Ultrapermeable Polymers of Intrinsic Microporosity (PIMs) Containing Spirocyclic Units with Fused Triptycenes

C. Grazia Bezzu,* Alessio Fuoco, Elisa Esposito, Marcello Monteleone, Mariagiulia Longo, Johannes Carolus Jansen,* Gary S. Nichol, and Neil B. McKeown*

1. Introduction

Gas separation by polymeric membranes is an energy efficient process used to effectively separate industrially and commercially important gas mixtures, such as N₂ or O₂ enrichment of air, CO₂ from CH₄ for sweetening of natural gas, or H₂ from N₂ for hydrogen recovery during ammonia preparation. In addition, their potential utilisation for the separation of CO₂ from N₂ for post-combustion carbon-capture is being considered, with promising results from pilot-plant scale demonstration studies. For gas separations on such a massive scale, good selectivity and high permeability are required to enhance both the process efficiency and productivity, by reducing the effective size and cost of the membrane system. However, polymeric membranes show a well-known empirical trade-off between permeability (P) and selectivity (α, P_b/P_a) for a given gas pair (a and b), with highly permeable polymers possessing low selectivity and vice versa. The upper bound of this inverse relationship in plots of log(P_a) versus log(P_b/P_a), representing the best performing polymeric membranes at the time for the separation of several important gas pairs, was defined for the first time in 1991 and then reviewed in 2008 by Robeson. Although these plots are based on single gas permeation data, the position of the gas permeability data for a newly prepared polymeric membrane relative to the 1991 and 2008 upper bounds is used to estimate its potential for gas separations. By a theoretical analysis of the Robeson upper bound, Freeman suggested that its slope results directly from the size-sieving nature of the highly rigid chain of the polymers used to define it, hence increasing both chain rigidity (i.e., diffusivity selectivity) and interchain distance would improve polymer gas permseleciivity. Polymers of Intrinsic Microporosity (PIMs) with their contorted and rigid structures, which inhibit efficient packing, adhere closely to this design concept. Indeed, permeability data for PIM-1 (Figure 1) and PIM-7 were used to revise the Robeson upper bound for most gas pairs in 2008. Recently, many PIMs with greater chain rigidity have shown improved permeability and selectivity properties with data exceeding the 2008 upper bounds. Their enhanced rigidity was achieved by replacing the relatively flexible spirobisindane (SBI) unit in PIM-1 with spirobifluorene (PIM-SBF-1; Figure 1) or by using stiff bridged bicyclic monomers such as Trögers base, ethanoanthracene, methanopentacene or triptycene (Trip). In particular, polymers based on the triptycene unit showed remarkable permselectivity so that Pinnau et al. proposed in 2015 to...
update the O₂/N₂, H₂/N₂, and H₂/CH₄ upper bounds using permeability data from aged films of highly selective triptycenc-based PIMs (e.g., PIM-Trip-TB[9b] and TPIM-1[12a]). Recently, we proposed the 2019 revision of the upper bounds for CO₂/N₂ and CO₂/CH₄,[14] using data from the ultrapermeable and selective 2D ribbon-shaped benzotriptycene polymers (e.g., PIM-BTrip, PIM-TMN-Trip,[15] and PIM-TFM-BTrip).

Herein we report the synthesis and properties of PIM-SBI-Trip and the copolymer PIM-1/SBI-Trip (Figure 1), derived from a novel structural unit consisting of a spirobisindane to which two rigid and bulky triptycenes have been fused, which was designed to enhance the rigidity of the spirocyclic site of contortion and greatly increase the distance between polymer chains.[16] In addition, we investigate in detail the mechanism of gas transport through the resulting PIMs under a wide range of operating conditions, with pure gases as a function of temperature and with CO₂/N₂ and CO₂/CH₄ gas mixtures as a function of pressure. This allows their potential application in important fields such as carbon capture from flue gas or biogas upgrading to be assessed. In addition, a detailed analysis of the temperature dependence of the permeability and diffusion provides fundamental insight into the energetic and entropic contribution to the high size-selectivity of the membranes.

2. Results and Discussion

2.1. Monomer and Polymer Synthesis

The newly designed monomer, 6,6′,7,7′-(9,10-dihydroanthracene-9,10-diyl)-2,2′,3,3′-tetrahydroxy-9,9′-spiro-bifluorene (SBI-Trip), was prepared by adapting the well-established route to spirobifluorenes[17] that is based on the addition of the biphenyl 2-lithio-anion to fluorenone followed by acid-mediated cyclisation (Figure 2).

