The single-particle mixing state and cloud scavenging of black carbon at a high-altitude mountain site in southern China

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Highlights

- In situ investigation of both the mixing state and cloud scavenging of BC in China was first reported.
- Measured BC-containing particles were activated into cloud droplets to the same extent as all the measured particles due to being predominantly mixed with sulfate.
- BC-containing particles with higher fractions of organics were activated relatively less than those with higher fractions of sulfate.
Abstract

In the present study, a ground-based counterflow virtual impactor (GCVI) was used to sample cloud droplet residual (cloud RES) particles, while a parallel PM$_{2.5}$ inlet was used to sample cloud-free or cloud interstitial (cloud INT) particles. The mixing state of black carbon (BC)-containing particles in a size range of 0.1-1.6 µm and the mass concentrations of BC in the cloud-free, RES and INT particles were investigated using a single particle aerosol mass spectrometer (SPAMS) and two aethalometers, respectively, at a mountain site (1690 m a.s.l.) in southern China. The measured BC-containing particles were internally mixed extensively with sulfate, and were activated into cloud droplets to the same extent as all the measured particles. The results indicate the preferential activation of larger particles and/or that the production of secondary compositions shifts the BC-containing particles towards larger sizes. BC-containing particles with an abundance of both sulfate and organics were activated less than those with sulfate but limited organics, implying the importance of the mixing state on the incorporation of BC-containing particles into cloud droplets. The mass scavenging efficiency of BC with an average of 33% was similar for different cloud events independent of the air mass. This is the first time that both the mixing state and cloud scavenging of BC in China have been reported. Since limited information on BC-containing particles in the free troposphere is available, the results also provide an important reference for the representation of BC concentrations, properties, and climate impacts in modeling studies.
Keywords: black carbon, cloud droplet residues, mixing state, scavenging efficiency, interstitial particle
1 Introduction

Black carbon (BC), also known as soot or elemental carbon, is primarily emitted from incomplete combustion processes (Petzold et al., 2013; Bond et al., 2013). Fresh BC-containing particles are generally hydrophilic due to the presence of thin coatings of inorganic or organic materials (Zuberi et al., 2005), and during transport they become more hydrophilic when further coated through coagulation, condensation and photochemical oxidation (Zuberi et al., 2005; Zaveri et al., 2010; Matsui, 2016). Hydrophilic BC-containing particles can act as cloud condensation nuclei (CCN) and thus modify cloud microphysical properties (Straub et al., 2012; Schroder et al., 2015; Roth et al., 2016). The increase in CCN activity enhances the in-cloud scavenging of BC and thus reduces its lifetime (Zaveri et al., 2010). Aerosol-cloud interactions represent one of the largest uncertainties in our current understanding of human-induced climate forcing (McFiggans et al., 2006; Andreae and Rosenfeld, 2008). Therefore, a more comprehensive understanding of how aerosol particles form cloud droplets is required in order to reduce the uncertainty of the impacts of aerosols on the climate (Furutani et al., 2008).

The abilities of particles to act as CCN are largely controlled by their sizes and chemical compositions or mixing state (Dusek et al., 2006; Cubison et al., 2008; Kammermann et al., 2010; Ching et al., 2012; Baustian et al., 2012). Larger aerosol particles were found to be more easily activated into cloud droplets (Drewnick et al., 2007). Zhou et al. (2009) reported higher scavenging rates for sulfate, nitrate and BC than those for organics at Mount Tai in northern China. At the same site, 92% of the cloud residues were attributed to sulfate-related...
salts (Li et al., 2011b). On the other hand, the chemical compositions of the original CCN would be altered after the evaporation of the cloud droplets through the effective formation of secondary aerosol compositions during cloud processing (Hayden et al., 2008; Herrmann et al., 2015; Roth et al., 2016). For BC-containing particles, the mixing state should be much more significant, since their activation as CCN is primarily attributed to the presence of secondary coatings (Lambe et al., 2015; Schroder et al., 2015). Additionally, the mixing state of BC-containing particles is complex and constantly changing in the atmosphere, and they are highly influenced by the particle size, sources, the formation of secondary species and transport processes (Cahill et al., 2012; Healy et al., 2012; Zhang et al., 2014).

