



Review

Origins and Evolution of Inorganic-Based and MOF-Based Mixed-Matrix Membranes for Gas Separations

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Abstract: Gas separation for industrial, energy, and environmental applications requires low energy consumption and small footprint technology to minimize operating and capital costs for the processing of large volumes of gases. Among the separation methods currently being used, like distillation, amine scrubbing, and pressure and temperature swing adsorption, membrane-based gas separation has the potential to meet these demands. The key component, the membrane, must then be engineered to allow for high gas flux, high selectivity, and chemical and mechanical stability at the operating conditions of feed composition, pressure, and temperature. Among the new type of membranes studied that show promising results are the inorganic-based and the metal-organic framework-based mixed-matrix membranes (MOF-MMMs). A MOF is a unique material that offers the possibility of tuning the porosity of a membrane by introducing diffusional channels and forming a compatible interface with the polymer. This review details the origins of these membranes and their evolution since the first inorganic/polymer and MOF/polymer MMMs were reported in the open literature. The most significant advancements made in terms of materials, properties, and testing conditions are described in a chronological fashion.

Keywords: gas separations; membranes; mixed-matrix membrane; MMM; zeolite; molecular sieves; metal-organic framework; MOF; ZIF; MOF-MMM

1. Introduction

Gas separations are important processes for a variety of applications. In the petrochemical, natural gas, and fuel industry [1,2], for example, the purification of commodity gases, like propane, propylene, hydrogen, methane, and ammonia, is vital for the production of other goods like fuels (e.g., propane, hydrogen, methane) [3–11], polymers (e.g., polypropylene from propylene) [12–15], and fertilizers (e.g., ammonium salts) [16,17]. Recently, environmental applications for gas separations have also been explored mainly for the capture and sequestration of carbon dioxide from the burning and gasification of coal in the generation of electricity and the production of hydrogen [18–20]. Another environmental application for gas separations targets the separation of oxygen from nitrogen (air separation) for the oxycombustion of coal, aiming to reduce the amount of carbon dioxide emitted. Owing to the physical properties of the gases to be separated (e.g., kinetic diameter, critical temperatures, shape), shown in Table 1, the processes designed for their separation (e.g., amine scrubbing, distillation, pressure and temperature swing adsorption) could become expensive due to the amount of energy spent in the separation [21], or even become not viable due to the lack of an efficient and cost effective technology [14,22].

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Gas	Kinetic Diameter (Å)	Critical Temperature (°C)	Critical Pressure (Bar)	Vapor Pressure (Bar at (°C))		
C_3H_6	4.5	91.75	45.55	8.4 (21.1)		
C_3H_8	4.3	96.74	42.51	8.0 (21.0)		
CH_4	3.8	-82.59	45.99	-		
CO	3.76	-140.45	34.94	-		
N_2	3.64	-146.95	33.51	-		
O_2	3.46	-118.55	49.7	-		
CO_2	3.3	30.98	73.0	60 (22.4)		
H_2	2.89	-232.60	12.69	0.025 (21.0)		
H_2O	2.65	373.80	217.7	8.76 (21.0)		
NH_3	2.6	132.25	113.3	-		
He	2.6	-267.96	2.24	-		

Table 1. Physical properties of permanent gases and organic vapors.

Therefore, the search for other separation methods has extended to the use of membranes in which the separation is afforded by the differential in concentration of the gases between the two sides of the membrane (Figure 1). This difference in concentration is typically expressed, and in practice is carried out, in terms of a differential pressure between the high pressure side (P_{feed}) and the low pressure side ($P_{permeate}$) of the membrane.

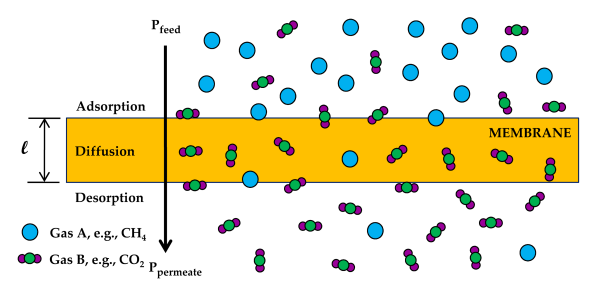


Figure 1. Gas transport in polymer membranes according to the solution-diffusion model showing the adsorption, diffusion, and desorption of the gas molecules through a membrane with thickness *l*.

Several mathematical expressions have been developed to describe gas transport in different types of membranes. For example, in a dense polymer membrane, gas transport is described by the solution-diffusion model (1) that combines the solubility (S) and diffusivity (D) properties of the gases in the membrane to calculate their permeability (P) [23–28]. A good discussion of how to calculate diffusivity and the interpretation of the units for permeability can be found in the work of H. Yasuda [29].

$$P = S \times D, \tag{1}$$

In this model, the gases are adsorbed on the high-pressure side of the membrane, diffuse through the membrane with thickness l, and then desorb from the low-pressure side of the membrane. For these type of membranes, it can be inferred that to increase the selectivity ($\alpha_{A/B} = P_A/P_B$) of a membrane for gas A, the properties of the polymer membrane need to be modified to increase the diffusivity, the solubility, or both properties of gas A in the membrane. The strategy to be adopted depends largely on the physical properties of the gases. For example, if H_2 is to be separated from CO_2 , keeping CO_2 in the retentate side, then the best strategy is to increase the diffusivity of H_2 in the

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membrane, taking advantage of its small size (2.89 Å) and low critical temperature (-232.60 °C) (Table 1). To further improve this strategy, eliminating functional groups from the polymer that adsorb CO₂ could reduce the solubility of CO₂ in the polymer.

The potential application of membranes to gas separations has become an important technological advance in recent years owing to their reduced energy consumption and ease of operation when compared to traditional separation processes (e.g., cryogenic distillation, pressure and temperature swing adsorption, amine absorption, etc.), which are typically energy intensive processes. The simplicity of a membrane-based gas separation plant results in a footprint smaller than that required for other separation processes, making it an attractive alternative. A variety of industrial, energy, and environmental membrane-based gas separation applications have been targeted, some already at the industrial scale. Separations such as H_2/N_2 , H_2/CO_2 , H_2/H_2 , CH_4/CO_2 , N_2/CO_2 , O_2/N_2 , $NH_3/H_2/N_2$, N_2/CH_4 , C_3H_6/C_3H_8 [11], C_3H_8/N_2 [5], $C_3H_6/C_3H_8/N_2$ [12], and C_2H_4/C_2H_6 [8,9], and other organic vapors [30-50] are some examples of where membranes could be employed to provide a cost effective separation process [51,52]. Owing to the nature of membranes as barrier materials, the operating conditions required in industry to separate a particular gas mixture impose constraints and challenges on the materials and manufacturing processes required for a membrane to be successfully used in industry. These limitations, however, open the possibility for improvements in the preparation and selection of materials for the fabrication of membranes resulting in a variety of membrane types as shown in Figure 2.

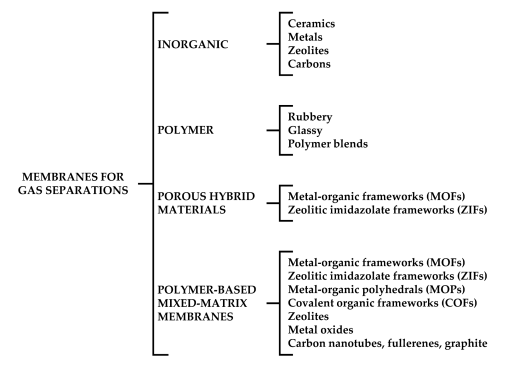


Figure 2. Types of membranes for gas separations and the materials used for their fabrication.

Inorganic membranes have been investigated extensively due to their high selectivity and flux. Unfortunately, these membranes are difficult to fabricate and are prone to mechanical failure. Polymer membranes, on the other hand, are more processable into different geometries (e.g., flat, spiral, hollow fiber) than inorganic membranes, but offer limited selectivity and flux. The advent of porous hybrid materials or MOFs presented a unique opportunity to tune the separation properties of the membranes by chemically modifying the organic component of the MOF by incorporating functional or bulky groups in the walls of the MOFs. This enabled control of the pore aperture of the MOF to perform molecular sieving. A pure MOF membrane, however, would still need a support

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on which to crystallize to form a continuous layer and, as in the case of molecular sieve membranes, the fabrication process is complicated and sensitive to synthesis conditions. A new type of membranes with the potential of producing a breakthrough is the mixed-matrix matrix membranes or MMMs. This membrane combines a polymer with different types of additives (generally one of the materials listed in Figure 2) with the aim of marrying the processability of the polymer matrix with the separation properties of the additive. These membranes simplify the fabrication process because they do not require the formation of a continuous layer of the additive to perform the separation. MMMs also allow for the testing of materials that cannot form continuous layers or are too brittle to form mechanically stable films. Although MMMs can be fabricated into different membrane geometries (e.g., flat, spiral, tubular, hollow fiber), their preparation and effectiveness is limited to the compatibility of the polymer and the additive. Early attempts to introduce porous inorganic additives (mostly zeolites) resulted in the formation of non-selective voids at the additive/polymer interface that degraded the separation properties of the membranes. Another problem this type of membrane faces is the limited loading of the additive in the matrix that can be achieved before agglomeration, non-selective voids (i.e., sieve in a cage), or defects are present in the membrane (Figure 3) [53,54]. Therefore, the combination of MOFs and polymers for the preparation of MMMs for gas separations has resulted in a natural step aiming to exploit the organic character of MOFs to improve interfacial compatibility.

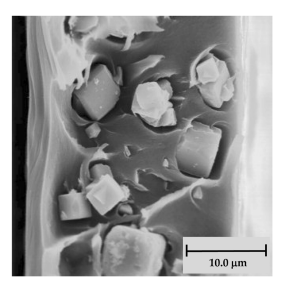


Figure 3. Non-selective voids at the zeolite 4A/UltemTM mixed-matrix membranes (MMM) interface formed by incompatibilities between the inorganic and organic materials (sieve in a cage) [53]. Adapted with permission from [53]. Copyright, 2007, Wiley.

Since a MMM is a blend of inorganic or hybrid organic-inorganic particles (additive) with a polymer, the fabrication techniques are similar to those of conventional polymeric membranes. First, by using one of the mixing procedures shown in Figure 4, a homogenous solution of the additive and the polymer is prepared [55]. To prepare the membranes, five fabrication techniques have been used (Figure 5) [56]: solution-casting, dry phase inversion, wet phase inversion, dry/wet phase inversion, and thermally induced phase separation [57–60]. The simplest and most common technique is the solution-casting method for small scale and laboratory membrane fabrications. In this technique, the dope solution containing the inorganic additive is evenly spread on a uniform substrate with the aid of a knife or a doctor blade to form a uniform film. Then, the solvent (s) is evaporated to obtain a thin film. This simple technique can be used to obtain crystalline and isotropic membranes. The phase-inversion methods are commonly used for the preparation of asymmetric membranes (e.g., hollow fibers). In the dry phase inversion technique, the membrane is exposed to air during the evaporation of the more volatile solvent (vs) whereas in the wet phase inversion the membrane is

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directly immersed in a non-solvent (ns) coagulant as a result of having the spinneret or casting knife immersed in the coagulation bath. In the dry/wet phase inversion technique or the Loeb-Sourirajan technique, the membranes are exposed to air to form the selective layer first and then immersed in a coagulation bath to form the support for the membrane [56]. In the thermally induced phase separation technique, a phase inversion is induced by lowering the temperature of the solution. For this technique to work, the solvent should be miscible with the polymer at high temperatures but not at low temperatures.

This review describes the initial work in the preparation of MMMs and the evolution and innovations of MOF-based MMMs since the first MOF MMM was reported in the open literature. Advances and new theories to understand the interfacial and gas transport properties of MOF MMMs are presented along with their gas separation properties.

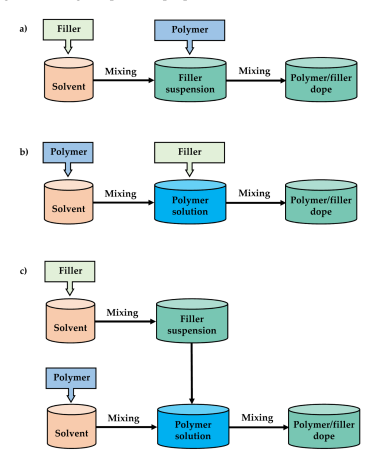


Figure 4. Methods for MMM solution preparation: (a) Polymer added to additive dispersion; (b) additive added to polymer solution; (c) mixing of additive suspension and polymer solution. Adapted with permission from [55]. Copyright, 2010, Elsevier.

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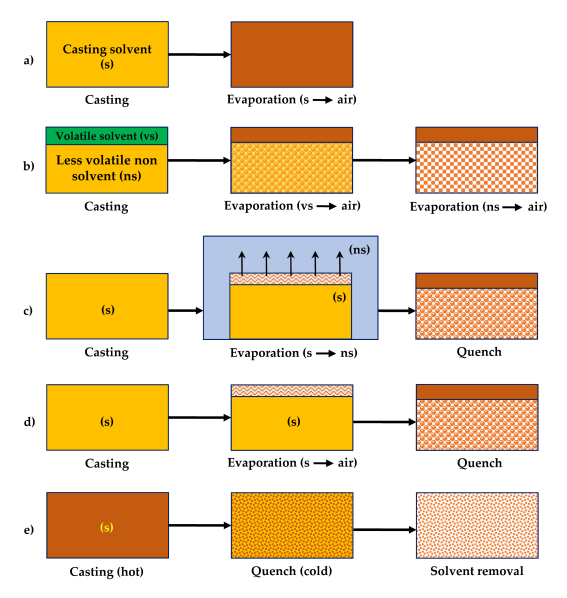


Figure 5. Techniques for membrane fabrication: (a) solution casting; (b) dry phase inversion; (c) wet phase inversion; (d) dry/wet phase inversion; and (e) thermally induced phase separation [56].

2. Origins of Inorganic-Based and MOF-Based MMMs

2.1. Membrane History

The early history of membrane development was well documented in the dedicated volume to the history of membrane science in volume 100 of the Journal of Membrane Science in 1995 [61] and in the work of Bödderek [62,63] about the history of membranes. In these articles, the most significant contributions to the development of membranes from 1748 to 1917 have been compiled and in some cases translated.

The earliest observations of the properties of a membrane can be traced back to experiments with pig bladders that covered flasks reported by J. A. (Abbé) Nollet in 1748 [64,65] and the observations of de la Hire in 1688 and R. A. de Reaumur in 1714 mentioned in Nollet's work. In Nollet's own words, "I shall end this report with a description of a phenomenon that I discovered by chance, and which at first struck me as being as particular as the lowering of mercury to below its own level", the permeation phenomenon was discovered by chance when Nollet observed that pig bladders allowed water to permeate into a flask when it was immersed in a water bath. Through a series of experiments with water and

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ethanol, he observed that the bladder only allowed water to permeate but not ethanol. Later on, in 1827, R. J. H. Dutrochet used the term osmosis to describe the diffusion of liquids (endosmosis and exosmosis) across a permeable partition (membrane) that he observed. In one experiment in which pure water and albumen were separated by a membrane, Dutrochet observed that water crossed the membrane and mixed with the albumen with which it would not mix in the absence of the membrane [66,67]. Dutrochet attributed this *endosmotic / exosmotic phenomenon* to a phenomenon other than the affinity between the liquids since he previously observed that albumen and pure water do not mix when water is poured in a glass containing albumen. Based on the observations of Becquerel on the generation of electricity when liquids are in contact with solid materials, Dutrochet attributed the phenomenon of osmosis to electricity and rationalized that the contact of the liquids with the membrane generated electricity, " ... although I admit that I did not succeed in obtaining a reading on the galvanometer . . . " Dutrochet stated. As a side note, Dutrochet used endosmosis and exosmosis in reference to the flows that occur in hollow organs made of organic tissue [66,67]. Then, T. Graham, in 1829, described briefly the concept of the solution-diffusion mechanism of gas transport across a pig bladder [23,24]. T. Graham observed that when a partially filled bladder with coal gas (methane) or air was enclosed in an atmosphere of carbonic acid (CO2) over water, the bladder would expand significantly. This phenomenon, however, was not observed when the bladder was partially filled with methane and enclosed in an atmosphere of air over water. T. Graham then theorized that CO₂ was solubilized in water in the outer sides of the capillary canals of the bladder in contact with water and then permeated through the canals to later rise into the air or methane in the bladder, this may have been the basis for his solution-diffusion model to explain gas transport in polymer membranes he stated years later in 1866. Two years later in 1831, J. K. Mitchell described the different rates of permeation of several gases through animal and rubber membranes [68,69]. His initial observations came from floating rubber balloons filled with hydrogen that descended after a few days. This led him to believe that the gas escaped the balloons by permeating through the dense walls of rubber. With this knowledge, he performed permeability experiments with several gases and sorted the gases according to their relative facility of transmission: $NH_3 > sulphuretted hydrogen (H_2S) > cyanogen (CN_2) >$ $carbonic\ acid\ (CO_2) > N_2O > arsenuretted\ hydrogen\ (AsH_3) > olefiant\ gas\ (C_2H_4) > H_2 > O_2 > carbonic\ oxide$ $(CO) > N_2$, an impressive feat considering the toxicity and flammability of the gases tested, how the apparatuses were assembled, and the conditions at which the experiments were performed (Figure 6).

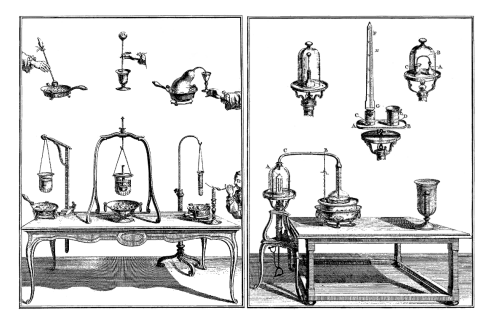


Figure 6. Nollet's illustrations of experimental setups and apparatuses for the study of water and ethanol permeation in pig bladders [64].

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Another two years passed, and T. Graham in 1833, published his work "On the diffusion of gases" in which he states his famous law of the diffusion of gases [70–73]. Working on the observations made by Doebereiner on the escape of hydrogen through a fissure or crack in glass receivers, Graham performed experiments to measure the rate of mixing of hydrogen and other gases with air when the gas diffused through a plug made of plaster, from which he was able to deduce that the rates at which gases diffuse are inversely proportional to the square root of their densities. The next major development came in 1855 when A. Fick published his work on the diffusion of salts in water that led him to develop a fundamental law to describe diffusion (Fick's diffusion law) [74–76]. Although Fick developed this law, his work followed Graham's extensive work on the diffusion of salts in water that was published years before, in Fick's own words: "A few years ago Graham published an extensive investigation on the diffusion of salts in water . . . It appears to me a matter of great regret, however, that in such an exceedingly valuable and extensive investigation, the development of a fundamental law, for the operation of diffusion in a single element of space, was neglected, and I have therefore endeavoured to supply this omission." Additional important progress in membrane science came a few years later, in 1866, from T. Graham's work, "On the absorption and dialytic separation of gases by colloid septa" [25-27]. Graham performed permeability tests (penetration of rubber by gases) with several gases using a simple glass tube 1 inch in diameter capped on one side with a thin plate of stucco that acted as a porous support for a rubber membrane that was placed and secured on top of it. The apparatus (diffusiometer) was then filled with mercury and inverted to create vacuum, and the membrane was exposed to air or other gases (e.g., H₂, CO₂, O₂, N₂, CH₄, CO). From the rates of the depression of the mercury column, T. Graham was able to postulate that the gases had to first condense or liquefy in the membrane due to chemical affinity with the membrane, diffuse, and later be converted into the gaseous form at the vacuum side of the membrane, the very solution-diffusion mechanism that is still used today to explain gas transport in polymer membranes. In the following years, more important developments were made by S. Wroblewski, who in 1879 studied the solubilities of gases in rubber [77], by J. H. van't Hoff's work in 1887 that introduced the idea of semipermeable membranes [78–81], and by S. L. Bigelow and A. Gemberling in 1907 with the preparation of synthetic "collodion" membranes made of pyroxylin (nitrocellulose) [82,83], the first non-rubber artificial membrane. In 1909, continuing on the works of A. Kundt [84], E. Warburg [85], and C. Christiansen [86] on the flow of gases through very narrow channels, M.H.C. Knudsen stated that viscous or molecular flow is achieved depending on the length of the mean free path of the gas and the dimensions of the channels through which the gas diffuses [87,88]. Another significant advance was made by F. G. Donnan in 1911 with his observations on the diffusion of ions in solution (water solutions of Congo Red sodium salt and sodium chloride) through parchment paper that gave rise to ion exchange membranes [89,90] and the development of electrodialysis. An important development in the area of membranes, specifically for MMMs, was made by F. Steinitzer in 1912 with his studies of the adsorption and permeation of CO₂ in rubbers compounded with mineral matter [91], probably the first gas permeation work with mixed-matrix membranes. Five years later, in 1917, P. A. Kober introduced the concept of pervaporation to describe the evaporation of water through a membrane upon the observation made by his assistant, C. W. Eberlein, that a liquid in a sealed "collodion bag" had evaporated [92,93]. Later on, in 1920, J. D. Edwards and S. F. Pickering concluded that the permeability (flux) of any gas was proportional to its partial pressure and that rubber membranes age with time and exhibit reduced permeabilities to gases [94]. This surprising early detection of membrane aging came from their gas permeation studies that compared permeabilities of fresh rubber films with films stored for up to 1 year. Although chemical aging along with physical aging may have also played a significant role in the increased brittleness and reduced permeability of their membranes, the association of decreased permeability to membrane aging was a significant observation that is still used these days to explain permeability reductions in membranes. A more rigorous analysis of the cause and effects of membrane aging would come 40 years later from the works of A. J. Kovacs (1963) [95], K. H. Illers (1969) [96], S. E. B. Petrie (1972, 1976) [97,98], L. C. E. Struik (1978) [99], and M. R. Tant (1981) [100,101] where it is extensively described that the properties of

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a polymer in the glassy state (e.g., permeability, free volume, enthalpy, entropy, mechanical strength) change over time and strongly depend on the thermal history of the polymer (e.g., finite rate of cooling from drying or annealing temperatures to glassy state). The result of the thermal treatment is the quenching of the polymer into a non-equilibrium conformation due a decrease in molecular mobility during the transition to the glassy state in the time frame of cooling. The effects of membrane aging on gas permeability reduction could be significant as seen in Figure 7 where CO₂ permeability in PVAc at 10 °C and 0.13 atm was measured at different time intervals for 20 days [102]. An interesting critical review about membrane aging that expands and complements this topic was published by D. Cangialosi, et al. in 2013 [103]. In this review the authors list the methods used to monitor physical aging and probe polymer segmental mobility and discuss the variations in the degree of aging by the presence of inorganic fillers in polymer nanocomposites, among other topics related to membrane aging.

