1	The density of ambient black carbon retrieved by a new method:
2	implications to CCN prediction
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25 Abstract.

The effective density of black carbon (BC) is a crucial factor relevant to its aging 26 degree that would add uncertainty in evaluating its climate effect. Here, we have 27 developed a new method to retrieve the effective density of internally mixed BC in the 28 atmosphere combining field observations conducted during 15 November -14 29 December 2016 in urban Beijing with the Köhler theory. The uncertainty of the retrieval 30 method was evaluated within ± 30 %, which is primarily caused by assumptions of the 31 32 hygroscopic parameter of organics and the fraction of primary organic aerosols in nonhygroscopic or hygroscopic mode. Using the method, we obtain that the ambient 33 internally mixed BC, accounting for 80±20 % of total BC aerosol particles, is retrieved 34 with a campaign mean density of 1.1±0.6 g cm⁻³ during the observed periods. The 35 retrieved result is comparable with that reported in the Literature. By applying a lower 36 (0.14 g cm⁻³) and upper (2.1 g cm⁻³) limit of the retrieved BC density in cloud 37 condensation nuclei (CCN) number concentrations (N_{CCN}) estimation, we derived that 38 39 neglect of such variations in BC density would lead to an uncertainty of -28 %~11 % 40 in predicting N_{CCN} at supersaturations of 0.23 % and 0.40 %. We also find that the N_{CCN} is more sensitive to the variations of BC density when it is <1.0 g cm⁻³. This illustrates 41 42 a necessity of accounting for the effect of BC density on CCN activity closer to source regions where the BC particles are mostly freshly emitted. The CCN closure achieves 43 when introducing the retrieved real-time BC density and mixing state. This study 44 provides a unique way of utilizing field measurements to infer ambient BC density and 45

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highlights the importance of applying variable BC density values in models when
 predicting CCN and assessing its relevant climate effect.

52 1 Introduction

53 Black carbon (BC) aerosols, as the major absorber of solar radiation, play a vital role in energy budget and climate of the earth-atmosphere system by affecting the 54 radiative forcing and cloud properties (Flanner et al., 2007; Ramanathan and 55 Carmichael, 2008). The light-absorbing capability induced by BC is related to its 56 density and morphology (Zhang et al., 2008; Rissler et al., 2014), which can be 57 modified after mixing with other atmospheric aerosol particles (Khalizov et al., 2009; 58 59 Xue et al., 2009). Changes in its physicochemical properties would also regulate its ability to serve as cloud condensation nuclei (CCN) and further indirectly affect the 60 61 radiative balance by affecting the clouds process (Yuan et al., 2008; Wang et al., 2011). Owing to the complex evolution of the mixing state, density and morphology of BC, 62 the contribution of BC particles to CCN budgets is still not well understood. 63

BC particles, with diesel vehicles, industrial and residential coal combustion as major sources, are ubiquitous in urban environments (Bond et al., 2013; Dameto et al., 2017; Li et al., 2017; Liu et al., 2019a). The mixing state of BC describes the distribution of the bare BC and coating masteries among the aerosol population. Typically, freshly generated BC exists in the form of chain aggregates and initially uncoated, which is known as externally mixed BC (Ex-BC). When the BC particles were emitted, they generally mix with other materials by condensation, coagulation,

71	and other processes (Riemer et al., 2004; Zhang et al., 2008; Liu et al., 2013; Zhang et
72	al., 2020a), forming the internally mixed BC (In-BC) particles consisting of BC core
73	and other chemical components (Cheng et al., 2006; Zhang et al., 2016). The BC
74	structure would be more compact with regular shapes (Pagels et al., 2009; Zhang et al.,
75	2008; Wang et al., 2017), and the effective density of internally mixed BC are changed
76	accordingly with the reconstruction (Liu et al., 2019b). The density and morphology of
77	BC particles are closely related to its sources, mobility size, coating thickness, coating
78	material and its chemical composition (Zhang et al., 2008; Pagels et al., 2009; Peng et
79	al., 2016; Zhang et al., 2022). A wide range of BC density has been reported in previous
80	studies (Lide 1992; McMurry et al., 2002; Park et al., 2004; Kiselev et al., 2010). Recent
81	field measurements have indicated that the average BC density is ~1.2 g cm ⁻³ in the
82	ambient atmosphere (Zhang et al., 2016). Field measurements have also indicated that
83	a considerable fraction of externally mixed/uncoated BC exists (Clarke et al., 2004;
84	Cheng et al., 2012), although a higher proportion of internally mixed/aged BC particles
85	in the ambient atmosphere were observed (Schwarz et al., 2008; Massoli et al., 2015;
86	Chen et al., 2020). In climate models, the BC was generally assumed completely
87	internally, mixed and treated to have a void-free spherical structure and a density value
88	of 1.8 g cm ⁻³ (Bond et al., 2013). This may lead to bias in estimating the climate effect
89	driven by BC.
90	Previous study based on a case study show that when the aging degree of ambient
91	particles is low, the BC density (~1.8 g cm ⁻³) under the spherical assumption will lead

92 to the overestimation of particle hygroscopicity by 40-50 % and the overestimation can

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95	be explained almost 100 % using the effective density of fresh BC (~0.45 g cm ⁻³) (Fan
96	et al. 2020). This indicates the importance of using reasonable BC density values in the
97	calculation of particle hygroscopicity. In addition, when estimating the CCN number
98	concentration, a significant bias of $-35 \% \sim +20 \%$ was found due to the assumption of
99	particle mixing state (Ren et al., 2018). However, these studies have not yet accounted
100	for such impact of BC density and mixing state on CCN prediction due to lack of real
101	time measurement data.

