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High Flux Thin-Film Nanocomposites with Embedded Boron Nitride Nanotubes for Nanofiltration

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10 Abstract

A novel thin film nanocomposite (TFN) membrane was obtained by incorporating boron nitride 11 nanotubes (BNNTs) into a polyamide (PA) thin selective layer prepared via interfacial 12 polymerisation. The addition of just 0.02 wt% of BNNTs led to a 4-fold increase in pure water 13 14 permeance with no loss in rejection for divalent salts, methylene blue or humic acid compared to the pure PA membrane. Loadings higher than 0.02 wt% of BNNTs led to agglomeration 15 16 with overall loss of performance. For the membranes containing 0.02 wt% BNNTs, the pure water permeance was 4.5 LMH@bar, with > 90% rejection of MgSO4 and > 80% rejection of 17 18 CaCl₂. Fouling tests with humic acid showed a flux recovery ratio of > 95% with ~50% lower flux loss during the fouling cycle compared to the polyamide only membrane. These values 19 20 represent a significant improvement over both commercial polyamide membranes and TFN 21 membranes incorporating carbon nanotubes. We assert that the very small quantity of BNNTs 22 needed to produce the enhanced performance opens the way to their use in water treatment 23 applications where nanofiltration membranes are subject to severe organic fouling.

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25

26 Keywords: Boron nitride nanotubes; Chemical vapour deposition; Interfacial Polymerisation; Nanofiltration

28 **1. Introduction**

Polymeric nanofiltration (NF) membranes have become a mainstay of water treatment processes, with high recovery rates [1], facile modular scale up [2] and economic viability across a broad range of feed [3]. NF membranes are particularly effective for the combinatorial rejection of salts, organic compounds, natural organic matter (NOM), and dyes. [4]. Industry, however, still faces challenges including high energy costs per unit volume of water purified [5], handling of retentate waste [6], membrane fouling, and a fundamental understanding of the mechanisms underlying the purification of complex feeds [7].

36 Thin-film composite (TFC) membranes composed of a very thin, dense selective layer supported by a porous support combine high flux and rejection with mechanical stability. 37 38 Although TFC membranes are successfully used commercially [8], there is still need to 39 increase efficiency, reduce energy consumption and extend chemical stability [9]. Many approaches have been explored to improve the performance of TFCs, including diverse 40 41 fabrication methods, and the tuning of precursors used to fabricate the membranes [10]. 42 Another promising strategy is the incorporation of inorganic nanomaterials in the thin selective layer to form so-called thin-film nanocomposite (TFN) membranes [7]. Nanomaterial additions 43 44 alter the structure of the selective layer by finely tuning properties such as hydrophilicity [11], porosity [12], surface zeta potential [13] and stability [14, 15]. Additionally, the nanomaterials 45 46 can introduce desired features such as fouling resistance [16], adsorption [17] and photocatalytic characteristics [18] into the membranes. 47

Amongst the wide range of nanomaterials tested to date [2], carbon nanotubes (CNTs) have 48 49 been considered for application in membrane technology due to their fast water transport and 50 low tortuosity [19]. The reduced friction of water on the defect-free carbon surface in these 51 nanotubes results in very high water permeances through the tubes [20], translating into higher efficiency (i.e. higher flux) per applied pressure [21]. However, efforts to create membranes 52 53 with the CNTs aligned perpendicularly to the membrane's surface in a commercially scalable 54 fashion have been have been, so far, unsuccessful [22-24]. On the other hand, the incorporation 55 of randomly aligned CNTs in polymer matrices has led to the successful formation of selective 56 membranes [25], but with only modest increases in water permeance and a decrease in 57 selectivity [26, 27]. While the former can be attributed to the small fraction of tubes directly 58 connecting feed and permeate, the latter is attributed to the formation of uncontrolled 59 permeation pathways at the interface between the CNTs and the polymer, due to poor chemical

60 compatibility [28]. This aspect has been addressed by introducing an additional functionalisation step with acid groups, to form hydroxyl and carboxyl groups on the tube 61 62 surface [29]. The functionalised CNTs have been added in relatively large amounts (1 wt% -10 wt%), with a corresponding increase in rejection of up to 5% and providing permeances up 63 64 to 50% higher than those of the starting polyamide membrane [30, 31]. CNTs have also been 65 functionalised with more complex zwitterion groups. However, these have penalised the water permeance when compared to pristine CNTs, though improving the rejection of NaCl from 66 67 97.6% to 98.5% thanks to the steric hindrance of the zwitterion functional groups [32]. Despite 68 these promising results, the potential for permeance increase is limited by the low loadings of 69 hydrophobic nanomaterials that can be incorporated into the membrane matrix before the onset of agglomeration, which leads to the formation of pinholes, with a subsequent loss of 70 71 performance [33].

