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Quantification of black carbon mixing state from traffic: implications for aerosol optical properties

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Abstract. The climatic impacts of black carbon (BC) aerosol, an important absorber of solar radiation in the atmosphere, remain poorly constrained and are intimately related to its particle-scale physical and chemical properties. Using particle-resolved modelling informed by quantitative measurements from a soot-particle aerosol mass spectrometer, we confirm that the mixing state (the distribution of co-emitted aerosol amongst fresh BC-containing particles) at the time of emission significantly affects BC-aerosol optical properties even after a day of atmospheric processing. Both single particle and ensemble aerosol mass spectrometry observations indicate that BC near the point of emission co-exists with hydrocarbon-like organic aerosol (HOA) in two distinct particle types: HOA-rich and BC-rich particles. The average mass fraction of black carbon in HOA-rich and BC-rich particle classes was < 0.1 and 0.8, respectively. Notably, approximately 90 % of BC mass resides in BC-rich particles. This new measurement capability provides quantitative insight into the physical and chemical nature of BC-containing particles and is used to drive a particle-resolved aerosol box model. Significant differences in calculated single scattering albedo (an increase of 0.1) arise from accurate treatment of initial particle mixing state as compared to the assumption of uniform aerosol composition at the point of BC injection into the atmosphere.

1 Introduction

Incomplete combustion emits teragram quantities of black carbon (BC) aerosol to the troposphere each year, resulting in a significant warming effect on climate that may be second only to carbon dioxide (Bond et al., 2013; Ramanathan and Carmichael, 2008; Jacobson, 2001). BC influences climate directly, by absorbing solar radiation, and indirectly, by changing cloud properties and altering snow and ice melt. BC impacts on the global scale remain poorly constrained and are intimately related to its particle-scale physical and chemical properties (Bond et al., 2013). The majority of BC emissions in North America, Europe, and Latin America are derived from traffic-related sources, though the specific physical and chemical properties of BC-containing particles at emission depend greatly on the source (Bond et al., 2013). BC-containing particles are generally hydrophobic near emission and become mixed over time with hydrophilic species through condensation and coagulation (Johnson et al., 2005; Moteki et al., 2007; Moffet and Prather, 2009), with resulting impacts on particle hygroscopicity (McMeeking et al., 2011b; D. Liu et al., 2013; Laborde et al., 2013) and optical properties (Zhang et al., 2008; Cappa et al., 2012; Luck et al., 2012; Knox et al., 2009; Liu et al., 2015). Despite extensive previous work, our understanding of the
role of mixing state in influencing the climate impacts of BC remains incomplete, in part because of instrumental challenges in particle characterization.

Studies assessing both the composition and amount of non-BC species in BC-containing particles are rare. A large body of evidence from urban, tunnel, and engine emission studies has shown that individual combustion particles are mixtures of BC, inorganic species, metals, and hydrocarbon-like organic species at the time of emission (e.g., Johnson et al., 2005; Toner et al., 2006; Schneider et al., 2006; Tritscher et al., 2011; Chirico et al., 2010; Zhang et al., 2013; Massoli et al., 2012; Dallmann et al., 2014). Among these studies, single particle mass spectrometry has directly demonstrated mixing of BC and non-BC species in traffic-derived particles. For example, Toner et al. (2006) observed that emissions from heavy duty diesel engines were dominated by particles containing BC, organic species, calcium, and phosphate, with one particle type dominated by BC and another with higher levels of organic species. In contrast, Healy et al. (2012) observed BC-dominated particles from traffic emissions in a European city. Other approaches to measure BC mixing state, including hygroscopicity and volatility differential mobility techniques, have highlighted the presence of an external mixture in terms of particle volatility with lower volatility, BC-containing aerosol being less hygroscopic (Kuwata et al., 2009; McMeeking et al., 2011a). Further, single particle soot photometer measurements of coating thickness in urban areas have shown that traffic-related BC-containing particles are largely uncoated or very thinly coated (Shiraiwa et al., 2008; McMeeking et al., 2011b; Laborde et al., 2013; Liu et al., 2014). In addition, microscopy studies have illustrated the dominance of bare or thinly coated BC-containing particles in traffic emissions (China et al., 2014) and the occurrence of coating, embedding and compaction of BC as it is aged (Adachi and Buseck, 2013; Adachi et al., 2014). While previous work has provided valuable insight into BC mixing state, the majority of past approaches have not allowed simultaneous quantification of BC mixing state on a mass basis and chemical characterization of non-BC species.

