

1 **Mixing state of black carbon at different atmospheres in north and southwest**
2 **China**

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12 **Abstract**

13 Large uncertainties remain when estimating the radiative forcing by black carbon
14 (BC) because the corresponding microphysical properties have not been well addressed.
15 In this study, the BC size distributions were studied based on three different field
16 campaigns at an urban site, a suburban site, and a background site in China using a
17 single particle soot photometer (SP2) in tandem with a differential mobility diameter.
18 Measurement results indicate that the BC particles were composed of either thinly or
19 thickly coated aerosols. The mean number fractions of the thinly coated BC aerosols
20 were 51%, 67%, and 21% for the urban, suburban, and background sites, respectively.
21 The corresponding thickly coated (thinly coated) core mass median diameters were 187
22 nm (154 nm), 182 nm (146 nm), and 238 nm (163 nm), respectively. The mean diameter
23 of the thickly coated BC-containing aerosols was larger than that of the thinly coated

24 BC-containing aerosols, while the mean BC core diameter of the thickly coated BC-
25 containing aerosols was smaller than that of the thinly coated BC-containing aerosols.
26 About 10% of the BC-containing aerosols with the BC core are attached to the other
27 non-BC components, which were mainly generated by coagulation between the BC and
28 non-BC components. The measurement results in our study can be further used in
29 modeling studies to help with constraining the uncertainties of the BC radiative effects.

30 **Introduction**

31 Black carbon (BC) plays an important role in the climate system by absorbing solar
32 radiation (Ramanathan and Carmichael, 2008), interacting with the cloud (Roberts et al.,
33 2008), and changing the albedo of the snow (Menon et al., 2002). It is the second most
34 important aerosol component after carbon dioxide, contributing to global warming
35 (Bond et al., 2013). The solar absorption of BC has a significant influence on the
36 development of the boundary layer and then aggravates the air pollution (Ding et al.,
37 2016). The turbulence in the atmospheric boundary layer can be suppressed due to the
38 existence of BC (Wilcox et al., 2016). The BC also plays a remarkable role in driving
39 the formation and trend of regional haze (Zhang et al., 2020).

40 BC is mainly generated by the incomplete combustion of biofuels and fossil fuels
41 (Bond and Bergstrom, 2006). After emission, the morphology of BC transforms from
42 fractal to spherical and subsequently grows to a fully compact particle with other
43 chemical components coating it (Peng et al., 2016). During the aging process, the BC
44 optical properties change significantly up to a factor of 3 and then the corresponding
45 magnitude of climate forcing contributed by BC is increased by up to a factor of 2
46 (Zhang et al., 2008). Large uncertainties remain in estimating the BC radiative effects
47 due to the large variation in BC microphysical properties, such as size distributions and
48 mixing states during the aging process (Zhao et al., 2019; Moffet et al., 2016; Matsui et

49 al., 2018). Therefore, characterizing the differences in size distributions and mixing
50 states between the thinly and thickly coated BC particles can help better constrain the
51 uncertainties of BC aerosol radiative effects. To our best understanding, few studies
52 have specified the mixing states and size distributions of both the thinly and thickly
53 coated BC aerosols.

54 The thickly coated BC particles can also be classified into two morphological types:
55 bare BC on the surface of non-BC particles or partially coated by non-BC particles
56 (attached type) and BC embedded within or coated by non-BC components (coated
57 type). With the same amount of non-BC components, the mass absorption cross-sections
58 of BC by the attached type are much smaller than those by the coated type (Moteki and
59 Kondo, 2008; Moteki and Kondo, 2010; Moteki et al., 2014). Therefore, the impact of
60 BC on climate can be better estimated when accurately identifying the two types of
61 ambient BC-containing particles. Observations are required to constrain the spatial and
62 temporal microphysical properties of the atmospheric BC.

63 The single-particle soot photometer (SP2) is always used to measure the mixing
64 states and size distributions of ambient BC particles. In the previous study, advanced
65 technology was used to study the coating over different BC core size diameters on the
66 ground (Liu et al., 2019a) and for vertical profiles (Ding et al., 2019). The measured
67 signals from SP2 can be used to distinguish the BC-containing aerosols as thinly and
68 thickly coated ones. The measured results can also be employed to distinguish the BC-
69 containing particles between attached and coated types, which were described in detail
70 in the methodology part.

71 In this study, the tandem SP2 and differential mobility analyzer (DMA) was
72 employed at an urban site, a suburban site, and a background site in China to investigate
73 the microphysical properties of the BC particles. The size distributions and mixing states

of both the thinly coated and thickly coated BC aerosols at different atmospheres were characterized. We also investigated the corresponding morphology properties of the BC-containing aerosols. The measured microphysical properties provide the basis for future modeling studies of the BC radiative effects in different environments in China.

2 Methodology

2.1 Measurement sites

The measurements were conducted at three different atmospheric sites in China, namely the urban site of Peking University Urban Atmosphere Environment Monitoring Station (PKU, 39.9°N, 116.1°E, 58m a.s.l) in Beijing between 20 January and 4 February 2016, the suburban site of Changping (CP, 40.3°N, 116.2°E, 70m a.s.l) in Beijing between 15 May and 5 June 2016, and the background site of Lijiang (LJ, 27.2°N, 100.2°E, 3410 m a.s.l) in Yunnan Province between 22 March and 4 April 2015. The PKU site is located northwest of Beijing. This site could characterize the air pollution of urban Beijing (Hu et al., 2017; Hu et al., 2021b). The CP site locates in the northwest of the Beijing urban area, representing a regional atmosphere (Zhao et al., 2021; Wang et al., 2019b). The LJ site represents the background areas, located in the Mountain Yulong, in the Yunnan Province of China (Shang et al., 2018; Zheng et al., 2017; Wang et al., 2019a). The aerosol optical depth at the wavelength of 550 nm during the year 2020 indicated that the LJ site was very clean and the PKU and CP sites were more polluted as shown in Fig. S1 in the supplement.

2.2 Instruments

2.2.1 DMA-SP2 system

As for the SP2, the continuous Nd: YAG laser beam with the wavelength of 1064 nm is generated intensively in the instrument chamber. When the BC-containing particles pass through the laser beam, they absorb the radiation and then are heated to around 3500-5000 K. The intensity of the emitted incandescent light from the heated BC particle is then transformed to the BC mass concentration. The scattering signals of the BC particle are recorded to estimate the BC particle mixing state.

