1 Article

2 **Glassy PEEK-WC vs rubbery Pebax**®1657 polymers:

³ effect on the gas transport in CuNi-MOF based mixed

4 matrix membranes

5 Elisa Esposito^{1*}, Rosaria Bruno², Marcello Monteleone¹, Alessio Fuoco¹, Jesús Ferrando-Soria³, 6 Emilio Pardo³, Donatella Armentano², Johannes Carolus Jansen^{1*}

⁷ ¹Institute on Membrane Technology, CNR-ITM, Via P. Bucci 17/C, 87036 Rende (CS), Italy.

8 <u>e.esposito@itm.cnr.it (E.E.); m.monteleone@itm.cnr.it (M.M.); a.fuoco@itm.cnr.it (A.F.)</u>

⁹ Dipartimento di Chimica e Tecnologie Chimiche (CTC), Università della Calabria, Rende 87036, Cosenza,
 ¹⁰ Italy. rosaria.bruno@unical.it<u>(R.B),</u> donatella.armentano@unical.it<u>(D.A.).</u>

- ³ Departamento de Química Inorgánica, Instituto de Ciencia Molecular (ICMol), Catedrático José Beltrán
 Martínez, 2, Universidad de Valencia, 46980 Paterna, Valencia, Spain. jesus.ferrando@uv.es
 (J.F.); emilio.pardo@uv.es (E.P.)
- 15 * Correspondence: <u>e.esposito@itm.cnr.it(E.E.), johannescarolus.jansen@cnr.it</u>(J.C.J.); +39-0984-492008 (E.E.) +39-0984-492031(J.C.J.)
- 17 Received: date; Accepted: date; Published: date
- 18

19 Abstract: Mixed matrix membranes (MMMs) are seen as promising candidates to overcome the 20 fundamental limit of polymeric membranes, known as the so-called Robeson upper bound, which 21 defines the best compromise between permeability and selectivity of neat polymeric membranes. 22 To overcome this limit, the permeability of the filler particles in the MMM must be carefully 23 matched with that of the polymer matrix. The present work shows that it is not sufficient to match 24 only the permeability of the polymer and the dispersed phase, but that one should consider also the 25 individual contributions of the diffusivity and the solubility of the gas in both components. Here 26 we compare the gas transport performance of two different MMMs, containing the metal organic 27 framework CuNi-MOF in the rubbery Pebax®1657 and in the glassy poly(ether-ether-ketone) with 28 cardo moiety, PEEK-WC. The chemical and structural properties of MMMs were investigated by 29 means of FT-IR spectroscopy, scanning electron microscopy and EDX analysis. The influence of 30 MOF on the mechanical and thermal properties of both polymers was investigated by tensile tests 31 and differential scanning calorimetry, respectively. The MOF loading in Pebax®1657 increased the 32 ideal H₂/N₂ selectivity from 6 to 8 thanks to an increased H₂ permeability. In general, the MOF had 33 little effect on the Pebax®165 membranes because an increase in gas solubility was neutralized by 34 an equivalent decrease in effective diffusivity. Instead, the addition of MOF to PEEK-WC increases 35 the ideal CO₂/CH₄ selectivity from 30 to ~48 thanks to an increased CO₂ permeability (from 6 to 48 36 Barrer). The increase in CO₂ permeability and CO₂/CH₄ selectivity is maintained under mixed gas 37 conditions.

38 Keywords: Mixed matrix membrane, glassy polymer, rubbery polymer, PEEK-WC, Pebax®1657,
 39 Gas separation, CuNi-MOF

- 40
- 41

42 **1. Introduction**

43 The research field on materials for gas separation membranes is constantly expanding due to the 44 pressing industrial request for more performing materials to employ in gas treatment, such as 45 hydrogen recovery (H₂/CO₂) [1], oxygen-enriched air production (O₂/N₂) [2,3], biogas up-grading 46 [4,5] and natural gas treatment (CO₂/CH₄) [6], and post combustion carbon capture from flue gas 47 (CO₂/N₂) [7]. This increasing interest is dictated by the advantages of membrane technology 48 compared to traditional gas separation techniques. Gas separation by means of membrane 49 technology is an economic process, it is easily scalable and it can be used in non-drastic temperature 50 and pressure conditions, which are more environmentally friendly. Despite numerous efforts to 51 develop new materials for gas separation, the membrane market still need to overcome some 52 challenges. In fact, higher permeable rubbery polymers present low selectivity and the high selective 53 glassy polymers are less permeable. To overcome this trade-off, extensively reported by Robeson in 54 1991 and in 2008 [8,9], research is focusing on the design and development of hybrid membranes 55 based on the combination of two different materials, in order to get advantages of both [10]. 56 According to this concept, a valid strategy is the embedding porous metal-organic frameworks 57 (MOFs) in the polymer matrix in order to obtain mixed matrix membranes (MMMs) with enhanced 58 gas transport properties [11–13]. MOFs are an attractive new class of microporous materials built by 59 the combination of metal atoms/clusters with a wide variety of organic ligands, which can be 60 specifically designed for improving the compatibility with the organic polymer phase and with high 61 specify for gases [14,15]. The gas transport in polymer membranes occurs following the solution-62 diffusion mechanism in which permeability and selectivity are determined by kinetic parameters 63 (diffusion) and thermodynamic factors (solubility), via P = D * S; $\alpha_{x/y} = D_x/D_y * S_x/S_y$. The 64 addition of MOFs can improve the gas transport properties of the neat polymer by influencing these 65 two parameters [10]. The diffusion, being a kinetic phenomenon, is strongly linked to the free volume 66 of polymer materials and to the molecular size of the penetrating species. In the MOFs, the link 67 between metals and organic units forms a three-dimensional structure with cages having well-68 defined shape and size, which can improve free volume elements of matrix, increasing gas transport 69 in terms of permeability. At the same time, the cages of MOF forming a preferential pathway for a 70 specific gas can act as a molecular sieve increasing membrane selectivity. On the contrary, the main 71 factor that determines the solubility in a polymer matrix is the ability of the penetrant gases to 72 condense, which in turn is correlated with the interactions that occur between the gas and the matrix 73 of membrane. MOFs chemical structure can be easily designed or functionalized synthetically in 74 order to improve its affinity for specific gases [16]. In this case, the membrane permeability is 75 expected to improve as a consequence of an increased contribution in solubility due to the gas 76 condensability in the MOF. Furthermore, the introduction of chemical groups with a specific affinity 77 to one gas in a mixture will also increase its selectivity. The choice of the materials combination for 78 MMMs preparation must be made on the basis of specific physical and chemical properties, in order 79 to tailor in advance, the membrane for a desired gas separation. For this reason, it is necessary to 80 understand how the addition of MOF can influence the gas transport properties of different 81 polymers, rubbery or glassy.

In this work, the gas transport properties for MMMs prepared by embedding the same oxamatebased MOF in two different polymers, Pebax®1657 rubbery polymer and PEEK-WC glassy polymer,
will therefore be investigated. The oxamate-based MOF, previously reported [17], with formula
NiII2{NiII4[CuII2(Me3mpba)2]3}·54H2O (where Me3mpba is the N,N'-2,4,6-trimethyl-1,3-phenylenebis
(oxamate) ligand), has already shown interesting gas separation properties [18].

