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### MINI REVIEW

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# Hierarchically microporous membranes for highly energy-efficient gas separations

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The implementation of synthetic polymer membranes in gas separations, ranging from natural gas sweetening, hydrogen separation, helium recovery, carbon capture, oxygen/nitrogen enrichment, etc., has stimulated the vigorous development of high-performance membrane materials. However, size-sieving types of synthetic polymer membranes are frequently subject to a trade-off between permeability and selectivity, primarily due to the lack of ability to boost fractional free volume while simultaneously controlling the micropore size distribution. Herein, we review recent research progress on microporosity manipulation in high-free-volume polymeric gas separation membranes and their gas separation performance, with an emphasis on membranes with hourglass-shaped or bimodally distributed microcavities. State-of-the-art strategies to construct tailorable and hierarchically microporous structures, microporosity characterization, and microcavity architecture that govern gas separation performance are systematically summarized.

Keywords: Gas separation membranes; Hierarchical microporosity; Micropore size distribution; Configurational free volume; Solution-diffusion mechanism.

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### 1 Introduction

Gas separation membranes, based predominantly on synthetic polymers, have attracted substantial attention in various chemical, environmental, and energy-demanding separations, such as natural gas sweetening (CO<sub>2</sub>/CH<sub>4</sub>), helium recovery from natural gas (He/CH<sub>4</sub>), carbon capture (CO<sub>2</sub>/N<sub>2</sub>), hydrogen recovery (H<sub>2</sub>/N<sub>2</sub>, H<sub>2</sub>/CH<sub>4</sub>), and oxygen/nitrogen enrichment (O<sub>2</sub>/ N<sub>2</sub>), due to the advantages of high energy efficiency, small footprint, relatively low investment, and ease of operation. 1-3 However, synthetic polymer membranes are subject to a tradeoff between permeability and selectivity, known as the Robeson upper bounds first introduced in 1991 and later in 2008, 4,5 which were redefined in 2015 for H<sub>2</sub>/N<sub>2</sub>, H<sub>2</sub>/CH<sub>4</sub>, and O<sub>2</sub>/N<sub>2</sub> gas pairs<sup>6</sup> and in 2019 for CO<sub>2</sub>/N<sub>2</sub> and CO<sub>2</sub>/CH<sub>4</sub> gas pairs.<sup>7</sup> Common synthetic polymer membranes feature a broad pore size distribution of free volume-based micropores; thus, the enhancement of gas permeability by manipulating a polymer structure often leads to a more significant permeability increase for larger gases than smaller ones, resulting in decreased selectivities. This phenomenon has been frequently observed in dense polymer membranes and could be attributed to the lack of ability to boost fractional free volume while simultaneously controlling the size distribution of free volumes.8 In this regard, microporosity manipulation plays a crucial role in obtaining attractive membranes with outstanding gas separation performance.

Ideal gas separation membranes feature precise and uniform pore sizes, i.e., very narrow pore size distribution,

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which enables both high gas permeability and selectivity due to high microporosity and low tortuosity in such membranes. Crystalline membrane materials, such as metal-organic frameworks (MOFs)9 and covalent organic frameworks (COFs),<sup>10</sup> offer effective ways to adjust pore architecture and obtain narrow pore size distributions, potentially resulting in high permeability and high selectivity. However, almost all synthetic polymer gas separation membranes exhibit amorphous structures with random chain packing and microcavity architecture; thereby, it remains a significant challenge to achieve uniform or very narrowly distributed micropores in synthetic polymer membranes via the pore fabrication methods used in inorganic membranes. Biological membranes, such as aquaporins and potassium ion channels, have extraordinarily high permeability-selectivity combinations (Fig. 1a).8 This is ascribed to the unique hourglass-shaped cavities providing open spaces for fast molecule transport and narrow neck for highly selective molecule sieving (Fig. 1b). The concept of introducing hourglass-shaped microcavities has also attracted much attention in synthetic polymer gas separation membranes, and some high-performance gas separation membranes were also observed to have this kind of microcavity architecture. However, the practical development of these gas separation membranes is still in its infancy and hasn't been systematically reviewed. Besides, most recently published reviews of gas separation membranes focused on the macromolecular design and polymer synthesis strategies of polymers of intrinsic microporosity; 11-13 few highlighted the construction and manipulation of microcavity structures.

