



1   **Variations of the density of ambient black carbon retrieved by a new**  
2   **method: importance to CCN prediction**

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24 **Abstract.**

25       The effective density of black carbon (BC) is a crucial factor relevant to its  
26 morphology and mixing state that would add uncertainty in evaluating its climate effect.  
27 Here, we develop a new method to retrieve the effective density of ambient BC  
28 combining field observations with the Köhler theory. The uncertainty of the new  
29 retrieval method was evaluated within  $\pm 30\%$ , which is primarily caused by  
30 assumptions of the hygroscopic parameter of organics and the fraction of primary  
31 organic aerosols in non-hygroscopic or hygroscopic mode. Using the new method, we  
32 obtain that the ambient BC density during the campaign varies widely from 0.14 to 2.1  
33  $\text{g cm}^{-3}$ , with a campaign mean density of  $1.11 \pm 0.54 \text{ g cm}^{-3}$  for internally-mixed BC that  
34 accounts for  $79 \pm 18\%$  of total BC particles. The retrieved values fall within the range  
35 of typical density of internally-mixed BC reported in the literatures. We further  
36 examined the sensitivity of cloud condensation nuclei (CCN) number concentrations  
37 ( $N_{\text{CCN}}$ ) prediction to variations of BC density, showing an uncertainty of  $-28\% \sim 11\%$   
38 in calculating  $N_{\text{CCN}}$  at supersaturations of 0.2 % and 0.4 % by varying the BC density  
39 within the retrieved ranges. We also find that the  $N_{\text{CCN}}$  is more sensitive to the variations  
40 of BC density when it is  $< 1.0 \text{ g cm}^{-3}$ , illustrating a necessity of accounting for such  
41 effect closer to source regions where the BC particles are mostly freshly emitted. The  
42 CCN closure achieves when introducing the retrieved real-time BC density and mixing  
43 state. This study provides a unique way of utilizing field measurements to infer ambient  
44 BC density and highlights the importance of applying varying BC density values in



45 models when predicting CCN and assessing its relevant climate effect.

## 46 **1 Introduction**

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

58 BC particles, with diesel vehicles, industrial and residential coal combustion as  
59 major sources, are ubiquitous in urban environments (Bond et al., 2013; Dameto et al.,  
60 2017; Li et al., 2017; D. Liu et al., 2019). Typically, freshly generated BC exists in the  
61 form of chain aggregates. While after aging, the BC particles generally mix with other  
62 materials by condensation, coagulation, and other processes (Riemer et al., 2004; Zhang  
63 et al., 2008; Liu et al., 2013; Zhang et al., 2020). The structure of BC would be more  
64 compact with regular shapes (Pagels et al., 2009; Zhang et al., 2008; Wang et al., 2017),  
65 and the effective density of BC are changed accordingly with the reconstruction (H. Liu



66 et al., 2019). The density and morphology of BC particles are closely related to its  
67 sources, mobility size after aging, coating thickness, coating material and its chemical  
68 composition (Zhang et al., 2008; Pagels et al., 2009; Peng et al., 2016; Zhang et al.,  
69 2022). A wide range of BC density has been reported in previous studies (Lide 1992;  
70 McMurry et al., 2002; Park et al., 2004; Kiselev et al., 2010). Recent field measurements  
71 have indicated that the average density of BC is  $\sim 1.2 \text{ g cm}^{-3}$  in the ambient atmosphere  
72 (Zhang et al., 2016). Field measurements have also indicated that a fraction of  
73 externally mixed/uncoated BC exists (Clarke et al., 2004; Cheng et al., 2012; Chen et  
74 al., 2020) and even higher than that internal/ aged BC particles in the ambient  
75 atmosphere (Schwarz et al., 2008; Massoli et al., 2015). While, in climate models, the  
76 BC was generally assumed internally-mixed and treated to have a void-free spherical  
77 structure and a density value of  $1.8 \text{ g cm}^{-3}$  (Bond et al., 2013). This may lead to bias in  
78 estimating the climate effects driven by BC.

79 Previous study based on a case study show that when the aging degree of ambient  
80 particles is low, the BC density ( $\sim 1.8 \text{ g cm}^{-3}$ ) under the spherical assumption will lead  
81 to the overestimation of particles hygroscopicity by 40-50 % and the overestimation  
82 can be explained almost 100 % by using the effective density of fresh BC ( $\sim 0.45 \text{ g cm}^{-3}$ )  
83 (Fan et al. 2020). This indicates the importance of using reasonable BC density values  
84 in the calculation of particle hygroscopicity. In addition, when estimating the CCN  
85 number concentration, a significant bias of  $-35 \% \sim +20 \%$  was found caused by the  
86 assumption of particle mixing state (Ren et al., 2018). However, these studies have not  
87 yet account for such impact of BC density and mixing state on CCN prediction due to



88 lack of real time measurement data.

89 The mixing state and the density of BC particles are usually directly measured by  
90 several techniques, such as an integrated system of a volatility tandem differential  
91 mobility analyzer and a single particle soot photometer (VTDMA-SP2) (Zhang et al.,  
92 2016), or a differential mobility analyzer with a SP2 (DMA-SP2) (Olfert et al., 2007;  
93 Rissler et al., 2014; Wu et al., 2019), and a differential mobility analyzer–centrifugal  
94 particle analyzer–single-particle soot photometer (DMA–CPMA–SP2) system (H. Liu  
95 et al., 2019; Yu et al., 2020), etc. However, such techniques or measurements are not  
96 available in many previously conducted field campaigns. In this study, we develop a  
97 novel method for retrieving the mixing state and effective density of ambient BC  
98 particles by combining field measured hygroscopic growth factor and aerosol chemical  
99 composition and Köhler theory (Petters and Kreidenweis, 2007). The uncertainty of the  
100 new retrieval method was evaluated. The retrieved results were also compared and  
101 validated with existing observations. In addition, the effect of BC density and mixing  
102 state on prediction of CCN number concentrations is further evaluated through a  
103 sensitivity and closure test by accounting for the retrieved real-time variations of BC  
104 density and mixing state.

