 5.57 mmol | mDCX | 0.98 g | 5.57 mmol | 4.00 g | 3.89 g | 62% |
| 4  | [TC I3A pX][Tf$_2$N] | TC I3A | 2.50 g | 5.57 mmol | pDCX | 0.98 g | 5.57 mmol | 4.00 g | 4.31 g | 70% |
| 5  | [IC I4A px][Tf$_2$N] | IC I4A | 2.00 g | 4.46 mmol | pDCX | 0.78 g | 4.46 mmol | 3.20 g | 2.76 g | 56% |
| 6  | [IC I4A mX][Tf$_2$N] | IC I4A | 2.00 g | 4.46 mmol | mDCX | 0.78 g | 4.46 mmol | 3.20 g | 2.41 g | 49% |
| 7  | [TC I2A oX][Tf$_2$N] | TC I2A | 1.50 g | 3.35 mmol | oDCX | 0.58 g | 3.35 mmol | 2.40 g | 2.84 g | 76% |
**2.5. Gas Separation Measurements**

Ideal (i.e., single gas) permeation measurements of membranes formed from select PA-ionenes were performed using a high-vacuum time lag apparatus based on the constant-volume/variable-pressure method, as described in our previous works [6,8,17,47]. To avoid membrane fracture at the edges due to the pressure of the O-ring within the permeation unit, the membranes were masked on both sides using an adhesive aluminum tape in order to confine gas permeation through a fixed membrane area of $\frac{1}{2}$" diameter ($A = 0.196 \text{ in}^2$). These masked membranes were prepared from folded aluminum tape which was punched using a 47 mm punch to create two discs. The edges of the two circles were aligned and a $\frac{1}{2}$" hole was punched in the center, and the membrane was sealed between adhesive layers. The aluminum ‘tabbing’ method has been shown to have no effect on the permeability versus using a full membrane [17,50]. All permeation measurements were conducted at 20 °C and the feed pressure was ~3 atm (~45 psia) against initial downstream vacuum (<0.01 psia). Before each measurement, both the feed and the permeate sides were thoroughly evacuated to remove any residual gases. Pressures and temperatures of both the upstream and downstream volumes were monitored and recorded using the most recent version of LabVIEW software (National Instruments). Using these data, the permeability coefficient of each membrane was determined from the linear slope of the plot of downstream pressure rise relative to time ($\frac{dp}{dt}$) according to the following equation:

$$P = \frac{273}{76} \times \frac{V l}{A T p_0} \times \frac{dp}{dt}$$

where $P$ is the permeability expressed in barrer (1 barrer = $10^{-10} \text{ cm}^3 \text{ (STP) cm cm}^{-2} \text{ s}^{-1} \text{ cmHg}^{-1}$); $V$ (cm$^3$) is the downstream volume; $l$ (cm) is the thickness of the membrane; $A$ (cm$^2$) is the effective gas permeation area of the membrane; $T$ (K) is the measurement temperature; $p_0$ (Torr) is the pressure of the feed gas at the upstream and $\frac{dp}{dt}$ is the rate of the steady-state pressure rise. The ideal permselectivity ($\alpha_{A/B}$) for a pair of gases (A and B) was calculated from the ratio of the individual gas permeability coefficients as, $P_A/P_B$.

**3. Results & Discussion**

**3.1. Characterization of Materials**

All PA-ionenes were thoroughly characterized, to quantify polymer properties prior to membrane formation and testing. This allows for correlation of repeat unit structure and backbone connectivity with polymer properties. As demonstrated in the supporting information, the purity and structures of all the monomers and polymers were first confirmed by $^1$H-NMR (Figures S1–S12). Density data were collected for each polymer and reported in Table 2. The density measurements were performed using a Mettler Toledo Density Kit (Product MS-DNY-54), paired with a Mettler Toledo analytical balance. This kit allows for solid state density determinations, which is based on Archimedes’ principle. Bulk samples were prepared (dried thoroughly) and tested in triplicate. The samples were weighed at room temperature in air and then in a liquid of known density, in this case, heptane. It should be noted that these ionenes are completely insoluble in heptane nor do they show any swelling/solvent...
interactions. The measurement was performed using the buoyancy method, and calculations were based on the following equation,

$$\rho_{\text{polymer}} = \frac{W_0}{W_0 - W_1} \times \rho_{\text{liquid}}$$

(2)

where $W_0$ and $W_1$ are the membrane weights in air and heptane, respectively. The uncertainty for these measurements ±0.01 g [51]. The molecular weight of these ionenes was determined using MALDI-TOF MS experiments, as summarized in Table 2 and included in Figure 2. The thermal transitions and degradation of these PA-ionenes was also investigated using DSC and TGA techniques, and obtained $T_g$ values are included in Table 2 (Figures S14–S16).

