Black carbon content of traffic emissions significantly impacts black carbon mass size distributions and mixing states

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Received: 27 February 2023 – Discussion started: 27 March 2023
Revised: 10 May 2023 – Accepted: 15 May 2023 – Published: 14 June 2023

Abstract. Both the size and mixing state of black carbon (BC)-containing aerosols are crucial in estimating the environmental, health, and climate impacts of BC. Traffic emissions are a major global source of BC; however, parameterization of BC mass size distributions and mixing states associated with traffic remains lacking due to its dependence on vehicle types and driving conditions. To investigate BC mass size distributions and mixing states associated with traffic emissions, a field campaign was conducted in the Guangzhou urban area during winter, which used a system coupling a differential mobility analyzer (DMA) and a single-particle soot photometer (SP2) to measure BC mass size distributions in the range of 100 to 700 nm. The resolved primary organic aerosols were hydrocarbon-like organic aerosols (HOA) and cooking-like organic aerosols (COA), as well as refractory BC (rBC), which was detected by the DMA–SP2 and correlated highly with HOA ($R^2 = 0.88$), confirming that traffic emissions are the dominant source of atmospheric BC during the observations. The BC mass size distribution was found to be best fitted by a lognormal distribution, with a geometric mean ($D_{g,BC}$) of 258 ± 16 nm, varying between 200 and 300 nm. During daytime, active formation of secondary nitrate and organic aerosols reduced both $f_{BCc}$ and $f_{ext}$, with the decrease in $f_{ext}$ being more pronounced for larger particles, possibly due to the higher relative
coating thickness. Variations in $f_{\text{ext}}$ during nighttime were mainly controlled by the emission conditions. For example, $f_{\text{ext}}$ for 600 nm particles decreased from 0.82 to 0.46 as rBC / HOA increased from 1 to 3.5, while the mass ratios of secondary aerosols to rBC varied little, demonstrating that the BC content also significantly affects the mixing states of freshly emitted BC from traffic emissions. This study suggests that BC content can be used as the key factor to parameterize both the BC mass size distribution and mixing states from traffic emissions, which warrants future comprehensive investigation. In addition, other sources such as biomass burning and coal combustion also contribute substantially to BC emissions, and it was important to investigate whether BC content of other major BC sources than traffic is also important in determining BC mass size distributions and mixing states. Overall, results of this study have significant implications for accurate representation of BC from different sources when modeling the impacts of BC.

1 Introduction

Aerosols significantly impact human health through deposition on human tissues, visibility, weather and climate by interacting with solar radiation and acting as cloud condensation nuclei (CCN). Most of the atmospheric aerosols scatter readily while absorbing negligibly or little solar irradiation, but one exception is black carbon (BC), which absorbs solar irradiation strongly and thus heats the atmosphere. This strong absorption makes BC the second atmospheric warming component (Bond et al., 2013) and plays a major role in climate and air pollution (Zhang et al., 2019). Menon et al. (2002) found that BC’s absorption could affect trends of droughts and floods in India and China by altering regional atmospheric stability and vertical motions. The heating effects of BC (Wang et al., 2013) in the atmospheric boundary layer can suppress boundary layer turbulence, impacting boundary layer development and meteorology (Wilcox et al., 2016), and consequently affecting local haze formation (Ding et al., 2016). Moreover, BC-containing aerosols can interact with clouds and serve as CCN (Zhang et al., 2017; Motos et al., 2019; Hu et al., 2021), thus indirectly impacting climate (Koch and Del Genio, 2010). These effects of BC can be quantitatively determined through its radiative forcing. In addition, inhalation of atmospheric BC also poses a threat to human health; evidence suggests that it may be associated with changes in subclinical cardiovascular health effects in individuals (Nichols et al., 2013).

The radiative and health effects of BC are highly dependent on its size and mixing state, as these factors determine the distribution of BC in aerosols and their optical properties (Bond et al., 2006), hygroscopic growth (Liu et al., 2013) and depositions in the human respiratory tract (Man et al., 2022). Detailed parameterizations of BC size and mixing states as well as aging of BC-containing aerosols in models are crucial for modeling environmental and climate effects of BC (Wang et al., 2018). Freshly emitted BC can be either almost bare or coated with other materials, usually organic aerosols, which can then undergo rapid aging processes through vapor condensation or serve as medium for reactions (Zhang et al., 2018a, 2021; F. Zhang et al., 2020). This leads to size and morphological changes in BC-containing aerosols (Zhang et al., 2008) and influences their physical properties, such as hygroscopicity (Liu et al., 2013) and activation abilities (Ding et al., 2019; Yu et al., 2022). The coating of other components on BC can also significantly affect the optical properties of BC-containing aerosols, such as the lensing effect that enhances light absorption (Bond et al., 2006; Peng et al., 2016). This effect is non-linear (Liu et al., 2017; Wang et al., 2021) and is highly dependent on the mixing state heterogeneity of BC-containing aerosols (Fierce et al., 2020; Zhao et al., 2021; Zhai et al., 2022). In addition, Zhao et al. (2019) showed that the BC mass size distributions also play a major role in the direct radiative effects of BC. The size and mixing states of BC, as well as the chemical composition of its coatings, vary significantly with sources (Y. Zhang et al., 2020). This results in marked differences in the aging processes of freshly emitted BC from different sources in the atmosphere, depending on the emission sources and meteorological conditions in a given location.

