Membrane materials targeting carbon capture and utilization

Guining Chen, Tianlei Wang, Guangru Zhang, Gongping Liu, Wanqin Jin

State Key Laboratory of Materials-Oriented Chemical Engineering, College of Chemical Engineering, Nanjing Tech University, 30 Puzhu Road (S), Nanjing, 211816, China

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ABSTRACT

Over the past decades, fossil fuel combustion has emitted large quantities of CO2 into the atmosphere, resulting in global climate change. Nowadays, it's considered a feasible strategy to capture CO2 from some significant point sources. Three main strategies have been developed, namely post-combustion, pre-combustion, and oxy-fuel combustion. Recent research indicates that the membrane technology for CO2 capture has become competitive compared with conventional technologies because of the improved separation performance in materials and process designs. This paper mainly reviews the progress and breakthroughs of membrane materials for the three gas separation systems corresponding to the CO2 capture strategies. Besides, the CO2 utilization by the membrane process has also been highlighted.

1. Introduction

The climate environment has been dramatically affected because of the rapid development of industry and the dependence on fossil fuels. According to the latest CO2 emission data supported by the global monitoring laboratory, the current concentration of CO2 sits at ~416 ppm, which is excessively higher than what it was at the beginning of the last century [1]. This phenomenon has attracted serious concern in government organizations in the past few decades. The Intergovernmental Panel on Climate Change (IPCC) reports that human activities will cause a temperature increase of 1.5°C by 2030–2052 [2]. Consequently, the United Nations Framework Convention on Climate Change (UN-FCCC) treaty was reached in the United Nation’s Earth held in 1992 to control the concentration of CO2 and other greenhouse gases in the atmosphere, which accelerated the research and development of CO2 capture technologies.

Among various CO2 emission sources, the power generation sector is considered to be a significant anthropogenic source. Therefore, the CO2 capture technology from power plants has aroused great interest for many governments and researchers. At present, three main strategies are summarized for mitigation of CO2 emission [3-6]: 1. The post-combustion process (CO2/N2): the CO2 is separated from the low-pressure exhaust gas of fuel combustion in the air. 2. The pre-combustion process: the CO2 is captured from syngas produced by gasification. 3. The oxy-fuel combustion: the fuel combustion is realized in a pure oxygen atmosphere, leading to mixed gases that mainly contain CO2 and H2O. Although the oxy-fuel combustion process does not involve the direct CO2 separation, the exhaust gas with high CO2 concentration will greatly facilitate the subsequent CO2 separation and improve the capture efficiency. The general outline of the CO2 capture process is illustrated in Fig. 1. In addition, natural gas purification and biogas sweetening (mainly involving CO2/CH4) will not be involved in this review, although membrane separation has achieved remarkable progress in this area.

Nowadays, the difficulty of CO2 capture is to reduce energy consumption and capital cost. As of 2020, most commercial-scale carbon capture projects are in early development [7]. In numerous processes studied for CO2 capture, membrane technology is known for its ease of installation, energy efficiency, flexibility in operation and maintenance, and the ability to overcome thermodynamic limitations [8]. It also requires very few chemicals compared to conventional separation processes. With the massive emission of CO2, the demand for CO2 capture provides an opportunity for the deployment and application of membrane materials.

This paper mainly reviews the materials selection and development for diverse scenarios. According to the three main carbon capture strategies, we reviewed popular materials including polymers, inorganics and hybrid materials and their separation performance, respectively. Subsequently, CO2 utilization by membrane reactor, as briefly discussed using some cases. Finally, we discussed the challenges and opportunities of membrane materials for CO2 capture and utilization.
2. Post-combustion

Post-combustion means the separation of CO₂ from flue gases produced by burning fossil fuels. Generally, flue gases are mainly composed of N₂, water vapor, and CO₂. The content of CO₂ is primarily controlled by the used combustibles. Due to the low CO₂ partial pressure and the enormous amount of flue gas, high CO₂ permeable membranes with moderate CO₂/gas selectivity are critical for the practical application of membrane technology in CO₂ capture.

2.1. Polymeric membranes

Among numerous membrane materials, polymer-based materials show intrinsic advantages in cost, processability, and variety. Generally, solution-diffusion and facilitated transport mechanism has been widely recognized and adopted to guide the design of new polymers. Polymers including polyacetylene, polyaniline, polyetherimides, polycarbonates, poly(phenylene oxide), poly(ethylene oxide), and polysulfone have been investigated for post-combustion. By manipulating polymer preparation and chemical composition, these membranes have highly tunable membrane permeability and selectivity. However, the swelling and plasticization caused by CO₂ adsorption cannot be neglected.

Among the efforts to enhance CO₂ performance, polymer blends and copolymers have shown high potential. In 1982, Kawakami et al. reported a pioneering work by blending a plasticizer polyethylene glycol (PEG) in a low permeable cellulose nitrate [9]. It was observed that membrane performance improved significantly with increasing PEG percentage and molecular weight. Subsequently, composite polymeric membranes with various groups or free volumes were developed for CO₂ separation to compensate for the defects of a single polymer or combine the advantages of multiple functional groups. Similarly, copolymers and cross-linked polymers have radiated greater vitality. One of the most famous copolymers is PEO-based materials.

Poly (ethylene oxide) (PEO) has been widely used to improve CO₂ permeability in membranes because of the high affinity towards CO₂ caused by the quadrupolar interactions of the polar ether group with CO₂ molecules. In 2012, Jiang group reported a new polyethylene oxide-b-polystyrene block copolymer which exhibited high CO₂ permeance over 20,000 Barrer [10]. The research demonstrated that the self-assembled microphase structures and functional block units played synergistic roles in the high membrane performance. In recent decades, a series of commercial membranes, e.g., Pebax®, Polaries®, and Polyactive® have been developed and proved to be promising candidates for post-combustion capture. The typical structure of Pebax® MH 1657 is shown in Fig. 2(a), where the amide groups (PA) account for 40%, and the ether linkages (PE) account for 60%. (1 Barrer = 1 × 10⁻¹⁰ cm³(STP)/cm²·s·cmHg)

![Diagram of CO₂ capture approaches from power plants](image)

Fig. 1. Three CO₂ capture approaches from power plants.