Considerable attention has been paid to BC-containing aerosol because of the potential for short-term climate mitigation through emission reduction (Shindell et al., 2012). Therefore, a quantitative understanding of BC mixing state is crucial for three reasons: first, to assess rates of BC processing and removal in the atmosphere; second, to assess the role that BC-containing particles play as cloud nuclei; and third, to assess direct effects on solar radiation. Aerosol optical properties are central parameters required to evaluate direct radiative forcing (DRF). One of the largest contributors to uncertainty in DRF calculations is the single scattering albedo (SSA) (McComiskey et al., 2008), defined as the ratio of aerosol scattering to total light extinction. Previous work has clearly demonstrated that calculations of aerosol optical properties depend upon assumptions about particle mixing state (e.g., Jacobson, 2000), with assumptions of uniform internal mixing producing overestimates of absorption efficiency and underestimates of single scattering albedo (Oshima et al., 2009; Zaveri et al., 2010; Matsui et al., 2013). The role of BC-aerosol mixing state at emission in affecting its mixing state in the atmosphere has also been highlighted in modeling studies explicitly treating detailed aerosol microphysical and chemical processes (e.g., Matsui et al., 2013). However, model assessments driven by quantitative measurements of aerosol mixing state remain rare.

In this work, we use a soot-particle aerosol mass spectrometer, equipped with a light-scattering module, to determine the mixing state of BC-containing particles from traffic-dominated sources in an urban environment. These measurements provide quantitative insight into the physical and chemical nature of BC-containing particles near emission, and they are used to drive a particle-resolved aerosol box model to assess the effect of accurately representing BC mixing on aerosol optical properties.

2 Methods

2.1 The soot-particle aerosol mass spectrometer (SP-AMS)

2.1.1 Two urban studies: locations and SP-AMS configuration

A soot-particle aerosol mass spectrometer (Aerodyne Research Inc., Billerica, MA, USA), equipped with a light scattering module, was deployed in two urban studies to assess the mixing state of refractory black carbon (rBC) containing particles derived from vehicle emissions (Lee et al., 2015a). The first study was conducted in downtown Toronto away from major roadways (referred to as the “non-roadside” study), and the second was performed at ground level, near a busy road in downtown Toronto (referred to as the “roadside” study), to investigate fresh vehicle emissions. The non-roadside site is described in detail in Lee et al. (2015a). The roadside study took place from 31 May to 24 June 2013 at the Southern Ontario Centre for Atmospheric Aerosol Research (SOCAAR) facility in downtown Toronto, Canada, located at ground level and adjacent to a road with traffic volumes ranging from 16 000 to 25 000 vehicles per day (Sabaliuskas et al., 2014). Ambient air was sampled at 170 L min$^{-1}$ through a 10 cm (inner diameter) stainless steel tube fitted with a 2.5 μm cut-off inlet, located 15 m from the roadside at a height of 3 m a.g.l.

Two main modes of operation are possible with the SP-AMS, depending on the vaporizer configuration. The instrument was operated such that both rBC-containing particles and non-rBC-containing particles were detected in the non-roadside study (i.e., a dual-vaporizer configuration), whereas exclusively rBC-containing particles were detected in the
roadside study (i.e., a laser-only configuration) (Lee et al., 2015a; Onasch et al., 2012).

