Physical Aging, Plasticization and Their Effects on Gas Permeation in “Rigid” Polymers of Intrinsic Microporosity

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Supporting Information

ABSTRACT: Long-term physical aging and plasticization, two mobility-based phenomena that are counterintuitive in the context of “rigid” polymers of intrinsic microporosity (PIMs), were evaluated using pure- and mixed-gas permeation data for representative ladder and semiladder PIMs. PIMs between 1 and 4 years old retained from 10- to 1000-fold higher H2 and O2 permeabilities than commercial membrane materials with similar or higher selectivities. A triptycene-based ladder polymer (TPIM-1) exhibited very large selectivity gains outweighing permeability losses after 780 days, resulting in unprecedented performance for O2/N2 (P(O2) = 61 Barrer, α(O2/N2) = 8.6) and H2/N2 (P(H2) = 1105 Barrer, α(H2/N2) = 156) separations. Interestingly, TPIM-1 aged more and faster than its more flexible counterpart, PIM-1, which exhibited P(O2) = 517 Barrer and α(O2/N2) = 5.0 at 1380 days. Additionally, the more “rigid” TPIM-1 plasticized more significantly than PIM-1 (i.e., TPIM-1 endured ~93% increases in mixed-gas CH4 permeability over pure-gas values compared to ~60% for PIM-1). A flexible 9,10-bridgehead (i.e., TPIM-2) mitigated the enhancements induced by physical aging but reduced plasticization. Importantly, intra-chain rigidity alone, without consideration of chain architecture and ultra-microporosity, is insufficient for designing aging- and plasticization-resistant gas separation membranes with high permeability and high selectivity.

INTRODUCTION

Polymer membrane technology constitutes a potentially cost-effective and less energy-intensive alternative to conventional gas separation technologies such as distillation and absorption.1,2 The criteria for deployment of a polymer into the industrial membrane arena are rigorous.3,4 Newly developed polymers should: (i) Demonstrate combinations of permeability and selectivity appropriate for the target separation, (ii) show performance stability over time (typically 3–5 years), (iii) be solution processable, and (iv) be manufacturable into mechanically robust asymmetric or composite membranes with thin and predominantly defect-free selective layers. The difficulty in satisfying all these criteria is reflected in a domination of 90% of the market by less than ten, mostly commercially available, polymers.3

It has been suggested for more than 20 years that polymer structures striking a balance between intrachain mobility and interchain spacing would offer superior gas separation membranes potentially defying state-of-the-art permeability/selectivity trade-offs.5–7 Recently developed amorphous and solution-processable polymers of intrinsic microporosity (PIMs), which trap microporosity (IUPAC, <2 nm “pores”) simply by virtue of torsion-resistant and contorted fused-ring backbones that pack inefficiently in the solid state8–25 may be one of the most promising classes of materials providing unprecedented gas separation performance. Specifically, previous reports showed that the integration of bridged-bicyclic contortion centers (i.e., ethanoanthracene, Tröger’s base, triptycene) into fully fused-ring ladder polymers26–29 and semiladder PIM-polymides (PIPs)30,31 resulted in highly ultramicroporous (<7 Å) PIMs with high selectivities matching those of commercial materials (e.g., polycarbonate, cellulose acetate, Matrimid) but with up to 3 orders of magnitude higher permeabilities, especially in the large-scale industrial applications of nitrogen production from air and hydrogen purification.3,4