Guangzhou is an expansive metropolis in the highly industrialized Pearl River Delta (PRD) region of China. Previous studies have shown that emissions from fossil fuel combustion are major sources of BC (Liu et al., 2014), while biomass burning emissions might also make certain contributions during autumn and winter (Sun et al., 2020). However, recent studies on source apportionment have not detected obvious signals attributed to biomass burning in autumn and winter, suggesting that other sources such as traffic activities are the main contributors to BC emissions (Guo et al., 2020; W. Chen et al., 2021; Liu et al., 2022; Zhai et al., 2023). Few studies have used the single-particle soot photometer (SP2) to measure bulk BC mass concentrations and mixing states in the Guangzhou urban area (Huang et al., 2011; Tao et al., 2021). Furthermore, no comprehensive measurements that characterize both size distribution and mixing states of BC in this region have been conducted, and the factors that control variations in BC mass size distributions and mixing states remain unknown. BC emissions from diesel vehicles dominate traffic BC emissions (Bond et al., 2013) and depend on many factors, such as fuel type, engine operating conditions, engine types, driving patterns, and environmental...
conditions (Adler et al., 2010). These conditions have a significant impact on the size distributions and mixing states of emitted particles (Lähde et al., 2011; Xu et al., 2014); therefore, BC mass size distributions and mixing states vary a lot in real traffic conditions. While numerous studies have examined the BC size and mixing states of emissions from various types of vehicles (Adler et al., 2010; Liu et al., 2017), only a few have directly investigated the BC size distribution and mixing states as a function of aerosol mobility diameter using the DMA–SP2 system, which couples differential mobility analyzer (DMA) and SP2 (Raatikainen et al., 2017), and reported the average coating characteristics of aerosol particles emitted from diesel vehicle exhaust (Han et al., 2019; Y. Zhang et al., 2020). However, the variations of BC mass size distributions and BC mixing states from real traffic emissions using the DMA–SP2 system have rarely been studied, and how to parameterize them remains elusive. This study carried out a field campaign and employed the DMA–SP2 system to investigate the dominant contribution of traffic emissions to atmospheric BC, which provided an ideal opportunity to evaluate how primary traffic emissions and their subsequent aging can affect BC mass size distributions and mixing states.

2 Materials and methods

2.1 Campaign information

The campaign was conducted to characterize BC mass size distributions and mixing states from 11 January to 27 February 2022 at the Haizhu wetland park (23°05′N, 113°22′E) in Guangzhou. The instruments used for characterizing aerosol chemical and physical properties included a quadrupole aerosol chemical speciation monitor (Q-ACSM) for monitoring aerosol chemical compositions, a DMA–SP2 system for measuring BC mass size distributions and mixing states and a scanning mobility particle sizer (SMPS) system for measuring aerosol particle number size distributions ranging from 13 to 800 nm. An AE33 aethalometer (Drinovec et al., 2015) was used to measure aerosol absorptions at multiple wavelengths and indirectly measure bulk BC mass concentrations. A PM$_{2.5}$ inlet (BGI, SCC 2.354) with a flow rate of 8 L min$^{-1}$ was used for aerosol sampling. The flow rates of the Q-ACSM, condensation particle counter (CPC), SP2 and AE33 instruments were set to 3, 0.3, 0.1 and 5 L min$^{-1}$, respectively, to meet the flow rate requirement of the impactor although there are some deviations. All instruments were housed in a temperature-controlled container (23–27°C) and placed downstream of a Nafion drier designed to lower the sample RH to less than 35% (placed outside of the container and vertically to ensure a straight line of the sampling route so that sampling loss of aerosols can be minimized). Meteorological parameters such as temperature, wind speed and direction, and relative humidity (RH) were measured using an automatic weather station. Further details about this site can be found in Liu et al. (2022). In addition, concentrations of PM$_{2.5}$ and nitrogen dioxide (NO$_2$) were obtained from the China National Environmental Monitoring Center network, which is publicly available (http://www.cnemc.cn/en/, last access: 7 June 2023); there is a site located within 5 km distance to our observation site.