**Table 2. Summary of thermal, structural, and molecular weight data for PA-ionenes.**

| ID | PA-Ionene                  | IL (Equivalents) | $T_g$ (°C) | $d$-Spacing (Å) | Density (g/cm$^3$) | $M_N$ (kDa) |
|----|---------------------------|------------------|------------|-----------------|-------------------|-------------|
| 1  | [TC I4A pX][Tf$_2$N]      | Neat             | 136.1      | 4.07            | 1.517             | 60.4        |
| 1+IL| [Bmim][Tf$_2$N] (1)       | -                | 4.26       | 1.575           | *                |
| 2  | [IC I3A mX][Tf$_2$N]      | Neat             | 133.5      | 4.35            | 1.443             | 62.5        |
| 2+IL| [Bmim][Tf$_2$N] (1)       | -                | 4.55       | 1.259           | *                |
| 3  | [TC I3A mX][Tf$_2$N]      | Neat             | 54.5       | 4.36            | 1.388             | 59.9        |
| 4  | [IC I4A pX][Tf$_2$N]      | Neat             | 74.7       | 4.39            | 1.427             | 75.4        |
| 5  | [IC I4A pX][Tf$_2$N]      | Neat             | 112.0      | 4.97            | 1.505             | 66.4        |
| 6  | [IC I4A mX][Tf$_2$N]      | Neat             | 115.7      | 5.09            | 1.462             | 76.1        |
| 7  | [TC I2A oX][Tf$_2$N]      | Neat             | 76.4       | 4.47            | 1.577             | 117.7       |

* MALDI-TOF MS analysis was not performed for derivatives containing IL, as the same polymer from 1 and 2 was used in the preparation of samples 1 + IL and 2 + IL and thus has the same $M_N$ value.

3.1.1. Structural Characterizations

$M_N$ values were determined via MALDI-TOF MS (Figure 2) and range from 60–118 kDa, confirming that these combinations of monomers are amenable to the formation of high molecular weight polymers. Given these $M_N$ values and utilizing the repeat unit molecular weight (1112.91 g/mol, including “free” anions), number average degree of polymerization ($X_N$) values of 54–104 repeat units were calculated. These $X_N$ values are consistent with those calculated by $^1$H-NMR end group analysis using the ratio of residual dichloro-xylene peaks relative to the polymerized xylyl linkage, which results in a shift of the benzylic CH$_2$ protons from ~4.3–4.6 ppm to ~5.5–5.6 ppm. These shifts are pronounced and clear, although integration and comparison of the shifting proton peaks for the imidazole to imidazolium transformation could also be used. From these integration-based ratios, conversion ($\rho$) and thus $X_N$ were derived using Carothers’ equation, with calculated $X_N$ values between 50–100.
weight polymers. Given these $M_N$ values and utilizing the repeat unit molecular weight (1112.91 g/mol, including “free” anions), number average degree of polymerization ($X_N$) values of 54–104 repeat units were calculated. These $X_N$ values are consistent with those calculated by $1H$-NMR end group analysis using the ratio of residual dichloro-xylene peaks relative to the polymerized xylyl linkage, which results in a shift of the benzylic $CH_2$ protons from ~4.3–4.6 ppm to ~5.5–5.6 ppm. These shifts are pronounced and clear, although integration and comparison of the shifting proton peaks for the imidazole to imidazolium transformation could also be used. From these integration-based ratios, conversion ($\rho$) and thus $X_N$ were derived using Carothers’ equation, with calculated $X_N$ values between 50–100.

Figure 2. MALDI-TOF MS data for all neat PA-ionene derivatives.

The structure was also probed using XRD, as the corresponding $d$-spacing values from these profiles have been associated with estimation of interchain spacing in polymers, which is influential with respect to film formation and gas permeability. Two broad halos are observed for all derivatives in the XRD profiles obtained in the 5°–70° 2$\theta$ spectrum, as shown in Figure 3. Typically, polymers do not show defined sharp peaks indicative of significant crystallinity in the bulk material, but instead, show a distribution in the form of an amorphous peak. The center of the distribution, or “peak”, of the main halo is centered between $2\theta = 20.2°–25.4°$, determined using Diffrac-EVA software. Using Bragg’s law, the corresponding $d$-spacing were calculated in the range of 4.07–5.09 Å. These diffraction profiles support that the distribution of $d$-spacings are similar for this set of PA-ionenes, and reasonable in comparison to other polyamides in literature [52]. It has been shown that $d$-spacing values are representative of interchain spacing and can be used as an approximation for the distance between polymer chains [53]. The intersegmental distances are very closely correlated with derivatives in which the same monomer is used, showing that the connectivity of the xylyl linkage is not as influential. The addition of IL shows a consistent increase in $d$-spacing and sharpening of the main halo peak, which suggests that the effects of IL may lead to greater regularity in the self-assembly and structuring of the matrix. Derivatives 3–6 have mixed connectivity and exhibit the highest $d$-spacing values, which is supported by increased gas permeability when compared to more linear derivatives with consistent connectivity such as 1. No apparent trend was observed when analyzing the density values, though all fell within a consistent range. However, the rigidity and inherent strain of the ortho-connectivity demonstrated in polymer 7 must greatly influence the regularity of chain packing, indicated by the highest density and narrowest XRD halo.
derivatives in which the same monomer is used, showing that the connectivity of the xylyl linkage is not as influential. The addition of IL shows a consistent increase in d-spacing and sharpening of the main halo peak, which suggests that the effects of IL may lead to greater regularity in the self-assembly and structuring of the matrix. Derivatives 3–6 have mixed connectivity and exhibit the highest d-spacing values, which is supported by increased gas permeability when compared to more linear derivatives with consistent connectivity such as 1.