The required triptycene-based 2-bromo-biphenyl precursor (2) was prepared by bromination of the corresponding biphenyl (1) which was obtained by Suzuki coupling of 3,4-dimethoxyphenylboronic acid with 2-bromotriptycene. Bromo derivative 2 was used to prepare the corresponding fluorenone derivative (4) via a one-pot, two-step procedure that involved treatment with n-butyl lithium and addition of CO₂ to give the corresponding carboxylic acid (3), followed by an acid mediated ring closure (i.e., intramolecular acylation). The lithio-anion derived from 2 was then combined with fluorenone 4 to give, after acid treatment, the tetramethoxy precursor 5. Demethylation with boron tribromide afforded the required bis-catechol monomer SBI-Trip 6. In addition to the...
usual spectroscopic methods (e.g., ESI), conformation of the structure of SBI-Trip 6 was obtained by XRD analysis using crystals grown by slow solvent evaporation from a diethyl ether solvent. The crystals belong to the monoclinic P21/c space group with each unit cell containing four SBI-Trip 6 molecules along with sixteen diethyl ether molecules, with which they interact via hydrogen bonding (Figure 3). Polymerisation was achieved using the now standard dioxin-forming aromatic nucleophilic substitution reaction between SBI-Trip and 2,3,5,6-tetrafluoroterephthalonitrile (TFTPN). The random copolymer, PIM-1/SBI-Trip, was prepared similarly by reacting an equimolar mixture of SBI-Trip 6 and commercially available 5,5″,6,6″-tetrahydroxy-3,3″,3‴-tetramethyl-1,1‴-spirobisindane (SBI) with TFTPN.

Both polymers proved soluble in chloroform, and were purified by recrystallisation from a non-solvent (acetone and methanol) to remove oligomeric impurities. Analysis by Gel Permeation Chromatography (GPC) of PIM-SBI-Trip and PIM-1/SBI-Trip confirmed their high molecular mass with Permeation Chromatography (GPC) of PIM-SBI-Trip and PIM-1/SBI-Trip respectively, which are within the upper range for solution processable PIMs,[9a,14,15] and greater than the values reported for PIM-1[8] and PIM-SBF-1.[8] Significantly, PIM-SBI-Trip possesses apparent microporosity that is similar to that of the best performing SBF-based PIMs.[19] This is in agreement with the predictions, based on molecular simulations, that larger substituents on rigid spiro-centre units both frustrate chain packing and increase interchain distances, resulting in enhanced porosity.[20] The shape of the N₂ isotherms are similar for both polymers and show a significant gas uptake at higher pressures, followed by a large hysteresis in the desorption isotherms. This can be attributed to the large triptycene substituents reducing cohesion between polymer chains, leading to swelling upon N₂ adsorption. Adsorption of CO₂ at 273 K (Figure 4b) shows similar uptakes for the two polymers (2.9 and 2.7 mmol g⁻¹, for PIM-SBI-Trip and PIM-1/SBI-Trip, respectively). Micropore size distributions calculated from CO₂ adsorption at 273 K using DFT (Figure 4c) are remarkably similar for the two polymers and show a large contribution from ultramicropores (<0.7 nm).

Films were prepared by solution casting, allowing slow solvent evaporation from 3% to 4% v/v polymers solutions in CHCl₃ and their Young’s moduli were investigated by AFM force spectroscopy measurements (Table 1 and Figure S11: Supporting Information). The modulus of the copolymer PIM-1/SBI-Trip falls between those of the homopolymers PIM-1 and PIM-SBI-Trip. Both polymers are much stiffer than PIM-1[21] but are not as stiff as the very rigid PIM-BTrip,[22] which does not contain spirocyclic units.

Freshly methanol treated self-standing films of PIM-SBI-Trip and PIM-1/SBI-Trip demonstrate ultrapermeability

![Figure 3. Single crystal XRD structure of monomer SBI-Trip 6. a) Face-on and b) edge-on molecular view. c) Molecular packing. The included diethyl ether solvent molecules have been removed for clarity. Color code C: grey, O: red, H: white. The unit cell is shown as a red box.](image)

Table 1. Properties of PIM-SBI-Trip and PIM-1/SBI-Trip.

| Polymer                  | PIM-SBI-Trip | PIM-1/SBI-Trip |
|--------------------------|--------------|----------------|
| Yield, %                 | 88           | 85             |
| Solubility               | CHCl₃        | CHCl₃          |
| $M_\text{w}$ (g mol⁻¹)   | 302 000⁶     | 225 000⁶       |
| $M_\text{w}$/$M_\text{a}$ | 2.8          | 2.1            |
| $S_{\text{BET}}$ (m² g⁻¹) | 930          | 858            |
| $V_{\text{tot}}$ (cm³ g⁻¹) | 0.72        | 0.60           |
| $V_{\text{micro}}$ (cm³ g⁻¹) | 0.33        | 0.31           |
| CO₂ uptake (mmol g⁻¹)    | 2.88         | 2.71           |
| Young’s modulus (MPa)    | 2004 (±53)   | 1790 (±83)     |

⁶BET surface area calculated from N₂ adsorption isotherm obtained at 77 K; ⁷Total pore volume estimated from N₂ uptake at $P/P_0 = 0.98$; ⁸Micropore volume estimated from N₂ uptake at $P/P_0 = 0.05$; ⁹CO₂ adsorption at 1 bar and 273 K; ¹⁰Measured by AFM force spectroscopy; ¹¹From recrystallised polymer relative to polystyrene standards.