This review summarizes the research progress on microporosity manipulation in high-free-volume polymer gas separation membranes and their gas separation performance, with an emphasis on membranes with hourglass-shaped or bimodally distributed microcavities. Here, we propose the idea of "hierarchically microporous membranes" featuring

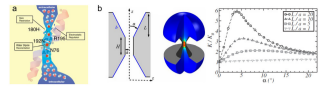


Fig. 1 (a) The sagittal cross-section of an aguaporin water channel (reprinted with permission from ref. 14, Copyright (2002) American Society for Clinical Investigation) and (b) the effect of the hourglass pore dimensional structure on water permeability (reprinted with permission from ref. 15, Copyright (2013) National Academy of Sciences).

both micropores (<2 nm) and ultra-micropores (<7 Å) with bimodal or multimodal microcavity size distributions, which are distinctly different from the broad microcavity size distributions conventional membranes. polymer Microporosity characterization methods, construction strategies of hierarchically microporous membranes, and structure-property relationships are discussed and reviewed. applications of hierarchically microporous membranes are also presented.

### 2 Characterization of free volume

Various techniques have been employed to characterize microporosity in synthetic polymer membranes, including positron annihilation lifetime spectroscopy (PALS),16 lowtemperature gas adsorption (LTGA),<sup>17</sup> spin probe method,<sup>18</sup> inverse gas chromatography, 19 and molecular modeling. 20 Among them, PALS is frequently used to probe the size distribution of synthetic polymer membranes. The relatively easy operation have recently promoted the use of LTGA in measuring micropore size and pore size distribution.



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industrial application of a number of green complete technologies. He has published over 600 peer-reviewed articles and his publications have received more than 32 000 citations with an H-index of 92 (Google Scholar), authored 11 monographs.

Mini review

### 2.1 Positron annihilation lifetime spectroscopy

Positron annihilation spectroscopy (PAS) has been demonstrated as an experimental instrumental technique to determine the free-volume (pore) properties of polymeric gas separation membranes. The positronium atom (Ps, a bound state of an electron and a positron, a quantum probe with a diameter of 0.106 nm) could be formed even in a subnanometer-scale pore with a diameter ranging from 1.5 to 7 Å (i.e., ultra-micropore).<sup>21</sup> As the most commonly used PAS technique, positron annihilation lifetime spectroscopy (PALS) provides direct measurements of pore size, size distribution, and total pore fraction in polymeric gas separation membranes. A relationship exists between the o-Ps (the spintriplet state of Ps) lifetime and the pore size, and the formation probability of o-Ps is correlated to the number of pores in a polymeric membrane. Using the CONTIN and MELT programs to analyze the PALS spectrum of a porous polymer, the size distribution of pores may also be obtained.

The age-momentum correlation (AMOC) of positron annihilation radiation provides valuable information on the chemical environment of the pore structure in the outermost layers of the membrane through analysis of the momentum distribution of long-lived o-Ps (the momentum of a thermalized positron is only approximately 0.025 eV at room temperature, far less than that of electrons). Additionally, using a variable mono-energetic positron beam (usually 0-30 keV), the depth-resolved porous structure (including porosity, pore connectivity, and chemical environment) of polymeric films in the depth range from the outermost layer to a maximum of  $\sim 10~\mu m$  can be revealed. Compared to the traditional gas absorption/desorption method, as discussed below, PAS possesses many apparent advantages in characterizing the microporous structures of polymeric membranes, as PAS experiments can be applied to polymeric membranes in various states: (1) as-prepared membranes but not broken powders (necessary for the gas absorption/ desorption method); (2) wet-state membranes; (3) membranes at different temperatures (fixed temperature for the gas absorption/desorption method); and (4) membranes in different gases at different pressures.