One major obstacle to the commercialization of glassy amorphous PIMs for membrane applications is physical aging and the ensuing instability of their permeation properties over time.1,2,27,28 Physical aging in glassy polymers involves a reversible densification driven by a dissipation of nonequilibrium free volume toward an equilibrium state of polymer chain packing. Struik has thermodynamically described the driving force for physical aging—and hence its extent and rate—as being proportional to the difference between the current specific volume (v) and the specific volume at equilibrium (v∞) (i.e., dv/dt ∝ (v – v∞)).28 Though still not fully understood, it is thought to occur via molecular reorganizations depending on chain mobility.27 For PIMs, this is rather counterintuitive in the context of (i) their “rigid”
chains, (ii) the lack of a measurable $T_g$, and (iii) the concept that they possess open “pores” with relative permanency on the time-scale of gas transport, unlike traditional solution-diffusion membranes in which transport occurs through transient pores forming via thermally activated chain motions. However, a body of literature suggests that neither intrachain rigidity (e.g., central to PIM design) nor interchain rigidity (e.g., covalently cross-linked polyimides) have successfully eluded the microstructural rearrangements of physical aging, and that instead the polymers with higher intra-chain rigidity aged more and faster. In the broadest rationalization of physical aging, the amount of free volume governs the rate and extent of aging. As in physical aging, plasticization is based upon molecular chain reorganization of a polymer induced by high sorption of a condensable gas. The resulting selectivity reductions increase product CH4-loss into a low-pressure permeate and reduce the membrane system’s competitiveness against standard amine absorption columns. There have been expectations that state-of-the-art PIMs boasting high intrachain rigidities could demonstrate resilience against plasticization. To date, unlike for physical aging, plasticization has successfully been mitigated with polymers engaging in interchain interactions via (i) high-temperature chemical cross-linking, thermal rearrangement, and carbonization, (ii) sub-$T_g$ annealing, and/or (iii) the introduction of hydrogen-bonding moieties. Several works have recently reached similar conclusions with ladder PIMs and PIM-type polyimides (PIM–PIs).

The main focus of this work was to investigate how the relatively extreme intra-chain rigidity and high ultramicroporosity in some of today’s most permeable and selective PIMs can potentially mitigate long-term physical aging and plasticization, two mobility-based phenomena that are counterintuitive in the context of the “rigid” backbones characteristic of PIMs. To address this matter, two sets of permeation data are presented for films of representative ladder PIMs, including the prototypical PIM-1 and semiladder PIM–PIs: (i) long-term

### Table 1. Structure, BET Surface Areas, Thicknesses and Casting Solvents for Representative PIM Films.

<table>
<thead>
<tr>
<th>Polymer</th>
<th>Structure</th>
<th>BET S.A. (N$_2$) (m$^2$ g$^{-1}$)</th>
<th>Thickness (μm)</th>
<th>Solvent</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>PIM-1</td>
<td><img src="image" alt="Structure" /></td>
<td>770</td>
<td>102</td>
<td>CHCl$_3$</td>
<td>This work</td>
</tr>
<tr>
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<td>860</td>
<td>103</td>
<td>CHCl$_3$</td>
<td>This work</td>
</tr>
<tr>
<td>TPIM-2</td>
<td><img src="image" alt="Structure" /></td>
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<td>46</td>
<td>CHCl$_3$</td>
<td>This work</td>
</tr>
<tr>
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<td>180</td>
<td>CHCl$_3$</td>
<td>18, 19</td>
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<tr>
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<td>132</td>
<td>CHCl$_3$</td>
<td>19</td>
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<tr>
<td>KAUST-PI-1</td>
<td><img src="image" alt="Structure" /></td>
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<td>77; 102</td>
<td>CHCl$_3$</td>
<td>This work</td>
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<tr>
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<tr>
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<td>123</td>
<td>CHCl$_3$</td>
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</table>
physical aging data gauged by pure-gas permeabilities and (ii) high-pressure pure- and mixed-gas CO$_2$/CH$_4$ separation properties. The permeation data are discussed in terms of relative changes and absolute separation performance to gauge the effects of physical aging and plasticization on the PIMs.

![Figure 1](https://example.com/figure1.png)

**Figure 1.** (a) Repeat unit chemical structures, (b) energy-minimized chain conformations (Materials Studio, Accelrys, 7.0), and (c) the degrees of torsional freedom at 35 °C in one repeat unit for several representative PIMs. Torsions were calculated about the color-coded, highlighted bonds in the repeat units.