No apparent trend was observed when analyzing the density values, though all fell within a consistent range. However, the rigidity and inherent strain of the ortho-connectivity demonstrated in polymer 7 must greatly influence the regularity of chain packing, indicated by the highest density and narrowest XRD halo.

![Figure 3. XRD profiles for all PA-ionene derivatives.](image)

3.1.2. Thermal Characterizations

The thermal behaviors of each material were analyzed using DSC to probe thermal transitions and TGA to determine stability/degradation. Full DSC cycles were run multiple times with polymers which had been dried for 72 h at 100 °C. Precise T_g and T_m values for these ionenes were difficult to extract due to relatively broadened transitions possibly due to the complex combination of intermolecular interactions. As indicated in Table 2, T_g values for these ionenes range from 54–136 °C. These T_g values were identified through analysis of multiple scans, and correlation of the endotherm on the ramp from 30–300 °C to the corresponding exotherm on the ramp from 300–30 °C (Figures S14 and S15). While the DSC scans were only run up to 300 °C to avoid approaching degradation, it can be seen that each ionene exhibits another broad transition, suggesting T_m values in the range of 200–320 °C. From TGA analysis, these PA-ionenes are stable at elevated temperatures, though certain cases indicate insufficient drying of these samples. Though some mass loss under 200 °C may have occurred due to the removal of residual trapped solvents, which would imply an underestimation of T_d, these polymers exhibit T_d, 10% values in the range of 290–360 °C (Figure S16). Generally, the more stable derivatives have consistent connectivity along the backbone (i.e., TC I4A pX, or p-p-p), rather than a mix of para-, meta-, and ortho-connectivity (i.e., TC I3A pX, or p-p-p).

3.2. Membrane Fabrication

Solutions (5 wt%) of each PA-ionene, neat or with 1 eq. of [Bnmim][Tf_2N] per repeat unit, were prepared using 0.4–0.6 g of neat polymer in DMAc. The solutions were prepared in 15 mL centrifuge tubes which were placed in a hot water bath for 2 h, to promote the dissolution and distribution of...
The resultant film thicknesses for these derivatives were 120–140 μm, as initially measured with digital calipers and later confirmed by SEM images of cross-sections (Figures S17–S19). It should be noted that derivative 1 was very mechanically durable and flexible when cast as much thinner films, contrary to other derivatives that required greater thicknesses for sufficient testing as unsupported membranes. Thus, [TC I4A pX][Tf2N] was solvent cast to as thin as 4.1 μm to demonstrate its mechanical stability and permeation consistency even as a thinner membrane. Casting of [IC I3A mX][Tf2N] and [TC I2A oX][Tf2N] derivatives (2, 7) was attempted but yielded brittle, paper-like materials which did not hold under pressure. This is likely due to poor entanglement or inability to properly align for H-bonding of consistent meta-connectivity for 2, the strain of the ortho-connectivity along the backbone for 7. Figure 4 shows the materials which were amenable to film formation, demonstrating flexibility and stability as thin, transparent membranes.

Figure 4. (a) Images of solvent cast films (d = 60 mm) of four PA-ionenes introduced in this work; (b) Comparison of [TC I4A pX][Tf2N] as a neat solvent cast film versus ionene-containing [Bnmim][Tf2N].

3.3. Gas Permeation Testing

Pure gas permeation measurements were performed to determine the gas separation performance of these aromatic PA-ionene membranes using a high-vacuum time lag measurement unit based on the constant-volume/variable-pressure method. Our previous work discusses the specifications and construction of the units [8,47]. All measurements were conducted at 20 °C and ~3 atm (~45 psia) feed pressure against initial downstream vacuum (<0.01 psia). Thin films were cast of derivatives 1–5 via the methods outlined above. The PA-ionenes with consistent connectivity along the backbone (i.e., [TC I4A pX][Tf2N] = para-para-para; [IC I3A mX][Tf2N] = meta-meta-meta) exhibited notably higher glass transitions and suitable mechanical stability in the para-case. The film of derivative 2 appeared opaque,
paper-like, and thus brittle. Derivatives 3–6 also yielded transparent thin films with good properties, despite the inconsistent connectivity, but these did possess lower \( T_g \) values and broader distributions seen in the MALDI-TOF MS data. Thus, 1 was selected for comparison when IL is incorporated into the matrix. The addition of \([Bnmm][Tf_2N] \) improved the flexibility of the membranes, and altered the optical qualities of the films. This induced opacity may be attributed to the additional ordering and coordination which occurs when IL is added to the ionene. The IL was expected to regulate and increase the free volume, as indicated by an increase in \( d \)-spacing. However, a stark decrease in permeability was observed from the addition of IL. Therefore, derivatives 3–5 were tested as neat membranes.