Guo *et al.* used PALS to investigate the micropore size distribution of novel pentiptycene-based polyimides, and a bimodal micropore size distribution with average cavity sizes of 7–8 and 3–4 Å was observed. The microporous structure of these pentiptycene-based polyimide membranes was highly tailorable through the "partial filling" of the substituent groups in the pentiptycene moieties, as demonstrated by the PALS results.<sup>22</sup> Lee *et al.* revealed that the microporosity changed from a unimodal to a bimodal pore size distribution after the thermal rearrangement of hydroxyl-containing polyimide membranes. It was believed that the smaller 3–4 Å microcavities provided interconnections between the larger 6–8 Å micropores, which significantly enhanced gas permeabilities.<sup>23</sup> Additionally, the evolution of microporosity in thermally crosslinked membranes was also monitored by PALS. Luo *et al.* observed that dual

thermally crosslinked polyimide membranes displayed a hierarchical micropore size distribution featuring ultramicropore and micropore sizes in the range of 2–6 Å and 7–10 Å, respectively, which was different from the unimodal pore size distribution of the precursors. The dual thermally crosslinked polyimide membranes showed excellent plasticization resistance and low-temperature gas separation performance. $^{24}$ 

### 2.2 Low-temperature gas adsorption

The low-temperature gas adsorption technique uses an inert gas (e.g., N2) that doesn't chemically adsorb onto an adsorbent as a probe to evaluate the surface area, average pore size, and pore size distribution of porous materials. 17,25,26 For most mesoporous materials with pore sizes in the range of 2-50 nm, the low-temperature N<sub>2</sub> adsorption can achieve reliable data due to their high enough adsorbing amounts. However, it usually takes too much time to obtain valuable data for pore size analysis of microporous (pore size <2 nm) materials. In this case, CO<sub>2</sub> is more frequently used to investigate the free volume and intrinsic microporosity in cellular materials due to the much shorter adsorption equilibrium time at 273.15 K.<sup>27</sup> Therefore, appropriate testing conditions and using adequate samples with repeated measurements are crucial for acquiring reliable pore structure data using this technique. Low-temperature adsorption has been widely used to probe micropore characteristics, including the intrinsic Brunauer-Emmett-Teller (BET) surface area, micropore size distribution, and gas storage capacity.27 Pinnau et al. studied the microcavity size distribution of a series of triptycene-based polymers of intrinsic microporosity using low-temperature nitrogen adsorption, and they found that both the TPIM and KAUST-PI-1 membranes exhibited a typical hierarchical bimodal pore size distribution using N<sub>2</sub> molecules as the probe. <sup>28,29</sup>

# 3 Hierarchically microporous gas separation membranes

### 3.1 Polymers of intrinsic microporosity

Polymers of intrinsic microporosity (PIMs), which exhibit unprecedentedly high gas separation performance, have attracted significant attention since they were first developed in 2004 due to their characteristic ladder-type, contorted, and rigid polymer architectures enabling high microporosity.<sup>30</sup> Different PIMs have since then been developed with various structural considerations, such as PIM-1-based ladder polymers, PIM-polyimide (PIM-PI) polymers, and Tröger's base-based (PIM-TB) polymers (Fig. 2). Benchmarking PIM membranes in Robeson's upper bound plots<sup>4,5</sup> highlights ultrahigh gas permeabilities compared with conventional polymer membranes; however, the gas selectivities of PIMs need to be improved. The relatively low gas selectivities of PIM membranes can be ascribed to the absence of well-defined microvoids, especially ultramicrocavities, which are crucial for size sieving. In this

Fig. 2 Representative chemical structures of ladder-type PIMs.

regard, the fabrication of hierarchically microporous PIMs via pore manipulation strategies is of great importance.

PIM-1 is the most studied polymer for gas separations, and various approaches were developed to overcome its relatively low gas selectivity. For example, amidoxime-functionalized PIM-1 (AO-PIM-1, Fig. 2) was synthesized through the reaction between the nitrile groups and hydroxylamine, leading to a hierarchical microcavity size distribution with ultramicroporosity (<7 Å) as analyzed by low-temperature nitrogen adsorption using NLDFT micropore analysis, which led to a three-fold increase in CO<sub>2</sub>/CH<sub>4</sub> selectivity compared to PIM-1.<sup>31</sup> A similar trend was observed for hydrolyzed PIM-1 membrane (hPIM-1, Fig. 2), which displayed a multimodal pore size distribution from the NLDFT analysis of N2 adsorption isotherms at -196 °C, 32 and the CO<sub>2</sub>/N<sub>2</sub> selectivity of the hPIM-1 membranes increased significantly with the increase of hydrolysis degree.<sup>33</sup> Additionally, direct fluorination of PIM-1 using a F<sub>2</sub>/N<sub>2</sub> (5/95) gas mixture significantly improved size sieving due to the fluorine atom-induced pore blocking effect, and the fluorinated membranes also showed outstanding aging resistance.34