### EXPERIMENTAL SECTION

**Polymers.** Table 1 is a tabulation of names, chemical structures, BET surface areas and the film thicknesses and casting solvents used for some representative ladder PIMs and semiladder PIM–PIs reported in the literature. This work presents new data for thick films of ladder polymers PIM-1, TPIM-1 and TPIM-2 as well as for semiladder PIM–PIs KAUST-PI-2 and KAUST-PI-7, which are shown in comparison to published data for the other polymers.

**Polymer Film Preparation.** Solutions containing 3 wt % polymer were syringe-filtered with 0.45 μm PTFE filters and cast onto clean, flat-bottomed glass dishes. Clear, isotropic polymer films were obtained by covering the dishes to induce slow evaporation of the casting solvent. The resulting films were dried at 120 °C under vacuum for 12 h, soaked in methanol for 24 h to “reset” their thermal histories and exchange any residual casting solvent, and finally dried at 120 °C under vacuum for 24 h. Removal of the casting solvent was confirmed with thermal gravimetric analysis (TGA, TA Q5000). Effective areas and thicknesses of the films were determined by calibrated scanning software and a digital micrometer, respectively. All polymers in Table 1 were treated this way except for PIM-EA-TB, PIM-Trip-TB and PIM-PI-EA, which were air-dried after the methanol soak. It is important to note that previous work has demonstrated thickness-dependent physical aging properties for PIMs.

**Molecular Simulations.** The repeat units of PIM-1, TPIM-1, PIM-EA-TB, and KAUST-PI-1 were constructed in Materials Studio (Accelrys, 7.0). The conformer module was used to incrementally vary select dihedral angles (bold and color-coded in Figure 1) from −180° to +180°. For each conformer, geometry optimizations were performed with the Forcite module using a COMPASS force field and Smart algorithm (fine quality and 50 000 iterations). The energy barriers (kJ/mol) to changes in the dihedral angles were calculated relative to the lowest energy attained (ΔE = Ei − Emin) over the 360° range. The degrees of torsional freedom were then defined as the total number of degrees available for torsion about the color-coded bonds at a temperature T, where the available energy at temperature T was approximated by the equipartition theorem (~3RT). Isolated polymer chains (usually of 10–20 repeat units) were constructed from the repeat units using the polymer builder function. Initially, the chains were constructed with random torsions between the repeat units. The Forcite module (COMPASS force field, Smart algorithm with fine quality and 50 000 iterations) was then used to obtain the energy-minimized chain conformation of each chain in isolation.

**Pure-Gas Permeation Measurements.** The pure-gas permeabilities of the films were determined using the constant-volume/variable-pressure method. After drying the methanol-soaked films at 120 °C for 24 h under vacuum, they were immediately masked between two aluminum-tape donuts. On one side, the interface between the film and tape was lightly epoxied using a two-part potting epoxy (GC Electronics) which showed excellent durability over long times. The masked film was degassed in the permeation test apparatus on both sides under high vacuum at 35 °C. Pure gases (grade 5 and above) were fed to the epoxied side at 35 °C and 2 bar. In addition, the CO$_2$ and CH$_4$ pure-gas permeabilities were measured at pressures from 2 to 15 bar for TPIM-1 and TPIM-2. The increase in permeate pressure with time was measured by an MKS Baratron transducer. The pure-gas permeabilities were calculated by

\[
P = D S = 10^{10} \frac{V_{d} \cdot dp}{P_{up} \cdot ART \cdot dt}
\]

where P is the gas permeability (Barrer) ($1 \text{ Barrer} = 10^{-10} \text{ cm}^3 \text{(STP)} \text{ cm} \text{ cm}^2 \text{ s}^{-1} \text{ cmHg}^{-1}$), $P_{up}$ is the upstream pressure (cmHg), $dp/dt$ is the steady-state permeate-side pressure increase (cmHg s$^{-1}$), $V_{d}$ is the calibrated permeate volume (cm$^3$), I is the membrane thickness (cm), A is the effective membrane area (cm$^2$), T is the operating temperature (K), and R is the gas constant (0.278 cm$^3$ cmHg cm$^{-3}$ (STP) K$^{-1}$). The ideal permeability ($\alpha_{A/B}$) for separation of A/B was calculated by

\[
\alpha_{A/B} = \frac{P_A}{P_B}
\]

