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Polyethyleneimine NH₂-UiO-66 nanofiller-based mixed matrix membranes for natural gas purification

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16 Abstract

Effective separation of CO₂ from CH₄ is crucial for the purification of natural gas, 17 which requires membrane materials with high permeability, selectivity, and stability of 18 under high pressures. In this work, a high CO2-affinity MOF-based nanofiller 19 (Pin@NH2-UiO-66) was prepared and integrated in polyetherimide (PEI) to prepare 20 mixed matrix membranes (MMMs) for enhanced CO₂/CH₄ separation capabilities. The 21 Pin@NH2-UiO-66 nanofiller comprised an in situ-formed polyethyleneimine-NH2-22 23 UiO-66 composite prepared via a one-pot synthesis. The polyethyleneimine was covalently attached to the H₂BDC-NH₂ ligand allowing for a stronger integration within 24 the PEI matrix. The polyethyleneimine affects the nucleation of MOF, leading to 25

miniaturization of the MOF particles. The smaller-sized filler is beneficial for improved 26 interaction at the filler-matrix interface. The numerous amino functionalities grafted 27 onto the NH2-UiO-66 increased the CO2 adsorption sites, thereby enhancing the affinity 28 for CO₂. Owing to the improved interaction, the elevated CO₂ attraction, and the 29 inherent properties of the porous NH2-UiO-66, the fabricated MMMs preformed 30 superior in separating CO₂/CH₄. The 30-P_{in}@NH₂-UiO-66-PEI membrane (containing 31 30 wt% nanofiller) exhibited a CO₂/CH₄ selectivity of 27.7 and a CO₂ permeability of 32 2498.9 Barrer. The CO₂ permeability was 21 times greater than that of the pristine PEI 33 membrane, and 4 times higher compared to P@NH2-UiO-66 MMM with 34 polyethyleneimine modified MOF filler prepared by a traditional wet impregnation 35 method. Additionally, the novel MMM demonstrates excellent separation stability 36 under conditions that mimic industrial settings, demonstrating its potential application 37 for natural gas purification. 38

Keywords: Mixed-matrix membrane; NH₂-UiO-66; MOF nano-filler modification;
CO₂ separation; in situ formation

41 **1. Introduction**

Recently, the development of technologies for CO₂ separation from natural gas has 42 attracted considerable attention [1, 2]. The membrane separation technique [3-5] is 43 recognized as one of the most feasible strategies to purify natural gas [6, 7], due to the 44 minimal operational expenses, energy-efficient operation, and decreased carbon 45 emissions. Industrial membranes for natural gas purification should operate effectively 46 at temperatures of about 50 °C (the outlet temperature of a natural gas pipeline) and at 47 48 elevated gas pressures, and consistently maintain high CO₂ permeability, selectivity and stability [2, 8]. Polymeric membranes currently predominate the membrane market due 49 50 to their low cost and large-scale manufacturing. The cellulose acetate membrane developed by UOP is the most widely used membrane material in all membrane natural 51 gas processing plants, with a selectivity of 10-15 [9]. However, there are several issues 52 with acid cellulose membranes, including susceptibility to hydrolysis, inadequate 53 permeability and selectivity. Freshly-made thin composite membranes often lose 25% 54

of their permeability within a week. The balance between permeability and selectivity
of polymeric membranes remains challenging, along with maintaining the membrane's
structural stability for effective CO₂ separation [10-12]. To tackle these challenges,
diverse permeable fillers have been integrated into the polymer matrix to create mixed
matrix membrane (MMM) [13-15].

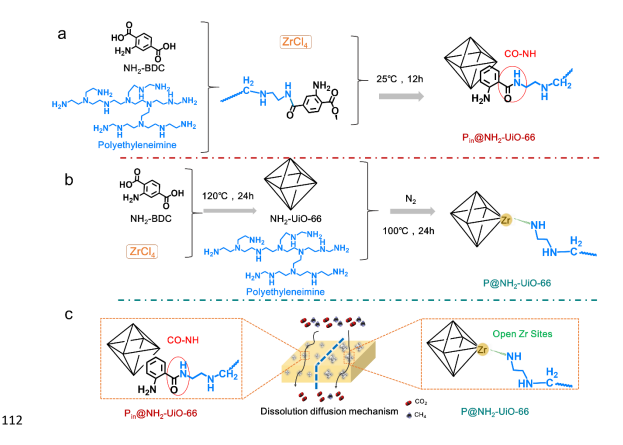
In MMMs, the porous filler is crucial for enhancing CO₂ separation performance by 60 61 creating pathways for low-resistance transport [15-19]. Among these fillers, metal-62 organic frameworks (MOFs) stand out as auspicious options, ascribed to the large surface areas and controllable porosity for CO₂ separation [20, 21]. However, 63 improving interaction at the filler-matrix interface is challenging due to the intrinsic 64 65 differences between organic polymers and MOFs. Moreover, the CO₂-binding property of the MOF fillers exerts a significant role in determining membrane performance. 66 Therefore, it is essential to design MOF fillers that possess superior CO₂ adsorption 67 properties and exhibit strong affinity with the polymer matrix [22]. 68

69 Plenty research have been dedicated to directed towards advancing the performance of new-generation membranes by tuning the chemical affinity towards CO₂ [14, 23-28]. 70 Amines interacting with open metal sites in MOFs effectively boost the CO₂ uptake at 71 low pressures via furnishing extra adsorption sites [20, 29]. Polyethyleneimine stands 72 out as a highly suitable choice for improving the CO₂ adsorption capabilities of MOFs, 73 as it is rich in amino groups, particularly the primary amine sites located at the chain 74 ends [26, 30, 31]. Additionally, the combination of organic polyethyleneimine and MOF 75 materials is beneficial for enhanced interaction at the filler-matrix interface [28, 32]. 76

The wet impregnation approach is a conventional method used to modify MOFs with polyethyleneimine. This approach involves the coupling of polyethyleneimine with the metal sites of the MOF through weak binding interactions. Liu et al. synthesized branched polyethyleneimine-functionalized UiO-66 via the wet impregnation approach [32], which served as filler in 6FDA-ODA to prepare MMMs for CO₂/CH₄ separating. This resulted in an enhanced CO₂/CH₄ selectivity of 56.49. In addition, the same group designed a nanocomposite filler (UiO-66-PEI@bmim[Tf₂N]) using UiO-66 decorated **3** / **27**

with branched polyethyleneimine via wet impregnation method [33], and the MMMs 84 prepared showed an optimum CO₂/CH₄ selectivity of 59.99. Although 85 polyethyleneimine-modified MOF fillers enhance the separation performance of 86 MMMs, the preparation process of modified MOF fillers usually involves high-87 temperature anhydrous and oxygen-free environment, which complicates the 88 membrane preparation process. Furthermore, a higher level of amine functionalization 89 would result in a higher CO₂ adsorption as well as an enhanced CO₂ permeance. 90 91 Notably, in comparison with the wet impregnation technique, the in situ covalent grafting method employs a stronger force, consequently facilitating the filler's ability 92 to bind to a greater number of amino groups. Therefore, seeking a simple strategy to 93 modify MOF fillers by introducing more amino groups without high-temperature 94 processing can significantly reduce operational steps and energy consumption. 95

