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# 2 Metal-free modified boron nitride for enhanced CO<sub>2</sub>

## 3 capture

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12 Abstract: Porous boron nitride is a new class of solid adsorbent with applications in CO<sub>2</sub> capture. In 13 order to further enhance the adsorption capacities of materials, new strategies such as porosity 14 tuning, element doping and surface modification have been taken into account. In this work, metal-15 free modification of porous boron nitride (BN) has been prepared by a structure directing agent via 16 simple heat treatment under N<sub>2</sub> flow. We have demonstrated that textural properties of BN play a 17 pivotal role in CO<sub>2</sub> adsorption behavior. Therefore, addition of a triblock copolymer surfactant 18 (P123) has been adopted to improve the pore ordering and textural properties of porous BN and its 19 influence on the morphological and structural properties of pristine BN has been characterised. The 20 obtained BN-P123 exhibits a high surface area of 476 m<sup>2</sup>/g, a large pore volume of 0.83 cm<sup>3</sup>/g with 21 an abundance of micropores. More importantly, after modification with P123 copolymer, the 22 capacity of pure CO<sub>2</sub> on porous BN has improved by about 34.5% compared to pristine BN (2.69 23 mmol/g for BN-P123 vs. 2 mmol/g for pristine BN under ambient condition). The unique 24 characteristics of boron nitride opens up new routes for designing porous BN, which could be 25 employed for optimizing CO<sub>2</sub> adsorption.

Keywords: Porous boron nitride; metal-free modification; structure directing agent; CO<sub>2</sub> capture.

### 28 1. Introduction

29 Over the last decades, climate change has become a global challenge for countries across the 30 world [1]. Carbon capture and storage (CCS) is expected to play a substantial role in meeting the 31 global warming targets set by the Inter-governmental Panel on Climate Change (IPCC) [2]. It is a 32 promising option to maintain fossil fuels as a global central energy contributor and is expected to 33 progress in lab scale, pilot scale, demonstration scale and commercial scale [3]. Chemical absorption 34 technology, which has been used to remove CO2 from natural gas, is reached to the commercial phase 35 of development and several pilot-scale projects have been conducted since decades ago [4,5]. To date, 36 the capture technologies that have been demonstrated in pilot plant scale or higher are classified as 37 (i) post-combustion, (ii) pre-combustion, and (iii) oxy-fuel combustion [6]. Among the current 38 available technologies, CO<sub>2</sub> capture from post-combustion emission gases is the most common 39 technology in power plants and industries. Effective methods to separate, capture, store and convert 40 CO2 have attracted increasing attention [7]. The current technology for CO2 capture is based on 41 absorption in aqueous organic amines [8]. However, the intensive energy consumption, equipment 42 corrosion and toxicity, make exploration of new materials for CO<sub>2</sub> capture highly demanding [9].

Solid adsorption is an alternative process to solvent-based chemisorption, which can be applied for a wide range of temperature and pressure conditions. Also, it can be designed for both postcombustion and pre-combustion applications. One of the challenges in this area relates to the manufacturing of new adsorbents that enables enhancing good diffusion kinetics and providing 47 improved performance [10]. In this regards, porous materials with high porosity, chemical and 48 thermal stability are promising solid adsorbents for capturing CO<sub>2</sub> [11,12]. Among various kinds of 49 porous materials, such as zeolites [13,14], porous polymers [15], metal-organic framework [16,17], 50 activated carbons [18], porous BN has become one of the new classes of CO<sub>2</sub> adsorbents. Due to 51 intrinsic properties of boron nitride such as high thermal and oxidation stability, large surface area 52 and polarity, porous boron nitride is a desired alternative for CO<sub>2</sub> capture [19,20]. However, 53 according to theoretical and experimental works, the interaction between pristine BN and CO<sub>2</sub> is very 54 weak due to electron deficient nature of boron atoms and the Lewis acidic nature of CO<sub>2</sub> [21,22,23]. 55 To overcome such disadvantages, some strategies in view of tuning the structure, pore size and 56 charge state of the BN have been studied. Nag et al. [24] made a few-layered BN by controlled 57 chemical synthesis. Similarly, the effects of different proportions of urea on the number of BN layers 58 indicate that the sample with the lowest layer thickness shows a high CO<sub>2</sub> uptake. Xiao et al. [25] 59 reported a few layered porous boron nitride nanosheets (BNNs) by using magnesium diboride 60 (MgB<sub>2</sub>) as a dynamic template. However, the CO<sub>2</sub> uptake of their work was relatively low (0.45 61 mmol/g), while it demonstrated excellent selectivity of CO2 over N2. Yang et al. [26] developed a three 62 dimensional (3D) functionalized porous BN with flower-like morphology and high specific surface 63 area ~1114 m²/g , and it shows 1.69 mmol/g CO2 capture capacity at 1 bar and 273 K. It is also 64 interesting to note that in most studies, element doping, and surface functionalization of boron nitride 65 is critical to increase the CO2 adsorption capacity. For instance, Chen et al. [27] produced ultrahigh 66 microporous volume of boron carbon nitride (BCN) by doping carbon into hexagonal structure of 67 BN. The performance of BCN material displays increased relative to pure BN for CO2 uptake and 68 reached up to 3.91 mmol/g at 298 K and ambient pressure. In addition, Huang et al. [28] enriched the 69 interaction between electron-deficient boron atoms in BN and CO<sub>2</sub> molecules by aminopolymer-70 functionalized BN nanosheets. Therefore, porous structure and surface chemistry features of BN-71 based materials play crucial roles in CO<sub>2</sub> adsorption performance and it remains a challenge to control 72 the parameters of boron nitride porosity and develop novel structures with high capacitive 73 performance for carbon capture.