In this study, the SP2 (Droplet Measurement Technology, Inc., USA) was placed after the DMA (Model 3081, TSI, USA) to measure the size-resolved BC mixing states, and the instrument setup is schematically shown in Fig. S2. The DMA was set to scan the aerosol over the size range between 12.3 and 697 nm every five minutes. The flow rate leading to the SP2 and the condensation particle counter (CPC, Model 3776, TSI, USA) were 0.12 and 0.28 L/min, respectively. The sheath flow of the DMA was 4 L/min.

The Aquadag was used to calibrate the measured incandescence signal of the SP2 using the DMA-SP2 system. The formula from Gysel et al. (2011) was used to convert the mobility diameter into the mass of Aquadag. A correction factor of 0.75 was applied to account for the different response sensitivity of SP2 to Aquadag and ambient BC (Moteki et al., 2010).

In this study, the coating thickness of the BC-containing aerosols was calculated by the difference between the total mobility diameter measured by the DMA and the optical equivalent diameters of the BC core. Details of calculating the optical equivalent coating thickness can refer to Zhang et al. (2018b) and can be found in section 3 in the supplementary material.

2.2.2 Other instruments

119 The submicron particles (PM_{10}) chemical compositions were measured using a high-
120 resolution time-of-flight aerosol mass spectrometer (AMS; Aerodyne Research Inc.,
121 Billerica, MA, USA). The data processing software PIKA (version 1.16) was used for
122 data analysis. The positive matrix factorization (PMF) analysis was conducted for the
123 source appointment of the organic aerosols (Ulbrich et al., 2009). More details on the
124 measurement of the aerosol chemical compositions and data processing can be found in
125 Zheng et al. (2017).

126 The mass concentrations of O_3 were measured using UV absorption (model 49i,
127 Thermo Fischer Inc. USA) with a time resolution of 1 minute. The mass concentrations
128 of NO and NO_2 were measured using the chemiluminescence technique (NO- NO_2 - NO_x
129 Analyzer, Model 42i, Thermo Scientific, USA). The mass concentrations of SO_2 were
130 measured using the ultraviolet fluorescence method (SO_2 analyzer, model 43i-TLE,
131 Thermo Scientific, USA). The temperature (T), relative humidity (RH), wind speed
132 (WS), and wind direction (WD) were monitored continuously during these campaigns.

133 2.3 Methodology

134 For the BC-containing aerosol, there is a lag between the peak time of the scattering
135 and the incandescence signal (Metcalf et al., 2012). The lag time between the peak
136 scattering signal and the peak incandescence signal can be employed to describe the
137 coating thickness (Moteki and Kondo, 2007; Schwarz et al., 2006) and further used to
138 distinguish the BC-containing aerosols as thinly and thickly coated ones. ~~Despite that,~~
139 ~~the time lag method may not effectively distinguish the BC particles between fresh or~~
140 ~~aged ones because some BC particles are sourced from biomass burning (Schwarz et al.,~~
141 ~~2008b) and solid fuel burning (Liu et al., 2014; Liu et al., 2019b) initially have a higher~~
142 ~~coating and were not aged ones. However, the~~ The lag-time probability distribution at
143 our measurement sites also shows two modes which will be shown in section 3.2, and

thus the lag-time can be used to efficiently distinguish the BC-containing aerosols as thinly and thickly coated ones here. It should be noted ~~Despite that, the time-lag method may not effectively distinguish the BC particles between fresh or aged ones because some BC particles are sourced from biomass burning (Schwarz et al., 2008b) and solid fuel burning (Liu et al., 2014; Liu et al., 2019b) initially have a higher~~ thick coating and but cannot be grouped into aged BC particles. ~~were not aged ones. However,~~

For the thickly coated BC particles, the measured scattering and incandescence signal can also be employed to distinguish the BC-containing particles as attached and coated types (Moteki et al., 2014) by calculating the time-dependent scattering cross-sections of BC-containing particles (Moteki and Kondo, 2007). For the coated type, all of the coating material will evaporate and the scattering cross-sections will decrease to zero after passing through the laser beam, while the scattering cross-section of the attached BC-containing aerosol will not decrease to zero (Moteki and Kondo, 2008). The method adopted by Dahlkötter et al. (2014) was employed here to characterize the morphology of the BC-containing aerosols. Details of distinguishing the BC-containing particles as attached and coated types can also refer to section 4 in the supplementary materials.

3 Results and discussions

3.1 Overview of the measurement results in different atmospheres

The time series of the measurement results are shown in Fig. S6, Fig. S7, and Fig. S8 for the PKU, CP, and LJ sites, respectively. For the PKU site, the wind was mainly from the north and the wind speed was low with a mean value of 2.2 m/s. The ambient atmosphere was very dry with a mean RH of 27.6%, with minimum and maximum values of 5.8% and 72.6%, respectively. The temperature in the winter in Beijing had a mean value of 0.8 °C between -5.9 °C and 9.2 °C. The mean mass concentration of PM_{2.5}

169 was $49.3 \pm 55.4 \mu\text{g}/\text{m}^3$. The concentration of SO_2 and NO_x ($\text{NO}_x = \text{NO} + \text{NO}_2$) had the
170 same trends as $\text{PM}_{2.5}$, with mean values of 16.3 ± 11.9 ppb and 68.2 ± 63.4 ppb,
171 respectively. The O_3 concentration is anti-correlated with $\text{PM}_{2.5}$. For the suburban site
172 CP, the wind showed obvious diurnal cycles with high-speed west wind during the day
173 and low-speed east wind during the night. The mean wind speed was 2.4 ± 1.6 m/s. The
174 RH during the campaign was $38.8 \pm 16.0\%$, with a maximum value of 80.5%. The
175 temperature during the campaign was 21.8 ± 5.2 °C with a maximum value of 33.2 °C.
176 As for the NO_x , the mean concentration was 21.4 ± 17.7 ppb. The mean concentration of
177 SO_2 was 2.89 ± 1.10 ppb. The measured mean O_3 concentration was 54.5 ± 38.8 ppb. The
178 mean $\text{PM}_{2.5}$ concentration was $22.6 \pm 16.8 \mu\text{g}/\text{m}^3$, with a maximum value of $71.8 \mu\text{g}/\text{m}^3$.
179 As for the background LJ site, The mean value of the wind speed, RH, and T were 3.13
180 m/s, 50.23%, and 6.5 °C, respectively. The mean $\text{PM}_{2.5}$ mass concentration was $6.2 \pm$
181 $5.7 \mu\text{g}/\text{m}^3$. The mean NO_x and SO_2 concentrations were 0.05 ppb and 0.97 ppb
182 respectively.