## 105 **2 Field measurements and methodology**

### 106 **2.1 Field measurements**

107 Measurements in this study were conducted from 15 November to 14 December  
108 2016 at a typical urban site of Beijing (39.97°N, 116.37°E, 49 m above sea level). The



109 site locates at the Institute of Atmospheric Physics, Chinese Academy of Sciences,  
110 which is mainly influenced by the surrounding cooking, road traffic and residential coal  
111 burning emissions during the home heating periods (Sun et al., 2016). The detailed  
112 information about the sampling site was presented in previous studies (Sun et al., 2015;  
113 Zhang et al., 2019). The number concentration of condensation nuclei (CN) at each size  
114 was measured by a scanning mobility particle sizer, which is equipped with a  
115 differential mobility analyzer (DMA; model 3081, TSI) and a condensation particle  
116 counter (CPC; model 3772, TSI). Then the mono-dispersed particles were introduced  
117 into a Droplet Measurement Technologies CCN counter (CCNc, DMT; Lance et al.,  
118 2006) to measure CCN number concentration. A hygroscopic tandem differential  
119 mobility analyzer (HTDMA) system was used to measure the hygroscopic growth  
120 factor (Gf) (Tan et al., 2013). Here four diameters of 40, 80, 110, 150, and 200 nm are  
121 selected in the campaign. Gf is defined as the ratio of the mobility diameter at the given  
122 RH to the dry diameter (Petters and Kreidenweis, 2007). The nonrefractory submicron  
123 aerosol chemical composition was measured by an Aerosol Chemical Speciation  
124 Monitor (ACSM; Sun et al., 2015), including sulfate, nitrate, ammonium, chloride, and  
125 organics. Two factors, including a non-hygroscopic primary organic aerosol (POA) and  
126 hygroscopic secondary organic aerosol (SOA) were classified by using positive matrix  
127 factorization (PMF) with PMF algorithm (v4.2) method (Paatero and Tapper, 1994) and  
128 followed the procedures reported in Ulbrich et al., 2009. The refractory black carbon  
129 mass loading was measured by an aethalometer (model AE33, Magee Scientific  
130 Corporation). Both the nonrefractory materials and BC mass concentration were



131 measured with diameters  $< 1.0 \mu\text{m}$ . The detailed description of the instrument operation  
132 and data process have been described in details elsewhere (Ren et al., 2018; Xu et al.,  
133 2019; Zhang et al., 2019; Fan et al., 2020).

## 134 2.2 Retrieving the mixing state and density of BC

135 The Gf probability distribution function (Gf-PDF) for a specified diameter can be  
136 retrieved firstly based on the TDMAinv algorithm (Gysel et al., 2009). The  $\kappa$ -PDF can  
137 be further calculated based on the Gf-PDF (Fan et al., 2020). Size-resolved  $\kappa$  is derived  
138 using  $\kappa$ -Köhler theory based on hygroscopic growth factor (Gf) (Petters and  
139 Kreidenweis, 2007),

$$140 \quad \kappa_{gf} = (Gf^3 - 1) \cdot \left[ \frac{1}{RH} \exp\left(\frac{4\sigma_{s/a}M_w}{RT\rho_w D_d Gf}\right) - 1 \right] \quad (1)$$

141 where Gf is hygroscopic growth factor, RH is the relative humidity in the HTDMA  
142 (90 %),  $D_d$  is the dry diameter,  $\sigma_{s/a}$  is assumed to be the surface tension of pure water,  
143  $R$  is the universal gas constant,  $T$  is the temperature,  $M_w$  and  $\rho_w$  is the molecular mass,  
144 and the density of water, respectively.

145 The  $\kappa$ -PDF patterns of particles in different sizes always present two modes: nearly  
146 hydrophobic (NH) mode with  $\kappa_{gf} \leq 0.1$  and more hygroscopic (MH) mode with  $\kappa_{gf} > 0.1$   
147 (Fig. S1). Firstly, based on the  $\kappa$ -PDF patterns, the number fraction (NF) of the total  
148 nearly hydrophobic group with the boundary of  $[0, 0.1]$  was calculated by using the  
149 following equation:

$$150 \quad NF = \int_0^{0.1} c(\kappa, D_p) d\kappa \quad (2)$$



151 Here, the  $\kappa$ -PDF, represented by  $c(\kappa, D_p)$ , was normalized as  $\int c(\kappa, D_p) d\kappa = 1$ ,  
152 where  $\kappa$  can be replaced by  $\kappa_{gf}$ ,  $D_p$  is the selected electrical mobility diameter in the  
153 campaign.

154 Considering the nearly hydrophobic mode consists of both external mixed POA  
155 (Ex-POA or bare POA) and externally mixed BC (Ex-BC). Here, the number fraction  
156 of POA in nearly hydrophobic mode ( $NF_{NH-POA}$ ) was assumed to be 70 % according to  
157 the simultaneous measurements (Liu et al., 2021), and the rest 30 % of POA is assumed  
158 mixed with the other hygroscopic aerosols. And thus, the number fraction of Ex-BC  
159 was calculated using the total number fraction of NH mode minus the number fraction  
160 of NH-POA. Then, the number size distribution of the external mixed BC ( $n_{Ex-BC}(\log$   
161  $D_p)$ ) can be calculated based on the particle number size distribution (PNSD) and the  
162 number fraction of the hydrophobic mode of BC ( $NF_{Ex-BC}$ ) as follows:

$$163 \quad n_{Ex-BC}(\log D_p) = NF_{Ex-BC} \times n(\log D_p) \quad (3)$$

164 where  $n(\log D_p)$  is the function of the aerosol number size distribution,  $D_p$  is the  
165 mobility diameter.

166 By assuming that the particles are spherical (Rader and McMurry, 1986), the mass  
167 size distribution of Ex-BC ( $M_{Ex-BC}$ ) was obtained as follows:

$$168 \quad M_{Ex-BC}(\log D_p) = \frac{\pi}{6} D_p^3 \rho n_{Ex-BC}(\log D_p) \quad (4)$$

169 where  $D_p$  is the mobility diameter,  $\rho$  is the effective density of Ex-BC, and  $n_{Ex-BC}$   
170 ( $\log D_p$ ) is the function of the number size distribution of Ex-BC, respectively. Here we  
171 assume that the Ex-BC effective density is  $0.40 \text{ g cm}^{-3}$  according to previous  
172 measurements reported in North China Plain (Peng et al., 2016, 2017; Wu et al., 2019;





173 Liu et al., 2020).

174 The Ex-BC mass size distributions was modelled as a single log-normal  
175 distribution as shown in Fig. S2 (Wu et al., 2017; D. Liu et al., 2019). Thus, the bulk  
176 mass concentration of Ex-BC can be calculated from the integration of the mass size  
177 distribution:

$$178 \quad m_{\text{Ex-BC}} = \int_{D_{\text{start}}}^{D_{\text{end}}} M_{\text{Ex-BC}}(\log D_p) d \log(D_p) \quad (5)$$

$$179 \quad m_{\text{In-BC}} = m_{\text{BC}} - m_{\text{Ex-BC}} \quad (6)$$

180 where  $D_{\text{start}}$  and  $D_{\text{end}}$  are the lower and upper size limit,  $M_{\text{Ex-BC}}(\log D_p)$  is the  
181 function of the Ex-BC mass size distribution. We then obtained the bulk mass  
182 concentration of internally mixed BC ( $m_{\text{In-BC}}$ ) by minus  $m_{\text{Ex-BC}}$  from the bulk BC mass  
183 concentration measured by AE33 in the equation 6.