2.2 DMA–SP2 system and data processing

The SP2 (Droplet Measurement Technologies) can measure aerosol scattering and incandescence signals of individual particles and identify if they contain detectable BC mass. It can also provide BC mass concentrations at the single-particle level, thus allowing for the determination of BC mixing states. The scattering signals can be used to estimate the particle size of each BC-free particle; however, a leading-edge-only method is required for sizing BC-containing particles (Schwarz et al., 2006), with the estimated optical-equivalent size potentially deviating substantially from the mobility size due to variations in aerosol refractive index and morphology. As such, the use of the DMA–SP2 system to measure BC mass size distributions and mixing states has been previously proposed (Raatikainen et al., 2017; Han et al., 2019; Sarangi et al., 2020; Zhao et al., 2021), which complements the size measurements, and additional size information can be used to derive physical properties such as morphology and effective densities (Zhang et al., 2018b; Wu et al., 2019). A similar system has also been used for other applications, such as investigation of the hygroscopic properties of BC-containing aerosols (McMeeking et al., 2011; Liu et al., 2013). The DMA–SP2 set-up of Zhao et al. (2021) employed a continuous scanning mode of the DMA, allowing for black carbon mass size distribution (BCMSD) measurements with a time resolution of 5 min. However, accurate matching of the time of particles in the DMA and SP2 is necessary. Some previous studies passed size-selected monodisperse aerosols to the SP2 at only a few diameters (Zhang et al., 2018b; Han et al., 2019), limiting the retrieval of BC mass size distributions and mixing states across the entire sub-micrometer diameter range. In this study, we developed a software which enables the DMA to scan at different diameters for different time periods depending on their number concentrations. For example, DMA scans at 100 nm last about 36 s, while scans at 700 nm last about 1.5 min, allowing for enough particles to be sampled at larger particle sizes. The diameter set points (18 points) of DMA scans are 100, 120, 160, 200, 235, 270, 300, 335, 370, 400, 435, 470, 500, 535, 570, 600, 635 and 700 nm, with a full scan taking 20 min.

With this set-up, the particle number size distributions of BC-containing and BC-free aerosols can be derived from the DMA–SP2 measurements using an inversion routine that mainly accounts for the effects of the DMA transfer function and multiple charge. Assuming a BC density of 1.8 g cm$^{-3}$, the BC volume equivalent diameter of each BC-
containing particle (termed as BC core diameter, $D_c$) can be calculated, assuming a core–shell structure. The particle number size distribution of BC-containing aerosol, containing information of BC mixing states, can then be described using a two-variable formulation:

$$\frac{dN}{\partial \log(D_p) \partial \log(D_c)} = \text{terms accounted for the impacts of multiple charge corrections}$$

and the multiple charge correction method proposed by Zhao et al. (2021) was used here to account for the impacts of multiple charge corrections for the BC mass of each $D_c$. Details about the inversion routines are introduced in Sect. S1 of the Supplement. Note that the effective density of bare BC or BC contained in a particle can vary substantially due to BC morphology and existence of air voids (Zhang et al., 2016; Zhao et al., 2020b). Therefore, a simple assumption of 1.8 g cm$^{-3}$ for BC density could bring uncertainties to $D_c$ calculations. In addition, the optical equivalent diameter of BC-containing aerosols cannot be retrieved in this study due to the failure of the SP2 split channel hardware during the campaign, which rendered the leading-edge-only method unusable.

The average particle number size distribution (PNSD) derived from the DMA–SP2 system was compared with the one obtained from independent SMPS measurements, as shown in Fig. 1a. The detection limitations of the SP2 scattering channels (Raatikainen et al., 2017) caused the PNSD from SP2 to be markedly lower than that from SMPS for diameters $<200$ nm. For diameters larger than 200 nm, the PNSD from SP2 was generally consistent with that from SMPS, with the average ratio of SMPS to SP2 measurements being 0.89 ± 0.05, which is similar to the phenomenon reported in Raatikainen et al. (2017), with an average ratio of 0.82. Additionally, the observed rBC mass concentrations correlated highly with the optically equivalent BC mass concentrations reported by the AE33 aethalometer ($R^2 = 0.96$ and an average ratio of 0.96), as shown in Fig. S4. The consistency tests between the DMA–SP2 system and SMPS measurements validated the number size distributions and BC mass concentrations inverted from DMA–SP2 measurements.

The number fractions of BC-containing aerosols of various diameters can be calculated using the DMA–SP2 measurements. Based on the time lag between the peak time of the scattering and the incandescence signal (Schwarz et al., 2006; Moteki and Kondo, 2007; Sedlacek et al., 2012), these aerosols can be roughly divided into two categories: bare/thinly coated BC particles and thickly coated BC particles. The time lag distribution of pure BC aerosols can be identified from the SP2 calibrations using bare BC aerosols. Consequently, bare/thinly coated BC particles can be identified using the calibrated critical lag time.