Herein, we propose a simple in situ one-pot synthesis strategy for the fabrication of 96 polyethyleneimine-modified NH2-UiO-66 (Pin@NH2-UiO-66) (Scheme 1). The 97 98 Pin@NH2-UiO-66 nanofillers were then physically blended with polyetherimide to prepare mixed matrix membranes with an enhanced CO₂ separation performance 99 compared to P@NH2-UiO-66 fillers prepared by conventional wet impregnation. 100 During the in situ synthesis process, polyethyleneimine and H₂BDC-NH₂ ligands are 101 covalently linked to form nano-sized Pin@NH2-UiO-66 fillers with a higher loading of 102 polyethyleneimine. In contrast, the wet impregnation method relies on weaker 103 interactions between NH₂-UiO-66 and polyethyleneimine, resulting in a lower loading 104 capacity of polyethyleneimine. Nano-fillers with high polyethyleneimine load and 105 small size should have better interfacial compatibility with the polymer matrix and 106 107 higher CO₂ adsorption capacity. Benefiting from these features of P_{in}@NH₂-UiO-66, the MMMs exhibited outstanding CO₂/CH₄ separation performance. The 30-P_{in}@NH₂-108 UiO-66-PEI membrane with a 30 wt% of the fillers had a CO₂ permeability of 2498.9 109 Barrer and a CO₂/CH₄ selectivity of 27.7, with 4 times greater permeability compared 110 to MMMs prepared using conventional wet impregnation P@NH₂-UiO-66 fillers. 111



Scheme 1: Schematic illustration for the two preparation methods of mixed-matrix membranes in
 this work: (a) P_{in}@NH₂-UiO-66 in situ one-pot fabrication process, (b) P@NH₂-UiO-66 wet
 impregnation synthesis process, (c) membrane processing.

116 **2. Experimental**

117 *2.1. Materials*

Zirconium (IV) chloride (ZrCl₄, 99.5%) powder, formic acid (HCOOH, 88%) and 118 dichloromethane (CH₂Cl₂, 99%) were supplied by Aladdin Biochemical Technology 119 Co., Ltd. (China, Shanghai). 2-Aminoterephthalic acid (H2BDC-NH2, 98%) was 120 sourced by Shanghai Macklin Biochemical Co., Ltd, N, N-Dimethylformamide (DMF, 121 99.8%), Ethanol (CH₃CH₂OH, AR) and methanol (CH₃OH, 99.7%) were supplied by 122 Sinopharm Chemical Reagent Co., Ltd. Polyethyleneimine (99%, W.M. =10000) was 123 supplied by Arkema, France. CH₄ and CO₂ (99.999%) were acquired from Nanjing 124 Special Gases Company. Polyetherimide (PEI) was obtained from SABIC. All 125 compounds employed as received without any other processing. 126

127 2.2. Synthesis of NH₂-UiO-66, P@NH₂-UiO-66 and P_{in}@NH₂-UiO-66 nano-fillers

128 The synthesis of NH₂-UiO-66 was conducted in accordance with established protocols

from previous studies [34]. A mixture of ZrCl₄ (4.91 mmol, 1.16 g) and 2aminoterephthalic acid (4.91 mmol, 0.89 g) was disbursed in 55 mL of dimethylformamide. Subsequently, the mixture was elevated to 120 °C and maintained for 24 h. The obtained NH₂-UiO-66 yellow solid was isolated by centrifugation with dimethylformamide, methyl alcohol and DI water at 5000 rpm for 10 minutes. Afterward, the powder was activated at 60 °C to remove any remaining solvent.

The obtained NH₂-UiO-66 was modified with polyethyleneimine using a wet 135 impregnation method. To eliminate coordinated water, NH2-UiO-66 particles were 136 heated at 150 °C overnight. Next, 30 mg of polyethyleneimine and 240 mg of NH₂-137 UiO-66 were separately disbursed in 24 mL methanol under sonication for 10 min. The 138 NH₂-UiO-66 solution was then disbursed to the polyethyleneimine solution slowly 139 while being sonicated. The mixture solution was agitated at 100 °C for 12 hours within 140 a nitrogen environment. Subsequently, the articles were centrifuged and cleaned with 141 142 absolute ethanol to eliminate solvents. The obtained product called P@NH₂-UiO-66 was dried at 110 °C for 12 h. 143

Based on the room temperature synthesis of NH₂-UiO-66 [35], 17.5 mL HCOOH was 144 pre-mixed with 160 mL deionized (DI) water containing 1.16 g ZrCl₄ under agitation 145 for 15 min. Following this, an additional 0.1 g of polyethyleneimine was incorporated 146 into 50 mL of ethanol with 0.95 g of H₂BDC-NH₂. The blend was then thoroughly 147 mixed for 12 hours at ambient temperature. Upon completion of mixing, a pinkish solid 148 was obtained through centrifugal separation and was further purified by washing with 149 ethanol and deionized water to eliminate contaminants. The product called Pin@NH2-150 UiO-66 was obtained after drying. NH2-UiO-66 was prepared in the same way but 151 without the addition of polyethyleneimine and the product was called NH₂-UiO-66-RT. 152

153 2.3. Fabrication of MMMs

154 0.5g Polyetherimide was added to 4.5 g CH₂Cl₂ at room temperature. Following this, 155 the MOF fillers (P@NH₂-UiO-66 and P_{in}@NH₂-UiO-66) were ground and dispersed 156 into the solution to yield a filler content from 10 to 40 wt%. Subsequently, the combined 157 process of 10 minutes of mechanical agitation and ultrasonic treatment were adopted. 158 The casting solutions were subsequently spread onto a clean glass substrate using an 159 automatic membrane applicator to regulate the membrane thickness. Following the 160 application, CH₂Cl₂ solvent was allowed to evaporate completely at ambient condition for 24 h, followed by additional evaporation periods at 60 °C and 120 °C for 24 h each.
Once the solvent was evaporated, the prepared mixed matrix membranes (MMMs) were