184 For retrieval of the density of BC, the principal idea is to use the measured  $\kappa_{\text{gf}}$  to  
185 calculate the density of BC based on the Zdanovskii–Stokes–Robinson (ZSR) mixing  
186 rule (Stokes and Robinson, 1966; Zdanovskii, 1948) with the chemical composition  
187 measured by ACSM (Petters & Kreidenweis, 2007). In the retrieval, several aspects are  
188 concerned. First, since the ZSR rule is with an assumption of the aerosol particles are  
189 internally mixed, the  $\kappa_{\text{gf}}$  value of the more MH mode ( $\kappa_{\text{gf-MH}}$ ) is thus applied for  
190 retrieving the density of internally-mixed BC. Second, since the size distribution of BC  
191 number concentration is usually with peaks between 100 and 200 nm (D. Liu et al.,  
192 2019; Yu et al., 2020; Zhao et al., 2022), the  $\kappa_{\text{gf-MH}}$  value of particles in accumulation  
193 mode, which showed an independence on particle size when the  $D_p > 100$  nm during the  
194 campaign period (Fan et al., 2020), was averaged and applied for the retrieval. In



195 addition, because the inversion including measurements from HTDMA and ACSM, a  
 196 total mass closure of the measured aerosol particles was conducted between the two  
 197 techniques by comparing the mass concentration of  $PM_{10}$  and the results are well  
 198 consistent (Fig. S3). The density of internally-mixed BC (In-BC),  $\rho_{In-BC}$  is then derived  
 199 from the following equations:

$$200 \quad \kappa_{gf-MH} = \kappa_{chem} = \sum_i \varepsilon_i \kappa_i = \frac{v_{inorg}}{v_{total}} \kappa_{inorg} + \frac{v_{SOA}}{v_{total}} \kappa_{SOA} + \frac{v_{In-POA}}{v_{total}} \kappa_{POA} + \frac{v_{In-BC}}{v_{total}} \kappa_{BC} \quad (7)$$

201 where  $\kappa_{gf-MH}$  is the hygroscopic parameter of the more hygroscopic (MH) mode,  
 202  $\kappa_{chem}$  is the hygroscopic parameter of aerosol particles in the mixed composition and  
 203 can be calculated based on chemical volume fractions by using a simple rule (Petters &  
 204 Kreidenweis, 2007),  $\kappa_i$  is the hygroscopic parameter of each pure composition and  $\varepsilon_i$  is  
 205 the volume fraction of the individual components in the internal-mixed particle.  $v_{inorg}$ ,  
 206  $v_{SOA}$  and  $v_{In-POA}$  are the volume of the inorganic, SOA and internally mixed POA species,  
 207 and can be calculated as follows:  $v_{inorg} = \frac{m_{inorg}}{\rho_{inorg}}$ ,  $v_{SOA} = \frac{m_{SOA}}{\rho_{SOA}}$ , and  $v_{In-POA} =$   
 208  $\frac{m_{In-POA}}{\rho_{POA}}$ .  $v_{total}$  is the total volume of all the species and can be written as  $v_{total} = \frac{m_{inorg}}{\rho_{inorg}} +$   
 209  $\frac{m_{SOA}}{\rho_{SOA}} + \frac{m_{In-POA}}{\rho_{POA}} + \frac{m_{In-BC}}{\rho_{In-BC}}$ . In the equation (7),  $\kappa_{BC}$  and  $\kappa_{POA}$  are assumed to be 0. Then, the  
 210  $\rho_{In-BC}$  can be calculated based on its mass concentration and volume as follows:

$$211 \quad \rho_{In-BC} = \frac{m_{In-BC}}{\left( \frac{m_{inorg}}{\rho_{inorg} \kappa_{gf-MH}} + \frac{m_{SOA}}{\rho_{SOA} \kappa_{SOA}} - \frac{m_{inorg}}{\rho_{inorg}} - \frac{m_{SOA}}{\rho_{SOA}} - \frac{m_{In-POA}}{\rho_{POA}} \right)} \quad (8)$$

212 where,  $m_{In-BC}$  is the mass concentration of internal-mixed BC,  $m_{inorg}$  and  $m_{SOA}$  are  
 213 the mass concentrations of the inorganic species and SOA, which are measured by the  
 214 ACSM.  $m_{In-POA}$  is the mass concentrations of internal-mixed POA and can be calculated  
 215 by using the total mass concentrations of POA minus the mass fraction of NH-POA.  
 216  $\rho_{inorg}$ ,  $\rho_{SOA}$  and  $\rho_{POA}$  are the density of the inorganic species, SOA and POA. Here



217 inorganic species were derived by applying a simplified ion pairing scheme (Gysel et  
218 al., 2007). Three inorganic salts including  $\text{NH}_4\text{HSO}_4$ ,  $(\text{NH}_4)_2\text{SO}_4$ , and  $\text{NH}_4\text{NO}_3$  were  
219 applied in our study. The densities for inorganic salts and organics were referred from  
220 the articles published (Gysel et al., 2007; Wu et al., 2016). Here the densities for three  
221 inorganics are 1.78, 1.79 and 1.72  $\text{g cm}^{-3}$ , respectively. And the  $\rho_{\text{SOA}}$  and  $\rho_{\text{POA}}$  are  
222 assumed as 1.4 and 1.0  $\text{g cm}^{-3}$ . The values of  $\kappa$  for each pure component was referred  
223 from Petters & Kreidenweis, 2007 and Gunthe et al., 2009. For the inorganic  
224 compounds, the  $\kappa$  values are 0.56 for  $\text{NH}_4\text{HSO}_4$ , 0.48 for  $(\text{NH}_4)_2\text{SO}_4$  and 0.58 for  
225  $\text{NH}_4\text{NO}_3$ . The  $\kappa_{\text{SOA}}$  is assumed to be 0.15 according to the field studies in urban areas  
226 (Chang et al., 2010; Kawana et al., 2016).