2.2. Gas Adsorption and Gas Transport Properties

The powder form of both PIM-SBI-Trip and PIM-1/SBI-Trip adsorbs a significant amount of N₂ at low relative pressure ($P/P_0 < 0.05$) at 77 K, indicating significant microporosity. Analysis of the N₂ adsorption isotherms (Figure 4) provides apparent Brunauer–Emmett–Teller (BET) surface areas ($S_{\text{BET}}$) of 930 and 858 m² g⁻¹ for PIM-SBI-Trip and PIM-1/SBI-Trip respectively, which are within the upper range for solution processable PIMs,[9a,14,15] and greater than the values reported for PIM-1[8] and PIM-SBF-1.[8] Significantly, PIM-SBI-Trip possesses apparent microporosity that is similar to that of the best performing SBF-based PIMs.[19] This is in agreement with the predictions, based on molecular simulations, that larger substituents on rigid spiro-centre units both frustrate chain packing and increase interchain distances, resulting in enhanced porosity.[20]
Table 2. Single gas permeabilities $P$, diffusivity $D$, solubility $S$, and related ideal selectivities $\alpha$ for methanol treated and aged films of PIM-SBI-Trip ($l = 248 \mu m$) and its copolymer PIM-1/SBI-Trip ($l = 180 \mu m$), and PIM-1 measured at 25°C and 1 bar of feed pressure.

| Polymer | $P$ [Barrer] | Ideal selectivity ($P_a/P_b$) | $D$ [10$^{-12}$ m$^2$ s$^{-1}$] | Ideal selectivity ($D_a/D_b$) | $S$ [cm$^3$ cm$^{-3}$ bar$^{-1}$] | Ideal selectivity ($S_a/S_b$) |
|---------|-------------|-------------------------------|-------------------------------|-------------------------------|--------------------------------|-------------------------------|
| **N$_2$** | **O$_2$** | **CO$_2$** | **CH$_4$** | **H$_2$** | **He** | **N$_2$/O$_2$** | **CO$_2$/O$_2$** | **O$_2$/N$_2$** | **CO$_2$/CH$_4$** | **N$_2$/O$_2$** | **CO$_2$/O$_2$** | **O$_2$/N$_2$** | **CO$_2$/CH$_4$** |
| PIM-SBI-Trip | 2315 | 6390 | 35 600 | 4600 | 13 170 | 45 800 | 5.69 | 15.4 | 2.76 | 7.74 | 34.2 | 1.12 | 2.70 | 2.66 |
| (501 days) | 471 | 2295 | 11 500 | 590 | 7700 | 2730 | 16.3 | 24.4 | 4.87 | 19.5 | 4.41 | 1.38 | 3.41 | 3.27 |
| PIM-1/SBI-Trip | 1298 | 4230 | 24 410 | 2519 | 8360 | 2820 | 6.44 | 18.8 | 3.26 | 9.69 | 88.2 | 1.55 | 4.22 | 5.00 |
| (401 days) | 338 | 1500 | 7700 | 430 | 4500 | 1730 | 13.3 | 22.8 | 4.87 | 19.5 | 48.1 | 1.22 | 2.96 | 2.82 |
| PIM-1 | 506 | 1580 | 9650 | 913 | 3830 | 1560 | 7.57 | 19.1 | 3.12 | 10.6 | 90.6 | 1.45 | 3.84 | 4.35 |
| (2219 days) | 227 | 917 | 5000 | 302 | 2830 | 1220 | 12.5 | 22.0 | 4.04 | 16.6 | 3.48 | 1.35 | 2.55 | 2.42 |