In addition to PIM-1 functionalization, novel ladder polymers of intrinsic microporosity were fabricated aiming at microcavity manipulation using either new PIM monomers or polymerization methods. Guo et al.36 synthesized two pentiptycene-based PIM copolymers (PPIMs) with permanent and configurational free volume instilled by the shapeconsistent pentiptycene moieties (Fig. 2). The as-prepared pentiptycene-based PIM membranes exhibited a typical bimodal micropore size distribution with two primary peaks at around 7-8 Å and 10-14 Å, respectively. The resulting hierarchical microcavity architecture enables superior gas separation performance far exceeding the 2008 upper bound and exceptional resistance to physical aging that often plagues high-free-volume PIMs. Luo et al.37 reported a new macromolecular design of full-ladder polymers of intrinsic microporosity containing 9H-xanthene units (SACP-3) by superacid-catalyzed Friedel-Crafts polymerization. SACP-3 exhibited a bimodal micropore size distribution with two

maxima at ~5 Å and ~8 Å (Fig. 3a). As a result, it showed a much higher CO2 permeability of 6497 Barrer due to the more interconnected microporous structure, and the super rigid ladder structure enabled superior plasticization resistance unattainable in PIM-1. Xia et al. reported a novel class of hydrocarbon ladder polymers that displayed outstanding gas separation performance and aging resistance compared to conventional PIM membranes.<sup>38</sup> Particularly, the CANAL-Me-DHP membrane exhibited high H<sub>2</sub>/CH<sub>4</sub> and CO<sub>2</sub>/CH<sub>4</sub> selectivities of 621 and 68, respectively, with high H<sub>2</sub> and CO<sub>2</sub> permeabilities of 860 and 94 Barrer, respectively, after aging for 158 days. The profound effect of aging behavior was ascribed to the bottlenecks between freevolume elements narrowed down over time, leading to a more significant decrease in diffusion coefficients for larger gases. McKeown et al. fabricated novel PIMs containing spirocyclic units with fused triptycene, and the membranes exhibited a bimodal pore size distribution with a predominant ultra-microporosity size in the range of 5-7 Å and ultrahigh gas permeabilities.<sup>39</sup>

Tröger's base (TB)-based PIMs represent another type of high-performance gas separation membrane material due to the unique three-dimensional, V-shaped structure of Tröger's base, which has a characteristic twisting angle of 117° and can effectively disrupt polymer chain packing and generate high microporosity (Fig. 2). 40,41 Compared to PIM-1, PIM-EA-TB showed a bimodal micropore size distribution with higher chain rigidity, as evidenced by simulating their torsional angles. 42 Therefore, PIM-EA-TB exhibited higher size sieving and gas selectivities. Hu et al. synthesized four norbornyl bisbenzocyclobutene-Tröger's base polymers, and it was found that the anti-CANAL-4-MeNH<sub>2</sub>-based TB polymer showed a unimodal pore size distribution, while the syn-CANAL-4-MeNH<sub>2</sub>-based TB polymer displayed a bimodal pore size distribution. 43 Consequently, the syn-CANAL-4-MeNH2-based TB polymer showed higher gas permeabilities and selectivities, and its gas separation performance for H<sub>2</sub>/CH<sub>4</sub>, H<sub>2</sub>/N<sub>2</sub>, and O<sub>2</sub>/N<sub>2</sub> gas pairs approached the 2015 upper bounds.6 Carta et al.44 systematically investigated the microporosity and gas separation properties of four PIM-TB polymers, e.g., PIM-Trip(H2)-TB, PIM-Trip(Me2)-TB, PIM-

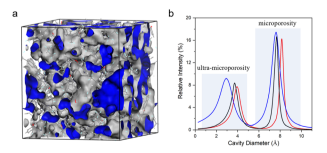


Fig. 3 (a) Modeling of micropore surfaces for a typical hierarchically microporous ladder polymer (reprinted with permission from ref. 35, Copyright (2022) Elsevier) and (b) pore size distribution of iptycenebased PI membranes (reprinted with permission from ref. 22, Copyright (2016) Elsevier).