2.3 Q-ACSM measurements and positive matrix factorization (PMF) analysis

The Q-ACSM measured non-refractory sub-micrometer (NR-PM$_1$) species, including organic aerosol (OA), sulfate (SO$_4$), nitrate (NO$_3$), ammonium (NH$_4$) and chloride (Cl), at a time resolution of 15 min. A more detailed description can be found in Liu et al. (2022) and Ng et al. (2011). The mass spectra measured by the Q-ACSM were analyzed using ACSM standard data analysis software (ACSM Local 1.5.10.0 released 6 July 2015), written in Igor Pro (version 6.37). The composition-dependent collection efficiency (CE) parameterization scheme proposed by Middlebrook et al. (2012) was used to calculate the mass concentrations of OA and inorganic species. This was also detailed in Liu et al. (2022). As calibration of the Q-ACSM was not available during this campaign, relative ionization efficiencies (RIE) of 5.15 and 0.7 for ammonium and sulfate from previous calibrations were used, while the default RIEs of 1.4, 1.1 and 1.3 were used for organic aerosol, nitrate and chloride, respectively. The quality assurance of the Q-ACSM measurements was first performed by comparing the mass concentrations of PM$_1$ (summation of measured NR-PM$_1$ concentrations and rBC concentrations measured by the DMA–SP2 system) with PM$_1$ mass concentrations calculated from the particle number size distribution measurements of SMPS, assuming an aerosol density of 1.6 g cm$^{-3}$. Good consistency (as shown in Fig. 1b) was achieved between SMPS and Q-ACSM measurements ($R^2 = 0.99$), with NR-PM$_1$ + rBC values slightly lower than PM$_1$ concentrations from SMPS (an average ratio of 1.19). Three reasons explain the deviations between SMPS and Q-ACSM plus rBC measurements: (1) the assumed average aerosol density may be biased from the real variations; (2) some aerosol species are not measured by the Q-ACSM, such as sub-micrometer dust; and (3) Q-ACSM has uncertainties of about 30 % (Ng et al., 2011; Fröhlich et al., 2013), and uncertainties of SMPS differ among different diameter ranges (Wiedensohler et al., 2012).

Following the same procedure of the PMF analysis for the Q-ACSM measurements introduced in Liu et al. (2022) and Zhai et al. (2023), the PMF technique with the multilinear engine (ME-2) (Canonaco et al., 2013, 2021) was applied to ACSM spectra for deconvolving OA into different factors and detailed in the Supplement. In total, four factors were identified, including two primary OA (POA) factors – a hydrocarbon-like OA (HOA, O/C ∼ 0.16) and a cooking-like OA (COA, O/C ∼ 0.14) – and two oxygenated OA factors – a less oxidized oxygenated OA (LOOA, O/C ∼ 0.89) and a more oxidized oxygenated OA (MOOA, O/C ∼ 0.94). SOA was represented by the summation of LOOA and MOOA as done in previous studies (Kuang et al., 2020). The mass spectra of these factors, the determination of the factor number, the selection of solutions and more details about the factor analysis can be found in Sect. S2 of the Supplement.
3 Results and discussion

3.1 Overview of aerosol chemical compositions

During the observation period, the PM\(_{2.5}\) mass concentration varied significantly (from 1 to 126 µg m\(^{-3}\)), with an average of 20 µg m\(^{-3}\), and several pollution episodes were observed during relatively stagnant conditions when wind speeds were near or below 1 m s\(^{-1}\). The time series of meteorological parameters as well as PM\(_{2.5}\), ammonium sulfate (AS), ammonium nitrate (AN), SOA, HOA, COA and rBC are shown in Fig. 2. The scheme proposed by Gysel et al. (2007) was used to identify AS and AN. On average, secondary aerosols including nitrate, sulfate, ammonium and SOA together accounted for about 80 % of non-refractory PM\(_1\) mass concentration, and secondary aerosols increased substantially during pollution episodes, demonstrating active secondary aerosol formations during the observations which might significantly impact BC mass size distributions as well as BC mixing states. The average air RH during the observations varied a lot from 42 % to 98 % with an average of 76 %, suggesting that the heterogeneous reactions that involve aerosol water were favored during this campaign, which is consistent with the quick formation of ammonium nitrate in pollution episodes. On average, AN, AS and SOA accounted for 33 %, 25 % and 42 % of identified NR-PM\(_1\) secondary components respectively, which is consistent with Zhai et al.'s (2023) conclusion that nitrate concentrations are higher than sulfate during winter in the Guangzhou urban area, especially under pollution conditions. The time series of rBC mass concentrations were shown in Fig. 2d, with rBC mass concentrations ranging from about 0.1 to 20 µg m\(^{-3}\) with an average of 2.3 µg m\(^{-3}\). Resolved POA factors were HOA and COA, which is consistent with results of previous studies in recent years that traffic emissions and cooking emissions are two main sources of primary aerosols in the Guangzhou urban area (Guo et al., 2020; W. Chen et al., 2021; C. Chen et al., 2021). The rBC correlated highly with HOA (\(R^2 = 0.88\)), demonstrating that traffic emissions contributed dominantly to atmospheric BC during the observations.