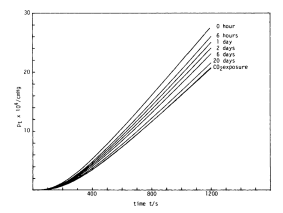


Figure 7. Time dependence of physical aging on the permeability of CO_2 in polyvinyl acetate (PVAc) at 10 °C and 0.13 bar. Reproduced with permission from [102]. Copyright, 1985, Wiley.

Back in 1920, H. A. Daynes introduced the concept of "time lag" (2) to express the time it takes for gases to achieve steady state (Figure 8) when permeating through a rubber membrane of thickness *l* [104]. Note that the *y* axis in Figure 8 represents the response (deflexion) of a galvanometer that measured the current from a katharometer (TCD detector) that was used to detect the flow of gases (hydrogen in Daynes' experiments) in the permeate side of the membrane. The aim was to separate the solubility component from diffusivity using permeability data. By relating the time lag (L) to the diffusivity (K), he succeeded in calculating the solubility and diffusivity constants from permeability data.

$$L = l^2 / 6K, \tag{2}$$

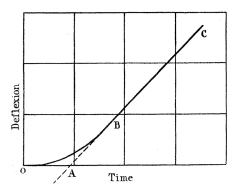


Figure 8. Dayne's time lag (OA) measurement from permeability data at the steady state (BC) in a rubber polymer [104].

The chronological listing of these advances and other significant progress in the field of membrane science is shown in Figure 10 at the end of this section.

2.2. Mixed-Matrix Membrane Origins

One of the first references of gas permeation in membranes that contained additives can be traced back to F. Steinitzer in 1912 with his work on CO₂ adsorption and permeation through rubber materials containing mineral matter [91]. Although the membranes Steinitzer used were not identified as MMMs, their composition, up to 58.60% mineral matter in rubber, resembles what is known today as an MMM. Subsequent work by C.S. Venable, et al. in 1922 on the solubility of gases in rubber containing ZnO and sulfur provided more accurate information from direct measurements of gas solubility [105]. Their work was motivated by the rapid development of rubberized balloon fabric that in those days led to the study of the factors influencing the gas transport of gases in rubber. In 1929, T. Ohya, et al. performed a more extensive study on the permeability of H_2 through compounded rubber looking for a rubber/additive combination that would reduce the diffusion of H₂ [106]. In his work, Ohya added ZnO, clays, carbon black, and softeners to rubber and performed permeability tests that yielded unwanted results: H₂ diffused faster in the compounded rubber than in pure rubber. Then, in 1931, V.N. Morris performed similar experiments with air [107]. His goal was to study the effects of several fabrication parameters (milling, thickness, stretching, nature of crude rubber, vulcanization temperature, and the addition of fillers to rubber) on the diffusion of air through rubber with the potential application of inner tubes for car wheels. Morris found results contradictory to Ohya's. When adding ZnO to rubber, his results showed a decrease in air's permeability through the membrane as the loading of ZnO increased from 5% to 20%. In an effort to explain the discrepancies between his results and Ohya's results when pigments were added to rubber, Morris introduced the possibility that aggregation of the additive could be making the membrane thinner in some regions thus increasing the permeability of the membrane. Extending on previous works on gas solubility in filled rubbers, G. J. Van Amerongen in 1955 performed an extensive study on the effects of fillers in gas permeation and sorption aiming to reduce the permeability of gases in rubber [108]. His findings indicated that lamellar fillers (e.g., mica, graphite, bentonite, alumina) were better in slowing the diffusion of gases than ordinary fillers (e.g., carbon blacks, iron oxide, barium sulfate). By 1959, N. A. Melikhova, et al. performed a series of nitrogen permeability experiments with natural and synthetic rubbers (e.g., sodium-polybutadiene, butadiene-styrene, butadiene-acrylonitrile) filled with minerals (e.g., ground mica, aluminum powder, silica, lamp black, channel black, kaolin) with loadings up to 20% by volume [109]. From permeability and time lag data, diffusivity and solubility values were calculated for each membrane, aiming at obtaining enough information to elucidate the nature of the interaction of synthetic rubbers with fillers and to estimate their effects on gas permeation. Results indicated that all membranes exhibited a reduction in permeability as the filler content increased. The results also indicated that the interactions were more complex than expected and depend on several factors, the volume of the filler in the membrane, the strength of the filler/rubber interactions, the wettability of the fillers by the rubber, and the shape, size, and orientation of the fillers. A key conclusion is that the filler should have a strong interaction with the rubber in order to control the permeation of gases. In 1971, G. Christen, et al. made significant progress in membrane preparation by introducing zeolites in polymers as was described in his patent [110]. G. Christen prepared mordenite/PDMS heterogeneous membranes (mixed-matrix membranes) and tested them in O_2/N_2 separations achieving a selectivity of 3.6 (Knudsen selectivity = 0.93). This is probably one of the first membranes containing porous fillers reported in the open literature that stimulated the development of zeolite-based MMMs for the next 45 years. By 1973, D. R. Paul and D. R. Kemp recognized the effect that zeolites would have in gas adsorption in zeolite/polymer membranes by immobilizing gas molecules in the membrane and therefore increasing the time lag [111,112]. By adapting their work on gas diffusion, using Henry's law to describe the adsorption of mobile species, and utilizing Langmuir sorption to describe sorption of the immobile species, they developed a modified time lag

model for gas transport in the transient state (i.e., before reaching the steady state), useful for the determination of diffusivity in zeolite/polymer membranes and in barrier membranes for coatings and packaging materials [113]. In 1987, H. J. C te Hennepe, et al. reported on pervaporation experiments with silicalite/silicone rubber membranes with loadings up to 60% (w/w) [114]. In this work, silicalite was used as an alcohol-selective sieve to improve the membrane's alcohol (e.g., methanol, ethanol, propanol) separation from water, achieving R-OH/H₂O selectivities of 38.3 with increasing silicalite loadings. The conclusion was that the zeolite particles were well coated by the polymer, and as the zeolite loading increased, the water molecules encountered a more tortuous path since they had to diffuse around the zeolites. A similar work was reported in 1988 by S. Kulprathipanja, et al. in patents for the separation of monosaccharides from polysaccharides in water solution using zeolite/cellulose acetate membranes [115,116]. This is one of the first reports that describes these membranes as mixed-matrix membranes. In a second patent, the utilization of this type of membrane (i.e., zeolite/polymer) was extended to fluids, either liquids or gases. The new patent also encompassed a larger variety of polymers (e.g., polysulfones, polycarbonates, polyethers, and polyamides) and additives (e.g., inorganic carbons, ion-exchange resins, silicalites, and aluminosilicates). In 1991, L. M. Robeson proposed an empirical limit on the permeability-selectivity tradeoff he observed when plotting permeability and selectivity data of several polymer membranes in a logarithmic plot [117,118]. Robeson realized that all polymer membranes lay below an imaginary bound that became known as Robeson's upper bound against which the performances of membranes are compared (Figure 9). By 2000, Mahajan, et al. addressed the problems encountered with the dispersion of zeolites in polymers [119]. Starting with the priming of zeolites through the addition of a small amount of the polymer solution to disperse the zeolites, the authors in this work suggest using polymers with selectivity-permeability properties close to the upper bound to match the transport properties of the materials and to promote polymer adsorption onto the surface of the sieve in order to achieve maximum performance. The use of zeolites in MMMs for gas separations proved to be promising but the low affinity of the polymer for the inorganic additive motivated the search for additives that were more compatible with the polymer matrix. The search for these materials lead to the dawn of the MOF-based mixed-matrix membranes.

In 2004, H. Yehia, et al. reported the first work on the use of porous hybrid materials (MOFs) in mixed-matrix membranes for gas separations using a copper-based MOF and a rubbery polymer (PAET) [120]. This MMM was loaded with up to 30% (w/w) MOF crystals and was tested for CH₄/CO₂ separations with promising results. A year later, in 2005, Li, et al. studied the interactions between porous additives and polymers and reported on their gas permeation experiments with zeolites 3A, 4A, 5A/PSf MMMs [121]. They concluded that polymer rigidification and partial pore blockage of the zeolite by the polymer chains occurred on the surface of the zeolite and resulted in a decrease in gas permeation with increasing zeolite loading. By 2006, A. Car, et al. prepared Cu(BTC)₂ and Mn(HCOO)₂ MOFs that were used in conjunction with PSf and PDMS in the preparation of MMMs [122]. They reported mixed results for permeability and selectivity. The same year, E. V. Perez, et al. incorporated for the first time nanocrystals of a MOF (MOF-5) into Matrimid[®], a polyimide with high T_g [123]. Their MMMs showed improved gas permeation with increased MOF loadings.

Three years later, in 2009, E. V. Perez, et al. reported on their extensive work on MOF-based MMMs for gas separations showing improved dispersion of MOF nanocrystals and strong interactions between the MOF and the polymer that were attributed to the presence of organic components in the MOF [124–126]. Gas transport in these MMMs improved with increased loadings, bringing the membranes to or above the upper bound. In 2010, T.-H. Bae, et al. prepared the first ZIF-based MMM incorporating ZIF-90 into UltemTM, Matrimid[®], and 6FDA-DAM [127]. A few years later, in 2015, E. V. Perez, et al. reported that they had successfully tested (Al) NH₂-MIL-53/VTECTM MMMs at pressures up to 30 bar and 300 °C, achieving H_2/CO_2 mixture selectivities of 8 [128]. The chronological progress for MMMs is shown in Figure 10.

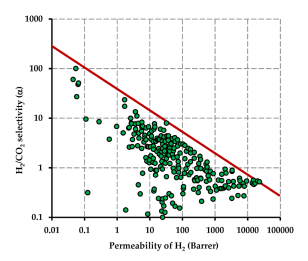


Figure 9. Updated (2008) Robeson's upper bound for H_2/CO_2 . The *x*-axis corresponds to the most permeable gas. Adapted with permission from [118]. Copyright, 2008, Elsevier.

Progress in membrane science until 1899	Year	Contributors			
First observations of permeation phenomena with water	1748	M. l'Abbé Nollet [64,65]			
Osmosis introduced to explain the diffusion of liquids across	1827	R. J. H. Dutrochet [66,67]			
permeable partitions (membranes)					
Solution-diffusion concept describing gas transport	1829	T. Graham [23,24]			
Rate of permeation of gases through membranes	1831	J. K. Mitchell [68,69]			
Graham's law of gas diffusion	1833	T. Graham [70-73]			
Fick's law of diffusion (diffusion of salts in water)	1855	A. Fick [74-76]			
Solution-diffusion model for gas transport in membranes	1866	T. Graham [25-27]			
Solubility of gases in rubber	1879	S. Wroblewski [77]			
Concept of semipermeable membranes	1887	J. H. van't Hoff [78-81]			
P 4000 4000	3/	0 1 11 1			
Progress 1900 - 1999 Preparation of synthetic non-rubber membranes: cellulose nitrate	Year 1907	Contributors			
Knudsen diffusion	1907	S. L. Bigelow, et al. [82,83] M. Knudsen [87,88]			
Ion exchange membranes	1911	F. G. Donan [89,90]			
CO ₂ adsorption and permeability in mineral matter-compounded	1911	F. Steinitzer [91]			
rubber membrane (inorganic-based MMM)	1714	r. Stentitzer [51]			
Pervaporation	1917	P. A. Kober [92,93]			
Effects of membrane aging on gas permeation and gas flux	1920	J. D. Edwards, et al. [94]			
proportional to partial pressure in the mixture	1720	j. D. Edwards, et al. [54]			
Time lag concept to calculate solubility and diffusivity from	1920	H. A. Daynes [104]			
permeability	1,20	11.11.2 dynes [101]			
CO ₂ adsorption in compounded rubber with ZnO	1922	C. S. Venable, et al. [105]			
Permeability to H ₂ of compounded rubbers	1929	T. Ohya, et al. [106]			
Gas permeation with fillers: additive aggregation	1931	V. N. Morris [107]			
Solubility and permeability in rubbers filled with minerals	1955	G. J. Van Amerongen [108]			
Interaction of rubber with fillers and effects on gas permeation	1959	N. A. Melikhova, et al. [109]			
Zeolites in polymers	1971	G. Christen, et al. [110]			
Time lags in zeolite/silicone measuring solubilities	1973	D. R. Paul, et al. [111,112]			
Extensive research on zeolite/polymer membranes	1987	H. J. C. te Hennepe, et al.			
		[114,129-132]			
Silicalite/cellulose MMMs	1988	S. Kulprathipanja, et al.			
		[115,116]			
Robeson's upper bound for selectivity-permeability correlations for	1991	L. M. Robeson [117,118]			
polymer membranes					
Progress 2000 to present day	Year	Contributors			
Factors controlling the successful formation of MMMs	2000	Mahajan, et al. [119]			
Cu-MOF/PAET MMMs for gas separations	2004	H. Yehia, et al. [120]			
Effects of polymer rigidification, zeolite pore size, and partial pore	2005	Li, et al. [121]			
blockage on MMMs	2003	Li, et al. [121]			
Cu ₃ BTC ₂ /PDMS and Mn(HCOO) ₂ /PSf MOF-MMMs	2006	A. Car, et al. [122]			
MOF-5/Matrimid® MMMs for gas separations using MOF	2006	E. V. Perez, et al. [123]			
nanocrystals and polyimide					
MOF-5, Cu-MOF, MOP-18/Matrimid® MMMs	2009	E. V. Perez, et al. [124-126]			
ZIF-90/Ultem™/Matrimid®/6FDA-DAM MMMs	2010	TH. Bae, et al. [127]			
NH₂-MIL-53/VTEC™ (MOF-based MMM) tested at high pressure	2015	E. V. Perez, et al. [128]			
and high temperature (30 bar and 300 °C)					

Figure 10. Chronological progress in mixed-matrix membrane science.

3. Evolution of Inorganic-Based MMMs

Researchers in the membrane-based gas separation field have done extensive research to substitute polymeric for metallic membranes due to the gas separation limitations of polymeric membranes. Metallic membranes show good gas separation properties, including high gas pair selectivities and high gas permeabilities, whereas, in contrast to metallic membranes, polymeric membranes are mechanically robust. Due to these reasons, MMMs received more attention and review articles emerged describing the advances in inorganic-based MMMs [133–136].

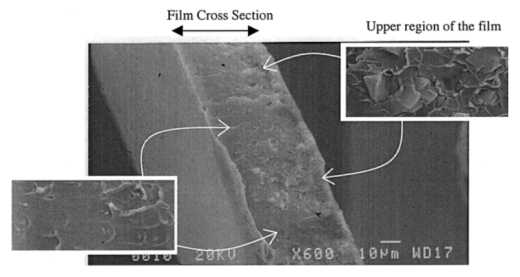
The history of inorganic materials as additives for MMMs goes back to 1971 in a patent published by G. Christen, et al. The authors used synthetic/natural zeolites in polysiloxane membranes to separate O_2 and N_2 from air [110]. After this first report of MMMs incorporating zeolites, no other reports appeared until a patent was published in 1985 by Sterzel and Sanner on the use of a poly(tetrafluoroethylene) membrane containing the cobalt-salicylaldehyde-ethylenediamine complex as the inorganic filler. The MMM was used to separate a O_2/N_2 mixture [137]. A year later, in 1986, R. S. Drago, et al. studied the facilitated transport of O_2 in polystyrene using $C_0(II)$ Schiff base complexes as additives for MMMs increasing O_2 permeability [138]. After that, a patent was published in 1988 based on MMMs containing cellulose acetate and silicate particles at ambient conditions. These membranes were used to separate O_2/N_2 and H_2/CO_2 [115]. From 1988 to 2000, there was a slight increase in the use of MMMs having inorganic fillers for gas separations. In the last two decades, an extensive number of research studies were carried out on MMMs containing inorganic additives, primarily molecular sieving zeolites and metal oxides. Interest in carbon molecular sieve-based MMMs emerged in the 2000 to 2002 period [133,134].

3.1. Zeolites Silicates and Metal Oxides in MMMs

After the first report of silicate-containing MMMs in 1988, MMMs consisting of different ratios of polyethersulfone (PES) and zeolites 13X and 4A were reported in 1994 [139]. Permeabilities were recorded for H₂, N₂, O₂, CO₂, and Ar. With a 50 wt % zeolite 13X loading, O₂/N₂, Ar/N₂, CO₂/N₂, H₂/N₂, and CO₂/O₂ ideal gas selectivities were 4.18, 1.68, 43.1, 70.8, and 10.4, respectively. Following the work carried out in 1994, Battal and co-workers were able to incorporate higher loadings of zeolite 4A into the PES matrix by using a special fabrication procedure that included the evaporation of excess solvent under a 0.2 atm pressure of N_2 at 60 °C followed by annealing at 80 °C. For ideal gas experiments, the H₂ permeability significantly increased with the incorporation of the zeolite, while the permeability values of the other gases increased slightly when compared to pristine PES. More importantly, this study was the first report of mixed gas experiments carried out on MMMs. With increasing the percentage of H_2 and CO_2 in the feed gas mixture, the permeabilities of H_2 and CO₂ increased [140]. In a research study carried out in 2000, the effect of zeolite particle size on the performance of polymer-zeolite MMMs was investigated. Zeolite silicates having particle sizes ranging from 0.1 to 8 µm were used to investigate CO₂, O₂, and N₂ permeabilities at 20 wt % and 40 wt % loadings. With increasing particle size, CO_2 , O_2 , and N_2 permeabilities increased significantly. For example, permeability increased approximately from 2000 to 4000 Barrer when the particle size increased from 0.1 to 8 µm [141].

There were just a handful of publications on inorganic material-based MMMs before 2001. However, the ten-year period from 2001 to 2010 turned out to be the emerging period for incorporating inorganic materials into MMMs for gas separation applications. Many publications and patents were published showing improved results for gas separations. Although higher permeabilities could be achieved by increasing the amount of inorganic material, the mechanical stability of the membranes was reduced. During this period, several different studies were carried out to address this issue by improving polymer-additive interactions in order to improve the mechanical characteristics of the respective MMMs. In one such study, the plasticizer di-butyl phthalate was used together with zeolite 4A in a Matrimid® polymer matrix [142]. In addition to incorporating plasticizers, the membranes were formed at elevated temperatures to ensure the reactivity of the plasticizer and the polymer.

As shown in Figure 11, the lower region of the membrane shows better contact between the zeolite particles and the polymer matrix.



Lower region of the film

Figure 11. Scanning electron microscopy (SEM) image of the cross-section of a 75/25 Matrimid[®]/di-butylphthalate polymer blend with 15% zeolite 4A. Reproduced with permission from [142]. Copyright, 2002, Wiley.

By 2002, T. C. Merkel, et al. reported the first reverse selective MMM using fumed silica as filler in PMP-2 for hydrocarbon separations [143]. The authors deduced that the fumed silica nanoparticles prevented a close packing of the chains which in turn increased the free volume of the polymer. The increase in free volume led to a decrease in diffusivity selectivity between larger and smaller molecules leaving the solubility selectivity to control gas transport in the MMM. Since larger molecules are more soluble than smaller molecules, larger hydrocarbons like *n*-butane would permeate faster than methane, as was seen in their permeability experiments. Based on the discovery made by T. C. Merkel, et al. a new theory was presented by R. J. Hill in 2006 [144]. The leading speculation in this work was based on the repulsive interactions between the non-porous additive and the polymer chains which results in free volume at the polymer-additive interface. As a result, the free volume of the entire system increases resulting in higher permeabilities for larger molecules over small molecules. These advances and the ability of additives to change drastically the gas transport properties of polymers propelled the research in the MMM field in the following years, extending into potential industrial applications for which the geometry of the MMM had to be changed from the traditional flat geometry to a hollow fiber to increase membrane area.

The use of hollow fiber membranes for gas separations emerged around 1973 [145]. Although many polymeric hollow fiber membranes were tested for gas separations, mixed-matrix hollow fiber membranes were not common mainly due to the challenges in fabrication. Mixed-matrix hollow fiber membranes were first reported in 2006 by Li, et al. who fabricated PES/BTDA-TDI/MDI co-polyimide (P84) hollow fibers with a 0.55 μ m PES-zeolite beta MMM dense, selective layer that showed good CO₂/CH₄ separation properties for pure gases (CO₂/CH₄ = 29.2) and gas mixtures (CO₂/CH₄ = 33.8). A very thin (0.55 μ m) PES-zeolite beta mixed-matrix dense selective layer was applied by manipulating the flow rates of the inner and outer layers during the spinning process. The interaction between the inner and the outer layer of the hollow fiber was improved through heat treatment at 235 °C [146].

Researchers incorporated many different zeolites into polymers to produce useful MMMs for gas separation. In addition to these, some have modified the chemical structure of zeolites in order to improve gas separation properties. To further improve the characteristics of MMMs, a third component

was sometimes added to the MMM mixture. Y. Zhang and co-workers were able to improve the permeabilities of gases, including H₂, N₂, O₂, CH₄, and CO₂, by 10% to 50% by combining carbon aerogel and zeolite in Matrimid[®] [147,148]. According to the N₂ adsorption data, the availability of combined micropores and mesopores improves the separation properties of these membranes. From SEM images of the cross-sections of the MMMs, the authors observed a good dispersion of the aerogel particles in the polymer which exhibited plastic deformation from the freeze fracture. Y. Zhang also performed mechanical analysis (stress-strain) of the MMMs to measure changes in Young's modulus and T_g, the resuls showed increased modulus from 2.4 GPa (Matrimid[®]) to 2.7 GPa (20% w/w aerogel/Matrimid® MMM) and increased T_g of the MMMs from 317 °C (Matrimid®) to 338 °C (20% w/w aerogel/Matrimid[®] MMM). These mechanical properties improvements were attributed to the penetration of the polymer chains into the mesopores of the carbon aerogel particles to achieve a good contact. These results show that a strong additive/polymer interaction could be achieved in MMMs and can be detected by routine mechanical analyses. Y. Zhang, et al. also reported similar results for ZSM-5/Matrimid® MMMs in 2008 [149], the mechanical analyses showed increased modulus and T_g for the MMMs with increasing ZSM-5 loadings up to 30% w/w. A study carried out in 2006 reported the effect of zeolite 4A nanocrystals and microcrystals on the gas separation of He/N_2 , H₂/N₂, He/CO₂ and H₂/CO₂ [150]. Although the gas permeabilities decreased with the increased percentage of zeolite particles in the matrix, gas selectivities increased significantly, especially with nanocrystalline zeolite 4A particles. According to another publication in 2007, Na ions in zeolite NaA were successfully replaced with Ag ions to improve the gas separation characteristics of PES-based MMMs [151]. The membranes with zeolite AgA showed higher permeability for CO₂ and lower CH₄ permeabilities (compared to NaA zeolite-MMMs) resulting in a higher CO₂/CH₄ selectivity of 59 at a 50 wt % loading. This same year, W. Yave, et al. [152] reported that 40 wt % TiO₂/PMP-2 MMMs also suffered from physical aging and probably chemical aging too atfter being stored for 4 months, the n-C₄H₁₀ permeability in these MMMs dropped by 40% and the n-C₄H₁₀/CH₄ selectivity dropped by 10%. Similar loss of permeability (up to 40%) over a period of 154 days were also observed by S. Matteuci, et al. [153] for 20 vol % TiO₂/PTMSP MMMs. An important progress in the reduction of aging in MMMs came in 2011 from the work of J. K. Ward, et al. [154]. Ward observed that molecular sieve SSZ-13/PDMC MMMs suppressed significantly physical aging of the MMMs with a loss of 5%–10% of permeability wereas the neat PDMC loosed 30% of permeability in the same time frame. MMMs using Matrimid[®] and mesoporous ZSM-5 nanoparticles to improve gas separations were reported in 2008 [149]. The results indicated a good contact between the nanoparticles and the polymer matrix since the polymer chains penetrated into the mesopores of ZSM-5. Meanwhile, the micropores of ZSM-5 improved the size and shape selectivity. At a 10% loading of ZSM-5, the H₂/N₂ selectivity increased from 79.6 to 143 compared to pure Matrimid[®]. At 20% loading, the O₂/N₂ and H₂/CH₄ selectivities increased from 6.6 to 10.4 and 83.3 to 169, respectively, when compared to pure Matrimid[®]. In 2011, the amine-functionalized FAU/EMT intergrown zeolites were used as the inorganic filler in 6FDA-ODA-based MMMs to improve CO₂/CH₄ separations [155]. Defect-free MMMs were obtained with amine-functionalized zeolites and, at 25 wt % modified zeolite loadings, the MMMs showed encouraging CO₂/CH₄ selectivities for both pure and mixed gas experiments. The MMMs achieved CO₂/CH₄ selectivity increments from the pure polymer of 118% (ideal) and 94% (gas mixture).