![Diagram of polymer structures](image)

Fig. 2. (a) Pebax® MH 1657; (b) thermal rearrangement protocol of polyimide; (c) PIM-1 and PIM-SBF structure.
cm$^{-2}$ s$^{-1}$ cmHg$^{-1} = 3.344 \times 10^{-16}$ mol m$^{-2}$ Pa$^{-1}$ s$^{-1}$)

In 2007, Park and coworkers prepared new polymers via thermal rearrangement of polyimides and polyamides [11]. The thermal rearrangement process can generate high free volume and narrow cavities in polymers, resulting in a substantial increase in permeance by at least two orders of magnitude. Then, extensive works about the design of precursor structures and heat treatment protocols have been reported to optimize the separation performance and increase the mechanical strength. Freeman's group studied the influence of temperature in a polyimide precursor during thermal treatments [12]. Fig. 2(b) gives the reaction scheme where the polyimide HAB-6FDA was converted to polybenzoxazoles. The sorption coefficients of membranes showed the same upward trend with the increase in temperature. Sanders et al. compared a series of functional polyimide precursors and the resulting TR polymers [13]. Studies have shown that large functional groups can effectively inhibit the chain packing and increase free volume. Since the superior thermal stability and outstanding performance, TR polymers demonstrate bright application prospects in post-combustion.

Another well-known polymer with high permeability is polymers of intrinsic microporosity (PIM) reported by McKeown et al. in 2004 [14]. Due to the conformation of rigid polymer chains, PIMs have numerous interconnected pores smaller than 2 nm. The studies on PIMs are mainly focused on introducing rigid units and changing conformation angles to increase free volume to improve CO$_2$ permeability further. McKeown group reported a highly gas-permeable polymer using spirobifluorene as the central structural unit [15] (Fig. 2(c)). Subsequently, a series of PIMs prepared by Tröger's base were reported with membrane performance approaching 2008 upper bound [16].

Polymeric membranes mentioned above all belong to solution-diffusion transport-based membranes. Facilitated transport membranes with reactive carriers are another kind of high CO$_2$ permeable membranes where CO$_2$ transport is enhanced by interaction with CO$_2$ molecules through reversible reactions. Liquid membrane, ion-exchange membrane, and fixed carrier membrane are three primary forms of facilitated transport membranes. The advantages and disadvantages are listed as follows (See Tables 1 and 2).

The fixed carrier membranes are widely favorable because of their excellent permeability, low cost, and simple fabrication process. As the name implies, the carriers in fixed carrier membranes are covalently bonded and cannot move, so the selectivity of this kind of membrane is not high. Groups, such as –NH$_2$, CO$_2$–, -COO–, are the conventional choice of CO$_2$ carriers. One typical fixed carrier membrane material is polyvinylamine (PVAm) which contains a high density of amine groups. In 2013, Hägg group reported that the pH control could remarkably enhance the PVAm performance for CO$_2$ capture [17]. It was found that the membranes prepared by higher pH casting solution had more amine groups, resulting in faster CO$_2$ transport. The facilitated transport mechanism is shown in Fig. 3.

Since the number of effective carriers plays an important role in membrane performance, different methods have been developed to increase the carrier concentration. In 2013, Qiao et al. reported a PVAm/PDMS/Polyimide (PS) composite membrane [18]. High PIP loading improved the concentration of carriers and effectively reduced the crystallinity of PVAm, bringing improved CO$_2$ permeance of 6500 GPU. Recently, they successfully synthesized metal-induced ordered mesoporous polymers (MMPs) via PVAm, divalent metal ions, and organic linkers [19]. The prepared membranes demonstrated excellent structural stability under humid and dry feed conditions, showing auspicious CO$_2$ capture performance. Recently, Winston Ho group synthesized an ultrahigh molecular weight PVAm by inverse emulsion polymerization, which demonstrated an improved transport performance due to the alleviation of penetration and a higher amine content in the membrane. The PVAm based membrane with 85 wt% piperazine glycinate (PG) loading achieved CO$_2$ permeance of 839 GPU and a CO$_2$/N$_2$ selectivity of 161 at a typical flue gas temperature of 57°C [20]. Furthermore, they have successfully realized the scale-up fabrication of PVAm/PDMS-containing membranes in a roll-to-roll manner using the thin-film coating (TFC) assembly (Fig. 4). A total of >2000 feet long and 14-in. wide membrane with a selective layer thickness of <200 nm was successfully fabricated which showed good agreement with lab-synthesized membranes in CO$_2$/N$_2$ separation [21]. (Gas permeance unit, 1 GPU = $1 \times 10^{-6}$ cm$^3$(STP) cm$^{-2}$ s$^{-1}$ Pa$^{-1}$)

### 2.2. Non-polymeric membranes

Compared with the remarkable progress of numerous polymeric membrane materials, non-polymeric materials such as zeolite, metal-organic framework, carbon molecule sieve have also been in an emerging stage of vigorous development in CO$_2$ separation. Usually, those membranes show better stability than polymeric membranes and thus are the leading candidates for gas mixtures separation, especially under harsh operating conditions.