Herein, boron nitride nanotubes (BNNTs) are investigated as nanomaterials for TFN 72 fabrication. We speculate that their physico-chemical characteristics will overcome some of 73 74 the limitations of CNT-based TFNs highlighted above. Hexagonal BNNTs (hBN) are isostructural to graphitic CNTs, but behaving as electrical insulators and showing higher 75 76 resistance to oxidation [34]. Molecular dynamics simulations on BNNTs in the subcontinuum 77 range, with diameters 0.8 nm, have shown faster pure water flux than in CNTs [35]. For such small diameters it was shown that increased van der Waals and electrostatic interactions 78 79 between the nanotube walls and the water molecules contribute to an easier filling of the bore 80 of BNNTs than for CNTs [36]. However, when the diameters of the tubes studied were larger, 81 CNTs outperformed BNNTs in terms of improved water fluxes [37]. This was ascribed to 82 differences in the electronic landscape in the two nanotube walls [38]. Simulations also showed 83 that BNNTs have tunable cation and anion selective properties due to the partial charge on the boron and nitrogen atoms of the nanotube [39] and osmotic energy storage capabilities [40]. 84 85 Additionally, when boron nitride nanosheets were recently embedded in mixed matrix 86 membranes, they showed improved fouling resistance [26]. Boron nitride nanotubes have 87 recently been used to fabricate ultrafiltration membranes with improved thermal resistance and 88 mechanical stability [41]. Moreover, CNTs have been shown to have antioxidant capabilities 89 that slow down chlorine attack on polymeric membranes [28], however this effect is as yet 90 unreported for BNNTs. Using BNNTs as membrane nanofillers is motivated, together with its 91 novelty, by the fact that materials with high negative zeta potentials allow for rejection of 92 pollutants not only by size but also by charge [42].

93 Despite the many potential advantages of BNNTs over CNTs, there has been notably less 94 published research on BNNTs than on their carbon counterparts, attributed mainly to the lack 95 of methods for the production of BNNTs at scale [34]. This obstacle has been overcome in this 96 work, optimising a known technique [43] for the production of BNNTs by chemical vapour 97 deposition (CVD). The nanomaterial was then embedded in the selective phase of an 98 interfacially polymerised polyamide (PA-BNNT) membrane. BN is negatively charged in 99 water over a broad pH range [44] and can adsorb OH- on its surface further increasing its 100 negative charge [40].

101 **2. Materials and Methods**

102 2.1 Materials

Boron (B, \geq 95%), magnesium oxide (MgO, \geq 99.99%), iron oxide (Fe₂O₃, \geq 99.9%) and MgO 103 104 nanopowder (average particle size ≤ 50 nm, measured by BET [45]) were purchased from 105 Sigma Aldrich. Ethanol (\geq 99%) was purchased from Fisher Scientific. P-type silicon wafers polished on one side were purchased by Agar Scientific. Hydrochloric acid (38%) was 106 purchased from Sigma Aldrich. Commercial polyether sulfone (PES) membrane Microdyn 107 108 Nadir PMUP010 with 10 kDa nominal molecular weight cut-off was purchased from Steriltech. Deionised (DI) water was used unless specified otherwise. Methanol (MeOH, anhydrous, 109 99.8%), Piperazine (PIP, 99%) ReagentPlus®, with MW86, n-hexane (anhydrous, 95%) and 110 111 trimesoyl chloride (TMC, 99%) were purchased from Sigma Aldrich. Diiodomethane (DIM, 112 99%) was purchased from Sigma Aldrich. Salts (e.g., NaCl, CaCl2 and MgSO4) were purchased 113 from Sigma Aldrich. Sodium hypochlorite technical solution was purchased from Fischer 114 Scientific.

115 2.2 BNNTs synthesis

BNNTs synthesis (Fig. 1) with ammonia gas and boron powder precursors was catalysed by Fe₂O₃ and MgO catalysts with a molar ratio of B:MgO:Fe₂O₃ = 2:1:1. B, MgO and Fe₂O₃ were pre-mixed at 250 r.p.m. for 12 h in a Fritsch Pulverisette P6 planetary ball mill, half filling a 45 ml stainless steel grinding bowl with 2.2 g of B, MgO and Fe₂O₃ in an ethanol suspension and 18 grinding balls with 5 mm diameter. After ball milling, 5 ml of precursor was poured in a CoorsTM alumina combustion boat, which was then capped with a silicon wafer previously seeded with MgO nanopowder. The closed boat was then placed in a 15 cm long quartz test 123 tube (ID 18 mm, OD 19 mm) facing the gas inlet in the centre of a quartz tube reactor (H-124 Baumbach, ID 20 mm, OD 22 mm) in a three sections horizontal TZF 12/38/850 type 125 CARBOLITE tubular furnace. Temperatures inside the furnace were monitored by external 126 thermocouples. Gas flows were controlled with Omega FMA 5400A/5500A series mass flow 127 controllers (MFCs) regulated by a LabVIEW program. The tube reactor was abundantly flushed with Ar and then let ramp up at 10 °C/min up to 1100 °C under a 200 sccm Ar flow. 128 129 Then, the gas flow was switched to 145 sccm NH3 and the temperature increased to 1200 °C at 130 the same heating rate. This maximum temperature was kept for 1 hour before letting the system 131 cool down to room temperature under a 200 sccm Ar stream. The exhaust NH₃ gas was neutralized with a sulphuric acid scrubber, generating ammonium sulphate salts. The BN 132 133 nanotubes white powder was gently removed with a stainless steel spatula by scratching it from 134 the silicon substrate and boat top and sides. The unreacted boron in the collected white powder 135 was removed in air at 700 °C for 2 hours, where it reacted to form boric anhydrite vapour. BNNTs have high thermal stability and are resistant to oxidation up to 950 °C [45]. Catalysts 136 were removed with a 3 hours 10% HCl (purity 36.5-38.0%, purchased from Sigma Aldrich) 137 water cleaning at 40 °C, followed by washing of the products in DI water by vacuum filtration 138 using a 0.45 µm pore size nylon membrane (Pall Corporation). 139



140

141 **Fig. 1** Schematic of the CVD of boron nitride nanotubes production.