Non-porous and ordered mesoporous silica based MMMs have also been significantly studied. Joly, et al. studied composite membranes made from polyimide and silica [156]. Unlike porous inorganic materials such as zeolites, addition of nonporous silica alters the polymer chain packing thus affecting the gas transport properties of the polymer. According to Gomes, et al. incorporation of silica particles in a polymer matrix could result in a higher permeability or a slight decrease in selectivity due to an increased free volume in the membrane that results from the formation of non-selective voids at the additive/polymer interface [136,157]. Ahn, et al. studied the addition of fumed silica into PIM-1 in terms of gas permeability and selectivity. By studying the density, thermal stability, and morphology of the MMMs, the authors of this work were able to describe the reasons for the

increments of gas permeabilities. Additional voids created in the polymer backbone by the interactions of the polymer chains and the silica particles resulted in higher permeabilities. For instance, the CO_2 permeability increased from 6000 Barrer (pure PIM-1) to 13400 Barrer upon the addition of 24% silica into PIM-1 [158]. Takashi, et al. also studied the effect of chemical modification of SiO_2 nanoparticle dispersion in poly(ether imide). MMMs containing sylil-modified SiO_2 fillers displayed reduced void volumes compared to unmodified MMMs. At 5 wt % loading, void volumes in MMMs containing silylated SiO_2 and pristine SiO_2 were 0.25 vol % and 1.00 vol %, respectively [159,160].

Most common molecular sieving materials such as MOFs, ZIFs, Zeolites show low aspect ratios compared to layered materials such as silicates. Therefore, the incorporation of a small volume fraction of such layered materials can improve the performance of MMMs [161,162]. Defontaine, et al. were able to improve the CO_2/CH_4 selectivity of PDMS by 145% with the incorporation of sepiolite [163]. The effect on the H_2/CO_2 separation in PBI with the addition of sheet like AMH-3 was studied by Choi, et al. in 2008. According to their observations, a 3 wt % AMH-3 MMM showed a lowering of CO_2 permeability which resulted in an increase in selectivity from 15 to 40 with respect to the polymer [164].

Modifications to the chemical structure of the polymer matrix were important to the mechanical properties of MMMs. In 2007, SPES was used as the precursor for MMMs with zeolite 4A [165]. Compared to PES membranes, SPES-based membranes displayed improved properties at the polymer-zeolite interface as visualized by SEM. In SPES MMMs, particle aggregation is minimal, which is considered an important characteristic for industrial gas separation MMMs. Figure 12 shows the morphological differences of PES-zeolite 4A membranes and SPES-zeolite 4A membranes from SEM micrographs of their cross-sections.

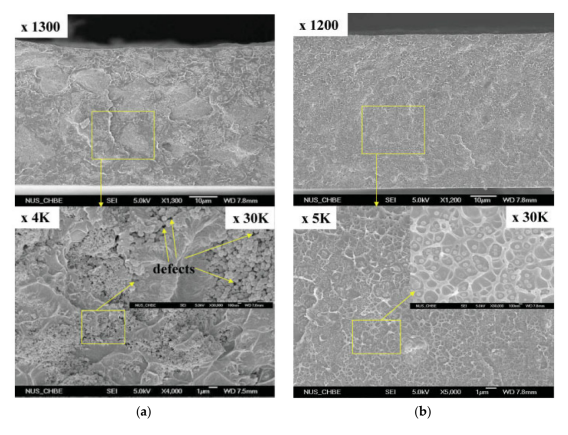


Figure 12. SEM images of the cross-sections of (a) polyethersulfone (PES)-zeolite 4A MMM; (b) sulfonated polyethersulfone (SPES)-zeolite 4A MMM. Reproduced with permission from [165]. Copyright, 2007, Wiley.

In another study, Jha and co-workers were able to obtain an increased CO_2/N_2 selectivity, from 43 to 54, by preparing MMMs with methoxy ethoxy ethanol (MME)-substituted PPZ and SAPO-34 [166]. PPZ's rubbery nature allows better interfacial contact with the additive. By converting to MME-substituted PPZ, the CO_2 selectivity of the MMMs was increased. Their MMMs showed gas separation properties that surpassed the Robeson upper bound for CO_2/N_2 separations. Another study carried out in 2010 was based on acrylate-derivatized polysulfone (PSf-acrylate) and zeolite 3A for the separation of H_2 from CO_2 [167]. Unmodified PSf has a H_2/CO_2 separation factor of 1.53, but the 40% zeolite 3A-PSF-acrylate MMM exhibited an improved selectivity of 3.57. With further increase in zeolite loading, however, the H_2 permeability increased and the selectivity ratio decreased.

In addition to chemically modifying the polymer structure or incorporating additives into the polymer matrix, the polymer and additive were chemically reacted together in a few occasions to obtain MMMs. In one study, the cross-linkable polymer PDMC was cross-linked with the dispersed inorganic additive SSZ-13 to prepare MMMs [154]. In this work, the non-crosslinked 25 wt % SSZ-13/PDMC MMM showed a CO₂ permeability of 153 Barrer and a CO₂/CH₄ ideal selectivity of 34.7, whereas the cross-linked MMM exhibited good separation properties for the CO₂/CH₄ gas pair with selectivity and permeability increments of 6.9% and 121%, respectively. In mixed gas experiments (10% CO₂ + 90% CH₄) at 700 psia and 35 °C, the CO₂/CH₄ selectivity and CO₂ permeability were increased by 13% and 47%, respectively. In this work, cross-linking the polymer with the inorganic particles ensured good contact between the continuous phase and the dispersed phase. In 2012, MMMs for CO₂ separations were fabricated based on SPEEK and $-SO_3$ functionalized mesoporous MCM-41 [168,169]. Covalent interactions between the inorganic filler and the polymer were visualized using SEM. Experiments were carried out at temperatures ranging from 25 to 65 °C and at pressures of 10 bar to test the membranes for industrial applications. CO₂/N₂ and CO₂/CH₄ selectivities of 40.5 and 22.9, respectively, were reported at a CO₂ permeability of 21 Barrer.

3.2. Carbon Materials in MMMs

A variety of carbon based materials, including CNTs, graphene, fullerenes, and CMSs, prepared by the pyrolysis of polymer precursors at high temperatures under controlled environments were used as fillers in MMM preparation [170,171]. CNTs and CMSs are known to have several orders of magnitude higher permeabilities than molecular sieves. In contrast, a graphitic material could be categorized as a selective material. Like other inorganic molecular sieving materials, such as zeolites and metal oxides, CMSs show better selectivity and good permeabilities at the same time. The main advantage of CMSs is the variety of pore sizes and structures that can be obtained by varying the precursor material and the pyrolysis temperature. Generally, CMSs can be used to improve gas flux through a polymeric membrane. A patent on mixed-matrix nanoporous carbon membranes published in 2001 by Corbin, et al. is the first report of carbon material-based mixed-matrix membranes [172]. Following this study, Koros and co-workers in 2002 made mixed-matrix membranes having carbon-based molecular sieving particles prepared by pyrolyzing a precursor polymer [173]. Successful preparation of MMMs using carbon molecular sieves was again reported in 2003 [174]. During this particular study, Matrimid® 5218 polyimide films were pyrolyzed at 800 °C following a published procedure and then ball-milled to get fine carbon molecular sieve (CMS) particles. These particles were then dispersed in Matrimid[®] 5218 and UltemTM 1000 precursors to form MMMs. SEM images showed poor polymer-sieve contact in UltemTM 1000 and good contact in Matrimid[®] 5218. CO₂/CH₄ selectivities of 50 and O₂/N₂ selectivities of 7 were obtained for these membranes. Matrimid[®] 5218 MMMs having 36% CMS displayed gas separation properties that surpassed the Robeson's upper bound for O_2/N_2 separation.

Gas separation properties for KY-zeolite/Matrimid $^{\odot}$ MMMs pyrolyzed at 800 $^{\circ}$ C were studied focusing on CO_2/CH_4 separations [175]. This is the first report of MMMs containing both zeolitic and

carbon materials. For the pyrolyzed membranes having KY zeolites, the measured CO_2 and CH_4 permeabilities were 266 Barrer and 2.2 Barrer, respectively and the CO_2/CH_4 selectivity was 124.

To improve the interaction between the polymer matrix and CMS particles, the addition of a third component was tested in a study carried out in 2015 [176]. Interactions between PES and CMS were enhanced by the addition of DEA as the third component in the MMMs. Their MMM, having 15 wt % DEA, showed more than a three-fold increment in CO_2 permeance (from 30 to 123.5 GPU) from the PES membrane. Results show that the MMMs reported in this work had good separation properties for CO_2/CH_4 that surpassed the Robeson's upper bound.

There are not many reports of bucky balls and other fullerenes being used as inorganic fillers. However according to a few studies, C_{60} fullerenes could increase the selectivities for the gases considered. MMMs based on benzylamine modified C_{60} as the filler in Matrimid[®] 5218 were reported in 2003 [177]. Although the permeabilities for all the gases tested (He, O_2 , N_2 , CH_4 and CO_2) decreased with the increasing amount of benzylamine modified C_{60} , the selectivity for He/ N_2 increased while the others remained constant. In their work carried out on MMMs of ethyl cellulose (EC) and C_{60} particles, Sun, et al. reported improved permselectivity for C_3H_6/C_3H_8 separations with a 16 wt % C_{60}/EC MMM [178]. C_3H_6 favors the π -electron cloud in C_{60} due to π - π interactions, which improves the solubility and permeability of C_3H_6 . Permeability and selectivity values increased 7.1 and 1.6 times, respectively, when compared to those of the pure EC membrane. To further emphasize their theory, the membrane was treated with UV radiation, which destroys the π -electron cloud in C_{60} . Afterwards, the permeability of C_3H_6 and the C_3H_6/C_3H_8 selectivity decreased compared to the initial values.

Unlike fullerenes and carbon molecular sieves, both CNTs and FCNTs were commonly used to fabricate MMMs [171,179]. CNTs can act either as a pathway for gas molecules to go through or as a sieve. More importantly, CNTs are known to be stiff particles having dimensions ranging from 1 to 100 nm. As a result, they can be used as rigidifying agents in membranes. The first report of CNT-based MMMs for gas separations, in 2006, used poly(imide-siloxane) polymer and SWCNTs with closed ends [180]. The gas permeation results showed a decrease in permeability for He and an increase in permeability for O_2 , O_2 , and O_3 , and O_4 . The drop in He permeability was attributed to the affinity of SWCNTs with the polymer matrix. All types of CNTs are electrically conducting materials, therefore, by applying an external electric field, CNTs can be aligned in one direction, which is important for gas transport and membrane rigidity. Based on this property, in 2012, Kumar, et al. prepared MMMs using PS and MWCNTs to enhance O_4 permeability [181]. MWCNTs were dispersed in the PS matrix either by mechanical sonication and stirring or by applying a high transverse O_4 electric field to orient the MWCNTs. According to their gas permeability data, MMMs made with oriented MWCNTs showed a high O_4 high O_4 gas permeability of 275 Barrer.

The next noticeable improvement in CNT-based MMMs incorporated functionalized CNTs. FCNTs were used either to improve the interactions between the polymer and the CNTs or to improve the separation properties of the membranes. Ismail and co-workers fabricated MMMs comprised of PES and FMWCNT that showed improved gas separation for CO_2 over CH₄ [182]. Before the MWCNTs were functionalized, they were purified by treating with strong acids. Then, 3-aminopropyltriethoxysilane was used to functionalize the surface of the MWCNTs, which improved their dispersion within the polymer matrix. MMMs containing 0.5 wt % functionalized MWCNTs showed a 388% increase in permselectivity for CO_2/CH_4 compared to pure PES. In 2014, A. L. Ahmad, et al. explored the effects of the addition of β -cyclodextrin-modified MWCNTs into a cellulose acetate (CA) matrix to prepare MMMs focusing on CO_2/N_2 separation [183]. Good gas permeance values were obtained for CO_2 during the single gas experiments, which were greater than the permeabilities of either MMMs containing pristine MWCNTs or the bare CA membranes. The MMMs with a 0.1 wt % modified MWCNT loading exhibited a CO_2 permeance of 741 GPU at 3 bar.

In most cases, the incorporation of CNTs improved gas permeabilities. However, graphene oxide showed better sieving properties when oriented perpendicularly to the direction of gas flow [184,185]. Based on these observations, Li, et al. studied the synergistic effect of GO and CNTs on the permeability

and selectivity of gases in MMMs made with both additives. MMMs based only on GO and Matrimid demonstrated high gas selectivities for CO_2/N_2 and CO_2/CH_4 but low gas permeabilities. The CNT-containing MMMs, however, showed higher permeabilities but lower selectivities. When the authors tested the MMMs containing both GO and CNT in Matrimid, they observed that both, permselectivity and permeability, increased. Numerically, compared to pure Matrimid, the MMM with 5 wt GO and 5 wt CNT showed increments of 331%, 149% and 147% in CO_2 permeability, CO_2/CH_4 selectivity, and CO_2/N_2 selectivity, respectively [171]. The chronological progress for the inorganic-based MMMs is shown in Figure 13.

	V	Comballing
Inorganic-based MMMs Progress Heterogenous membrane for separating fluid mixtures	Year 1971	Contributors G. Christen, et al. [110]
Polymer membranes containing crystalline compounds	1985	Sterzel, et al. [137]
Facilitated transport of O ₂ in Co(II)/polystyrene MMMs	1986	R. S. Drago, et al. [138]
Gas separation by means of mixed-matrix membranes	1988	S. Kulprathipanja, et al.
Gas permeation characteristics of polymer-zeolite MMMs	1994	[115] Sueer, et al. [139]
Effect of feed composition on the performance of polymer-zeolite MMMs	1995	Battal, et al. [140]
Factors controlling the successful formation of MMMs	2000	Mahajan, et al. [119]
Effect of zeolite particle size on the performance of MMMs	2000	Tantekin, et al. [141]
n-Pentane/i-pentane separation with zeolite-PDMS MMMs	2001	Tantekin, et al. [186]
First reverse-selective fumed silica/PMP-2 MMMs	2002	T. C. Merkel, et al. [143]
Preparation and characterization of a glassy fluorinated polyimide/zeolite MMMs	2002	Pechar, et al. [187]
The effects of polymer chain rigidification, zeolite pore size, and pore blockage PES-zeolite A MMMs	2005	Li, et al. [121]
Gas separation of nanostructured PES-zeolite 4A MMMs	2006	Huang, et al. [150]
Diffusive permeability and selectivity of nanocomposite membranes	2006	R. J. Hill [144]
Mechanical analysis of MMMs	2006	Y. Zhang, et al. [147,148]
Removal of thiophenes from n-octane with PDMS-AgY zeolite MMMs	2007	Qi, et al. [188]
Poly(4-methyl-2-pentyne)/TiO ₂ hybrid nanocomposite membrane for	2007	Yave, et al. [152]
butane/methane separation Ag*-zeolite/polymer MMMs with a high CO2/CH4 selectivity	2007	Li, et al. [151]
Primer to prevent nanoparticle agglomeration in MMMs	2007	Li, et al. [165]
CO2 selective MMMs formulation and characterization using rubbery substituted PPZ	2008	Jha, et al.[166]
H ₂ separation using PSf acrylate-zeolite MMMs	2010	Khan, et al. [167]
Thin Pd-silica MMMs with nano-dispersion for improved H2 permeability	2010	Kanezashi, et al. [189]
Cross-linkable MMMs with surface modified molecular sieves and aging reduction	2011	Ward, et al. [154]
MMMs based on polymerizable room-temperature ionic liquids and SAPO-34 particles to improve CO ₂ separation	2011	Hudiono, et al. [190]
Cross-linked polyvinylalcohol-polysiloxane/fumed silica MMMs	2011	Xing, et al. [191]
containing amines for CO ₂ /H ₂ separation Amine-functionalized zeolite FAU/EMT-polyimide MMMs for CO ₂ /CH ₄	2011	Nik, et al. [155]
separation SPEEK and functionalized mesoporous MCM-41 MMMs for CO2	2012	Khan, et al. [169]
separations Amine-functionalized SBA-15 in poly(styrene-b-butadiene-b-styrene) nanostructured MMMs	2013	Buonomenna, et al. [192]
PSf MMMs incorporating Pd nanoparticles dispersed in PVPy for H ₂ separation	2014	Suhaimi, et al. [193]
separation		

Figure 13. Chronological progress for inorganic-based MMMs.

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4. Evolution of MOF-Based MMMs

4.1. Metal-Organic Frameworks

There is still an ambiguity regarding the distinction between coordination polymers (CP) and MOFs that muddies the classification of hybrid organic-inorganic materials [194]. For simplicity and following IUPAC recommendations [195], we will use the term MOFs to refer to the organic-inorganic compounds that form 1D, 2D, or 3D frameworks. MOFs offer great potential for gas separations owing to their large surface areas, gas sorption capacities, pore apertures suitable for molecular sieving, and high temperature stability. These materials can also undergo chemical functionalization of their organic components before or after their synthesis to tune their gas affinity, pore size, pore aperture, and affinity to polymers. MOFs, ZIFs, MILs, MOPs, MOCs, and their variations are materials that can form crystalline frameworks or single molecular cages (MOPs and MOCs) that result from combining metal ions (SBUs) with organic linkers (Figure 14a,b). It is important to note that MOPs (Figure 14b) and MOCs are single porous molecular cages that offer the potential to be dissolved in organic solvents while retaining their porosity and 3D structure in solution. MOFs, on the other hand, grow into particles of sizes that range from the nanometer scale to the micrometer scale, depending on the synthesis conditions, and cannot be dissolved into single unit cells without destroying the framework. MOFs can be dispersed in organic solvents (Figure 14a).

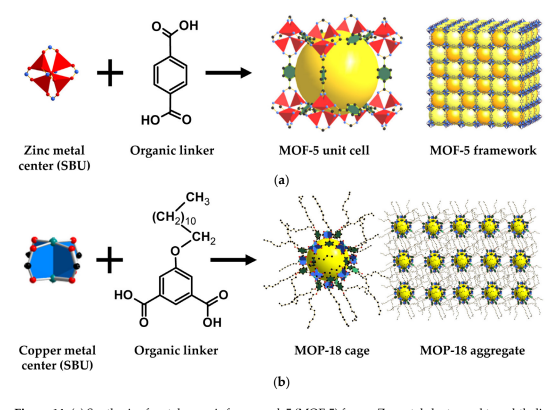


Figure 14. (a) Synthesis of metal-organic framework 5 (MOF-5) from a Zn metal cluster and terephthalic acid. The figure shows the unit cell and the framework drawn using Diamond and crystallographic data from [196]; (b) Synthesis of metal-organic polyhedra 18 (MOP-18) from a copper dimer and alkylated isophthalic acid. Note that the resulting cage does not form frameworks. Drawn using Diamond and crystallographic data from [197]. Yellow spheres show the available volume in the pore.

ZIFs are a subfamily of MOFs that are characterized by the similitude of their Zn–N–Zn angle (145°) to that of sodalite's Si–O–Si bonds. The Zn–N bond in ZIFs is resistant to high temperatures and to water, making them attractive for MMM preparation. Due to the variety of available imidazolate

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linkers, a large number of ZIFs with different chemical functionalities and pore apertures have been synthesized and tested for gas adsorption and gas separation in MMMs (Figure 15) [198].

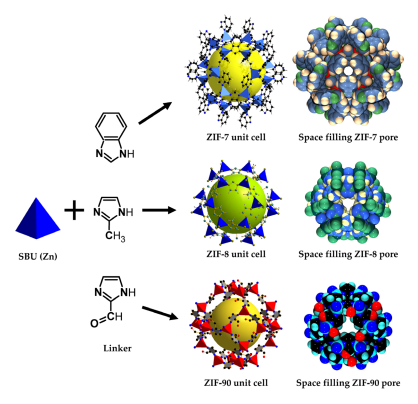


Figure 15. Synthesis of zeolitic imidazolate 7 (ZIF-7), ZIF-8, and ZIF-90 from a Zn ion and imidazolates. Unit cells and space filling models drawn using Diamond with crystallographic data for ZIF-7 [199], ZIF-8 [200], and ZIF-90 [201]. Note that depending on the linker, different pore aperture shapes can be obtained.

4.2. MOFs in MMMs

One of the first reports on the use of MOFs for the preparation of MMMs for gas separations emerged from the work of H. Yehia, et al. in 2004 [120]. In this work, a copper-based MOF (Cu-MOF) was synthesized from copper acetate, 4,4'-biphenyl dicarboxylate, and triethylenediamine [202]. The obtained crystals were then mixed with the rubbery polymer PAET to form MMMs with loadings up to 30% (w/w). The MMMs were tested for CH₄/CO₂ separations at 35 °C and 2.0 bar and resulted in an increase in CH₄ permeation and a decrease in CO₂ permeation that were attributed to the hydrophobicity and pore size (0.12 nm) of the MOF. Two years later, in 2006, A. Car, et al. synthesized Cu(BTC)₂ and Mn(HCOO)₂ MOFs that were used at loadings up to 40% (w/w) in conjunction with PSf and PDMS for the preparation of MMMs [122]. The membranes were tested for H₂, N₂, O₂, CH₄, and CO₂ permeability. The authors reported mixed results for permeability and selectivity, but no further characterization of the MOFs or the MMMs was attempted. The same year, E. V. Perez, et al. incorporated for the first time nanocrystals of MOF-5 (Figures 14a and 16) into Matrimid[®], a polyimide with high T_g [123]. Their preliminary results showed that, at 30% (w/w) loadings, MOF-5 improved the gas permeability of the MMMs by 120% over the pure polymer without significant variation in gas selectivity.