3.2 Overview of DMA–SP2 measurements

The observed average BC mass size distribution, as shown in Fig. 3a, exhibits a single lognormal mode for diameters greater than 100 nm, with a fitted geometric mean (\(D_{g, BC}\)) of 258 (±16) nm ranging from 200 to 300 nm. The formula form of fitting is introduced in Eq. (S1) of the Supplement, and the mean of fitted geometric standard deviation (\(\sigma_g\)) is 1.69. A small mass mode might exist for diameters less than 100 nm; however, it cannot be characterized due to the detection limitation of the SP2, which measures BC-containing particles with a \(D_c\) larger than 80 nm. Previous studies have reported BC mass size distribution as a function of rBC core diameter (Kompalli et al., 2020; Liu et al., 2019). The retrieved \(D_{g, BC}\) is higher than most mass median diameters of rBC core measured in urban environments, near 200 nm in urban Beijing Liu et al. (2019), which is reasonable due to intrinsic coatings (Adler et al., 2010). Some prior studies reported BC mass size distribution as a function of mobility diameter \(D_p\) measured by coupling the DMA with aethalometer (Stabile et al., 2012; Ning et al., 2013; Zhao et al., 2019). A few studies also reported BC mass size distribution as a function of aerodynamic diameter using the size-segregation filtering method (Hu et al., 2012). Zhao et al. (2019) reported bimodal characteristics of BC mass size distribution in the North China Plain, with the second mode accounting for most of the rBC mass and a \(D_{g, BC}\) of the coarse mode ranging from 430 to 580 nm, which is much higher than the average one reported here. This difference may be attributed to the markedly different sources of rBC (Y. Zhang et al., 2020) and the different roles of BC-containing aerosols in the formation of secondary aerosols.

The BC mixing state is an essential factor for determining BC’s climate effects. Figure 3b displays the average size-dependent fractions of BC-containing aerosols (\(f_{BCc}\))
and the fraction of identified externally mixed (bare/thinly coated) BC particles ($f_{\text{ext}}$) in all BC-containing aerosols. For diameters below 200 nm, not all BC-free aerosols were detected by the SP2, so the number concentration of all aerosols from SMPS measurements was used to calculate $f_{\text{BCc}}$. The results show a decrease in $f_{\text{BCc}}$ from 200 to 100 nm, reaching nearly 0.01 at 100 nm. This may be due to the lower fraction of BC-containing aerosols at smaller diameters, as well as the detection limit (∼80 nm) of the SP2, which may fail to detect many BC-containing aerosols below 80 nm. This can be further explained based on the measurements from the volatility tandem differential mobility analyzer (V-TDMA) in the Guangzhou urban area in previous studies. Cheung et al. (2016) and Tan et al. (2016) used the number fraction of remaining aerosols at 300°C in V-TDMA measurements to represent $f_{\text{BCc}}$, assuming that all BC-free aerosols had completely evaporated at this temperature. Their results showed an increasing trend in $f_{\text{BCc}}$ from 0.62 to 0.86 for diameters ranging from 40 to 300 nm, which is much higher than the values reported in this study (ranging from 200 to 700 nm, with an average of 0.24). Both methods may be biased in $f_{\text{BCc}}$ measurements due to the detection limit of BC mass in the SP2, which may underestimate $f_{\text{BCc}}$ (Zhao et al., 2020b), and the assumption that all BC-free aerosols have evaporated at 300°C in V-TDMA may overestimate $f_{\text{BCc}}$ by miscount-
ing some aerosols with extremely low volatility components (Tasoglou et al., 2020). Nevertheless, the low \( f_{\text{BCc}} \) values obtained from previous V-TDMA measurements confirm that \( f_{\text{BCc}} \) is smaller for smaller diameters (\( D_p < 200 \) nm). The size-dependent \( f_{\text{BCc}} \) is critical for simulating aerosol optical properties (Li et al., 2019) and CCN predictions (Ren et al., 2018). The facts that most BC masses reside in particles larger than 100 nm and most rBC masses would be detected by SP2 suggest that the \( f_{\text{BCc}} \) values reported from SP2 measurements are more suitable for use in aerosol optical simulations. As for \( f_{\text{ext}} \), the average \( f_{\text{ext}} \) shows a decreasing trend from 200 to 700 nm, with an average of 0.59 and with the \( f_{\text{ext}} \) at 100 nm being significantly affected by the detection limit. This suggests that BC is generally externally mixed during the observations, with a higher degree of aging for larger particles. The average distributions of rBC core at different diameters are shown in Fig. 3c, with a single mode at all diameters, deviating more from the 1 : 1 line at larger diameters, again indicating a higher degree of aging for larger particles. The estimated average coating thickness (CT) at different diameters is shown in Fig. 3d, with CT increasing from 29 nm at 200 nm to 164 nm at 700 nm and a relative coating thickness (RCT, \( D_p/D_C \)) ranging from 1.27 at 200 nm to 1.88 at 700 nm. The RCT of 200 nm is similar to that of BC-containing aerosols freshly emitted from diesel vehicles (Y. Zhang et al., 2020), consistent with diesel vehicle emissions being the dominant source of BC traffic emissions (Bond et al., 2013).