**Mixed-Gas Permeation Measurements.** The mixed-gas permeation properties for ~15-day-old films were measured at 35 °C with a 50-50 CO$_2$/CH$_4$ binary mixture using a setup similar to that previously described by O’Brien et al. The films were evaluated at total pressures ranging from 4 to 30 bar such that the mixed-gas CO$_2$ partial pressures were equal to those used in the pure CO$_2$ permeation experiments. This methodology permits direct comparison of the pure- and mixed-gas data to elucidate any nonidealities including plasticization and competitive sorption. The stage-cut, that is, the ratio of permeate to feed flow rates, was less than 0.01 to ensure that the residue composition was essentially equal to that of the feed gas. CO$_2$ and CH$_4$ concentrations were measured with a calibrated gas chromatograph (Agilent 3000A Micro GC) equipped with thermal conductivity detectors. The mixed-gas permeability of component i was determined by
Macromolecules

\[ P_i = 10^{10} \frac{\gamma_{Vf}^i}{x_{p}^{\text{ART}}} \frac{dp}{dt} \]

where \( y \) and \( x \) are the mole fractions in the permeate and feed, respectively, and \( p_u \) is the upstream partial pressure of component \( i \). Because the downstream pressure was negligible, the mixed-gas selectivity of component \( i/j \) was obtained from

\[ \alpha_{i/j} = \frac{y_i/y_j}{x_i/x_j} \]

Steady-state permeation was identified at each feed pressure when regular measurements of permeability and selectivity both changed negligibly with time.

## RESULTS AND DISCUSSION

### Molecular Simulations: Intra-Chain “Rigidity” and Chain Architecture

Polymers of intrinsic microporosity are often recognized for their high “rigidity,” a rather general term which is more appropriately referred to here as “intra-chain” rigidity—describing specifically short-range motions including, for example, local torsions within the backbone—to distinguish from “inter-chain” rigidity. Figure 1 qualitatively investigates the intrachain rigidities and chain architectures for several representative polymers as determined via molecular simulation (Materials Studio, Accelrys, 7.0). As expected, the ladder polymers (e.g., PIM-1, TPIM-1, PIM-EA-TB) exhibit considerable reductions in backbone flexibility (i.e., degrees of torsional freedom) over the semiladder polymide KAUST-PI-1, primarily due to the presence of a torsionable imide-linkage in the latter. Moreover, PIM-EA-TB, a completely fused-ring polymer lacks any flexible xylene or phenazyl moieties present in the other polymers, shows the highest intrachain rigidity (i.e., lowest number of degrees of torsional freedom). In addition, Figure 1 highlights key differences in the chain architectures of the PIMs (Figure 1b). Most interestingly, TPIM-1 possesses a unique chain conformation that resembles a two-dimensional ribbon-like propagation through space (Figure 1b). This is in stark contrast to the highly contorted and erratic structure of the spiro-centered PIM-1 and even the structures of Tröger’s-base-containing ladder polymers such as PIM-EA-TB, all which exhibit a three-dimensional propagation. KAUST-PI-1 chains, as for other polyimides, also propagate in three dimensions owing to a larger amount of torsional freedom about the N-phenyl imide bonds (Figure 1c). These observations will be addressed in discussion of the transport properties below.

### Pure-Gas Permeation Properties: Long-Term Physical Aging

Long-term physical aging of the polymers presented in Table 1 was characterized with pure-gas permeabilities and summarized in Table 2. Details of testing and storage conditions are provided in the footnotes to Table 2. All the polymers were soaked in methanol to “reset” their thermal histories and remove residual casting solvent. The main differences in the handling of the films were as follows: (i) Data for PIM-EA-TB, PIM-Trip-TB, and PIM–PE–IA were previously reported and collected at 25 °C after ambient drying of the methanol-soaked films; (ii) the remainder of the polymers were examined in this work at 35 °C after methanol soaking and then drying for 24 h at 120 °C under vacuum.