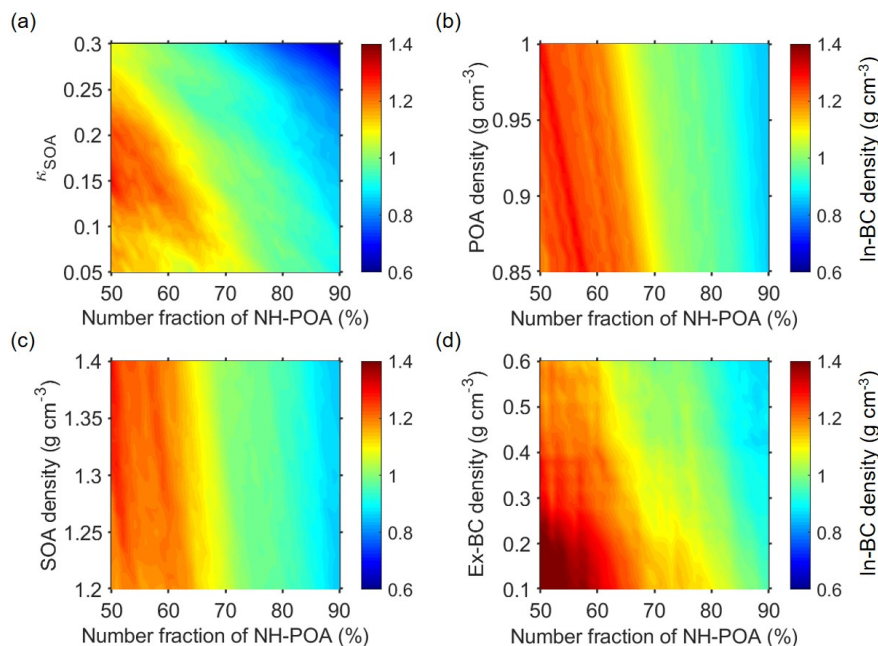
### 227 2.3 Uncertainties and limitations

228 For the retrieval, the assumptions on the values of  $\kappa_{\text{SOA}}$ ,  $\rho_{\text{POA}}$ ,  $\rho_{\text{SOA}}$  and  $\rho_{\text{Ex-BC}}$  as  
229 well as the fraction of primary organic aerosols in non-hygroscopic or hygroscopic  
230 mode would add uncertainty in the inferred values of ambient internally-mixed BC  
231 density. Therefore, we examined the sensitivities of In-BC density to the variations of  
232 these factors, as exhibited in Fig. 1. Within a typical atmospheric observed range of 50-  
233 90 % for the  $NF_{\text{NH-POA}}$  (Liu et al., 2021), the assumption on  $NF_{\text{NH-POA}}$  can lead to  
234 relative deviations (uncertainty) of -20 %-+30 % for the retrieved BC density (Fig. 2a).

235 In addition, unlike inorganics, which the hygroscopicity has been already well-  
236 understood (Petters and Kreidenweis, 2007), the hygroscopicity of organic species  
237 varies largely due to the complexity in organic aerosol constituents. Therefore, the



238 assumption of the values of  $\kappa_{\text{SOA}}$  will add the uncertainty in the calculation of BC



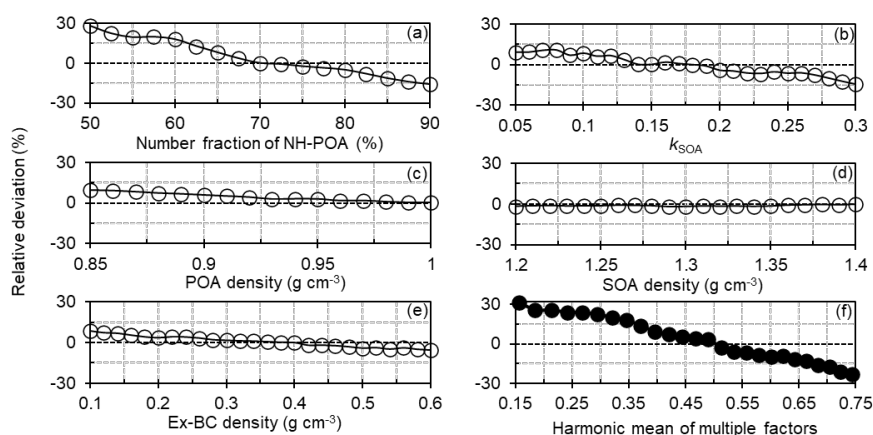
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240 **Figure 1.** Sensitivities of In-BC density to the variations in the number fraction of  
241 nearly hydrophobic (NH) POA and hygroscopic parameter of OA ( $\kappa_{\text{SOA}}$ ) (a), POA  
242 density (b), SOA density (c) and the externally-mixed BC density (d).

243 density. Previous studies have suggested that the organics has a wide range of  $\kappa$  values  
244 ranging from 0.05 to 0.3 (Jimenez et al., 2009; Mei et al., 2013). Thus, the sensitivity  
245 test has also been done to examine the effect due to changes in  $\kappa_{\text{SOA}}$  on calculating the  
246 density of BC (Fig. 1a). The result shows that the assumption of  $\kappa_{\text{SOA}}$  value can cause  
247 an average relative deviation of -20 % - +10 % in calculating the density of In-BC (Fig.  
248 2b). However, the sensitivity test shows that the impact of both the  $\rho_{\text{POA}}$  and  $\rho_{\text{SOA}}$   
249 variations on the BC density estimation is very small or even negligible (Fig. 1b, c). By  
250 varying the  $\rho_{\text{POA}}$  from 0.85 to 1.0  $\text{g cm}^{-3}$  and the  $\rho_{\text{SOA}}$  from 1.2 to 1.4  $\text{g cm}^{-3}$  according



251 to the literatures (Noureddini et al., 1992; Alfarrá et al., 2006; Reyes-Villegas et al.,  
 252 2018), the retrieval uncertainties in the BC density are within  $\pm 10\%$  and  $\pm 5\%$   
 253 respectively (Fig. 2c, d). Similarly, it exhibits that the variations of the  $\rho_{\text{Ex-BC}}$  would



254 **Figure 2.** Relative deviations of the number fraction of nearly hydrophobic (NH) POA  
 255 to the In-BC density (a), the hygroscopic parameter of OA to the In-BC density (b), the  
 256 POA density to the In-BC density (c), the SOA density to the In-BC density (d), the  
 257 externally-mixed BC density to In-BC density (e) and the combined deviations based  
 258 on multiple factors mentioned above (f).