As shown in Table 3, PA-ionene 1 which contained consistent \textit{para}-connectivity exhibited the best gas permeation performance as well as stability. This material had the highest \( T_g \) and was robust mechanically even as the thinnest film cast in this work (4.1 \( \mu \)m). To ensure consistency of permeation experiments, this permeation data was also collected with a membrane sample with a thickness in the same range as 3–6. The CO\(_2\) permeability was 11 barrer, which yielded high selectivity relative to N\(_2\) and CH\(_4\). The stoichiometric incorporation of IL into the matrix drastically reduced permeability for all gases, particularly for CO\(_2\) which decreased by nearly 95\%. Derivative \( 1 + \text{IL} \) exhibited very low gas permeability, which may be due to increased interactions between chains tightening the matrix. While other similar PI-ionenes and PA-ionenes have shown improved performance with the addition of IL, these PA-ionenes form a denser polymer matrix when neat due to their primarily aromatic character and combined H-bonding, ionic, and stacking intermolecular interactions. Rather than pushing chains apart, in this case, the introduction of IL may improve coordination of the ionene and tighten the interchain spacing or fill free volume, and inhibiting gas permeation. Derivatives 3 and 4 exhibited comparable \( d \)-spacing yet 4 was notably denser, which is reflected by greater gas permeability in 3 versus 4. As both have \textit{para}-connectivity from the terephthalamide segment, 3 has adjacent \textit{meta}-connectivity in the imidazole-amide connection with the \textit{meta}-xylyl linkage, while 4 contains adjacent \textit{meta}-connectivity in the imidazole-amide connection with the \textit{para}-xylyl linkage. This suggests that further deviation from consistent \textit{para}-connectivity corresponds to decreasing permeability and that connectivity within these various repeat unit segments does influence the efficiency of chain packing. Comparing the three TC-based ionenes (1, 3, 4), the initial [TC 14A pX][Tf\(_2\)N] ionene exhibited CO\(_2\) permeability which was nearly 3\( \times \) or 4.5\( \times \) higher than 3 and 4, respectively. However, these all showed high selectivity for CO\(_2\), as \( N_2 \) and CH\(_4\) permeability was not promoted by this system. The spacing between the amide groups via \textit{para}-connectivity may improve H-bonding and chain interactions, demonstrated by the greater mechanical stability of these materials. Shifting to derivative 5, which contains the isophthalimide linkage adjacent to \textit{para}-connectivity between the imidazolium and amide groups, obtained relatively low permeabilities for all the gases with superior CO\(_2\) selectivities over \( N_2 \) and CH\(_4\) gases, plausibly due to the high density despite the increased interchain \( d \)-spacing value compared to 3 and 4.

### Table 3. Pure gas permeability and permselectivity data for PA-ionenes at 20 °C and ~3 atm.

| ID | PA-Ionene | \( P_{CO_2} \) | \( P_{N_2} \) | \( P_{CH_4} \) | \( a_{CO_2/N_2} \) | \( a_{CO_2/CH_4} \) |
|----|-----------|----------------|----------------|----------------|-----------------|-----------------|
| 1  | [TC 14A pX][Tf\(_2\)N] | Neat | 11.00 ± 0.02 | 0.34 ± 0.02 | 0.23 ± 0.01 | 32.6 ± 1.9 | 46.9 ± 2.0 |
| 2  | [Bnmm][Tf\(_2\)N] | [TC 14A pX][Tf\(_2\)N] | 4.043 ± 0.06 | 0.112 ± 0.002 | 0.087 ± 0.031 | 3.6 ± 0.5 | 4.6 ± 1.8 |
| 3  | [TC 14A mX][Tf\(_2\)N] | Neat | 3.91 ± 0.04 | 0.096 ± 0.004 | 0.122 ± 0.003 | 40.9 ± 1.8 | 30.5 ± 0.08 |
| 4  | [TC 14A pX][Tf\(_2\)N] | Neat | 2.46 ± 0.03 | 0.082 ± 0.005 | 0.103 ± 0.003 | 30.2 ± 1.4 | 23.9 ± 0.08 |
| 5  | [IC 14A pX][Tf\(_2\)N] | Neat | 1.69 ± 0.05 | 0.046 ± 0.004 | 0.039 ± 0.005 | 36.6 ± 3.4 | 43.3 ± 5.7 |