- 163 meticulously stripped from the glass surface. All membranes were reactivated at 120 °C
- for 24 h prior to gas permeability tests. $P@NH_2$ -UiO-66-based and $P_{in}@NH_2$ -UiO-66-
- based membranes with different MOFs loadings were labelled as wt%-P@NH₂-UiO-
- 166 66-PEI and wt%-P_{in}@NH₂-UiO-66-PEI, respectively. As a comparison, 30wt % NH₂-
- 167 UiO-66 was added as a filler to PEI to make 30-NH₂-UiO-66-PEI membrane.
- 168 2.4. Characterization methods

The microstructure and size of the MOF fillers, with the surface properties of the 169 membranes, were investigated using a scanning electron microscopy (SEM, JEOL-170 7900F) and a transmission electron microscopy (TEM, JEOL JEM-2100UHR). 171 Additionally, X-ray powder diffraction (XRD, Bruker D8 Advance) and Fourier 172 transform infrared (FTIR) spectrometry (Bruker Vertex 70V) were employed to analyze 173 the crystallographic characteristics of both the MOF fillers and the membranes. X-Ray 174 photoelectron spectroscopy (XPS) analysis was carried out via a PH 5000 Versaprobe 175 spectrometer. A Quantachrome Autosorb IQ instrument was carried out to measure N2 176 adsorption isotherms at -196 °C. The CO₂ adsorption capacities of MOFs and 177 membranes were also measured by a Quantachrome, Auto sorb IQ instrument at 25°C. 178 The CO₂ and CH₄ adsorption isotherms for the membranes were measured with the 179 Micromeritics ASAP-2020 surface area analyzer at a temperature of 25°C. 180 Thermogravimetric (TGA) and differential scanning calorimetry (DSC) analyses were 181 conducted using NETZSCH (STA 449 F5) to assess the mass loss and glass transition 182 temperature T_g of MMMs, respectively. The nitrogen elemental analysis was performed 183 with a Vario Elementar cube. The kinetic viscosity of the casting solutions was 184 measured by Rheometer (Haake Mars 60). 185

186 *2.5. Gas-separation tests*

The membranes were integrated into a stainless-steel chamber for conditioning, prior to the gas separation experiments. The gas permeability of the membranes was determined using the Wicke-Kallenbach technique, maintaining a feed pressure of 1 bar and a delta pressure across the membrane of 1 bar. The inlet gas was f supplied at a rate of 100 mL min⁻¹ (equivalent to 50 cm³ min⁻¹ for each of the binary gas mixtures). A sweeping argon of 40 cm³ min⁻¹ was applied. For analyzing the composition of the permeated gases, a gas chromatography setup (456-GC, Scion) was employed. Average permeation data were calculated based on measurements from at least three different membranes. The permeability, which is also referred to permeance P_i in Barrer (1 Barrer $=1 \times 10^{-10}$ cm³ (STP)·cm·cm⁻² s⁻¹·cmHg⁻¹), was calculated using Equation (1) for the membranes:

$$P_i = \frac{10^{-10} l N_i}{A \Delta p_i} \tag{1}$$

In the specified formula, N_i (cm³ (STP)·s⁻¹) is the permeate flux, l (cm) denotes the membrane thickness, Δp_i (cmHg) is the pressure difference across the membrane, and A (cm²) indicates the membrane area.

202 The separation factor $(\alpha_{i/j})$ of the membrane was calculated based on Equation (2):

203
$$\alpha_{i/j} = \frac{X_i/X_j}{Y_i/Y_i}$$
(2)

204 Where X_i/X_j and Y_i/Y_j represent the volume fraction in the permeate gas and feed gas.

The solution-diffusion model was applied to study the gas permeance through the membranes. The calculation formula is shown in Equation (3):

$$P_i l = S_i \times D_i \qquad (3)$$

where the permeability $(P_i l)$ is equal to the product of the solubility $(S_i, \text{ mol } m^{-3} Pa^{-1})$ and the diffusivity $(D_i, m^2 s^{-1})$, and *l* is the thickness of membranes.

210 The S_i can be calculated by Equation (4):

$$S_i = \frac{c_i}{p} \qquad (4)$$

where C_i (mmol cm⁻³) and P are the adsorption amount of the membrane for component *i* and pressure, respectively. D_i was calculated according to Equation (3).

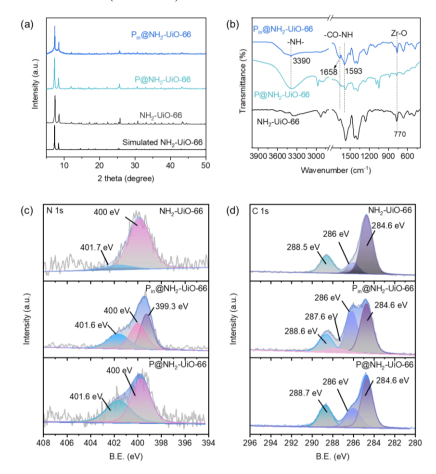
214 **3. Results and discussion**

215 *3.1. NH*₂-*UiO*-66, *P*@*NH*₂-*UiO*-66 and *P*_{in}@*NH*₂-*UiO*-66 nano-fillers

XRD analysis (Fig. 1a) showed that NH₂-UiO-66 and P@NH₂-UiO-66 samples exhibit 216 characteristic peaks 7.3° and 8.4° two theta, matching well the simulated NH₂-UiO-66 217 XRD pattern documented in the previous research [36]. This suggested that the wet 218 impregnation method used for synthesizing P@NH₂-UiO-66 preserves the crystalline 219 220 arrangement of NH₂-UiO-66. In the case of P_{in}@NH₂-UiO-66 prepared by in situ, onepot synthesis, the typical NH₂-UiO-66 peaks at 7.3° and 8.4° two theta were observed, 221 222 together with an amorphous peak in the range 3-12° two theta, indicating the presence of polyethyleneimine in the composite. The crystallinity of NH2-UiO-66 in Pin@NH2-223 UiO-66 was calculated via the background constant method [37, 38]. The crystallinity 224 of NH₂-UiO-66 was assessed to be 100%, and the relative crystallinity of P_{in}@NH₂-225 UiO-66 was found to be 60.4%. During the in situ preparation of P_{in} (2)NH₂-UiO-66, the 226 227 introduction of polyethyleneimine interfered with the coordination between the carboxyl groups and the metal zirconium (Zr), thereby affecting the nucleation 228 procedure of NH₂-UiO-66 and leading to a reduced crystallinity. 229