183 The characteristics of the measurement sites are summarized and shown in Fig. 1.
184 The differences in the temperature and RH among these sites mainly resulted from the
185 that the measurements were conducted in different seasons. The concentrations of SO_2 ,
186 NO_x , and $\text{PM}_{2.5}$ indicated that the urban site PKU was the most polluted. The suburban
187 site CP was slightly polluted and the background LJ was the cleanest.

188 The air mass back trajectories as shown in Fig. S9 during the measurement at PKU
189 show that the measurement site was mainly influenced by the polluted air from the south
190 and southeast, and the relatively clean air from the northwest. The CP site was mainly

191 influenced by the clean air from the northwest and the polluted air from the southeast.
192 The air mass of the LJ site was mainly from the southwest and west.

193 **3.2 Mixing states of the thinly coated and thickly coated BC-containing aerosols**

194 The measured lag time probability distributions for the PKU, CP, and LJ sites are
195 shown in Fig. 2 (a), (b), and (c), respectively. The lag time had two modes for each
196 measurement site. The BC particles are sorted as thinly or thickly coated BC. A two
197 log-normal distribution was used for the probability distribution of the lag time for BC-
198 containing particles as:

$$199 \quad \text{PDF}(\Delta t) = \sum_{i=1,2} \frac{A_i}{\sqrt{2\pi} \log(\sigma_{g,i})} \exp \left[-\frac{\log(\Delta t) - \log(\Delta t_i)}{2 \log^2(\sigma_{g,i})} \right],$$

200 Where Δt is the lag time, A_i , $\sigma_{g,i}$, Δt_i are the scale factor, geometric standard
201 deviation, and geometric mean lag time of mode i respectively. The critical lag time that
202 distinguishes the thinly and thickly BC particles was determined by calculating the value
203 when the probability distribution values of mode 1 and mode 2 are equal. In this study,
204 the BC-containing aerosols with a lag time larger than $1.4 \mu\text{s}$ were classified as thickly
205 coated particles for the LJ site. The other BC-containing aerosols were classified as
206 thinly coated particles. Our critical lag time of $1.4 \mu\text{s}$ is smaller than the previous studies
207 that distinguished the BC-containing aerosols between thinly coated BC and thickly
208 coated BC with a lag time of $2 \mu\text{s}$ (Moteki and Kondo, 2007; Metcalf et al., 2012), 1.8
209 μs (Metcalf et al., 2012), and $4.2 \mu\text{s}$ (Liu et al., 2010), which was determined by the
210 internal setup up of the SP2. The critical lag time for the PKU and CP sites were $1.3 \mu\text{s}$
211 and $1.7 \mu\text{s}$, respectively.

212 For each type of BC-containing aerosols, we calculated the coating thickness
213 probabilities and the results are shown in Fig. 2(d), (e) and (f) for the PKU, CP, and LJ
214 sites, respectively. Results showed that the BC-containing aerosols were mainly
215 composed of thickly coated BC aerosols and thinly coated BC aerosols. The coating
216 thickness for the thinly coated BC-containing aerosol was smaller than that of the thickly
217 coated BC-containing aerosols. However, the coating thickness of the thickly coated
218 BC-containing aerosols spread wider than that of the thinly coated ones.

219 The number fractions of the thickly coated BC-containing aerosols were
220 significantly different for different atmospheres as shown in Fig. 2 (g), (h), and (i). At
221 the polluted urban site, the number concentration of the thickly coated BC-containing
222 aerosols was comparable to that of the thinly coated BC-containing aerosols with the
223 number fractions of 56% and 44% for the thinly coated and thickly coated BC particles,
224 respectively. The number fraction of the thickly coated BC aerosols at the CP site was
225 67 %. However, the BC-containing aerosols at the background LJ site were dominated
226 by thickly coated ones with a number fraction of 81%.

227 The difference in the number fraction of the thickly coated BC particles was
228 synthetically influenced by the ambient pollution levels and the sources of the BC
229 aerosols. The suburban site CP had the largest number fraction of the thinly coated BC
230 particles. The CP site is not far from the urban, and thus the thinly coated BC particles
231 from the traffic contribute a large amount of the total ones. The urban site PKU had a
232 larger number fraction of the thickly coated BC than that of the CP site. This might be
233 resulted from the PKU site being more polluted than the CP site and then the aging
234 processing at the PKU site was faster than that at the CP site. The LJ site is far from the
235 traffic sources. The measured BC particles at the LJ site were mainly from long-range
236 transportation and experienced a long time of aging process than that at the CP and PKU

237 sites. Therefore, the BC-containing aerosols were dominated by the thickly coated ones
238 at the LJ sites.

239 We compared the number fraction of the thickly coated BC at different measurement
240 sites from literature (Ueda et al., 2016; Schwarz et al., 2008a; Wang et al., 2017c; Wang
241 et al., 2017a; Wu et al., 2017; Wang et al., 2017b; Wang et al., 2014; Huang et al., 2012;
242 Metcalf et al., 2012; Wang et al., 2016; Shiraiwa et al., 2007; Mcmeeking et al., 2012;
243 Subramanian et al., 2010; Schwarz et al., 2008b; Saha et al., 2018; Krasowsky et al.,
244 2018; Holder et al., 2014) and the results are shown in Fig. 3. The number fraction
245 values were divided into three different kinds of groups, namely the roadside, urban or
246 suburban, and background. Results from Fig. 3 show that the number fractions at the
247 roadside tend to be the lowest. These sites were close to the traffic sources and the
248 measured BC-containing aerosols were mainly from the traffic. The left part of the green
249 circles corresponds to the relatively clean urban or suburban sites with the number
250 fractions of the thickly coated BC around 30%. However, the number fractions of the
251 relative polluted urban or suburban sites had a larger number fraction of the thickly
252 coated BC around 50%. The number fractions of the thickly coated BC at the
253 background sites were the largest. Therefore, the number fractions of the thickly coated
254 BC-containing aerosols were synthetically influenced by the distance from the primary
255 source and the pollution levels of the ambient atmosphere. The number fraction of the
256 thickly coated BC-containing aerosols increased with the distance from the primary
257 emission sources and the pollution levels. Our results were consistent with the aerial
258 measurement by Metcalf et al. (2012), who found that the number fraction of the thickly
259 coated BC was 29%~41% at the top of the Los Angeles city and 47%-54% for the out
260 plume of this city.