On the basis of Table 2, the overall losses in O₂ permeability, P(O₂), and gains in O₂/N₂ selectivity, α(O₂/N₂), over the times investigated are expressed percentage wise in Figure 2. The same is done for H₂ permeability and H₂/N₂ selectivity in Figure 3.

### Table 2. Long-Term Physical Aging of PIMs Tracked by Pure-Gas Permeabilities

<table>
<thead>
<tr>
<th>polymer</th>
<th>age (days)</th>
<th>H₂</th>
<th>O₂</th>
<th>N₂</th>
<th>H₂/N₂</th>
<th>O₂/N₂</th>
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</thead>
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<tr>
<td>PIM-1</td>
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<td>3623</td>
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<td>308</td>
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<td></td>
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<td>64</td>
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<td></td>
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<td>780</td>
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<td></td>
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<td>737</td>
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<td>850</td>
<td>226</td>
<td>13</td>
<td>3.8</td>
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*Film immersed in methanol, dried at 120 °C in vacuo for 24 h.
Storage: days 1–15, in vacuo, 35 °C; days 15+, in a sealed plastic bag at ambient conditions. Permeation properties measured at 35 °C and 2 bar.
*Film immersed in methanol, dried at 120 °C in vacuo for 24 h.

Figure 2. Effects of long-term physical aging on O₂ permeability, P(O₂), and O₂/N₂ selectivity for various PIMs. The number of days aged is shown in parentheses.
than oxygen ($k_0 = 3.46 \text{ Å}$) due to its smaller size, and thus oxygen permeability dropped more with time than hydrogen permeability for any polymer. The gains in $H_2/N_2$ selectivity thus outweighed the losses in $H_2$ permeability (Figure 3), as expected. It is important to note that permeability loss in PIMs appears to proceed in two phases. The first phase is documented and characterized by a rapid permeability decline over $\sim 15$ days and driven by excess nonequilibrium free volume created by methanol soaks. At this point, previously termed an “aging-knee,” a slower, more gradual permeability decline occurs that is attributed to a reorganization of the “intrinsic” microporosity. The latter phase is slow due to the restricted mobilities of PIM chains deep in their glassy states (i.e., often no observable $T_g$). However, as indicated by extents of permeability loss and selectivity gain over long times (Table 2 and Figures 2 and 3), significant redistributions in free volume occur in this latter phase. The resulting densification of the microstructure during physical aging reduces both gas diffusivity and solubility coefficients, thereby reducing permeability, but the overall process is diffusion-governed as indicated by the larger changes in both diffusivity and diffusivity selectivity (Table S1, Supporting Information).

There are two particularly interesting observations to be made from Figures 2 and 3, namely (i) an enhanced aging response for TPIM-1 relative to the other PIMs and (ii) a significant effect of the 9,10-triptycene-bridgehead substituent on aging. The first observation is apparent in a comparison of TPIM-1 and PIM-1 films of nearly identical thickness ($\sim 100 \mu m$) and treatment history. TPIM-1 aged more extensively and faster than PIM-1. PIM-1 lost $\sim 70\%$ in $O_2$ permeability and gained $\sim 40\%$ in $O_2/N_2$ selectivity over 1380 days. In comparison, TPIM-1 lost $\sim 95\%$ in $O_2$ permeability and gained $\sim 115\%$ in $O_2/N_2$ selectivity over a shorter period of 780 days. Additionally, $H_2/N_2$ selectivity increased more than 10-fold for TPIM-1, compared to 2-fold for PIM-1. This trend is consistent with prior literature discussed above demonstrating intrachain rigidity to be insufficient in the mitigation of physical aging, and free volume to be the main driving force: That is, TPIM-1 has a more rigid backbone (Figure 1) than PIM-1 and initially higher free volume, the latter indicated by higher $N_2$--BET surface areas (860 vs 780 m$^2$ g$^{-1}$) and higher $O_2$ permeabilities (1200 vs $\sim 1000$ Barrer) (Table 2).