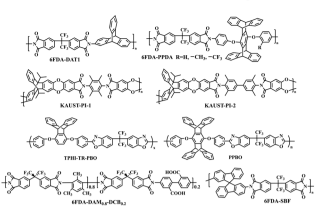
EA(Me<sub>2</sub>)-TB, and PIM-EA(H<sub>2</sub>)-TB (Fig. 2). Interestingly, all PIM-TB membranes revealed a trimodal pore size distribution with the main pore sizes centered at around 3.5, 5.5, and 8.3 Å, respectively, confirming the enhancement of pore connectivity and consequently the superior gas separation performance. Pinnau *et al.* reported the synthesis of catalytic arene–norbornene annulation–Tröger's base ladder polymers (CANAL-TBs), which showed high BET surface areas of 900–1000 m<sup>2</sup> g<sup>-1</sup> and a hierarchically bimodal pore size distribution. The CANAL-TBs exhibited promising  $O_2/N_2$  separation performance.

### 3.2 Polyimides and their derivatives

Polyimides (PIs) are arguably the most widely used membrane materials for gas separation, due to their outstanding mechanical strength, excellent thermal stability, high structural tunability, and attractive gas selectivities. However, conventional polyimide membranes suffer from relatively low fractional free volumes and gas permeabilities, hindering their wide applications. Thus, various strategies were employed to manipulate the micropore structure to boost polyimides' gas separation performance, including novel bulky and rigid PI monomers, thermal rearrangement (TR), and thermal crosslinking. This section focuses on the recent progress in the development of microporous polyimides and their derivative membranes with tailored hierarchical microporous structures and the fundamental structure–property relationship.

Iptycenes are a family of bulky and rigid molecules composed of multiple benzene rings fused into a 3D configuration, among which triptycene and pentiptycene are the most frequently studied structure units to develop microporous membranes.46 Incorporating iptycene moieties efficiently inhibit chain packing, enhancing the fractional free volume and micropore connectivity. 47,48 Additionally, the permanent configurational free volume elements delineated by the adjacent benzene rings exhibited comparable dimensions with the kinetic diameters of gas molecules, providing a unique strategy to finely tune the microporosity and enable high gas selectivities.49,50 For instance, Luo et al. 47 synthesized a series of pentiptycene-based polyimides, which displayed bimodal micropore size distributions (6FDA-PPDA, Fig. 4). Compared to the corresponding triptycenecontaining polyimides (6FDA-DAT1, Fig. 4), pentiptycenebased polyimide membranes showed higher FFVs and correspondingly higher gas permeabilities, and the microcavity could be finely modulated by different substituents (Fig. 3b). Notably, these iptycene-based polyimide membranes exhibited excellent resistance to physical aging or unusual aging-enhanced separation performance<sup>51</sup> due to the "stable" and accessible configurational free volumes generated by the shapepersistent iptycene units (Fig. 5a). 22,52

Compared with PIM-1-based ladder polymers, PIM-PI polymers have also attracted much attention due to their



**Fig. 4** Representative chemical structures of microporous polyimides and thermally rearranged (TR) polymers.

enhanced processability and excellent gas separation performance. Pinnau et al. fabricated KAUST-PIs (Fig. 4) containing the three-dimensional rigid triptycene moieties, and KAUST-PI-1 showed a super high BET surface area of 752 m<sup>2</sup> g<sup>-1</sup>.<sup>29</sup> Notably, pore size distributions obtained from NLDFT analysis demonstrated a bimodal pore size distribution for both KAUST-PI-1 and KAUST-PI-2, and it was believed that interconnectivity induced by ultra-microporosity assisted molecule transport and united the high permeabilities with high selectivities for H<sub>2</sub>/N<sub>2</sub>, H<sub>2</sub>/CH<sub>4</sub>, and O<sub>2</sub>/N<sub>2</sub> separations. This hierarchically microporous structure provided excellent gas separation performance with an H2 permeability of 3983 Barrer and H<sub>2</sub>/N<sub>2</sub> selectivity of 37 for KAUST-PI-1. Later on, KAUST-PI-1' was fabricated by replacing 9,10-diisopropyl-substituted triptycenes with 9,10dipropyl-substituted triptycenes, and the obtained KAUST-PI-1' showed a similar bimodal pore size distribution.<sup>53</sup>