2.1.2 SP-AMS operation and calibration

Details of the SP-AMS (and SP-AMS with light scattering) have been described elsewhere (Lee et al., 2015a; Onasch et al., 2012). The SP-AMS detects black carbon, which evaporates at \( \sim 4000 \) K, as \( C_2^+ \) fragments (predominantly \( m/z \) 12, 24, 36, 48, and 60) and is referred to as refractory black carbon (rBC) (Onasch et al., 2012). We use the term “black carbon” when referring generally to the concept of “soot”-type aerosol, while rBC is used when referring to quantities measured by the SP-AMS. In the SP-AMS, rBC and associated species are volatilized in an infrared laser beam (1064 nm) and are ionized using 70 eV electron impact ionization followed by detection in a time-of-flight mass spectrometer (ToF-MS) operated in “V mode.” For the majority of the study the SP-AMS was operated at 1 min time resolution alternating between bulk mass spectrum (MS), particle time-of-flight (pToF), and single particle modes. The SP-AMS was operated at high time resolution (1 Hz) in MS mode for a total of 5 days during the roadside study. SP-AMS data were analyzed using the Igor Pro based analysis tool PIKA (Seupher, 2010). The single particle categorization procedure and \( k \)-means clustering algorithm that were used to analyze the single particle data followed the description in Lee et al. (2015a). Positive matrix factorization (PMF) analysis of ensemble data was performed to identify the sources of rBC and organics and to speciate different types of organic species (i.e., oxygenated organic aerosol (OOA), biomass burning organic aerosol (BBOA), and hydrocarbon-like organic aerosol (HOA)), based on procedures described previously (Paatero and Tapper, 1994; Zhang et al., 2011; Ulbrich et al., 2009).

Without the tungsten vaporizer, direct calibration of the ionization efficiency for \( \text{NH}_4\text{NO}_3 \) (IE\(_{\text{NO}_3} \)) is not possible. Therefore, size-selected (300 nm) Regal black (Regal 400R pigment, Cabot Corp.) particles were used to determine the mass-based ionization efficiency of rBC (mIE\(_{\text{rBC}} \)) (Onasch et al., 2012; Willis et al., 2014). The relative ionization efficiency for rBC (RIE\(_{\text{rBC}} = \text{mIE}_{\text{rBC}}/\text{mIE}_{\text{NO}_3} \)) was 0.2 ± 0.05, as experimentally determined before removal of the tungsten vaporizer. Assuming that RIE\(_{\text{rBC}} \) is constant, IE\(_{\text{NO}_3} \) could be calculated based on known values of mIE\(_{\text{rBC}} \) and RIE\(_{\text{rBC}} \). The average mIE\(_{\text{rBC}} \) was 189 ± 20 ions pg\(^{-1} \) in the roadside study. The calculated IE\(_{\text{NO}_3} \) was then used with recommended relative ionization efficiencies to quantify other aerosol species associated with rBC (Jimenez et al., 2003). Collection efficiency for rBC particles was determined in the roadside study using beam width probe (BWP) measurements described in Willis et al. (2014). Ambient rBC-containing particles had an average beam width \( \sigma = 0.46 ± 0.03 \) mm, which is close to, but wider than, that of 300 nm Regal black particles (\( \sigma = 0.40 ± 0.08 \)) (Willis et al., 2014). Therefore, a collection efficiency (CE) of 0.6 was applied for absolute quantification of rBC and associated species. A time-varying collection efficiency was not possible with BWP measurements available here; the assumption of a constant CE over periods of local and long-range transport influence in this study provided a good linear correlation with photoacoustic soot spectrometer (PASS) absorption measurements (405 and 781 nm) and SP-AMS rBC (Healy et al., 2015, Fig. 4). Note that the CE applied will not impact calculations of the mass fraction of rBC (mf\(_{\text{rBC}} \)). However, two additional uncertainties in SP-AMS measurements may affect calculation of mf\(_{\text{rBC}} \). First, there are uncertainties in the recommended RIE for organic species evaporating from rBC in the SP-AMS of up to \( \sim 50 \% \) (Lee et al., 2015a; Willis et al., 2014), which could cause an overestimation in the mass of coating material and a corresponding underestimation in mf\(_{\text{rBC}} \). Second, it is possible for rBC-containing particles to pass through the edges of the laser vaporizer, thus producing a heating effect sufficient to evaporate some fraction of the coating materials but not evaporate the rBC itself. This effect may also lead to an underestimation in mf\(_{\text{rBC}} \). SP-AMS CE and quantification are discussed in further detail in Supplement Sect. S1.