The second observation, regarding a significant effect of the 9,10-triptycene-bridgehead substituent on the redistribution of free volume during aging, is apparent in a comparison of TPIM-1 (isopropyl bridgeheads) and TPIM-2 (propyl bridgeheads) aged for $\sim 750$ days. TPIM-1 and TPIM-2 both lost $\sim 95\%$ in $O_2$ permeability, but TPIM-1 gained $\sim 60\%$ more $O_2/N_2$ selectivity. On the other hand, both lost $\sim 70\%$ in $H_2$ permeability and acquired similar 9- to 10-fold gains in $H_2/N_2$ selectivity. These two observations collectively indicate that physical aging in the isopropyl-substituted TPIM-1 significantly affected a narrow range of ultramicroporosity between 3.46 Å (i.e., $O_2$ kinetic diameter) and 3.64 Å (i.e., $N_2$ kinetic diameter). This is further corroborated by the $\sim 2$-fold higher $O_2$ diffusivity and $30\%$ higher $O_2/N_2$ diffusivity selectivity in TPIM-1 compared to TPIM-2 (Table S1, Supporting Information). The effect was not as specific in PIM-1 and TPIM-2 where aging more greatly impeded the transport of oxygen, hence the differences in selectivities. The profound role of triptycene’s 9,10-isopropyl-bridgehead on gas sorption in covalent networks and gas separation in amorphous polymer films has been previously noted, and here it is established as a critical factor in enhancing the molecular sieving function of TPIM-1 during physical aging.

Mechanistically rationalizing the relatively enhanced aging responses of TPIM-1 and TPIM-2 is not trivial. Without overinterpreting the data, the TPIMs trend—more than other similarly aged polymers—toward the upper right corners of Figures 2 and 3, where permeability losses and selectivity gains are quite extensive. Despite only being air-dried after a methanol soak (which should increase the apparent effects of aging), PIM-EA-TB, which has a notably higher chain rigidity (Figure 1), higher free volume due to higher $N_2$--BET surface areas (1000 vs 860 m$^2$ g$^{-1}$) and nearly 2-fold higher initial $O_2$ permeability (Table 2), does not demonstrate such a drastic aging behavior even by 470 days. Similar behavior was observed for a 660-day-old KAUST-PI-7 and 870-day-old KAUST-PI-2. We hypothesize that chain architecture and conformation also play a role in physical aging, potentially by altering the target $v_\infty$ in the context of Struik’s model. That is, the unique ribbon-like two-dimensional geometry of energy-minimized TPIM chains (Figure 1) may be more amenable to efficient packing than the more contorted and three-dimensional chain propagations of PIM-EA-TB, PIM-1, etc. That is, a lower $v_\infty$ translates into an increased driving force for aging toward a more efficiently packed equilibrium structure. Along the same lines, what preliminarily appears to be a “mitigated” aging in PIM-EA-TB may be the result of the high concentration of bulky bridged-bicyclic moieties in a fused ring structure that contort the macromolecule in three dimensions and simply increase the $v_\infty$. The impact of physical aging on the absolute separation performance of the PIMs is shown in Figures 5 and 6 relative to the 2008 upper bounds for $O_2/N_2$ and $H_2/N_2$ separations, respectively. The smaller symbols under the gray shade represent data taken initially for the “fresh” methanol-treated PIMs (details in footnotes to Table 1). For both separations, the ensuing increases in selectivity and concomitant losses in permeability locate the PIMs well above the 2008 upper bounds to polymer separation performance. Favorably, however, the performance generally transitions from being primarily permeability driven to having a more favorable, balanced contribution of permeability and selectivity. This balance is exemplified by TPIM-1, which exhibits an unprecedented combination of $O_2$ permeability of 61 Barrer and $O_2/N_2$ selectivity of 8.6 at 780 days. Relative to a commercially employed air separation material, tetrabromo-poly carbonate...
(TB-PC) \( P(O_2) = 1.4 \text{ Barrer}, \alpha(O_2/N_2) = 7.8 \), \(^{61}\) TPIM-1 has higher \( O_2/N_2 \) selectivity along with \( \sim45 \)-fold higher \( O_2 \) permeability. Given the current membrane infrastructure and employed membranes and a permeability that is orders of magnitude higher. \(^{62}\) Accordingly, \textit{intra}-chain rigidity alone—which is a main target of standard PIM design—is insufficient in the design of highly permeable \textit{and} highly selective gas separation membranes with PIMs.