142 2.3 Polyamide membrane fabrication

143 The PA-BNNTs membranes were synthesized by interfacial polymerisation following an 144 established vacuum filtration technique [46], which has been recently used for the fabrication 145 of nanocomposite membranes [47]. The PES support membrane (Microdyn Nadir PMUP010) 146 was cut in discs with 5.5 cm in diameter, and then flushed with 20 ml of water in a filtration 147 setup prior to synthesis. Then, 1 wt% PIP aqueous solution (MeOH:H2O 50:50 v/v%) solution 148 was prepared by rapidly dissolving the PIP flakes. For the PA-BNNT membranes, 0.01, 0.02 149 or 0.03 wt% BNNTs were dispersed in the amine solution by ultrasonication for 1 h (Table 1). 150 Then, 25 ml amine solution or amine solution with dispersed BNNT was pumped through the 151 support membrane until the entire amount of solution was filtered, while ensuring the membrane remained wet. Subsequently, any residual drops on the side of the wetted membrane 152 153 were removed by using an air gun to avoid the formation of defects in the resulted amine-rich 154 film. In the interfacial polymerisation, a 0.8 wt% TMC n-hexane solution statically contacted 155 the amine-saturated support in the filtration setup. The reaction time was 3 min. The residual 156 organic solution was discarded, and the membrane was quenched with n-hexane for 1 min. 157 After reaction, the membrane was left to dry at room temperature for 24 h. Similarly, a thin film was formed by contacting 1 ml of PIP solution with 1ml of TMC solution, specifically for 158 159 the analysis of a free standing film at the interface.

Membranes with different concentration of BNNTs in the starting solution were prepared; their nomenclature, PIP solution composition, and the estimated weight of nanofiller deposited per unit area by filtering 25 ml of amine solution are reported in Table 1. The concentrations to be investigated were chosen in a range where no obvious large agglomeration could be observed on the membrane top surface with the naked eye. It should be noted that there is an uncertainty in the estimation of the amount of BNNTs per unit area, due to the possibility that some minor fraction of nanomaterial filtered through the PES support.

167	Table 1 Composition of the PIP s	olutions in M	eOH:H2O 50:50 v	v/v% for the PA	A-BNNTs membranes
	1				

Membrane	PIP (wt%) MeOH/H2O (wt%)	BNNTs (wt%)	<i>c</i> _{<i>S</i>} (mg/cm ₂)
PA-BARE	1.00 99.00	0.00	0.000
PA-BNNTs0.01	1.00 98.99	0.01	0.096
PA-BNNTs0.02	1.00 98.98	0.02	0.193
PA-BNNTs0.03	1.00 98.97	0.03	0.283

¹⁶⁸

169 2.4 Characterisation of BNNTs powders

170 The produced nanotubes were coated with 5 nm of chromium and positioned on carbon tape 171 for analysis with a JEOL JSM-6301F FESEM at 5kV. JEOL JSM-2100Plus TEM samples were prepared by dispersing the nanotubes in ethanol. Two to five drops of the sample were then 172 173 placed on a TEM window (Lacey carbon purchased from EM Resolutions) until a desirable 174 concentration was reached. Analysis of structural features with ImageJ was done on a minimum 175 of 10 measurements. Optical Images of the substrates with BNNTs grown on them were taken 176 with a Digital Microscope VHX-6000 series. Raman spectroscopy was carried out on the as-177 synthesized samples on a glass slide in a Renishaw Raman Microscope series 1000 using a 178 frequency-doubled argon ion laser (wavelength 244 cm-1, 5.08 eV) with spectral resolution of 179 5 - 10 cm⁻¹ and a 40 × UV objective lens. XRD tests of products of the synthesis were reduced 180 into fine powders dispersed on a silicon wafer were carried on 1 mg of product with a Bruker 181 D8 Advance for $10 < 2\theta < 70$ with a Vantec detector with Cu K-alpha radiation.

182 XPS was performed on powdered BNNTs samples using a Thermo Fisher Scientific K-alpha+ 183 spectrometer. Samples were analysed using a micro-focused monochromatic Al x-ray source 184 (72 W) over an area of approximately 400 microns. Data was recorded at pass energies of 150 185 eV for survey scans and 40 eV for high resolution scan with 1 eV and 0.1 eV step sizes 186 respectively. Charge neutralisation of the sample was achieved using a combination of both 187 low energy electrons and argon ions. Data analysis was performed in CasaXPS using a Shirley 188 type background and Scofield cross sections, with an energy dependence of -0.6.