*Prior to the permeability measurements, the films were soaked in methanol for 24 h to remove residual solvent, and then dried for 24 h in air. Values between parentheses give the sample age in days of the aged samples; $^a$ B Barrer $= 10^{-9}$ cm$^3$ STP cm$^{-2}$ s$^{-1}$ cm Hg$^{-1}$; $^b$ This work; $^c$ Associated measured time-lag data are reported in Table S1 (Supporting Information); $^d$ The calculated values for H$_2$ and He slightly lose in precision due to the fast membrane time-lag.
The order of gas permeability is $\text{CO}_2 > \text{H}_2 > \text{O}_2 > \text{He} \approx \text{CH}_4 > \text{N}_2$, and the similar values for He and CH$_4$ show that the permeability is strongly influenced by both the diffusion and the sorption coefficients. The high CH$_4$ permeability is due to its remarkably high solubility coefficient, which is among the highest known for any polymer (Table 2).\textsuperscript{[9a,14,15]} The PIM-1/SBI-Trip 1:1 copolymer demonstrates gas permeability data that lie neatly between those of the two parent homo polymers PIM-SBI-Trip and PIM-1 (Figure 5). The very high gas permeabilities of these two PIMs measured from freshly methanol treated films decrease over time (Table 2) due to physical ageing, a well-known feature of all high free volume glassy polymers.\textsuperscript{[23]} The decrease in permeability is mainly due to lower diffusion coefficients (Figure 6 and Figure S13a: Supporting Information).

![Figure 5](image_url)

**Figure 5.** Robeson plots for the a) CO$_2$/CH$_4$, b) CO$_2$/N$_2$, c) O$_2$/N$_2$, and d) H$_2$/N$_2$ gas pairs with the upper bounds represented by blue lines for 1991,\textsuperscript{[3]} red lines for 2008,\textsuperscript{[4]} yellow lines for 2015 (O$_2$/N$_2$ and H$_2$/N$_2$),\textsuperscript{[3]} purple lines for 2019\textsuperscript{[6]} (CO$_2$/N$_2$ and CO$_2$/CH$_4$) and a black dotted line for the upper bound proposed in 2018\textsuperscript{[24]} for the CO$_2$/CH$_4$ mixture, at 1 bar CO$_2$ partial pressure at 35 °C. The gas permeability data for PIM-SBI-Trip are reported in red (freshly MeOH treated ○; and aged 501 days ◆), and in blue for the PIM-1/SBI-Trip copolymer (freshly MeOH treated ○; and aged 401 days ◆). Measurements performed on the aged samples at 35, 45, and 55 °C are reported in triangles ▲ with the same color as the corresponding samples tested at 25 °C ◆. Data measured in mixed conditions on the aged films are indicated as empty symbols with the borders of the same color ◇. For comparison, the data measured under the same conditions by Longo et al. for PIM-1 after MeOH treatment ○ and 2219 days of ageing ◆ are indicated in green.\textsuperscript{[22]} Other literature data are reported in grey symbols for PIM-1 (+), SBF-based PIMs (○), ultrapermeable triptycene based PIMs (■) and PTMSP (×).
while solubility is not as significantly influenced by ageing (Figure S13b, Supporting Information).

The decrease in permeability is combined with a strong increase in selectivity, which places the data of aged PIM-SBI-Trip on the 2015 upper bound for O₂/N₂, with similar selectivity to that of commercial polymers used to produce membranes but with two-orders of magnitude higher permeability, and close to the 2019 upper bounds for CO₂/N₂ and CO₂/N₂ gas pairs (Figure 5).

For the H₂/N₂ gas pair, the trend upon ageing is also much steeper than the Robeson upper bound. The decrease in permeability is connected to the loss of larger elements of free volume over time resulting in a greater degree of ultramicroporosity and enhanced size-selectivity. The effect on permeability is stronger for bulkier gases with respect to smaller ones, for example, after 500 days of ageing, CH₄ loses 88% of its diffusion coefficient while He decreases only 17%, resulting in a marked increase in He/CH₄ permselectivity. This is clearly illustrated by the steeper correlation between the effective gas diameter, dₑff, and the diffusion coefficient, D (Figure S13a, Supporting Information).

Additional measurements on the aged films of PIM-SBI-Trip and PIM-1/SBI-Trip were carried out at temperatures of 25, 35, 45, and 55 °C (Figure S5 and Figure S14: Supporting Information). The gas diffusion coefficients increase with temperature (Figure S15, Supporting Information) with the aged membrane demonstrating very similar diffusivities as the freshly MeOH treated membrane at 25 °C (Figure 6).

Permeabilities (Figure S14, Supporting Information) also increase as a function of temperature due to the increasing diffusion coefficients, but they are not fully restored (Figure 5) since solubility decreases (Figure S16, Supporting Information). At higher temperature, the selectivities for all gas pairs decrease so that at 55 °C they are roughly similar to those of the freshly methanol treated samples at 25 °C.