261 For a better understanding of the source of the thinly coated and thickly coated BC,
262 we compared the number concentrations of the BC-containing aerosols with the source

apportionment results from the AMS data for the CP site. Among the PMF results, the factor of hydrocarbon-like organic aerosol (HOA) is mainly composed of long-chain hydrocarbon, and oxygenated organic aerosol (OOA) is mainly from the secondary formation. HOA is mainly from the diesel exhaust, gasoline exhaust, and lubricating oil emission. From Fig. 4(a), the number concentration of the thinly coated BC and mass concentration of HOA showed good consistency, with R^2 equaling 0.69 as shown in Fig. S10, which further proved the evidence that the thinly coated BC-containing aerosols were from the traffic sources. The time series of the thickly coated BC and OOA showed good consistency as shown in Fig. 4 (b), with R^2 equaling 0.87. Therefore, the aging processing of the ambient BC was accompanied by the ambient OA. The mass concentration of OOA and the number concentration of thickly coated BC can be used as good indicators for each other.

3.3 Size distributions of the thinly coated and thickly coated BC-containing aerosols

The size distributions of the BC-containing aerosols exert a significant influence on their corresponding radiative effects (Zhao et al., 2019; Matsui et al., 2018). We calculated the number size distribution (NSD) of BC-containing aerosols for the thinly coated and thickly coated ones at different sites, and the results are shown in Fig. 5. It should be noted that the D_p in Fig. 5 corresponds to the mobility diameter from the DMA. The BC-containing aerosol NSD was further fit using the log-normal distribution.

As for the thinly coated BC-containing aerosols, the geometric mean diameters were 193, 161, and 162 nm for the PKU, CP, and LJ sites, respectively. The geometric standard deviations (GSD) of the BC-containing aerosol NSD were 1.50, 1.63, and 1.91 for the PKU, CP, and LJ sites, respectively. The GSD to some extent reflects the diversity of the BC sources. The LJ site had the largest GSD, which indicated multiple

sources of thinly coated BC-containing aerosols. The LJ site was highly influenced by atmospheric transportation, due to the high altitude of this location (Zheng et al., 2017; Tan et al., 2021). Therefore, the thinly coated BC-containing aerosols could be originated from different orientations. As for the urban site PKU, the thinly coated BC aerosols were mainly from urban lifestyle emissions. Therefore, the thinly coated BC aerosols at the PKU site had the lowest value of the GSD. However, the thinly coated BC aerosols at the suburban site CP were influenced synthetically by urban lifestyle sources and some other sources from suburban, and thus had a larger value of GSD than that of PKU.

As for the thickly coated BC, it is obvious that they had larger diameters than those of the thinly coated BC due to the coating of other non-BC components. The geometric mean D_p values of the thickly coated BC were 294, 244, and 257 nm for the PKU, CP, and LJ sites, respectively. The corresponding GSD values were 1.37, 1.41, and 1.46.

Based on the above results, the geometric mean D_p values of the thickly coated BC aerosols were larger than that of the thinly coated BC aerosols by 52%, 52%, and 59% for the PKU, CP, and LJ sites, respectively. The GSD values of the thickly coated BC were consistent with that of the thinly coated BC with the lowest value at the PKU site and highest value at the LJ site, which is consistent with the diversity of the sources of BC-containing aerosols. For each site, the GSD values of the thickly coated BC aerosols were smaller than that of the thinly coated ones. The GSD of BC-containing aerosols tends to be smaller during the aging processing because the increment of the diameter should decrease with the diameter.

3.4 Size distribution of the thinly coated and thickly coated BC core

The number and mass concentrations of the BC core under different mass equivalent diameters were calculated and the results are shown in Fig. 6 and Table 1. It should be

313 noted here that, when it comes to the BC size distribution, the mass-equivalent diameter
314 of BC cores (D_{me}) (assuming a density of 1.8 g/cm³) was adopted in this study for direct
315 comparison with previous studies.

316 As for the number size distribution of the BC core, the geometric mean D_{me} of the
317 thinly coated BC particles were 115, 107, and 127 nm, for the PKU, CP, and LJ sites
318 respectively. The corresponding GSD values are 1.58 1.53 and 1.68, respectively. The
319 D_{me} for the thickly coated BC particles were 114, 95, and 111 nm for the PKU, CP,
320 and LJ sites respectively and the corresponding GSD values were 1.40, 1.45, and 1.43,
321 respectively. Both the GSD and the D_{me} of the thickly coated BC were smaller than
322 that of the thinly coated BC. The overall geometric mean diameter of the BC core
323 number size distributions are 114, 100, and 111 nm for the PKU, CP, and LJ sites
324 respectively.

325 There are mainly three possible reasons that may lead to the rather smaller geometric
326 mean diameter for the thinly coated BC than the thickly coated BC. First, the smaller
327 BC core tends to have a higher time lag as a smaller BC core will take a longer time to
328 evaporate the coating on it and thus the thinly coated particles tend to have smaller core
329 diameters. Second, it takes less time for the smaller BC particles to grow the same
330 amount of coating thickness when the increment of the BC particles was dominated by
331 condensation Thirdly, the small BC particles may have a longer life than the large BC
332 particles.

333 As for the mass size distribution of the BC core, the geometric mean D_{me} of the
334 thinly coated BC were 187, 182, and 238 nm for the PKU, CP, and LJ sites respectively
335 and the corresponding GSD values were 1.35, 1.48, and 1.47. The overall geometric
336 mean diameter of the BC core mass distributions are 172, 169, and 181 nm for the PKU,

337 CP, and LJ sites respectively. The geometric mean diameter of the BC core mass
338 distributions of 172 nm in PKU was slightly smaller than that of Liu et al. (2019a), with
339 a geometric mean diameter of 195 nm in another measurement in the urban environment
340 in Beijing and comparable to ~~the of~~ Zhang et al. (2018a) with a geometric mean diameter
341 of the BC core around 180 nm.

342 **3.5 Morphology of the BC-containing aerosols**

343 The time series of the number fractions of the attached BC-containing aerosols to
344 the total BC- containing aerosols (f_{attached}) are shown in Fig. 7. From Fig. 7, the f_{attached}
345 ranged between 0 and 0.21 with a mean value of $7.2 \pm 3.7\%$, $11.0 \pm 3.7\%$, and $10.1 \pm$
346 4.1% . Moteki et al. (2014) found that the f_{attached} was generally less than 0.1 in Tokyo.
347 The f_{attached} ranged between 3% and 16% in suburban London (Liu et al., 2015). A mean
348 value of 12% was found for biomass burning particles using electron microscopy (China
349 et al., 2013). Our measurement results were consistent with the previous studies. The
350 f_{attached} tend to increase with the $\text{PM}_{2.5}$ for different sites, which may indicate that the
351 attached BC-containing aerosols were generated from the coagulation of BC and non-
352 BC aerosols.