2.2 Photoacoustic soot spectrometer

A photoacoustic soot spectrometer (PASS-3, Droplet Measurement Techniques, Boulder, CO) was used to measure aerosol absorption \( (b_{\text{abs}}) \) and scattering \( (b_{\text{abs}}) \) coefficients \( (\text{Mm}^{-1}) \) at 405 and 781 nm. A 532 nm laser is not installed in this particular unit. The PASS determines aerosol absorption \( (\text{Mm}^{-1}) \) in a cavity which acts as an acoustic resonator. The absorption of incoming radiation heats the particles, which in turn heat the surrounding air in the cavity (Arnott et al., 1999). The aerosol-laden air thus expands, resulting in a pressure disturbance. By modulating the laser power at the resonance frequency of the cavity, the pressure disturbance is amplified and the resulting acoustic wave is measured using a microphone. Light scattering at both wavelengths is concurrently measured using reciprocal nephelometry (Moosmüller et al., 2009; Flowers et al., 2010; Chan et al., 2011). Scattering signals were not corrected for truncation; however, since mass-based particle size distributions at the roadside site generally peaked near or below 100 nm a negative bias in the scattering measurement is unlikely. Since a 532 nm laser was not present in this unit an NO\(_2 \) calibration was not possible, and the instrument was calibrated using a propane soot generator (miniCAST, 6203A, Jing). PASS measurements of the bulk single scattering albedo at 405 nm (selected due to superior signal-to-noise ratio for scattering relative to the 781 nm channel) are used here only to illustrate differences in optical properties in vehicle plumes with varying composition.
2.3 Particle-resolved box model simulations

We used the stochastic particle-resolved box model PartMC-MOSAIC (Particle Monte Carlo – Model for Simulating Aerosol Interactions and Chemistry) (Riemer et al., 2009; Zaveri et al., 2008) to quantify the importance of the observed mixing state information for the calculation of cloud condensation nuclei (CCN) and aerosol optical properties. PartMC-MOSAIC is suited for this task, as it explicitly tracks the composition of individual aerosol particles (in our case about $10^5$ particles) in a population of different particle types within a well-mixed computational volume. Particle emissions, dilution with the background, and Brownian coagulation are simulated stochastically with PartMC, by generating a realization of a Poisson process using weighted particles in the sense of DeVille et al. (2011) and an accelerated binned sampling strategy (Michelotti et al., 2013). Coupled to PartMC is the MOSAIC chemistry code, which simulates gas chemistry (Zaveri and Peters, 1999), particle-phase thermodynamics (Zaveri et al., 2005a, b), and dynamic gas-particle mass transfer (Zaveri et al., 2008) in a deterministic manner.

Here we compare two simulations that differ in the way the mixing state of the aerosol initial conditions and the aerosol emissions are prescribed. The setup of the particle-resolved simulations is similar to the urban plume scenarios in Riemer et al. (2009) and Zaveri et al. (2010). Common to both simulations are the following specifics. The number of computational particles used in the simulation is about $10^5$. The total simulation time is 24 h, with a simulation start of 06:00 local standard time. The initial gas-phase concentrations are the same as in Zaveri et al. (2010). In contrast to Zaveri et al. (2010), we prescribe a constant mixing height (400 m), relative humidity (70%), and temperature (298.15 K) to simplify the interpretation of the results. Since the entrainment of background aerosol can modify the aerosol mixing state substantially (Liu et al., 2011), we include constant dilution with the background at a rate of $1.5 \times 10^{-5}$ s$^{-1}$. The background aerosol is non-absorbing and consists of ammonium sulfate mixed with biogenic secondary organic aerosol. This dilution rate corresponds to about 75% of the aerosol being replaced in a 24 h period, comparable to a diurnal mixing height increase of 500 to 2000 m (e.g., Liu et al., 2011), although imposed uniformly over the day for simplicity. We also reduced the gas-phase emissions by a factor of 2 compared to Zaveri et al. (2010) to create more moderately polluted conditions. The initial aerosol population was sampled from log-normal size distributions fitted from the measurement data, as shown in Table S1 in the Supplement. The derivation of model inputs from measurement data is described in Supplement Sect. S3.