From the FTIR spectra (Fig. 1b), all samples exhibit a peak at 770 cm⁻¹, associated with 230 the of the Zr-O vibration in NH₂-UiO-66. The -NH- vibration of polyethyleneimine can 231 be seen at 3390 cm⁻¹, indicating the effective grafting of polyethyleneimine onto NH₂-232 UiO-66. In contrast to P@NH2-UiO-66, new characteristic peaks of the amide bond 233 appear in P_{in}@NH₂-UiO-66 at 1658 cm⁻¹ [39] and 1593 cm⁻¹ [33], primarily arising 234 from the condensation reaction between the carboxyl group in H₂BDC-NH₂ and the 235 amino group in polyethyleneimine [28, 39]. In P@NH₂-UiO-66, the amine groups can 236 bind to open Zr sites as reported [33, 40], however, this interaction is relatively weak, 237 238 resulting in only modest quantity of polyethyleneimine being grafted onto the NH₂-UiO-66. The robust interaction enables P_{in} (∂NH_2 -UiO-66 to combine a greater quantity 239 of polyethyleneimine compared to P@NH2-UiO-66. The nitrogen content was 240 determined by elemental analyzer to compare the polyethyleneimine content of the 241 three samples. The nitrogen contents of Pin@NH2-UiO-66, P@NH2-UiO-66, and NH2-242

UiO-66 were 6.56%, 4.78%, and 4.51%, respectively, indicating that Pin@NH2-UiO-66 243 binds enhanced polyethyleneimine compared to P@NH2-UiO-66. The interaction was 244 further examined by XPS (Fig. 1 c and d). The N1s peaks at 400 eV and 401.8 eV 245 correspond to the -NH₂ and -NH groups of NH₂-UiO-66, respectively (Fig. 1c). The 246 emergence of a distinct peak at 399.3 eV in the XPS spectrum corresponds to the amide 247 bonds in Pin@NH2-UiO-66 [41]. Further, the C1s XPS spectra (Fig. 1d) indicate that 248 NH₂-UiO-66 reveals three distinct peaks at 284.5eV, 286 eV and 288.5eV, which are 249 respectively belong to sp² C, C-N, and O-C-O groups. The C-N ratio of the two 250 composites increases compared with that of NH2-UiO-66, with the C-N ratio in 251 P_{in} @NH₂-UiO-66 being higher, confirming that more polyethyleneimine is present in 252 P_{in}@NH₂-UiO-66. Additionally, the characteristic peak of amide carbon was observed 253 at 287.6 eV for Pin@NH2-UiO-66 [42]. The XPS results show that there is a 254 condensation reaction between the amino functional group and the carboxyl functional 255 group to form an amide (-CO-NH-). 256

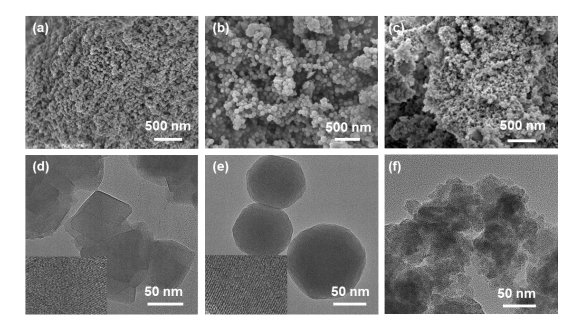


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Fig. 1. (a) XRD, (b) FTIR, and (c) XPS N1s analysis and (d) XPS C1s analysis of the as-made NH₂-258 10 / 27

259 UiO-66, P@NH₂-UiO-66, and P_{in}@NH₂-UiO-66 powders.

Fig. 2 exhibits the SEM and TEM images of the as-made NH2-UiO-66, P@NH2-UiO-260 66 and Pin@NH2-UiO-66 powders. The NH2-UiO-66 nanoparticles exhibit dimensions 261 between 50 and 75 nm, exhibiting a cubic-like shape (Fig. 2a and d), while the P@NH₂-262 UiO-66 are spherical with particle sizes around 100 nm (Fig. 2b and e). The differences 263 in the shape and size of P@NH₂-UiO-66 nanoparticles are considered to be a result of 264 the encapsulation of polyethyleneimine. The SEM image of NH₂-UiO-66-RT prepared 265 in the absence of polyethyleneimine indicates a cubic-like mosphology of this sample 266 with particle sizes of about 100 nm, and an XRD pattern that matches with the simulated 267 NH₂-UiO-66 (Fig. S1). Relatively, P_{in}@NH₂-UiO-66 consisted of significantly smaller 268 nanoparticles (~ 20 nm) (Fig. 2c and f). The condensation reaction to form amide groups 269 competes for the carboxyl group that should be coordinated with the metal Zr, thus 270 affecting the nucleation process of MOF and leading to the miniaturization of the MOF 271 particles [43]. The reduced particle size of the latter filler enhances the surface area and 272 273 could improve compatibility with the polymer matrix.



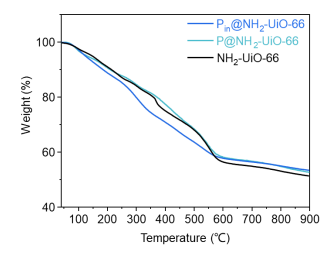
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Fig. 2. SEM and TEM images of the as-made (a, d) NH₂-UiO-66, (b, e) P@NH₂-UiO-66, and (c, f)

- 276 P_{in} @NH₂-UiO-66. The lower left corner of d and e is a high-resolution image.
- 277 The N₂ adsorption isotherms of the NH₂-UiO-66, P@NH₂-UiO-66 and P_{in}@NH₂-UiO-

66 all followed hybrid type I and IV isotherms with H4 hysteresis (Fig. S2). Calculated 278 by the t-plot method, the micropore volume of NH2-UiO-66, P@NH2-UiO-66 and 279 P_{in} @NH₂-UiO-66 were 0.45 cm³/g, 0.34 cm³/g and 0.21 cm³/g (Table S1). The pore 280 size distribution reveals that the majority of pores measure approximately 0.4-0.6 nm 281 and 1.1 nm. The latter represents the mesopores accumulated by micropores. There are 282 also mesopores that are generated by nanoparticle accumulation in Pin@NH2-UiO-66. 283 While for the P@NH₂-UiO-66, the pore size distribution reveals lower value at around 284 1.1 nm, indicating less intra-particle mesopores. In addition, there were less mesopores 285 that accumulated by the nanoparticles due to the larger particle size of P@NH2-UiO-286 66. The mesopores allow rapid gas diffusion into the pores of fillers, further 287 significantly decrease the trans-membrane mass transfer resistance of the MMMs, 288 which facilitates diffusion processes. Therefore, the diffusion coefficient of P@NH2-289 UiO-66-PEI membrane will be lower. The micropore diameters of NH₂-UiO-66 was 290 0.55 nm, being close to the MOF structure windows of about 6 Å, and the pore size of 291 P@NH₂-UiO-66 and P_{in}@NH₂-UiO-66 reveal a slightly decrease (0.50 and 0.44 nm). 292 293 The both reduction of pore volume and pore size indicates the occupation of the pores by polyethyleneimine. The Pin@NH2-UiO-66 binds more polyethyleneimine than the 294 P@NH₂-UiO-66, leading to more severe pore plugging. 295