**Pressure Dependence of Pure- and Mixed-gas \( CO_2/CH_4 \) Permeation Properties: Plasticization.** To date, very few reports exist on the plasticization behavior of ladder PIMs\(^{49}\) and semiladder PIM–PIs\(^{13,14,51}\) under \( CO_2 \)-containing mixed-gas feeds. These studies indicated that structural changes increasing interchain rigidity (i.e., between chains) were required to improve plasticization resistance. This was established either via the introduction of (i) hydrogen bonding moieties, (ii) intrachain flexibility that facilitates coplanarization over the \( N \)-phenyl imide bonds of polyimides and thus the formation of interchain charge-transfer-complexes (CTCs), and (iii) a combination thereof. Moreover, it has been recommended that polymers having lower free volume—typically induced by the aforementioned changes—are better choices for \( CO_2/CH_4 \) separation because they sorb less of the plasticizing \( CO_2 \) gas.\(^{63}\)

It remains interesting, however, to investigate the plasticization behavior of TPIM-1, which possesses a significantly higher intrachain rigidity and molecular sieving character than previously reported PIM-1 and KAUST-PI-1 (Figure 1). Given the large impacts on gas permeation properties of the 9,10-bridgehead substituent on triptycene, as shown above and elsewhere,\(^{12}\) TPIM-2 was also evaluated. Pure- and mixed-gas (1:1 feed) permeability isotherms spanning (partial) \( CO_2 \) pressures of 15 bar are presented for \( CO_2 \) and \( CH_4 \) in Figures 6 and 7, respectively. In general, lower \( CO_2 \) permeabilities under the mixed-gas than the pure-gas conditions are expected due to competitive sorption with \( CH_4 \). The \( CO_2 \) permeabilities in PIM-1 exhibited pressure-dependent drops due to dual-mode reductions in the solubility coefficient of \( CO_2 \) with pressure, whereas those in the TPIMs were nearly constant or only slightly increased with pressure, preliminarily indicating a small degree of plasticization. However, the only unambiguous indicator of plasticization is the mixed-gas permeability of the slower \( CH_4 \) gas (Figure 6). At 10 bar \( CO_2 \) partial pressure, TPIM-1 and PIM-1 endured 93% and 62% increases in mixed-gas \( CH_4 \) permeability over the pure-gas permeability. TPIM-1 plasticized appreciably more than PIM-1, indicating that greatly
increased intrachain rigidity does not inhibit CO₂-induced plasticization, which remains consistent with the literature. Accordingly, the CO₂/CH₄ mixed-gas selectivity in TPIM-1 dropped ~60% in the mixed-gas from ~30 at 2 bar to ~13 at 10 bar CO₂ partial pressure (Figure 8).

![Figure 8. Pressure-dependence of CO₂/CH₄ selectivity isotherms for PIM-1, TPIM-1, and TPIM-2 at 35 °C using a 1:1 CO₂:CH₄ binary feed. Open points, pure gas. Closed points, mixed gas.](image)