The mixing state and the density of BC particles are usually directly measured by 102 several techniques, such as an integrated system of a volatility tandem differential 103 mobility analyzer and a single particle soot photometer (VTDMA-SP2) (Zhang et al., 104 2016), or a differential mobility analyzer with a SP2 (DMA-SP2) (Olfert et al., 2007; 105 106 Rissler et al., 2014; Wu et al., 2019), and a differential mobility analyzer-centrifugal 107 particle analyzer-single-particle soot photometer (DMA-CPMA-SP2) system (Liu et 108 al., 2019b; Yu et al., 2020), etc. However, such techniques or measurements are not available in many previously conducted filed campaigns. In this study, we develop a 109 novel method for retrieving the mixing state and effective density of ambient BC 110 particles by combining field measured hygroscopic growth factor and aerosol chemical 111 composition and Köhler theory (Petters and Kreidenweis, 2007). The uncertainty of the 112 113 new retrieval method was evaluated. The retrieved results were also compared and validated with existing observations. In addition, the effect of BC density and mixing 114 state on prediction of CCN number concentrations is further evaluated through a 115 116 sensitivity and closure test by accounting for the retrieved real-time variations of BC

117 density and mixing state.

118 2 Field measurements and methodology

119 2.1 Field measurements

Measurements in this study were conducted from 15 November to 14 December 120 121 2016 at a typical urban site of Beijing (39.97°N, 116.37°E, 49 m above sea level). The 122 site locates at the Institute of Atmospheric Physics, Chinese Academy of Sciences, which is mainly influenced by the surrounding cooking, road traffic and residential coal 123 124 burning emissions during the home heating periods (Sun et al., 2016). The detailed information about the sampling site was presented in previous studies (Sun et al., 2015; 125 Zhang et al., 2019). The number concentration of condensation nuclei (CN) at each size 126 was measured by a scanning mobility particle sizer, which is equipped with a 127 differential mobility analyzer (DMA; model 3081, TSI) and a condensation particle 128 129 counter (CPC; model 3772, TSI). Subsequently, the mono-dispersed particles were introduced into a Droplet Measurement Technologies CCN counter (CCNc, DMT; 130 131 Lance et al., 2006) to measure CCN number concentration. A hygroscopic tandem differential mobility analyzer (HTDMA) system was used to measure the hygroscopic 132 growth factor (Gf) (Tan er al., 2013). Here, four diameters of 40, 80, 110, 150, and 200 133 nm are selected in the campaign. Gf is defined as the ratio of the mobility diameter at 134 the given RH to the dry diameter (Petters and Kreidenweis, 2007). The nonrefractory 135 submicron aerosol chemical composition was measured by an Aerodyne high-136 resolution time-of-flight aerosol mass spectrometer (HR-AMS; Xu et al., 2019), 137 6

138 including sulfate, nitrate, ammonium, chloride, and organics. Two factors, including a non-hygroscopic primary organic aerosol (POA) and hygroscopic secondary organic 139 aerosol (SOA) were classified by positive matrix factorization (PMF) with PMF 140 algorithm (v4.2) method (Paatero and Tapper, 1994) and followed the procedures 141 reported in Ulbrich et al. (2009). The refractory black carbon mass loading was 142 143 measured by an aethalometer (model AE33, Magee Scientific Corporation). Both the nonrefractory materials and BC mass concentration were measured with diameters < 144 145 $1.0 \,\mu\text{m}$. The detailed description of the instrument operation and data process have been described in details elsewhere (Ren et al., 2018; Xu et al., 2019; Zhang et al., 2019; Fan 146 et al., 2020). 147

148 2.2 Retrieving the mixing state and density of BC

149 2.2.1 Retrieving the mixing state of BC

150 The Gf probability distribution function (Gf-PDF) for a specified diameter can be 151 retrieved firstly based on the TDMAinv algorithm (Gysel et al., 2009). The κ -PDF can 152 be further calculated based on the Gf-PDF (Fan et al., 2020). Size-resolved κ is derived 153 using κ -Köhler theory based on hygroscopic growth factor (Gf) (Petters and 154 Kreidenweis, 2007),

155
$$\kappa_{gf} = (\mathrm{Gf}^3 - 1) \cdot \left[\frac{1}{\mathrm{RH}} \exp\left(\frac{4\sigma_{s/a}M_{\mathrm{w}}}{\mathrm{RT}\rho_{\mathrm{w}}\mathrm{D}_{\mathrm{d}}\mathrm{Gf}}\right) - 1 \right]$$
(1)

where Gf is hygroscopic growth factor, RH is the relative humidity in the HTDMA (90 %), $D_{\rm d}$ is the dry diameter, $\sigma_{\rm s/a}$ is assumed to be the surface tension of pure water,

158 R is the universal gas constant, T is the temperature, M_w and ρ_w is the molecular mass, and the density of water, respectively. 159

The *k*-PDF patterns of particles in different sizes always present two modes: nearly 160 161 hydrophobic (NH) mode with $\kappa_{gf} \leq 0.1$ and more hygroscopic (MH) mode with κ_{gf} > 0.1 (Fig. S1). Firstly, based on the κ -PDF patterns, the number fraction (NF) of the total 162 163 nearly hydrophobic group with the boundary of [0, 0.1] was calculated according to the following equation: 164

165
$$NF = \int_0^{0.1} c(\kappa, D_p) d\kappa$$
(2)

166 here, the κ -PDF, represented by c (κ , D_p), was normalized as $\int c(\kappa, D_p) d\kappa = 1$, where κ can be replaced by κ_{gf} , D_p is the selected electrical mobility diameter in the campaign. 167 The nearly hydrophobic mode consists of both externally mixed POA (Ex-POA or 168 169 bare POA) and externally mixed BC (Ex-BC). Since the number fraction of the nearly-170 hydrophobic POA would change with the emission and aging processes, in this study, 171 we have applied different values for the number fractions of hydrophobic POA (NH-172 POA) under clean (91 %), moderately polluted (70 %), and heavily polluted conditions (31 %) by referring the literature (Liu et al., 2021a), as shown in Fig. S2. The number 173 concentration of Ex-BC was then calculated using the total number fraction of NH 174 mode minus the number of NH-POA. 175

176
$$N_{POA-containing} = N_{total} \times NF_{POA-containing}$$

$$N_{POA-containing} = N_{total} \times NF_{POA-containing}$$

$$N_{bare-POA} = N_{POA-containing} \times NF_{bare-POA}$$

$$N_{Ex-BC} = N_{NH} - N_{bare-POA} \tag{3}$$

179 where NPOA-containing and NFPOA-containing are the number concentration and fraction of