In this work, with the aim of achieving high CO<sub>2</sub> adsorption capacity of porous BN through tuning the porosity, BN structure has been modified in the presence of P123 as a structure directing agent. The synthesized BN-P123 possesses high specific surface area and pore volume compared to pristine BN, which results in an enhanced CO<sub>2</sub> adsorption capacity.

#### 78 2. Materials and Methods

79 Metal-free modified BN was synthesized on a two-step method of solvent evaporation and high 80 thermal decomposition [29]. Boric acid (ACS reagent, Sigma-Aldrich), urea (for synthesis, Sigma-81 Aldrich) and Pluronic P123 triblock copolymers (Mn ~5800, Sigma-Aldrich) based on poly(ethylene 82 glycol)-poly(propylene glycol)-poly(ethylene glycol) were used as raw materials. In detail, boric acid 83 and urea with molar ratio 1:30 were added into 50 ml deionized water. Then 0.5 g P123 was 84 introduced into the mixture solution and heated up at 338 K under vigorous stirring. A white 85 precipitate was obtained after complete evaporation of water at that temperature. Afterwards, the 86 precursor was dried into the oven for 24 hours and annealed at 1173 K under N<sub>2</sub> gas (the flow rate is 87 150 ml/min) for 3 hours. Finally, modified porous BN was collected and marked as BN-P123. For 88 comparison, pristine BN was also prepared following the same procedure without adding any 89 surfactant.

The morphology of samples was collected scanning electron microscope (SEM, PEMTRON PS-230) in secondary electron mode (SE detector) at 10 kV. The microscope was equipped with an energy-dispersive X-ray (EDX, OXFORD X-act) with energy resolution at 5.9 keV for the local elemental analysis. The samples were ground, deposited on carbon tape, and coated (BIO-RAD sputter coater) with gold to reduce charging in the microscope.

95X-ray diffraction (XRD) was performed to evaluate the structural properties using an X-ray96diffractometer (Bruker D8 Advance) in reflection mode. The diffraction patterns run at an anode97voltage of 40 kV and an emission current of 40 mA using monochromatic Cu Kα radiation ( $\lambda$ = 1.5419878 Å). The intermolecular bonding and chemical properties of the materials were characterized by

99 Fourier-transform infrared (FTIR) spectroscopy. The spectra were recorded on a Nicolet Avatar 370

- 103 thermal stability of samples was performed using thermal gravimetric analysis (TGA), Ultra micro
- balance, Mettler Toledo. An alumina pan was loaded with 5-10 mg of sample material and heated
- 105 from room temperature to 1273 K at a rate of 10 K/ min in air atmosphere.
- 106 Nitrogen isotherms were obtained using a porosity analyzer (Micromeritics ASAP2060) at 77 K.
- 107 The surface areas of the samples were calculated using the Brunauer–Emmett–Teller (BET) method.
- 108 The total volume of pores was calculated using the software from the BJH adsorption calculation.
- The CO<sub>2</sub> adsorption and desorption performance were evaluated with a TGA/DSC 3+ micro balance,
   Mettler Toledo. The CO<sub>2</sub> capture capacity of the sorbents were determined by measuring the mass
- 111 uptake of the sample during the CO<sub>2</sub> adsorption.

## 112 3. Results and discussion

## 113 3.1. Sample characterisation

The morphology of samples was studied using scanning electron microscopy (SEM). The pristine BN exhibited an ultrathin flake-like morphology, as displayed in (Figures 1a–b). BN-P123 presented a cloud-like sheet structure with interconnected network of porous structure (Figures 1c– d). Moreover, the formation of pristine BN and BN-P123 were studied by EDX and elemental mapping (Figures 2a–b). It is obvious that the product is mainly comprised of boron (B), nitrogen (N) and oxygen (O). As shown in Figure 2b, BN-P123 has very little increase in oxygen content compared to pristine BN. This is due to the introduction of P123 into BN formation.