353 We calculated the f_{attached} under different aerosol diameters and the results are shown
354 in Fig. 8. There were few attached BC-containing aerosols when the diameter was
355 smaller than 250 nm with f_{attached} lowing than 2%. The f_{attached} increased with the diameter
356 for all of the measurement sites. It could reach 30% for the LJ sites. Based on the results
357 from the electron microscopy, the BC volume fractions are smaller than those of the
358 non-BC volume fractions in the attached BC aerosols (Moteki et al., 2014). The
359 increment of f_{attached} with D_p is essentially consistent with the results from Hu et al.
360 (2021a) that larger D_p contains more fractal BC, which is hard to be enveloped by
361 coatings. Our results further indicate that the attached BC aerosols were formed from

coagulation, as the coagulation efficiency of the two particles increased with the difference between their sizes (Cai and Jiang, 2017; Kim et al., 2016; Mahfouz and Donahue, 2021).

Under the heavier pollution, more secondary aerosol forms and more condensation process would on one hand increase the coating of the previously coated BC particles, which would not increase the number fraction of coated BC. On the other hand, the condensation process would coat on the attached BC particle and to some content would lead to the transformation from the attached BC to coated BC particles. Based on our measurement results, the above process of transformation from attached BC to coated BC may not be comparable to the process of coagulation between thinly coated BC and non-BC particles, which would lead to the increment of the fraction of attached BC with the pollution levels.

The f_{attached} under different aerosol number concentrations (N) and different ratios of the BC-free aerosol number concentrations to the BC-containing aerosol number concentrations are shown in Fig. 9. Results showed that the f_{attached} increased with the above two factors. The results were consistent with the fact that the coagulation between BC and non-BC components is more likely to happen with the increment of the BC-free aerosol number concentrations. Based on the analysis above, we concluded that the attached BC- containing aerosols are mainly formed through coagulation.

4 Conclusions

In this study, the BC microphysical properties were studied based on field measurement using the DMA-SP2 system at the urban site PKU, suburban site CP and a background site LJ in China.

385 The number fractions of the thickly coated BC-containing aerosols were 49%, 33%,
386 and 79% for the PKU, CP, and LJ sites respectively. The mass concentrations of the
387 thinly coated BC-containing aerosols showed good consistency with that of HOA,
388 which indicated that the thinly coated BC-containing aerosols were mainly generated
389 from the emission of vehicles. The thickly coated BC-containing aerosols are highly
390 correlated with the OOA.

391 The geometric diameter of the thinly coated BC-containing aerosols ranged between
392 160 nm and 200 nm, while the corresponding range was 240~300 nm for the thickly
393 coated BC-containing aerosols. The GSD of the BC-containing aerosols decreased
394 during the aging process. The corresponding mobility diameters of these thickly coated
395 (thinly coated) BC-containing aerosols were 294 (193), 244 (161), and 257 (162) nm.
396 The measured thickly coated (thinly coated) BC core number median diameters were
397 115 (114), 107 (95), and 127 (111) nm for the urban, suburban, and background sites,
398 respectively. The corresponding thickly coated (thinly coated) core mass median
399 diameters were 187 (154), 182 (146), and 238 (163) nm respectively. The mean diameter
400 of the thickly coated BC-containing aerosols was larger than that of the thinly coated
401 BC-containing aerosols, while the mean BC core diameter of the thickly coated BC-
402 containing aerosols was smaller than that of the thinly coated BC-containing aerosols.
403 There are about 10% of the BC-containing aerosols with the BC core attached to the
404 other non-BC components. We concluded that these attached BC-containing aerosols
405 were mainly generated by coagulation between the BC and non-BC components even
406 though the aging of the ambient BC aerosols was driven by condensation.

407 **Data availability.** The data is available at <https://doi.org/10.5281/zenodo.5816310>.

408 **Author contributions.** **Gang Zhao:** Conceptualization, Writing - Original Draft,
409 Visualization, Software, **Tianyi Tan:** Data Curation, Conceptualization, Visualization,

410 **Shuya Hu:** Data Curation, Conceptualization, **Zhuofei Du:** Data Curation, **Dongjie**
411 **Shang:** Data Curation, **Zhijun Wu:** Data Curation, Conceptualization, **Song Guo:** Data
412 Curation, Conceptualization, **Jing Zheng:** Data Curation, Conceptualization, **Wenfei**
413 **Zhu:** Data Curation, Conceptualization, **Mengren Li:** Data Curation, Conceptualization,
414 **Limin Zeng:** Data Curation, Conceptualization, **Min Hu:** Resources, Supervision, Data
415 Curation, Conceptualization, Revision.

416 ***Competing interests.*** The authors declare that they have no conflict of interest.