Single-crystal X-ray diffraction measurements unveil the crystal structure of CuNi-MOF. The anionic Ni^{II}₆Cu^{II}₆ open-framework structure, exhibits a pillared square/octagonal layer architecture, where nickel(II) and copper(II) ions are located on the vertices and midpoints of the edges, respectively, featuring three types of pores, different in size and shape, propagating along the c axis and enfolding up to 60% of the total lattice volume. Free nickel(II) cations are further accommodated within pores of the MOF. It consists of regularly spaced, small almost square sized pores (virtual diameter of ca 0.4 nm) and two kinds of hydrophobic and hydrophilic octagonal pores (Figure 1)

94 resulting from the different disposition of the trimethyl-substituted phenylene spacers, pointing 95 inwards or outwards of the voids, which accounts for their virtual diameters of ca. 1.5 and 2.2 nm, 96 respectively (Figure 1).

97 PEEK-WC is a glassy poly(ether-ether-ketone) with a cardo group in the polymer backbone, 98 which makes it soluble in various common organic solvents. It has been used extensively in 99 membrane preparation and characterization under the names PEEK-WC, PEEKWC or PEK-C [19]. 100 Membranes have been prepared in the form of hollow fibres for liquid filtration [20] and as flat films 101 for liquid filtrations [21] or for gas separation [22–24] and numerous other applications [19]. In this 102 work, we take advantage of its good solubility in chloroform for the preparation of mixed matrix 103 membranes by the solvent evaporation method. PEEK-WC presents a high selectivity but a too low 104 permeability to be interesting as a material for industrial application. For this reason, different 105 researchers have tried to improve its permeability without loss in selectivity by means of nanoparticle 106 addition. An extensive protocol to obtain MMMs based on PEEK-WC, embedding NaA (LTA) zeolite, 107 was developed in order to improve gas separation parameters [25]. However, no one procedure have 108 enhanced gas transport properties of neat polymer and a decrease in selectivity was observed 109 probably due to a presence of defects at interface between polymer and zeolite. Some methods have 110 been developed in order to further improve the adhesion of PEEK-WC for 3D-mesoporous 111 nanoparticles. For example, the MIL-101 was functionalized with sulfonic acid group for further 112 increase the affinity for polymer matrix. The resulted membranes have shown an improved CO₂ 113 permeability and CO₂/gas selectivity. The increased selectivity was mainly attributed to the increased 114 polar interaction between CO₂ and sulfonic acid groups as well as the good filler–polymer interface 115 compatibility [26].

116



Pebax®1657, x=0.6, y=0.4, density = 1.14 g cm⁻³ (a)



PEEK-WC, density = 1.25 g cm⁻³ (**b**) **CuNi-MOF** structure



Density = 0.54 g cm^{-3} (c)

Figure 1. Chemical structure and density of (**a**) Pebax®1657, (**b**) PEEK-WC; and (**c**) CuNi-MOF. Cu and Ni atoms from the network are represented by cyan and green polyhedra, respectively, whereas organic ligands are depicted as sticks. Green spheres represent Ni²⁺ metal ions in pores.

117

Pebax®1657 is a thermoplastic elastomer multiblock-copolymer containing linear chains of rigid polyamide segments interspaced with flexible polyether segments. The hard polyamide (PA) blocks supply mechanical strength and the presence of the polar ethylene oxide (PEO) group, increases the

121 affinity for CO₂, allowing a good CO₂/non-polar (H₂, N₂, and CH₄) species separation and

122 permselectivity. Several membranes have been prepared in the form of composite hollow fibre 123 membranes [27–29], hybrid and mixed matrix membranes for gas separation [30,31,40,32–39]. Mixed 124 matrix membranes based on amine-functionalized and pristine MIL-53(Al) in Pebax® MH-1657 have 125 shown an increased CO₂ separation and the higher permeability and selectivity (PCO₂ 100 Barrer and 126 α CO₂/CH₄=17; CO₂/N₂=50), was attributed to high porosity introduced by the presence of the MOF, 127 as well as the selective adsorption of CO₂ inside the MOF [37]. Composite hollow fiber membranes 128 with a selective layer of Pebax®1657 and different functionalized Uio-66 MOF have demonstrated a 129 simultaneous improvement in gas permeance and selectivity. The enhanced selectivity was 130 attributed to an increased rigidity of polymer matrix due to formation of hydrogen bond between 131 MOF and Pebax®1657 polymer chains [41]. Pebax®1657 containing Cu₃BTC₂-MOF have shown an 132 increased CO₂/CH₄ ideal selectivity about 15% compared to the neat Pebax®1657 [42]. The 133 improvement in selectivity was attributed to the improvement in the CO₂ solubility, which having a 134 strong quadrupole moment presents higher affinity with unsaturated Cu sites than CH₄, leading to a 135 higher CO₂ permeability. In this work, the CuNi-MOF was dispersed in the Pebax®1657 rubbery 136 polymer and in the PEEK-WC glassy polymer, which presenting Cu site and well-defined cages (1.5 137 nm and 2.2 nm), is expected to influence at the same time solubility and diffusion coefficient. The aim 138 is to understand the main factors that determine the behavior when a MOF is dispersed within a 139 glassy or rubbery polymeric matrix. FT-IR, EDX, DSC and SEM analysis, investigate the potential 140 chemical interaction between CuNi-MOF and polymers. Besides pure gas permeability tests with six 141 different gases were carried out to obtain a general understanding of the MMM performance, while 142 permeation tests were also carried out with binary CO₂/N₂ and CH₄/CO₂ gas mixtures to evaluate its 143 potential performance in a real separation process. These two gas pairs simulate flue gas in view of 144 potential use of the membranes in CO_2 capture, and biogas in view of the potential exploitation of 145 renewable energy in a strongly emerging market.

146 2. Materials and Methods

Pebax®1657, a poly(ethylene-oxide) (PEO) and poly[imino(1-oxohexamethylene)] (PA6) multiblock co-polymer in the molar ratio 60/40, was kindly provided by Arkema in the form of pellets.

PEEK-WC was provided as a powder by the Institute of Applied Chemistry, Changchun, China,and was used as received, without further purification.

151 The CuNi-MOF was obtained as crystalline phase through a double cation-exchange reaction in 152 the solid state by immersing crystals of $Mg^{II_2}{Mg^{II_4}[Cu^{II_2-}(Me_3mpba)_2]_3}\cdot45H_2O$ in saturated aqueous 153 solutions of Ni(NO₃)₂·6H₂O for several weeks [43]. Alternatively, a large scale synthesis of 154 Ni₂^{II}{Ni^{II_4}[Cu^{II_2}(Me_3mpba)_2]_3}·54H_2O (2) was carried out by direct reaction of two aqueous solutions 155 of Na₄[Cu₂(Me_3mpba)_2]·4H₂O (0.1 mol) and Ni(NO₃)₂·6H₂O (0.13 mol) in water and subsequent 156 addition, after filtration and re–suspension in water of the resulting compound, of 0.067 mol of 157 Ni(NO₃)₂·6H₂O (Yield 99 %) [17].

158 2.1. Membranes preparation

159 Mixed matrix membranes were prepared loading different concentrations of CuNi-MOF (9 wt%, 160 17 wt%, 23 wt%) in Pebax®1657 and in PEEK-WC. The Pebax®1657 solution at 10 wt% was prepared 161 by swelling polymer pellets in a mixture of distilled water and ethanol (ratio 30:70 wt/wt) at room 162 temperature overnight. Then a homogeneous dope solution was obtained by heating to 80 °C under 163 magnetic stirring for at least 10 min. The CuNi-MOFs were dispersed and sonicated in water-ethanol 164 mixture for 30 min and subsequently added to the Pebax®1657 solution under magnetic stirring. 165 While, homogenous solution at 5 wt% of PEEK-WC was obtained by dissolving the polymer powder 166 in chloroform at room temperature for 24 h. Concurrent the CuNi-MOFs were separately dispersed 167 in chloroform and sonicated in an ultrasonic bath for 30 min before filtering the PEEK-WC solution 168 through glass wool into the suspension. One hour of mechanical stirring and 30 minutes of sonication 169 make homogeneous each solution. The resulting solutions were cast into levelled flat Teflon petri-170 dish and left to evaporate over 2-3 days. The MMMs produced were dried in oven at 70°C under 171 vacuum condition for 24 h.