The average diurnal variations of POA, rBC and secondary aerosol components including SOA, AN and AS are depicted in Fig. 4a. In the morning, rBC and POA (HOA + COA) decrease due to dilution effects associated with boundary layer development, whereas SOA concentrations increase since 08:00 (all times are local time). However, POA and rBC begin to increase after 14:00, with the diurnal pattern of rBC being generally consistent with that of HOA. The rapid increase in COA after 17:00 does not lead to a significant rise in rBC, confirming that activities associated with cooking contribute negligibly to BC emissions. AN and AS begin to decrease after noon, with SOA continuing to increase until 20:00, which is in line with the findings reported in Zhai et al. (2023) that the highest SOA mass concentrations result from the coordination of daytime and nighttime SOA formation. The substantial decrease in \( f_{\text{BCc}} \) was observed in the morning when the prominent SOA formation occurs, with the decrease in \( f_{\text{BCc}} \) being greater as the particle size increases, suggesting that secondary aerosols are formed more efficiently on larger BC-free particles, which are then migrated to larger sizes. This is further supported by the diurnal variations of \( f_{\text{ext}} \), which revealed that secondary aerosol formation is more efficient in larger BC-containing aerosols (Fig. 4c). A decrease in \( f_{\text{ext}} \) from the morning to the afternoon was most prominent for aerosols at 500 nm (0.17), while a small decrease in \( f_{\text{ext}} \) (0.05) was observed at 200 nm. This is consistent with the findings from the coating thickness results in Fig. 3d, which showed that larger particles have higher coating thickness and are therefore likely to contain more aerosol water, thus favoring secondary aerosol formation via multiphase reactions.

3.3 Impacts of primary emissions and secondary aerosol formation on BC mass size distributions

The diurnal variations of \( f_{\text{ext}} \) revealed that secondary aerosols are formed on BC-containing aerosols, thus impacting mixing states, and this might also result in changes in BC mass size distribution. The average BC mass size distributions are shown in Fig. 5a and are normalized with rBC during the night (from local time 20:00 to 06:00 the next morning) and during the afternoon when active secondary aerosol formation is at its final stage and the impacts of accumulations of primary emissions are relatively small (local time 12:00 to 17:00). Figure 5a indicates that daytime secondary aerosol formation does not modify the shape of BC mass size distributions, which is confirmed by the variations of \( D_{BC} \) as a function of the SA / rBC ratio shown in Fig. 5b (where SA includes mass concentrations of sulfate, nitrate, ammonium and SOA). The DMA–SP2 measurements distinguish BC-free and BC-containing aerosols, allowing for the volume variations of BC-free and BC-containing aerosols with contributions of secondary aerosol formation to be differentiated. As seen in Fig. 5c, as SA / rBC increases, the total volume of BC-containing aerosols increases very slightly, while secondary aerosol formation mainly adds mass to BC-free aerosols, explaining the same BC mass size distribution shape during both daytime and nighttime. Many previous studies have demonstrated that BC can serve as sites for heterogeneous reactions (Khalizov et al., 2010) and may even promote secondary aerosol formation, thereby playing a significant role in haze formation (F. Zhang et al., 2020).

Results from Zhang et al. (2021) pointed out that BC promotes sulfate formation in the urban area of Guangzhou during summer. Our results indicate that BC plays a minor role in haze formation in Guangzhou during winter, which might have significant implications for haze formation mechanisms in this region.

The BC primary emissions and their subsequent aging in the air determine the observed BC mass size distributions. The aforementioned little influences of secondary aerosol formation on BC mass size distribution evolution suggest that primary emissions played a significant role in the observed variations in the BC mass size distributions. Traffic emissions dominate BC emissions during the observations as discussed before, and diesel vehicles contribute dominantly to BC emissions from traffic activities (Bond et al., 2013). The results from previous studies indicate that the ratio of elemental carbon (EC) to organic carbon (OC) changed significantly depending on the external factors (Adler et al., 2010; Lu et al., 2012), such as vehicle type, engine load and driving conditions. This ratio represents the emission conditions of
Figure 4. Average diurnal variations of (a) POA, SOA, ammonium nitrate (AN), ammonium sulfate (AS) and rBC; (b) \( f_{BCc} \) at different diameters; (c) \( f_{ext} \) at different diameters.