189 2.5 Polyamide membrane characterisation

Membranes were coated with 10 nm of Cr before imaging in a JEOL JSM-6301F FESEM. 190 191 Micrographs were taken at 5kV. Atomic Force Microscopy (AFM) Nanosurf EasyScan 2 Flex 192 scans were taken by using the Dynamic Force Mode with a 190Al-G tip on areas $5 \mu m \times 5 \mu m$ with 256 points/line. The data was analysed with Gwyddyion and the software's internal 193 194 functions were used to assess the membrane's average roughness (R_a) . Dataphysics Optical 195 Contact Angle (OCA) Measuring Device with 0.5 µl wetting liquid drops was also used to 196 characterise the membranes. The Young's contact angle (θ_Y) on a flat smooth surface is related 197 to the measured contact angle (θ_w) via the Wenzel equation [48]:

$$\cos\theta_W = r \cos\theta_Y,\tag{1}$$

198 where r is the ratio between the membrane surface area and the projected area, obtained by 199 AFM.

The surface zeta potential of samples with dimensions 2×1 cm or 1×1 cm was measured in a SurPASS electrokinetic analyser with adjustable gap cell in a pH range between 3 and 10. For each pH value, the measurement was repeated four times. A Perkin Elmer Spectrum FTIR-ATR Spectrometer was used to characterise the membranes' selective layer surface, with 16 scans per run between 600 and 4000 cm-1 and a spatial resolution of 2 cm-1.

Sections of the polyamide membranes top surface measuring $1 \text{ cm} \times 1 \text{ cm}$ were also analysed by XPS. The ratio O/N from the XPS analysis was measured to assess the degree of crosslinking in relation to loading percentage. This was calculated from

crosslinking (%) =
$$\left(\frac{(O/N)_{XPS} - (O/N)_{fully\ cross-linked}}{(O/N)_{fully\ linear} - (O/N)_{fully\ cross-linked}}\right)$$
, (2)

where $(O/N)_{XPS}$ is the oxygen to nitrogen ratio obtained experimentally, neglecting the 398 eV contribution of the *h*BN nitrogen [49]. $(O/N)_{fully \ linear}$ and $(O/N)_{fully \ crosslinked}$ are the ratios for a fully linear and fully crosslinked polyamide, respectively [50].

- 211 Phillips CM200 TWIN TEM samples were prepared by gently depositing a thin film on a TEM
- 212 window and imaging it at 250 and 25k magnification.

213 **2.6** *Membrane filtration performance*

- 214 Pure water flow was tested in cross flow mode (schematics in [26]) on a minimum of three
- 215 membranes per composition, with 24 h of compaction at 7 bar and three days of testing at 3
- 216 bar.
- Dye and salts rejection was tested for 7 h with a 45 L h-1 pump flow rate. Rejection of methylene
 blue was assessed using a UV spectrophotometer (UV Cary 100, Agilent, U.K.), while rejection
- 219 of the salts was measured using an Orion Versastar ThermoScientific conductivity meter. The
- feed concentration for the dye tests was 0.01 g L-1, whilst for salts rejection it ranged from 500
 to 2000 ppm.
- Humic acid fouling tests were performed on pre-compacted membranes in cross flow mode. The 2.5 L feed of 1 g L-1 humic acid in water was prepared beforehand and mixed with a magnetic stirrer for 24 hours before the test. The fouling test consisted of two fouling cycles of 15 hours and two cleaning cycles of one hour each. The flow rate was set as 175 ml min-1 (Reynolds number, Re = 130) and 750 ml/min (Re = 550) for fouling and cleaning respectively. Flux recovery ratio (*FRR*) was measured before each cycle as follows:

$$FRR(\%) = \left(\frac{J_{AF}}{J_{BF}}\right) \times 100, \qquad (3)$$

- where J_{BF} and J_{AF} are the two fluxes before fouling and after cleaning, calculated for each cycle.
- 230 The membrane total resistance (R_t) after 15 hours of fouling test was also calculated:

$$R_t = R_m + R_r + R_{ir}, (4)$$

with R_m , R_r and R_{ir} being the intrinsic membrane resistance and the reversible and irreversible fouling resistance, respectively. These are calculated by using:

$$R_m = \frac{\Delta P}{\mu \times J_{BF}};\tag{5}$$

$$R_{ir} = \frac{\Delta P}{\mu \times J_{AF}} - R_m; \tag{6}$$

$$R_r = \frac{\Delta P}{\mu \times J_F} - R_m - R_{ir}.$$
⁽⁷⁾

233

The transmembrane pressure is indicated with ΔP and the viscosity with μ , while J_F is the flux in m₃ s-1 of humic acid after 15 hours of fouling. The term $R_{\%}$ was used to assess the extent of reversible fouling in each membrane during fouling tests:

237

$$R_{\%} = \frac{R_r}{R_t} \times 100.$$
⁽⁸⁾

238

The first 10 ml of permeate were discarded in every test, to account for dead volume and permeate tubing. The mass balance for the rejection of component i is so calculated:

mass balance (%) =
$$\frac{V_{P,i}c_{P,i} + V_{R,i}c_{R,i}}{V_{F,i}c_{F,i}} \times 100$$
, (9)

241 where the subscripts P, R and F stand for permeate, feed and retentate respectively; V is the 242 volume and c the concentration.