To fully exploit the information supplied by the observations, we introduced a new method of sampling the aerosol composition for the aerosol initial conditions and emissions. We now allow for stochastic composition variation at each di-
Unit mass resolution (UMR) spectra of HOA-rich (a) and rBC-rich (b) particle classes identified by k-means cluster analysis of single particle SP-AMS data acquired in the non-roadside environment. UMR spectra of HOA-rich (c) and rBC-rich (d) particle classes identified from single particle data in the roadside environment. High-resolution mass spectra of HOA-rich (e) and rBC-rich (f) PMF factors from ensemble SP-AMS data from the roadside study (i.e., for rBC-containing particles only). Examples of UMR spectra collected at high time resolution during HOA-rich (g) and rBC-rich (h) plumes (corresponding time series are shown in Fig. 6) during the roadside study. Insets in the lower panels illustrate the UMR spectrum of organic species (without CO\textsubscript{x} fragments) in the rBC-rich mass spectra. Black and colored sticks represent the fragments of rBC (C\textsubscript{x}H\textsubscript{y}O\textsubscript{w}) and organic species (C\textsubscript{x}H\textsubscript{y}+), respectively.

The two simulations differ in the assigned mixing state of the initial aerosol populations and the aerosol emissions. For the measurement-constrained case we distinguish between rBC-rich and HOA-rich particle classes, as the observations indicate. Both classes consist of two modes, which amounts to four modes in total, as listed in Table S1 in the Supplement. Note that the modes BC1 and HOA1 have the same geometric mean diameter and geometric standard deviation, but they differ in their abundance and their emission rate, respectively. The same applies to modes BC2 and HOA2. This specific choice is guided by the observations as shown in Table S1.

For the uniform case only two modes are prescribed, with the same geometric mean diameter and standard deviation as modes BC1/HOA1 and BC2/HOA2, respectively (Table S2). The mass fractions of BC and HOA in these two modes are identical, and they are equal to the mass fractions of the bulk concentrations for the measurement constrained case. Hence the uniform case represents conditions for which the bulk mass concentrations and the size distribution are the same as for the measurement-constrained case, but for which the detailed mixing state information is not available. A comparison of the two cases therefore quantifies the importance of mixing state information at emission for properties of interest, such as optical properties and CCN activation properties.

Figure S10 shows the temporal evolution of the bulk aerosol species over the course of the 24 h simulation. The primary species black carbon and organic carbon increase initially owing to emissions and then decrease after emissions are discontinued at \( t = 12 \) h as a result of dilution with the background. The model simulations also show the production of inorganic and organic secondary aerosol species, which condense on the primary particles, continuously modifying the composition of each particle in the aerosol population. Not shown in this figure is the aerosol water content. While the particles start out dry, they take up water around \( t = 6 \) h when nitrate formation begins.

We calculate the optical properties for ambient conditions (including aerosol water content) at a wavelength of 550 nm and the critical supersaturations of each particle in a post-processing step according to Zaveri et al. (2010). For the optical properties we assume spherical particles with a “core-and-shell” configuration in which BC forms the particle core and the other substances compose the shell. We then use Mie calculations (Ackerman and Toon, 1981) to determine the extinction and scattering cross sections for each particle in the population. From these values, the volume extinction, scattering, and absorption coefficients can be reconstructed. Note that we assume all primary and secondary organic substances to be non-absorbing; hence we do not consider any effects due to brown carbon.