Zeolites are crystalline microporous aluminosilicate frameworks built of corner-sharing AIO$_4$ or SiO$_4$ tetrahedra, employed in numerous applications such as purification, adsorption, separation, catalysis. Zeolite membranes used for CO$_2$ capture can offer high selectivity because of their well-defined pore structures. Several zeolites such as Na–Y [22], SAPO-34 [23], and MFI [24] membranes have been investigated for post-combustion. Kusakabe and coworkers reported excellent results of Na–Y membrane with CO$_2$/N$_2$ separation factor of 100 at 303 K [22]. Recently, Zhou group prepared defect-free SAPO-34 membranes on tubular alumina substrates [25]. At 243 K, the CO$_2$ permeability reached about 4100 GPU with selectivity about 110 with a CO$_2$/N$_2$ volume ratio of 15/85. Currently, SAPO-34 membranes show excellent separation performance far beyond the polymeric upper bound and demonstrate potential applications in industrial post-combustion.

Porous metal-organic framework (MOF) materials are also prevalent for CO$_2$ separation. In 2016, a high selective IRMOF-1 membrane was reported with CO$_2$ permeance over 600 GPU and CO$_2$/N$_2$ selectivity of 410 with a CO$_2$ mole fraction of 0.874 [26]. Noticeably, the membrane was nonselective when the CO$_2$ mole fraction was smaller than 0.65. Jomekian et al. tried coating a PDMS layer on ZIF-8 film, and the membrane selectivity was successfully improved from 5.7 to 13.5 with a slight CO$_2$ permeance reduction from 1044 GPU to 925 GPU [27]. Some other examples of the best results reported on CO$_2$/N$_2$ separation using zeolites and MOF membranes are listed below.

A carbon molecular sieve (CMS) was prepared with rich and rigid gas permeation channels by pyrolysis of polymer precursors. Koros group has done pioneer works on the control of pyrolysis conditions to explore its relationship and membrane performance [34,35]. Generally, the higher pyrolysis temperature means higher gas permeability and lower selectivity. Besides, the structural properties of precursors may also significantly affect the microstructure of CMS membranes, resulting in different gas separation performances. For example, Hou and coworkers investigated several CMS membranes made of polyimides with varying parts of diamine [36]. In Fig. 5, the CMS membrane with turbostratic carbon was fabricated by pyrolysis of polyimide precursor. The PPD-PMDA with linear structure and low fraction of free volume (FFV) resulted in a dense carbon structure where molecular sieving dominated the gas separation.

<table>
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<tr>
<th>Carriers work form</th>
<th>Advantages</th>
<th>Disadvantages</th>
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<tr>
<td>Liquid membrane</td>
<td>move freely</td>
<td>Excellent performance, Poor stability</td>
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<tr>
<td>Ion-exchange</td>
<td>held by</td>
<td>Favorable stability, Low permeability</td>
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<tr>
<td>membrane</td>
<td>electrostatic forces</td>
<td>high selectivity, Low permeability</td>
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<tr>
<td>Fixed carrier</td>
<td>vibrate in a confined space</td>
<td>High permeability, Moderate selectivity</td>
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While carbon membrane prepared by BDAF-PMDA precursor with helical structure and high FFV showed looser structure accumulation, causing a CO₂ permeability about 3100 Barrer and moderate selectivity 21.8. Except for the category materials mentioned above, some other inorganic materials have also performed well in CO₂/N₂ separation. In 2017, Yu group fabricated an ultrathin graphene oxide (GO)-based hollow fiber membrane using a facile coating process [37]. Under simulated flue gas conditions, it exhibited outstanding separation performance with a CO₂/N₂ selectivity as high as 680.

2.3. Mixed-matrix membranes

Mixed-matrix membranes (MMMs) formed by doping inorganic fillers into a polymer matrix received much attention due to the potential high separation performance and simple fabrication method. In 2013, Long group demonstrated that the CO₂ separation performance could be significantly enhanced by incorporating Mg₂(dobdc) nanocrystals within polyimide membrane [38]. Then, Xin et al. reported SPEEK based MMMs by incorporating PEI-decorated MIL-101(Cr) MOFs [39]. In Fig. 6(a), the PEI was impregnated into MOF fillers via a facile vacuum-assisted method. Compared with MMMs doped unfilled MIL-101(Cr), the CO₂/N₂ selectivity of SPEEK/PEI@MIL-101(Cr) MMMs increased by 102.4%. Our group reported polyether block amide (PEBA) MMMs with Uio-66 NH₂ MOF fillers for CO₂ separation [40]. The amine-functionalized MOF particles showed a stronger affinity with CO₂ molecules and better dispersion in PEBA than unmodified MOFs. The MMMs with 10% Uio-66 NH₂ loading remarkably increased CO₂/N₂ selectivity from 43.2 to 72.2. Later, Yang group prepared MMMs with ionic liquid (IL)-modified ZIF-8 particles [41]. The confinement of ionic liquids in nanocages gave ZIF-8 filler an effective alteration of the molecular sieving properties for CO₂ separation. The IL@ZIF-8 MMMs demonstrated remarkable separation performance that transcend the upper bound of polymeric membranes.

In addition to porous MOF fillers, we reported a kind of MMMs by incorporating layered GO nanosheets for CO₂ capture (Fig. 6(b)) [42]. The GO nanosheets were assembled into regular GO laminates, showing proper channels for CO₂/N₂ separation. Amorphous polymer chains from TEM images surrounded the GO nanosheets, and the d-spacing of the two neighboring nanosheets was about 0.7 nm. This membrane exhibited preferential CO₂ permeability of 100 Barrer and excellent CO₂/N₂ selectivity of 91. Qiao and coworkers prepared a highly permeable aligned montmorillonite (AMT) MMM with a layered structure (Fig. 6(f)) [43]. It was found that the interlayer gaps of AMT were consistent with the orientation of the polymer chains and served as efficient CO₂ transport channels. From Fig. 6(g) and (h), the regular cuboid shape of MTs and highly orientated Na-exchanged MT were observed. This membrane exhibited CO₂ permeance of 800 GPU and good stability over 600 h. Overall, the variety of polymers and fillers such as zeolites, MOFs, MXene, etc., have extensively enriched the MMMs family for CO₂ capture, and some typical MMMs are summarized in Fig. 6(i).