Figure 5. (a) Normalized average BCMSD during night and daytime; (b) average variations of \( D_{g,BC} \) as a function of the ratio SA / rBC – error bars are standard deviations; (c) scatter plots of SA / rBC and the ratio total volume (Vtot) to rBC of BC-free and BC-containing aerosols; (d) correlations between \( D_{g,BC} \) and HOA / rBC variations during night.

diesel vehicles, which also influences the size distributions of diesel exhaust particles (Lähde et al., 2011; Han et al., 2019). Here, we use the ratio rBC / HOA to represent different emission conditions related to traffic activities and investigate the potential effects of rBC / HOA variations on BC mass size distributions. To avoid the potential effects of secondary aerosol formation and daytime evaporation of HOA due to dilution, only data points from nighttime (from 18:00 to 06:00 the next morning) are used, the results of which are displayed in Fig. 5d. Our results showed for the first time that variations of \( D_{g,BC} \) are moderately correlated with rBC / HOA, and a linear relationship of \( D_{g,BC} = 34 \times \text{rBC} / \text{HOA} + 177 \) can be derived, indicating that a larger particle diameter of BC-containing aerosols is associated with a higher BC content.

Even though rBC and HOA during nighttime in this study are accumulated from different vehicle sources, the relationship between \( D_{g,BC} \) and rBC / HOA still holds, suggesting that the black carbon content might be used for parametrizing BC mass size distributions in traffic-related emissions.

3.4 Impacts of primary emissions and secondary aerosol formation on BC mixing states

As introduced in Sect. 3.1, both \( f_{BCc} \) and \( f_{ext} \) decreased during daytime due to secondary aerosol formation. Here, the variations of daytime (from 08:00 to 18:00) \( f_{BCc} \) and \( f_{ext} \) under different SA / rBC conditions were directly investigated and shown in Fig. 6. The \( f_{BCc} \) values ranging from 200 to 600 nm decreased from about 0.3 to around 0.175 as SA / rBC increased from 5 to 15 (Fig. 6a), highlighting significant impacts of secondary aerosol formation on \( f_{BCc} \). As discussed in Sect. 3.1, the decrease in \( f_{BCc} \) should be associated with the fact that secondary aerosols are formed much more quickly on BC-free aerosols than on BC-containing aerosols, which is consistent with the conclusion in Sect. 3.2 that secondary aerosol formation mainly adds mass to BC-containing aerosols. New particle formation also increases the number concentration of BC-free aerosols; however, its impact is limited for aerosol particles beyond 200 nm (Zhang et al., 2012). The evolution of particle number size distribution shape beyond 200 nm is mainly associated with vapor condensation, although coagulation also plays a role (Seinfeld and Pandis, 2016).

The variations in \( f_{ext} \) under different SA / rBC conditions are presented in Fig. 6b. Larger particles exhibited a more significant decrease in \( f_{ext} \), with SA / rBC increasing from 5 to 17.5; for example, \( f_{ext} \) of 200 nm particles only decreased from 0.75 to 0.6, while that of 600 nm particles decreased from 0.6 to 0.24. Aside from secondary aerosol formation, the emission conditions are also important factors that influence BC mixing states, particularly due to the contribution of the co-emitted intrinsic organic aerosols (Adler et al., 2010). At night, the shallow boundary layer facilitates the accumulation of freshly emitted aerosols, leading to increases
in the mass concentrations of rBC and POA (Fig. 4a) and the mixing of freshly emitted and aged aerosols. Even so, the variations of $f_{ext}$ during night under different rBC/HOA conditions might shed some lights on the impacts of primary emissions on BC mixing states, which is shown in Fig. 6c. As rBC/HOA increased from 1 to 3.5, $f_{ext}$ generally decreased, especially for particles larger than 300 nm. The $f_{ext}$ of 600 nm decreased from 0.82 to 0.46, which is higher than the degree of variations influenced by secondary aerosol formation as shown in Fig. 6b, and most importantly SA/rBC decreased from 9.4 to 6.9 as rBC/HOA increased from 1 to 3.5, demonstrating that the change in emission conditions has dominated the nighttime variations of BC mixing states. The increased internal mixing degree of BC observed under higher rBC/HOA conditions is also reflected in the variations of the average coating thickness of BC-containing aerosols. As seen in Fig. 6d, the coating thickness of 600 nm BC-containing aerosols increased from 118 to 134 nm as rBC/HOA increased from 1 to 3.5. Interestingly, as rBC/ HOA increases, the relative amount of coating to rBC decreases; however, both fractions of internally mixed BC and coating thickness increase, suggesting that a higher fraction of co-emitted intrinsic OA resides in BC-containing aerosols. This section demonstrates the significant impacts of emission conditions of traffic sources on BC mixing states.