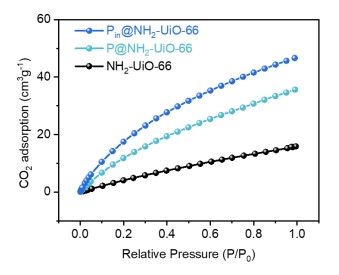
The thermal stability of P@NH₂-UiO-66 and P_{in} @NH₂-UiO-66 was tested under N₂ atmosphere (Fig. 3). The initial mass decrement observed above 100 °C is assigned to the evaporation of adsorbed H₂O and gases. The step mass loss before 280 °C is attributed to dehydroxylation of the zirconium oxo clusters. The subsequent mass reduction occurring beyond 280 °C is ascribed to the structural collapse of NH₂-UiO-66 [35, 44]. These results show that the P@NH₂-UiO-66 and P_{in}@NH₂-UiO-66 fillers have good thermal stability.



303

Fig. 3. TGA in N₂ of the as-made NH₂-UiO-66, P@NH₂-UiO-66, and P_{in}@NH₂-UiO-66.

Pin@NH2-UiO-66 and P@NH2-UiO-66 exhibited CO2 saturation adsorption capacities 305 of 46 and 35 cm³g⁻¹, respectively, surpassing that of NH₂-UiO-66 (15 cm³g⁻¹) (Fig. 4). 306 The increase of CO₂ adsorption was mainly related to the modification of MOF by 307 polyethyleneimine. The grafting of polyethyleneimine with rich amino groups to NH2-308 UiO-66 generates additional sites for CO₂ adsorption. Furthermore, from the elemental 309 analysis and XPS results (Fig. 1c, d), Pin@NH2-UiO-66 has a higher polyethyleneimine 310 content, therefore it has the highest CO₂ adsorption capacity. This high CO₂ adsorption 311 capacity is beneficial for the CO₂/CH₄ separation performance of the MMMs. 312



313

Fig. 4. The CO₂-adsorption curves of the as-made NH₂-UiO-66, P@NH₂-UiO-66, and P_{in}@NH₂UiO-66.

316 *3.2. Membrane characterization*

SEM images in Fig. 5, Fig. S3 and Fig. S4 were applied to characterize the top and 317 cross-sectional structure of the pure PEI, P@NH2-UiO-66-PEI, and Pin@NH2-UiO-66-318 PEI membranes. The fabricated MMMs exhibited a substantial surface roughness, 319 which tends to increase with the loading, compared to the PEI membrane (Fig. 5 a-c, 320 Fig. S3 a-e and Fig. S4 a-d). Agglomeration was noticed on the surface of the P@NH2-321 UiO-66-PEI membrane (Fig. 5c and Fig. S3), indicating a less-than-ideal distribution 322 of particles throughout the PEI matrix. In the cross-section SEM images (Fig. S3 f-j 323 and Fig. S4 f-j), the thickness of the PEI, Pin@NH2-UiO-66-PEI, and P@NH2-UiO-66-324 PEI membranes were determined to be in the range 68-91 µm. For Pin@NH2-UiO-66-325 PEI, it is evident that P_{in}@NH₂-UiO-66 is enveloped by the polymer chains, forming 326 defect-free microstructures (Fig. 5e and Fig. S4). The emergence of polymer veins 327 serves as additional evidence supporting the existence of robust interfacial interactions 328 329 [45], while this polymer vein was invisible in P@NH₂-UiO-66 based MMMs (The upper right corner of Fig. 5e and f). The outcome of this effect could be attributable to 330 the presence of interaction forces between the imine groups (-N(R)-H) of the 331 polyethyleneimine and the carbonyl groups (C=O) of the PEI matrix. The EDS mapping 332 in Fig.5g-i shows that the nitrogen content of Pin@NH2-UiO-66-PEI membrane is 333 higher than that of P@NH2-UiO-66-PEI and pure PEI membranes, further proving the 334 nitrogen content of Pin@NH2-UiO-66-PEI is the highest. 335

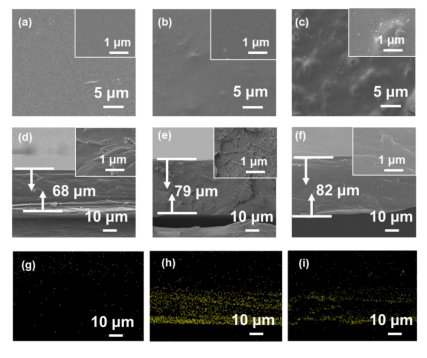


Fig. 5. Top-view SEM images of (a) pure PEI membrane, (b) 30-P_{in}@NH₂-UiO-66-PEI, (c) 20P@NH₂-UiO-66-PEI, and cross-section SEM images of (d) pure PEI membrane, (e) 30-P_{in}@NH₂-UiO66-PEI and (f) 20-P@NH₂-UiO-66-PEI (The upper right corner of a-f is the enlarged image),
Corresponding elemental mapping images of N ka of (g) pure PEI membrane, (h) 30-P_{in}@NH₂-UiO66-PEI, (i) 20-P@NH₂-UiO-66-PEI

XRD characterization was conducted to further characterize the structure of as-made 342 membranes. Pristine PEI is a copolymer that features broad peaks centered at 14° two 343 theta (Fig. 6a) [46]. The MOFs preserve their crystallinity and phase within the PEI as 344 seen by the peaks at 7.37° and 8.44° two theta. Fig. S5 a, b displays the XRD patterns 345 of P@NH2-UiO-66-PEI and Pin@NH2-UiO-66-PEI membranes prepared with different 346 loadings. As the loading of the MOF increased, the intensity of the MOF characteristic 347 peaks (especially at 7.37° and 8.44° two theta) progressively enhanced, indicating that 348 the crystallinity of the nano-filler can be maintained during the membrane preparation 349 process. The d-spacing values were calculated by using the Bragg's law. This value was 350 lower for MMMs (d=5.7-5.9Å) than pristine PEI membrane (d=6.4 Å), as shown in the 351 352 Fig .S5a, b. Thus form the result, the interaction between filler and polymer matrix increased the rigidity of the membrane, reducing the inter-segmental mobility and d-353 spacing, and enabling the membrane to allow only small molecules to penetrate through 354 it. Given that the CO₂ molecule (3.6 Å) is smaller in size compared to the CH₄ molecule 355 (3.8 Å), the MMMs potentially facilitates the passage of CO₂ through the membrane, 356 thus enhancing CO₂ permeability. 357