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420

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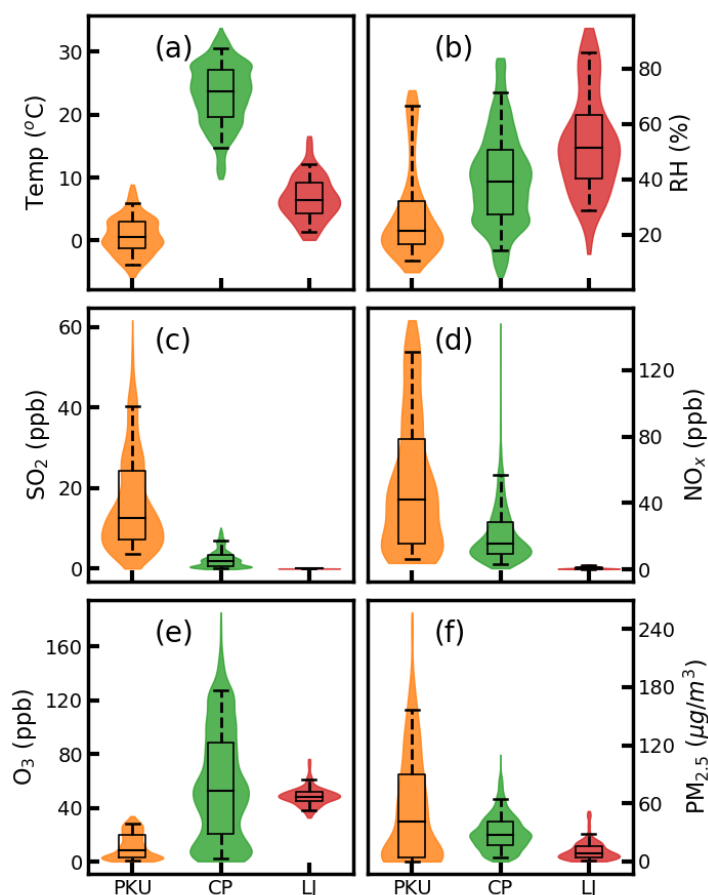
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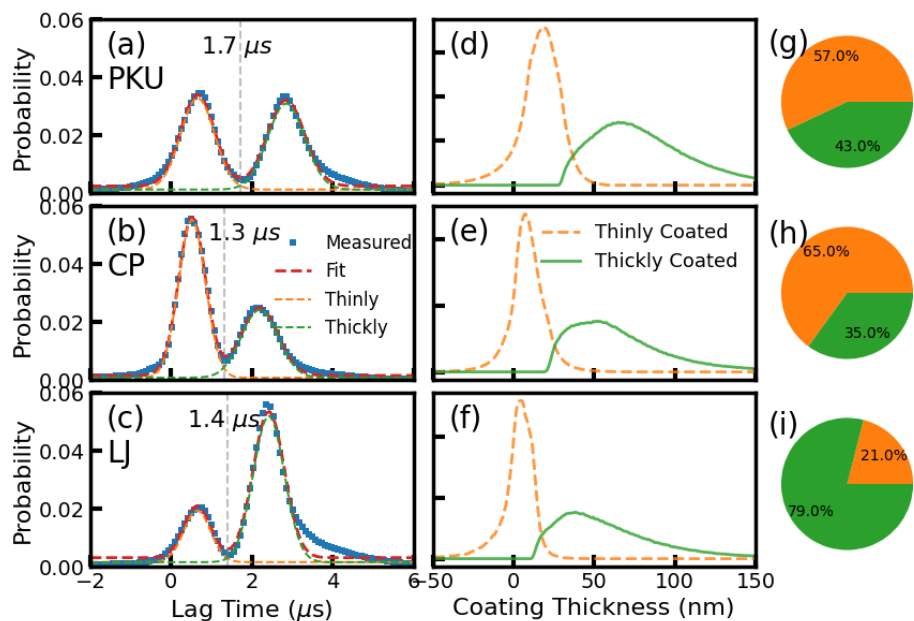
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674 **Figure 1.** The measured distribution of (a) temperature, (b) RH, (c) SO₂, (d) NO_x, (e)
 675 O₃ and (f) PM_{2.5} for PKU (orange), CP (green) and LJ (red) sites, respectively. The box
 676 and whisker plots represent the 5th, 25th, 75th, and 95th percentiles. The width of the
 677 filled colors represents the probability distributions of the corresponding measured
 678 values.

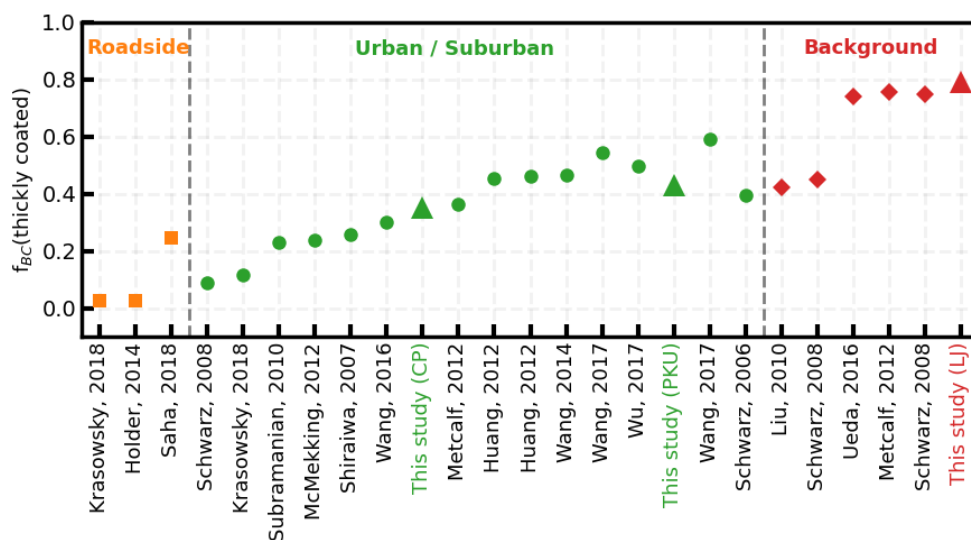
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681 **Figure 2.** (a) The measured probability distribution of the lag time for the PKU site.
 682 Panel (d) shows the corresponding coating thickness distributions of thinly coated
 683 (orange) and thickly coated (green) BC-containing aerosols. Panel (g) gives the number
 684 fraction of the thinly coated (orange) and thickly coated (green) BC-containing aerosols.
 685 Panel (b), (e), and (h) are the corresponding values for the CP site. Panel (c), (f), and (g)
 686 give the results for LJ sites.

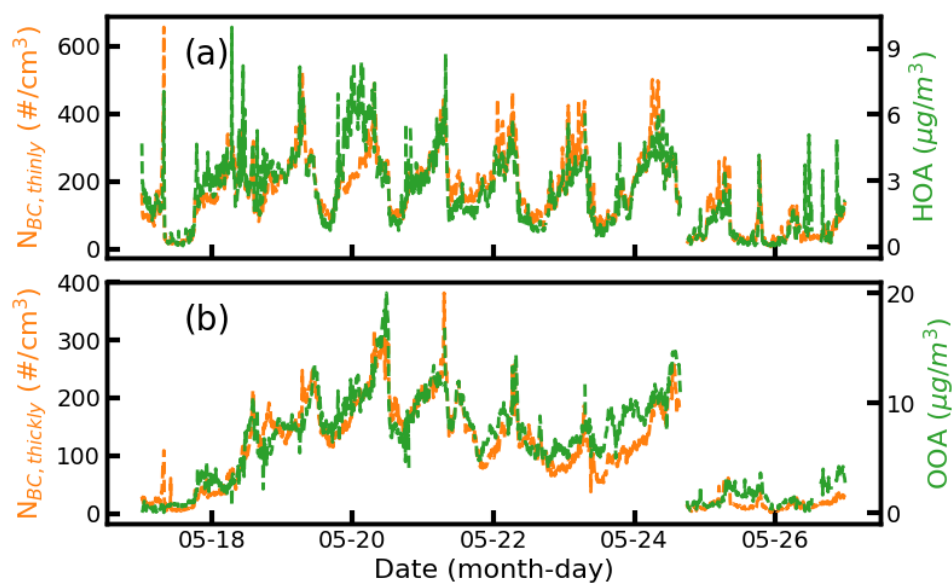
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689 **Figure 3.** Measured number fraction of the thickly coated BC under different
 690 atmospheric environments based on literature. Our measured values are shown as
 691 triangles.