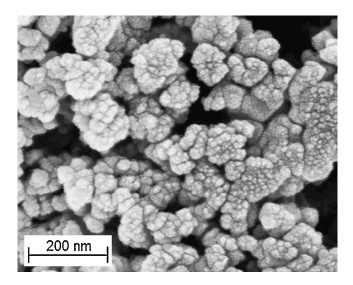


Figure 16. SEM image of nanocrystals of MOF-5. Reproduced with permission from [124]. Copyright, 2009, Elsevier.

Also in 2006, C. Barcena prepared PAET, POT, and PDMS MMMs of Cu-MOF with loadings up to 30% (w/w) [203]. Gas permeation (H_2 , N_2 , O_2 , CH_4 , and CO_2) results for these membranes resulted in no significant changes in permeability or selectivity compared to the pure polymers. In 2008, working on the progress made by Yehia, et al., Car, et al., and Perez, et al., in 2008 Y. Zhang, et al. reported on the preparation of Matrimid®-based MMMs that contained a submicron, water stable Cu-MOF (Cu-BPY-HFS) [204]. The MOF exhibited a thermal stability up to 100 °C that limited the drying temperature of the membranes to 50 °C. An increase in CH₄ permeability up to 180% was observed at 35 °C for a 40 wt % MOF loading. Ideal selectivities also decreased as the MOF content increased, but did so more markedly for those gas pairs containing CH_4 and N_2 . The authors attributed the declines to the increased affinity of the MMM for CH₄ due to the presence of the MOF. Y. Zhang, et al. also reported the mechanical properties of their MOF-based MMMs by performing stress-strain tests at room temperature. The results indicated an increased modulus from 2.41 GPa (Matrimid®) to 2.67 GPa (MMM with 20 wt % MOF loading). At 30 wt %, the authors observed a decrease in modulus from 2.67 GPa to 2.54 GPa. These improvements in modulus were attributed to a good contact and affinity between the polymer and the MOF. T_g measurements were not performed in this work due to the lower thermal stability of the MOF (up to 200 $^{\circ}$ C) and high T_g of the plymer (316 $^{\circ}$ C). Similar measurements of mechanical properties of MOF-based MMMs were also performed by E. V. Perez, et al. in 2009 [125,126] (increased modulus of MOP-18/Matrimid® MMMs), by T. Yang, et al. in 2011 [205] (increased Tg of ZIF-7/PBI MMMs), by E. V. Perez, et al. in 2015 [128] (mechanical stability of NH₂-MIL-53/VTEC MMMs at 300 °C by thermomechanical analyses), and in MOF-based MMM reviews [206,207]. Recently, in 2016, S. J. D. Smith, et al. [208] and E. M. Mahdi, et al. [209] reported more detailed works on mechanical analyses of MMMs which are discussed at the end of this section, following the chronological progress.

Continuing their work on MOF-based MMMs, E. V. Perez, et al. reported complete studies of MOF-5/Matrimid® and Cu-MOF/Matrimid® MMMs in 2009 [124,125]. The same year, they also reported their initial work on MOP-18/Matrimid® MMMs that later was reported in more detail [126]. E. V. Perez, et al. found that plastic deformation of the polymer (e.g., polymer veins) occurred around the MOF-5 particles due to the strong interaction between the MOF and the polymer (Figure 17) [124]. Their detailed observations of gas permeation at 35 °C and 2 bar showed that up to a 120% increase in permeability with no significant variation in ideal selectivity could be achieved at a 30% (w/w) MOF-5 loading without introducing nonselective voids in the membrane. They also concluded that diffusivity was significantly enhanced due to the porosity introduced by the uniform channels of

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MOF-5. With their work on Cu-MOF/Matrimid[®] MMMs, E. V. Perez, et al. provided support for the case that the uniformity of the MOF channels and the pore size may play a significant role in gas permeation in MMMs [125]. In their work, the highly porous and thermally stable Cu-MOF [202] (BET surface area of 3000 m²/g and pore size of about 1 nm) was loaded into Matrimid[®] up to 80% (w/w). Compared to pure Matrimid[®], MMMs with a 30% (w/w) Cu-MOF loading exhibited permeabilities enhanced by 1000% (v/w) or 1400% (v/w), while suffering only a 17% decrease in v/w0 selectivity (v/w0 matrimid[®]; 5.07 for MMM; Permeability of Matrimid[®] to v/w0 arrer, MMM = 17.2 Barrer).

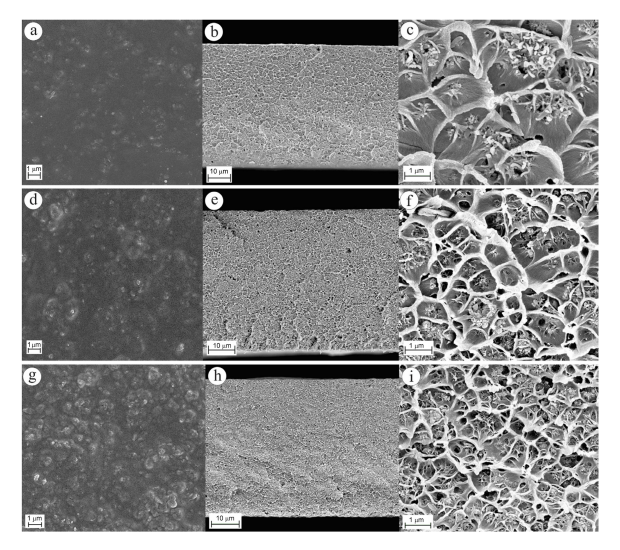


Figure 17. SEM images of the surface (\mathbf{a} , \mathbf{d} , \mathbf{g}) and the cross-sections at low (\mathbf{b} , \mathbf{e} , \mathbf{h}) and at high (\mathbf{c} , \mathbf{f} , \mathbf{i}) magnifications of 10% (w/w), 20% (w/w), and 30% (w/w) MOF-5/Matrimid[®] MMMs, respectively. Cross-sections showing plastic deformation of the polymer matrix due to the presence of MOF-5 nanoparticles. Reproduced with permission from [124]. Copyright, 2009, Elsevier.

Although the use of MOFs for MMM preparation significantly mitigated interfacial rejections and resulted in membranes free of defects, E. V. Perez, et al. recognized that the limiting factor in achieving a perfect dispersion was the aggregation of the nanocrystals. Aiming to solve this problem, they introduced MOP-18 molecular cages (Figure 14b) into Matrimid[®] to study the effect this type of hybrid material would have on the gas separation properties of MMMs [125]. Their work showed that MOP-18 was dispersed at the molecular level in the polymer matrix since it both dissolved in

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the casting solvent and exhibited affinity for the polymer. Gas permeability, however, did not change significantly possibly due to pore blockage by the alkyl chains of the MOP.

Following on the success of previous works of Perez, et al. [124] and Zhang, et al. [204], R. Adams, et al., in 2010, prepared defect-free MMMs of CuTPA (a Cu-MOF) and PVAc with a 15% MOF loading [210]. Their permeability experiments showed that the CO₂/CH₄ selectivity of the MMM increased by as much as 15% over the pure polymer owing to the presence of the MOF in the membrane. By comparing the apparent diffusivities of the gases in the MMM with those in the pure polymer, the authors concluded that the gases were diffusing through the MOF and not through voids or defects at the MOF-polymer interface, and that a true mixed-matrix effect was achieved. This same year, T.-H. Bae, et al. reported on the first ZIF-based MMM incorporating ZIF-90 crystals (Figure 15) into Ultem™, Matrimid®, and 6FDA-DAM polyimides [127]. Submicrometer-sized ZIF-90 crystals synthesized using methanol and deionized water as nonsolvents (Figure 18) were incorporated in these polyimides (Figure 19) a 15 wt % ZIF loadings. The SEM images of the cross-sections showed good adhesion of the polymer to the ZIF and no gross defects at the interface. The SEM images also showed different degrees of polymer plasticization around the crystals, with UltemTM showing the least plasticization and 6FDA-DAM showing the most. Gas permeation studies showed that ZIF-90/UltemTM and ZIF-90/Matrimid® MMMs only increased permeability and retained CO₂/CH₄ selectivities. This result, according to the authors, was due to a mismatch between the high gas permeability in the pores of ZIF-90 (estimated to be thousands of Barrers) and the low permeability in the polymers (Ultem[™] and Matrimid[®], 1 to 10 Barrer). In the case of ZIF-90/6FDA-DAM, the mismatch was less because 6FDA-DAM exhibits higher permeabilities (CO₂ = 390 Barrer) than UltemTM and Matrimid[®], resulting in an MMM with improvement in both permeability and selectivity. The ZIF-90/6FDA-DAM MMM showed an increase in CO₂ permeability of 720 Barrer and a higher CO₂/CH₄ selectivity of 37 compared to 24 for 6FDA-DAM.

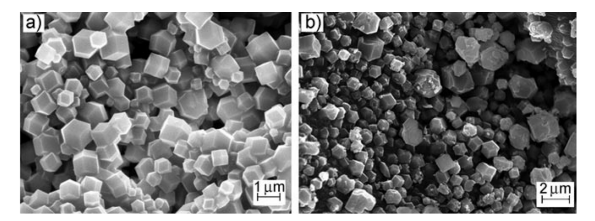


Figure 18. SEM images of submicrometer-sized ZIF-90 particles synthesized using nonsolvents: (a) methanol (ZIF-90A); (b) deionized water (ZIF-90B) [127]. Reproduced with permission from [127]. Copyright, 2010, Wiley.

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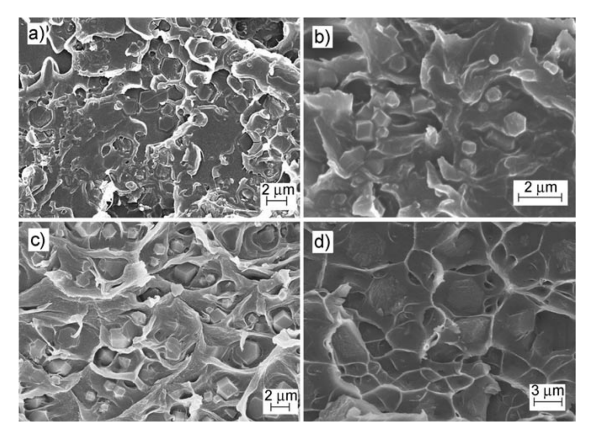


Figure 19. SEM images of cross-sections of 15 wt % ZIF-90/polyimide MMMs: (a) ZIF-90A/Ultem™; (b) ZIF-90A/Matrimid®; (c) ZIF-90A/6FDA-DAM; and (d) ZIF-90B/6FDA-DAM. Reproduced with permission from [127]. Copyright, 2010, Wiley.

By 2010, K. Diaz, et al. studied [13C]O₂ gas diffusion in ZIF-8/PPEES MMMs containing up to 30 wt % ZIF-8 [211]. The authors used macroscopic (e.g., gas adsorption, time lag method) and microscopic techniques (e.g., PFG-NMR) to study gas transport in the membranes. Gas solubility experiments were carried out using the dual chamber method and the dual-mode solubility model to identify Langmuir (partially immobilized gas) and Henry sorption. Permeability experiments were also performed to obtain solubility and diffusivity data using the time lag method. From these studies, the authors concluded that ZIF-8 crystals provided Langmuir sorption sites for CO₂ that resulted in an increase in gas adsorption in the MMMs when the ZIF content was increased. From PFG NMR, the authors also observed chemical shifts, corresponding to ¹³C, for both free (124.2 ppm) and sorbed (120.7 ppm) [13C]O₂ molecules in the membranes. The authors also found that the self-diffusion coefficient of CO₂ in the MMMs was about 2 times larger than the effective diffusion. This same year, J. Hu, et al. reported on the preparation of Cu₃(BTC)₂/PI (ODA-PMDA) hollow fiber MMMs [212]. In this work, the MOF was first synthesized and then its size reduced to submicrometer size using a ball mill. Though this method for reducing the crystal size is rather crude and prone to destroy the crystals, the authors loaded 6 wt % of the MOF into the polyamic acid form of the ODA-PMDA polymer, and then, using the dry/wet spinning method, prepared the hollow fiber MMMs. The fibers were solvent exchanged in water for 1 d, washed with methanol and hexanes 3 times, and dried to induce imidization using a temperature program up to 300 °C. Although most Cu-MOFs are thermally labile and sensitive to water, the authors obtained X-ray diffraction patterns for the hollow fiber MMMs. Gas permeation at 10 bar and 25 $^{\circ}\text{C}$ showed improved flux, from 876 (PI) to 1270 (MMM) GPUs, and improved H₂/CO₂ selectivity, from 10.0 (PI) to 27.8 (MMM). Another significant contribution to this field in 2010 came from the theoretical work of S. Keskin, et al. on the application of MOFs in MMMs for gas separations [213,214]. S. Keskin used

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atomistic and continuum modeling to select the best MOF for a particular separation [213]. The authors first validated their models using gas permeation data for MOF-5/Matrimid® MMMs reported by E. V. Perez, et al. [124] and then predicted the CO₂/CH₄ separation performance for 6 MOF/Matrimid[®] MMMs. From the series of Matrimid[®]-based MMMs containing IR-MOFs (e.g., IRMOF-1, IRMOF-8, IRMOF-10, IRMOF-14) and copper MOFs (CuBTC and Cu(hfipbb)(H₂hfipbb)_{0.5}), the authors found that Cu(hfipbb)(H₂hfipbb)_{0.5}/Matrimid[®] MMMs possessed the best CO₂/CH₄ separation performance with selectivity and permeability that improved with increasing MOF loading. In another work reported the same year, S. Keskin, et al. performed Grand Canonical Monte Carlo and equilibrium molecular dynamics simulations to calculate adsorption isotherms and self-diffusivities of H₂/CH₄ mixtures in a Zn(bdc)(ted)_{0.5} MOF [214]. The results revealed that the MOF strongly adsorbed CH₄ and reduced hydrogen's diffusivity. It was also concluded that when the MOF was exposed to a H_2/CH_4 mixture, H_2 dragged the slowly diffusing CH_4 molecules through the pore. The authors also performed atomistic and continuum modeling of the Zn(bdc)(ted)_{0.5}/polymer (Hyflon[®], sulfonated and 6FDA-based polyimides, TeflonTM, and polytrimethylsilylpropyne) MMMs for this separation and concluded that the MMMs greatly enhanced the permeability of H2 without lowering the selectivity. More recently, in 2016, I. Erucar and S. Keskin [215] reviewed the progress made in this field and observed that there is still significant amount of work to be done to understand the effects of the incorporation of MOFs in MMMs for gas separations. The authors determined that the effects of the physical and chemical properties of MOFs (e.g., pore size, pore volume, surface area, type of organic linker, and compatibility between the MOF and the polymer) on MMM properties need to be studied in order to develop a structure-performance relationship to predict MMM performance from its constituents (MOFs and polymers). Size, dispersion, orientation of MOF particles in the MMM and their interaction with the polymer (e.g., pore blockage, polymer rigidification around MOF) are also areas that need more development to improve performance. In 2010, M. J. C. Ordoñez, et al. [216] and I. H. Musselman, et al. [217] report on the use of ZIF-8 (Figure 15) for the preparation of ZIF-8/Matrimid® MMMs for gas separations. The authors found that, as the ZIF loading increased up to 60% (w/w), the MMMs became more rigid and difficult to handle. From SEM images of the MMMs with 30% and 50% (w/w) ZIF-8 loadings (Figure 20), it was concluded that there was a strong interfacial contact between the ZIF-8 crystals and the polymer. However, as the loading increased to 80% (w/w), reduced interfacial contact lead to aggregation of the ZIF. For all the gases tested (H₂, CO₂, CH₄, N₂, O₂, and C₃H₈), the permeabilities increased with increasing ZIF loading up to 40% (w/w). At this loading, the permeabilities increased by 53% for C_3H_8 , 147% for H_2 , 158% for CO₂, 170% for O₂, 239% for N₂, and 271% for CH₄ with no significant changes in selectivity, which was attributed to the increase in free volume in the membrane due to the increase in distance between the polymer chains. Above 40% (w/w) loading, the permeabilities decreased significantly, up to 94% relative to the 40% loading, due to the increase in tortuosity and increased polymer density that forced the smaller molecules to diffuse around the ZIF crystals.

Progress in the preparation of MOF-based MMMs continued in 2011, when S. Basu, et al. prepared dense and asymmetric MMMs of $Cu_3(BTC)_2$, ZIF-8, and (Al) MIL-53 in Matrimid [218]. This was the first time that a MOF (MIL-53) with a breathing or gating effect was used for the preparation of MMMs. At 30 wt % loading, the dense MMMs exhibited permeability increments of 196% for $Cu_3(BTC)_2$, 209% for ZIF-8, and 218% for (Al) MIL-53 with a 15% increase in CO_2/CH_4 separations. The authors also noted that, although the structure varied significantly from MOF to MOF, the gas permeation properties of their MMMs were quite similar at any loading. The PDMS-coated asymmetric MMMs of $Cu_3(BTC)_2$ and (Al) MIL-53 showed increased CO_2/CH_4 mixture selectivities with increasing MOF loadings at any given CO_2/CH_4 feed composition. The ZIF-8 MMMs, however, did not show changes in selectivity with respect to the Matrimid. The increased selectivities were attributed to strong interactions of CO_2 with the MOF frameworks.

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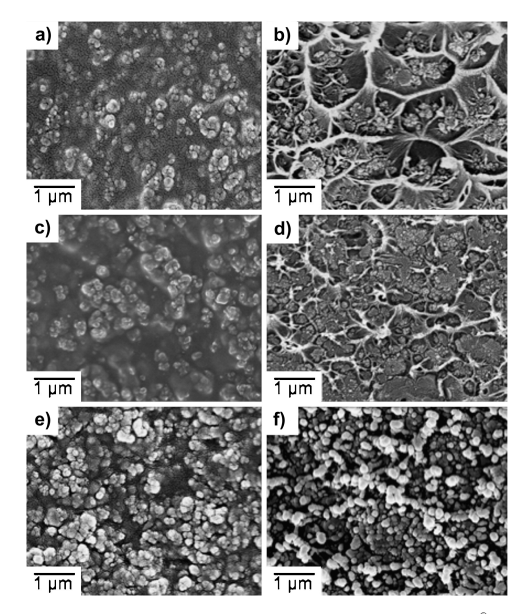


Figure 20. SEM images of surfaces (**a**,**c**,**e**) and cross-sections (**b**,**d**,**f**) of ZIF-8/Matrimid[®] MMMs: (**a**,**b**) 30%; (**c**,**d**) 50%; and (**e**,**f**) 80% (w/w). Adapted with permission from [216]. Copyright, 2010, Elsevier.

Another significant development in MOF-based MMMs came also in 2011, where X. L. Liu, et al. extended the application of MOF-based MMMs to the recovery of C_2 – C_5 alcohols from water by organophilic pervaporation [38]. The incorporation of the superhydrophobic and flexible framework of ZIF-8 into PMPS, a silicone rubber polymer, resulted in thin, defect-free ZIF-8/PMPS MMMs (Figure 21a) that were supported on alumina capillaries and showed a isobutanol/water selectivity of 40.1 and an isobutanol permeance of 7000 GPU. The authors also concluded that the energy required for pervaporation per unit of isobutanol was only half that of distillation, making the membrane-based separation more efficient. Membrane separation performance as a function of increased ZIF-8 loading was also explored and showed that the MMMs increased both selectivity and flux as the ZIF-8 loading increased. ZIF-8/PMPS MMMs also showed improved separation and flux properties for ethanol, n-propanol, n-butanol, and n-pentanol separations from water (Figure 21b). The authors also used ZIF-7 (Figure 15) to prepare ZIF-7/PMPS MMMs for the separation of isobutanol from water, but their results showed membranes with lower separation factors and lower isobutanol permeances. The reduced

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performance of the ZIF-7/PMPS MMMs was attributed to the reduced pore aperture of ZIF-7 (0.30 nm) and to its more rigid framework.

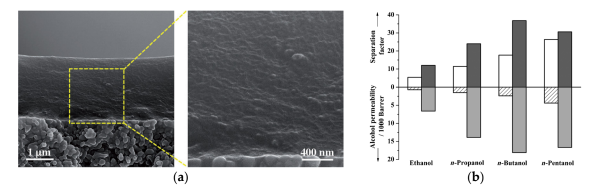


Figure 21. (a) SEM image of the cross-section of a 10% (w/w) ZIF-8/polymethylphenylsiloxane (PMPS) MMM on an alumina capillary support; (b) Alcohol/water separation factors and alcohol permeability of PMPS (open and line filled) and ZIF-8/PMPS MMMs (gray). Adapted with permission from [38]. Copyright, 2011, Wiley.

A new procedure to incorporate ZIFs into polymers came from the work of T. Yang, et al. in 2011 with the dispersion of colloidal ZIF-7 in PBI [205]. By keeping the ZIF-7 nanocrystals in a DMF solution, the authors avoided the aggregation of crystals when dried and eliminated the lengthy MOF dispersion steps that usually include sonication and stirring. With this strategy, MMMs with up to 50 wt % loadings were prepared and tested for gas mixture permeation from 35 to 180 °C at 7 bar. TEM and SEM images of a 50/50~(w/w) ZIF-7/PBI MMM (Figure 22a) showed a uniform dispersion of ZIF-7 nanocrystals with no significant agglomeration due to their coating by PBI that acts as a coating agent.

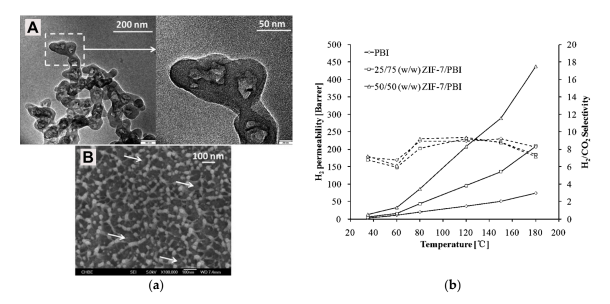


Figure 22. (a) Transmission electron microscopy (TEM) images (A) of the MMM suspension precipitated in methanol and SEM images (B) of the cross-section of a 50/50 (w/w) ZIF-7/Polybenzimidazole (PBI) MMM; (b) H_2/CO_2 mixture permeability and selectivity of PBI and ZIF-7/PBI MMMs with increasing temperatures. Reproduced with permission from [205]. Copyright, 2011, The Royal Society of Chemistry.