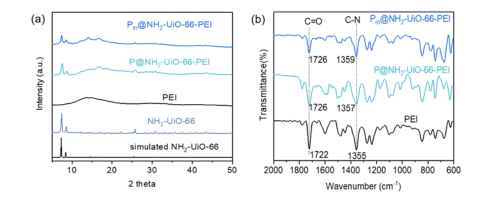


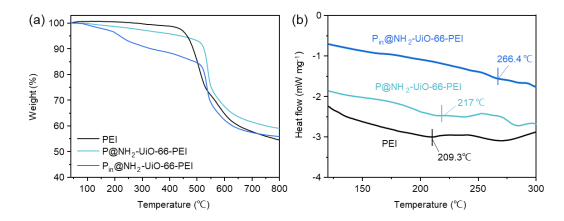
Fig. 6. XRD patterns (a) and FTIR spectra (b) of pristine PEI, P@NH₂-UiO-66-PEI and P_{in}@NH₂-

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360 UiO-66-PEI membranes.

FTIR spectra of the membrane samples (Fig. 6b) were obtained to acquire insight of 361 the interaction between incorporated P@NH2-UiO-66 or Pin@NH2-UiO-66 MOFs and 362 PEI. The FTIR spectrum of the PEI matches with that described in published studies 363 [47]. The characteristic peaks corresponding to the pristine PEI polymer remain intact 364 in all MMMs, showing no noticeable changes in intensity. As the filler loading 365 increased, the peaks observed at 1780 cm⁻¹ (C=O asymmetric), 1722 cm⁻¹ (C=O 366 symmetric stretching vibration), and 1355 cm⁻¹ (C-N stretch of imide groups) (Fig. S5 367 c, d), which were slightly red-shifted, and their intensity gradually increased. This could 368 be can be ascribed to the interaction between the amide groups and the carbonyl (C=O) 369 groups of the PEI via hydrogen bonding. Pin@NH2-UiO-66-PEI exhibited enhanced 370 shift, suggesting significantly intensified hydrogen bonds. 371

The XRD pattern of 30-NH₂-UiO-66-PEI membrane matches with the simulated NH₂UiO-66 and pure PEI (Fig. S6). The top-view SEM images of 30-NH₂-UiO-66-PEI
indicates a rough surface morphology with particles agglomerated. The cross-section
SEM images shows obvious interface defects and voids, with membrane thickness of
59 μm.



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Fig. 7. TGA curves (a) and DSC curves (b) of pristine PEI, P@NH₂-UiO-66-PEI and P_{in}@NH₂-UiO66-PEI membranes.

The thermal stability of mixed matrix membranes was investigated using
thermogravimetric measurements (Fig. 7a). Starting from 200 °C, MMMs experienced

weight loss due to MOF degradation and the decomposing of polyethyleneimine of the 382 fillers. The mass loss begin at 450-500 °C occurred as a result of PEI decarboxylation 383 [46]. The MMMs with Pin@NH2-UiO-66 and P@NH2-UiO-66 fillers demonstrated a 384 larger residual mass than the pristine PEI, suggesting filler decomposition. The glass 385 transition temperature (Tg) determined by DSC measurements (Fig. 7b and Table.S2) 386 387 was performed to assess the interaction between the fillers and the PEI. Pristine PEI membrane exhibited a T_g of 209.3 °C which matched well with the literature [48]. The 388 30-Pin@NH2-UiO-66-PEI exhibited a much higher Tg value (266.4 °C) compared to 20-389 P@NH₂-UiO-66-PEI (217 °C). The reason is that more amine groups on the P_{in}@NH₂-390 UiO-66 show stronger hydrogen bonding with the PEI matrix, which indicates that 391 Pin@NH2-UiO-66-PEI have higher chain rigidity with robust interfacial interactions 392 within the membranes. In addition, with the increase of filler content, the Tg of MMM 393 394 increased, while the T_g of 40-P_{in}@NH₂-UiO-66-PEI membrane decreased slightly, which was due to the fact that overmuch filler was prone to agglomeration in the matrix 395 and weakened the interfacial compatibility. 396

The viscosity of the casting solutions was measured to further investigate the interactions between the filler and the polymer (Fig. S7). The $P_{in}@NH_2$ -UiO-66-PEI solution exhibited superior viscosity compared to that of the P@NH_2-UiO-66-PEI solution and the PEI polymer, confirming the enhanced interfacial compatibility [49, 50]. The enhanced interface compatibility is beneficial for the selective separation effect of CO₂.

403 *3.3. Gas separation performance*

Mixed gas permeation measurements (with $CO_2/CH_4 = 50/50$ by volume) were performed to evaluate the separation capabilities of prepared membranes. Initially, the impact of P@NH₂-UiO-66 and P_{in}@NH₂-UiO-66 loading was examined. (Fig. 8a, b and Table S3). As exhibited in Table S3, the modified Zr-MOF MMMs exhibit higher CO₂ permeability and CO₂/CH₄ selectivity compared to NH₂-UiO-66-PEI. The aggregation caused by the larger crystal size of NH₂-UiO-66 and inhomogeneous distribution results in nonselective voids, leading to the lower selectivity for CO₂/CH₄.