Recent in situ studies of cloud droplets have provided the most direct information on the incorporation of BC into clouds. The mass scavenging efficiency was observed to be in a range of 33–74% for BC, which was higher with increasing particle sizes at the Puy de Dome (1465 m a.s.l.), France (Sellegri et al., 2003), while it ranged from 13% to 50% within different air masses at a coastal Chilean hill (450 m a.s.l.) (Heintzenberg et al., 2016). Cozic et al. (2007) reported a scavenging rate of BC similar to those of bulk aerosols due to its internal mixing state with soluble materials. Wang et al. (2012) showed a higher scavenging efficiency for BC than those for organics. Roth et al. (2016) found an enhanced contribution of BC-containing particles in cloud residues compared to that in interstitial particles. However, Zelenyuk et al. (2010) observed negligible BC in cloud droplet residues above Alaska, USA. Therefore, an in-depth study on the composition, size and mixing state of BC in cloud droplets and interstitial particles is requisite for a better understanding of the
interactions between BC and cloud droplets and the influences of anthropogenic emissions on cloud formation in the free troposphere.

Single-particle mass spectrometry (SPMS) studies on fog interstitial particles and droplet residual particles were performed previously at an urban site in southern China (Zhang et al., 2012; Bi et al., 2016). The predominance of BC-containing particles that serve as effective fog condensation nuclei highlights the important influence of anthropogenic emissions on the public environment and regional climate (Bi et al., 2016). However, there are no direct observations of the cloud scavenging of BC or the mixing states of cloud interstitial (cloud INT) and droplet residual (cloud RES) BC-containing particles in the high-altitude atmosphere or the free troposphere above China to date. Therefore, the size-resolved mixing state and the scavenging efficiency of BC-containing particles were investigated at a high-altitude site to further our knowledge of (1) the mixing state of BC-containing particles, (2) the influence of the mixing state on the incorporation of BC into cloud droplets, and (3) the influence of anthropogenic activities on cloud formation in the free troposphere above southern China.

2 Methods

2.1 Sampling setup

The observations of cloud events were conducted at the National Atmospheric Background Monitoring Station in Nanling of Guangdong Province, which is located on the top of Mount Tianjing (24°41′56″N, 112°53′56″E, 1690 m a.s.l.) in southern China, from 16
to 26 Jan 2016. The site is located in a natural preserve isolated from anthropogenic activities. A map of the location and terrain of the site can be found elsewhere (Lin et al., 2017).

Aerosols were introduced into the instruments through two parallel sampling lines. The first inlet is a ground-based counterflow virtual impactor (GCVI) (Model 1205, Brechtel Mfg., Inc., USA) (Bi et al., 2016). The GCVI employs a compact wind tunnel upstream of the CVI inlet (Model 1204) (Shingler et al., 2012) to accelerate fog and cloud droplets into the CVI inlet tip. Similar methodologies have been extensively applied to collect fog/cloud RES particles (e.g., Sorooshian et al., 2013; Roth et al., 2016; van Pinxteren et al., 2016).

Upstream of the CVI sampling tip, only the droplets that exceed 8 μm can pass through the counterflow. The sampled cloud droplets enter the evaporation chamber (with an airflow temperature of 40 °C), where the droplets are dried, thereby leaving behind cloud RES particles that are capable of acting as CCN. A testing before measurements demonstrates that the influence of background aerosols on the collection of cloud droplets could be negligible (Zhang et al., 2017). The ambient inlet is a PM$_{2.5}$ sampling line that delivers ambient particles during cloud-free periods or cloud INT particles during cloud events. Cloud INT particles were regarded as PM$_{2.5}$ during the cloud events. More detailed description on the sampling can be found in the companion papers (Lin et al., 2017; Zhang et al., 2017).

Cloud events were characterized by a sudden drop in visibility and a sharp increase in the relative humidity (RH) measured by the GCVI. An upper-limit visibility threshold of 5 km and a lower-limit RH threshold of 95% were established to identify the cloud events and
trigger the sampling of the cloud RES particles. A rainfall detector within the GCVI system was used to exclude rainy periods. The enhancement factor (EF) for the particles collected by the GCVI is 5.25 (Shingler et al., 2012). Therefore, the reported mass concentrations for the cloud RES particles in the following text were first divided by 5.25.