259 cause an average deviation of  $\pm 10\%$  (Fig. 2e) in retrieved In-BC density when  
 260 increasing the values of  $\rho_{\text{Ex-BC}}$  from 0.10 to 0.60  $\text{g cm}^{-3}$ , which represents a typical range  
 261 of  $\rho_{\text{Ex-BC}}$  in ambient atmosphere (Wu et al., 2019; Liu et al., 2020). A combined  
 262 uncertainty ( $\delta$ ) caused by the multiple factors ( $\delta_i$ ), which is calculated by the equation  
 263 9, is  $-23\%$ – $+31\%$  as shown in Fig. 2f.

$$264 \quad \delta = \sqrt{\sum_{i=1}^n \delta_i^2} \quad (9)$$

265 Note that, this method fails to retrieve the BC density when organics account for



266 a large fraction (>60 %). This is because that a higher fraction of OA usually  
267 corresponds to lower total volume of all the species (Fig. S4), yielding negative values  
268 for  $v_{In-BC}$  introduced in equation 10. As a result, 62 % of the data observed during the  
269 campaign were valid for calculating the BC density.

$$270 \quad v_{In-BC} = \frac{v_{inorg} \kappa_{inorg} + v_{SOA} \kappa_{SOA}}{\kappa_{gf-MH}} - v_{inorg} - v_{SOA} - v_{In-POA} \quad (10)$$

271 Similarly, the bulk density of BC ( $\rho_{bulk-BC}$ ) is calculated with the same method as  
272 that for calculation the  $\rho_{In-BC}$ . When calculating the  $\rho_{bulk-BC}$ , the bulk  $\kappa_{gf}$  value measured  
273 by HTDMA is applied with the assumption of all the aerosol particles are internally  
274 mixed.

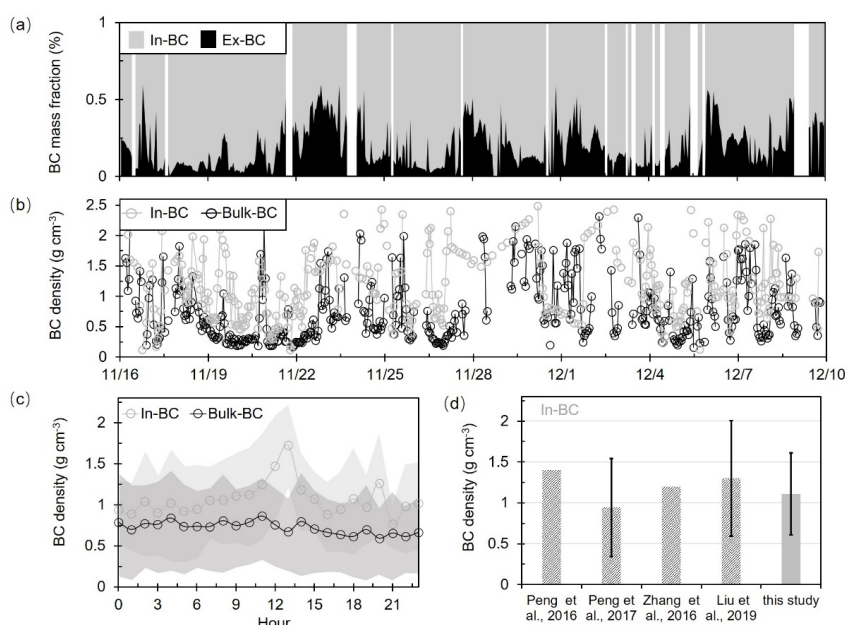
## 275 **3 Results and Discussion**

### 276 **3.1 Retrieved mixing state and density of BC: comparison and validation**

277 Figure 3a shows retrieved time series of the mixing state of ambient BC during the  
278 campaign. Large temporal variations of the mass fraction of internal and external mixed  
279 BC are presented during the observed period at the sites. The temporal changes should  
280 be related to the atmosphere aging process or diurnal variations of emissions (D. Liu et  
281 al., 2019; Fan et al., 2020). Statistically, the average mass fraction of externally and  
282 internally-mixed BC is  $21 \pm 18$  % and  $79 \pm 18$  % respectively, showing that most of the  
283 BC particles were aged and internally-mixed with other components. Previous studies  
284 at urban sites have shown that the co-existence of the externally mixed BC in the  
285 ambient atmosphere (Schwarz et al., 2008; Cheng et al., 2012; Chen et al., 2020) due  
286 to continuous combustion processes (e.g., vehicle exhaust and residential sector) (Wang



287 et al., 2017; D. Liu et al., 2019). Our results are basically comparable with both those  
288 directly measured or indirectly retrieved previously reported results. For example, Chen  
289 et al., 2020 found that the mass fraction of internal mixed BC particles was nearly to be



290  
291 **Figure 3.** (a) Time series of the mass fraction of the retrieved internal- and external-  
292 mixed BC; (b) Time series of the retrieved density of the bulk and internal- mixed BC  
293 (In-BC); (c) Diurnal variation of the retrieved density of bulk and In-BC; (d)  
294 Comparison of the results of the derived In-BC density in this study with that reported  
295 in literatures.