181POA are the number concentration and fraction of bare POA partice number of nearly, hydrophobic group.182number of nearly, hydrophobic group.183The number size distribution of the externally mixed BC (n_{Ex} 184calculated based on the particle number size distribution (PNS)185fraction of the hydrophobic mode of BC (NF_{Ex-BC}) as follows:186 $n_{Ex-BC}(\log Dp) = NF_{Ex-BC} \times n (\log Dp)$ 187where $n (\log D_p)$ is the function of the aerosol number size distribution188mobility diameter.189By assuming that the particles are spherical (Rader and McMu190size distribution of Ex-BC (M_{Ex-BC}) was obtained as follows:	icles, and $N_{\rm NH}$ is the $_{\rm x-BC}$ (log $D_{\rm p}$)) can be D) and the number	删除了: -
182number of nearly_hydrophobic group.183The number size distribution of the externally mixed BC (n_{Ex} 184calculated based on the particle number size distribution (PNSI185fraction of the hydrophobic mode of BC (NF_{Ex-BC}) as follows:186 $n_{Ex-BC}(\log Dp) = NF_{Ex-BC} \times n (\log Dp)$ 187where $n (\log D_p)$ is the function of the aerosol number size distribution188mobility diameter.189By assuming that the particles are spherical (Rader and McMu190size distribution of Ex-BC (M_{Ex-BC}) was obtained as follows:	_{x-BC} (log D_p)) can be D) and the number	删除了:-
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184 calculated based on the particle number size distribution (PNS) 185 fraction of the hydrophobic mode of BC (NF_{Ex-BC}) as follows: 186 $n_{Ex-BC}(\log Dp) = NF_{Ex-BC} \times n (\log Dp)$ 187 where $n (\log D_p)$ is the function of the aerosol number size distribution of the aerosol number size distribution of the particles are spherical (Rader and McMu 190 size distribution of Ex-BC (M_{Ex-BC}) was obtained as follows: 181 $M_{ex-BC}(\log Dp) = \frac{\pi}{2} p_{ex}^{3} m_{ex-BC} (\log Dp)$	D) and the number	
185fraction of the hydrophobic mode of BC (NF_{Ex-BC}) as follows:186 $n_{Ex-BC}(\log Dp) = NF_{Ex-BC} \times n (\log Dp)$ 187where $n (\log D_p)$ is the function of the aerosol number size dis188mobility diameter.189By assuming that the particles are spherical (Rader and McMu190size distribution of Ex-BC (M_{Ex-BC}) was obtained as follows:101 $M_{ex} = (l_{Ex}-BC) = \frac{\pi}{2} D_{ex}^{3} = m_{ex} = (l_{Ex}-BC)$		
186 $n_{\text{Ex-BC}}(\log Dp) = NF_{\text{Ex-BC}} \times n (\log Dp)$ 187 where $n (\log D_p)$ is the function of the aerosol number size dis 188 mobility diameter. 189 By assuming that the particles are spherical (Rader and McMu 190 size distribution of Ex-BC ($M_{\text{Ex-BC}}$) was obtained as follows: 101 $M_{\text{Ex-BC}} = Dr$)		
187 where <i>n</i> (log D_p) is the function of the aerosol number size dis 188 mobility diameter. 189 By assuming that the particles are spherical (Rader and McMu 190 size distribution of Ex-BC (M_{Ex-BC}) was obtained as follows: 101 $M_{Ex-BC} = D_{Ex}$	(4)	
188 mobility diameter. 189 By assuming that the particles are spherical (Rader and McMu 190 size distribution of Ex-BC (M_{Ex-BC}) was obtained as follows:	stribution, D_p is the	
By assuming that the particles are spherical (Rader and McMu size distribution of Ex-BC (M_{Ex-BC}) was obtained as follows:		
190 size distribution of Ex-BC ($M_{\text{Ex-BC}}$) was obtained as follows:	urry, 1986), the mass	
$M = (1 - D_{T})^{\pi} D_{3}^{3} \cdots (1 - D_{T})$		
$M_{\text{Ex-BC}}(\log Dp) = -\frac{1}{6}D_p \cdot \rho n_{\text{Ex-BC}}(\log Dp)$	(5)	
192 where D_p is the mobility diameter, ρ is the effective density of Ex-	-BC, and $n_{\text{Ex-BC}}$ (log	
193 $D_{\rm p}$) is the function of the number size distribution of Ex-BC, respect	tively. By reviewing	
and summarizing the existing results, we show that typical value	es of density for the	删除了: abou
195 freshly emitted or externally mixed BC observed in the winter of urb	ban Beijing or North	
196 China Plain span over 0.14-0.50 g cm ⁻³ , with mean of $\sim 0.40\pm0.10$) g cm ⁻³ (Fig. S3), in	删除了: spans
197 the size range of 100 to 300 nm, where the mass concentration of e	externally mixed BC	
198 <u>mostly concentrated</u> (Geller et al., 2006; Peng et al., 2016, 2017; V	Wu et al., 2019; Liu	删除了: accour
et al., 2020; Zhao et al., 2022). Therefore, an average $\rho_{\text{Ex-BC}}$ of 0.4	g cm ⁻³ was used for	
200 calculating the mass concentration of externally <u>mixed BC in our</u>	r study. Uncertainty	删除了: -mixed exhibits that th
analyses due to the variations of $\rho_{\text{Ex-BC}}$ were given in section 2.3.		average deviati

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:7: accounted for a large proportion in urban Beijing

:-mixed BC in our study. The uncertainty analysis bits that the variations of the $\rho_{\text{Ex-BC}}$ could lead to an age deviation of ± 10 % in the calculating In-BC density . 3e) by increasing the $\rho_{\text{Ex-BC}}$ from 0.1 to 0.6 g cm⁻³, showing a small impact on the retrieved result.