3. Pre-combustion capture

Pre-combustion means removing CO₂ from the fuel before combustion. Pre-combustion is mainly used in the IGCC system, where H₂ and CO₂ can then be separated via a membrane separation system. In the case
Fig. 5. Illustration of CMS membranes derived from different polyimide precursors. Reprinted with permission from ref. 36. Copyright (2021) Elsevier.

Fig. 6. (a) Illustration of the fabrication of PEI@MIL-101(Cr) based MMMs. (b) Schematic representation of GO/PEBA MMM. (c) Digital photograph of free-standing GO/PEBA MMM. (d–e) TEM image of GO/PEBA MMM [42]. (f) AMT/PSf membrane, (g) surface SEM, and (h) AMT/PSf membrane TEM [43]. (i) Robeson 2008 CO₂/N₂ upper bound line [44] and the separation properties of some typical MMMs including Pebax-based MMMs [39, 41, 45–47], PIM-1/ZIF-8 [48], AMT/PSf [44] and PEI/MIL-101 [39]. (b–e) Reproduced from ref. 42, copyright 2014 John Wiley & Sons. (f–h) Reproduced from ref. 43, copyright 2016 John Wiley & Sons.
of pre-combustion, the mixtures of CO₂ and H₂ are at high pressure and high temperature, which provide sufficient driving force for membrane separation. The IGCC process allows the application of two kinds of membranes: H₂-selective and CO₂-selective membranes. In this review, membranes that permeate H₂ are the focus since they are the most available.

3.1. H₂/CO₂ separation membranes

3.1.1. Non-polymeric membranes

Among H₂-selective membranes, metallic membranes are widely investigated because of the high H₂ purity up to 99.99% when using palladium (Pd) as the membrane materials [49]. For Pd-based membranes, the hydrogen separation process is realized by three steps: First, hydrogen is dissociated and chemically adsorbed on the surface to produce atomic hydrogen. Then, atomic hydrogen diffused to the opposite side of the Pd membrane through the metal lattice driven by the partial pressure drop. Finally, atomic hydrogen was recombined into H₂ molecules again, and the H₂ transmission was completed. The H₂ permeability rate is greatly affected by the membrane surface roughness, metal purity, and metallic lattice during this process. Besides, the Pd membrane exhibits improved performance at higher temperatures. Generally, ultra-thin metallic layers were coated on porous inorganic supports to obtain high H₂ flux and reduce costs. Ma group studied the Pd membrane performance with porous stainless substrates under different conditions [50]. The membrane showed high H₂ permeance of 34 m³/m² h bar⁻¹, at 400°C under H₂/H₂O (10–37%) mixed gas conditions. Since the poor H₂ permeance of most Pd membranes, binary and complex alloys of Pd have been proven to be an effective method to improve H₂ permeability. Tong group reported a defect-free Pd–Ag membrane with submicron thickness. The membrane exhibited high separation fluxes of up to 4 mol of H₂/m²·s at 450°C [51]. However, Pd-based membranes are vulnerable in many aspects, including hydrogen embrittlement, the presence of contaminants, limited lifespan, and high membrane costs. These drawbacks have primarily reduced their attractiveness in H₂ separation.

Except for metallic membranes, silica, zeolites, and MOFs have also been investigated for H₂/CO₂ separation. In 1998, Renate and coworkers reported a series of defect-free silica membranes [52]. Silica membrane calcined at 600°C showed H₂ permeance of 1474 GPU at 200°C and H₂/CO₂ selectivity about 71. However, researchers found that silica-based membranes were susceptible to humidity, and thus apparent performance decline was observed when exposed to high water vapor concentrations. This phenomenon was also observed in CMS membranes, where the water vapor blocks the membrane’s pores, decreasing gas permeance. Surprisingly, the cellulose-derived CMS membrane showed excellent stability under high humidity due to its high hydrophilicity [53]. Lei and coworkers recently reported cellulose-based carbon hollow fiber membranes (CHFMs) with ultra-microporous of 3–4 Å for superior H₂ separation [54]. The prepared carbon membranes exhibited apparent asymmetric structural morphology (Fig. 7(a and b)). From Fig. 7(c), an

![Fig. 7. (a–b) Cross-sectional SEM images of CHFM-700. (c) Single-gas permeances of CHFM-850 membrane at 130°C and 2 bar. (d–f) Schematic diagram of the Zn₂(bim)₃ membrane fabrication processes. (g–h) SEM pictures of delaminated MXene nanosheets and membrane surface. (i) Illustration of two adjacent MXene nanosheets. (a–c) Reproduced from ref. 54. Copyright 2021 Nature Publishing Group. (d–f) Reproduced from ref. 59, copyright 2017 John Wiley & Sons. (g–i) Reproduced from ref. 61. Copyright 2021 Nature Publishing Group.](https://example.com/fig7)
explicit cutoff between H₂ and CO₂ was observed. The membrane exhibited not only high separation performance but also good water vapor stability.