172 2.2. Membranes characterization

173 2.2.1 Structural characterization

174 Chemical and morphological analysis of membranes were performed by scanning electron 175 microscopy (SEM) (Phenom Pro X desktop SEM, Phenom-World. The samples for cross-section SEM 176 characterization were prepared by freeze-fracturing in liquid nitrogen. Samples were analyzed 177 without sputter coating with gold. Elemental analysis was performed with the Phenom- Pro X 178 desktop SEM and which is equipped with an energy dispersive X-ray spectroscopy detector (EDX).

- 179 Infrared spectroscopy (FT-IR) analyses were performed by means of Spectrum Spotlight Chemical
- 180 Imaging Instrument (PerkinElmer) in ATR mode.

181 2.2.2. Thermal and mechanical characterization

182 DSC analysis was carried out using a Pyris Diamond Differential Scanning Calorimeter (Perkin 183 Elmer, USA) equipped with an intracooler refrigeration system. Samples of 15-20 mg. Unless 184 specified otherwise, three cycles were performed. The PEEK-WC samples were first heated from -60 185 °C to 300 °C, kept at this temperature for 1 min, and cooled down to 0 °C where they were kept for 5 186 min. In the second run, the samples were heated up again to 300 °C. The DSC runs were performed 187 at a scan rate of 15 °C min⁻¹. Before the measurements, the samples were kept at 50°C under vacuum 188 for one night in order to remove possibly adsorbed water. The Pebax®1657-based samples were tested 189 by an identical method in the range from $-40^{\circ}C \rightarrow 220^{\circ}C \rightarrow -75^{\circ}C \rightarrow 225^{\circ}C$.

190 Tensile tests were carried out at room temperature on a single column Universal Testing 191 Machine, Zwick/Roell model Z2.5, equipped with a 50 N load cell and pneumatic clamps. Surface of 192 one flat clamp was covered with adhesive rubber to avoid slipping or damage of the samples, while 193 second clamp had convex surface to increase the local pressure and to avoid the extraction of the 194 sample. The average value and the standard deviation of the Young's modulus, the tensile strength 195 and the maximum deformation were determined on a series of at least four samples. The sample 196 width was 5 mm, the grip-to-grip distance 40 and 30 mm for Pebax®16 and PEEK-WC, respectively. 197 The test were carried out at a deformation rate 20 mm min⁻¹ (200 % min⁻¹) for Pebax®1657-based 198 samples and 6 mm min⁻¹ (20 % min⁻¹) for PEEK-WC-based samples.

199 2.2.3 Single gas permeation method

Single gas permeation tests were carried out at 25 °C and at a feed pressure of 1 bar, using a fixed-volume pressure increase instrument (ESSR), described elsewhere [44]. Permeability coefficients, *P*, and diffusion coefficients, *D*, were determined by the time-lag method [45]. The permeability coefficient, *P*, is calculated from the permeation curve in steady state described by:

204
$$P_t = P_0 + (dp/dt)_0 \cdot t + \frac{RT \cdot A}{V_P \cdot V_m} \cdot \frac{p_f \cdot p}{\ell} \left(t - \frac{\ell^2}{6D} \right)$$
(1)

205 the last term in (Eq. 1) represents the so-called permeation time lag, Θ , which is inversely 206 proportional to the diffusion coefficient of the gas:

$$207 \qquad \qquad \Theta = \frac{\ell^2}{6D} \tag{2}$$

the approximate gas solubility coefficient, S, was obtained indirectly as the ratio of the permeability to the diffusion coefficient by assuming the solution-diffusion transport mechanism:

$$S = P/D \tag{3}$$

211 Permeabilities are reported in Barrer [1 Barrer=10⁻¹⁰ cm³ (STP) cm cm⁻² s⁻¹ cm Hg⁻¹].

212 2.2.3 Mixed gas permeation measurement

213 Mixed gas permeation tests were carried out using a custom made constant pressure/variable 214 volume instrument, described in detail elsewhere [46], equipped with a quadrupole mass filter (HPR-

- 215 20 QIC Benchtop residual gas analysis system, Hiden Analytical). Measurements were performed
- from 1 to 6 bar(a) with two binary gas mixtures of CO_2/N_2 (15:85 vol%) and CO_2/CH_4 (35:65 vol%),
- simulating flue gas and biogas, respectively. Argon was used as a sweeping gas and as the internal
- 218 standard for the calculation of the permeate gas composition.

219 3. Results and discussion

220 3.1. Chemical and morphological characterization

221 The MOF loading in polymer matrix, was evident by distinctive colour of obtained membranes, 222 and it was accurately confirmed by the EDX analysis, which reveals the attendance of Cu and Ni 223 metals in both Pebax®1657 and PEEK-WC polymers (See SI, Figure S1-a and b, respectively). The 224 chemical characterization of neat membranes and MMMs, as well the interaction between polymer 225 chains and nanoparticles were investigated by FTIR-ATR. Figure 2b and Figure 2d shows the 226 spectrum for neat Pebax®1657 and for Pebax®1657/CuNi-MOF membranes. The functional groups 227 of Pebax®1657 give characteristic peaks at 3297 cm⁻¹ for N-H group, at 2943 and 2859 cm⁻¹ attributed 228 to the asymmetric and symmetric stretching of the C-H bound, 1731 cm⁻¹ for the symmetric stretches 229 of carboxylate R-O-C=O, while asymmetric R-O-C=O at ca. 1430, 1636 cm⁻¹ for H-N–C=O and 1099 230 cm⁻¹ for -C-O-C.

231







Figure 2. SEM image of the samples showing the interface between MOF and polymer phase (**a**) for Pebax®1657 and (**c**) PEEK-WC. FT-IR of neat polymer and MMMs with 23 wt% of CuNi, for Pebax®1657 (**b**) and PEEK-WC (**d**).