It is worth noting the effect of the bridgehead on TPIM plasticization. TPIM-2 plasticized significantly less than TPIM-1 given less than 10% increases in CH₄ mixed-gas permeability uniquely increases in di-plasticization, which remains consistent with the literature. The same hypothesis is corroborated here and described schematically in Figure 9. Briefly, given the combination of high O₂/N₂ selectivity and high O₂ permeability in TPIM-1, a significant portion of its ultramicroporosity is likely dimensioned around O₂ and N₂ at 3–4 Å. This is additionally supported by the unique increases in diffusivity-governed (Table S1) O₂/N₂ sieving behavior observed during physical aging (e.g., time-dependence of pure-gas permeabilities). The envisioned dilation of these 3–4 Å pores during pressure-dependent CO₂/CH₄ testing would also impact CO₂ permeability, since CO₂ has an effective size (based on diffusivity, ∼3.5–3.6 Å) between that of O₂ (3.46 Å) and N₂ (3.64 Å). This is indicated by the strong rises in CO₂ permeability isotherms (Figure 6 and 7) with pressure for TPIM-1, but not for PIM-1 or TPIM-2 (i.e., in which O₂/N₂ transport data indicate such pores do not contribute as significantly to gas transport as in TPIM-1). Furthermore, an even larger effect is envisioned on the diffusion of the larger CH₄ (kᵣ = 3.8 Å) gas especially in a highly ultramicroporous polymer sieve that contains more of the 3.4–3.6 Å pores (Figure 9).

Finally, the CO₂/CH₄ separation performance of notable PIMs is presented in Figure 10 on a permeability/selectivity performance map. Data from 2 bar pure-gas measurements (open points) are compared against data from 20 bar 1:1 mixed-gas measurements (closed points) to emphasize the impact of nonideal phenomena including plasticization and competitive sorption on performance. Recently, Baker has defined a mixed-gas selectivity of 40 as critical for membrane technology to outcompete standard amine absorption columns. The solid line represents the 2008 upper bound. The dashed line represents a practically relevant “selectivity” of 40. Open points, pure-gas feeds at 2 bar, 35 °C. Closed points, mixed-gas feeds at 20 bar, 35 °C.

![Figure 10. Effects of CO₂-induced plasticization on pure- and mixed-gas CO₂/CH₄ separation performance for PIMs. The solid line represents the 2008 upper bound. The dashed line represents a practically relevant “selectivity” of 40. Open points, pure-gas feeds at 2 bar, 35 °C. Closed points, mixed-gas feeds at 20 bar, 35 °C.](image)

**CONCLUSIONS**

The results reported herein provide a critical perspective on the effects of physical aging and plasticization on a rapidly emerging class of promising polymers, PIMs. Apparently, the very intrachain rigidity that offers PIMs their attractive high-free-volume, ultramicroporous gas sieving microstructures also increases their vulnerability to physical aging and plasticization, both viewed as obstacles for commercialization. However, physical aging induces a favorable transition from permeability-driven gas transport properties to balanced combinations of permeability and selectivity that are outstanding relevant to both the “upper-bound” metric and the performance of...
commercially employed polymers. The major conclusions of the work are summarized as follows:

1 Physical aging occurs in all PIMs despite their high intrachain rigidities. In fact, the initially high free volume created by greater backbone rigidity serves as a driving force for more extensive and rapid physical aging of the bulk microstructure.

2 The inherent chain architecture may be an important consideration in physical aging. Simulations indicate that TPIM macromolecules propagate through space in two-dimensional (i.e., in one plane) ribbon-like conformations, starkly different from the more erratic conformations PIMs usually assume. TPIMs may be able to reach a denser equilibrium chain packing, which would be consistent with the significantly improved molecular sieving behaviors of aged TPIMs.

3 Higher intrachain rigidity, central to PIM design and exemplified in the TPIMs studied herein, does not alone prevent plasticization. On the contrary, a highly O₂/N₂ sieving ultramicroporous structure—to which intrachain rigidity is key—appears to be more vulnerable to CO₂-induced plasticization (i.e., as gauged by increases in mixed-gas CH₄ permeabilities under 1:1 CO₂/CH₄ feeds).

Work is currently underway to develop PIM-like structures that are more resilient against physical aging and plasticization. For cases where aging is unavoidable, techniques to accelerate it and improve the stability of the permeation properties with time are needed.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.macromol.5b01581.

Table of long-term aging data and figures showing the effects of long-term physical aging (PDF)

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Notes

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