The trends of P, D, and S as a function of temperature follow Arrhenius behavior described by Equations (1–3)

\[ P = P_0 \exp\left(\frac{-E_p}{RT}\right) \]  
\[ D = D_0 \exp\left(\frac{-E_d}{RT}\right) \]  
\[ S = S_0 \exp\left(\frac{-H_s}{RT}\right) \]  

Where, R is the universal gas constant; T the absolute temperature; \( P_0, D_0, \) and \( S_0 \) are the pre-exponential factors and define the intercept on the vertical axis in the Arrhenius plot, \( E_p \) is the activation energy of permeation, \( E_d \) the activation energy of diffusion, and \( H_s \) the heat of sorption. For both PIM-SBI-Trip and in PIM-1/SBI-Trip, \( E_p \) is higher than that of the triptycene-based ultrapermeable PIMs. Both polymers show among the highest values of \( H_s \) reported for PIMs (Table 3), indicating a strong polymer-sorbent interaction and high sorption capability.

The large values of \( E_d \) for larger gases (i.e., \( N_2 \) and \( CH_4 \)) correlate to the strong size-sieving properties of the polymers that result in their high diffusion selectivity (Table 2). Although there are some differences in the like-to-like comparison between specific gas pairs, the two polymers have comparable size-sieving properties, as highlighted by the similar slope of the activation energy of diffusion as a function of the square effective diameter of the penetrant gases (Figure S17, Supporting Information). According to the transition theory of diffusion,[27] the diffusion selectivity between two gases \( a \) and \( b \) \((D_a/D_b)\) can be defined as:

\[ \frac{D_a}{D_b} = \frac{\lambda_a^2 \exp\left(\frac{\Delta S_{d(a,b)}}{R}\right) \exp\left(\frac{-\Delta E_{d(a,b)}}{RT}\right)}{\lambda_b^2 \exp\left(\frac{\Delta S_{d(a,b)}}{R}\right) \exp\left(\frac{-\Delta E_{d(a,b)}}{RT}\right)} \]  

where \( \lambda_a \) and \( \lambda_b \) are the average diffusive jumps for gas \( a \) and \( b \); \( \Delta S_{d(a,b)} \) is the difference in the activation entropy of diffusion, i.e., \( \Delta S_{d(a,b)} = (S_{d(a)} - S_{d(b)}) = (S_{d(a)} - S_{d(b)}) - (S_{d(a)} - S_{d(a)}) \) where \( S_{d(a)} \) and \( S_{d(b)} \) are the entropies of the gas \( a \) and \( b \) in the activated (in the transition region) and normal (in the micropores) states respectively and it defines the entropic selectivity; and \( \Delta E_{d(a,b)} \) is the difference in the activation energies of diffusion, which influences the energetic selectivity. \( \lambda_a \) and \( \lambda_b \) are generally unknown for a specific matrix, but the ratio \( \lambda_a^2/\lambda_b^2 \) can be approximated as \( d_a^2/d_b^2 \), where \( d \) is the effective diameter of the penetrant gas allowing the estimation of the entropic and energetic contributions. The entropic selectivity is related to the changes in the degree of freedom of one gas during its diffusion process from the micropores (normal state) to the motion-enabled zone (transition state), and it is a characteristic of molecular sieve materials. The entropic selectivity is below 1 for O₂/N₂, CO₂/N₂, H₂/N₂, and He/N₂, which means that in the normal state
(i.e., during the diffusion within a micropore) the smaller gas has more translational freedom, while in the transition state (i.e., diffusion between two micropores), all degrees of freedom are suppressed. For a similar reason, the entropic selectivity (i.e., diffusion between two micropores), all degrees of freedom has more translational freedom, while in the transition state (i.e., during the diffusion within a micropore) the smaller gas molecules lose more entropy, which is a less favored thermodynamic process. The high value for CH\textsubscript{4}/N\textsubscript{2} entropic selectivity, when related to N\textsubscript{2} since their diffusion requires only small thermal motions of the polymer chains. Remarkably, the energetic selectivity for O\textsubscript{2}/N\textsubscript{2} in PIM-SBI-Trip is about 9 times higher than the best performing semi-rigid polymer 6FDA/PMDA(10/90)-TAB\cite{28} and that of the ultrapermeable PIM-BTrip.\cite{26} PIM-BTrip and PIM-SBI-Trip have similar BET surface areas, i.e., 911 and 930 m\textsuperscript{2} g\textsuperscript{-1} respectively. As the overall rigidities of these two PIMs are expected to be similar, the higher energetic selectivity of PIM-SBI-Trip may be related to packing of its 3D contorted chains resulting in a higher proportion of smaller micropores relative to that of the 2D chains of PIM-BTrip.