\( a \) Permeability reported in barrer, where 1 barrer = \( 1 \times 10^{-10} \) [cm\(^3\) (STP) cm]/(cm\(^2\) s cm Hg). \( b \) Error analysis for permeability was calculated from variation between multiple gas permeation runs. The propagation of this error was calculated for the selectivity ratios.
4. Conclusions

A series of seven bis-imidazole amide monomers that were polymerized with xylyl linkages to form imidazolium PA-ionene isomers with varied connectivity. The synthetic methods were simple and each product was obtained in high yields. The influence of connectivity in each segment was probed through thermal and structural characterization, and four of the seven polymers provided suitable very thin, unsupported membranes. [TC I4A pX][Tf2N] with consistent para-connectivity corresponded to the most stable, robust material with the highest CO2 permeability and also yielded superior CO2/CH4 and CO2/N2 selectivities. Nonetheless, the addition of free IL significantly reduces the permeability of [TC I4A pX][Tf2N], plausibly because of dense amorphous chain packing regardless to the free IL content, which was in good agreement with the density measurements. Derivatives 3–5, which incorporated varied connectivity along the PA-ionene backbone, exhibited moderate CO2 permeabilities without significant loss in selectivities. While certain isophthalamide-based systems were less mechanically stable and permeable, derivative 5, which contains the isophthalamide linkage adjacent to para-connectivity, displayed enhanced CO2/N2 and CO2/CH4 selectivities. This work highlights the beneficial features of different amide linkages in the ionene backbone. Therefore, future investigations will explore more on the incorporation of other functional groups and spacing of the alternating ionic and functional groups.

Supplementary Materials: The following are available online at http://www.mdpi.com/2077-0375/10/3/51/s1. Figure S1: 1H-NMR spectrum for TC I4A monomer, Figure S2: 1H-NMR spectrum for IC I4A monomer, Figure S3: 1H-NMR spectrum for TC I3A monomer, Figure S4: 1H-NMR spectrum for IC I4A monomer, Figure S5: 1H-NMR spectrum for TC I2A monomer, Figure S6: 1H-NMR spectrum for [TC I4A pX][Tf2N], Figure S7: 1H-NMR spectrum for [IC I3A mX][Tf2N], Figure S8: 1H-NMR spectrum for [TC I3A pX][Tf2N], Figure S9: 1H-NMR spectrum for [IC I3A mX][Tf2N], Figure S10: 1H-NMR spectrum for [IC I4A mX][Tf2N], Figure S11: 1H-NMR spectrum for [IC I4A pX][Tf2N], Figure S12: 1H-NMR spectrum for [IC I2A oX][Tf2N], Figure S13: MALDI-TOF spectra for all neat polyamide-ionenes, Figure S14: DSC cycles for all neat derivatives, Figure S15: Section of DSC cycles where Tg values are observed, Figure S16: TGA data for a selection of neat polyamide-ionenes, Figure S17: Cross-section of freeze-fractured, solvent-cast membrane of [TC I4A pX][Tf2N] at 9500X magnification. Thickness was measured to be 4.17 microns, Figure S18: Cross-section of freeze-fractured, solvent-cast membrane of [IC I3A mX][Tf2N] at 300X magnification. Thickness was measured to be 68.7 microns, Figure S19: Cross-section of freeze-fractured, solvent-cast membrane of [IC I3A mX][Tf2N] at 150X magnification. Thickness was measured to be 68.7 microns.

Author Contributions: Conceptualization: K.E.O. and J.E.B.; Methodology, K.E.O. and I.K.; Synthesis, K.E.O., E.M.T. and D.M.N.; Characterization K.E.O., E.M.T. and D.M.N.; Membrane Fabrication: K.E.O., I.K.; Data Curation, K.E.O. and I.K. and G.P.D.; Formal Analysis, K.E.O.; Gas Permeation Experiments, I.K.; Preparation of Figures, K.E.O.; Preparation of Tables, K.E.O. and I.K.; Thermal Analysis, K.E.O. and E.M.J.; Resources, J.E.B.; Writing—Original Draft Preparation, K.E.O.; Writing—Review & Editing, K.E.O., I.K., and J.E.B.; Supervision, J.E.B.; Project Administration, J.E.B.; Funding Acquisition, J.E.B. All authors have read and agreed to the published version of the manuscript.

Funding: This material is based upon work supported by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences, Separation Science program under Award Number DE-SC0018181. Thermal characterization were supported by a grant from NASA (CAN 80MSFC18M0041). Support from the National Science Foundation from the Major Research Instrumentation Program (CHE-1726812) for the purchase of the MALDI-TOF MS is gratefully acknowledged. K.E.O. is supported by the U.S. Department of Education as a GAANN Teaching Fellow.

Acknowledgments: The authors acknowledge Christopher S. Brazel of the University of Alabama ChBE department for allowing use of the DSC in his laboratory.

Conflicts of Interest: The authors declare no conflict of interest.