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Gas permeation with H₂/CO₂ mixtures (Figure 22b) showed that the MMMs retained H₂/CO₂ selectivity and increased H₂ permeability significantly with both ZIF loading and temperature increments. This same year, 2011, B. Zornoza, et al. reported on the preparation of a MMM using PSf (Udel®) and a functionalized MOF, (Al) NH₂-MIL-53 [219]. The aminated MOF was the first attempt to control the separation properties of the MOF-based MMM by decorating the channels of the MOF with amine functional groups (-NH₂) covalently bonded to the organic linker. SEM images of the cross-sections of MMMs containing 8, 16, 25, and 40 wt % NH₂-MIL-53 revealed that the polymer effectively wetted the micrometer-sized crystals (Figure 23a). Gas mixture (CO₂/CH₄) permeation experiments at 3 bar and 35 °C showed that the membranes had moderate CO₂ permeation increments and a small CH₄ permeability decrease that resulted in an increase in CO₂/CH₄ selectivity. At 10 bar, the MMM selectivity to CO₂/CH₄ (Figure 23b) improved significantly from 55 (PSf) to 110 (MMM) due to the opening of the MOF pores in the MMM (gating effect) that allows for more CO₂ to be adsorbed in the pores. A similar work comparing the CO₂/CH₄ separation properties of -NH₂ functionalized UiO-66 and MOF-199/6FDA-ODA MMMs to that of the corresponding neat MOF/6FDA-ODA MMMs was reported by O. G. Nik, et al. in 2012 [220]. The authors observed CO₂/CH₄ selectivity increments of 35% for the 25 wt % NH₂-MOF-199/6FDA-ODA MMM and 17% for the 25 wt % NH₂-UiO/6FDA-ODA MMM over the pure polymer. The increase in selectivity was attributed to polymer rigidification induced by the amine moiety in the MOFs.

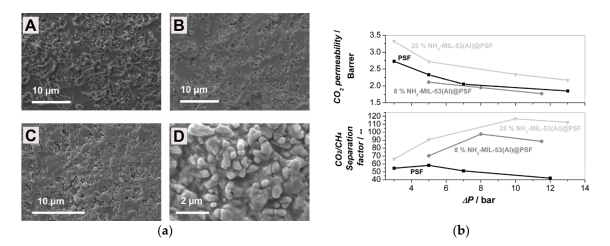


Figure 23. (a) SEM images of the cross-sections of 8 (A); 16 (B); 25 (C); and 40 (D) wt % NH₂-MIL-53/polysulfone (PSf) MMMs; (b) CO_2/CH_4 mixture separation performance of NH₂-MIL-53/PSf MMMs at -10 °C with increasing pressures from 3 to 13 bar. Reproduced with permission from [219]. Copyright, 2011, The Royal Society of Chemistry.

In this same year, B. Zornoza, et al. [221] tested MMMs comprised of PSf, silicalite-1 (S1C), and HKUST-1 or ZIF-8. The SEM images of the cross-sections of the MMMs (Figure 24) showed an even dispersion for the ZIF-8/PSf MMM and some polymer veins for the HKUST-1/PSf MMM. The addition of the S1C zeolite to both systems introduced regions with a slightly different morphology but not a significant S1C/PSf affinity in the form of polymer veins or plastic deformation was observed. Permeability experiments showed mixed results depending on the type of MOF used and the gas separation performed (Figure 25). For the ZIF-8/S1C/PSf MMM, the combination of the ZIF and S1C improved the MMM performance for separations based on gas diffusivity differences. For the HKST-1/S1C/PSf MMM, the performance of the MMM improved for CO₂ separations probably due to the affinity of the MOF for CO₂. The authors conclude that the incorporation of the zeolite in the MMMs may have a synergistic effect on the MMM that leads to improved separation properties. Although the mechanism of this effect is not clear, the results suggest S1C has an effect on the properties of the MMM.

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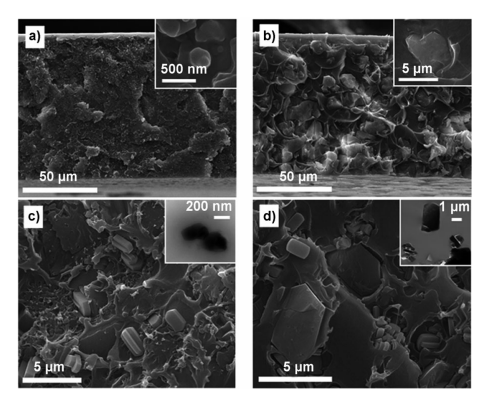


Figure 24. SEM images of the cross-sections of MMMs: (a) 16 wt % ZIF-8/PSf; (b) 16 wt % HKUST-1/PSf; (c) 8 wt % ZIF-8 + 8 wt % S1C/PSf; and (d) 8 wt % HKUST-1 + 8 wt % S1C/PSf. Reproduced with permission from [221], Copyright, Year, Wiley.

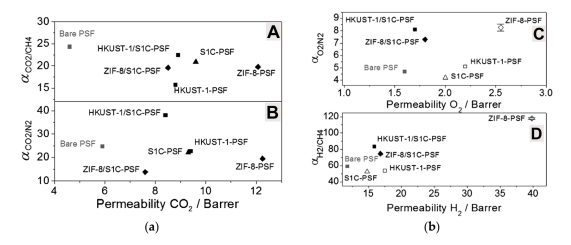


Figure 25. (a) CO_2/CH_4 (A) and CO_2/N_2 (B) selectivities and CO_2 permeabilities for PSf and the MMMs; (b) O_2/N_2 (C) and H_2/CH_4 (D) selectivities and H_2 permeabilities for PSf and the MMMs. Reproduced with permission from [221]. Copyright, 2011, Wiley.

The development of MOF-based MMMs was boosted in 2012 when, H. B. T. Jeazet, et al. reported on their work with MIL-101, a hydrolytically stable MOF, used in conjunction with PSf in the preparation of MMMs for O_2/N_2 separations [222]. The low hydrothermal stability of MOFs (e.g., MOF-5, IRMOF series, HKUST-1) was recognized to be a potential problem for industrial gas separations and was addressed by the authors with the use of a water stable MOF (MIL-101). MMMs with 8, 16, and 24 wt % MIL-101 exhibited increased permeability with increasing MOF loading, while retaining the O_2/N_2 selectivity of the membrane. Oxygen permeabilities increased from 1.5 Barrer (PSf) to 6 Barrer (24 wt % MIL-101/PSf MMM), while the O_2/N_2 selectivity remained

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between 5 and 6. Another important development in 2012 towards the potential industrial utilization of MOF-based MMMs came from the work of Y. Dai, et al. with the preparation of the first MOF-based hollow fiber MMMs [223]. The 13 wt % ZIF-8/UltemTM hollow fiber MMMs were coated with PDMS to seal surface defects and then tested for CO₂/N₂ separations (flue gas) over a range of temperatures and pressures. The hollow fiber MMMs exhibited an increase in the CO_2/N_2 separation performance at 25 °C and 7 bar over that for pure gases (MMM: $CO_2/N_2 = 44$, $CO_2 = 18$ GPUs; PSf: $CO_2/N_2 = 36$, $CO_2 = 11$ GPUs) and a decrease in CO_2/N_2 selectivity for mixed (20% CO_2 in N_2) gases (MMM: ideal $CO_2/N_2 = 44$, $CO_2 = 18$ GPU; MMM: gas mixture $CO_2/N_2 = 32$, $CO_2 = 26$ GPU). Nevertheless, the preparation of MOF-containing hollow fiber MMMs was a significant milestone towards the implementation of MMMs in large-scale gas separations. Another important development in 2012 came from the work of T. Yang, et al. with the testing of hollow fiber ZIF-8/PBI MMMs with ZIF-8 loadings up to 33 wt % for the separation of H_2/CO_2 (50/50) gas mixtures at 7 bar and temperatures up to 180 °C [224]. The SEM images of the cross-sections of the hollow fiber MMMs (Figure 26a) showed a thin ZIF-8/PBI selective layer on the shell side of the fiber bearing a macroporous support. Gas permeation data with H₂/CO₂ mixtures at temperatures from 25 to 180 °C (Figure 26b) showed a continuous increase in H₂ permeance with increasing ZIF-8 loading, from 60 GPU (10 wt % ZIF-8 loading) to 200 GPU (33 wt % ZIF-8 loading). Selectivities for H₂/CO₂ at 180 °C ranged from 8 (33 wt % ZIF-8 loading) to 12 (10 wt % ZIF-8 loading). The decrease in selectivity was attributed to the larger pore aperture of ZIF-8.

Although most of the work performed until 2012 concentrated on the separation of permanent gases, the separation of volatile organic compounds with MOF-based MMMs was explored this year by C. Zhang, et al. with the preparation of 16.4 wt % to 48.0 wt % ZIF-8/6FDA-DAM MMMs and the testing of these MMMs in propylene/propane separations with pure gases and gas mixtures (1:1 C_3H_6 : C_3H_8) [11]. To test the MMMs, the temperature was set to 35 °C and the pressure was increased from 0.7 to 2.8 bar. The membrane stage cut (θ) was kept to <1% to minimize concentration polarization of the less permeable gas (i.e., the retentate) when gas mixtures were tested. The omission of removing the retentate results in a non-constant feed composition enriched with the retentate that reduces the selectivity of the membrane. For this reason, a portion of the feed close to exposed membrane area is removed at a rate that is determined by the flux of the gases across the membrane. The MMMs showed a significant improvement in gas separation over the pure polymer for both pure and gas mixtures that put the MMMs above the Robeson [118] upper bound for C_3H_6/C_3H_8 separations. A gas mixture selectivity of about 18 for C_3H_6/C_3H_8 and a C_3H_6 permeability of 42 Barrer were obtained for the 48.0 wt % ZIF-8/6FDA-DAM MMMs, whereas a C_3H_6/C_3H_8 selectivity of about 8 and a C_3H_6 permeability of about 11 Barrer was obtained for the 6FDA-DAM membrane.

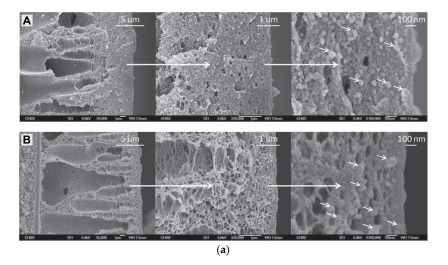


Figure 26. Cont.

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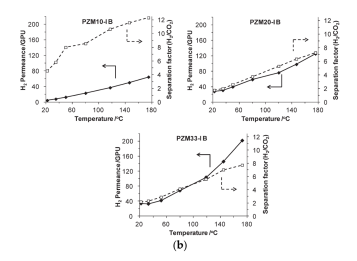


Figure 26. (a) SEM images of the cross-sections of 10 (A) and 33 (B) wt % ZIF-8/PBI hollow fiber MMMs (arrows point to ZIF-8 nanoparticles); (b) H_2/CO_2 (50/50) gas mixture separation of 10 wt % (PZM10-IB), 20 wt % (PZM20-IB), and 33 wt % (PZM33-IB) hollow fiber MMMs at 7 bar with temperature increasing from 25 to 180 °C. Reproduced with permission from [224]. Copyright, 2012, Wiley.

Until 2013, most polymers used for MOF-based MMMs were dense, with limited porosity. This approach changed when A. F. Bushell, et al. prepared 11 vol %, 28 vol %, 36 vol %, and 43 vol % ZIF-8/PIM-1 MMMs [225]. PIM-1 (Figure 27) [226,227] is a polymer with intrinsic microporosity that results from the combination of a spirobisindane [228] as co-monomer with tetrafluorophthalonitrile.

Figure 27. Synthesis of the polymer of intrinsic microporosity 1 (PIM-1) from 3,3,3',3', -tetramethyl-1,1''-spirobisindane-5,5',6,6'-tetrol (**a**) and 2,3,5,6-tetrafluorophthalonitrile (**b**).

The polymer is soluble in common organic solvents and possesses permeabilities on the order of thousands of Barrer for H_2 (1300 Barrer) and CO_2 (2300 Barrer) that surpass the upper bound. The polymer also exhibits good H_2/N_2 and CO_2/N_2 selectivities of 14 and 25, respectively, that make it attractive for membrane-based gas separations [227]. Permeability experiments with the MMMs at 1 bar and 22 °C showed that the MMMs improved the permeability and selectivity of smaller gases (e.g., H_2 , H_2 , H_2 , H_2 , H_3 , H_4 , H

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Table 2. Permeabilities and selectivities of 0, 11, 28, 36, and 43 vol % ZIF-8/PIM-1 MMMs. Adapted with
permission from [225]. Copyright, 2013, Elsevier.

ZIF-8 (vol %)	Permeability (Barrer)					Selectivity					
	H ₂	He	O ₂	N ₂	CO ₂	CH ₄	H ₂ /N ₂	H ₂ /CH ₄	He/N ₂	O ₂ /N ₂	CO ₂ /CH ₄
0	1630	760	580	180	4390	310	9.1	5.3	4.2	3.2	14.2
11	2560	1310	820	250	4815	320	10.2	8.0	5.2	3.3	15.0
28	2980	1430	870	195	4270	230	15.2	13.0	7.3	4.5	18.6
36	5745	2930	1640	380	6820	510	15.1	11.3	7.7	4.3	13.4
43	6680	3180	1680	350	6300	430	19.1	15.5	9.1	4.8	14.7

A second PALS study on the correlation of gas permeability with free volume came also in 2013 from H. B. T. Jeazet, et al. using 7.5, 14, 19, and 24 wt % (Cr) MIL-101/PSf MMMs for O_2 , N_2 , CO_2 , and CH_4 separations [229]. SEM images (Figure 28a) of the MMMs showed even MOF distributions and good wetting of the particles by the polymer. As with other MOF-based MMMs, permeabilities at 30 °C and 3 bar increased with increasing MOF loading, but selectivities remained constant.

Using PALS studies and the Tao and Eldrup standard model that correlates the pick-off lifetime of ortho-positronium to the free volume hole size in polymers, the authors attributed the increase in permeability to the free volume in the polymer and the added large free volume in the MIL-101 particles (Figure 28b). PALS also indicated that the polymer chains did not penetrate into the pores and that both the polymer and the MOF structures were not affected. The results also suggested that neither defects nor additional free volume in the polymer and at the MOF/polymer interface was introduced. This same year (2013), T. Singh, et al. [230] performed a finite element analysis on single component permeation in MMMs that resulted in contradictions in the fundamental assumptions of the Maxwell, Bruggeman, Pal, Lewis-Nielsen, and other models [231,232] that describe or predict gas transport (permeability and selectivity) in MMMs. The hypothesis for their work was that the effects of the interfacial adsorption equilibrium constant (K) on permeability were not taken into account in the models by implicitly considering $K \approx 1$, when in reality most filler/polymer interfaces may not have interfacial equilibrium constants of 1 ($K \neq 1$). The assumption that $K \approx 1$ ultimately leads to an erroneous interpretation that deviations of experimental results from the models are due to "non-idealities", polymer rigidification, or defects at the filler/polymer interface. These observations about the models are even more important for MOF-based MMMs because MOFs have better compatibility with the polymer, which greatly reduces defects in the membrane. Singh, et al. concluded that there was a need to perform a "rigorous" prediction of permeation in ideal MMMs to separate the true dependence of the permeability on the diffusivities in the matrix and filler phases, the filler/polymer interface adsorption equilibrium, and the volume fraction of the filler. With these predictions, it would then be possible to reliably interpret MMM data and separate the effects of the filler properties from non-idealities or defects. Singh, et al. developed an "exact model" that takes into account the effects of interfacial equilibrium between the filler and the matrix and the differences in diffusivity between the filler and the polymer. The solution to their model was performed by finite element analysis using COMSOL Multiphysics. From their model, they conclude that the individual values of K and the diffusivity ratio D_f/D_m determine the MMM permeability and not the combined permeability ratio $P_f/P_m = KD_f/D_m$. In this work, the authors also back-calculated the permeability of CO₂ in ZIF-8 from the experimental data of 6 different ZIF-8/polymer (Torlon[®], UltemTM, Matrimid[®], 6FDA-DAM:DABA 4:1, 6FDA-DAM, and PDMS) MMMs using the Maxwell and Lewis-Nielsen models. The results show that the models fail to predict the permeability of CO₂ in ZIF-8 at different loadings (Figure 29).

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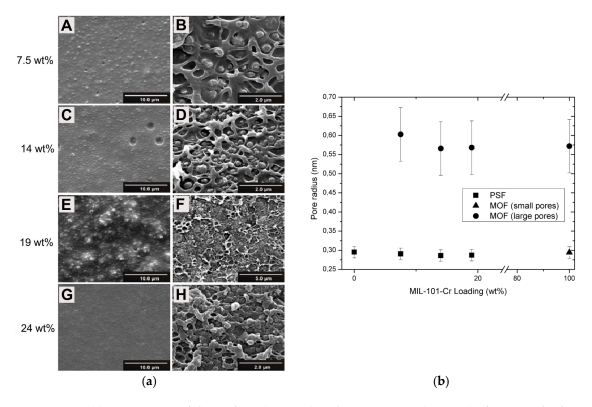


Figure 28. (a) SEM images of the surfaces (A,C,E,G) and cross-sections (B,D,F,H) of MIL-101/PSf MMMs; (b) Pore radii of the free volume with increasing MIL-101 loading in PSf. Adapted with permission from [229]. Copyright, 2013, MDPI under the Creative Commons Attribution License (CCBY).

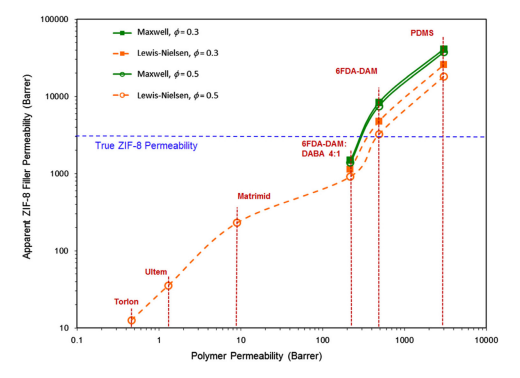


Figure 29. Back-calculated apparent CO_2 permeabilities of ZIF-8 according to polymer permeability and ZIF loading (ϕ). True ZIF-8 permeability obtained from experimental data. Reproduced with permission from [230]. Copyright, 2013, Elsevier.

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A new approach into the preparation of MOF-based MMMs was also introduced in 2013 by L. Hao, et al. [233] with the preparation of RTILs/ZIF-8/polyRTILs MMMs aiming to improve the performance of RTILs/polyRTILs membranes with the addition of ZIF-8. RTILs have been shown to have a good CO₂ affinity making them attractive for CO₂/N₂ and CO₂/CH₄ separations, but for their application in gas separations they needed to be immobilized. An approach to immobilize RTILs is the preparation of UV-polymerizable RTILs to prepare polyRTILs membranes. Although polyRTILs retain much of the selectivity of RTILs, the resulting membranes exhibit lower CO₂ permeabilities than the RTILs, for this reason polyRTILs are doped with RTILs, but further permeability improvements could be achieved with the incorporation of porous fillers (e.g., SAPO-34, ZIF-8). The authors prepared ZIF-8/polyRTILs MMMs and RTILs/ZIF-8/polyRTILs MMMs with 0, 10, 20, and 25 wt % ZIF-8 loadings keeping the RTILs/polyRTILs ratio to 1:2 for all the RTILs/ZIF-8/polyRTILs MMMs. The free RTILs in the MMMs were [emim][NTf₂], [emim][BF₄], and [emim][B(CN)₄] and the UV-polymerizable RTIL was [vbim][NTF₂]. Gas permeation experiments were performed at 35 °C and 3.5 bar with CO₂, N₂, CH₄, and CO₂/N₂ and CO₂/CH₄ mixtures at 50/50 mol %. The results showed a retention of CO₂/N₂ and CO₂/CH₄ selectivities for the ZIF-8/polyRTILs at 20 and 12, respectively, with ZIF-8 loadings up to 18.9 wt %. For these MMMs, CO₂ permeabilities increased from 101 (0 wt % ZIF-8) to 199 Barrer (18.9 wt % ZIF-8), a 100% increase in CO₂ permeability. The incorporation of the free RTILs in the ZIF-8/polyRTILs MMMs (RTILs/ZIF-8/polyRTILs) increased CO₂ permeability in all cases with a retention or small decrease in selectivity. The maximum CO₂ permeability achieved was with the addition of free [emim][B(CN)₄] to the 25 wt % ZIF-8/polyRTILs MMM, this membrane provided a CO₂ permeability increase of 190% from the RTILs/polyRTILs MMM (at 0 wt % ZIF-8 loading: $CO_2 = 365 \text{ Barrer}, CO_2/N_2 = 29.9, CO_2/CH_4 = 15.8; \text{ at 25 wt } \% \text{ ZIF-8 loading: } CO_2 = 1062 \text{ Barrer},$ $CO_2/N_2 = 24.2$, $CO_2/CH_4 = 12.3$). Gas mixture separations for this membrane showed a slight decrease in selectivities ($CO_2/N_2 = 20.1$, $CO_2/CH_4 = 11.6$), as is the case with most polymer membranes. The results of this work matched very well with the predictions of the Maxwell model and proved that the combination of free RTILs and ZIF-8 improve the separation properties of the polyRTILs in the form of RTILs/ZIF-8/polyRTILs MMMs.

Although the most common mode of operation for a polymer membrane is to permeate the smaller gas molecule rather than the larger one, some types of membranes allow larger molecules to permeate faster than smaller molecules, reversing the selectivity of the membrane. In 2014, R. Abedini, et al. reported on the use of MIL-53 and PMP in the preparation of MIL-53/PMP MMMs for reverse CO_2/H_2 separations [234]. The MMMs showed increased CO_2 solubility with increasing MOF loading up to 40 wt %, which led to an increase in CO_2/H_2 separation and CO_2 permeability. The CO_2/H_2 selectivity increased from 8.86 for PMP to 16.92 for 35 wt % MIL-53/PMP, and the CO_2 permeability increased from 98.74 for PMP to 225.21 for 35 wt % MIL-53/PMP. Diffusivities, however, experienced a small decreased with MOF loading. Also in 2014, L. Diestel, et al. performed CO_2/CH_4 , CO_2/N_2 , H_2/CO_2 , and O_2/N_2 pure and gas mixture separation studies on pure ZIF-8, pure polymer (PMPS), and the 8.3 wt % ZIF-8/PMPS MMM prepared on α -alumina supports [235]. In all cases, the alumina support (70 nm particles) was immersed in the synthesis (ZIF-8 membrane from seeding and secondary growth) or casting solutions (PMPS and MMM) to coat the surface. SEM images (Figure 30) show that thin membranes (<20 μ m) were deposited onto the substrate.

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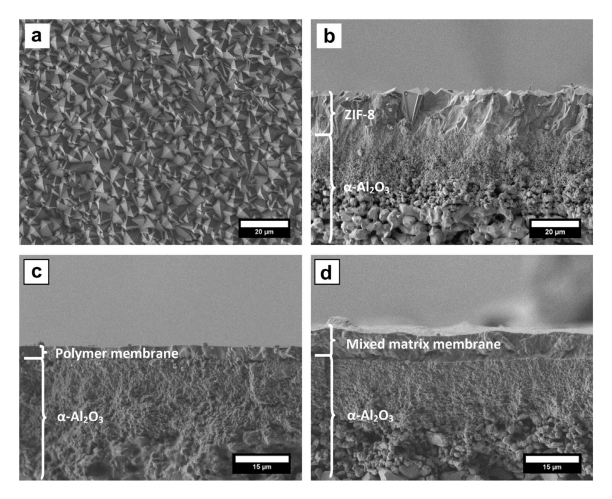


Figure 30. SEM images of the surface (a) and cross-section (b) of a ZIF-8 membrane on an α -Al₂O₃ support. The cross-sections of the PMPS and the 8.3 wt % ZIF-8/PMPS MMM are shown in (c) and (d), respectively. Reproduced with permission from [235]. Copyright, 2014, Elsevier.