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The mass size distribution of Ex-BC was fit using the log-normal distribution as shown in Fig. S4 (Wu et al., 2017; Liu et al., 2019a; Zhao et al., 2022). Thus, the bulk 212 mass concentration of Ex-BC can be calculated from the integration of the mass size 213 distribution: 214

215
$$m_{\text{Ex-BC}} = \int_{D_{start}}^{D_{end}} M_{\text{Ex-BC}}(\log D_p) \ d \log(D_p) \tag{6}$$

$$m_{\rm In-BC} = m_{\rm BC} - m_{\rm Ex-BC} \tag{7}$$

where D_{start} and D_{end} are the lower and upper size limit, $M_{\text{Ex-BC}}$ (log D_{p}) is the function 217 218 of the Ex-BC mass size distribution. We then obtained the bulk mass concentration of internally mixed BC (m_{In-BC}) by subtracting m_{Ex-BC} from the bulk BC mass 219 220 concentration measured by AE33 in equation 7. It should be noted that the mass concentration of BC obtained from AE33 based on aerosol light absorption may lead 221 222 some uncertainty, as has been further addressed in Section 2.3.

2.2.2 Retrieving the density of BC 223

224 For retrieval of the density of BC, the principal idea is to use the measured κ_{gf} to calculate the density of BC based on the Zdanovskii-Stokes-Robinson (ZSR) mixing 225 226 rule (Stokes and Robinson, 1966; Zdanovskii, 1948) with the chemical composition measured by AMS (Petters & Kreidenweis, 2007). In the retrieval, several aspects are 227 228 concerned. First, since the ZSR rule assumes the aerosol particles are internally mixed, 229 the κ_{gf} value of the more MH mode (κ_{gf-MH}) is thus applied for retrieving the density of 230 internally mixed BC. Second, since the size distribution of BC number concentration is usually with peaks between 100 and 200 nm (Liu et al., 2019a; Yu et al., 2020; Zhao et 231

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233	al., 2022), the κ_{gf-MH} value of particles in accumulation mode was averaged and applied	
234	for the retrieval. Previous studies showed an independence of $\kappa_{gf\text{-}MH}$ on particle size	
235	when the $D_p > 100$ nm during the campaign period (Fan et al., 2020). Therefore, the	
236	average of κ_{gf-MH} in accumulation mode is reasonable for the determination of the In-	
237	BC density. In addition, because the inversion including measurements from HTDMA	
238	and HR-AMS, a total mass closure of the measured aerosol particles was conducted	
239	between the two techniques by comparing the mass concentration of $\ensuremath{PM}\xspace_1$ and the results	
240	are well consistent (Fig. <u>\$6</u>). The density of internally mixed BC (In-BC), $\rho_{\text{In-BC}}$ is then	删除了: S5
241	derived from the following equations:	
242	$\kappa_{gf-MH} = \kappa_{chem} = \sum_{i} \varepsilon_{i} \kappa_{i} = \frac{v_{inorg}}{v_{total}} \kappa_{inorg} + \frac{v_{SOA}}{v_{total}} \kappa_{SOA} + \frac{v_{In-POA}}{v_{total}} \kappa_{POA} + \frac{v_{In-BC}}{v_{total}} \kappa_{BC} $ (8)	
243	where κ_{gf-MH} is the hygroscopic parameter of the more hygroscopic (MH) mode, κ_{chem}	
244	is the hygroscopic parameter of aerosol particles in the mixed composition and can be	
245	calculated based on chemical volume fractions using a simple rule (Stokes and	
246	Robinson, 1966; Petters & Kreidenweis, 2007), κ_i is the hygroscopic parameter of each	
247	pure composition and $\boldsymbol{\varepsilon}_i$ is the volume faction of the individual components in the	
248	internally mixed particle. v_{inorg} , v_{SOA} and v_{In-POA} are the volume of the inorganic, SOA	删除了: internal-
249	and internally mixed POA species, and can be calculated as follows: $v_{inorg} = \frac{m_{inorg}}{\rho_{inorg}}$,	
250	$v_{SOA} = \frac{m_{SOA}}{\rho_{SOA}}$, and $v_{In-POA} = \frac{m_{In-POA}}{\rho_{POA}}$. v_{total} is the total volume of all the species and can be	
251	written as $v_{total} = \frac{m_{inorg}}{\rho_{inorg}} + \frac{m_{SOA}}{\rho_{SOA}} + \frac{m_{In-POA}}{\rho_{POA}} + \frac{m_{In-BC}}{\rho_{In-BC}}$. In equation (8), κ_{BC} and κ_{POA} are	
252	assumed to be 0. So, the total volume v_{total} can be further written as v_{total} =	移动了(插入) [1]
253	$\frac{v_{inorg}\kappa_{inorg} + v_{SOA}\kappa_{SOA}}{\kappa_{gf-MH}}$. The volume of internally mixed v_{In-BC} can be calculated as follows,	
254	$v_{in-BC} = \frac{v_{inorg}\kappa_{inorg} + v_{SOA}\kappa_{SOA}}{\kappa_{gf-MH}} - v_{inorg} - v_{SOA} - v_{In-POA}$	移动了(插入) [2] 移动了(插入) [3]
·		

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$$= \frac{\frac{m_{inorg}}{\rho_{inorg}}\kappa_{inorg} + \frac{m_{SOA}}{\rho_{SOA}}\kappa_{SOA}}{\kappa_{af-MH}} - \frac{m_{inorg}}{\rho_{inorg}} - \frac{m_{SOA}}{\rho_{SOA}} - \frac{m_{In-POA}}{\rho_{POA}}$$
(9)

258 Then, the $\rho_{\text{In-BC}}$ can be calculated based on its mass concentration and volume as follows:

$$\rho_{In-BC} = \frac{m_{In-BC}}{\left(\frac{m_{inorg}}{\rho_{inorg}} + \frac{m_{SOA}}{\rho_{SOA}} - \frac{m_{inorg}}{\rho_{inorg}} + \frac{m_{SOA}}{\rho_{SOA}} - \frac{m_{In-POA}}{\rho_{POA}}\right)}$$
(10)
(10)

where, $m_{\text{In-BC}}$ is the mass concentration of internally mixed BC, m_{inorg} and m_{SOA} are the 260 mass concentrations of the inorganic species and SOA, which are measured by the AMS. 261 $m_{\text{In-POA}}$ is the mass concentrations of internally mixed POA and can be calculated 262 263 through subtracting the mass fraction of NH-POA from the total mass concentrations 264 of POA. ρ_{inorg} , ρ_{SOA} and ρ_{POA} are the density of the inorganic species, SOA and POA. Since the AMS measures the concentrations of the organic and inorganic ions, including 265 SO₄²⁻, NO₃⁻, NH₄⁺, Cl⁻. Here inorganic species were derived by applying a simplified 266 ion pairing scheme (Gysel et al., 2007) to convert mass concentrations of ions to the 267