And the low CO₂ saturated adsorption capacity of NH₂-UiO-66 leads to the lower CO₂ 411 permeability. As the increase in both filler loadings, the CO₂ permeability increases, 412 mainly because the introduction of more MOFs provides more gas transport channels 413 for MMMs. Correspondingly, the CO₂/CH₄ selectivity demonstrated a tendency of 414 initial of incremental rise initially, which was succeeded by a decline. The increase of 415 416 polyethyleneimine load in the membrane introduced more CO₂ adsorption sites, which resulted in higher gas selectivity. When the filler loading was too high, non-selective 417 voids were generated due to nanofiller aggregation (for instance Fig. S3h), reducing the 418 CO₂/CH₄ selectivity. For P_{in}@NH₂-UiO-66-PEI membrane, the MMM with 30% filler 419 loadings exhibited the best performance. The 30-Pin@NH2-UiO-66-PEI membrane 420 exhibited the CO₂ permeability of 2498.9 Barrer with CO₂/CH₄ selectivity of 27.7, 421 respectively. The 20-P@NH2-UiO-66-PEI demonstrated best results for the P@NH2-422 UiO-66 filler with a CO₂ permeability of 484.2 Barrer with CO₂/CH₄ selectivity of 11.4, 423 respectively. The significant difference in separation performance of Pin@NH2-UiO-66 424 and P@NH2-UiO-66 based membranes was mainly related to the amount of 425 426 polyethyleneimine modification. The membranes with higher polyethyleneimine loading capacity showed superior CO₂/CH₄ separation performance. 427

Fig. 8c revealed that the incorporation of P@NH2-UiO-66 and Pin@NH2-UiO-66 nano-428 fillers improved the selectivity and permeability for both gases over the pure PEI 429 membranes. The Pin@NH2-UiO-66-PEI exhibited enhanced CO2/CH4 performance 430 compared to P@NH₂-UiO-66-PEI, which was due to the improved CO₂ affinity of 431 Pin@NH2-UiO-66 and better interfacial compatibility the PEI matrix. The CO2 432 permeability and CO₂/CH₄ selectivity of the 30-P_{in}@NH₂-UiO-66-PEI membrane was 433 respectively 4 and 2.7 times of the P@NH₂-UiO-66-PEI membrane and much higher 434 than the pure PEI membrane. 435

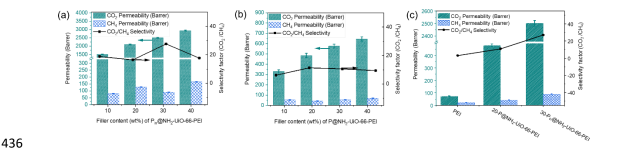


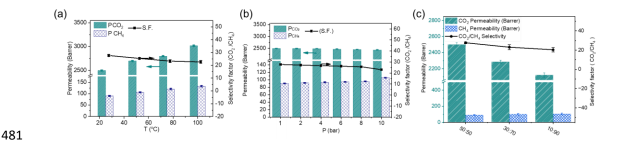
Fig. 8. The effect of fillers loading on the CO₂/CH₄ separation efficiencies of (a) P_{in}NH₂-UiO-66-PEI
and (b) P@NH₂-UiO-66-PEI membranes. (c) Contrast of CO₂/CH₄ separation efficiencies of PEI, 20P@NH₂-UiO-66-PEI and 30-P_{in}NH₂-UiO-66-PEI membranes.

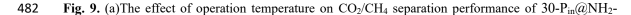
To illustrate the improvement of CO_2 adsorption capacity of the membranes, the gas 440 adsorption experiments were performed on the P@NH2-UiO-66-PEI membrane and 441 Pin@NH2-UiO-66-PEI membrane with CO2 and CH4 at 298 K (Fig. S8). The adsorption 442 capacities of the P@NH2-UiO-66-PEI membrane for CO2 and CH4 were 10.20 and 3.31 443 cm³ g⁻¹ at 298 K, while those of P_{in}@NH₂-UiO-66-PEI membrane for CO₂ and CH₄ 444 were 25.33 and 4.12 cm³ g⁻¹ at 298 K. The CO₂ adsorption ability of the P_{in} @NH₂-445 446 UiO-66-PEI membrane is higher than that of the P@NH2-UiO-66-PEI membrane, proving that the in-situ modification of polyethyleneimine to the fillers can effectively 447 improve the CO₂ adsorption capacity of the membrane. The solubility (S) and 448 diffusivity (D) for the 30-P_{in}@NH₂-UiO-66-PEI and P@NH₂-UiO-66-PEI membranes 449 are summarized in Table S4. The result shows that the selectivity of the membrane is 450 more dependent on solubility ($S_{CO2}/S_{CH4}=6.4 > D_{CO2}/D_{CH4}=4.3$). It is much easier to 451 adsorb CO₂ than CH₄ by the 30-P_{in}@NH₂-UiO-66 membranes, declining the adsorption 452 and diffusion of CH₄, thereby enhancing the separation selectivity. The CO₂ solubility 453 of 30-P_{in}@NH₂-UiO-66-PEI membrane shows higher value $(1.2 \times 10^{-8} > 8.2 \times 10^{-9})$ 454 than 30-P@NH₂-UiO-66-PEI membrane due to the higher CO₂ adsorption ability of 455 membrane. Furthermore, in contrast to the 30-Pin@NH2-UiO-66-PEI membrane, the 456 CO_2 diffusion of the 30-P@NH₂-UiO-66-PEI membrane shows lower value (5.3 × 457 $10^{-6} < 1.1 \times 10^{-5}$), attributed to the reduced presence of mesopores in P@NH₂-UiO-66. 458

Fig. 9a assessed the impact of operational temperature within the range of 25–100 °C. For 30-P_{in}@NH₂-UiO-66-PEI membrane, the permeability of CO₂ and CH₄ exhibited a notable enhancement with increasing operational temperature, while the selectivity of CO₂/CH₄ remained at a high level of 19.4. This shows that P_{in} @NH₂-UiO-66-PEI membrane can maintain excellent CO₂/CH₄ separation performance even at high operating temperatures (100 °C).

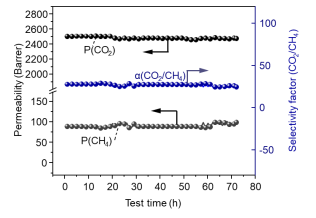
Examining the impact of transmembrane pressures on the performance of the produced 466 membranes is essential. The observed trend in Fig. 9b indicates that the CO₂ 467 permeability gradually decreases with the transmembrane pressure increases from 1 bar 468 to 10 bar. This can be attributed to that the CO_2 affinity of the membrane approaches 469 470 saturation with the feed pressure increasing. As a result, the available sites for CO₂ adsorption become limited, leading to a decline in the overall permeability and 471 472 CO₂/CH₄ selectivity [11, 16]. The CO₂/CH₄ selectivity of the MMMs remains substantially superior to that of the PEI membrane under elevated pressures (e.g. 10 bar, 473 Fig. 9b), emphasizing the enormous potential of these as-made MMMs under 474 operational conditions. 475

The different feed gas pressure of CO_2/CH_4 mixed gases (50/50, 30/70, and 10/90 by volume) was investigated and the results are shown in Fig. 9c. The CO_2/CH_4 selectivity decreased with the increase of CH_4 permeability due to higher CH_4 feed partial pressure. 30-P_{in}@NH₂-UiO-66-PEI still maintained the CO_2/CH_4 selectivity of 21.3 while a permeability of up to 2116 Barrer in the case of a feed volume ratio of 10/90.