The membranes were tested at $\Delta P = 0.2$ bar, and the permeates were analyzed using a GC. Gas permeation results indicated that the ZIF-8 membrane performed molecular sieving, but without a sharp cut off due to the flexibility of the ZIF. The authors also observed that gas separations for the PMPS membrane and the MMM were driven mainly by a solubility process. The observed gas permeabilities (in Barrer) were: ZIF-8 ($H_2 = 2598$, $CH_4 = 234$, $CO_2 = 623$, $N_2 = 260$, $O_2 = 468$), PMPS $(H_2 = 34, CH_4 = 37, CO_2 = 305, N_2 = 35, O_2 = 58)$, and MMM $(H_2 = 103, CH_4 = 114, CO_2 = 827, N_2 = 118, CH_4 = 114, CO_2 = 827, CH_4 = 114, CH_4 = 114, CO_2 = 827, CH_4 = 114, CH_4$ O₂ = 190). From the three membranes tested, the authors found that the MMM had identical selectivity to the polymer, but with higher fluxes, which was attributed to the increased free volume in the MMM by the presence of the MOF. The authors also did not observe significant variations in ideal and gas mixture selectivities among the membranes tested. In another work carried out by L. Diestel, et al. in 2015 [236] the authors performed a comparison study using experimental data of H₂ and CH₄ gas permeation properties and the Maxwell model predictions. The authors prepared membranes of ZIF-8, ZIF-90, Matrimid[®], and the 10, 20, and 30 vol % ZIF-8/Matrimid[®] and ZIF-90/Matrimid[®] MMMs on porous α alumina supports with a pore size of 70 nm. Permeability data for both ZIF-8 and ZIF-90 MMMs showed that at 10 and 20 vol % ZIF loadings the MMMs exhibited lower permeabilities and higher H₂/CH₄ selectivities than the polymer and the respective ZIF membrane and were significantly different to what the Maxwell model would predict. At 30 vol % loadings the MMMs showed less selectivity and reduced permeability than the polymer and still deviated significantly from the Maxwell model. The authors attributed the improved selectivity and reduced permeability of the MMMs to the

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limited flexibility of the ZIF framework resulting from the blockage of linker distortions in the ZIFs caused by the polymer/ZIF interactions.

More progress was made when, in 2014, C. Duan, et al. reported on the effect of thermal treatment on gas permeation [237]. The authors prepared Cu₃(BTC)₂ MMMs (0 to 50 wt % loadings) from DMAc using Matrimid[®] and ODPA-TMPDA polymers as well as ZIF-8 MMMs with ODPA-TMPDA. The membranes were dried for 1 day at temperatures ranging from 150 °C to 220 °C before gas permeation tests were performed. The authors observed that, for both the polymer and the MMMs, an increase in permeability was achieved when the membranes were dried above the boiling point of the solvent (DMAc) and that above this temperature no significant increase in gas permeability was observed. It was concluded that incomplete removal of solvent from the membranes and from the MOFs reduced gas permeability in MMMs. Another contribution in the MOF-based MMM field came from N. A. Nordin, et al. in 2014 [238]. The authors prepared asymmetric 5 wt % ZIF-8/PSf membranes from NMP/THF using ZIF-8 crystals that they synthesized with sizes of 100, 300, and 500 nm. Testing the membranes at 27 °C and 4 bar with CH₄ and CO₂ revealed that the MOF particle size played an important role in the performance of the MMMs. ZIF-8 particles of 100 nm suffered from partial pore blockage, increased tortuosity due to the large number of small particles, and polymer rigidification that resulted in increased CO₂/CH₄ selectivity, compared to the MMMs with 300 and 500 nm ZIF-8 particles, and a reduction in CO₂ permeability. The performance of this membrane, however, was still inferior to that of PSf, with 58% reduction in permeability and 38% reduction in selectivity with respect to PSf. Larger ZIF-8 particles (300 and 500 nm) introduced non-selective voids at the interface that increased the permeability of the gases and reduced the selectivity of the MMM.

Another contribution came in 2014 from the work of E. V. Perez, et al. with MOP-18/Matrimid® MMMs [126]. This was the first time MOP-18 nanocages (Figure 14b) were incorporated into a polymer matrix and tested for gas separation at high pressures and elevated temperatures. Since the crystal or particle size is the limiting factor for the degree of dispersion of MOFs and ZIFs in polymers, independent of the procedure taken for its dispersion, the use of molecular cages in MMMs was the next logical step to follow. MOPs and MOCs are molecular cages that can be dissolved in organic solvents and, therefore, dispersed in polymer matrices at the molecular level, achieving greater particle (molecule)/polymer contact than with ZIFs and MOFs. Perez, et al. synthesized MOP-18 and prepared 0, 23, 33, 44, and 80 wt % MOP-18/Matrimid[®] MMMs that did not show aggregation or gross defects. SEM images of the membranes showed smooth and even morphologies on both the surface and cross-sections of the MMMs (Figure 31) without signs of aggregation or gross defects, even at the extremely high MOP-18 loading of 80 wt %. Compared to MOF-based MMMs, the MOP-18-based MMMs did not show macroscopic plastic deformation or rigidification at the MOP/polymer interface. The membranes were tested with H₂ and CO₂ at 35 and 70 °C and from 3 to 50 bar. Other gases, like N₂, O₂, CH₄, C₃H₆, and C₃H₈, were also tested but at lower pressures (2.6 bar). Gas permeation results (Figure 32a) showed that the 44 wt % MOP-18/Matrimid[®] MMM increased the permeability for small molecules, such as H₂, CO₂, O₂, and N₂, by up to 100%. Larger molecules, however, experienced higher increases in permeability, with CH₄, C₃H₈, and C₃H₆ increasing by 330%, 433%, and 770%, respectively, indicating the MOP's affinity for hydrocarbons that was attributed to the presence of new sorption sites associated with the MOP-18 molecules like the pore, the surface of the core, and the alkyl chains that extend from the organic linker. Selectivities (Figure 32b) decreased for most gas pairs except for C_3H_8/N_2 and C_3H_6/N_2 , which experienced significant increase in selectivity relative to the polymer. High pressure experiments with H₂ up to 30 bar and CO₂ up to 50 bar at 35 and 70 °C also demonstrated that the MMMs were mechanically robust and that no defects were present at the MOP-18/Matrimid[®] interface that would reduce the selectivity since H₂/CO₂ selectivites at 30 bar and 70 °C decreased only slightly from 2.42 (Matrimid®) to 2.04 (23 wt % MOP-18/Matrimid® MMM). High pressure CO₂-induced plasticization studies performed with the polymer and the MMMs also showed that

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the degree of plasticization depends largely on temperature and that, at higher temperatures, the plasticization pressure increases for both the polymer and the MMM (Figure 33).

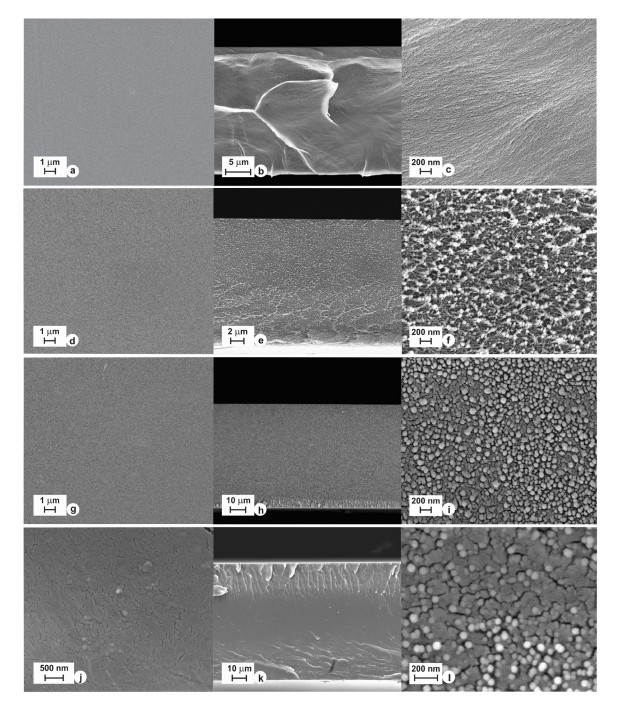


Figure 31. SEM images of the surfaces (a,d,g,j), cross-sections at low magnification (b,e,h,k), and cross-sections at high magnification (c,f,i,l) of 0 (a-c); 23 (d-f); 44 (g-i); and 80 (j-l) wt % MOP-18/Matrimid[®] MMMs. Reproduced with permission from [126]. Copyright, 2014, Elsevier.

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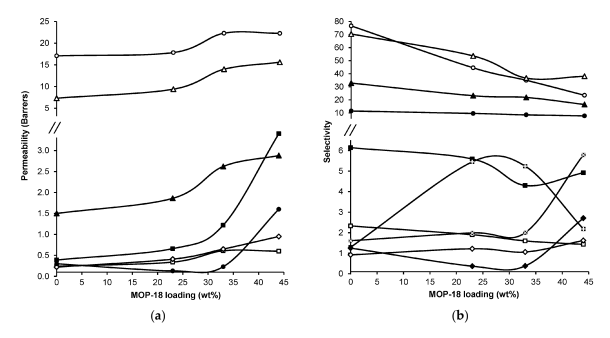


Figure 32. (a) Permeabilities of MOP-18/Matrimid[®] MMMs at 35 °C and 2.6 bar: H_2 (○), CO_2 (△), O_2 (▲), C_3H_6 (■), N_2 (□), CH_4 (♦), and C_3H_8 (•); (b) Ideal selectivities of MOP-18/Matrimid[®] MMMs: O_2/N_2 (■), H_2/CO_2 (□), CH_4/N_2 (♦), C_3H_8/N_2 (♦), C_3H_6/N_2 (x), C_3H_6/C_3H_8 (+), H_2/CH_4 (○), H_2/N_2 (△), CO_2/CH_4 (▲), and H_2/O_2 (•). Reproduced with permission from [126]. Copyright, 2014, Elsevier.

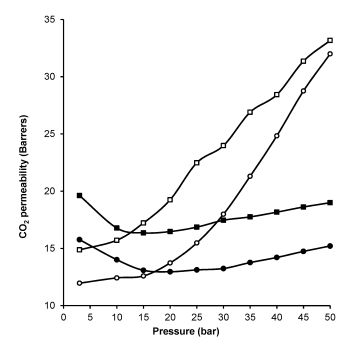


Figure 33. Permeability-pressure isotherms of CO_2 in Matrimid[®] (○ and •) and the 23 wt % MOP-18/Matrimid[®] MMM (□ and ■). Open symbols for 35 °C and solid symbols for 70 °C. Reproduced with permission from [126]. Copyright, 2014, Elsevier.

The authors also performed high pressure and high temperature gas adsorption studies on MOP-18 crystals, Matrimid $^{\text{@}}$, and the 23 wt % MOP-18/Matrimid $^{\text{@}}$ MMM. The results showed an impressive CO₂ uptake at high pressure (30 bar) and elevated temperatures (Figure 34) that was comparable to that of MOFs and activated carbons. The sudden uptake of CO₂ was attributed to

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the opening of the pores of the MOF that allowed the penetration of CO_2 molecules into the pore. The results also showed that MOP-18 in the MMM was able to adsorb more CO_2 than in its crystalline state. This behavior was attributed to the blockage of the pores by the long alkyl chains when MOP-18 is the crystalline form, whereas incorporation of the MOP into the polymer extends the chains from the surface of the MOP into the matrix, thereby exposing the pores.

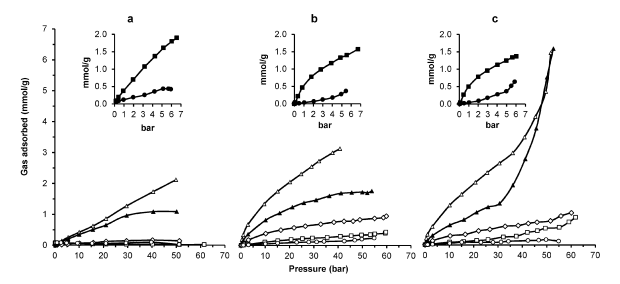


Figure 34. Gas adsorption isotherms of N_2 (\square), CH_4 (\lozenge), H_2 (\bigcirc), CO_2 (\triangle), C_3H_6 (\blacksquare , insert) and C_3H_8 (\bullet , insert) at 35 °C and CO_2 (\blacktriangle) at 70 °C in: (a) MOP-18 crystals; (b) Matrimid[®] membranes; and (c) 23 wt % MOP-18/Matrimid[®] MMMs. Reproduced with permission from [126]. Copyright, 2014, Elsevier.

Another step in the development of MOF-based- MMMs for industrial applications came also in 2014, when S. Shahid, et al. reported on the testing of MOF-based MMMs at low and high pressures (40 bar) with pure and CO_2/CH_4 gas mixtures at 35 °C [239,240]. The testing of MMMs at high pressures is important since it puts to the test the strength of the MOF/polymer interface. S. Shahid, et al. prepared flat MMMs of MIL-53, ZIF-8, and Cu_3BTC_2 with Matrimid® with loadings up to 30 wt % and dried them at 200 °C, the MMMs showed an increased CO_2/CH_4 selectivity due to the interactions of CO_2 with the MOFs that increased its permeability in the membranes and to the sieving properties of the MMMs for CH_4 that decreased its permeability. Mixed gas separation, however, showed a significant decrease in the CO_2/CH_4 selectivity for the polymer with increasing pressure due to plasticization. The effects of the CO_2 -induced plasticization on the polymer were reduced with MOF loading increments that, in turn, increased the MMM's CO_2/CH_4 selectivity from 15 to 40.

Although SEM images provide a good representation of the distribution of the MOF in the membrane, the spatial distribution of the MOF particles in MMMs was not performed until 2014 when T. Rodenas, et al. performed tomographic SEM imaging of the MMM cross-sections milled with Ga⁺ ions (FIB-SEM) [241]. The aim of the work was to assess the distribution of the filler and its contact with the polymer by ion milling the MMMs, imaging the exposed cross-sections, and reconstructing a 3D image from a set of 2D SEM images. Ion milling (FIB) effectively avoids the introduction of artifacts and rough surfaces that generally result from freeze-fracture, allowing visualization of the spatial distribution of the MOF in the polymer as well as large defects (i.e., voids), if present, that may originate from the casting process. The authors synthesized (Al) NH₂-MIL-53 crystals and then prepared 25 wt % NH₂-MIL-53/Matrimid[®] MMMs that were FIB milled for SEM imaging in the BSE mode (Figure 35). In the BSE mode, the intensity scales with the atomic number of the elements present in the sample, making it suitable for imaging areas with different chemical compositions (e.g., the MOF and the polymer). Another advantage of this mode is that it provides a more uniform contrast than the

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in-lens mode. The images show a random and even distribution of the MOF in the matrix (Figure 35b) with the presence of voids at the MOF agglomerates (Figure 35c). Upon alignment of the stack of 2D SEM images, the 3D structure of the analyzed volume was reconstructed (Figure 36a,c,d) and segmentation of each phase (polymer, MOF, voids) by image thresholding generated the volumes corresponding to the MOF (Figure 36b) and voids (Figure 36e).

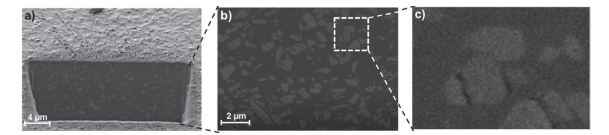


Figure 35. SEM images of a 25 wt % NH₂-MIL-53/Matrimid[®] MMM cross-section: (a) low magnification image of the hole milled with focused ion beam (FIB); (b) representative image of the cross-section obtained in backscattered electron (BSE) mode; (c) enlarged view showing the contrast between the polymer, MOF, and voids. Reproduced with permission from [241]. Copyright, 2014, Wiley.

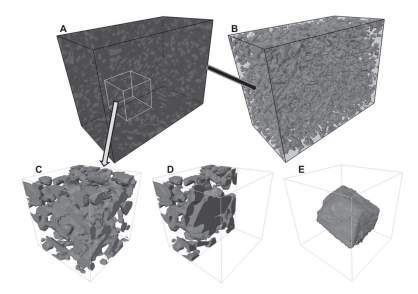


Figure 36. A 25 wt % NH₂-MIL-53/Matrimid[®] MMM imaged with FIB-SEM: (**A**) 3D reconstructed volume of the studied portion; (**B**) volume corresponding to MOF crystals; (**C**) magnification of selected volume in (**A**); (**D**) half volume of (**C**); (**E**) surface-rendered view of volume corresponding to voids. Reproduced with permission from [241]. Copyright, 2014, Wiley.

In 2014, the application of MOF-based MMMs was expanded by the work of Z. Y. Yeo, et al. with the utilization of ZIF-8/PES MMMs as supports for the growth of ZIF-8 membranes on the surface of the MMM via secondary growth [242]. Due to the limited affinity of a MOF for an inorganic support (e.g., Al_2O_3 or TiO_2), the use of a MMM as an intermediary support facilitates the adhesion of the selective layer (e.g., ZIF-8) due to the favorable interactions between the organic ligands of the MOF and the polymer. The authors prepared 5 and 10 wt % ZIF-8/PES MMMs as supports and performed rub seeding and dip coating in a 1% ZIF-8 solution to provide nucleation sites. The ZIF-8 membrane was then grown by solvothermal synthesis in a methanol solution containing $ZnCl_2$, 2-methylimidazole, and sodium formate. SEM images of the cross-sections of the ZIF-8 membranes (Figure 37a) showed no noticeable interfaces between the ZIF-8 layer and the MMM, indicating that good adhesion of the ZIF to the MMM was achieved. Flexible, continuous, and homogeneous ZIF-8 layers about 6–8 μ m

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in thickness were obtained after 6 h. SEM of the surface of the ZIF-8 layer (Figure 37b) showed a continuous, dense, well inter-grown, and compact layer with no gross defects. Gas permeation for $\rm CO_2$ and $\rm CH_4$ obtained for the 5 and 10 wt % ZIF-8/PES MMMs showed an increase in gas flux with constant ideal $\rm CO_2/CH_4$ selectivity. These properties were greatly modified when the ZIF-8 layer was grown onto the polymer and the MMM. For these membranes, the CH₄ flux decreased 50%, and the ideal $\rm CO_2/CH_4$ selectivity increased by 60%, from 9 for PES to 15 for the ZIF-8 layer on PES and the MMM. Gas mixture (50/50 $\rm CO_2/CH_4$) separation showed a slight increase in $\rm CO_2/CH_4$ separation for the ZIF-8 layer grown on the MMM with respect to the layer grown on the polymer (ZIF-8 on PES, $\rm CO_2/CH_4 = 13.8$; ZIF-8 on MMM, $\rm CO_2/CH_4 = 14.6$).

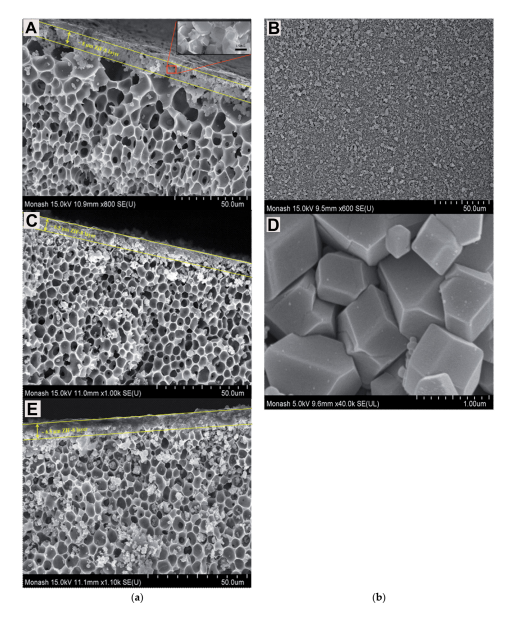


Figure 37. (a) SEM images of the cross-sections of continuous ZIF-8 membranes grown on PES (A); 5 wt % ZIF-8/PES MMM (C), and 10 wt % ZIF-8/PES MMM (E); (b) SEM images of the surface of the ZIF-8 membrane grown on an MMM: low (B) and high (D) magnification. Adapted with permission from [242]. Copyright, 2014, The Royal Society of Chemistry.

Another important development in 2014 was the suppression of aging in MMMs that was reported in the work of C. H. Lau, et al. [243]. In this work, three highly permeable super glassy polymers

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(PTMSP, PMP, and PIM-1) and their PAF-1 MMMs were tested for CO₂ permeation over a period of 240 days. Remarkably, the MMMs exhibited less than 7% decrease in permeability over the 240 days period while the pristine polymers experienced between 38% and 62% loss in permeability due to aging in the same period of time (Figure 38). This work shows that additives in MMMs have the capability to tune and freeze the packing of the polymer chains to prevent loss of free volume and gas permeability.

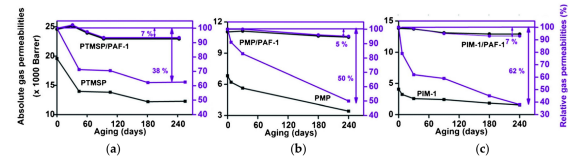


Figure 38. Absolute (black) and relative (purple) CO₂ permeabilities of: (a) PAF-1/poly(1-trimethylsilil-1-propyne) (PTMSP), (b) PAF-1/poly(4-methyl-1-pentyne) (PMP), and (c) PAF-1/ and PIM-1. Vertical arrows reperesent the aging degree of the pristine polymers (■) and the MMMs (●). Reproduced with permission from [243]. Copyright, 2014, Wiley.

The fight against membrane aging continued in 2015 with the incorporation of a series of MOPs into PTMSP to reduce aging and permeability losses. Extending on the work of E. V. Perez, et al. on the dispersion of single molecular cages in a polymer to prepare MOP-18/Matrimid[®] MMMs [126], M. Kitchin, et al. [244] prepared a series of MOPs to reduce membrane aging. In the work of M. Kitchin, et al., *t*-Bu-MOP (*tert*-butyl MOP), DEG-MOP (diethylene glycol MOP), TEG-MOP (triethylene glycol MOP), and MOP-18 (dodecane MOP) (Figure 39) were used to prepare MOP/PTMSP MMMs at 20 wt % MOP loadings. The pristine polymer as well as the MMMs were tested immediately and after 365 days for CO₂ and N₂ permeation.