Unlike silica and CMS membranes, where microporous structures mainly depend on the calcination or pyrolysis step, the porous properties of zeolites and MOFs materials depend on the connection methods. So far, many zeolites and MOFs materials have been reported, which dramatically expands the candidates for H₂/CO₂ separation. For example, Dong and coworkers reported an on-stream modified MFI zeolite membrane. The membrane obtained H₂ single gas permeance of 555 GPU with H₂/CO₂ selectivity of 17.5 [55]. Caro group fabricated a ZIF-95 membrane which displayed H₂ permeances of about 5700 GPU and H₂/CO₂ selectivity of about 25.7 [56]. Later, Huang et al. used ZIF-95 nanosheets as seeds for ZIF-95 membrane preparation [57]. Although the membrane showed an evident decline in H₂ permeance (~500 GPU), the selectivity was improved to 41.6. In 2014, Yang group adopted Zn(bim)₂ nanosheets as building units to prepare molecular sieving membranes [58]. The prepared membrane achieved H₂ permeance of about 3000 GPU, and selectivity greater than 200, far beyond 2008 polymeric upper bound. Later, they developed a similar Zn(bim)₂ membrane with a static honeycomb-like aperture of 0.29 nm [59]. The detailed membrane fabrication processes were represented in Fig. 7(d–f). The prepared ultrathin membranes showed a remarkable separation performance owing to the size-exclusion effect of Zn(bim)₂ structure.

Besides MOF nanosheets, other 2D materials, including GO and MXene have been reported for H₂/CO₂ separation. In 2016, Our group reported the manipulation of GO-assembled 2D channels by external forces [60]. The prepared membrane with an interlayer height of about 0.4 nm demonstrated higher H₂ permeance and enhanced H₂/CO₂ selectivity. In 2018, Wang group reported lamellar MXene membranes with ordered nanochannels [61]. The membrane exhibited excellent H₂ permeability of more than 2200 Barrer and H₂/CO₂ selectivity over 160, better than most advanced membranes. From Fig. 7(g), the MXene nanosheets were very thin, and the Tyndall scattering effect was visible. The free spacing of the adjacent MXene nanosheets was about 0.35 nm (Fig. 7(h)), serving as an effective molecular sieve for H₂ and CO₂ gas mixtures.

3.1.2. Polymer-based membranes and mixed-matrix membranes

In terms of molecular kinetic sizes (H₂: 0.289 Å, CO₂: 0.33 Å), the diffusivity of H₂ is greater than that of CO₂. However, most polymers show affinity with CO₂, leading to the solubility of H₂ over CO₂ smaller than one. Generally, polymeric membranes with H₂ preferential permeation mainly depend on diffusion selectivity while the CO₂ solubility is inhibited. Because of size-selective affinity, a wide range of glassy polymers such as polysulfone (PSf) and polyimide have shown H₂/CO₂ separation performance. However, they suffer from low permeability (10–50 Barrer) and/or low H₂/CO₂ selectivity (1–10) and the trade-off restriction between permeability and selectivity. MMPMs based on PSf and polyimide have been considered an effective way to achieve a breakthrough in permeance. For example, Leo group incorporated palladium (Pd) nanoparticles into PSf matrix where Pd nanoparticles were stabilized using polyvinylpyrrolidone (PVP). The MMP with 2% Pd loading exhibited high H₂ permeability of 5779 Barrer and satisfactory H₂/CO₂ selectivity of 6.2 [62]. Musselman and coworkers reported (Al) NH₂-MIL-53/V-TEC™ polyimide (PI)-1388 MMPMs H₂/CO₂ mixture separations at high pressure and high temperature. At 300 °C, the MMPMs demonstrated simultaneous enhancement in permeability and selectivity (H₂ permeability: 85 Barrer to 144 Barrer, H₂/CO₂ selectivity: from 4.0 to 5.6) [63].

Polybenzimidazole (PBI) is another promising candidate among those polymer materials due to its high H₂/CO₂ selectivity, superior thermal and moisture stability, and excellent chemical resistance. However, the PBI membrane exhibits low H₂ permeability resulting from dense polymer chain packing. Therefore, strategies to improve H₂ permeability while simultaneously keeping selectivity are required to make it more industrially attractive. It was found that modification of main chain structure could disrupt PBI chain packing and improve H₂ permeability (up to 997.2 Barrer), which was much higher than commercially available poly(2,2-(m-phenylene)-5,5-bifenzimidazole) (m-PBI) (76.81 Barrer) [64]. However, lower H₂/CO₂ selectivity of about 5–7 was also observed versus m-PBI membrane of 23. Yang and coworkers prepared ZIF-8/PBI MMMs with higher solubility and diffusion coefficients [65]. The ZIF-8/PBI membrane with 30 wt% loadings had H₂ permeability of 470.5 Barrer with an H₂/CO₂ separation factor of 26.3. Recently, Lin group demonstrated a new kind of PBI membranes doped with polyprotic acids, which showed unprecedented size-selective ability in H₂/CO₂ separation [66]. In Fig. 8(a), the cross-linking of PBI chains was achieved by transferring protons from H₃PO₄ to imidazole rings of PBI. A series of H₃PO₄ doped PBIs were prepared by immersing PBI membranes in solutions containing H₃PO₄ and methanol. When the molar ratio of PBI and H₃PO₄ was 1:1, the membrane exhibited an H₂/CO₂ separation factor of 140, far beyond the previously known polymeric materials [67].

TR polymers and PIMs have also been widely investigated for pre-combustion. For example, Lee and coworkers prepared different TR poly(benzoxazole-co-amide) membranes derived from monomers [68]. Among those membranes, the best performing membrane exhibited an H₂ permeability of 26.8 Barrer and H₂/CO₂ separation factor of 8.0 at 210 °C. Later, a series of Tröger’s Base based copolyimides were synthesized [69]. Although high H₂ permeabilities (249–667 Barrer) were observed, they all suffered from low H₂/CO₂ selectivity (highest: 2.8).