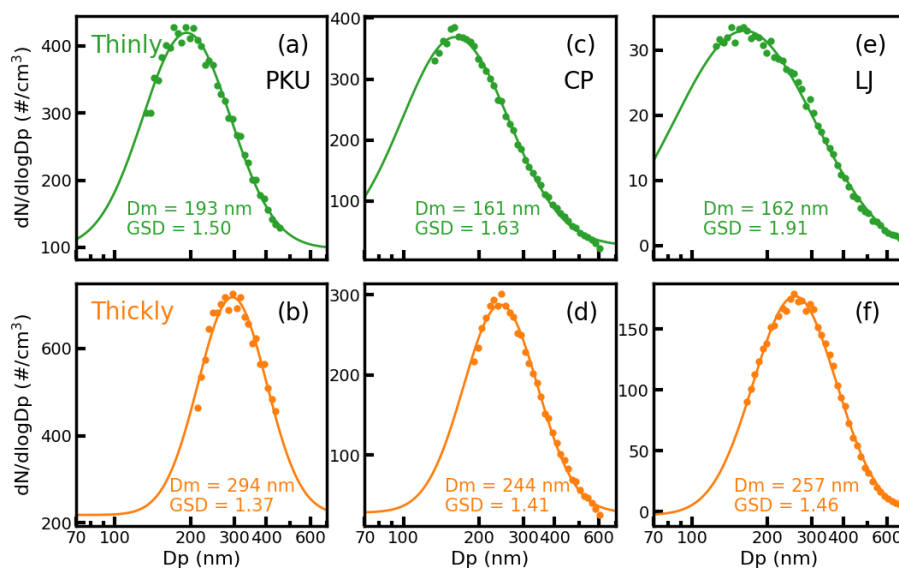
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693

694 **Figure 4.** The time series of (a) the number concentration of the thinly coated BC
 695 (orange) and the mass concentration of HOA (green), (b) the number concentration of
 696 thickly coated BC (orange), and the mass concentration of OOA (green) for the CP site.

697



698

699 **Figure 5.** The number size distributions of the thinly coated BC-containing aerosols
700 at (a) PKU, (c) CP, and (e) LJ sites. Panels (b), (d), and (f) are the number size
701 distributions of the thickly coated BC-containing aerosols for the PKU, CP, and LJ sites,
702 respectively. The dots in the figure are the measurement results and the lines are the
703 corresponding fit results with a log-normal distribution.

704

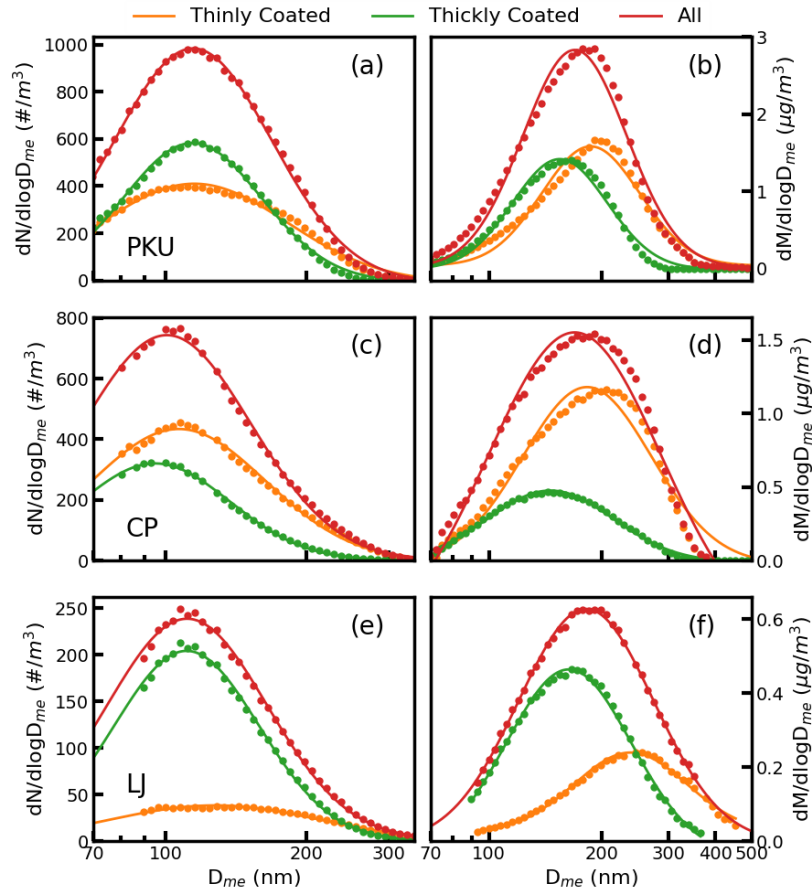
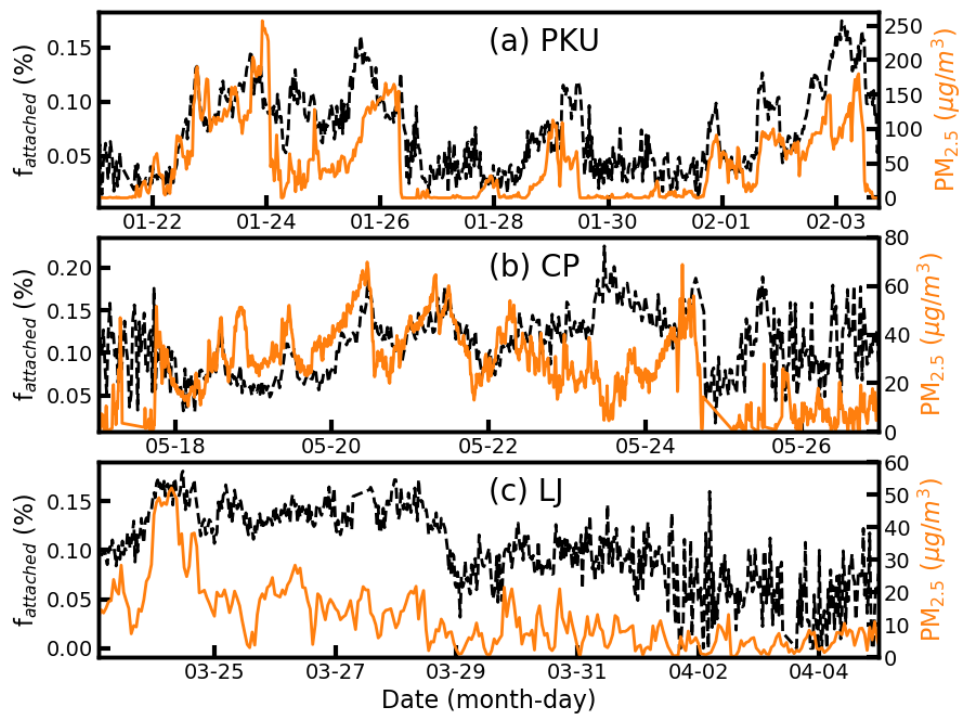


Figure 6. The BC core number size distributions of the thinly coated (orange), thickly coated (green), and overall (red) BC aerosols for the (a) PKU, (c) CP, and (e) LJ sites. Panel (b), (d) (f) show the BC core mass distributions of the thinly coated (orange), thickly coated (orange), and overall (red) BC aerosols for the PKU, CP, and LJ sites, respectively.