268 inorganic salts as follows:

270

259

 $n_{\rm NH_4NO_3} = n_{\rm NO_3^-}$

271
$$n_{\rm NH_4HSO_4} = \min(2n_{\rm SO_4^{2-}} - n_{\rm NH_4^+} + n_{\rm NO_3^-}, n_{\rm NH_4^+} - n_{\rm NO_3^-})$$

$$n_{(\mathrm{NH}_4)_2\mathrm{SO}_4} = \max(n_{\mathrm{NH}_4^+} - n_{\mathrm{NO}_3^-} - n_{\mathrm{SO}_4^{2^-}}, 0)$$

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272

$n_{\rm NH_4^+} + n_{\rm NO_3^-})$ (11)
$n_{\rm NH_4^+} + n_{\rm NO_3^-}$) (11)

where *n* represents the number of moles, and the mass concentrations were obtained by the number of moles times the molar mass of each inorganic salts. Because the value of the $n_{H_2SO_4}$ was zero in this campaign. Three inorganic salts including NH₄HSO₄, (NH₄)₂SO₄, and NH₄NO₃ were applied in our study. The densities for inorganic salts were taken from previous studies (Gysel et al., 2007; Wu et al., 2016). Here the densities for three inorganics are 1.78, 1.77 and 1.72 g cm⁻³, respectively. By summarizing the

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283	previous studies (Gysel et al., 2007; Dinar et al., 2006), 1.4 g cm ⁻³ was selected as the		
284	density of SOA (ρ_{SOA}). The density of POA (ρ_{POA}) is assumed to be 1.0 g cm ⁻³ for urban		
285	environments, which is similar to the <u>that of the</u> lubricating oil (Wu et al., 2016). <u>Since</u>		删除了: Co
286	the cooking organic aerosols represent a high contribution to POA in urban		
287	environments, we choose the mean density of the rapeseed oil and oleic acid (~0.85 g		删除了:ad
288	<u>cm⁻³)</u> (Reyes-Villegas et al., 2018), to evaluated the result as shown in section 2.3. The		删除了: for
289	values of κ for inorganic components are 0.56 for NH ₄ HSO ₄ , 0.48 for (NH ₄) ₂ SO ₄ and		删除了: w
290	0.58 for NH ₄ NO ₃ , along with the best-fit values for the three inorganic salts (Petters &		
291	Kreidenweis, 2007 and Gunthe et al., 2009). The κ_{SOA} is assumed to be 0.15 according		
292	to the field studies in urban areas (Chang et al., 2010; Kawana et al., 2016).		
293	Note that, the method fails to retrieve the BC density when organic accounts for a		删除了: ,th
294	large fraction (>60 %). This is because that a higher fraction of OA usually corresponds		删除了: org
295	to lower total volume of all the species (Fig. <u>\$7</u>), yielding negative values for $v_{\text{In-BC}}$		删除了: S6
296	introduced in equation <u>2</u> . As a result, 61 % of the data observed during the campaign		删除了:11
297	were valid for calculating the BC density.		
298	$v_{In-BC} = \frac{v_{Inorg} \kappa_{inorg} + v_{SOA} \kappa_{SOA}}{\kappa_{SOA}} - v_{In-POA}$ Similarly, the bulk density of BC ($\rho_{\text{bulk-BC}}$) is		上移了 [2]:
299	calculated with the same method as that for calculating the $\rho_{\text{In-BC}}$ When calculating the		上移了 [1]:
300	$\rho_{\text{bulk BC}}$ the bulk κ_{sf} value measured by HTDMA is applied assuming that all the aerosol		上移了 [3]:
000		$\langle \rangle $	删除了:
301	particles are internally mixed.	$\langle \rangle$	删除了: −≀
			删除了:
302	2.3 Uncertainties and limitations		删除了: wit

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	上移了 [2]: v _{In-BC}
	上移了 [1]: = $\frac{v_{inorg} + v_{SOAKSOA}}{\kappa_{gf-MH}}$
$\left(\right)$	上移了 [3]: -v _{In-POA}
$\left(\right)$	删除了:
	删除了: -v _{inorg} - v _{SOA}
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For the retrieval, the assumptions on the values of $\kappa_{\rm SOA}$, $\rho_{\rm POA}$, $\rho_{\rm SOA}$ and $\rho_{\rm Ex-BC}$ as

well as the fraction of primary organic aerosols in non-hygroscopic or hygroscopic mode would add uncertainty in the inferred values of ambient internally mixed BC density. For example, the freshly emitted POA particles might consistently be coated with the secondary particles during the aging process, resulting in changes of the $NF_{\rm NH}$ -POA. However, a real-time variation of the $NF_{\rm NH-POA}$ is not yet available due to the lack



et al., 2019). In addition, the value of κ_{SOA} spans largely due to the variability in the emissions of gas precursors and formation processes under different atmospheric conditions (Zhang et al., 2015; Liu et al., 2021b). Therefore, we examined the sensitivities of In-BC density to the variations of these factors, as exhibited in Fig. 1



352 decreases slightly as $\rho_{\text{Ex-BC}}$ increases (Fig. 2e), suggesting applying a larger $\rho_{\text{Ex-BC}}$ would derive smaller values for In-BC density. The In-BC density is insensitive to the 353 changes of the density of POA and SOA, showing an almost negligible effect on the 354 retrieved results (Fig. 2c and d). 355

356 The uncertainty analysis shows that, by comparing the results based on the mean fractions of the $NF_{\rm NH-POA}$ with a typical atmospheric observed range of 50-90 % for the 357 NF_{NH-POA} (Liu et al., 2021a), we show that the assumption on NF_{NH-POA} can lead to 358 relative deviations (uncertainty) of -17 <u>%-+</u>27 % for the retrieved BC density (Fig.3a).