### 2.3. Mixed Gas Permeation

Gas mixtures of CO\textsubscript{2}/CH\textsubscript{4} (35/65 vol\%, Figure 7a,b), simulating biogas, and CO\textsubscript{2}/N\textsubscript{2} (15/85 vol\%, Figure 7c,d), simulating typical flue gas, were used to assess the performance of the PIM-SBI-Trip and PIM-1/SBI-Trip membranes, according to

| Property                  | Polymer          | N\textsubscript{2} | O\textsubscript{2} | CO\textsubscript{2} | CH\textsubscript{4} | H\textsubscript{2} | He   |
|---------------------------|------------------|--------------------|--------------------|--------------------|-------------------|-------------------|------|
| E\textsubscript{a} [kcal mol\textsuperscript{-1}] | PIM-SBI-Trip\textsuperscript{a,b)} | 5.79 ± 0.18 | 2.23 ± 0.21 | 1.91 ± 0.21 | 7.61 ± 0.04 | 0.44 ± 0.12 | 1.08 ± 0.31 |
| E\textsubscript{a} [kcal mol\textsuperscript{-1}] | PIM-1/SBI-Trip\textsuperscript{a,b)} | 4.74 ± 0.17 | 1.47 ± 0.13 | 1.36 ± 0.15 | 6.88 ± 0.06 | 0.33 ± 0.06 | 0.65 ± 0.06 |
| H\textsubscript{s} [kcal mol\textsuperscript{-1}] | PIM-SBI-Trip\textsuperscript{a,b)} | −3.92 ± 0.01 | −3.60 ± 0.66 | −5.32 ± 0.09 | −4.71 ± 0.06 | −1.93 ± 0.32 | −1.02 ± 1.65 |
| H\textsubscript{s} [kcal mol\textsuperscript{-1}] | PIM-1/SBI-Trip\textsuperscript{a,b)} | −3.86 ± 0.09 | −4.03 ± 0.10 | −5.03 ± 0.12 | −3.96 ± 0.04 | −1.47 ± 0.29 | −1.32 ± 1.13 |
| E\textsubscript{d} [kcal mol\textsuperscript{-1}] | PIM-SBI-Trip\textsuperscript{a,b)} | 9.71 ± 0.18 | 5.83 ± 0.51 | 7.23 ± 0.11 | 12.33 ± 0.08 | 2.37 ± 0.30 | 2.70 ± 1.14 |
| E\textsubscript{d} [kcal mol\textsuperscript{-1}] | PIM-1/SBI-Trip\textsuperscript{a,b)} | 8.60 ± 0.10 | 5.50 ± 0.04 | 6.40 ± 0.06 | 10.84 ± 0.09 | 1.80 ± 0.23 | 1.98 ± 1.09 |
| Diffusion selectivity D\textsubscript{N\textsubscript{2}}/D\textsubscript{G} | PIM-SBI-Trip\textsuperscript{a)} | 6 | 5.3 | 4.2 | 7.8 | 3.2 | 3 |
| Diffusion selectivity D\textsubscript{N\textsubscript{2}}/D\textsubscript{G} | PIM-1/SBI-Trip\textsuperscript{a)} | − | 4.72 | 1.54 | 0.266 | 107 | 176 |
| Energetic selectivity (see Equation (4)) | PIM-SBI-Trip\textsuperscript{a,b)} | − | 716 ± 127 | 66.8 ± 7.9 | 0.012 ± 0.001 | 2.45 · 10\textsuperscript{-5} | 3.88 · 10\textsuperscript{-5} |
| Energetic selectivity (see Equation (4)) | PIM-1/SBI-Trip\textsuperscript{a,b)} | − | 188 ± 8.5 | 41.4 ± 3.1 | 0.023 ± 0.002 | 9.79 · 10\textsuperscript{-5} | 7.26 · 10\textsuperscript{-5} |
| Entropic selectivity (see Equation (4)) | PIM-SBI-Trip\textsuperscript{a,b)} | − | 3.26 | 20.9 | 0.048 | 113 | 159 |
| Entropic selectivity (see Equation (4)) | PIM-1/SBI-Trip\textsuperscript{a,b)} | − | 0.007 ± 0.001 | 0.024 ± 0.003 | 0.207 ± 2.00 | 0.9 · 10\textsuperscript{-4} | 1.3 · 10\textsuperscript{-4} |
| PIM-1/SBI-Trip\textsuperscript{a,b)} | − | 0.025 ± 0.001 | 0.038 ± 0.003 | 12.4 ± 1.0 | 1.8 · 10\textsuperscript{-4} | 4.7 · 10\textsuperscript{-4} |
| PIM-1/SBI-Trip\textsuperscript{a,b)} | − | 0.99 | 0.077 | 6.37 | 0.81 | 0.81 |

\textsuperscript{a}This work; \textsuperscript{b}Experimental errors associated to these data are calculated via linear regression of the diffusion coefficients as a function of the temperature and via the error propagation theory; \textsuperscript{c}The calculated values for H\textsubscript{2} and He may slightly lose in precision due to the fast membrane time-lag.