An illustrative scheme of the instrumentation setup is provided in Fig. S1 in the Supporting Information (SI). Downstream of the GCVI, an aethalometer (Model AE-33, Magee Scientific, USA), a single particle aerosol mass spectrometer (SPAMS, Hexin Analytical Instrument Co., Ltd.) and a scan mobility particle sizer (SMPS, MSP Corporation, USA) were used to measure the concentration of BC, the size-resolved mixing state of the collected particles, and the number size distribution of submicron particles, respectively. Downstream of the ambient inlet, an SMPS (Grimm 5.041, Germany), an aethalometer (Model AE-31, Magee Scientific, USA), and a tapered element oscillating microbalance (Model 1405, TEOM) were used to determine the number size distribution of submicron particles and the mass concentrations of BC and PM$_{2.5}$, respectively. During the cloud-free periods, the instruments downstream of the GCVI were manually shifted and connected to the ambient PM$_{2.5}$ inlet. During the present study, three cloud events (Cloud I, II, III, each with a RH constantly above 95% for more than 12 hours) were encountered and identified by the GCVI (Lin et al., 2017), as shown in Fig. 1. During Cloud I and II, the cloud RES particles provided by the GCVI were measured by the instruments downstream of the GCVI. During Cloud III, the cloud RES and cloud INT particles were intermittently measured by these instruments at approximately one-hour intervals.
2.2 Determinations of the mass concentrations of BC

The AE–31 and AE–33 measured the BC concentration at seven different wavelengths (370, 450, 520, 590, 660, 880 and 950 nm), which are typically represented as equivalent BC (EBC) (Petzold et al., 2013). The measurements are based on the light beam attenuation (ATN) on a filter, which is proportional to the loading of the BC deposit. In this study, the ATN was converted to an EBC concentration at the wavelength of 880 nm. The EBC concentration reported in the present study was measured using the AE–33, which was described in great detail (Drinovec et al., 2015). The limitations and uncertainties of the AE-31 in measuring BC and the necessary corrections were well documented (Weingartner et al., 2003; Arnott et al., 2005; Backman et al., 2016). A brief description of this issue is provided in the Supplement.

2.3 Identification of BC-containing particles by the SPAMS

Both the vacuum aerodynamic diameter ($d_{va}$) and the chemical compositions of the individual particles were analyzed by the SPAMS, as briefly described in the supplement. A detailed description of the performance of the SPAMS can be found elsewhere (Li et al., 2011a). The mass spectra for approximately 75000 particles with $d_{va}$ values in the range of 0.1-1.6 µm were obtained by the SPAMS over the study. The diameter is represented herein as $d_{va}$ rather than the equivalent volume diameter ($d_{ve}$), the conversion for which can be found in the supplement (DeCarlo et al., 2004; Hu et al., 2012). An adaptive resonance
theory-based neural network algorithm (ART-2a) was applied to cluster the individual particles based on the presence and intensities of ion peaks (Song et al., 1999) with a vigilance factor of 0.7, a learning rate of 0.05, and 20 iterations. Three BC particle types were obtained: the mass spectra of particles with more carbon cluster ions ($C_n^{+/-}$, $n > 6$) and sulfate (BC-sul1), those with fewer carbon cluster ions ($C_n^{+/-}$, $n \leq 6$) and more intense sulfate (BC-sul2), and those with an abundance of both sulfate and organics (BC-OC-sul). The relative amount of OC to BC for the BC-OC-sul particles is significantly larger than the relative amount of OC in the BC-sul1 and BC-sul2 particles, as indicated in Fig. S2. Over all of the detected BC-containing particles, the BC-sul2 type is the most abundant (63%) particle type, followed by the BC-sul1 (21%) and BC-OC-sul (16%) types. More detailed information regarding the other particle types can be found elsewhere (Lin et al., 2017).

3 Results and Discussion

During the sampling periods, the temperature and RH generally varied between −9.9 - 11.4 °C and 6.7 - 100%, respectively. The sampling durations for the cloud-free, cloud RES and cloud INT (only detected in Cloud III) particles were approximately 109, 123, and 26 hours, respectively. The detected numbers of the cloud-free, cloud RES, and cloud INT particles by the SPAMS were 48835, 23616, and 1063, respectively. The average number fractions of BC-containing particles in the cloud-free, cloud RES, and cloud INT particles in the size range of $d_{vn} = 0.1$-1.6 µm were 44%, 49%, and 53%, respectively. The number
fractions of BC-containing particles that were incorporated within the cloud droplets ranged between those observed at an urban site (70%) in southern China (Bi et al., 2016) and those observed at a mountain site (~30%) in Germany (Roth et al., 2016). While some mineral dust might trigger heterogeneous ice nucleation at temperatures below –7 °C (Atkinson et al., 2013), this would not influence the discussion on the number fraction and chemistry of the cloud RES BC-containing particles. The sampling duration of the cloud RES particles with an average temperature of ~ –7 °C only lasted 2 hours, during which the detected cloud RES BC-containing particles only accounted for ~0.1% of all the detected ones.