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122 Figure 1. SEM images of (a, b) pristine BN; (b, c) BN-P123. The insert shows the interconnected porous

123 structure.





(b)

125 Figure 2. EDX spectra of (a) pristine BN; (b) BN-P123 and the insert show the elemental mapping and

126 corresponding atomic % of elements.

127 The structure of samples was analysed and Figure 3a depicts the XRD patterns of the prepared 128 samples. The results show two characteristic peaks around ~  $^{\circ}25$  and ~  $^{\circ}42$  which are attributed to 129 the (002) and (100) crystal planes of hexagonal boron nitride, respectively [30]. Additionally, the 130 peaks of both samples confirmed the poor crystallization with the presence of turbostratic material. 131 Compared with pristine BN, the (002) diffraction peaks of BN-P123 shifted to a lower angle, which 132 promote more disordered hexagonal boron nitride (h-BN) structure. More details of the chemical 133 features were supported by Fourier transform infrared (FT-IR) spectroscopy and the results are 134 shown in Figure 3b. All peaks exhibited two main characteristic bands of boron nitride at ~ 1365 and 135 800 cm<sup>-1</sup> corresponding to in-plane B-N transverse optical mode and out-of-plane B-N-B bending 136 mode, respectively [31]. No major chemical differences could be observed between the samples. 137 Thermal stability of samples was performed in air atmosphere (rate=10 K/min) and shown in Figure 138 3c. The TG thermograms reflected the weight loss of 19% for both samples, representing the removal 139 of moisture adsorbed on the material surface. After 573 K the lines show the thermal stability nature 140 of BN at high temperature and there was no significant weight loss up to 1273 K. The UV-vis 141 absorption collected in Figure 3d indicates a highly transparent from visible to UV wavelength [32]. 142 The maximum absorption peak around 209 nm, is attributed to intrinsic excitation absorption band 143 of h-BN. Besides, small humps around 265 and 352 nm were detected in BN-P123, corresponding to 144 impurities. So far, above all characteristic results on both samples, it was found that the structure and 145 chemical nature of BN has not changed after modifying BN with the surfactant compound and both 146 samples revealed almost similar features. We assumed that the utilisation of block copolymer (P123) 147 might affect the textural properties of pristine BN.

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#### 150 3.2. Textural Analysis

151 The textural properties of pristine BN and BN-P123 adsorbent were obtained by nitrogen 152 adsorption/desorption isotherm at 77 K and the results are summarized in Table 1. As indicated in 153 Figure 4a, the resulting isotherms display a typical type II curve based on IUPAC classification and 154 type H3 hystersis loop in the partial pressure range 0.4-1.0, which indicates the presence of mesopores 155 and slit shape pores [33]. There is a trivial rise in N2 adsorption-desorption isotherm of BN-P123 at 156 lower pressure ( $P/P^{\circ} < 0.25$ ), which is caused by presence of micropores [34]. The non-local density 157 functional theory (NLDFT) was used to calculate pore size distributions. As expected, the *dV/dW* pore 158 volume of BN-P123 increases dramatically with abundance of micropores (Figure 4b). Meanwhile, 159 using the Brunauer-Emmett-Teller (BET) method [35], the specific surface areas are calculated 160 showing that the BET surface area of BN-P123 is much higher than pristine BN (Table 1). It is clear 161 from the above measurements that the P123 introduced into precursors during the fabrication process 162 is highly effective on the microscale structure of porous BN and improves surface area and pore 163 volume of the sample.



164 Figure 4. Nitrogen adsorption-desorption isotherm of the pristine BN and BN-P123 (a); NLDFT pore size

165 distribution curves of the same samples (b).

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Table 1. Textural properties and CO<sub>2</sub> capacity (298 K) of prepared materials.

sample	Sвет <sup>1</sup> (m <sup>2</sup> /g)	V <sub>total<sup>2</sup></sub> (cm <sup>3</sup> /g)	CO2 Uptake (mmol/g)	
Pristine BN	102	0.46	2.00	
BN-P123	476	0.83	2.69	

167 <sup>1</sup> Specific surface area (m<sup>2</sup>/g) obtained by Brunauer-Emmett-Teller (BET) method. <sup>2</sup> Total pore volume (cm<sup>3</sup>/g)

168 calculated at  $P/P_0 = 0.99$ .