Chlorine resistance tests were performed with 4 L feed water containing 2000 ppm NaOCl and
2000 ppm CaCl₂. Rejection and permeance were monitored over a period of 5 hours and 30
minutes, while the membranes were tested in cross flow mode with a pump velocity of 45 L h1 at 3 bar. Membranes were pre-compacted before rejection tests.

3. Results and Discussion

248 3.1 BNNTs synthesis

249 BNNTs were synthesized via chemical vapour deposition, resulting in a dense product (Fig. 250 2a), with the BNNTs having open ends (Fig. 2b) and homogenously grown over the entire 251 substrate area (Fig. 2c) with a vertical orientation (inset cross-sectional HRTEM). The BNNTs powder was collected with a yield of ~10 mg per run by simple scratching of the silicon wafer 252 253 (W x L 14 x 70 mm) substrate. The diffraction pattern generated by a single tube (Fig. 2d) 254 confirmed the hBN (002) crystal structure of boron nitride multiwalled (wall thickness ~ 4.5 255 nm) nanotubes [51], with a straight inner channel of *c.a.* 8 nm in diameter and an outer diameter of c.a. 17 nm (Fig. 2e) where the interspace layer distance of hBN is 0.328 nm as expected 256 257 from literature [34]. Combining the open ends and penetrating inner voids, the as-synthesized BNNTs are ideal candidates for creating nanoscale channels in TFN membranes [52]. The 258 259 morphology of the BNNTs showed a relatively wide size distribution with outer diameters in the range of 5 to 105 nm and tube lengths in the range of 1 to 5 µm, as statistically counted by 260 261 200 tubes in TEM micrographs (Fig. 2f).





Fig. 2 (a) FESEM micrograph of BNNTs on the Si wafer taken at 3000 x magnification and (b) collected from the alumina boat taken at 5000 x magnification with 8 mm working distance. (c) Optical microscope image (500 x) with a TEM micrograph inset showing BNNTs grown on a piece of silicon wafer; TEM micrographs and (d) corresponding diffraction pattern and (e) measures for inner and outer tube diameter of a selected BNNT tube where the *h*BN interlayer spacing can be observed. (f) Counts of outer diameter intervals for 200 tubes randomly imaged with TEM.

269 The BNNTs powder samples present the typical Raman hBN peak at 1369 cm-1 (Fig. 3a). The XRD spectra in Fig. 3b shows four main BN peaks at $2\theta = 10.5_{\circ}$ (*h*BN 001), 29_{\circ} (*h*BN 002), 270 40_o (*h*BN 100) and 53_o (*h*BN 004) [53], alongside some iron and MgO peaks in the region 20_o 271 272 $< 2\theta < 40_{\circ}$, identified as catalyst impurities. Boron nitride FTIR peaks are identified in Fig. 3c for the vibration mode along the tubes' longitudinal axis at 1367 cm-1, in the tangential 273 274 direction at 1537 cm-1 and the out of plane buckling mode at 795 cm-1 [43]. The spectra also 275 presents peaks attributed to Si-O in the 1130-1000 cm-1 region and Si-H in the 850-900 cm-1 276 region, as the spectra was taken before the BNNTs material was scraped from the Si substrate. 277 XPS results (Fig. 3d) identify a B/N atomic ratio of 1.18 in the powder BNNT sample, very close to the theoretical value of 1. Peaks for *h*BN are identified at 190.41 eV and 398.00 eV 278 for B and N respectively, and a 3.75 at.% of N-B-O bonds can be observed in the B high 279 280 resolution spectra [49].

281



Fig. 3 (a) Raman analysis of BNNTs with the characteristic *h*BN peak at 1369 cm-1 and (b) XRD pattern of the
as produced BNNTs powder, showing four characteristic *h*BN peaks; (c) FTIR of BNNTs on silicon wafer; and
(d) B1s and N1s high resolution XPS spectra of the BNNTs.

286 3.2 Characterisation of PA-BNNTs membranes

The surface and cross-sectional morphologies of the produced polyamide membranes are shown in Fig. 4. The irregular morphology increased with BNNTs loading, which is consistent with an increase in the average surface roughness (R_a , nm) in Table 2, and therefore the ratio between the membrane surface area and the projected area, r, and morphological changes measured by AFM (Fig. 5). The crumpled areas observed in the PA-BNNTs membranes showed similar material stiffness as the rest of the membrane (see phase plot analysis in **Error! Reference source not found.**), indicating that no BNNTs protrude out of the membrane from





295

Fig. 4 FESEM top and cross section of membranes prepared from solutions containing different percentages of nanofiller: (a,e) bare, (b,f) 0.01 wt%, (c,g) 0.02 wt% and (d,h) 0.03 wt% PA-BNNTs membranes.