Air masses from the southwestern continental and marine areas dominated throughout the sampling period, carrying relatively warmer and wetter air masses that benefited the formation of clouds based on the back-trajectory analysis (Lin et al., 2017). Cloud II was strongly influenced by a northeastern air mass in contrast to the southwestern air mass that dominated during Cloud I and III. As shown in Fig. 2, the air mass during Cloud II represents relatively polluted conditions. The mass concentration of EBC during Cloud II was approximate 200 ng m\(^{-3}\), which is four times that (~50 ng m\(^{-3}\)) observed during the other two events. Similarly, the number fraction of the BC-containing particles in the cloud RES particles during Cloud II (~60%) was higher than those during the other two cloud events (< 30%). These results provide direct evidence for the influence of anthropogenic emissions and atmospheric transport on the formation of clouds at the remote high-altitude site in southern China.
3.1 Mixing state of BC for cloud-free, residual, and interstitial particles

The dominant ion peaks for the cloud-free, cloud RES, and cloud INT BC-containing particles were those of carbon cluster ions ($C_n^{+/−}$, $n = 1, 2, 3, \ldots$), OC fragments (m/z $27[C_2H_3]^+$, $-26[CN]^−$, $37[C_3H]^+$, and $43[C_2H_3O]^+$), and secondary inorganic species, such as sulfate ($−97[HSO_4]^{−}$), nitrate ($−62[NO_3]^{−}$ and $−46[NO_2]^{−}$), and ammonium ($18[NH_4]^+$) (Zhang et al., 2014). The cloud-free BC-containing particles were internally mixed to a great extent with detectable sulfate (97% in number), nitrate (50%), oxidized organics (72%), and/or ammonium (79%), as presented in Fig. S3. The oxidized organics were generally represented as the peak at m/z $43[C_2H_3O]^+$ in SPMS studies (Qin et al., 2012). A similar mixing state of the BC-containing particles has been observed at both urban and mountain sites (Moffet and Prather, 2009; Li et al., 2011c; Cahill et al., 2012). The overwhelming association of BC with sulfate strongly indicates a substantial influence of anthropogenic emissions of sulfate precursors (e.g., SO$_2$) on the aging of BC (Huang and Yu, 2008; Khalizov et al., 2009; Guo et al., 2012; Peng et al., 2016), which directly enhances the incorporation of BC into clouds as discussed in section 3.2. Compared to the BC-containing particles at urban and suburban sites that are situated close to emission sources, the relative amounts of sulfate and ammonium substantially increased for those at the mountain site, as shown in Fig. S4. The relative peak area (RPA) of each m/z relative to the sum of the peak areas in a mass spectrum was applied herein to represent the relative amount of a species in a particle (e.g., Xing et al., 2011; Jeong et al., 2011; Healy et al., 2013). The enhancement of sulfate in the atmosphere above southern China is reasonable since sulfate accounts for the
largest portion of the compositions in this region and should be mainly associated with ammonium (Zhang et al., 2013). As expected, the temporal variations of the RPAs were significantly correlated ($p < 0.01$) between ammonium and sulfate (Fig. S5). These species were generally regarded as secondary components, and thus, such high fractions of the internal mixing state and the enhancement of ammonium and sulfate at the high-altitude site indicates a highly aged state of the BC-containing particles.