233 In the spectrum of Pebax®1657/CuNi-MOF characteristic peaks appear at 1578 cm⁻¹ due to C–C 234 aromatic band of MOF's phenylene, at 1608 cm⁻¹ for stretching of oxamate ligand of MOF and a peak 235 at 1330 cm⁻¹, which could be considered diagnostic of the presence of bridging oxamate between Cu 236 and Ni [47]. Finally, the characteristic peaks between 2900 cm⁻¹ and 2800 cm⁻¹ due to the symmetric 237 and asymmetric C-H stretching are slight shifted to lower wavelengths confirming the interaction of 238 hydrogen bonds at the interface between MOF and Pebax®1657, as it was also see by Khosravi et al. 239 for Pebax1657/CuBTC mixed matrix membranes [48]. Figure 2d offers a comparison of PEEK-WC 240 membrane and PEEK-WC/CuNi spectrum. The functional groups of PEEK-WC give characteristic 241 peaks at 3052 cm⁻¹ and 3075 cm⁻¹ for aromatic C–H stretch, at 1767 cm⁻¹ for ketonic and esteric C=O 242 stretching, and at 1589 cm⁻¹ for C–C aromatic band [49]. In PEEK-WC/CuNi the peak for C-C aromatic 243 band is shift on the lower wavelengths probably due to the $\pi \cdots \pi$ stacking interaction between 244 benzene rings of polymers and MOF. In fact, when the ring is very conjugated, a weak band can be 245 observed at around 1580 cm⁻¹, such as it is visible from the PEEK-WC/CuNi spectrum. While, the 246 peaks due to the stretch of aromatic C-H appear to be totally fused with the water peak (3370 cm⁻¹) 247 that reveals the relatively hydroscopic nature of the MOFs. The interaction between the Pebax®1657 248 and CuNi-MOF was also confirmed by SEM analysis, and in the enlarged particular of Figure 2a it is 249 possible to see that polymer phase of Pebax®1657creates a sort of circular compressed polymer region 250 around the MOF, which Koros defined as "Case I matrix rigidification". On the other hand, the 251 distribution of MOF in the PEEK-WC polymer produces a defined and regular network structure 252 (Figure 2c) with a homogenous dispersion of nanoparticles without significant sedimentation across 253 the membranes (See SI Fig. S-2). Instead, the addition of MOFs in Pebax®1657 shows no significant 254 agglomeration at lowest 9 wt% that becomes more significant at highest 29 wt% concentration.

255 3.2 Mechanical and thermal properties

256 All membranes are dense mixed matrix membranes with good mechanical resistance for 257 handling. The results of the tensile tests, performed on the nanocomposite films obtained on the series 258 with different concentration of CuNi-MOFs in Pebax®1657 and in PEEK-WC are shown in Figure 3. 259 The PEEK-WC presents a Young's module about 1.14 GPa [50] and Pebax®1657 has a Young's module 260 about 0.10 GPa [51]. In both polymers, the loading of CuNi-MOF increases the Young's modulus. The 261 increase of Young's modulus indicates an increase of stiffness for both polymers increasing the MOF 262 concentrations. The glassy PEEK-WC presents a higher modulus compared to the rubbery 263 Pebax®1657 as expected and it remains higher even when MOF concentration was increased. On the 264 other hand, the maximum deformation is higher in the more flexible MMMs based on the rubbery 265 Pebax®1657 compared to the MMMs based on the rigid glassy PEEK-WC.

266



Figure 3. Young's modulus (a), tensile strength (b) and maximum deformation (c) as a function of the CuNi-MOF concentration in Pebax[®]1657 and PEEK-WC. Values of the neat PEEK-WC [50] and Pebax [51] samples from the literature. Trend lines are given as a guide to the eye.

267

268 The break strength decreases for Pebax®1657/CuNi membranes indicating that the membranes 269 become weaker compared to the neat polymer. On the contrary, the break strength increases for 270 PEEK-WC/CuNi membranes, suggesting a good adhesion between MOF and polymer that makes the 271 MMMs stronger than neat PEEK-WC membrane. The thermal properties of the membranes were 272 studied by DSC analysis and the results are displayed in Figure 4. For neat Pebax®1657 the two 273 dominant endothermal peaks at 19 °C and 200 °C are attributed to the fusion of the crystalline fraction 274 of the soft poly(ethylene oxide) (PEO) blocks and the hard polyamide (PA) blocks, respectively. In 275 the presence of MOF, the melting enthalpy of both PEO and PA decrease. However, the peak 276 maximum shifts to higher temperature, indicating that the MOF stabilizes the crystalline PEO and 277 that it mainly interacts with this phase. On the other hand, no significant changes occur in the glass

278 transition temperature of PEEK-WC at 230 °C as function of the MOF concentration.



Figure 4. DSC thermograms for membrane of (**a**) neat Pebax1657 and (**b**) neat PEEK-WC and MMMs with different concentration of CuNi-MOF during the second eating.

279 3.3 Pure gas transport properties

280 Single gas permeation experiments were carried out in the order H2, O2, N2, CH4 and CO2 at 25°C 281 but repeated experiments with N_2 and O_2 at the end of the cycle were identical and revealed no 282 structural changes in the material. An overview of the results of the Pebax®1657-based MMMs and 283 PEEK-WC based MMMs with different filler loadings is given in Figure 5 and Figure 6, respectively. 284 All permeability, diffusivity and solubility data, and their respective selectivities, are given in the 285 supporting information (See SI Table 2 and 3). The dispersion of CuNi-MOF does not produce a 286 substantial change in permeability of the rubbery Pebax[®]1657. The most permeable species is CO₂, 287 confirming a solubility controlled transport (CO₂ > H₂ > CH₄ > O₂ > N₂), typical for the rubbers (Figure 288 5a). A remarkably strong decrease in the effective diffusion coefficients of all gases as function of the 289 MOF loading (Figure 5b), is balanced by a proportional increase in their solubility coefficients (Figure 290 5c). This results in a similar permeability and selectivity (Figure 5d) for MMMs as in the neat 291 Pebax®1657, with the only exception of the H₂/CH₄ gas pair, which suggests a favourable effect of 292 the MOF on the size-selectivity. This is confirmed by the slight increase in the diffusion selectivities 293 (Figure 5e), apparently because the MOF's provide preferential diffusion paths for smaller gases.

294 While the gas solubility upon addition of the MOF to Pebax increases as expected, given the 295 generally high sorption capacity of MOFs for gases, it was not expected that this increase was more 296 or less similar for all gases, and it was even less expected the porous fillers would effectively lead to 297 a decrease in the diffusion coefficient. The explanation is that the filler does not effectively decrease 298 the diffusion coefficient in the membrane, but the highly sorbing MOF transforms the permeation 299 experiment in a sort of breakthrough experiment. The reason for the apparently slower diffusion is 300 the accumulation of the gas inside the pores of the filler, similar to the phenomenon of immobilizing 301 adsorption [52,53], but in this case it has virtually negligible net effect on the permeability.



Figure 5. (a) Permeability, (b) diffusivity and (c) solubility coefficients, and their respective selectivity towards N_2 (d, e, f) for each gas as functions of the weight percentage of CuNi-MOF in Pebax[®]1657.

303

304 In contrast to the effect in Pebax®1657, CuNi-MOF causes a drastic increase in the gas 305 permeability for all gases as function of MOFs loading in the glassy PEEK-WC, and the order of 306 permeation obeys that of the diffusion controlled gas transport mechanism with $H_2 > CO_2 > O_2 > N_2$ 307 > CH₄, typical for glassy polymers (Figure 6a). The increase in permeability is a direct consequence 308 of an increase in diffusion coefficient (Figure 6b), whereas there is no substantial change in solubility 309 coefficient (Figure 6c). The enhanced diffusion clearly indicates transport within the pore structure 310 of CuNi-MOFs, which increases the total free volume of MMMs promoting the gas diffusion for all 311 gases. The presence of MOF with chemical groups having a high affinity for CO2, increase its 312 solubility, its CO₂/CH₄ and CO₂/N₂ solubility selectivity (Figure 6f), and ideal selectivity for these two 313 gas pairs (Figure 6d). Instead, the diffusion selectivity is not strongly affected by CuNi-MOF. 314



Figure 6. (a) Permeability, (b) diffusivity, (c) solubility coefficients and respective N₂ selectivity (d, e, f) for each gas as functions of weight percentage of MOF loaded in PEEK-WC.