References
1. Chiappetta, G.; Clarizia, G.; Drioli, E. Design of an integrated membrane system for a high level hydrogen purification. Chem. Eng. J. 2006, 124, 29–40. [CrossRef]
2. Bernardo, P. Membrane gas separation: A review/state of the art. Ind. Eng. Chem. Res. 2009, 48, 4638–4663. [CrossRef]
3. Koros, W.J.; Fleming, G.K. Membrane-based gas separation. J. Membr. Sci. 1993, 83, 1–80. [CrossRef]
4. Koros, W.J. Polyamides and Polypyrrolones for Fluid Separation Membranes. U.S. Patent 5,262,056, 16 November 1993.

5. Qiu, W.; Xu, L.; Chen, C.-C.; Paul, D.R.; Koros, W.J. Gas separation performance of 6FDA-based polyimides with different chemical structures. Polymer 2013, 54, 6226–6235. [CrossRef]

6. O’Harra, K.E.; Kammakakam, I.; Devriese, E.M.; Noll, D.M.; Bara, J.E.; Jackson, E.M. Synthesis and performance of 6FDA-based polyimide-ionsenes and composites with ionic liquids as gas separation membranes. Membranes 2019, 9, 79.

7. Xiao, Y.; Low, B.T.; Hosseini, S.S.; Chung, T.S.; Paul, D.R. The strategies of molecular architecture and modification of polyimide-based membranes for CO2 removal from natural gas—A review. Prog. Polym. Sci. 2009, 34, 561–580. [CrossRef]

8. Mittenthal, M.S.; Flowers, B.S.; Bara, J.E.; Whitley, J.W.; Spear, S.K.; Roveda, J.D.; Wallace, D.A.; Shannon, M.S.; Holler, R.; Martens, R.; et al. Ionic polyimides: Hybrid polymer architectures and composites with ionic liquids for advanced gas separation membranes. Ind. Eng. Chem. Res. 2017, 56, 5055–5069. [CrossRef]

9. Zhuang, Y.; Seong, J.G.; Lee, Y.M. Polyimides containing aliphatic/acyclic segments in the main chains. Prog. Polym. Sci. 2019, 92, 35–88. [CrossRef]

10. Kammakakam, I.; Nam, S.; Kim, T.-H. Ionic group-mediated crosslinked polyimide membranes for enhanced CO2 separation. RSC Adv. 2015, 5, 69907–69914. [CrossRef]

11. McKeown, N.B. Polymers of intrinsic microporosity. ISRN Mater. Sci. 2012, 2012, 513986. [CrossRef]

12. Budd, P.; McKeown, N.; Ghanem, B.; Msayib, K.; Fritsch, D.; Starannikova, L.; Belov, N.; Sanfirova, O.; Yampolskii, Y.; Shantarovich, V. Gas permeation parameters and other physicochemical properties of a polymer of intrinsic microporosity: Polybenzodioxane pim-1. J. Membr. Sci. 2008, 325, 851–860. [CrossRef]

13. Smith, J.D.; Hou, R.; Lau, C.H.; Konstas, K.; Kitchin, M.; Dong, G.; Lee, J.; Lee, W.H.; Seong, J.G.; Lee, Y.M.; et al. Highly permeable thermally rearranged mixed matrix membranes (tr-mmm). J. Membr. Sci. 2019, 585, 260–270. [CrossRef]

14. Li, S.; Jo, H.J.; Han, S.H.; Park, C.H.; Kim, S.; Budd, P.M.; Lee, Y.M. Mechanically robust thermally rearranged (tr) polymer membranes with spiribisindane for gas separation. J. Membr. Sci. 2013, 434, 137–147. [CrossRef]

15. Shamsipur, H.; Dawood, B.A.; Budd, P.M.; Bernardo, P.; Clarizia, G.; Jansen, J.C. Thermally rearrangeable pim-polyimide membranes for gas separation. Macromolecules 2014, 47, 5595–5606. [CrossRef]

16. Kim, S.; Lee, Y.M. Thermally rearranged (tr) polymer membranes with nanoengineered cavities tuned for CO2 separation. J. Nanopart. Res. 2012, 14, 949. [CrossRef]

17. Kammakakam, I.; O’Harra, K.E.; Dennis, G.P.; Jackson, E.M.; Bara, J.E. Self-healing imidazolium-based ionene-polyamide membranes: An experimental study on physical and gas transport properties. Polym. Int. 2019, 68, 1123–1129. [CrossRef]

18. Hoehn, H.H. Aromatic polycarbonate membranes. Mater. Sci. Synth. Membr. 1985, 269, 81–98.

19. Kirsh, Y.E. Ion-exchange, pervaporation and reverse-osmosis membranes based on aromatic polyamides: Structure, selectivity, permeability and mechanisms of function. Ross. Khimicheskii Zhurnal 1998, 42, 117–129.