## 169 3.3. Gas adsorption analysis

170 Due to the relatively large porosity of samples, the modified sample BN-P123 and pristine BN 171 were assessed for  $CO_2$  capture under ambient conditions. The adsorption capacities of pure  $CO_2$  on 172 pristine BN and BN-P123 were determined by thermogravimetric analysis. Prior to the sorption test, 173 samples were dried at 393 K with a heating rate of 10 K/min from 298 K under flowing N<sub>2</sub> (150 174 ml/min) for 6 hours and then allowed to cool to the temperature at which the sorption is carried out 175 at 298 K with a heating rate of 10 K/min. When the sorption temperature was reached, samples were 176 stabilized under flowing N<sub>2</sub> (99,999%,150 ml/min) for 15 min and then N<sub>2</sub> flow was switched to pure 177 CO<sub>2</sub> (99,999%, 50 ml/min) flow for 6 hours. The mass uptake during this stage was interpreted as the 178 CO<sub>2</sub> capture capacity. Figures 5 and 6 (a-c) illustrate the setup of experiments and the TG profiles of 179 CO<sub>2</sub> uptake along with weight change results respectively.





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Figure 5. Illustration of CO<sub>2</sub> adsorption experiment.



182 Figure 6. TG profiles of CO<sub>2</sub> adsorption on Pristine BN and BN- P123 (a); TG profiles of weight change on the

184 The experiments reveal that pristine BN shows a CO2 capacity of 2 mmol/g at 298 K, which 185 enhanced with the addition of P123 copolymer (2.69 mmol/g for BN-P123 at 298 K) (Table 1). The 186 augmented CO<sub>2</sub> adsorption on BN-P123 sample is undoubtedly as a result of the porous morphology 187 of modified BN, whereas pristine BN exhibits lower porosity. It is well-known that higher 188 microporosity provides more active sites and storage space to boost the adsorption performances 189 [36,37,38]. To the best of our knowledge, the result of our synthetic method is higher than other 190 modification methods such as (3D) BCNO structure (1.8 mmol/g) at 298 K as reported by Lopez-Salas 191 et al. [39] and closer to porous BCN (2.49 mmol/g) at 298 K and porous BN fiber (2.85 mmol/g) at 273 192 K [40,41]. However, the use of melamine in porous BCN as well as BN fiber leads to the presence of

<sup>183</sup> same samples (b,c).

193 carbon content and consequently decreases the stability upon exposure to ambient air. The effect of 194 high-temperature treatment in different gases was studied in [42], where porous BN fibers treated in 195 NH<sub>3</sub> gas at 1673–1773 K enhanced CO<sub>2</sub> adsorption capacity (from 0.45 mmol/g to 1.6 mmol/g). 196 Nonetheless, this outcome is not a satisfactory result. One possible insight of this finding is that 197 pyrolysis temperature and carrier gas does not lead to a remarkable change in CO<sub>2</sub> uptake. In 198 addition, directing triblock co-polymer was also used to design a hierarchical carbon sorbent [43]. 199 This hierarchical structure is desirable for  $CO_2$  capture as it shows superior capacity (4.5 mmol/g) 200 under ambient conditions. It is noteworthy that at lower pressures (bar  $\leq$  1), the density of pore 201 volume, especially the micropores volume plays a critical role in capturing CO<sub>2</sub> [44]. Therefore, high 202 surface area and porosity of BN-P123 has brought about a positive adsorption interaction.

It is to be noted that for the pristine BN (without adding surfactant) the CO<sub>2</sub> adsorption capacity on our sample exceeds those materials reported by Marchesini et al. [45,46]. Although they accomplished highly porous boron nitride with a high surface area (>1900 m<sup>2</sup>/g), the CO<sub>2</sub> adsorption capacity of their sample was up to 1.6 mmol/g under ambient condition. Given the results of our approach, one can conclude that though the specific surface area (SSA) is one of the main factors to increase the CO<sub>2</sub> capacity, obtaining a high SSA e.g. (>1000 m<sup>2</sup>/g) does not lend itself to an increased CO<sub>2</sub> capacity.

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### 211 5. Conclusions

212 In summary, the porosity of boron nitride has been successfully tuned by introducing the 213 triblock copolymer surfactant (P123) during the preparation process of BN precursors. In particular, 214 we found that by utilizing the P123 into boron and nitrogen precursors leads to improved CO<sub>2</sub> 215 adsorption capacity up to 2.69 mmol/g, as compared with pristine porous BN which was found 2.00 216 mmol/g. Furthermore, based on the structural properties and morphology results obtained in this 217 work, we determine that both samples share almost similar chemical features. However, the porosity 218 of samples revealed a remarkable change and for the sample BN-P123 (0.83 cm<sup>3</sup>/g), it was virtually 219 quadrupled as a result of the modification. This significant change is attributed to the formation of 220 more gases during the decomposition process, thereby creating higher porosity levels. A natural progression of this work is to describe how textural property parameters of BN lead to higher 221 222 interaction between BN and CO2 molecules. Our work could be extended by exploring other 223 parameters (e.g. electronic features or surface chemistry of BN), which could influence on BN to 224 capture more CO<sub>2</sub>. All in all, our analyses demonstrate the applicability of metal-free modification of 225 BN for enhanced capacity of pure CO<sub>2</sub>.

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