296 ~80–90 % in summer of Beijing based on VTDMA measurements. Liu et al., (2020),  
297 using a tandem system with an aerodynamic aerosol classifier and SP2, reported that  
298 the mass fraction of internal BC-containing particles would increase with increasing  
299 size and reach ~70 % in Beijing. Overall, the mass fraction obtained in our study is

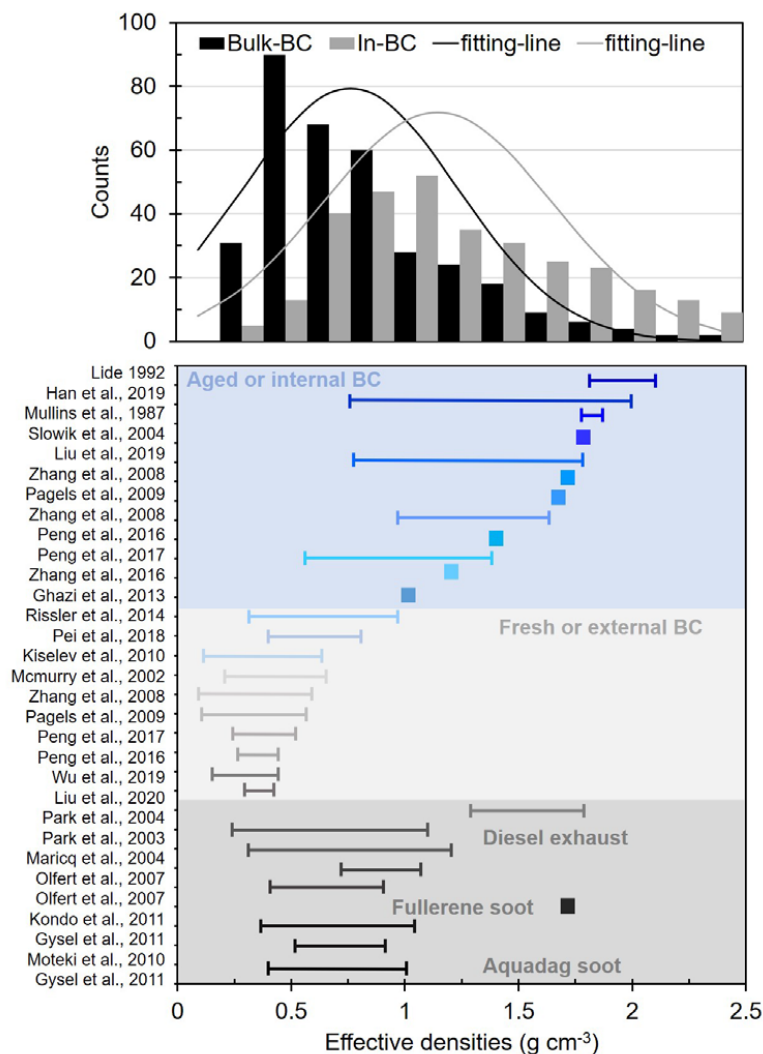


300 comparable with those reported in urban Beijing. Previous studies also displayed that  
301 the significant diversity of the BC mixing state among emission conditions and coating  
302 process (Shiraiwa et al., 2008; Pan et al., 2017; Zhang et al., 2020). Accordingly, the  
303 densities of the bulk and internal mixed BC present apparent fluctuations as shown in  
304 Fig. 3b, which is significantly affected by the variations of BC emission sources and its  
305 rapid aging process. The density of the In-BC during daytime were generally higher  
306 than that at night (Fig. 3c). The elevated BC density during daytime is likely due to that  
307 the strong photochemical processes promote the aging of BC particles, which resulted  
308 in a conversion from uncompact structure to compact and regular spherical shapes of  
309 BC (Qiao et al., 2018; H. Liu et al., 2019; Zhou et al., 2022). The slightly decreases  
310 were observed both in the bulk BC and In-BC density during traffic hours. This is likely  
311 associated with the continues emissions (e.g., vehicle exhaust) that lead to more  
312 uncoated or uncompact BC particles in urban regions. The diurnal cycles in BC  
313 density are consistent with that measured by a tandem CPMA-SP2-DMA-SP2 (Liu et  
314 al., 2020), demonstrating that the new method can derive the density of ambient BC  
315 particles reasonably. Averagely, the bulk and internal-mixed BC densities are with  
316 campaign averaged values of  $0.73 \pm 0.46$  and  $1.11 \pm 0.54$  g cm<sup>-3</sup> respectively, which are  
317 much less than 1.8 g cm<sup>-3</sup>, implying that the BC particles is not a void-free spheres in  
318 the urban atmosphere. The results of In-BC density are comparable with that observed  
319 at the other sites in North China Plain (NCP) as shown in Fig. 3d, illustrating that the  
320 BC effective density retrieved by this method is within the range of field measurements.  
321 Based on both field measurements (e.g. Lide 1992; Zhang et al., 2016; Wu et al., 2019;





322 H. Liu et al., 2019) and laboratory studies (e.g. McMurry et al., 2002; Park et al., 2003,  
 323 2004; Olfert et al., 2007; Kiselev et al., 2010; Gysel et al., 2011, 2012), the BC density  
 324 from diverse combustion sources or representing different aging degree has been



325  
 326 **Figure 4.** The probability distribution function (PDF) of the retrieved density of bulk  
 327 and In-BC and the measured density distribution spectrum of BC from different sources  
 328 reported in literatures.



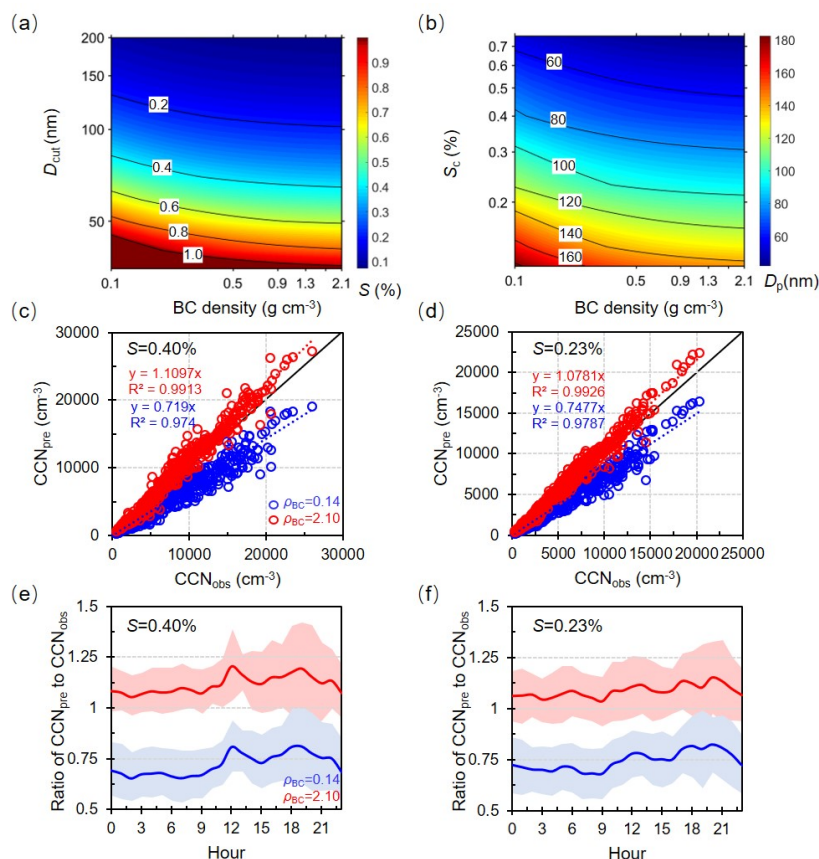
329 obtained and ranges widely from 0.14 to 2.1 g cm<sup>-3</sup>, as has been summarized and shown  
330 in Fig. 4. Mean probability distribution function (PDF) of the density of bulk and In-  
331 BC retrieved by this study are also presented in Fig. 4. It shows that the retrieved density  
332 of bulk BC exhibits a dominant mode with peak value of 0.70 g cm<sup>-3</sup>, which is situated  
333 between the typical density range of those externally-mixed and internally-mixed BC  
334 measured previously. For the In-BC, the PDF is with a peak value at 1.10 g cm<sup>-3</sup>, but  
335 ranges widely from ~0.5 to 2.4 g cm<sup>-3</sup>, which indicates various morphologies, different  
336 aging degree and compositions of ambient BC particles due to the complex impact of  
337 multiple local sources and aging processes during the observed period in urban Beijing.  
338 Overall, the retrieved values for In-BC fall within the range of typical internal mixing  
339 BC reported in the literatures, verifying the reliability of our inversion results.