This is the main factor that governs the size-sieving character of PIM-SBI-Trip and PIM-1/SBI-Trip and both polymers show values that are among the highest reported for PIMs (Table 3). This is ascribed to the overall rigidity of these polymers (Figure S12, Supporting Information), provided by the phenyl ring of the fused triptycene units reducing the flexibility about the spirocentre, as found for the spirobisfluorene PIMs (e.g., PIM-SBF-1)\cite{8,19} and the inherent rigidity provided by these bridged bicyclic substituents. He and H\textsubscript{2} have high energetic selectivity when related to N\textsubscript{2} since their diffusion requires only small thermal motions of the polymer chains. Remarkably, the energetic selectivity for O\textsubscript{2}/N\textsubscript{2} in PIM-SBI-Trip is about 9 times higher than the best performing semi-rigid polymer 6FDA/PMDA(10/90)-TAB\cite{28} and that of the ultrapermeable PIM-BTrip.\cite{26} PIM-BTrip and PIM-SBI-Trip have similar BET surface areas, i.e., 911 and 930 m\textsuperscript{2} g\textsuperscript{-1} respectively. As the overall rigidities of these two PIMs are expected to be similar, the higher energetic selectivity of PIM-SBI-Trip may be related to packing of its 3D contorted chains resulting in a higher proportion of smaller micropores relative to that of the 2D chains of PIM-BTrip.
a previously reported procedure. Ultrapermeable polymers easily lead to a high stage-cut, where the separation factor depends on the process conditions as much as the material properties. The experiments in this work were therefore carried out at low stage-cut (<1%, see Figure S18, Supporting Information), using a small membrane area, a high feed flow rate and a relatively high sweep flow rate, in order to focus mainly on the materials properties. The gas permeability in both PIM-SBI-Trip-based films shows a modest feed pressure dependence, similarly to that prepared from PIM-SBF-1. CO₂ permeability decreases with increasing feed pressure due to the saturation of Langmuir sites, which is a typical feature for materials showing dual mode sorption without undergoing strong swelling and/or plasticization. There is only weak hysteresis between mixed gas permeability measurements at increasing and then decreasing pressures in PIM-1/SBI-Trip. The slightly higher permeability during the pressure-decrease steps indicates a weak dilation of the polymer matrix at a higher partial pressure of CO₂ in the feed at higher total pressures or in CO₂ rich mixtures (Figure 7b). For both PIM-SBI-Trip and PIM-1/SBI-Trip, the mixed gas permeability data for the CO₂/CH₄ gas pair at 1–6 bar are similar to that obtained from single gas measurements and close to the 2019 upper bound. Although obtained at lower partial pressure, these data are also

Figure 7. a,c) Mixed gas permeation in the aged films of PIM-SBI-Trip and in its copolymer b,d) PIM-1/SBI-Trip as a function of the total feed pressure. Mixture CO₂/CH₄ (35/65 vol%) simulating biogas (a,b), and Mixture CO₂/N₂ (15/85 vol%) simulating a typical flue gas composition (c,d). Closed symbols and full lines represent the stepwise increasing pressure; open symbols and dashed lines represent the subsequent stepwise decreasing pressure. Lines are plotted as a guide to the eye and the arrows indicate the axis where to read the data. The analogous figures as a function of the CO₂ partial pressure are given in Figure S19 (Supporting Information).
well above the proposed 2018 CO$_2$/CH$_4$ upper bound for mixed gases at a CO$_2$ partial pressure of 10 bar\textsuperscript{[24]} (Figure 5a).

For both polymers, the CO$_2$/N$_2$ mixed gas permeability data lie in a more favorable position of the Robeson plot as compared to the single gas data (Figure 4b), and for PIM-SBI-Trip they are even above the 2019 upper bound. Remarkably, the mixed gas performance of PIM-SBI-Trip and PIM-1/SBI-Trip shows systematically higher selectivities than aged PIM-TMN-Trip and with similar CO$_2$ permeability (Figure S20, Supporting Information), demonstrating the potential of these ultrapermeable PIMs for biogas upgrading\textsuperscript{[32]} and CO$_2$ capture from flue gases.\textsuperscript{[34]} Previous work with benchmark polymer PIM-1 shows significantly improved selectivity in hollow fiber membranes,\textsuperscript{[15]} suggesting that also for the present polymers even better separation performance may be expected when prepared in the form of asymmetric or thin film composite membranes.