20. Yuan, J.; Mecerreyes, D.; Antonietti, M. Poly(ionic liquid)s: An update. Prog. Polym. Sci. 2013, 38, 1009–1036. [CrossRef]

21. Tome, L.C.; Marrucho, I.M. Ionic liquid-based materials: A platform to design engineered CO2 separation membranes. Chem. Soc. Rev. 2016, 45, 2785–2824. [CrossRef]

22. Bara, J.E.; O’Harra, K.E.; Durbin, M.M.; Dennis, G.P.; Jackson, E.M.; Thomas, B.; Odutola, J.A. Synthesis and characterization of ionene-polyamide materials as candidates for new gas separation membranes. MRS Adv. 2018, 3, 3091–3102. [CrossRef] [PubMed]

23. Tiwari, S.C.; Shimamura, K.; Mishra, A.; Shimojo, F.; Nakano, A.; Kalia, R.K.; Vashishta, P.; Branicio, P.S. Hydrogen bond preserving stress release mechanism is key to the resilience of aramid fibers. J. Phys. Chem. B 2019, 123, 9719–9723. [CrossRef] [PubMed]

24. Murthy, N.S. Hydrogen bonding, mobility, and structural transitions in aliphatic polyamides. J. Polym. Sci. Part B Polym. Phys. 2006, 44, 1763–1782. [CrossRef]

25. Ji, C.; Xue, S.; Tang, Y.-J.; Ma, X.-H.; Xu, Z.-L. Polyamide membranes with net-like nanostructures induced by different charged moieties for elevated nanofiltration. ACS Appl. Polymer Mater. 2019. [CrossRef]

26. Guo, Y.-S.; Mi, Y.-F.; Ji, Y.-L.; An, Q.-F.; Gao, C.-J. One-step surface grafting method for preparing zwitterionic nanofiltration membrane via in situ introduction of initiator in interfacial polymerization. ACS Appl. Polymer Mater. 2019, 1, 1022–1033. [CrossRef]
27. Yu, H.-Y.; Kang, Y.; Liu, Y.; Mi, B. Grafting polyzwitterions onto polyamide by click chemistry and nucleophilic substitution on nitrogen: A novel approach to enhance membrane fouling resistance. *J. Membr. Sci.* **2014**, *449*, 50–57. [CrossRef]

28. Rahaman, M.S.; Thérien-Aubin, H.; Ben-Sasson, M.; Ober, C.K.; Nielsen, M.; Elimelech, M. Control of biofouling on reverse osmosis polyamide membranes modified with biocidal nanoparticles and antifouling polymer brushes. *J. Mater. Chem. B* **2014**, *2*, 1724–1732. [CrossRef]

29. Perreault, F.; Tousley, M.E.; Elimelech, M. Thin-film composite polyamide membranes functionalized with biocidal graphene oxide nanosheets. *Environ. Sci. Technol. Lett.* **2013**, *1*, 71–76. [CrossRef]

30. Ma, W.; Chen, T.; Nanni, S.; Yang, L.; Ye, Z.; Rahaman, M.S. Zwitterion-functionalized graphene oxide incorporated polyamide membranes with improved antifouling properties. *Langmuir* **2019**, *35*, 1513–1525. [CrossRef]

31. Tiraferri, A.; Vecitis, C.D.; Elimelech, M. Covalent binding of single-walled carbon nanotubes to polyamide membranes for antimicrobial surface properties. *ACS Appl. Mater. Interfaces* **2011**, *3*, 2869–2877. [CrossRef]

32. Duong, P.H.H.; Hong, P.-Y.; Musteata, V.; Peinemann, K.V.; Nunes, S.P. Thin film polyamide membranes with photosensitive antibacterial activity. *ChemistrySelect* **2017**, *2*, 6612–6616. [CrossRef]

33. Gohil, J.M.; Ray, P. A review on semi-aromatic polyamide tfc membranes prepared by interfacial polymerization: Potential for water treatment and desalination. *Sep. Purif. Technol.* **2017**, *181*, 159–182. [CrossRef]

34. Kong, C.; Kanemoto, T.; Shintani, T.; Tsuru, T. Controlled synthesis of high performance polyamide membrane with thin dense layer for water desalination. *J. Membr. Sci.* **2010**, *362*, 76–80. [CrossRef]

35. Morisato, A.; Ghosal, K.; Freeman, B.D.; Chern, R.T.; Alvarez, J.C.; de la Campa, J.G.; Lozano, A.n.E.; de Abajo, J. Gas separation properties of aromatic polyamides containing hexafluoroisopropylidene groups. *J. Membr. Sci.* **1995**, *104*, 231–241. [CrossRef]

36. Ren, L.; Liu, J. Synthesis and gas transport properties of polyamide membranes containing pdms groups. *RSC Adv.* **2019**, *9*, 9737–9744. [CrossRef]

37. Rabiee, H.; Ghadimi, A.; Mohammad, T. Gas transport properties of reverse-selective poly(ether-b-amide6)[emim][bf4] gel membranes for co2/light gases separation. *J. Membr. Sci.* **2015**, *476*, 286–302. [CrossRef]