The development of high-performance polyimide membranes using other bulky structures was also reported. A spirobifluorene-containing polyimide (6FDA-SBF, Fig. 4) featuring intrinsic microporosity with a surface area of 550  $\rm m^2~g^{-1}$  was produced, and a multimodal pore size distribution was observed. Yan *et al.* designed and synthesized dibenzodioxane-containing polyimides, and they all exhibited a bimodal pore size distribution characterized

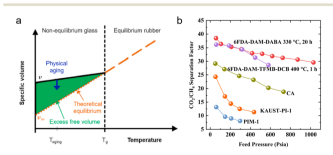


Fig. 5 (a) Physical aging of amorphous polymers (reprinted with permission from ref. 66, Copyright (2020) Elsevier) and (b) plasticization of typical crosslinked polyimide and PIM membranes (reprinted with permission from ref. 67, Copyright (2023) American Chemical Society).

by low-temperature N<sub>2</sub> adsorption.<sup>55</sup> Similarly, the 3D rigid dibenzodioxane moieties contributed to ultra-micropores (<7 Å), boosting their gas selectivities. Additionally, intrinsically microporous spirobifluorene-containing polyimides were designed and synthesized.56 A significant increase in the number of ultra-micropores due to the shrinkage of larger micropores after aging for 200 days was observed, resulting in a dramatic enhancement of gas separation performance for O<sub>2</sub>/N<sub>2</sub> and H<sub>2</sub>/N<sub>2</sub> gas pairs.

Thermally rearranged (TR) polymers, which are generally produced by thermal treatment of ortho-functional (e.g., hydroxyl, thiol, and amine) polyimide precursor membranes at elevated temperatures (300-450 °C) under inert conditions, are a highly attractive group of membrane materials that exhibit unprecedentedly high gas separation performance because of highly rigid and contorted backbone structures microporosity with hourglass-shaped architectures. 57 During the thermal rearrangement, an interconnected microcavity structure featuring a bimodal pore size distribution is generated. This leads to a significant enhancement of both permeability and selectivity compared to the polyimide precursors.<sup>58</sup> The microporosity and gas transport properties of TR polymers have been demonstrated to be highly tailorable via adjusting the precursor chemical structure or altering thermal treatment protocols.

Recent progress in TR polymers focuses on incorporating shape-persistent iptycene moieties to further microporosity by introducing configurational free volume. 49,50,59,60 For example, triptycene-based TR polymers (TPHI-TR-PBO in Fig. 4) derived from THPI and ether-free triptycene-containing HPI exhibited significantly enhanced gas permeabilities for  $H_2/N_2$ ,  $H_2/CH_4$ , and  $CO_2/CH_4$  gas pairs.<sup>50,59</sup> More recently, a series of pentiptycene-based TR polymers (PPBO in Fig. 4) were reported, which demonstrated highly tailorable microporosity and, consequently, a broad range of separation performances via simply altering the TR protocols.<sup>61</sup> The impressive boosted gas separation performances were ascribed to the formation of well-defined microcavities during TR conversion in combination with the configurational free volume of iptycene units, demonstrated by the increase in the population of microcavities (>7 Å) and negligible decrease in the population of ultra-microcavities (<7 Å). Later, new TR-able diamines such as adamantane<sup>62</sup> and fluorene-containing<sup>63</sup> hydroxyl diamines were designed to increase the TR conversion ratio and gas transport properties. Crosslinked TR polymers, which displayed the typical bimodal pore size distribution, were developed through simultaneous thermal rearrangement and crosslinking. 64,65 Notably, the population of micropores increased with increasing crosslinking degree, rendering a dramatic enhancement of gas permeabilities.