315

316 For a better understanding of the transport phenomena of gas separation membranes, the 317 diffusion and solubility coefficients are often correlated with the molecular properties of gases [54]. 318 Both neat polymers show a linear dependency of the diffusivity (D) on the squared gas diameter (d^{2}_{eff}) 319 (Figure 7a and b), suggesting that the gas transport in neat polymers follows the diffusion solution 320 model. In the presence of the MOF, both polymers show non-linearity of the D vs d^2_{eff} correlation 321 (Figure 7), which becomes more evident at higher MOF loading. A similar trend was recently 322 observed for PIMs [55], where it was attributed to a different size-selectivity for small and large gas 323 species due to the highly interconnected free volume. These results suggest that the CuNi-MOF 324 particles with their internal void structure have a similar effect on the gas diffusion of Pebax®1657 325 and PEEK-WC, introducing preferential diffusion pathways into the system, which change the main 326 transport mechanism especially for the small molecules. As described above, the remarkably lower 327 effective diffusion in Pebax®1657 is due to the much higher sorption capacity of the MOFs, than of 328 the neat Pebax®1657 [39]. Only for the CO₂/N₂ gas pair, there is a remarkable inversion from 329 preferential N₂ diffusion in the neat pebax, to preferential CO₂ diffusion for the sample with the 330 highest MOF concentration. Instead, in the glassy PEEK-WC, with much lower intrinsic diffusion 331 coefficients in the neat polymer, the MOF has a strong positive effect on the total permeability of the 332 mixed matrix membrane due to the far more rapid diffusion through the cages of the MOFs. In PEEK-333 WC, after an initial decrease compared to the neat polymer, the solubility of all gases increases with 334 the CuNi-MOF concentration. This is similar to what was observed in Pebax, but only in PEEK-WC 335 this leads to an increasing permeability, due to the positive effect on the diffusivity as well. 336



Figure 7. Correlation of the effective diffusion coefficient as a function of the molecular diameter of six light gases in **(a)** Pebax®1657/CuNi MMMs and **(b)** PEEK-WC/CuNi MMMs. Correlation of the effective solubility coefficient of the gases as a function of their critical temperature in **(c)** Pebax®1657/CuNi MMMs and **(d)** PEEK-WC/CuNi MMMs. The legend is identical in all individual graphs.

337 3.4 Mixed gas transport properties

338 Mixed gas permeation measurements were carried out on the representative membranes with 339 the highest MOF concentration in each polymer in view of two relevant industrial gas separations, 340 namely biogas upgrading and CO₂ capture from flue gas, involving CO₂ separation from CH₄ and 341 N₂, respectively (Figure 8). For this propose, measurements were performed from 1 to 6 bar(a) with 342 simulated flue gas (CO₂/N₂, 15/85 vol%) and simulated biogas (CO₂/CH₄, 35/65 vol%). The glassy 343 PEEK-WC MMM exhibits typical dual mode behaviour with the CO₂/CH₄ mixture, showing a 344 decrease of CO₂ permeability as a function of the increased feed pressure, which causes a 345 simultaneous and slightly smaller decrease of CO₂/CH₄ selectivity. This indicates that the free volume 346 in the PEEK-WC phase and in the MOFs' pores is gradually becoming occupied by CO₂, and as a 347 result, the CH4 permeability slightly decreases as a function of pressure. Moreover, some hysteresis 348 of CO₂ and CH₄ permeability is observed when reducing the pressure after the initial pressure 349 increase steps. This is ascribed to a slight CO2-induced dilation of the polymer, leaving a higher free

volume when the pressure is decreased, and thus a higher CH_4 permeability and a slightly lower selectivity. In the separation of the CO_2/N_2 mixture no further hysteresis takes place because of the lower CO_2 partial pressure, for the same reason the CO_2 permeability is slightly higher than in the biogas mixture.

Contrary to the glassy PEEK-WC, the rubbery Pebax®1657/CuNi-MOF (23 wt%) membrane shows essentially pressure-independent permeability and selectivity (Figure 8a), after only a slight increase in permeability with pressure in the first run, apparently due to a certain conditioning of the

- 357 sample. The transport properties of the Pebax-based membrane are dominated by the rubbery phase,
- 358 and the dispersed MOFs only provide a higher gas sorption capacity in the membrane but do not 359 significantly affect the overall performance.
- 360



Figure 8. Pressure dependence of CO₂ and CH₄ permeabilities and CO₂/CH₄ selectivity using the binary mixture of CO₂/CH₄ (35:65 vol%) for Pebax®1657_23 wt% CuNi-MOF (**a**) and PEEK-WC_23% CuNi-MOF (**b**). Pressure dependence of CO₂ and N₂ permeabilities and CO₂/N₂ selectivity in binary mixture conditions for CO₂/N₂ (15:85 vol%) of Pebax®1657_23 wt% CuNi (**c**) and PEEK-WC_23% CuNi (**d**). Closed symbols for stepwise increase of the pressure and open symbols for the subsequent stepwise decrease of the pressure. Trend lines are plotted as a guide to the eye.

362 3.5 Robeson's plots and performance overview

The gas transport properties for all membranes are summarized in the Robeson diagrams in Figure 9 for four gas pairs representing industrially important separations: CO₂/N₂, CO₂/CH₄, O₂/N₂ and H₂/N₂.



Figure 9. Robeson's plots for CO₂/N₂ (**a**) CO₂/CH₄ (**b**) O₂/N₂ (**c**) H₂/N₂ (**d**) showing the data of Pebax1657/CuNi (blue) and PEEK-WC/CuNi at different MOF loadings of 0 wt% (= neat polymers, •, •), 9 wt% (\blacktriangle , \bigstar),17 wt% (\blacksquare , \blacksquare) and 23 wt% (•, •). Blue line 1991 upper bound, red line 2008 upper bound [8,9]; yellow line 2015 [56]; purple line 2019 upper bound [57]. The arrows qualitatively indicate the direction of increasing MOF loading. The clouds of blue "X" and red "X" symbols represent the mixed gas data in the pressure range of 1-6 bar for Pebax1657/CuNi-MOF at 29 wt% and PEEK-WC/CuNi-MOF at 23 wt% of MOF loading. Please refer to the electronic version for colour figures.

366 For all gases except H₂, the neat rubbery Pebax®1657 is more permeable than neat glassy PEEK-

367 WC. On the other hand, PEEK-WC is more selective than Pebax®1657 for all gas pairs, except for the

368 CO₂/N₂ separation (Figure 9a). For the CO₂/N₂ gas pair, with nearly identical diameter of the

369 molecules, the selectivity is almost entirely due to solubility selectivity, which is higher in Pebax then

370 in PEEK-WC. However, whereas CuNi-MOF in Pebax®1657 has a marginal effect, CuNi-MOF in the

glassy PEEK-WC, increases drastically not only the CO₂/N₂ selectivity (from 25 to 37), but also the
CO₂ permeability (from 6 to ~50 Barrer), near the CO₂ permeability of the rubbery Pebax®1657 (PCO₂
~69 Barrer) (Figure 9a).

For CO_2/CH_4 separation, the PEEK-WC-based MMMs show far superior properties than the Pebax®-based membranes, which hardly change with the MOF concentration. A strong contribution of size selectivity between CO_2 and the much bulkier CH_4 molecules (Figure 7b) and a further increase of both solubility and diffusivity with increasing MOF concentration leads to enhanced selectivity (from 30 to ~ 50, above that of Pebax) and CO_2 permeability (from 6 to ~50 Barrer), exceeding the 1991 Robeson upper bound. For both mixtures, the performance is very similar to the ideal selectivity and pure gas permeability, indicating the absence of significant coupling effect (Figure 9b).