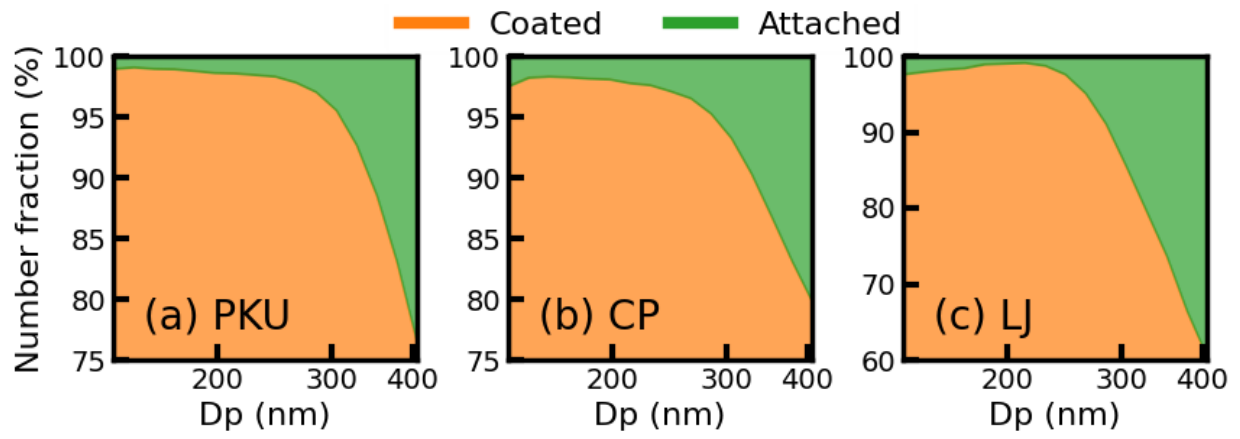


712

713 **Figure 7.** The time series of the number fractions of the attached BC (black) and PM_{2.5}
 714 mass concentrations (orange) for the (a) PKU, (b) CP, and (c) LJ sites.

715

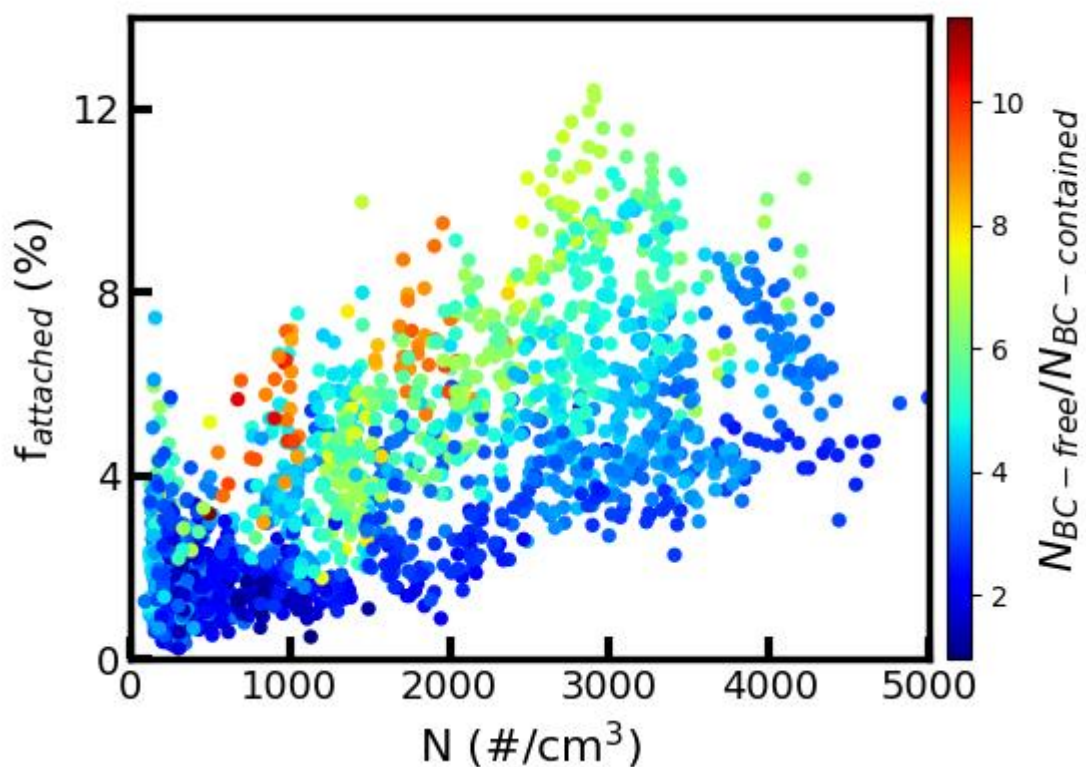
716



717

718 **Figure 8.** The number fractions of the coated and attached BC under different
719 diameters for the (a) PKU, (b) CP, and (C) LJ sites.

720



721

722 **Figure 9.** The number fractions of the attached BC aerosols under different total
 723 aerosol number concentrations for the CP sites. The filled colors represent the ratios
 724 between the BC-free aerosol number concentrations to the BC-containing aerosol
 725 number concentrations.

726

727 **Table 1.** The D_{me} and GSD values of the BC core at different sites.

Site	Value	Number Distribution			Mass Distribution ⁷²⁸		
		thinly coated	thickly coated	All	thinly coated	thickly coated	All
PKU	D_{me} (nm)	115	114	114	187	154	172
	GSD	1.58	1.40	1.47	1.35	1.34	1.37
CP	D_{me} (nm)	107	95	100	182	146	169
	GSD	1.53	1.45	1.51	1.48	1.47	1.47
LJ	D_{me} (nm)	127	111	112	238	163	181
	GSD	1.68	1.43	1.48	1.47	1.41	1.42