298

299 Fig. 5 AFM maps of membranes (a) bare, (b) 0.01 wt%, (c) 0.02 wt% and (d) 0.03 wt% PA-BNNTs membranes.

Measured contact angle varies as a function of BNNTs concentration (Table 2). Water contact angle increases by approximately 20_{\circ} from PA-BARE to PA-BNNTs0.03, in agreement with both the increase in roughness R_a already observed in Fig. 5 and an observed reduced material hydrophilicity. When the measured contact angle and roughness are used to calculate values for the Young's contact angles *via* the Wenzel equation [48], an observable although not drastic increase in material hydrophilicity is observed, with Young's water contact angles increasing 306 by 15-20% for each 0.01 wt% of BNNT added. In contrast, the non-polar diiodomethane 307 contact angle (θ_{Y_DIM}) decreased over the four membranes tested, in line with the loss of 308 hydrophilicity of the membrane.

- 309 **Table 2** Measured water (θ_{WATER}) and diiodomehtane (θ_{DIM}) contact angles, Young water (θ_{Y_WATER}) and
- 310 diiodomehtane ($\theta_{Y_{DIM}}$) contact angles obtained applying the Wenzel equation, average surface roughness R_a
- 311 results on PA-BARE and PA-BNNTs membranes and ratio *r* between the membrane surface area and the
- 312 projected area, obtained by AFM.

Membrane	θ water	θdim	θ_{Y_WATER}	θ_{Y_DIM}	R _a	r	
		(de	g, ± 2°)		(nm)	(-)	
PA-BARE	25	30	35	39	19	1.11	
PA-BNNTs0.01	32	29	40	38	49	1.11	
PA-BNNTs0.02	30	28	49	48	172	1.32	
PA-BNNTs0.03	45	10	59	44	181	1.37	

- 313 A free-standing film was placed in the TEM to observe the embedment of the BNNTs in the
- polyamide (Fig. 6a). Wrinkles in the thin layer are created when this is transferred to the TEM
- grid, but these formations can be clearly differentiated from the BNNTs as these show a hollow
- 316 nature as previously shown in Fig. 2 d,e and in the inset of Fig. 6a. Fig. 6b shows a picture of
- 317 the polyamide thin film formed at the interface between the PIP H₂O/MeOH solution and TMC
- 318 in hexane solution.



Fig. 6 (a) TEM micrograph of a free-standing film loaded with 0.01wt% BNNTs, observable in the magnified
 inset. Wrinkles generate in the film when it is transferred to the TEM grid. (b) Picture of a thin film produced at
 the interface between the TMC and PIP solutions.

As expected, the introduction of a negatively charged material in the texture of the IP membrane slightly decreased its surface zeta potential (Fig. 7). Although statistically significant (p-value=0.002), this change is not as dramatic as might be expected by the introduction of negatively charged nanomaterial, leading to the hypothesis that the vast majority of nanomaterial particles are surrounded by the selective polymer layer, and do not protrude from the top surface, consistent with the top layer stiffness results in **Error! Reference source not found.**

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Fig. 7 Surface zeta potential (ζ_m) vs. pH of a PA-BARE (IEP=4.40) and PA-BNNTs0.01 (IEP=4.32).





Fig. 8 FTIR spectra for (a) bare, (b) 0.01 wt%, (c) 0.02 wt% and (d) 0.03 wt% PA-BNNTs membranes.

Fig. 8a reports the FTIR-ATR spectra for the IP BARE membrane [54], and Fig. 8b-c the scans 336 for membranes with increasing BNNT loading. The beam penetrates the sample for $0.5 - 2 \,\mu m$ 337 during testing [55], therefore representing the whole thickness of the membrane top layer and 338 not just the very top surface. Thus, at the highest BNNTs concentration, the predominance of 339 340 the 1375 cm-1 BNNT' peak, corresponding to the longitudinal vibration mode of the tube, is observed [43, 56], together with a B-O functionalisation peak at 1721 cm-1 [57], showing a new 341 342 bond not observable in the FTIR for the BNNTs in powder form (Fig. 3c), and potentially 343 indicating some interaction between the boron nitride and the polymeric matrix already visible 344 for BNNTs concentrations of 0.01wt.%.

The O/N ratio for each membrane is reported in Table 3. A fully cross-linked polyamide, $(O/N)_{fully \, cross-linked}$, has a O/N ratio of 1 and a theoretically fully linear polyamide, $(O/N)_{fully \, linear}$, has a O/N ratio of 2 [50]. From these values, the degree of crosslinking was calculated from XPS results using Eq. (2). While the addition of 0.01wt% BNNTs increases the degree of crosslinking from 0.7 to 1.0, this declines moving to 0.02wt% and 0.03wt%, seemingly increasing the density of transport pathways available in selective layer [58].

Membrane	$(O/N)_{XPS}$	crosslinking (%)
PA-BARE	1.3	0.7
PA-BNNTs0.01	1.0	1.0
PA-BNNTs0.02	1.6	0.4
PA-BNNTs0.03	1.7	0.3

Table 3 Experimental *O*/*N* from XPS and degree of crosslinking.