- 483 UiO-66-PEI at 1 bar. (b)The effect of operation pressures on CO₂/CH₄ separation performance of 30484 P_{in}@NH₂-UiO-66-PEI at 25 °C. (c) The effect of different feed gas ratios of the CO₂/CH₄ separation
- 485 performance of $30-P_{in}@NH_2-UiO-66-PEI$ at 25 °C and 1 bar.



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Fig. 10. 72 h operation test of 30-P_{in}@NH₂-UiO-66-PEI.

488 3.5. Long-term stability

Fig. 10 presents long-term stability test of CO_2/CH_4 separation to assess the stability of the membrane structure. During a 72 h operation test, the $30-P_{in}@NH_2-UiO-66-PEI$ membrane remained a stable CO_2 permeability of 2498.9 Barrer and a consistent CO_2/CH_4 selectivity of 27.7. Additionally, it confirms that the fabricated $30-P_{in}@NH_2-$ UiO-66-PEI membrane exhibited robust structural stability, indicating its potential suitability for CO_2 capture from natural gas applications.

495 *3.6. Simulation of industrial environment*

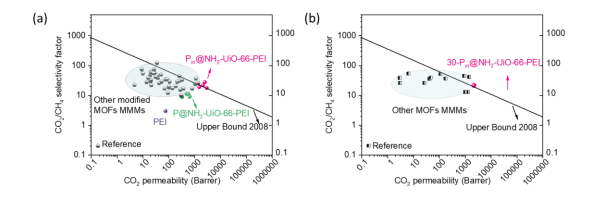
To further simulate natural gas industrial conditions, we conducted tests to evaluate the 496 separation performance with feed gas CO_2 : $CH_4 = 10$: 90 at various temperatures and 497 pressures (Table S5 and Table S6). The CO₂ permeability increases with increasing 498 temperature (25-50 °C) at 1 bar (Table S5). The CO₂ permeability reached 2363 Barrer 499 with the selectivity of 19.3 at 50 °C. As shown in Table. S6, when the pressure further 500 increased, the mixed-gas selectivity slightly decreased. The CO₂ permeability remained 501 at 2133 Barrer with CO₂/CH₄ selectivity of 15.1 at 50 °C and 10 bar, proving that the 502 Pin@NH2-UiO-66-PEI membrane has excellent CO2 permeability. MOFs-based 503 MMMs with mixed CO₂/CH₄ (volume ratio 10/90) reported in the literature are detailed 504 in Table S6, with 30-Pin@NH2-UiO-66-PEI showing superior CO2 permeability. 505

Therefore, the high-performance P_{in}@NH₂-UiO-66-PEI membrane holds significant
 potential for industrial applications.

The 30-P_{in}@NH₂-UiO-66-PEI was subjected to a pressure of 60 bar for 3 hours using 508 a preforming machine to emulate the stability under high-pressure environment. A 509 comparison between the optical photographs and the surface SEM images of the 510 membrane taken before and after the application of pressure (Fig. S9). The membrane's 511 surface retained its initial morphology with some indentations and wrinkles after the 512 pressure exposure. Subsequently, the gas separation efficiency of mixed CO₂/CH₄ 513 (volume ratio 10/90) of this membrane was tested at 50 °C and 10 bar, and the findings 514 indicated that the CO₂/CH₄ separation performance were maintained at a high level 515 (Table. S7). 516

517 *3.7. Comparison with upper bound*

Fig. 11 a shows the CO₂/CH₄ separation performances for the P_{in}@NH₂-UiO-66-PEI, 518 P@NH2-UiO-66-PEI membranes compared with reported other state-of-the-art 519 functional groups modified MOFs MMMs and 2008 Robeson upper bounds. It can be 520 indicated that CO₂ permeability and CO₂/CH₄ selectivity of P_{in}@NH₂-UiO-66-PEI 521 exhibited significant enhancements compared to other functional group-modified 522 MOFs membranes. Moreover, Fig. 11 b and Table S7 illustrates the comparison of 523 Pin@NH₂-UiO-66-PEI and reported other MMMs that were measured with a mixed 524 feed gas of CO₂/CH₄=10/90. The P_{in}@NH₂-UiO-66-PEI membrane stands out due to 525 its exceptionally high CO₂ permeability compared to other membranes. In addition, 30-526 Pin@NH2-UiO-66-PEI demonstrated enhanced CO2/CH4 separation performance, 527 exceeding the upper bound established in 2008, demonstrating the developed 528 529 membrane with high potentiality for natural gas purification.



530

Fig. 11. Contrast of CO₂/ CH₄ separation performances of 30-P_{in}@NH₂-UiO-66-PEI with (a) Robeson
upper bound and other modified MOFs based MMMs, (b) other MOFs MMMs in feed gas CO₂/CH₄
=10/90.

534 Conclusions

In summary, this study reported the synthesis of polyethyleneimine-modified MOF 535 fillers via an in situ one-pot method, and then incorporated into PEI to construct mixed 536 matrix membranes for efficient CO₂/CH₄ separation. The separation performance of as-537 synthesized MMMs (Pin@NH2-UiO-66-PEI) was also compared with that of MMMs 538 with fillers prepared via conventional wet impregnation method (P@NH₂-UiO-66-PEI). 539 Owing to the presence of polyethyleneimine in the in situ one-pot synthesis, P_{in} @NH₂-540 UiO-66 fillers have smaller particle size (~20 nm) and higher CO₂ adsorption capacity, 541 improving the interfacial compatibility of MMM and the CO₂ transfer capability. The 542 CO2 permeability and CO2/CH4 selectivity of 30-Pin@NH2-UiO-66-PEI MMMs (30 wt% 543 loading) reached 2498.9 Barrer and 27.7, which were remarkably increased than that of 544 pristine PEI membranes (75.1 Barrer and 3.3, respectively) and P@NH₂-UiO-66-PEI 545 membranes (484.2 Barrer and 11.4, respectively). In addition, we simulated the 546 industrial natural gas purification conditions (50 °C, 10 bar and CO₂: $CH_4 = 10$: 90 feed 547 gas), and the Pin@NH2-UiO-66-PEI membranes still have CO2 permeability of 2133 548 Barrer and CO₂/CH₄ selectivity of 15.1 under these conditions. This study provides a 549 novel perspective on the reasonable design of nano-fillers with unique properties and 550 solve the interfacial compatibility of the MMMs for the development of natural gas 551 purification membranes. 552

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