In 2018, Gascon group reported a facile interfacial polymerization to prepare supported benzenimadazole-linked polymer membranes (Fig. 8(b)) [70]. A brown layer was quickly observed at the liquid interface due to the reaction of amine groups with aldehyde moieties (Fig. 8(d)). The growing film served as a barrier and confined the reaction to defective areas, guiding the formation of a dense and defect-free membrane.

3.2. CO₂/H₂ separation membranes

When H₂ production is of high priority, the CO₂-selective membrane is more favorable because it reduces H₂ loss during the separation process and helps to reach almost 100% CO conversion in an equilibrium-dominated low-temperature water-gas shift reactor. For polymer membranes with CO₂ selectivity, it is essential that the membrane strongly favors CO₂ and minimizes H₂ diffusion. Some polymeric membranes mentioned in CO₂/N₂ separation can also be used in CO₂/H₂ separation. For example, Freeman group investigated the mixed gas permeation of syngas components in PDMS and PTMSP. The CO₂ permeability and CO₂/H₂ selectivity of the PDMS membrane are 3200 Barrer and 3.4, respectively [71]. Bondar and coworkers demonstrated that the PEBA membrane showed higher CO₂/H₂ selectivity when polar group concentration was increased [72]. These high selectivity values may attribute to the large solubility selectivity in favor of CO₂.

As an extension of polymeric membranes, facilitated transport membranes usually demonstrate high CO₂ permeability and excellent CO₂/H₂ selectivity greater than 100 due to the complexion reaction between CO₂ and facilitator. Generally, amino species, polar groups and ionic liquids are popular choices. Winston Ho group has done much work on facilitated transport mechanisms containing amino species. They induced amines in crosslinked poly(vinyl alcohol) and the CO₂ permeability and CO₂/H₂ selectivity of the membrane at 120 °C reached 8200 Barrer and 450, respectively [73]. Recently, they reported amino-containing MMMs incorporating amino-functionalized multi-walled carbon nanotubes (AF-MWNTs) for CO₂/H₂ separation. The MMM showed a selectivity of 205 and a CO₂ permeability of 3196 Barrer at 107 °C and 0.2 MPa. However, reduced transport performance was observed due to the carrier saturation [74].

4. Oxy-fuel combustion

Oxy-fuel combustion is burning hydrocarbon fuel in an environment close to pure oxygen instead of air. It can generate flue gas containing
very high CO2 and water vapor concentrations, facilitating the post-combustion process. Besides, this process can significantly reduce burnout time and NOx emissions, and exhaust gas volumes. Currently, various membranes have been studied for O2/N2 separation.

4.1. Polymer-based membranes and mixed-matrix membranes

For polymer membranes, the solution-diffusion mechanism is dominant in O2/N2 separation [75]. Polysulfone (PSf), polyamide (PA), polyurethane (PU), polydimethylsiloxane (PDMS) and polyimide (PI) have been studied for O2 preferential permeation. Among them, PSf has great potential because of its good mechanical strength and balanced permeability and selectivity. In 2002, Wang and coworkers reported PSf hollow fiber membranes with a dilute silicone coating layer [76]. The prepared membrane exhibited O2 permeance of 20–30 GPU and O2/N2 selectivity of 5–6.5 at 25°C. Another popular O2/N2 separation material is polyimide. Koros and coworkers reported a kind of 6FDA-IPDA PI hollow fiber membrane, which led O2 permeance of 65 GPU and O2/N2 selectivity of 6.3 [77]. Later, Kawakami group reported asymmetric 6FDA-FAP PI membrane with higher O2 permeance of 794 GPU and low O2/N2 selectivity of 5.3 [78]. As to microporous polymers, although PIM-1 shows high O2 permeability, the poor O2/N2 selectivity hinders its practical application [79].

In 2011, Zornoza and coworkers prepared PSf-based MMMs combining MOFs and zeolites fillers. The results indicated that the combination of HKUST-11 and silicalite-1 produced significant O2/N2 selectivity achievements and ZIF-8-PSf MMM exhibits the best O2 permeability and O2/N2 selectivity which are increased by 63% and 80%, respectively [80]. Zhang group reported a composite membrane with PIM and cobalt IL Bmim2Co(NCS)4 exceeding 0.4 wt%, the membrane exhibited O2/N2 separation performance beyond the 2008 Robeson upper bound.

4.2. Microporous inorganic membranes

Since the kinetic diameters of O2 (3.46 Å) and N2 (3.64 Å) molecules are very close, selectivity supported by Knudsen-diffusion can be ignored in microporous membranes. Similar kinetic diameters also bring difficulties for inorganic membranes based on molecular sieving. Compared to the mediocre performance of zeolites and MOFs, CMS membrane performs quite well [82]. For example, Koros group investigated 6FDA/DETDA:DABA(3:2) derived CMS membranes with different pyrolysis temperature [83]. The CMS membrane pyrolyzed at 800°C showed desirable O2 permeability and selectivity of 683 Barrer and 8.0, respectively. Liu et al. doped ZSM-5 nanoparticles into a carbon matrix [84]. A prominent O2 permeability increment from 2.21 to 431 Barrer and a slight O2/N2 rise from 13.8 to 14.4 was observed when ZSM-5 content was 9.1 wt%.

4.3. Mixed ionic-electronic conducting (MIEC) membrane

MIEC membrane is a good choice for pure oxygen separation. Unlike the membranes mentioned above, MIEC membrane is dense membranes with 100% theoretical selectivity to oxygen. In MIEC membrane, oxygen in the form of ions diffuses from one side to the other via oxygen vacancies in the lattice (Fig. 9(a)). Such a permeation process requires the membrane material to have both oxygen ion conductivity and electronic conductivity. Perovskite, with the generic composition ABO3–δ, is the most studied MIEC for oxygen permeable material. The partial substitution of A or B sites can generate oxygen vacancies which can induce the adsorption of oxygen and act as an oxygen carrier to migrate oxygen ions to the other side of the membrane surface (Fig. 9(b and c)) [85].