The O₂/N₂ separation (Figure 9c) is mainly due to diffusion selectivity. Neither the permeability, nor the selectivity changes much in Pebax®1657 upon addition of CuNi-MOF, but in PEEK-WC the permeability of both gases increases 5-fold in the presence of 23 wt% CuNi-MOF, maintaining an almost constant O₂/N₂ selectivity.

Finally, the trend for H₂/N₂ separation is similar to that of O₂/N₂, but in this case for both polymers the MOFs also have a slightly higher H₂/N₂ selectivity than the neat polymers, as a result of increased diffusion selectivity. The H₂ permeability in Pebax® 1657 slightly increases from 8 to 11 Barrer, and the H₂/N₂ selectivity and permeability in the PEEK-WC/CuNi-MOF MMMs both strongly increase, surpassing the 1991 upper bound.

390 4. Conclusions

391 Development of novel, well performing gas separation membranes requires good 392 understanding of their transport properties. This is even more important for complex systems such 393 as mixed matrix membranes in which a porous filler is dispersed in the polymer matrix. Therefore, 394 this work describes the comparison of the performance of two sets of membranes, glassy PEEK-WC 395 membranes and rubbery Pebax®1657 membranes, with different concentrations of an oxamate-based 396 CuNi-MOF. An increase of the Young's modulus for both membrane sets confirms that the MOF 397 increases the stiffness of the polymer. In the case of the Pebax®1657 membrane, this is also 398 accompanied by a slight shift of the melting peak of the PEO phase to higher temperatures, which 399 suggests a good interaction of the MOF with the PEO phase. On the other hand, the presence of the 400 MOF reduces the melting enthalpy and thus the overall crystallinity of both the PEO phase and the 401 PA phase of Pebax®, whereas it does not affect the glass transition temperature of PEEK-WC. Only 402 for PEEK-WC, along with Young's modulus also the tensile strength increases, but for both polymers 403 the maximum deformation decreases with the addition of the MOF.

404 In terms of transport properties, the permeability of PEEK-WC strongly increases with 405 increasing MOF content, mostly due to an increase in the effective diffusion coefficient, whereas 406 unexpectedly, the effective diffusion coefficient in Pebax®1657 drastically decreases upon addition 407 of the MOF. This is almost completely compensated by an increase in solubility, so that the 408 permeability remains nearly constant. The increase in diffusivity favours especially the smaller gas 409 species, and as a result, the PEEK-WC MMMs show a simultaneous increase in CO₂ permeability and 410 CO₂/CH₄ selectivity, and the membranes with the highest MOF loading approach Robeson's 2008 411 upper bounds for these gas pairs. In the range of 1-6 bar(a), the mixed gas permeation tests with 412 CO₂/N₂ 15/85 vol% and CO₂/CH₄ 35/65 vol% mixtures show only a weak pressure-dependence for 413 the CO₂/CH₄ mixture in PEEK-WC with 30% MOF, typical for materials with dual mode sorption 414 behavior, but not for the CO₂/N₂ mixture in PEEK-WC and for both mixtures in the Pebax-based 415 MMMs.

Summarizing, the detailed analysis of the gas transport properties of the two series of MMMs highlights the enormous impact of the polymer matrix on the effectiveness of the same MOF when it is embedded in a rubbery or a glassy polymer. Even if the MOF improves significantly the gas solubility, this may not have a positive effect on the permeability if the diffusivity is not increased simultaneously. Successful development of better-performing MMMs can therefore not rely on studies of only the overall permeability, but necessitates knowledge of the individual solubility and diffusion coefficient. For both parameters, the values of the polymer and the porous filler should becarefully matched to yield optimum performance.

424

425 Acknowledgments: Phenom-World B.V., Eindhoven (NL), is gratefully acknowledged for providing a 426 Phenom Pro X desktop SEM for evaluation. R.B. thanks the MIUR (project PON R&I FSE-FESR 2014-2020) for 427 predoctoral grants.

428

429 Supplementary Materials: The following are available online at: , Table SI-1. List of MMMs prepared. Fig. S-1 430 EDX of Pebax®1657/CuNi-MOF (a) and PEEK-WC/CuNi-MOF (b) at an accelerating voltage of 15Kv. Fig. S-2 431 SEM images of cross section for MMMs of Pebax®1657/CuNi-MOF and PEEK-WC/CuNi-MOF. Table S-2 Pure 432 gas permeability, solubility and diffusion coefficients, and respective selectivity for neat Pebax1657 and 433 Pebax1657/CuNi MMMs. Table S-3 Pure gas permeability, solubility and diffusion coefficients, and respective 434 selectivity for neat PEEK-WC and PEEK-WC/CuNi MMMs. Table S-4 Mixed gas permeabilities and selectivities 435 of PEEK-WC/CuNi 23 wt% membrane using binary mixture CO2/CH4 (35/65) at pressure of 1-6 bar. Table S-5 436 Mixed gas permeabilities and selectivities of PEEK-WC/CuNi 23 wt% membrane using binary mixture CO2/N2 437 (15/85) at pressure of 1-6 bar.

- 438 Author Contributions: E.E. and R. B. conceived, designed and performed the membrane preparation and 439 characterization experiments under the supervision of J.C.J and D.A.; J.F. S and E.P synthetized and provided 440 CuNi-MOF; E.E. and A. F. performed the single gas permeation and DSC experiments; E. E. and J.C.J. performed 441 mechanical tests. M.M. performed the mixed gas permeation experiments under the supervision of J.C.J. All 442 authors analyzed the data and wrote the paper with similar effort.
- 443

Funding: This work was supported by Ministero dell'Istruzione, dell'Università e della Ricerca (Italy).
Phenom-World B.V., Eindhoven (NL), is gratefully acknowledged for providing a Phenom Pro X desktop SEM for evaluation. Dr. C. Cantoni (Arkema Italy) is gratefully acknowledged for providing the Pebax®1657 pellets.

447 **Conflicts of Interest:** The authors declare no conflict of interest.

448 References

- 449
- 450 1. Ockwig, N.W.; Nenoff, T.M. Membranes for hydrogen separation. *Chem. Rev.* 2007.
- 451 2. Himma, N.F.; Wardani, A.K.; Prasetya, N.; Aryanti, P.T.P.; Wenten, I.G. Recent progress and challenges
 452 in membrane-based O2/N2 separation. *Rev. Chem. Eng.* 2019.
- 453 3. Coombe, H.S.; Nieh, S. Polymer membrane air separation performance for portable oxygen enriched
 454 combustion applications. *Energy Convers. Manag.* 2007.
- 4. Chen, X.Y.; Vinh-Thang, H.; Ramirez, A.A.; Rodrigue, D.; Kaliaguine, S. Membrane gas separation
 456 technologies for biogas upgrading. *RSC Adv.* 2015.
- 457 5. Esposito, E.; Dellamuzia, L.; Moretti, U.; Fuoco, A.; Giorno, L.; Jansen, J.C. Simultaneous production of
 458 biomethane and food grade CO 2 from biogas: an industrial case study. *Energy Environ. Sci.* 2019, *12*,
 459 281–289.
- Galizia, M.; Chi, W.S.; Smith, Z.P.; Merkel, T.C.; Baker, R.W.; Freeman, B.D. 50th Anniversary
 Perspective: Polymers and Mixed Matrix Membranes for Gas and Vapor Separation: A Review and
 Prospective Opportunities. *Macromolecules* 2017, *50*, 7809–7843.
- 463 7. Brunetti, A.; Scura, F.; Barbieri, G.; Drioli, E. Membrane technologies for CO2 separation. *J. Memb. Sci.*464 2010.
- 465 8. Robeson, L.M. The upper bound revisited. J. Memb. Sci. 2008, 320, 390–400.
- 466 9. Robeson, L.M. Correlation of separation factor versus permeability for polymeric membranes. *J. Memb.*467 *Sci.* 1991, *62*, 165–185.
- 468 10. Chuah, C.Y.; Goh, K.; Yang, Y.; Gong, H.; Li, W.; Karahan, H.E.; Guiver, M.D.; Wang, R.; Bae, T.-H.