Conventional polyimide membranes susceptible to plasticization due to polymer swelling upon adsorption of condensable gases, which results in severe deterioration of the mixed-gas CO2/CH4 selectivity for natural gas sweetening (Fig. 5b).68 Thermally-induced chemical crosslinking of polyimide membranes has been widely implemented to suppress plasticization because it can appreciably regulate their microporosity and gas separation properties. Luo et al. developed a dual-thermally crosslinked strategy using a novel 4,4'-diamino-2,2'-biphenyldicarboxylic acid monomer.<sup>24</sup> The dual thermally crosslinked polyimide membranes displayed a hierarchically bimodal pore size distribution with ultra-micropores (2.0-6.0 Å) and micropores (6.5-10.0 Å) because of the collapse of segments and evolution of CO2 during thermal treatment. The crosslinked membrane has superior plasticization resistance and mixedgas separation performance to those of conventional noncrosslinkable polyimide membranes (Fig. 5b). Controllable thermal oxidative crosslinking was also recently reported to construct well-defined microcavities and crosslinked gas to boost separation networks and diminish plasticization. 40,69 Notably, a new crosslinked membrane design involving a model network structure formed by thermal end-linking of telechelic oligomers of precisely controlled molecular weight was reported. 70,71 In sharp contrast to traditional randomly crosslinked membranes, these crosslinked model network membranes showed an even higher gas permeability than their linear counterparts due to the model network design, and their separation performance was demonstrated to be highly tunable via simply adjusting the chain length of the telechelic oligomers, from which they were prepared.

### 3.3 Carbon molecular sieve membranes (CMSMs)

CMSMs, which are generally fabricated by carbonization of polymer precursors, display characteristic hierarchical microcavities and ultra-microcavities, thus exhibiting molecular sieving while allowing high gas permeabilities.<sup>72</sup> The representative turbostratic structure of CMSMs comprises graphene layers packed imperfectly, leading to slitlike micropores and ultra-micropores.<sup>73</sup> Pyrolysis has been considered the critical process for CMSM fabrication, which generates the microporous carbon structure necessary for molecular sieving. Various polymer precursors, including polybenzimidazoles, polyimides, phenolic polyacrylonitrile (PAN), polyfurfuryl alcohol (PFA), PIMs, cellulose, polysulfones, etc., have been fabricated into CMSMs via pyrolysis.<sup>74</sup> During this process, H<sub>2</sub> evolution forms a graphite-like structure and heteroatom elimination results in polymer chain rearrangement.<sup>75</sup> Therefore, the microporosity of CMSMs is primarily manipulated by the polymer precursors and the pyrolysis conditions such as the pyrolysis atmosphere, temperature, pyrolysis time, and ramp rate.

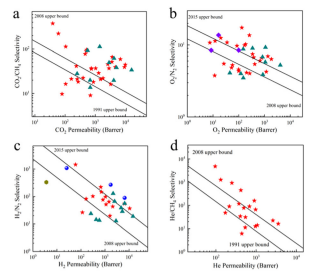
Among the pyrolysis conditions, the pyrolysis temperature plays a crucial role in regulating the microcavities. Generally, increasing the pyrolysis temperature increases crystallinity and reduces the average interplanar spacing, resulting in lower gas permeabilities and higher selectivities. This was evidenced by the weakening of the micropore peak (>5 Å) and the increase of the ultra-micropore peak (<5 Å),

providing evidence for the formation of ultramicropores at higher temperatures. Besides, high ramping rates usually lead to pinholes and cracks, while low ramping rates result in higher carbon crystallinity and ultra-micropores. Additionally, the pyrolysis atmosphere also affects the micropore structures. An oxidizing atmosphere typically enhances microporosity by degrading the precursors, yielding more open structures; an inert environment retains the microstructure and results in a tightly packed structure. Particularly, it was demonstrated that high-temperature H<sub>2</sub> exposure increased the CMS ultra-micropores, indicating that the high-temperature H<sub>2</sub> exposure can be used as post-synthesis modification to adjust the pore structure and the separation performance of CMS membranes.

Besides the pyrolysis conditions, various post-treatment including post-oxidation, chemical strategies, deposition, and post-pyrolysis, were employed to finely tune the microcavity size and distribution for demanding separation requirements.<sup>79</sup> Typically, post-oxidation or activation is frequently applied to increase the average micropore size when exposed to an oxidizing atmosphere. On the other hand, gas selectivities or molecule sieving of CMSMs can be enhanced by chemical vapor deposition (CVD), which introduces organic compounds into the micropore system and the pyrolytic decomposition. 80,81 Additionally, post-pyrolysis is another approach employed after post-oxidation to reduce micropore dimensions, and post-oxidation and post-pyrolysis are sometimes repeated alternately to achieve the desired micropore distribution.