352

353 3.3 Filtration performance of PA-BNNTs membranes: permeance, rejection and fouling

The permeance of the IP membranes evaluated increases with BNNT loading (Fig. 9a), from 354 355 an average of 1.1 LMH/bar for the bare membrane to 2.7 LMH/bar for the PA-BNNTs0.01, 4.5 LMH/bar for the PA-BNNTs0.02 and 4.1 LMH/bar for PA-BNNTs0.03. The permeance 356 values hints at a convex profile, often found in literature as a function of loading, as the initial 357 addition of nanomaterial generates an initial increase in pore size, and then a decrease [29]. 358 359 The former is ascribed to the higher free volume provided by the BNNTs [59], whereas the 360 latter is due to increasing agglomeration [60]. The increase in water flow pathways however does not impact the membrane's rejection performance up to 0.02 wt% loading, most likely 361 because the polyamide layer is still the main contribution to rejection up to that concentration 362 value [59]. This suggests that the addition of the BNNT fillers might have created additional 363 pathways for facile transport of water but not solutes [61, 62], and the slight increase in 364 365 membrane surface charge might also have contributed to maintain a high rejection [2]. The 366 slight decrease observed in the permeance value from PA-BNNTs0.02 to PA-BNNTs0.03 does 367 not, however, show a statistically significant difference (Student's *t*-test, p-value > 0.05).

368 Two batches of PA-BNNTs membranes (empty and filled symbols in Fig. 9a) were tested for 369 monovalent NaCl and divalent (CaCl₂, MgSO₄) salts rejection. Calibration curves for these 370 measurements are reported in Error! Reference source not found.-4, and ionic concentrations 371 in Error! Reference source not found. The rejection for NaCl remains low (20 - 40 %) for 372 the whole concertation range investigated with 0.03 wt% BNNTs being the worst performing 373 case. However the membranes perform well for divalent salts rejection, with the rejection for 374 MgSO4 is above 90 % for loading up to 0.02 wt% BNNTs, whilst it decreases to ~ 80 % for 375 PA-BNNTs0.03. CaCl₂ rejection raises from 75 – 80 % for the bare PA membrane to 97% for 376 the PA-BNNTs0.01 and then decreases to around 40 % with further addition of nanofiller. The 377 mass balance for the rejection of salts was \geq 96% for CaCl₂ and NaCl, and \geq 90% for MgSO₄. 378 In all cases, the addition of 0.03 wt% of BNNTs notably penalises the membranes rejection,

whilst the membranes show highly desirable performances for loadings ≤ 0.02 wt%, with PA-BNNTs0.02, in particular, combining the highest permeance (4.5 LMH/bar) with the highest divalent salt rejection. This is conceivably due to the additional free volume and thus water pathways offered by the presence of the BNNTs in the matrix, while the polyamide enveloping the nanomaterial provides salt rejection.

384 3.6

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Fig. 9 (a) Salt rejection (left axis) and permeance (right axis) of the bare and TFN membranes containing BNNTs; rejection tests have been repeated on two batches of membranes indicated with full and empty dots for MgSO4, CaCl₂ and NaCl: (**b**) First and (**c**) second fouling cycle; (**d**) Rejection of methylene blue (\blacklozenge) and (\triangleleft , \triangleleft) humic acid together with flux recovery ratio (FRR) performances in the two fouling cycles in orange and red respectively; (**e**) first and (**f**) second fouling cycle, as displayed in the top panel, but normalised by initial pure water permeance for each membrane tested.

The observed behaviour showed little change in terms of recovering initial flux after two long 392 fouling sequences in Fig. 9b and c, with high FRR: 97%, 100%, 95%, 97% for the first cycle 393 394 and 100%, 100%, 96, 92% in the second cycle for PA-BARE, PA-BNNTs0.01, PA-395 BNNTs0.02 and PA-BNNTs0.03, respectively. The membranes, possessing a white, opaque 396 colour at the start, could be cleaned completely by vigorous water flushing and no irreversible 397 contamination could be observed visually after the test or by the FRR results in the fouling 398 cycles. The pump flow rate chosen in this study (175 ml min-1) is a common setting for 399 membrane fouling tests [26]. However, this behaviour might change if the test was carried out at higher flows, where some irreversible fouling might be observed. However, Fig. 9e and f 400

401 explicitly show that PA-BARE reversibly fouls to a greater extent, decreasing to 40% of the 402 initial flux, compared to 80-90% for the TFN membranes. This is possibly due to the absence 403 of repulsion between the fouling material and nanofiller, which may lead to a higher percentage 404 flux decline during tests, even though this is eventually recovered after cleaning. The 405 contribution to total resistance during fouling can be observed in Fig.10.



407 **Fig. 10** Calculated membrane (R_m) , irreversible (R_{ir}) , reversible (R_r) and total (R_t) resistances for the 408 membranes under analysis during the first fouling cycle.

406

409 The fouling layer resistance contribution to the total resistance decreased with increased BNNT loading, as indicated by the decrease in value of the parameter $R_{\%}$ for the first fouling cycle 410 from 60.5% for PA-BARE to 8.0% for PA-BNNTs0.03 (Error! Reference source not 411 412 found.). This can be observed in Error! Reference source not found. for the second fouling 413 cycle as well, where, however, the benefit in terms of lower fouling layer resistance at high 414 BNNTs loading is attenuated by a slightly lower FRR in PA-BNNTs0.02 and PA-BNNTs0.03. 415 Fig. 9d shows high rejection (80-90%) of humic acid in all the membranes tested (UV-vis 416 calibration curve reported in Error! Reference source not found.). In addition to being able to effectively reject the foulant, all membranes could recover $\geq 95\%$ of their initial flux, with 417 418 PA-BNNTs0.03 recovering $\geq 90\%$ of its flux after physical cleaning, indicating that the 419 increased membrane roughness can make the removal of the formed fouling layer harder [4]. 420 Fig.9d also reports data on the rejection of methylene blue dye, with the addition of 0.01 wt%

421 BNNTs improving rejection by 17% compared to PA-BARE. Rejection then decreased with

422 increasing amounts of BNNTs added. As observed in the case of salts, the addition of 0.03 wt% 423 of BNNTs to the thin film worsens rejection performances. The mass balance for the rejection 424 of methylene blue was \geq 96% for all samples.