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Figure 3. Relative deviations of the number fraction of nearly hydrophobic (NH) POA 361 to the In-BC density (a), the hygroscopic parameter of OA to the In-BC density (b), the 362 POA density to the In-BC density (c), the SOA density to the In-BC density (d), the 363 externally mixed BC density to In-BC density (e) and the combined deviations based 364 on multiple factors mentioned above (f). 365



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上移了 [4]: Figure 1. Sensitivities of In-BC density to the variations in the number fraction of nearly hydrophobic (NH) POA and hygroscopic parameter of OA (ksoa) (a), POA density (b), SOA density (c) and the externally mixed BC density (d).

上移了 [5]: Figure 2. Sensitivity of the In-BC density to variations in the number fraction of nearly hydrophobic (NH) POA (a), the hygroscopic parameter of SOA (b), the POA density (c), the SOA density (d), the externally mixed BC density (e) and the harmonic mean of multiple factors (f).



the hygroscopicity has been already well-understood (Petters and Kreidenweis, 2007), 384 385 the hygroscopicity of organic species varies largely due to the complexity in organic aerosol constituents. Therefore, the assumption of the values of κ_{SOA} will add the 386 uncertainty in the calculation of BC density. Previous studies have suggested that the 387 organics has a wide range of κ values ranging from 0.05 to 0.3 (Jimenez et al., 2009; 388 Mei et al., 2013). Thus, the sensitivity test has also been done to examine the effect due 389 to changes in κ_{SOA} on calculating the density of BC (Fig. 1a). The result shows that the 390 391 assumption of κ_{SOA} value can cause an average relative deviation of $-10 \frac{}{2}$ % in 392 calculating the density of In-BC (Fig. 3b).

393 However, the sensitivity test shows that the impact of both the ρ_{POA} and ρ_{SOA} variations on the BC density estimation is very small or even negligible (Fig. 1b, c). By 394 varying the ρ_{POA} from 0.85 to 1.5 g cm⁻³ and the ρ_{SOA} from 0.9 to 1.65 g cm⁻³ according 395 396 to the literatures (Noureddini et al., 1992; Alfarra et al., 2006; Reyes-Villegas et al., 397 2018; Cai et al., 2020; Kostenidou et al., 2007), the retrieval uncertainties in the BC 398 density are <u>both</u> within ± 5 % (Fig. 3c, d). For ρ_{Ex-BC} , it exhibits that the evolution of the $\rho_{\text{Ex-BC}}$ could lead to an average deviation of $\frac{-16\%}{-19}\%$ in calculating In-BC density 399 (Fig. 3e) when increasing the values of ρ_{Ex-BC} from 0.1 to 1.0 g cm⁻³, which represents 400 a typical range in ambient atmosphere (Wu et al., 2019; Liu et al., 2020). A combined 401 402 uncertainty (δ) caused by the multiple factors (δ_i), which is calculated by equation 12, 403 is -<u>26 %~+</u>29 % as shown in Fig. 3f.

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$$\delta = \int_{1}^{n} \sum_{i=1}^{n} \delta_i^2$$

In addition, it should be noted that the mass concentration of BC obtained from



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Figure 3. Relative deviations of the number fraction of nearly hydrophobic (NH) POA to the In-BC density (a), the hygroscopic parameter of OA to the In-BC density (b), the POA density to the In-BC density (c), the SOA density to the In-BC density (d), the externally-mixed BC density to In-BC density (e) and the combined deviations based on multiple factors mentioned above (f).

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422	AE33 based on aerosol light absorption may lead some uncertainty. However, the
423	comparison of the simultaneously measured data by SP2 with that by AE33 during the
424	campaign shows that the temporal variations of BC mass concentrations measured by
425	the two techniques are well consistent (Fig S5). Note that the BC mass measured by
426	SP2 is occasionally low probably because of the low detection efficiency in small size
427	(McMeeking et al., 2010; Schwarz et al., 2006). In addition, the SP2 is unable to
428	quantify the BC mass beyond a certain limit because of the saturation of electronic
429	devices recording signals (Pileci et al., 2021). We show that, compared the results that
430	retrieved if applying the BC mass measured by SP2, the BC density retrieved based on
431	AE33 can be 18% higher. Given that the measurement bias from SP2, this
432	overestimation indicates an upper limit of the uncertainty.

433 3 Results and Discussion

434 3.1 Retrieved mixing state and density of BC: comparison and validation

435 Figure 4a shows retrieved time series of the mixing state of ambient BC during the 436 campaign. Large temporal variations of the mass fraction of internally and externally mixed BC are presented during the observed period at the sites. The temporal changes 437 should be related to the atmospheric aging process or diurnal variations of emissions 438 (Liu et al., 2019a; Fan et al., 2020). Statistically, the average mass fraction of externally 439 and internally mixed BC is 20±18 % and 80±20 % respectively, showing that most of 440 441 the BC particles were aged and internally mixed with other components. Previous studies at urban sites have shown that the co-existence of the externally mixed BC in 442 18





the ambient atmosphere (Schwarz et al., 2008; Cheng et al., 2012; Chen et al., 2020)
due to continuous combustion processes (e.g., vehicle exhaust and residential sector)

that the mass fraction of internally BC-containing particles would increase with 459 increasing size and reach ~70 % in Beijing. Overall, the mass fraction obtained in our 460 study is comparable with those reported in urban Beijing. Previous studies also 461 displayed that the significant diversity of the BC mixing state among emission 462 463 conditions and coating process (Shiraiwa et al., 2008; Pan et al., 2017; Zhang et al., 2020b). Accordingly, the densities of the bulk and internally mixed BC present apparent 464 fluctuations as shown in Fig. 4b, which is significantly affected by the variations of BC 465 466 emission sources and its rapid aging process. The density of the In-BC during daytime was generally higher than that at night (Fig. 4c). The elevated BC density during 467 daytime is likely due to that the strong photochemical processes promote the aging of 468 BC particles, which resulted in a conversion from uncompacted structure to compact 469 470 and regular spherical shapes of BC (Qiao et al., 2018; Liu et al., 2019b; Zhou et al., 471 2022). The lift in BC density around 20:00 LT might indicate that the BC particles 472 would be rapidly coated with the secondary inorganic aerosol (SIA) particles and 473 continuously aged in the polluted period due to the heterogeneous reactions of SIA in 474 urban regions (Zhang et al., 2016; Peng et al., 2017). Actually, following the haze evolution, the fraction of nearly hydrophobic group reduced rapidly (Fig. <u>58</u>). 475 476 Consequently, the average density of In-BC increased obviously from the clean 477 conditions to the polluted periods (Fig. <u>\$9</u>). A slight decrease was observed in the bulk 478 BC density during traffic hours. This is likely associated with the continues emissions (e.g., vehicle exhaust) that lead to uncoated or uncompacted BC particles in this period. 479 480 The diurnal cycle in In-BC density is consistent with the coating thickness measured