38. González-Díaz, M.O.; Sulub-Sulub, R.; Herrera-Kao, W.; Vázquez-Torres, H.; Zolotukhin, M.G.; Aguilar-Vega, M. Enhanced gas transport performance of polyamide membranes by postpolymerization modification. *Ind. Eng. Chem. Res.* **2018**, *57*, 8989–8996. [CrossRef]

39. Bisoi, S.; Mandal, A.K.; Singh, A.; Banerjee, S. Gas separation properties of troeger’s base-bridged polyamides. *e-Polymers* **2017**, *17*, 283–293. [CrossRef]

40. Amoli-Diva, M.; Irani, E.; Pourghazi, K. Photocatalytic filtration reactors equipped with bi-plasmonic nanocomposite/poly acrylic acid-modified polyamide membranes for industrial wastewater treatment. *Sep. Purif. Technol.* **2020**, *236*, 116257. [CrossRef]

41. Gu, J.-E.; Lee, J.S.; Park, S.-H.; Kim, I.T.; Chan, E.P.; Kwon, Y.-N.; Lee, J.-H. Tailoring interlayer structure of molecular layer-by-layer assembled polyamide membranes for high separation performance. *Appl. Surf. Sci.* **2015**, *356*, 659–667. [CrossRef]

42. Rajesh, S.; Zhao, Y.; Fong, H.; Menkhaus, T.J. Nanofiber multilayer membranes with tailored nanochannels prepared by molecular layer-by-layer assembly for high throughput separation. *J. Mater. Chem. A* **2017**, *5*, 4616–4628. [CrossRef]

43. Yuan, S.; Stroble, D.; Li, X.; Kruth, J.-P.; Van Puyvelde, P.; Van der Bruggen, B. 3d printed chemically and mechanically robust membrane by selective laser sintering for separation of oil/water and immiscible organic mixtures. *Chem. Eng. J.* **2020**, *385*, 123816. [CrossRef]

44. Chowdhury, M.R.; Steffes, J.; Huey, B.D.; McCutcheon, J.R. 3d printed polyamide membranes for desalination. *Science 2018*, **361**, 682–686. [CrossRef] [PubMed]

45. Chai, G.-Y.; Krantz, W.B. Formation and characterization of polyamide membranes via interfacial polymerization. *J. Membr. Sci.* **1994**, *93*, 175–192. [CrossRef]

46. Sridhar, S.; Smitha, B.; Mayor, S.; Prathab, B.; Aminabhavi, T.M. Gas permeation properties of polyamide membrane prepared by interfacial polymerization. *J. Mater. Sci.* **2007**, *42*, 9392–9401. [CrossRef]
47. Kammakakam, I.; O’Harra, K.E.; Bara, J.E.; Jackson, E.M. Design and synthesis of imidazolium-mediated 
tröger’s base-containing ionene polymers for advanced co2 separation membranes. ACS Omega 2019, 4, 
3439–3448. [CrossRef]
48. Bara, J.E.; O’Harra, K.E. Recent advances in the design of ionenes: Toward convergence with high-performance 
polymers. Macromol. Chem. Phys. 2019, 220, 1900078. [CrossRef]
49. Matthews, R.P.; Welton, T.; Hunt, P.A. Hydrogen bonding and pi-pi interactions in imidazolium-chloride 
ionic liquid clusters. Phys. Chem. Chem. Phys. 2015, 17, 14437–14453. [CrossRef]
50. Li, P.; Paul, D.R.; Chung, T.-S. High performance membranes based on ionic liquid polymers for co2 
separation from the flue gas. Green Chem. 2012, 14, 1052–1063. [CrossRef]
51. Kammakakam, I.; Kim, H.W.; Nam, S.; Park, H.B.; Kim, T.-H. Alkyl imidazolium-functionalized cardo-based 
poly(ether ketone)s as novel polymer membranes for o2/n2 and co2/n2 separations. Polymer 2013, 54, 
3534–3541. [CrossRef]
52. Gómez-Valdemoro, A.; San-José, N.; García, F.C.; De La Peña, J.L.; Serna, F.; García, J.M. Novel 
aromatic polyamides with main chain and pendant 1,2,4-triazole moieties and their application to the 
the extraction/elimination of mercury cations from aqueous media. Polym. Chem. 2010, 1, 1291–1301. 
[CrossRef]
53. Halasa, A.F.; Wathen, G.D.; Hsu, W.L.; Matrana, B.A.; Massie, J.M. Relationship between interchain spacing of 
amorphous polymers and blend miscibility as determined by wide-angle x-ray scattering. J. Appl. Polym. Sci. 
1991, 43, 183–190. [CrossRef]

© 2020 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access 
article distributed under the terms and conditions of the Creative Commons Attribution 
(CC BY) license (http://creativecommons.org/licenses/by/4.0/).