As expected, the permeability of PTMSP decreased by as much as 73% whereas the permeability of the MMMs decreased between 20% and 50% (Figure 40a). The best results were obtained with t-Bu-MOP which was tested at different loadings (Figure 40a) and showed that MMMs experienced reduced aging at different rates that depended on the MOP loading. From this study it was concluded that a 20 wt % t-Bu-MOP loading was the optimal loading which reduced aging by 20% over a period of 365 days. The authors concluded that the pore windows of t-Bu-MOP were large enough to intercalate the trimethylsilil side groups of PTMSP (Figure 40b) and to form an interlocked configuration that reduced chain mobility. More reports on the effects of additives in membrane aging can be found in [208,245–249].

More work on MOP-containing MMMs was reported in 2015 when J. Ma, et al. [250] prepared functionalized MOPs (e.g., MOP-SO₃Na and MOP-OH) for MOP/PSf MMM fabrication. Owing to the solubility of MOPs in organic solvents that facilitates the preparation of MMMs [126], J. Ma, et al. decorated the MOP pores (Figure 41a) with -SO₃Na and -OH groups to improve the CO_2/CH_4 selectivity of the membrane by increasing the solubility for CO_2 . For the preparation of the MMMs, the functionalized MOPs were dissolved in NMP and then mixed with PSf solutions in NMP. MMMs with 8, 12, and 18 wt % MOP loadings were then dried in a vacuum oven at 50 °C for 2 days before gas permeation experiments were performed with a $CO_2:CH_4$ mixture of 1:1 at 3 bar. With increasing MOP loading, MOP-SO₃Na/PSf MMMs exhibited gas permeability increments up to 113% (CO_2) and 76% (CH_4) that were attributed to the porosity of the MOP (pore size = 1.6 nm), which provided new diffusion paths in the MMM, to the larger pore apertures (0.45 and 0.62 nm), and to the presence of the -SO₃Na polar groups that interacted more strongly with CO_2 than with

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 CH_4 . The CO_2/CH_4 separation was also increased by as much as 60%, from 28 (PSf) to 45 (MMM with 12 wt % MOP loading) due to the preferential sorption of CO_2 in the membrane. To further examine the effect of the polar - SO_3 Na group in the MMM, MOP-OH/PSf MMMs were also tested for CO_2/CH_4 separations. The results from this control experiment (Figure 41b) indicated that MOP-OH did not increase the CO_2/CH_4 selectivity of the MMMs, but it did increase gas permeability in a similar fashion.

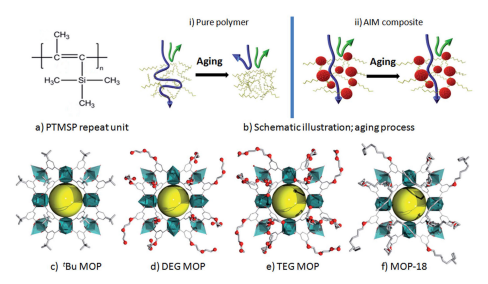


Figure 39. (a) PTMSP repeat unit; (b) schematic illustrations of (i) pure polymer exhibiting packing of polymer chains after aging; and (ii) MOP–MMM that prevents chain relaxation, maintaining membrane permeability; (c) *t*-Bu-MOP; (d) diethylene glycol-MOP (DEG-MOP); (e) triethylene glycol-MOP (TEG-MOP); and (f) MOP-18. Reproduced with permission from [244]. Copyright, 2015, The Royal Society of Chemistry.

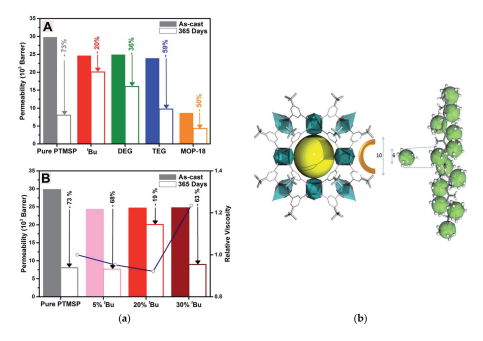


Figure 40. (a) CO₂ permeability properties of 20 wt % MOP-loaded PTMSP membranes as cast vs. 365 days (**A**) and CO₂ permeability properties of *t*-Bu–PTMSP membranes of different dopant loadings as cast vs. 365 days (**B**); (**b**) The largest *t*-Bu-MOP pore windows are large enough (13.671 °A) to allow intercalation of PTMSP trimethylsilyl groups into the MOP pore. Reproduced with permission from [244]. Copyright, 2015, The Royal Society of Chemistry.

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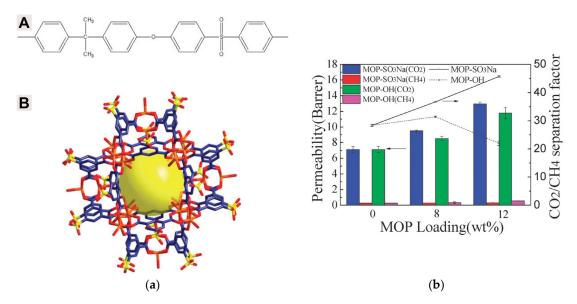


Figure 41. (a) Structures of PSf (**A**) and a porous metal-organic polyhedron (**B**); (b) CO_2/CH_4 (1:1) gas mixture separation performance of functionalized MOP/PSf MMMs at 25 °C and 3 bar. Adapted with permission from [250]. Copyright, 2015, The Royal Society of Chemistry.

This same year, T. Rodenas, et al. prepared Cu-BDC MOF (Figure 42) and high aspect ratio (>20) Cu-BDC MOF nanosheets (ns) from 1,4-benzenedicarboxylic acid and Cu(NO₃)₂ in DMF [251] to study the effects of filler morphology on MMM performance. From AFM images, the synthesized 2D MOF crystals were measured to be 0.5–4 µm in the x–y dimension and 5–25 nm in the z-dimension. The 2D MOF crystals were added to a PSf solution to prepare 2–12 wt % 2D ns-Cu-BDC/PSf MMMs, which were then dried under vacuum at 180 °C. An identical procedure was followed to prepare control materials (e.g., PSf and isotropic, bulk type b-Cu-BDC/PSf or sub micrometer-sized nc-Cu-BDC/PSf MMMs).

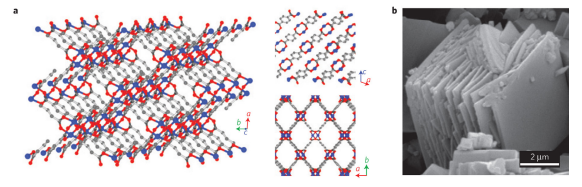


Figure 42. 3D crystalline structure of Cu-BDC MOF (a) and SEM image of bulk type Cu-BDC crystals showing a plate-like morphology (b). Adapted with permission from [251]. Copyright, 2015, Nature Publishing Group.

FIB-SEM images and tomograms (Figure 43a–d) revealed that the 8 wt % *ns*-Cu-BDC/PSf MMMs exposed one order of magnitude larger surface areas than the 8 wt % *b*-Cu-BDC/PSf MMMs, largely increasing its interaction with gas molecules. The surface-rendered views after segmentation of the different phases showed a significant difference of the nanostructure between the bulk and the nanosheet Cu-MOF/PSf MMMs. The bulk crystals left a significant portion in the MMM unoccupied (Figure 43e), whereas the nanosheet Cu-MOF was evenly distributed in the MMM (Figure 43f) and was oriented perpendicularly to the gas flux.

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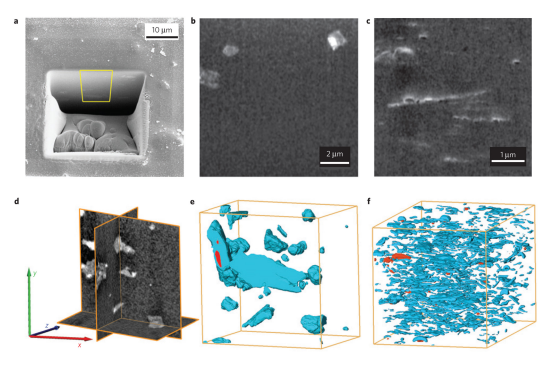


Figure 43. (a) SEM image of the trench carved in a 8 wt % MOF/PSf MMM with FIB milling; (b) SEM image of the cross-section of a 8 wt % *b*-Cu-BDC/PSf MMM containing bulk Cu-MOF; (c) SEM image of the cross-section of a 8 wt % *ns*-Cu-BDC/PSf MMM containing 2D nanosheets of Cu-MOF; (d) orthogonal SEM images of the cross-sections through the 3D FIB-SEM tomogram of a Cu-MOF/PSf MMM; (e,f) surface-rendered views of the segmented FIB-SEM tomograms for *b*-Cu-BDC/PSf and *ns*-Cu-BDC/PSf MMMs, respectively. The x:y:z directions are 11.2:11.2:7.6 for (e) and 4.9:4.9:6.6 μm for (f) showing the MOF in blue and the voids in red. Reproduced with permission from [251]. Copyright, 2015, Nature Publishing Group.

Gas mixture separations with CO₂:CH₄ (1:1) blends (Figure 44) showed that the bulk-type Cu-BDC crystals worsened the separation properties of the MMM (b-Cu-BDC/PSf) with respect to the polymer, most likely due to the generation of non-selective voids at the MOF/polymer boundary. The use of sub-micrometer-sized crystals resulted in a slight improvement in the performance of the MMM (nc-Cu-BDC/PSf), but was still inferior to that of the polymer. The best performance of the MMMs was achieved with the use of nanosheets (8 wt % ns-Cu-BDC/PSf) that significantly improved the CO₂/CH₄ selectivity of the PSf membrane in the range of pressures tested (3 to 7.5 bar). Selectivity increments of 30% to 80% (relative to PSf) and 75% to 8 times higher (relative to the b-Cu-BDC/PSf MMM) were observed for the ns-Cu-BDC/PSf MMM, which also showed a trend of increasing selectivity with increasing pressure, opposite to the polymer's selectivity trend that decays with increasing pressure due to CO₂-induced plasticization of the PSf membrane. In terms of permeability, the 8 wt % ns-Cu-BDC/PSf MMMs exhibited a continuous decrease in CO₂ permeability from 4.1 (3 bar) to 2.8 (7.5 bar) Barrer with increasing pressure, whereas the polymer retained the CO₂ permeability at 5.8 Barrer in the range of pressures tested. Similar studies on filler morphology effects on MMM performance were also performed by Z. Kang, et al. [252] this same year and by A. Sabetghadam, et al. in 2016 [253]. Z. Kang, et al. observed that MMMs of nanosheets of the [Cu₂(NDC)₂(DABCO)]_n MOF in PBI had better H₂/CO₂ gas separation performance than the MMMs of nanocubes of the MOF in PBI. A. Sabetghadam, et al. also observed that MMMs of nanoparticles of NH₂-MIL-53 in Matrimid[®] exhibited better CO₂/CH₄ separation performance than the MMMs of the nanorods or microneedles of the MOF in Matrimid[®]. Undoubtedly, the results of these independent works indicate that the morphology of the filler plays an important role in MMM performance and it should be considered and explored in more detail in future works.

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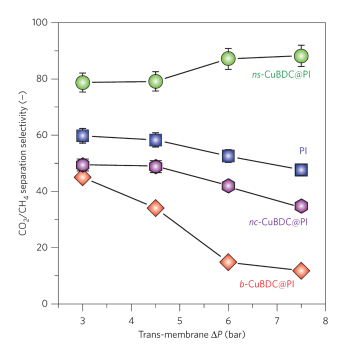


Figure 44. CO₂:CH₄ (1:1) gas mixture separation performance at 25 °C of PSf and the *b*, *nc*, *ns*-Cu-MOF/PSf MMMs as a function of pressure. Reproduced with permission from [251]. Copyright, 2015, Nature Publishing Group.

Work to improve the MOF/polymer interface was continued in 2015 by S. R. Venna, et al. when they modified the surface of (Zr) UiO-66-NH₂ MOF particles to improve its interaction with the polymer matrix, Matrimid[®] [254]. The MOF is composed of Zr₆O₄(OH)₄ clusters bridged with 2-amino-1,4-benzenedicarboxylates that form triangularly shaped pore apertures of 0.6 nm. (Zr) UiO-66-NH₂ was used due to its water, chemical, and thermal stability, and its ability to undergo post-synthesis chemical modification without degradation. The authors rationalized that the MOF/polymer interface could be improved by analyzing the structure of the polymer to determine the functional groups that could form an ideal interface with the MOF and then chemically modify the surface of the MOF with such functional groups. The authors prepared four MOFs (Figure 45) having polar, non-polar, or aromatic surface functional groups: neat (Zr) UiO-66-NH2 (I), aromatic-modified (Zr) UiO-66-NH₂ (I_{PA}), aliphatic C₁₀-modified (Zr) UiO-66-NH₂ (I_{C10}), and acid-modified (Zr) UiO-66-NH₂ (I_{SA}). The synthesized MOFs were activated with Soxhlet extraction for 7 day with Cl₂CH₂. The degree of functionalization of the MOFs was calculated from LCMS, XPS, and ¹H NMR data: the percentages of functionalized ligands obtained were 2%-6% for I_{PA} , 2%-5% for I_{C10} , and 16%-32%for I_{SA}. Owing to the size of the phenyl and alkyl chains that limit their diffusion into the channels of the MOF, the authors expected the surface functionalization for I_{PA} and I_{C10} to be low. The higher level of I_{SA} functionalities, however, indicated that some internal functionalization occurred along with surface functionalization. MMMs of Matrimid[®] and (Zr) UiO-66-NH₂ (and its functionalized derivatives) with loadings of 12 to 40 wt % were prepared with an initial priming of the MOF with a polymer solution. The MMMs were cast on glass, dried at 100 °C, and then annealed at 225 °C for 2 h in a vacuum oven. Pure CO₂ and N₂ gas permeation studies were performed at room temperature and 1.38 bar. The permeabilities and CO₂/N₂ selectivities of the polymer and the 23 wt % (Zr) UiO-66-NH₂ (I, I_{PA}, I_{C10}, I_{SA})/Matrimid[®] MMMs are shown in Figure 46a. The results indicated that ${
m CO_2}$ permeability increased largely for all MMMs with respect to the polymer, but only the MMMs containing the neat and the I_{PA}-functionalized MOF exhibited increased CO₂/N₂ selectivity relative to the polymer, which was attributed to the formation of a defect-free interface between the MOF and the polymer and the improved surface diffusion of CO₂, since it adsorbed more favorably to the MOF than did N_2 .

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Figure 45. Post-synthesis modification of (Zr) UiO-66-NH₂ (I) for functionalization at the $-NH_2$ functional group of the linker with aromatic (I_{PA}), aliphatic C_{10} (I_{C10}), and acid (I_{SA}) terminal functional groups. Reproduced with permission from [254]. Copyright, 2015, The Royal Society of Chemistry.

The authors also concluded that MMMs containing the I_{C10} and I_{SA} functionalities did not increase CO_2/N_2 selectivities because of the defects created at the MOF/polymer interface due to incompatibilities between the polymer and the I_{C10} and I_{SA} functional groups on the MOF. Additional testing with 12, 23, and 40 wt % (Zr) UiO-66-NH₂ (I, I_{PA})/Matrimid® MMMs revealed that the membranes increased the permeability for CO_2 with increasing MOF loadings (Figure 46b). CO_2/N_2 selectivities, however, reached a maximum at 23 wt % and then declined at 40 wt % signaling the presence of interfacial defects induced by the agglomeration of the MOF crystals at high loadings.

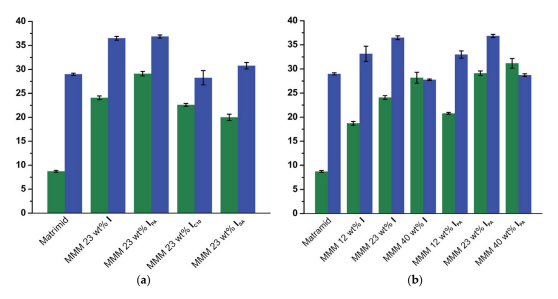


Figure 46. (a) CO_2 permeabilities (green) and CO_2/N_2 ideal selectivities for the polymer and the 23 wt % (Zr) UiO-66-NH₂ (I, I_{PA}, I_{C10}, I_{SA})/Matrimid[®] MMMs; (b) CO_2 permeabilities (green) and CO_2/N_2 ideal selectivities for the polymer and the 12, 23, and 40 wt % (Zr) UiO-66-NH₂ (I, I_{PA})/Matrimid[®] MMMs. Reproduced with permission from [254]. Copyright, 2015, The Royal Society of Chemistry.

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A novel approach for the preparation of polymers that inherently incorporate controlled porosity came from the work of Z. Zhang, et al. in 2015 with the synthesis of polyMOFs [255]. Since the first examples of the preparation of MOF-based MMMs, the aim was to disperse perfectly the crystallites in the polymer matrix without aggregation and voids at the interface. Several strategies have been used to attempt to achieve this goal, e.g., priming the crystals with dilute polymer solutions [165], extensive sonication, chemical functionalization of the organic ligands of the MOFs [250,254], and the use of porous single molecular cages in the form of metal organic polyhedras or MOPs [126]. Each of them achieved a certain degree of success that ultimately relied on the physical interaction between the distinct phases (MOF-polymer). In the work of Zhang, et al., polymer chains that include the carboxylated organic ligands of MOFs in the backbone are reacted with metal ions (Figure 47a and 47b-top), generating MOF unit cells in situ that are chemically bound to the polymer. The hybrid material effectively incorporates porosity from the in situ synthesized MOF and is termed a polymer-MOF hybrid or a polyMOF. In this approach, the polymer chains are effectively the organic linkers or polymer ligands of the MOF. The pore size of Zn-pbdc-7a (a polyMOF with 7 methylene groups between terephthalates, Figure 47b-top) and Zn-pbdc-8a (a polyMOF with 8 methylene groups between terephthalates), calculated by DFT methods from N2 sorption isotherms, were 0.7 nm and 0.9 nm, respectively. The authors were also able to synthesize Zn-pbdc-7a and Zn-pbdc-8a into 20 µm thick crystalline films that showed an intergrowth network of crystallites with morphologies distinctive of the polymer ligands, as was observed by SEM (Figure 47b-bottom). Although the preparation of this new type of material for gas separation is promising and provides an exciting new path in the quest for a perfect membrane, the mechanical properties and the performance of these membranes in gas separations are still to be determined.

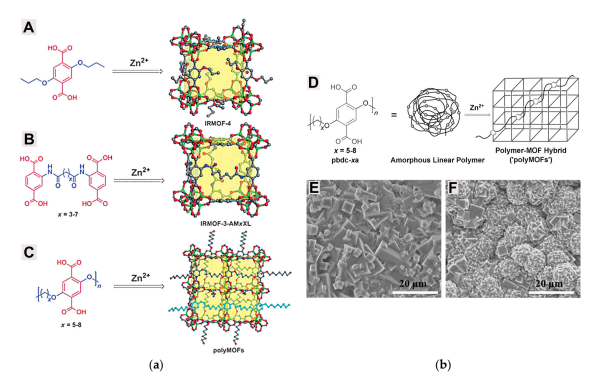


Figure 47. (a) Isoreticular metal-organic framework (IRMOF) derivatives showing the evolution of the organic ligands from simple (A) to crosslinked (B) to polymeric ligands (C); (b) Strategy to convert the linear polymer into a porous crystalline polyMOF (D) and SEM images (E,F) of the film morphology of Zn-pbdc-7a (E) and Zn-pbdc-8a (F). Adapted with permission from [255]. Copyright, 2015, Wiley.

Another contribution made in 2015 towards the industrial utilization of MMMs came from the work of E. V. Perez, et al. with the testing of 20 wt % (Al) NH₂-MIL-53/VTECTM MMMs at pressures

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that ranged from 5 to 30 bar and at temperatures that ranged from 35 to 300 °C, achieving H₂/CO₂ mixture selectivities of 7.5 [128]. Their work provided evidence that MMMs could be sufficiently mechanically robust to perform gas separations under combined extreme conditions of pressure and temperature, increasing permeability and selectivity (relative to the polymer) for H₂:CO₂ (1:1) mixtures and also retaining selectivity for measurements using pure gases of H₂ and CO₂. Gas permeation of the polymer and MMMs using pure gases showed that the membranes retained a H_2/CO_2 selectivity between 5.3 and 8.3 at up to 250 °C, whereas at 300 °C the selectivity dropped to 4.0 for the polymer and to 5.8 for the MMM. H₂ permeability increased from 5 (35 °C) to 84 (300 °C) Barrer for VTECTM and remained constant within the pressure range tested (5 to 30 bar) at each temperature. The MMMs exhibited increased H₂ permeability from 5 (35 °C) to 150 Barrer (300 °C), which remained constant within the pressure range (5–30 bar) at each testing temperature. Testing the MMMs with H_2 and CO₂ at these conditions (30 bar and 300 °C) is already remarkable progress for MOF-based MMMs because it proves that the MOF/polymer interface is strong enough to perform molecular sieving and to prevent the formation of defects. The fact that the MMMs retained H₂/CO₂ selectivities of 5.8 (Knudsen selectivity $H_2/CO_2 = 4.7$) at temperatures above the T_g of the polymer indicates that the MOF contributes to or enhances the mechanical strength of the polymer. Gas mixture experiments at 250 °C and 30 bar showed that the H_2/CO_2 selectivity depended largely on the membrane stage cut (θ). At a θ of 1.0 (no retentate removal or 100% H₂ recovery), the CO₂ polarization on the surface of the membrane changed the feed composition dramatically, which reduced the H₂/CO₂ selectivity of the polymer from 8.1 (ideal) to 2.9 (mixture) and the MMM's from 8.3 (ideal) to 4.4 (mixture). Reducing θ to <0.2 (<20% H₂ recovery) resulted in an improvement in the H₂/CO₂ separation of the membranes for gas mixtures. The H_2/CO_2 selectivities for the polymer increased from 2.9 ($\theta = 1.0$) to 7.2 ($\theta = 0.1$) and for the MMM from 4.4 ($\theta = 1.0$) to 7.5 ($\theta = 0.1$). The benefits of incorporating the MOF into the polymer were evident since the MMM outperformed the polymer in both its permeability and selectivity properties with pure gases and gas mixtures.