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by a tandem CPMA-SP2-DMA-SP2 (Liu et al., 2020), demonstrating that the new 483 method can derive the density of ambient BC particles reasonably. Averagely, the bulk 484 and internally mixed BC densities are with campaign averaged values of 0.7±0.5 and 485 1.1±0.6 g cm⁻³ respectively, which are much less than 1.8 g cm⁻³, implying that the BC 486 particles is not a void-free spheres in the urban atmosphere. The results of In-BC density 487 are comparable with that observed at the other sites in North China Plain (NCP) as 488 shown in Fig. 4d, illustrating that the BC effective density retrieved by this method is 489 within the range of field measurements. 490



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and In-BC and the measured density distribution spectrum of BC from different sources
 reported in literatures.

Based on both field measurements (e.g. Lide 1992; Zhang et al., 2016; Wu et al., 496 2019; Liu et al., 2019b) and laboratory studies (e.g. McMurry et al., 2002; Park et al., 497 498 2003, 2004; Olfert et al., 2007; Kiselev et al., 2010; Gysel et al., 2011, 2012), the BC density from diverse combustion sources or representing different aging degree has 499 been obtained and ranges widely from 0.14 to 2.1 g cm⁻³, as has been summarized and 500 shown in Fig. 5. Mean probability distribution function (PDF) of the density of bulk 501 and In-BC retrieved by this study is also presented in Fig. 5. It shows that the retrieved 502 density of bulk BC exhibits a dominant mode with a peak value of 0.7 g cm⁻³, which is 503 situated between the typical density range of those externally mixed and internally 504 mixed BC measured previously. For the In-BC, the PDF is with a peak value at 1.1 g 505 cm⁻³, but ranges widely from ~0.5 to 2.5 g cm⁻³, which indicates various morphologies, 506 different aging degree and compositions of ambient BC particles due to the complex 507 508 impact of multiple local sources and aging processes during the observed period in 509 urban Beijing. Overall, the retrieved values for In-BC density fall within the range of typical internal mixed BC reported in the literatures, verifying the reliability of our 510 inversion results. 511

512 **3.2 Sensitivity of predicted** *N*_{CCN} **to changes of BC density**

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513 A previous study showed that the use of an inaccurate density value of BC particles 514 would result in large bias in estimating κ of ambient aerosol particles with the ZSR

515	mixing rule (Fan et al., 2020), as would further lead to uncertainties in prediction of
516	$N_{\rm CCN}$ and relevant climate effects. Considering the large variation range of BC density
517	during the campaign, which is closely associated with its morphology or degree of its
518	aging, we further examine the sensitivity of critical supersaturation (S_c), critical
519	diameter (D_{cut}) and predicted N_{CCN} to variations of BC density (Fig. 6). Here, we use
520	the critical diameter and particle number size distribution to calculate $N_{\rm CCN}$. The method
521	to derive the critical diameter is based on Köhler theory and ZSR rule.



Figure 6. Sensitivity of critical supersaturation (S_c) (a) and diameter (D_{cut}) (b) to the

variations in BC density; Predicted $N_{\rm CCN}$ as a function of measured $N_{\rm CCN}$ by varying the density from 0.14 to 2.1 g cm⁻³ at *S*=0.40 % (c) and *S*=0.23 % (d), the black solid line is the 1:1 line; Diurnal variations in the ratio of predicted-to-measured $N_{\rm CCN}$ at *S*=0.40 % (e) and *S*=0.23 % (f).

528 The results show that, by varying the value of density from 0.14 to 2.1 g cm⁻³ that represents the lower and upper limit of BC density in the atmosphere, the D_{cut} reduces 529 apparently at a given supersaturation (S) (Fig. 6a), or similarly, the S_c decreases rapidly 530 531 for a given particle size (Fig. 6b). The results show that the changes of the D_{cut} and S_c are more sensitive when the BC density is below 1.0 g cm⁻³. And the effects on the D_{cut} 532 and S_c both gradually weakened with the increase of BC density. This shows that it is 533 critical to apply more accurate BC density for the aerosol particles with low aging 534 degree in predicting CCN and its climate effect. Accordingly, the ratios of predicted-535 536 to-measured N_{CCN} ranged from 0.72 to 1.11 by varying the BC density from 0.14 to 2.1 537 g cm⁻³ at the typical S of 0.23 % and 0.40 % (Fig. 6c, 6d), showing an estimation 538 uncertainty of -28 % $_$ 11 % in $N_{\rm CCN}$ prediction.

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The diurnal variations in the ratio of predicted-to-measured $N_{\rm CCN}$ at *S*=0.40 % and 0.23 % are shown to examine the response of the BC density on $N_{\rm CCN}$ prediction at different time periods (Fig. 6e, 6f). By applying the lower limit of density value of 0.14 g cm⁻³, the prediction is much worse compared to the use of the density of 2.1 g cm⁻³ at nighttime (00:00-06:00 LT), when the latter is much closer to the real density of ambient BC (Fig. 4c). The prediction is improved substantially by applying the value of 0.14 g cm⁻³ during evening rush hours (18:00-20:00 LT), during which the ambient BC particles is disturbed by the traffic emissions (Fig. 4c). And now, the prediction becomes worse by applying the value of 2.1 g cm⁻³, and an obvious overestimation by up to ~40 % is shown. The results further illustrate that it is critical to account for the real-time mixing state and density of BC particles in $N_{\rm CCN}$ prediction, particularly in those regions with heavy traffic and residential coal emissions.