To improve O2 flux, numerous new perovskite materials have been reported. The main strategies of developing new perovskite membrane materials are focused on the changes in the A or B metal elements and the variations in the stoichiometric coefficient of the composition. For example, Sr, Ba, and La at the A site and transition elements Fe, Cr, and Co at the B site have become the popular choice [86]. However, the application of perovskite membranes was hindered due to poor structure stability, oxygen flux, and high operating temperature. To overcome these shortcomings, much work has been done to design new perovskite materials. For example, our group found that ZrO2 doped SrCo0.4Fe0.6O3−δ, with remarkable structural stability under CO2 or H2 containing atmospheres [87]. Yang et al. successfully synthesized a series of BaCo2Fe1.6O4.8 perovskite membranes. Under a reductive atmosphere, these membranes showed both high permeability and stability [88].
Later, we reported unprecedented perovskite oxyfluoride membranes with F⁻ doping on O-site [89]. It was found that F⁻ could effectively weaken the chemical bonds of the perovskite structure, leading to enhanced oxygen permeability and reduced oxygen permeation activation energy. At low temperatures, the membrane exhibited extraordinary oxygen permeation performance, which completely fulfilled the commercial requirements.

In addition to the characteristics of the perovskite itself, the membrane configuration and preparation method, which determines the specific surface area and mechanical strength, also plays an important role in oxygen flux and structural stability. Researchers have designed many methods to prepare membranes, including multilayer structures with various architectures. Jin et al. first put forward the idea of synergetic shrinkage (i.e., the porous support and film shrink simultaneously in the sintering process) and developed a new method for preparing supported oxygen permeable membrane [90]. This idea was adopted and applied by the internationally famous company Air Products. Recently, our group developed a one-step thermal processing approach for efficient, high-performance perovskite hollow fibers production [91]. Compared with the traditional process, including mixing raw materials, two-step heating, repeated grinding, and phase inversion, the new approach was greatly simplified. Besides, multichannel hollow fiber membranes were designed and fabricated (Fig. 10(b)). Multichannel tubular membrane and hollow fiber membrane provided both high flux and high mechanical strength [92]. The 19-channel hollow fibers showed a mechanical strength of more than 50 N and could directly build an oxygen-permeable device or membrane reactor [93].

5. CO₂ utilization

As the main driver of climate change, CO₂ utilization technologies have recently received increasing attention. CO₂ utilization routes include enhancing oil and gas recovery, chemical conversion, mineralization, etc. Notably, converting CO₂ into fuels is considered the best way in CO₂ utilization. For example, Chen and coworkers reported an effective catalytic reverse water-gas shift (RWGS) process to produce CO with simultaneous CO₂ capture by a ceramic-carbonate dual-phase membrane reactor [94]. A CO₂ conversion of 56.8% and a CO production rate of
2.41 ml min$^{-1}$ cm$^{-2}$ was obtained at 750°C with La$_0.9$Ce$_0.1$NiO$_{3.6}$ catalyst. In addition to CO, methanol is another attractive product of CO$_2$ hydrogenation. Recently, Yu group reported NaA zeolite membrane with sodium ion (Na$^+$)-gated water-conduction nanochannels for CO$_2$ hydrogenation [95]. The in-situ water removal through the membrane continuously promoted the CO$_2$ conversion and methanol yield. Compared with a traditional reactor with copper-zinc-alumina catalysts (CO$_2$ conversion: 23.0%, methanol space-time yield (STY): 339 mg g$_{cat}^{-1}$ hour$^{-1}$), the membrane reactor incorporating the water-conduction membrane greatly boosted CO$_2$ conversion and methanol STY to 61.4% and 809 mg g$_{cat}^{-1}$ hour$^{-1}$, respectively. Followed by this work, Huang group reported a similar bifunctional catalytic membrane reactor with different zeolite and catalytic [96]. The CO$_2$ conversion and methanol selectivity reached 36.1% and 100%, respectively.

A membrane reactor has also been applied in the thermal decomposition of CO$_2$. Here, our group proposed integrating the thermal decomposition of CO$_2$ with the partial oxidation of CH$_4$ (POM) using dense mixed-conducting membrane reactors [97]. Such a membrane reactor could in situ remove oxygen and promote the conversion rate of CO$_2$ to 11%, which was much higher than the original CO$_2$ conversion rate (0.00052%). Then, the lifespan of the membrane reactor was significantly extended by modifying the porous protective layer on the surface of the membrane [98]. We prepared a coupled reaction condition (a porous-dense-porous triple-layer composite membrane) [99]. Due to the catalytic modification layer on the oxygen permeable side and the protective layer on the methane side, the CO$_2$ conversion rate reached 20.58%. It was operated stably for more than 500 h.

Generally, various industries have great potential to use CO$_2$ as a renewable resource rather than isolating it underground or in the oceans. Integrated systems like membrane reactors have provided good solutions for CO$_2$ utilization.

6. Summary and conclusions

In recent decades, people have shown increased interest in carbon capture and utilization. Some typical performance comparison was summarized in Fig. 12. As an alternative technology, membrane development has been in an emerging stage. In this article, the development of materials from post-combustion, pre-combustion, and oxy-fuel combustion is reviewed. Key achievements and challenges of membrane materials on each process are concluded as below:

(1) Post-combustion (CO$_2$/N$_2$): for polymeric materials including PEO-contained membranes, TR membranes and PIMs membranes, high CO$_2$ affinity functional groups and high fractional free volume are preferred in designing polymers based on solution-diffusion mechanism. However, performance degradation caused by plasticization requires special attention and a more detailed investigation for industrial application scenarios. As to facilitated transport membranes, they generally exhibit a CO$_2$/N$_2$ selectivity above upper bound lines, but carrier saturation will cause a significant drop in performance, making it difficult for practical application. When it comes to non-polymeric membranes, some zeolite membranes such as SAPO-34 membrane have shown good mechanical stability and high separation performance beyond the upper bound. Considering the scale-up preparation difficulty and structure vulnerability to water vapor, it is still a long way from laboratory to industry application. MMMs are considered an effective way to approach industrial goals, but the dilemma encountered by polymers and inorganic fillers also needs to be solved in MMMs.