As shown in Fig. 3, the secondary components were enhanced in the cloud RES BC-containing particles relative to the cloud INT BC-containing particles. The enhancement was more obvious for sulfate rather than for ammonium, oxidized organics or nitrate. The enhancement of sulfate in cloud RES particles has been broadly observed (Zelenyuk et al., 2010; Kamphus et al., 2010; Hiranuma et al., 2011). A comparison of the size distributions of the cloud RES and cloud INT BC-containing particles (Fig. 4) further suggests that the in-cloud addition of secondary components shifted the BC-containing particles towards larger sizes, which is discussed in the following section. Overall, our observations suggest that the BC-containing particles were heavily coated at the high-altitude site before they were incorporated into the cloud droplets and that the in-cloud production of coating materials (e.g., ammonium sulfate) was present. An abundance of BC-coated materials was also observed at Mt. Soledad (Schroder et al., 2015). Unfortunately, their chemical compositions cannot be ruled out by a single particle soot photometer. Therefore, our analysis reflects the importance of the chemical mixing state on the cloud processing of BC.
3.2 Fractions of BC incorporated into cloud droplets

3.2.1 Size-resolved activation of BC-containing particles

The normalized number size distributions of the cloud-free, cloud RES, and cloud INT BC-containing particles are shown in Fig. 4. While these distributions do not represent the actual particle number size distributions, they could reflect the importance of the particle size on the incorporation of BC-containing particles into cloud droplets (Dusek et al., 2006).

The cloud RES BC-containing particles had the largest size mode, followed by the cloud-free BC-containing particles with the cloud INT BC-containing particles in the smallest size mode. These size distribution patterns are consistent with those of previous studies (Zelenyuk et al., 2010; Drewnick et al., 2007; Roth et al., 2016), which are indicative of the preferential activation of larger particles and/or the addition of secondary species during in-cloud processing (Schroder et al., 2015). As expected, the BC-containing particles were internally mixed with increasingly higher intensities of sulfate, ammonium and oxidized organics with increasing size (Fig. S6). These results are consistent with the observations by Healy et al. (2012) and Zhang et al. (2014) insomuch that larger BC-containing particles were more thickly coated. The BC-containing particles detected by the SPAMS could track the variations of the BC mass concentration in the present study based on a correlation analysis of the time series of the unscaled number of BC-containing particles and the concentration of EBC (Fig. S7). A detailed discussion on the comparison of
these two measurements can be found in the supplement (Yu et al., 2010; Huang et al., 2011; Huang et al., 2012).

The size-resolved activated fractions ($N_{f\text{act}}$) of the BC-containing particles and all the detected particles were further investigated as a function of their size (Fig. 5). The $N_{f\text{act}}$ generally increased with an increase in the size, and those of the BC-containing particles were activated to a similar (or slightly lower) extent as those of all the detected particles. This indicates that the coating materials on the BC-containing particles enhanced their ability to act as CCN (Khalizov et al., 2009; Henning et al., 2012; Roth et al., 2016), which is also consistent with the enhanced internal mixing with secondary soluble species with an increase in the size (Fig. S6) discussed above. The increase of $N_{f\text{act}}$ with the particle size also suggests that nucleation scavenging is the dominant mechanism for the incorporation of BC-containing particles into cloud droplets (Schroder et al., 2015). The number fractions of the BC-containing particles incorporated into cloud droplets varied between 0.05–0.45 (Fig. 5). These fractions represent a rough estimate because the BC-containing particles in the cloud RES and cloud INT particles were measured intermittently rather than simultaneously. Henning et al. (2002) showed that particles with $d_{ve} = 700$ nm were only half activated in less dense clouds (with a liquid water content or LWC < 0.1 g m$^{-3}$), while their diameter decreased to ~100 nm when the LWC > 0.2 g m$^{-3}$ at a high-alpine site. Similarly, Hammer et al. (2014) showed that only particles with a $d_{ve}$ larger than 300 - 500 nm could be activated under low-LWC conditions (LWC < 0.1 g m$^{-3}$), which is a common condition for the formation of fog at the ground level. With an LWC of approximately 0.1
17 g m\(^{-3}\), Schroder et al. (2015) reported even lower activated fractions (0.01–0.1) of BC-containing particles at Mt. Soledad closer to the source region in California, USA. From this perspective, the relatively higher activation fractions of the BC-containing particles in the present study compared to those at Mt. Soledad (Schroder et al., 2015) could be mainly attributed to the long-range transport that resulted in the highly aged BC and possibly the higher LWC.