4 Atmospheric implications

In this study, characterizations of BC mass size distributions and mixing state and their influencing factors were first investigated using measurements of the DMA–SP2 system and aerosol mass spectrometer in the Guangzhou urban area. Traffic emissions are the dominant source of atmospheric BC during the observations. The lognormal distribution represents well the BC mass size distribution, with the geometric mean varying between 200 and 300 nm with an average of 258 ± 16 nm. On one hand, the evidence that the secondary aerosol formation mainly adds mass to BC-free particles suggests that reactions occurring on or within BC-containing particles play a limited role in the formation of haze in the winter of Guangzhou. The urban area in Guangzhou is quite representative of most urban regions where traffic emissions are the primary source of BC emissions. Hence, the above finding has important implications for the haze formation mechanisms, particularly in southern China where primary aerosol emissions and meteorological conditions are similar to those in Guangzhou. On the other hand, it was found that the daytime secondary aerosol formation reduced both $f_{BC}$ and $f_{ext}$, with the decrease in $f_{ext}$ being more pronounced for larger particles, suggesting that secondary aerosols actually formed on BC-containing aerosols though their contribution to haze formation is small. The size and mixing states of BC-containing particles determine their optical and hygroscopic properties and are therefore critical factors for evaluating the environmental, health and climate effects of BC; however, these factors are not adequately considered in both chemical transport and climate models (Bond et al., 2013; Saleh, 2020). The finding that secondary aerosol formation has little effect on the BC mass size distribution suggests that the study provides an excellent case scenario to investigate how changes in traffic emissions affect the BC mass size distribution and mixing states. It further shows that BC content in traffic emissions has a significant impact on both the BC mass size distribution and mixing states, and the almost linear trends between $D_{g,BC}$ and rBC/ HOA suggest that BC content can be used as the key factor to parameterize both the BC mass size distribution and mixing states from traffic emissions, which warrants future comprehensive investigation.

Traffic is a major global contributor to atmospheric BC concentrations, but other major sources, such as biomass burning and coal combustion, as well as off-road diesel engines, also contribute significantly to BC emissions and play an even more important role than traffic in some regions (Bond et al., 2013). It is hence also important to investigate whether BC content of other major BC sources than traffic is important in determining BC mass size distributions and mixing states. Saleh et al. (2014) found that BC content has a major effect on the brownness of organic aerosols emitted from biomass burning, and results from several later studies further confirmed this finding (Luo et al., 2022). Re-
cently, Saleh (2020) discussed the potential for parameterizing the optical properties of brown carbon using BC content in climate models. Moreover, the results of Luo et al. (2022) demonstrated that BC content can also be used to parameterize the volume size distributions of aerosols emitted from biomass burning, indicating that BC content plays an important role in both traffic emissions and biomass burning emissions. These findings suggest that more comprehensive experiments should be designed in the future to investigate the factors that control variations in BC mass size distributions and mixing states, as well as to discuss how to use BC content from different major BC sources for parameterizing BC mass size distributions and mixing states.

Data availability. All data needed are presented in time series of the figures and Supplement figures; raw datasets of this study are available from the corresponding author Ye Kuang (kuangye@jnu.edu.cn) upon request.

Supplement. The supplement related to this article is available online at: https://doi.org/10.5194/acp-23-6545-2023-supplement.

Author contributions. YK and LL planned this campaign; YK conceived and led this research. FL performed the data analysis and wrote the manuscript together with YK. BL performed the DMA–SP2, SMPS and AE33 measurements together with FL and performed the post-processing of the SP2 data with help of GZ; MZ maintained the Q-ACSM during the observations and performed the PMF analysis. JZ acquired funding and supervised. TD, XD and HT helped with the data acquisition and revised the manuscript. All authors reviewed and edited the manuscript.

Competing interests. The contact author has declared that none of the authors has any competing interests.

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Acknowledgements. We thank all the reviewers for their helpful comments and thank Gang Zhao for sharing the SP2 code.

Financial support. This work was supported by the Guangdong Major Project of Basic and Applied Basic Research (grant no. 2020B0301030004), the Natural Science Foundation of Fujian Province (2021J01463), the National Natural Science Foundation of China (42175083 and 42105092), the Guangdong Provincial Key Research and Development Program (2020B1111360003), the Guangdong Basic and Applied Basic Research Foundation (2019A1515110791), and the Special Fund for Science and Technology Innovation Strategy of Guangdong Province (grant no. 2019B121205004).

Review statement. This paper was edited by Yuan Wang and reviewed by Zezhen Cheng and two anonymous referees.

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