### 340 **3.2 Sensitivity of predicted $N_{CCN}$ to changes of BC density**

341 It has been studied that an inappropriate use of the density value of BC particles  
342 will result in large bias in estimating  $\kappa$  of ambient aerosol particles with the ZSR mixing  
343 rule (Fan et al., 2020), as would further lead to uncertainties in prediction of  $N_{CCN}$  and  
344 relevant climate effects. Considering the large variation range of BC density during the  
345 campaign, which is closely associated with its morphology or degree of its aging, we  
346 further examine the sensitivity of critical supersaturation ( $S_c$ ), critical diameter ( $D_{crit}$ )  
347 and predicted  $N_{CCN}$  to variations of BC density (Fig. 5). Here, we use the critical  
348 diameter and particle number size distribution to calculate  $N_{CCN}$ . The method to derive  
349 the critical diameter is based on Köhler theory and ZSR rule. Three closure studies were



350 used to evaluate the effect of BC density and mixing state on prediction of CCN number  
 351 concentrations. The detailed calculation methods are presented in the supporting  
 352 information (SI: Methods) or referenced from Ren et al., 2018.



353  
 354 **Figure 5.** Sensitivity of critical supersaturation ( $S_c$ ) (a) and diameter ( $D_{cut}$ ) (b) to the  
 355 variations in BC density; Predicted  $N_{CCN}$  as a function of measured  $N_{CCN}$  by varying the  
 356 density from 0.14 to 2.1  $\text{g cm}^{-3}$  at  $S=0.40\%$  (c) and  $S=0.23\%$  (d), the black solid line  
 357 is the 1:1 line; Diurnal variations in the ratio of predicted-to-measured  $N_{CCN}$  at  $S=0.40\%$   
 358 (e) and  $S=0.23\%$  (f).

359 The results show that, by varying the value of density from 0.14 to 2.1  $\text{g cm}^{-3}$  that



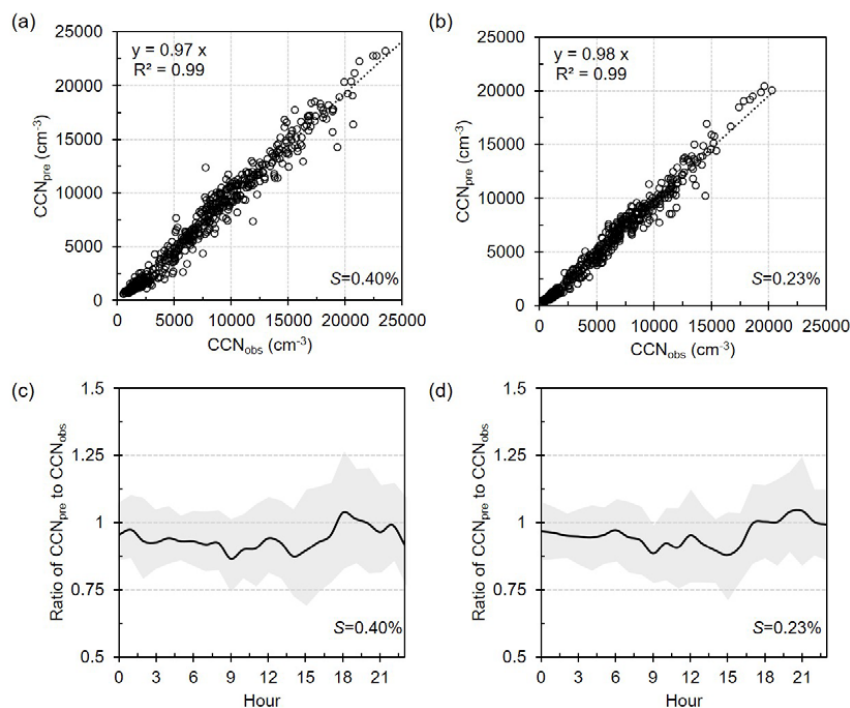
360 represents the lower and upper limit of BC density in the atmosphere), the  $D_{\text{cut}}$  reduces  
361 apparently at a given supersaturation ( $S$ ) (Fig. 5a), or similarly, the  $S_c$  decreases rapidly  
362 for a given particle size (Fig. 5b). The results show that the changes of the  $D_{\text{cut}}$  and  $S_c$   
363 are more sensitive when the BC density is below  $1.0 \text{ g cm}^{-3}$ . And the effects on the  $D_{\text{cut}}$   
364 and  $S_c$  both gradually weakened with the increase of BC density. This shows that it is  
365 critical to apply more accurate BC density for the aerosol particles with low aging  
366 degree in predicting CCN and its climate effect. Accordingly, the ratios of predicted-  
367 to-measured  $N_{\text{CCN}}$  ranged from 0.72 to 1.11 by varying the BC density from 0.14 to 2.1  
368  $\text{g cm}^{-3}$  at the typical  $S$  of 0.2 % and 0.4 % (Fig. 5c, 5d), showing an estimation  
369 uncertainty of -28 %–11 % in  $N_{\text{CCN}}$  prediction.