469		Harnessing Filler Materials for Enhancing Biogas Separation Membranes. Chem. Rev. 2018, 118, 8655–
4/0		8769.
4/1 472	11.	Zhu, H.; Liu, D. The synthetic strategies of metal–organic framework membranes, films and 2D MOFs and their applications in devices <i>L Mater Chem</i> A 2019
473	12	Najari S. Saeidi S. Gallucci F. Drioli F. Mixed matrix membranes for hydrocarbons separation and
474	12.	recovery: a critical review. Rev. Chem. Eng. 2019 0
л75	12	Processing N. Himme, N.E. Deddy, Sutriano, D. Monton, L. Ladowig, B.D. A. Boviow, on Empiring
476	15.	Organic containing Microporous Material Mombranes for Carbon Capture and Separation Cham Eng
477		J. 2019, 123575.
478	14.	Ebadi Amooghin, A.; Mashhadikhan, S.; Sanaeepur, H.; Moghadassi, A.; Matsuura, T.; Ramakrishna, S.
479		Substantial breakthroughs on function-led design of advanced materials used in mixed matrix
480		membranes (MMMs): A new horizon for efficient CO2 separation. Prog. Mater. Sci. 2019, 102, 222–295.
481	15.	Review, A. Performance of Mixed Matrix Membranes Containing Three-Dimensional (3D) Fillers for
482		CO 2 Separation : 2018.
483	16.	Cohen, S.M. The Postsynthetic Renaissance in Porous Solids. J. Am. Chem. Soc. 2017 , 139, 2855–2863.
484	17.	Fortea-Pérez, F.R.; Mon, M.; Ferrando-Soria, J.; Boronat, M.; Leyva-Pérez, A.; Corma, A.; Herrera, J.M.;
485		Osadchii, D.; Gascon, J.; Armentano, D.; et al. The MOF-driven synthesis of supported palladium
486		clusters with catalytic activity for carbene-mediated chemistry. <i>Nat. Mater.</i> 2017 , <i>16</i> , 760–766.
487	18.	Mon, M.; Tiburcio, E.; Ferrando-Soria, J.; Gil San Millán, R.; Navarro, J.A.R.; Armentano, D.; Pardo, E. A
488		post-synthetic approach triggers selective and reversible sulphur dioxide adsorption on a metal-organic
489		framework. <i>Chem. Commun.</i> 2018 , <i>54</i> , 9063–9066.
490	19.	Jansen, J.C.; Drioli, E. Poly(ether ether ketone) derivative membranes—a review of their preparation,
491		properties and potential. <i>Polym. Sci. Ser. A</i> 2009 , <i>51</i> , 1355–1366.
492	20.	Tasselli, F.; Jansen, J.C.; Drioli, E. PEEKWC Ultrafiltration Hollow-Fiber Membranes: Preparation,
493		Morphology, and Transport Properties. J. Appl. Polym. Sci. 2004, 91, 841–853.
494	21.	Buonomenna, M.G.; Figoli, A.; Jansen, J.C.; Drioli, E. Preparation of Asymmetric PEEKWC Flat
495		Membranes with Different Microstructures by Wet Phase Inversion. J. Appl. Polym. Sci. 2004, 92, 576–
496		591.
497	22.	Jansen, J.C.; Buonomenna, M.G.; Figoli, A.; Drioli, E. Asymmetric membranes of modified poly(ether
498		ether ketone) with an ultra-thin skin for gas and vapour separations. J. Memb. Sci. 2006, 272, 188–197.
499	23.	Iulianelli, A.; Algieri, C.; Donato, L.; Garofalo, A.; Galiano, F.; Bagnato, G.; Basile, A.; Figoli, A. New
500		PEEK-WC and PLA membranes for H2 separation. Int. J. Hydrogen Energy 2017, 42, 22138–22148.
501	24.	Iulianelli, A.; Clarizia, G.; Gugliuzza, A.; Ebrasu, D.; Bevilacqua, A.; Trotta, F.; Basile, A. Sulfonation of
502		PEEK-WC polymer via chloro-sulfonic acid for potential PEM fuel cell applications. Int. J. Hydrogen
503		Energy 2010 , 35, 12688–12695.
504	25.	Clarizia, G.; Algieri, C.; Regina, A.; Drioli, E. Zeolite-based composite PEEK-WC membranes: Gas
505		transport and surface properties. <i>Microporous Mesoporous Mater.</i> 2008 , 115, 67–74.
506	26.	Xin, Q.; Liu, T.; Li, Z.; Wang, S.; Li, Y.; Li, Z.; Ouyang, J.; Jiang, Z.; Wu, H. Mixed matrix membranes
507		composed of sulfonated poly(ether ether ketone) and a sulfonated metal-organic framework for gas
508		separation. J. Memb. Sci. 2015, 488, 67–78.
509	27.	- Esposito, E.; Clarizia, G.; Bernardo, P.; Jansen, J.C.; Sedláková, Z.; Izák, P.; Curcio, S.; Cindio, B. de;
510		Tasselli, F. Pebax®/PAN hollow fiber membranes for CO2/CH4 separation. Chem. Eng. Process. Process
511		Intensif. 2015 , 94, 53–61.