It should be noted that the assumption of the surface tension of water would 552 overestimate the critical diameter and underpredict CCN number concentration. While 553 554 the surface tension depression might be more obvious for the small size particles (<60 nm), as the fraction of organics are higher at small particles size (Meng et al., 2014; Cai 555 et al., 2018). Here, in this study, we calculated the critical diameters at supersaturations 556 of 0.40 % and 0.23 %, typical values in cloud, corresponding to larger sizes (> 70 nm 557 and 90 nm) of aerosols. Therefore, the uncertainties from the application of the surface 558 559 tension of pure water should be negligible (< 10 %). Here, three schemes were assumed to evaluate the effect of BC density and mixing state on prediction of CCN number 560 561 concentrations. The detailed calculation methods are presented in the supporting information (SI: Methods) or referenced from Ren et al. (2018). 562

563 **3.3 Using the real-time variations of BC density and mixing state to predict** N_{CCN}

Figure 7 exhibits the comparisons between predicted and measured $N_{\rm CCN}$ at *S* of 0.23 % and 0.40 % by accounting for the retrieved real-time variations of BC density and mixing state. It shows that the $N_{\rm CCN}$ can be well predicted with a slope of 1.01 and 1.02 at *S* of 0.23 % and 0.40 % respectively (Fig. 7a, 7b), only presenting a slight 删除了:., 删除了:. deviation. The slight deviation is primarily due to the fixed value of the density for the
externally mixed BC caused by the retrieved method, especially during noontime and
evening rush periods (Fig. 7c and 7d).



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Figure 7. Prediction CCN number concentration using the mixing state and In-BC density derived from HTDMAs at S=0.40 % (a) and S=0.23 % (b). Diurnal variations in the ratio of predicted-to-measured N_{CCN} at S=0.40 % (c) and S=0.23 % (d).

577 The diurnal variations in the ratio of predicted-to-measured $N_{\rm CCN}$ shows the $N_{\rm CCN}$ 578 can be underestimated by up to 15 % at *S*=0.40 % during those periods. While, a slightly 579 overrated during the evening traffic hours and nighttime may be due to the 580 underestimation of the number fraction of Ex-BC. Overall, the dependence of the CCN 581 prediction on *S* is due to the size dependence of κ and mixing state (Zhang et al., 2017; 26 Liu et al., 2020; Xu et al., 2021). The better closure at S=0.23 % is because that the bulk κ of particles is closer to that the critical diameter corresponding to S=0.23 %, with D_p of 100-150 nm. Similarly, the effect on CCN prediction induced by the bulk mixing state would be more critical for smaller particles, corresponding to the critical diameter at high *S*.

Overall, when considering the effective density of BC relevant to its mixing state, 587 the CCN closure achieves. Previous studies have shown that the fresh emitted BC 588 589 particles may convert from fractal-like aggregates to a compact structure and its density would increase with the aging process (Pagels et al., 2009; Rissler et al., 2014; Peng et 590 al., 2016; Liu et al., 2019b; Zhang et al., 2020a, 2022), but the actual density of In-BC 591 may be lower than 1.8 g cm⁻³ in the ambient atmosphere according to this study. 592 Therefore, the currently applied value represents a density of the void-free structure of 593 594 BC particles may cause an overestimation in CCN prediction.

595 4 Conclusions

The mixing state and effective density of BC changed through heterogenous chemistry process and thus would cause uncertainty in evaluating its CCN activity. In this study, we develop a new method to retrieve the mixing state and effective density of ambient BC using field measurements and the Köhler theory. The uncertainty of the new retrieval method was evaluated within ± 30 %, which is primarily caused by assuming the value of κ_{SOA} and the fraction of primary organic aerosols in nonhygroscopic mode. The retrieved results show that most of the BC particles were aged

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and internally mixed with other components, with mean mass fraction of 80 ± 20 %. Averagely, the retrieved densities of the bulk and <u>internally mixed BC are 0.7\pm0.5 and</u> 1.1±0.6 g cm⁻³ respectively, but ranges widely from ~0.1 to 2.5 g cm⁻³, indicating various morphologies, different aging degree and compositions of ambient BC particles due to the complex impact of multiple local sources and aging processes during the observed period. The retrieved results are basically comparable with the previous observations in North China Plain.

612 Further examination shows the $N_{\rm CCN}$ prediction is with uncertainties of -28 % _11 % at the typical S of 0.23 % and 0.40 % by varying the BC density from 0.14 to 2.1 g cm⁻ 613 ³ that represents the lower and upper limit of ambient BC particles. Moreover, the 614 prediction is found more sensitive to the variability of BC density when it is <1.0 g cm⁻ 615 616 ³, suggesting a great significance to account for the effect of BC density for the aerosol 617 particles with low aging degree when evaluating the climate effect. The CCN closure achieves when introducing the retrieved real-time BC density relevant to its mixing 618 619 state. This work provides a unique way of utilizing field observations to infer ambient BC density and highlights the current assumption of a void-free structure of BC 620 621 particles in models would cause large uncertainties in CCN prediction and in the relevant climate effect evaluation. 622

623 The method used to derive the ambient BC density has limitations. Since the 624 assumptions on the values of κ_{SOA} , ρ_{POA} , ρ_{SOA} and ρ_{Ex-BC} as well as the fraction of 625 primary organic aerosols in non-hygroscopic or hygroscopic mode would add 626 uncertainty in the inferred values of ambient internally mixed BC density. It is thus 删除了: internal-

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629	necessary to examine observational data to verify this methodology in further studies.
630	However, the method and results of this study could provide the way for a more
631	comprehensive understanding of the variability in BC density in Beijing. Additionally,
632	it has the potential to reveal the uncertainties of usage of void-free structure of BC
633	density in accessing the climate effects.

634 Data availability.

All data needed to evaluate the conclusions in the paper are present in the paper and/or
the Supplement. All data used in the study are also available from the corresponding
author upon request (zhangfang2021@hit.edu.cn).

638 Author contributions.

FZ and JR conceived the conceptual development of the manuscript. JR directed and
performed of the experiments with JL, LC, and FZ. JR conducted the data analysis and
wrote the draft of the manuscript. All authors edited and commented on the various
sections of the manuscript.

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648 Competing interests.

- 649 The contact author has declared that neither they nor their co-authors have any
- 650 competing interests.

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