The role of the mixing state on the activation of the BC-containing particles was further investigated through a comparison of the individual particle types of the cloud-free, cloud RES, and cloud INT BC-containing particles. As shown in Fig. 6, the number fraction of BC-OC-sul was much lower in the cloud RES (~8%) than those (~25%) in the cloud-free and cloud INT BC-containing particles. Despite the different distributions of the BC particle types, the BC-sul1 and BC-sul2 types were dominant, while the BC-OC-sul type contributed only a limited fraction to the cloud RES BC-containing particles during each of the cloud events. Consistently, the \(N_{\text{act}}\) of the BC-OC-sul particles was generally lower than 0.1 over the detected size range, which is much lower than those for the BC-sul1 and BC-sul2 types (Fig. S8). Furthermore, the organic-dominated particle types (organics internally mixed with sulfate and negligible BC (Lin et al., 2017)) in all the detected particles was activated to a lesser extent, as presented in Fig. S9. Distinct differences in the mixing state accompanied the observations of cloud RES BC-containing particles. BC-containing residuals with more sulfate and fewer organics were observed more frequently than those with more organics and less sulfate.
3.2.2 Mass scavenging efficiency of EBC

The concentration of EBC (5th - 95th) obtained using the AE33 for cloud-free air varied over a wide range of 57 - 812 ng m$^{-3}$ with a mean value of 418 ± 248 ng m$^{-3}$, which accounted for ~2% of the PM$_{2.5}$ on average. The average concentrations of cloud RES and INT EBC were 84 ± 75, and 198 ± 125 ng m$^{-3}$, respectively. A relatively lower contribution of EBC to the aerosol population supports a substantial addition of secondary aerosols during transport to the high-altitude site, given that EBC represents far more than ~2% of the fine particles near the source regions of southern China (Lan et al., 2013; Wu et al., 2013; Zhang et al., 2013). The observed relatively lower fraction of EBC is consistent with the highly aged state of BC-containing particles at the high-altitude site rather than at urban and suburban sites, as discussed in section 3.1. The mean concentration is much lower than those observed for urban (6000 ng m$^{-3}$) and rural (2600 ng m$^{-3}$) areas (Huang et al., 2012), it is similar to those observed at an oceanic site (540 ng m$^{-3}$) in southern China (Wu et al., 2013) and at the high-altitude Mt. Rax site (430-720 ng m$^{-3}$) (Hitzenberger et al., 2001), and it is several times higher than those at a marine boundary layer site (70 ng m$^{-3}$) in California, USA (Schroder et al., 2015), the mid-altitude regions (~60 ng m$^{-3}$) of Nova Scotia, Canada (Chýlek et al., 1996), and the high alpine Jungfraujoch station (50-60 ng m$^{-3}$) in Switzerland (Cozic et al., 2007). Additional detailed information on the sampling sites and BC concentrations can be found in Table S1. These comparisons suggest that...
Anthropogenic activities have a relatively large impact on the concentration of EBC at the high-altitude site. It can be seen from Fig. 1 that cloud scavenging could have a strong effect on the decreased particle concentrations (i.e., of EBC and PM$_{2.5}$). A sharp reduction in the particle concentrations were observed at the beginning of the cloud events. The mass-scavenging efficiency of BC ($M_{scav,EBC}$), defined as the fraction of EBC incorporated into cloud droplets relative to the total amount of EBC (Cozic et al., 2007), was evaluated as

$$M_{scav,EBC} = \frac{EBC_{RES}}{EBC_{RES} + EBC_{INT}} \times 100\%$$  \hspace{1cm} (R1)

Since the $EBC_{RES}$ and $EBC_{INT}$ were not simultaneously obtained using the AE-33, the $EBC_{INT}$ measured concurrently by the AE-31 was applied in the calculation. The EBC measured using the AE-31 is significantly correlated ($R^2 = 0.9$, $p < 0.001$) with and only slightly lower than that measured by the AE-33, as shown in Fig. S10. This validates the calculation in R1. The overall uncertainty in the $M_{scav,EBC}$ is within 10%, as assessed in the supplement. The measurements of EBC and the sampling of the cloud RES particles were regarded as the main influential factors.