New developments in the preparation of MMMs came in 2016 from the work of X. Cao, et al. with the preparation of COF-LZU1/PVAm MMMs on porous PSf ultrafiltration membranes for the separation of CO₂/H₂ [256]. Although COFs are not hybrid organic-inorganic frameworks, but rather pure covalently-bound organic frameworks that may contain boron, triazines, or imines [257], they share similar properties with MOFs. Both materials form 2D or 3D crystalline frameworks with defined porosities and gas sorption capacities that depend on the nature of the organic component. COF-LZU1, an imine-based and water stable 2D COF, was made from the reaction of 1,3,5-triformylbenzene with 1,4-diaminobenzene and mixed with PVAm in an aqueous solution. The polymer and the 10 wt % COF-LZU1/PVAm MMM were tested for gas separation with H₂:CO₂ (60:40 volume) mixtures at 25 °C and at pressures from 1.5 to 10 bar. The gas separation performance of the MMM was superior to that of PVAm in both flux and CO₂/H₂ selectivity. CO₂ flux increased from 100 GPU (PVAm, 1.5 bar) to 396 GPU (MMM, 1.5 bar) and declined to 286 GPU (MMM) as the pressure increased to 10 bar. The CO₂ flux remained unchanged for PVAm over the range of pressures studied. The CO₂/H₂ selectivity for the MMM decreased from 15 to 12.5 with increasing pressure, but it remained higher than for PVAm ($CO_2/H_2 = 11$). The authors attribute the improved performance of the MMM to the porosity introduced by the COF and the amino functionality in the pores that preferentially adsorbs CO₂. Another COF-based MMM study released this year came from Z. Kang, et al. with the preparation of MMMs containing exfoliated 2D COFs (NUS2 and NUS3, Figure 48a) [258]. NUS2 and NUS3 are layered, porous materials made from 1,3,5-triformylphloroglucinol (NUS2) or 1,3,5-triformylphloroglucinol and 5-diethoxy-terephthalo-hydrazide (NUS3), as depicted in Figure 48a. The layered COFs were exfoliated into nanosheets or even monolayers with high aspect ratios by suspending them with stirring and sonication in CHCl₃ or NMP before their incorporation into UltemTM or PBI. The 10, 20, and 30 wt % COF MMMs were then dried at 200 °C in a vacuum oven before permeability tests with H₂, CH₄, CO₂, and a (1:1) H₂:CO₂ mixture were performed at 35 °C and at pressures from 2 to 5 bar. The authors attempted to determine the degree of dispersion and orientation

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of the COF nanosheets in the MMMs, but due to the similarities in chemical composition between the COFs and the polymers, they could not obtain conclusive evidence. SEM images of the membrane cross-sections, however, revealed the presence of polymer veins due to plastic deformation from the strong COF/polymer interaction for both NUS2 and NUS3-based MMMs (Figure 48b). Pure gas permeation indicated that the MMMs exhibited their best separation and permeability performance at 20 wt % loading. Higher loadings negatively affected the performance of the MMMs for both CO_2/CH_4 and H_2/CO_2 separations (Figure 49). At 20 wt % NUS2 and NUS3/UltemTM MMMs, the H_2 and CO_2 permeabilities increased significantly with respect to the polymer, increasing the CO_2/CH_4 and H_2/CO_2 selectivities (Figure 49a–d). The increased permeability was attributed to the porosity introduced by the COF in the MMM, which is clearly observed in Figure 49a–d, where MMMs containing NUS3 (1.8 nm pore aperture) exhibit superior CO_2 permeabilities than MMMs containing NUS2 (0.8 nm pore aperture) and UltemTM. The 20 wt % NUS3-PBI MMMs exhibited a H_2 diffusivity 17-fold that of PBI (Figure 49e,f).

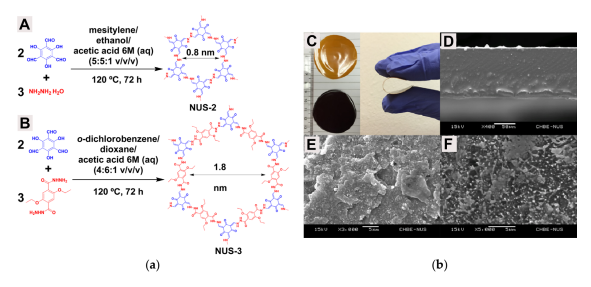


Figure 48. (a) Synthesis procedures and structures of the 2D covalen organic frameworks (COFs), NUS2 (A) and NUS3 (B); (b) Optical images (C) of a 20 wt % NUS2/UltemTM (brown) and a 20 wt % NUS3/UltemTM (black) MMMs, and SEM images of the cross-sections of UltemTM (D); 20 wt % NUS2/UltemTM (E); and a 20 wt % NUS3/UltemTM (F). Adapted with permission from [258]. Copyright, 2016, American Chemical Society.

Gas mixture separation properties of the MMMs with H_2/CO_2 blends showed that H_2 permeability decreased by 20% to 30% for all MMMs relative to the pure gases. The H_2/CO_2 selectivity of the NUS2-based MMMs decreased from 5.8 to 5.0 for the UltemTM MMMs and from 31.4 to 18.8 for the PBI MMMs (Figure 49).

As the amount of work related to MOF-based MMMs increased in recent years, the measurement of the mechanical properties of MOF-based MMMs became more important for the understanding of the MOF/polymer interactions. In 2016, E. M. Mahdi, et al. [209] and S. J. D. Smith, et al. [208] reported on the thermomechanical, stress-strain, and viscoelastic properties of MMMs and the effects of aging on mechanical properties. E. M. Mahdi, et al. performed a detailed study of these properties on ZIF-8/polyurethane MMMs and concluded that for this system the interactions between the ZIF and the polymer were tenue since the hyperelastic and viscoelastic characteristics of polyurethane were retained upon ZIF-8 incorporation. These interactions were mostly a collection of uniform and weak H-bonds, van der Waals forces, and π - π stacking interactions that produced a "mobile" free volume in the MMM due to the mobility of the ZIF-8 particles in the MMM when the ZIF loading was less than 20 wt %. At higher loadings of ZIF-8, however, the increased amount of the weak interactions formed more rigid composite that suppressed the mobility of the ZIF/polymer interface

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causing a decline in ductility and toughness. S. J. D. Smith, et al. [208] extended the mechanical analyses of MMMs by studying the effects of aging on the mechanical properties of MMMs as seen in Figure 50. In this study, PIM-1, Matrimid[®], and PTMSP were used as polymer matrices to prepare MMMs with fumed silica, PAF-1, UiO-66, and Ti₅UiO-66 additives at 10 wt % loadings. As is shown in Figure 51, the incorporation of the nano-additives reduced the stress and strain at failure of the as-cast MMMs. MMMs containing PAF-1 and fumed silica showed identical losses in mechanical performance regardless of the polymer matrix used. It was concluded that polymer-additive interactions strongly influence local physical aging and determine the properties of the MMMs, additionally, they can slow physical aging too. By comparing the properties of the additives in different polymers, it was observed that the polymer largely determined the mechanical properties of the MMM since Matrimid[®] MMMs were stronger than PIM-1 and PTMSP MMMs containing the same additives. The mechanical properties of the pure polymer membranes decreased with aging, specifically, the membranes became more brittle as they aged. As the membranes aged, Young's modulus increased with time but differed with the type of additive used, for example, Ti₅UiO-66/PIM-1 MMMs exhibited higher modulus than PAF-1/PIM-1 MMMs and PIM-1 membranes but PAF-1/PIM-1 MMMs showed less modulus values than PIM-1 membranes.

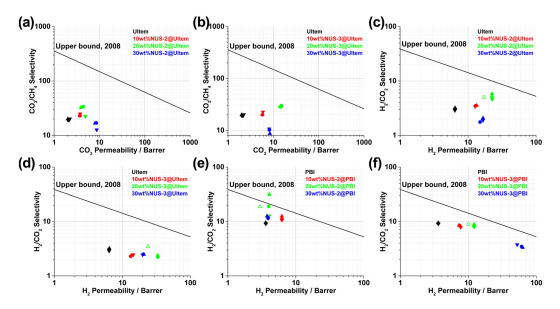


Figure 49. Pure gas (solid symbols) and gas mixture (open symbols) selectivities of NUS2 and NUS3/UltemTM and PBI MMMs: CO_2/CH_4 separation with NUS2/UltemTM (**a**) and with NUS3/UltemTM (**b**); H_2/CO_2 separation with NUS2/UltemTM (**c**) and with NUS3/UltemTM (**d**); H_2/CO_2 separation with NUS2/PBI (**e**) and NUS3/PBI (**f**). ∇ = 2 bar, \triangle = 5 bar. Reprinted with permission from [258]. Copyright, 2016, American Chemical Society.

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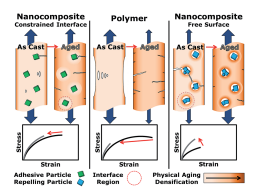


Figure 50. Fracture mechanisms of aged MMMs with different additive interface environments compared to freshly cast polymer embrittlement. Adhesive interaction reduces crack propagation, leading to better performance over the embrittled aged polymer. Repulsive interaction initially compromise film properties; however the localized increased aging rate isolates defects and leads to mechanical stability. Reproduced with permission from [208]. Copyright, 2016, The Royal Society of Chemistry.

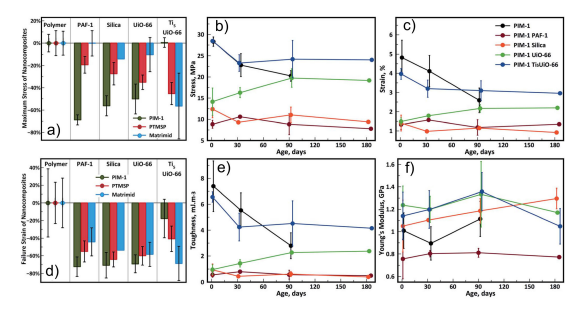


Figure 51. Effect of nanoparticle addition on maximum tensile stress (**a**) and strain at failure (**d**) of as-cast polymer composite films. Comparison of stress (**b**); strain (**c**); toughness (**e**); and modulus (**f**) of PIM-1 based nanocomposites with age. Pure PIM-1 films became too fragile for testing after 90 day. Adapted with permission from [208]. Copyright, 2016, The Royal Society of Chemistry.

The previously described advances made in the MOF-based MMM field are shown in Figure 52. The chronological evolution of this type of MMM is certainly promising and exciting and shows the attention that this type of material has received from the scientific community. Additionally, the reader is directed to several reviews of MOF-based MMMs [134,206,207,259–265] that present the most significant developments and challenges in this field.

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MOF-based MMMs Progress	Year	Contributors
Cu-MOF/PAET MMMs for gas separations	2004	H. Yehia, et al. [120]
Cu ₃ BTC ₂ /PDMS and Mn(HCOO) ₂ /PSf MOF-MMMs	2006	A. Car, et al. [122]
MOF/polyimide MMM for gas separations: MOF-5/Matrimid® MMMs	2006	E. V. Perez, et al. [123]
Cu-MOF/PDMS/POT/PAET MMMs for gas separations	2006	C. Barcena [203]
Cu-BPY-HFS/Matrimid® MMMs for gas separations and mechanical	2008	Y. Zhang, et al. [204]
analysis of Cu-BPY-HFS/Matrimid® MMMs		
MOF-5/Cu-MOF/MOP-18/Matrimid® MMMs for gas separations	2009	E. V. Perez, et al. [124,125]
Cu-TPA/PVAc MMMs for gas separation	2010	R. Adams, et al. [210]
	2010	
ZIF-90/Ultem [™] /Matrimid®/6FDA-DAM MMMs for gas separations		TH. Bae, et al. [127]
CO ₂ diffusion in ZIF-8/PSf MMMs by pulsed field gradient NMR	2010	K. Diaz, et al. [211]
Cu-based MOF hollow fiber MMM	2010	J. Hu, et al. [212]
Atomistic and continuum modeling of MOF-based MMMs	2010	S. Keskin, et al. [213,214]
Tortuosity in ZIF-8/Matrimid® MMMs for gas separations	2010	Ma. J. Ordoñez, et al. [216
Asymmetric MIL-53/Matrimid® MMMs for gas separations	2011	S. Basu, et al. [218]
ZIF-8/PMPS MMMs for C ₂ -C ₅ alcohols recovery from water by	2011	X. L. Liu, et al. [38]
organophilic pervaporation		
Colloidal ZIF-7/PBI MMMs for H ₂ /CO ₂ separations: improved	2011	T. Yang, et al. [205]
dispersion of ZIF in polymer		
NH2-functionalized MIL-53/PSf MMMs for gas separations	2011	B. Zornoza, et al. [219]
Combination of MOF and zeolites in MMMs: synergistic effects	2011	B. Zornoza, et al. [221]
Water stable MIL-101/PSf MMMs for O ₂ /N ₂ separations	2012	H. B. T. Jeazet, et al. [222]
ZIF-8/Ultem™ hollow fiber MMMs for CO ₂ /N ₂ separations	2012	Y. Dai, et al. [223]
H ₂ /CO ₂ separation with ZIF-8/PBI hollow fiber MMMs at elevated	2012	T. Yang, et al. [224]
temperatures		
C3H6/C3H8 separation with ZIF-8/6FDA-DAM MMMs	2012	C. Zhang, et al. [11]
ZIF-8/PIM-1 MMMs for gas separations and PALS study	2013	A. F. Bushell, et al. [225]
PALS study of MIL-101/PSf MMMs for the correlation of permeability	2013	H. B. T. Jeazet, et al. [229]
with free volume	2013	11. D. 1. Jeazer, et al. [229]
Interfacial MOF/polymer adsorption equilibrium effects on gas	2013	T. Singh, et al. [230]
	2013	1. 311gH, et al. [230]
permeation models: Finite element analysis	2012	I II-o et al [222]
Room temperature ionic liquid/ZIF-8 MMMs for gas separations	2013	L. Hao, et al. [233]
Reverse CO ₂ /H ₂ selective MIL-53/PMP MMMs	2014	R. Abedini, et al. [234]
Comparative gas permeation study of ZIF-8, polymer (PMPS), and the	2014	L. Diestel, et al. [235]
ZIF-8/PMPS membranes		
Drying temperature effects on MMM gas permeabilities	2014	C. Duan, et al. [237]
Impact of ZIF particle size on MMM CO ₂ /CH ₄ separation	2014	N. A. Nordin, et al. [238]
MOP-18/Matrimid® MMMs for gas separations: molecular level	2014	E. V. Perez, et al. [126]
dispersion, high pressure gas permeation, and mechanical analysis		
High pressure gas permeation with MOF-based MMMs	2014	S. Shahid, et al. [239,240]
FIB-SEM and 3D visualization of MOF dispersion in MMM	2014	T. Rodenas, et al. [241]
ZIF-8 membrane growth on ZIF-8-based MMM supports	2014	Z. Y. Yeo, et al. [242]
Aging cessation in PAF-1/PTMSP, PMP, PIM-1 MMMs	2014	C. H. Lau, et al. [243]
Aging reduction in MOP/PTMSP MMMs	2015	M. Kitchin, et al. [244]
Functionalized MOPs MMMs for gas separations	2015	J. Ma, et al. [250]
Two-dimensional Cu-BTC MOF nanosheet MMMs: effect of filler	2015	T. Rodenas, et al. [251]
morphology on gas permeation		,
MOF surface functionalization for improved compatibility	2015	S. R. Venna, et al. [254]
H ₂ /CO ₂ separations with NH ₂ -MIL-53/VTEC [™] MMMs at high pressure	2015	E. V. Perez, et al. [128]
and high temperature (30 bar and 300 °C) and thermomechanical	2010	2. v. r erez, et al. [120]
analyses of MMMs up to 300 °C	2015	7 7hong of -1 10551
Polymer-MOF hybrid materials: polyMOFs	2015	Z. Zhang, et al. [255]
COF/poly(vinyl amine) MMMs for gas separations	2016	X. Cao, et al. [256]
Exfoliated 2D COFs in Ultem™ and PBI for H ₂ separations	2016	Z. Kang, et al. [258]
Thermomechanical, strain-stress, and viscoelastic properties of ZIF-	2016	E. M. Mahdi, et al. [209]
8/polyurethane MMMs		
Effects of aging on mechanical properties of MMMs	2016	S. J. D. Smith, et al. [208]

Figure 52. Chronological progress for MOF-based MMMs.

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5. Future Directions

By focusing on reported MMMs based on inorganic fillers, the following future directions of the field can be envisioned. Although gas separation membranes are usually commercialized as hollow fiber membranes, reports of inorganic materials based hollow fiber membranes are just a handful. Extensive research on improving mechanical properties, chemical inertness of such MMMs will be the focus of the researchers in the field. Moreover, the combination of zeolites and carbon based materials in the preparation of MMMs with improved gas separation capabilities could be another research topic in the future.

The challenges for MOF-based MMMs to overcome are more evident and specific as the evolution of this type of membranes progresses. In contrast to inorganic-based MMMs where additive/polymer incompatibilities are a major concern, in MOF-based MMMs polymer rigidification around the MOF particles, presumably due to strong interactions, is a concern since it isolates the MOF and reduces its interactions with gas molecules. The ability to control and tune the MOF/polymer interface is a necessity for this type of membranes. The effects of filler size, morphology, and particle aspect ratio on membrane performance is also an emerging topic that could lead to improvements in membrane performances once their effects are understood. Another topic that deserves attention is suppression of membrane aging for industrial applications, extending the lifetime of the separation properties of membranes over a reasonable period of time could make polymer-based membranes attractive in industrial gas separations. Initial work on this topic has started but more development is needed.

6. Conclusions

From the early observations of the water and CO_2 permeation phenomena in membranes made of pig bladders in the mid-1700s, to the first report of an MMM using dense inorganic additives in 1912, to the incorporation of porous molecular sieves in 1971, and to the incorporation of porous MOFs in 2004, the field of MMMs has experienced an enormous progress.

To improve gas separation characteristics, researchers have fabricated polymer-based MMMs containing inorganic additives, including zeolites and carbon-based materials. Carbon molecular sieves, carbon nanotubes, and graphene were studied as carbon-based additives for MMMs. However, owing to poor interactions between the polymers and the additives, these membranes showed poor mechanical properties. Therefore, researchers have tried to improve the interactions by using different strategies. Some have explored the interactions of several different polymers with a certain zeolite. Modification of the chemical structures of the additives as well as the polymers has been employed in order to improve the interactions. Both zeolites and carbon-based additives have been chemically modified in fabricating MMMs, which showed improvements not only in mechanical properties but also in gas separation properties.

The MOF-based MMM technology has evolved rapidly owing to the potential these materials bring to the MMM field. Significant work was oriented to the understanding and improvement of the MOF/polymer interface. Of paramount importance is the effect this interface has on gas transport, and this is where there is still work to be done. Polymer rigidification at the interface, pore blockage, non-selective voids, and particle aggregation are some of the main challenges MMMs face before industrial application. In this review, the work of several scholars that have addressed these challenges has been presented. The authors of this work believe there is still more work to be done to fully understand and control these parameters before the full potential of MOFs can be realized in an MMM. The majority of the works presented herein showed an increase in gas permeation and relatively constant selectivity with increasing MOF loadings that was attributed to the porosity of the MOFs in the MMMs. The selectivities, however, seemed to still be controlled by the polymer (with a few exceptions), and little contribution was observed from the MOF, which may be the next area of concerted focus. Another important aspect of the art is the testing of MOF-based MMMs under real world conditions. Gas permeability and selectivity properties generally change when the membranes are exposed to gas mixtures and contaminants in the feed as well as to increasing temperatures that

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could render the most promising membranes unsuitable for industrial applications. Additionally, as higher pressures may be needed in these applications, the mechanical robustness of the membranes needs to be addressed without overlooking required fluxes. As it can be seen, there is still much to be done before MMMs can be employed in industrial gas separations, but the growing interest and dedication of membranologists is catching up.

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Abbreviations

The following abbreviations are used in this manuscript:

[emim][NTf₂] [1-ethyl-3-methyl imidazolium] (cation) [bis(trifluoromethanesulfonyl) imide] (anion)

[emim][BF₄] [1-ethyl-3-methyl imidazolium] (cation) [tetrafluoroborate] (anion) [emim][B(CN)₄] [1-ethyl-3-methyl imidazolium] (cation) [tetracyanoborate] (anion)

[vbim][NTF₂] Polymerizable RTIL (1-vinyl-3-butylimidazolium-bis(trifluoromethylsulfonyl)imidate)

θ Membrane stage cut

Barrer Non-international system unit for gas permeation

(1 Barrer = $1 \times 10^{-10} \text{ cm}^3_{(STP)} \cdot \text{cm} \cdot \text{cm}^{-2} \cdot \text{s}^{-1} \cdot \text{cmHg}^{-1}$)

BSE Backscattered electron
CA Cellulose acetate
CMS Carbon molecular sieve
CNT Carbon nanotube

 $\begin{array}{ccc} {\sf COF} & {\sf Covalent\ organic\ framework\ CP} & {\sf Coordination\ polymer\ DEA} & {\sf Diethanolamine\ } \\ D_f & {\sf Filler\ diffusion\ coefficient\ } \end{array}$

 D_f Filler diffusion coefficient D_m Matrix diffusion coefficient DFT Density functional theory FCNT Functionalized carbon nanotube

FIB-SEM Focused ion beam scanning electron microscopy FMWCNT Functionalized multi-walled carbon nanotube

GO Graphene oxide

GPU Gas permeation unit (1 GPU = $1 \times 10^{-6} \text{ cm}^3_{\text{(STP)}} \cdot \text{cm}^{-2} \cdot \text{s}^{-1} \cdot \text{cmHg}^{-1}$)

IRMOF Isoreticular metal-organic framework

IUPAC International union of pure and applied chemistry *K* Interfacial adsorption equilibrium constant LCMS Liquid chromatography mass spectrometry

MIL Materials Institute Lavoisier MMM Mixed-matrix membrane MOC Metal-organic cube MOF Metal-organic framework MOP Metal-organic polyhedra **MWCNT** Multi-walled carbon nanotube NMP N-methyl-2-pyrrolidone ODA 4,4-Oxydianiline

ODPA 4,4'-Oxydiphthalic anhydride PAET Poly(3-acetoxyethylthiophene) PAF-1 Porous aromatic framework 1

PALS Positron annihilation lifetime spectroscopy

PBI Polybenzimidazole
PDMS Polydimethylsiloxane
PES Polyethersulfone

 P_f Filler permeability coefficient PFG NMR Pulsed field gradient nuclear magnetic resonance

PI Polyimide

PIM-1 Polymer of intrinsic microporosity P_m Matrix permeability coefficient PMDA Pyromellitic dianhydride

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PMP Poly(4-methyl-1-pentyne)
PMP-2 Poly(4-methyl-2-pentyne)
PMPS Polymethylphenylsiloxane

PolyRTILs Polymerizable room temperature ionic liquids PPEES Poly(1,4-phenylene ether-ether-sulfone)

PPZ Polyphosphazene
PS Polystyrene
PSf Polysulfone

PTMSP Poly(1-trimethylsilil-1-propyne)

PVAc Poly(vinyl acetate)
PVAm Poly(vinyl amine)
PVPy Polyvinylpyrrolidone

RTILs Room temperature ionic liquids SAPO-34 Silicoaluminophosphate

SBS Poly(styrene-b-butadiene-b-styrene)

SBU Secondary building unit SEM Scanning electron microscopy SPEEK Sulfonated poly(ether ether ketone) SPES Sulfonated polyethersufone **SWCNT** Single-walled carbon nanotube TEM Transmission electron microscopy T_g Glass transition temperature **TGA** Thermogravimetric analysis

TMPDA 2,4,6-Trimethyl-1,3-phenylenediamine

XRD Powder X-ray diffraction
ZIF Zeolitic imidazolate framework
XPS X-ray photoelectron spectroscopy

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