425 The relation between rejection and CaCl₂ concentration in the operational range 500 - 2000 426 ppm was also studied (Fig. 11), with a stable selectivity between 500 and 1500 ppm. Above 427 this value, when Donnan type rejection becomes predominant [63] a decrease in rejection is 428 observed for the PA-BARE and PA-BNNTs0.03, but not for the PA-BNNTs0.01 and PA-BNNTs0.02 for a concentration of CaCl2 of 2000 ppm. When the PA layer is the major 429 contributor to rejection, however, a constant rejection over different concentrations is expected 430 431 [32]. We can therefore conclude that the PA layer is the major contributor to the rejection of the membranes analysed, for all membranes but PA-BNNTs, where hypothesized defects due 432 433 to the high loading demonstrate a crucial concern particularly at high concentrations (2000 434 ppm).



435

436 **Fig. 11** Dependence of CaCl₂ rejection on salt concentration for the membranes analysed in this work.

Membranes based on a polyamide linkage are prone to attack by chlorine in the feed, as free chlorine radicals tend to be attracted by the N-H electron density [2]. Thus, NaOCl exposure is studied here for the membranes under analysis. As observed in Fig. 12, the exposure to the chlorinating agent is more adverse for all PA-BNNTs membranes compared to the PA-BARE, indicating that the introduction of BNNTs in the polyamide structure exposes the amide bonds prone to chlorine attacks. Previously it was found that increasing the density of amide bonds in the PA membrane is a successful strategy for increasing chlorine resistance [64], and this is 444 consistent with the decrease in crosslinking in the PA-BNNTs membranes leading to premature 445 failure in presence of NaOCl. In fact, at an exposure of 5000 ppm over 1 hr, while CaCl₂ 446 rejection of PA-BARE decreased from 89% to 73%, PA-BNNT 0.01wt% plummeted from 447 97% to 32%. The permeance of PA-BNNTs 0.02wt% increased from 3.67 LMH/bar to 4.68 448 LMH/bar after 11000 ppm over 1 hr chlorine exposure. However, it remained fairly steady for 449 the other membranes, indicating that the maximum exposure tested did not dissolve the PA 450 layer, but was enough to perturb it and decrease notably its ion rejection.



452 Fig. 12 Rejection (*R*) of CaCl₂ and permeance (*K*) of PA-BARE, PA-BNNTs0.01, PA-BNNTs0.02 and PA453 BNNTs0.03 as a function of the exposure to sodium hypochlorite.

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454 The best performing PA-BNNTs membrane in this work could provide 4 times higher 455 permeance than PA-BARE membranes with only 0.193 mg cm-2 of nanofiller on the membrane 456 area. For their part, hydroxyl functionalised CNT membranes were reported to show 2 times 457 higher pure water permeance than thin film polyamide membranes, but required 13.3 mg cm-2 458 of filler [65]. An increase of permeance up to 2.7 times was reported in PA-CNTs, but this 459 required the use of 0.458 mg/cm² of modified MWCNTs. When compared with TFN 460 membranes based on CNTs, the membranes in this work have therefore the capability to 461 notably improve the permeance of pure polyamide using a limited amount of nanofiller (Error! 462 **Reference source not found.**) and without requiring any additional functionalisation step.

463 **Conclusions**

In this work, a known synthesis route for the production of boron nitride nanotubes was 464 465 optimised and deployed for the production of larger amounts of boron nitride nanotubes, which were then used as a nanofillers in nanofiltration thin film nanocomposite membranes prepared 466 467 via interfacial polymerisation of PA. BNNTs homogeneously integrate in the polyamide layer, forming a B-O bond between the nanofiller and the polymer. Rejection of divalent and 468 469 monovalent salts is not compromised for up to 0.02wt% BNNTs added to the aqueous phase 470 in interfacial polymerisation, while the average permeance at this concentration goes up four times compared to the permeance of a membrane with no nanofiller. This is ascribed to an 471 472 increase in water transport pathways given by the boron nitride nanochannels enveloped by the selective layer, with no appreciable loss of selectivity compared to the bare PA membrane. A 473 permeance 4.5 times higher than in a bare PA membrane can be observed for low amounts of 474 BNNTs, thus considerably limiting costs of adding nanofillers. Nonetheless, potential 475 476 nanofiller leaching and recycling will have to be further investigated prior to large scale application, as BNNTs may be hazardous for the environment. In addition, the BNNTs 477 478 membranes show a high resistance to irreversible fouling. This is a desirable condition for applications in, for example, the food industry, where standard operations take place in highly 479 fouling environments. 480

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