The $M_{scav,EBC}$ ranged between 15 - 54% (5$^{th}$ - 95$^{th}$) with an average of approximately 33%. The $M_{scav,EBC}$ in this study is within the range of those values (33-54%) reported for mid-altitude (approximately 1500 m) mountain sites, generally lower than those reported (45-74%) for high-altitude (approximately 3000 m) mountain sites, and higher than those reported (6-15%) for ground sites (Cozic et al., 2007 and references therein). The differences among the various observations are generally attributed to the water content.
and the sizes and mixing state of the BC-containing particles (Cozic et al., 2007). The 
M$f_{scav,EBC}$ was not so different for the cloud events (Fig. S11) impacted by different air 
masses, which is consistent with the highly aged state of the BC observed in this study. 
These results indicate that the incorporation of BC into clouds was dominantly controlled by 
its mixing state rather than other factors (e.g., the air mass or the concentration of EBC) 
under low-LWC conditions (e.g., < 0.1 g m$^{-3}$).

4 Conclusions

The influences of the size and mixing state on the incorporation of BC in clouds were 
investigated at a remote high-altitude mountain site in southern China. On average, the mass 
concentration of EBC was 418 ± 248, 84 ± 75, and 198 ± 125 ng m$^{-3}$ for cloud-free, cloud 
RES, and cloud INT particles, respectively. The BC was highly aged through the 
predominant accumulation of sulfate during transport. BC-containing particles were found 
to be scavenged in the cloud phase to a similar extent as bulk aerosols. The size-resolved 
activation fraction of BC-containing particles was estimated to be in the range of 0.05–0.45; 
it increased with an increase in the size and was mainly controlled by the mixing state with 
secondary soluble species. Our results are restricted to particles in the size range of 0.1-1.6 
µm, and thus, particles with sizes smaller than 0.1 µm that might serve as CCN are beyond 
the scope of this study. The mass-scavenging efficiency of BC varied between 15–54% and 
was independent of the air mass. This paper provides the first direct evidence on the 
substantial contribution of BC-containing particles to cloud droplet residues in the free
troposphere of southern China. The data are also useful for constraining models used for predicting BC concentrations in the free troposphere.

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Figure captions

Fig. 1. Temporal profiles (with a 1 hour resolution) of PM$_{2.5}$, EBC mass concentrations, number of BC-containing particles by SPAMS, RH and visibility. Three cloud events are illustrated with black bars above the figure.

Fig. 2. Box and whisker plots of (a) concentration of EBC and (b) number fraction of BC-containing particles for each cloud event. In a box and whisker plot, the lower, median and upper lines of the box denote the 25th, 50th, and 75th percentiles, respectively, and the lower and upper edges of the whisker denote the 10th and 90th percentiles, respectively.

Fig. 3. (a) Average mass spectrum of cloud RES BC-containing particles, and (b) the RPA ratios of ammonium, sulfate, nitrate, oxidized organic markers, and other organic markers (i.e., m/z 27[C$_2$H$_3$]+, -26[CN], 37[C$_3$H]+, 50[C$_4$H$_2$]+, 51[C$_4$H$_3$]+, 61[C$_5$H]+, and 63[C$_5$H$_3$]+) to BC (carbon ion clusters (C$_n^{+/-}$, n ≤ 5)), and the RPAs of BC for cloud RES and INT particles, respectively. Error bars represent the standard deviation in the hourly average RPA or the RPA ratios within a 95% confidence interval.

Fig. 4. Normalized number size distributions of the cloud-free, INT, and RES BC-containing particles. The data were averaged throughout the sampling period. The size distribution obtained from the SPMS is not representative of the actual aerosol size distribution due to the decreasing detection efficiencies at smaller sizes (Allen et al., 2000; Wenzel et al., 2003; Qin et al., 2006). A representative comparison between the size distributions measured by the SPAMS and the SPMS can be found in Fig. S12.
Fig. 5. Size-resolved $N_{\text{f act}}$ estimated for the BC-containing particles and all the detected particles. The $N_{\text{f act}}$ is calculated as the ratio of the average number size distribution for the cloud RES particles to the sum of the average number size distributions for the cloud RES and INT particles. Errors were estimated assuming that the particle numbers detected by the SPAMS follow a Poisson distribution.

Fig. 6. (a) Number fraction of each BC particle type in the cloud-free, INT, and RES BC-containing particles, and (b) the number fraction of each BC particle type in the cloud RES BC-containing particles separated for the three cloud events.
Fig. 1.
Fig. 2.

(a)

cloud III
cloud II
cloud I

concentration of EBC (ng m⁻³)

(b)

cloud III
cloud II
cloud I

Nf of BC-containing particles
Fig. 3.
Fig. 4.
Fig. 5.
Fig. 6.