In 2010, Merkel gave a Roselson-type trade-off plot of CO$_2$/N$_2$ selectivity versus CO$_2$ permeance for membranes reported in the literature as useful for flue gas treatment [101]. Membranes with CO$_2$ permeance over 1000 GPU and CO$_2$/N$_2$ selectivity over 20 are expected to be economically feasible in industrial applications. Some membranes have successfully reached or surpassed this goal. In this condition, problems that may be encountered in more realistic scenarios need to be considered to further clarify the choice of materials. For example, flue gas generated from coal-fired power plants and CH$_4$-fired power plants are different in composition. The former contains a high concentration of CO$_2$ about 15% and a variety of contaminants including fly ash, SO$_2$ and NOx. So, electrostatic precipitator and desulfurization pretreatment process are needed and the resistance of membrane materials to sulfides also needs to be considered and evaluated. The latter contains a relatively low concentration of CO$_2$ (<5%) which is bound to impose higher permeance demand on membrane materials. Besides, a high concentration of O$_2$ about 12% puts forward different requirements on the stability of membranes. In conclusion, the development of high performed membrane materials is a prerequisite for industrial application of membrane process, and the real industrialization still need more pilot plant projects and more detailed evaluation.

(2) Pre-combustion: for H$_2$/CO$_2$ separation: metallic membranes, especially Pd-based membranes, showed great attraction of high-purity H$_2$ production. The disadvantages of hydrogen embrittlement, performance degradation to sulfur, limited lifespan, and high membrane costs largely restricted Pd-based membranes in large-scale H$_2$ separation. Some new membrane materials such as GO, MOFs and MXenes have demonstrated high H$_2$/CO$_2$ separation performance, far beyond the upper bound line. However, the repeatability and industrial reliability still need to be investigated. As to polymeric membranes, glassy polymers with high FFV and mechanical stability demonstrate good H$_2$ permeability, but low-temperature operation requirements and poor H$_2$/CO$_2$ selectivity make them less attractive. Although PBIs exhibit relatively high H$_2$/CO$_2$ selectivity, the H$_2$ permeability is inferior because of polymer chain packing. Since PBI shows increased permeance at high temperatures, PBI-based membranes and mixed-matrix membranes (e.g., MMMs) can be suitable for high-temperature syngas purification. For CO$_2$/H$_2$ separation, facilitated transport membranes have shown high CO$_2$/H$_2$ selectivity at a temperature over 100°C and are not restricted by the upper bound line, which is very attractive in pre-combustion. It is worth noting that no matter what kind of membrane materials, it is necessary to consider the competitive permeability and possible adverse effects of minor components such as CO and steam.

(3) Oxy-fuel combustion (O$_2$/N$_2$): polymeric membranes, such as PSf and PI are widely used due to their excellent stability. Since the low permeability of polymeric membranes, the permeability...
Innumerable existing CO₂ capture technologies, membrane technology does not occupy a dominant position. For example, amine adsorption has been proved successful in treating industrial gas streams and is considered a leading candidate for post-combustion. However, research shows that amine adsorption is costly and energy-intensive to meet the U.S. Department of Energy target. The CO₂ separation in the IGCC plant (pre-combustion) is more accessible and cheaper due to the high CO₂ concentration and operating pressure. Although physical absorption is a promising process and some commercial product such as Selexol has been widely used, the regeneration of sorbents by temperature swing or pressure swing is energy-intensive. Besides, the increased viscosity of absorbent solvent after interaction with CO₂ is also a problem. In the short term, membrane separation is more suitable to couple with other technologies such as adsorption and adsorption and works as an auxiliary process. But from a long-term perspective, the membrane process has a significant opportunity to be an environmentally friendly, energy-saving and low-cost separation technology with the breakthrough in membrane performance and the development in membrane technologies.

In conclusion, membrane performance is an essential parameter of the membrane separation process, which is greatly affected by membrane materials, manufacturing process and operating conditions. We noticed that most of the existing studies on CO₂ capture membrane materials are focused on improving performance and lacking sufficient attention to other vital requirements. From the perspective of industrial application, membrane materials should be evaluated under complicated and stringent separation situations in the real world. Moreover, the feasibility and competitiveness of the membrane process hinge not only on membrane performance but also on membrane cost, efficient integration, and process intensification with other facility systems, which requires deeper collaboration between power plants and membrane developers.

Finally, CO₂ utilization related to membranes is briefly summarized, and recent studies on membrane reactors for CO₂ hydrogeneration are highlighted. Although there is still a massive gap between laboratory research and industrial application. The development of new materials and the progress of material properties will bring more vitality to membrane separation. Overall, membrane technology is considered a promising technology for carbon capture and utilization, and more efforts should be made to promote the industrial application of the membrane process.

**Declarations of competing interest**

None.

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**References**


Guining Chen received her bachelor degree in chemical engineering and technology from Nanjing Tech University in 2017. She then joined Prof. Wanqin Jin’s research group in the same year in Nanjing Tech University. She is now a Ph.D. candidate in the Department of Chemical engineering. Her research is focused on the metal-organic framework membranes for gas separation.