512	28.	Kim, K.; Ingole, P.G.; Kim, J.; Lee, H. Separation performance of PEBAX/PEI hollow fiber composite
513		membrane for SO2/CO2/N2 mixed gas. Chem. Eng. J. 2013, 233, 242–250.
514	29.	Asghari, M.; Mosadegh, M.; Riasat Harami, H. Supported PEBA-zeolite 13X nano-composite membranes
515		for gas separation: Preparation, characterization and molecular dynamics simulation. Chem. Eng. Sci.
516		2018 , <i>187</i> , 67–78.
517	30.	Dai, Z.; Bai, L.; Hval, K.N.; Zhang, X.; Zhang, S.; Deng, L. Pebax®/TSIL blend thin film composite
518		membranes for CO2 separation. Sci. China Chem. 2016, 59, 1–9.
519	31.	Ehsani, A.; Pakizeh, M. Synthesis, characterization and gas permeation study of ZIF-11/Pebax® 2533
520		mixed matrix membranes. J. Taiwan Inst. Chem. Eng. 2016.
521	32.	Jomekian, A.; Bazooyar, B.; Behbahani, R.M.; Mohammadi, T.; Kargari, A. Ionic liquid-modified Pebax®
522		1657 membrane filled by ZIF-8 particles for separation of CO2 from CH4, N2 and H2. J. Memb. Sci. 2017,
523		524, 652–662.
524	33.	Liu, G.; Chernikova, V.; Liu, Y.; Zhang, K.; Belmabkhout, Y.; Shekhah, O.; Zhang, C.; Yi, S.; Eddaoudi,
525		M.; Koros, W.J. Mixed matrix formulations with MOF molecular sieving for key energy-intensive
526		separations. Nat. Mater. 2018, 17, 283–289.
527	34.	Jomekian, A.; Behbahani, R.M.; Mohammadi, T.; Kargari, A. High speed spin coating in fabrication of
528		Pebax 1657 based mixed matrix membrane filled with ultra-porous ZIF-8 particles for CO2/CH4
529		separation. <i>Korean J. Chem. Eng.</i> 2016 , 34, 1–14.
530	35.	Jomekian, A.; Behbahani, R.M.; Mohammadi, T.; Kargari, A. CO2/CH4 separation by high performance
531		co-casted ZIF-8/Pebax 1657/PES mixed matrix membrane. J. Nat. Gas Sci. Eng. 2016, 31, 562–574.
532	36.	Atash Jameh, A.; Mohammadi, T.; Bakhtiari, O. Preparation of PEBAX-1074/modified ZIF-8
533		nanoparticles mixed matrix membranes for CO2 removal from natural gas. Sep. Purif. Technol. 2020, 231,
534		115900.
535	37.	Meshkat, S.; Kaliaguine, S.; Rodrigue, D. Mixed matrix membranes based on amine and non-amine MIL-
536		53(Al) in Pebax® MH-1657 for CO2 separation. Sep. Purif. Technol. 2018, 200, 177–190.
537	38.	Noroozi, Z.; Bakhtiari, O. Preparation of Amino Functionalized Titanium Oxide Nanotubes and Their
538		Incorporation within Pebax/PEG Blended Matrix for CO2/CH4 Separation. Chem. Eng. Res. Des. 2019.
539	39.	Mon, M.; Bruno, R.; Tiburcio, E.; Grau-Atienza, A.; Sepúlveda-Escribano, A.; V Ramos-Fernandez, E.;
540		Fuoco, A.; Esposito, E.; Monteleone, M.; Carolus Jansen, J.; et al. Efficient Gas Separation and Transport
541		Mechanism in Rare Hemilabile Metal-Organic Framework. Chem. Mater. 2019, 0, null-null.
542	40.	Song, C.; Li, R.; Fan, Z.; Liu, Q.; Zhang, B.; Kitamura, Y. CO2/N2 separation performance of Pebax/MIL-
543		101 and Pebax /NH2-MIL-101 mixed matrix membranes and intensification via sub-ambient operation.
544		Sep. Purif. Technol. 2020 , 238, 116500.
545	41.	Sutrisna, P.D.; Hou, J.; Zulkifli, M.Y.; Li, H.; Zhang, Y.; Liang, W.; D'Alessandro, D.M.; Chen, V. Surface
546		functionalized UiO-66/Pebax-based ultrathin composite hollow fiber gas separation membranes. J.
547		Mater. Chem. A 2018, 6, 918–931.
548	42.	Kim, J.; Choi, J.; Soo Kang, Y.; Won, J. Matrix effect of mixed-matrix membrane containing
549		CO <inf>2</inf> -selective MOFs. J. Appl. Polym. Sci. 2016, 133, 1–8.
550	43.	Grancha, T.; Ferrando-Soria, J.; Zhou, H.C.; Gascon, J.; Seoane, B.; Pasán, J.; Fabelo, O.; Julve, M.; Pardo,
551		E. Postsynthetic Improvement of the Physical Properties in a Metal-Organic Framework through a Single
552		Crystal to Single Crystal Transmetallation. Angew. Chemie - Int. Ed. 2015, 54, 6521–6525.
553	44.	Jansen, J.C.; Friess, K.; Drioli, E. Organic vapour transport in glassy perfluoropolymer membranes: A
554		simple semi-quantitative approach to analyze clustering phenomena by time lag measurements. J.

555		Memb. Sci. 2011, 367, 141–151.
556	45.	Crank, J. The mathematics of diffusion; 2nd ed.; Clarendon Press: Oxford, 1975; ISBN 0198533446.
557	46.	Fraga, S.C.; Monteleone, M.; Lanč, M.; Esposito, E.; Fuoco, A.; Giorno, L.; Pilnáček, K.; Friess, K.; Carta,
558		M.; McKeown, N.B.; et al. A novel time lag method for the analysis of mixed gas diffusion in polymeric
559		membranes by on-line mass spectrometry: Method development and validation. J. Memb. Sci. 2018, 561,
560		39–58.
561	47.	Fernandes, T.S.; Melo, W.D.C.; Kalinke, L.H.G.; Rabelo, R.; Valdo, A.K.; Da Silva, C.C.; Martins, F.T.;
562		Amorós, P.; Lloret, F.; Julve, M.; et al. 2D and 3D mixed MII/CuII metal-organic frameworks (M = Ca
563		and Sr) with: N, N '-2,6-pyridinebis(oxamate) and oxalate: Preparation and magneto-structural study.
564		Dalt. Trans. 2018, 47, 11539–11553.
565	48.	Khosravi, T.; Omidkhah, M.; Kaliaguine, S.; Rodrigue, D. Amine-functionalized CuBTC/poly(ether-b-
566		amide-6) (Pebax® MH 1657) mixed matrix membranes for CO2/CH4 separation. Can. J. Chem. Eng. 2017,
567		95, 2024–2033.
568	49.	Luo, H.; Vaivars, G.; Mathe, M. Cross-linked PEEK-WC proton exchange membrane for fuel cell. Int. J.
569		Hydrogen Energy 2009 , 34, 8616–8621.
570	50.	Li, W.; Galiano, F.; Estager, J.; Monbaliu, J.C.M.; Debecker, D.P.; Figoli, A.; Luis, P. Sorption and
571		pervaporation study of methanol/dimethyl carbonate mixture with poly(etheretherketone) (PEEK-WC)
572		membrane. J. Memb. Sci. 2018, 567, 303–310.
573	51.	Longo, M.; De Santo, M.P.; Esposito, E.; Fuoco, A.; Monteleone, M.; Giorno, L.; Jansen, J.C. Force
574		spectroscopy determination of Young's modulus in mixed matrix membranes. Polymer (Guildf). 2018,
575		156, 22–29.
576	52.	Paul, D.R. Effect of immobilizing adsorption on the diffusion time lag. J. Polym. Sci. Part A-2 Polym. Phys.
577		1969 , 7, 1811–1818.
578	53.	Grzywna, Z.; Podkowka, J. Effect of immobilizing adsorption on mass transport through polymer films.
579		J. Memb. Sci. 1981, 8, 23–31.
580	54.	Yampolskii, Y. Polymeric Gas Separation Membranes. 2012, 45, 3298–3311.
581	55.	Fuoco, A.; Rizzuto, C.; Tocci, E.; Monteleone, M.; Esposito, E.; Budd, P.M.; Carta, M.; Comesaña-
582		Gándara, B.; McKeown, N.B.; Jansen, J.C. The origin of size-selective gas transport through polymers of
583		intrinsic microporosity. J. Mater. Chem. A 2019, 7, 20121–20126.
584	56.	Swaidan, R.; Ghanem, B.; Pinnau, I. Fine-Tuned Intrinsically Ultramicroporous Polymers Redefine the
585		Permeability/Selectivity Upper Bounds of Membrane-Based Air and Hydrogen Separations. ACS Macro
586		<i>Lett.</i> 2015 , <i>4</i> , 947–951.
587	57.	Comesana-Gandara, B.; Chen, J.; Bezzu, C.G.; Carta, M.; Rose, I.; Ferrari, MC.; Esposito, E.; Fuoco, A.;
588		McKeown, N.B.; Jansen, J.C. Redefining the Robeson upper bounds for CO2/CH4 and CO2/N2
589		separations using a series of ultrapermeable benzotriptycene-based Polymers of Intrinsic Microporosity.
590		Submitt. to Energy Environ. Sci. 2019.
591		