Ascribed to the hierarchically microporous architecture, CMSMs exhibit unprecedentedly high permselectivities far beyond the Robeson upper bounds for various crucial industrial gas separations such as natural gas sweetening ( $CO_2/CH_4$ ), hydrogen recovery ( $H_2/N_2$ ,  $H_2/CH_4$ ), nitrogen removal from natural gas ( $N_2/CH_4$ ), oxygen enrichment ( $O_2/N_2$ ), and helium recovery from natural gas ( $He/CH_4$ ),  $He/CH_4$ ), as depicted in Fig. 6. Analysis of gas separation performance indicates that the ultrahigh gas selectivities originate from the constricted ultra-microcavities, which allow the permeation of small gases and exclude the transport of bulkier molecules. Herefore, the precise control of micropore dimensions of CMSMs renders an efficient separation of gas mixtures even with slight size differences.

However, CMSMs always suffer from undesirable permeability/permeance loss, recognized as physical aging.<sup>100</sup> Physical aging is primarily caused by the initial imperfections of the graphene-like layers, making the CMSMs undergo gradual rearrangements to reach their thermodynamic equilibrium state. This is further evidenced by the decrease of the D/G ratio in Raman spectroscopy, revealing that the graphene-like layers of CMSMs are prone to rearrangement into a more ordered architecture.<sup>101</sup> Thereafter, various approaches, such as pre-crosslinking, post-crosslinking,<sup>102</sup> and polydimethylsiloxane (PDMS) coating,<sup>103</sup> were developed to mitigate the physical aging of CMSMs. For instance, the



**Fig. 6** Gas separation performance of CMS membranes for (a)  $CO_2/CH_4$ , (b)  $O_2/N_2$ , (c)  $H_2/N_2$ , and (d)  $He/CH_4$  gas pairs. The red stars represent PI-based CMSMs,  $^{78,85-94}$  navy spheres represent PIM-based CMSMs,  $^{95,96}$  violet diamonds represent PBI-based CMSMs,  $^{97}$  dark yellow hexagon represents PFA-based CMSMs,  $^{98}$  and turquoise triangles represent polypyrrolone-based CMSMs.  $^{99}$ 

pre-crosslinking of polyimide precursors inhibited intersegmental motion and prevented the collapse of micropores, resulting in higher graphitic carbon content and less permeability reduction for the resulting CMSMs.<sup>100</sup> Despite numerous studies, limited success has been achieved in overcoming physical aging, and novel strategies are highly demanded to guarantee long-term stability. Additionally, low membrane strength has been the main obstacle limiting the wide application of CMSMs at the industrial level; therefore, novel CMSM fabrication methods, such as supported CMSMs, are needed to enhance membrane strength.

## 4 Conclusion and perspectives

This mini review systematically summarizes the recent progress on hierarchically microporous polymers, including highly microporous polymers of intrinsic microporosity, relatively less porous polyimides and their derivatives, and carbon molecular sieves, for high-performance separations. PIM functionalization, incorporation hierarchical moieties such as iptycene, and chemical crosslinking are efficient approaches to constructing welldefined hierarchical architectures featuring both microporosity and ultra-microporosity with characteristic bimodal or multimodal pore size distributions. Hierarchical microporosity provides narrow necking for molecule sieving and micropore connectivity for fast gas transport, rendering significantly enhanced gas separation performance for various gas pairs compared to conventional polymer membranes with generally broad pore size distributions.

Despite the advantages of hierarchically microporous membranes, many novel materials and design approaches are highly demanded to control micropore structures better and overcome the conventional upper bounds, providing directions for future research. To begin with, simulation and modeling at length scales from atomistic to continuum are crucial for understanding the basic structure–property relationship and offering rational guidance for the molecular-level design of future membranes. Secondly, pore engineering strategies such as anchoring functional groups to the micropore wall and incorporating crystalline structures to precisely manipulate the micropore size and architecture are highly demanded in future development. Developing novel membrane materials with diminished plasticization and physical aging would also expand their upscaling applications and boost membrane stabilities.

### Conflicts of interest

The authors declare no conflict of interest.

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