3. Conclusion

Fusing two bulky and rigid triptycene units to the SBI site of contortion provides a PIM with increased intrinsic microporosity, as demonstrated by enhanced gas adsorption and permeability. The resulting PIM-SBI-Trip and PIM-1/SBI-Trip are the first ultrapermeable PIMs in which the polymer chains are contorted in 3D. Their excellent permeselectivity data, in particular for the CO$_2$/CH$_4$ and CO$_2$/N$_2$ gas pairs, and the modest pressure dependence in mixed gas conditions, albeit demonstrated only at relatively low pressures, make these particularly promising materials for CO$_2$-related separations. In addition, PIM-SBI-Trip shows similar selectivity as traditional polymers used in industrial O$_2$/N$_2$ separation, but at much higher permeability, paving the way to the development of compact O$_2$ production or enrichment systems. This may be relevant to the development of small-scale oxygen generators for medical use in home or hospital by patients with respiratory diseases.

Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.

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Conflict of Interest

The authors declare no conflict of interest.

Data Availability Statement

The data that supports the findings of this study are available in the supplementary material of this article.

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gas separation, membranes, microporosity, polymers, size-sieving

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[19] C. G. Bezzu, M. Carta, M.-C. Ferrari, J. C. Jansen, M. Monteleone, E. Esposito, A. Fuoco, K. Hart, T. P. Liyana-Arachchi, C. M. Colina, N. B. McKeown, J. Mater. Chem. A 2018, 6, 10507.

[20] K. E. Hart, J. M. Springmeier, N. B. McKeown, C. M. Colina, Phys. Chem. Chem. Phys. 2013, 15, 20161.

[21] a) Q. Song, S. Cao, R. H. Pritchard, B. Ghalei, S. A. Al-Muhtaseb, E. M. Terentjev, A. K. Cheetham, E. Sivaniah, Nat. Commun. 2014, 5, 4813; b) K. Polak-Kraśna, C. Fuhrhop, S. Rochat, A. D. Burrows, A. Georgiadis, C. R. Bowen, T. J. Mays, Int. J. Hydrogen Energy 2017, 42, 23915; c) K. Polak-Kraśna, R. Dawson, L. T. Holyfield, C. R. Bowen, A. D. Burrows, T. J. Mays, J. Mater. Sci. 2017, 52, 3862.

[22] M. Longo, M. P. De Santo, E. Esposito, A. Fuoco, M. Monteleone, L. Giorno, B. Comesaña-Gándara, J. Chen, C. G. Bezzu, M. Carta, I. Rose, N. B. McKeown, J. C. Jansen, Ind. Eng. Chem. Res. 2020, 59, 5381.

[23] Z.-X. Low, P. M. Budd, N. B. McKeown, D. A. Patterson, Chem. Rev. 2018, 118, 5871.

[24] Y. Wang, X. Ma, B. S. Ghanem, F. Alghunaimi, I. Pinna, Y. Han, Mater. Today Nano 2018, 3, 69.

[25] L. M. Robeson, Z. P. Smith, B. D. Freeman, D. R. Paul, J. Membr. Sci. 2014, 453, 71.

[26] A. Fuoco, B. Comesaña-Gándara, M. Longo, E. Esposito, M. Monteleone, I. Rose, C. G. Bezzu, M. Carta, N. B. McKeown, J. C. Jansen, ACS Appl. Mater. Interfaces 2018, 10, 36475.

[27] W. J. Koros, C. Zhang, Nat. Mater. 2017, 16, 289.

[28] C. M. Zimmerman, W. J. Koros, Macromolecules 1999, 32, 3341.

[29] a) J. Crank, The Mathematics of Diffusion, Clarendon Press, Oxford 1975; b) A. Singh-Ghosal, W. J. Koros, Ind. Eng. Chem. Res. 1999, 38, 3647.

[30] P. Li, T. Chung, D. Paul, J. Membr. Sci. 2014, 450, 380.

[31] S. C. Fraga, M. Monteleone, M. Lanč, E. Esposito, A. Fuoco, L. Giorno, K. Pilnáček, K. Friess, M. Carta, N. B. McKeown, P. Izák, Z. Petrusová, J. G. Crespo, C. Brazinha, J. C. Jansen, J. Membr. Sci. 2018, 561, 39.

[32] P. Stanovsky, M. Karasova, Z. Petrusova, M. Monteleone, J. C. Jansen, B. Comesaña-Gándara, N. B. McKeown, P. Izak, J. Membr. Sci. 2021, 618, 118694.

[33] M. Monteleone, E. Esposito, A. Fuoco, M. Lanč, K. Pilnáček, K. Friess, C. G. Bezzu, M. Carta, N. B. McKeown, J. C. Jansen, Membranes 2018, 8, 73.

[34] P. Stanovsky, A. Zitkova, M. Karasova, M. Šyc, J. C. Jansen, B. C. Gándara, N. B. McKeown, P. Izak, Sep. Purif. Technol. 2020, 242, 116814.

[35] M. L. Jue, V. Breedveld, R. P. Lively, J. Membr. Sci. 2017, 530, 33.