370 The diurnal variations in the ratio of predicted-to-measured  $N_{\text{CCN}}$  at  $S=0.40$  % and  
371 0.23 % are shown to examine the response of the BC density on  $N_{\text{CCN}}$  prediction at  
372 different time periods (Fig. 5e, 5f). By applying the lower limit of density value of 0.14  
373  $\text{g cm}^{-3}$ , the prediction is much worse compared to the use of the density of  $2.1 \text{ g cm}^{-3}$  at  
374 nighttime (00:00-06:00 LT), when the latter is much closer to the real density of ambient  
375 BC (Fig. 3c). The prediction is improved substantially by applying the value of  $0.14 \text{ g}$   
376  $\text{cm}^{-3}$  during evening rush hours (18:00-20:00 LT), during which the ambient BC  
377 particles is more externally-mixed with smaller densities (Fig. 3c). And now, the  
378 prediction becomes worse by applying the value of  $2.1 \text{ g cm}^{-3}$ , and an obvious  
379 overestimation by up to ~40 % is shown. The results further illustrate that it is critical  
380 to account for the real-time mixing state and density of BC particles in  $N_{\text{CCN}}$  prediction,  
381 in particular in those regions with heavy traffic and residential coal emissions.



382 **3.3 Using the real-time variations of BC density and mixing state to predict  $N_{CCN}$**

383 Figure 6 exhibits the comparisons between predicted and measured  $N_{CCN}$  at  $S$  of  
384 0.2 % and 0.4 % by accounting for the retrieved real-time variations of BC density and  
385 mixing state. It shows that the  $N_{CCN}$  can be well predicted with a slope of 0.98 and 0.97  
386 at  $S$  of 0.2 % and 0.4 % respectively (Fig. 6a, 6b), only presenting a slight deviation.  
387 The slight underestimation is primarily due to an underestimation of the density for the  
388 externally mixed BC caused by the retrieved method, especially during noontime (Fig.  
389 6c and 6d).



390

391 **Figure 6.** Prediction CCN number concentration by using the mixing state and In-BC  
392 density derived from HTDMAs at  $S=0.40\%$  (a) and  $S=0.23\%$  (b). Diurnal variations  
393 in the ratio of predicted-to-measured  $N_{CCN}$  at  $S=0.40\%$  (c) and  $S=0.23\%$  (d).



394           The diurnal variations in the ratio of predicted-to-measured  $N_{CCN}$  shows the  $N_{CCN}$   
395 can be underestimated by up to 20 % at  $S=0.4$  % during those periods. While, a slightly  
396 overrated during the evening traffic hours and nighttime may be due to the  
397 underestimation of the number fraction of Ex-BC. Overall, the dependence of the CCN  
398 prediction on  $S$  is due to the size dependence of  $\kappa$  and mixing state (Zhang et al., 2017;  
399 Liu et al., 2020; Xu et al., 2021). The better closure at  $S=0.2$  % is because that the bulk  
400  $\kappa$  of particles is closer to that the critical diameter corresponding to  $S=0.2$  %, with  $D_p$   
401 of 100-150 nm. Similarly, the effect on CCN prediction induced by the bulk mixing  
402 state would be more critical for smaller particles, corresponding to the critical diameter  
403 at high  $S$ .

404           Overall, when considering the effective density of BC relevant to its mixing state,  
405 the CCN closure achieves. Previous studies have shown that the fresh emitted BC  
406 particles may convert from fractal-like aggregates to a compact structure and its density  
407 would increase with the aging process (Pagels et al., 2009; Rissler et al., 2014; Peng et  
408 al., 2016; H. Liu et al., 2019; Zhang et al., 2020, 2022), but the actual density of In-BC  
409 may be lower than  $1.8 \text{ g cm}^{-3}$  in the ambient atmosphere according to this study.  
410 Therefore, the currently applied value represents a density of the void-free structure of  
411 BC particles may cause an overestimation in CCN prediction.

## 412 **4 Conclusions**

413           The mixing state and effective density of BC changed through heterogenous  
414 chemistry process and thus would causes uncertainty in evaluating its CCN activity. In



415 this study, we develop a new method to retrieve the mixing state and effective density  
416 of ambient BC using field measurements and the Köhler theory. The uncertainty of the  
417 new retrieval method was evaluated within  $\pm 30\%$ , which is primarily caused by  
418 assumptions of the  $\kappa_{\text{SOA}}$  and the fraction of primary organic aerosols in non-hygroscopic  
419 or hygroscopic mode. The retrieved results show that most of the BC particles were  
420 aged and internally-mixed with other components, with mean mass fraction of  $79 \pm 18\%$ .  
421 Averagely, the retrieved densities of the bulk and internal mixed BC are  $0.73 \pm 0.46$  and  
422  $1.11 \pm 0.54 \text{ g cm}^{-3}$  respectively, but ranges widely from  $\sim 0.1$  to  $2.4 \text{ g cm}^{-3}$ , indicating  
423 various morphologies, different aging degree and compositions of ambient BC particles  
424 due to the complex impact of multiple local sources and aging processes during the  
425 observed period. The retrieved results are basically comparable with the previous  
426 observations in North China Plain.

427 Further examination shows the  $N_{\text{CCN}}$  prediction is with uncertainties of  $-28\% - 11\%$   
428 at the typical  $S$  of  $0.2\%$  and  $0.4\%$  by varying the BC density from  $0.14$  to  $2.1 \text{ g cm}^{-3}$   
429 that represents the lower and upper limit of ambient BC particles. Moreover, the  
430 prediction is found more sensitive to the variability of BC density when it is  $< 1.0 \text{ g cm}^{-3}$ ,  
431 suggesting a great significance to account for the effect of BC density for the aerosol  
432 particles with low aging degree when evaluating the climate effect. The CCN closure  
433 achieves when introducing the retrieved real-time BC density relevant to its mixing  
434 state. This work provides a unique way of utilizing field observations to infer ambient  
435 BC density and highlights the currently assumption of a void-free structure of BC  
436 particles in models would cause large uncertainties in CCN prediction and in the



437 relevant climate effect evaluation.

438 **Data availability.**

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

442 **Author contributions.**

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

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452 **Competing interests.**

453 The contact author has declared that neither they nor their co-authors have any





454 competing interests.

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