To determine the critical supersaturations, we employ \( \kappa \) Köhler theory (see Zaveri et al., 2010 for details). Based on the critical supersaturation values of each particle in the population, we construct CCN spectra, as shown in Fig. S11 for different times throughout the simulation. As the simulation
progresses, the CCN/CN fraction at low values of critical supersaturation increases, since the particles become more hygroscopic due to the condensation of secondary aerosol material. Since both primary organic carbon (κ = 0.001) and black carbon (κ = 0) are model species with similar and low hygroscopicity parameters, the differences between the uniform case and the measurement-constrained case are expected to be small, and the two CCN spectra are almost indistinguishable, as we will show in Sect. 3.3.

3 Results and discussion

3.1 Two classes of fresh black carbon particles

Based on single particle SP-AMS measurements in both roadside and non-roadside studies, we identified two types of particles originating from vehicle exhaust, which are primarily composed of rBC and hydrocarbon-like organic aerosol (Fig. 1a–d). One type is dominated by HOA mass (referred to as the “HOA-rich” particle class) and the other is dominated by rBC (“rBC-rich” particle class). The mass spectra of HOA-rich particles (Fig. 1a and c) exhibit fragmentation patterns associated with hydrocarbon structures (e.g., m/z 43 (C3H7+), 57 (C4H9+), 71 (C5H11+)) and are consistent with previous aerosol mass spectrometer observations of gasoline/diesel vehicle exhaust and unburned lubricating oil (Massoli et al., 2012; Canagaratna et al., 2004; Mohr et al., 2009). In conjunction with differences in mass spectra, clear differences in particle size distributions (Fig. 2c and d) demonstrate that the division of traffic-related rBC-containing particles into HOA-rich and rBC-rich classes is physically meaningful.

The average mass fractions of rBC (mf_{rBC}) in HOA-rich particles during the non-roadside and roadside studies are low: 0.03 and 0.05, respectively. The narrow distribution of mf_{rBC} (Fig. 3a) suggests that most of the HOA-rich particles are either mixed with a small amount of rBC or do not contain detectable rBC mass, while a very small number of particles contain larger mf_{rBC} (Fig. 3, split axis). Mass spectra of rBC-rich particles are dominated by C\textsuperscript{n+} fragments (i.e., m/z 12, 24, 36, 48, and 60 in Fig. 1b and d) arising from black carbon. Smaller signals associated with HOA-like material in these particles are similar to the primary organic materials derived from diesel engine exhaust observed in laboratory studies (Sage et al., 2008). In contrast to the HOA-rich particles, Fig. 3c illustrates a wider range of mf_{rBC} values in rBC-rich particles. In the non-roadside and roadside environments the average mf_{rBC} values (±1 standard deviation) in this particle class are 0.72 (±0.18) and 0.86 (±0.14), respectively, highlighting the possibility that condensation and/or coagulation of HOA material on rBC particles might occur within a short time of emission.

The mass fraction of rBC derived from SP-AMS measurements may be best regarded as a lower limit. For the reasons detailed in Sect. 2.1.2, including the potential for incomplete vaporization and uncertainty in SP-AMS sensitivity to rBC coating materials, mf_{rBC} values presented here may be un-
derestimated. A 50 % overestimation in the mass of HOA would increase \( m_{f_{BC}} \) in rBC-rich particles to 0.85 (±0.16) and 0.92 (±0.10) in the non-roadside and roadside studies, respectively (Fig. 3c, dashed lines). In addition to these uncertainties, in the roadside study only rBC-containing particles are detected by the SP-AMS, while in the non-roadside study it is possible that the HOA-rich class includes HOA particles that do not contain rBC (i.e., “externally mixed” HOA).

### 3.1.1 Differences in particle coating thickness

With single particle detection capability, SP-AMS observations can be used to quantify the thickness of non-refractory particulate matter on individual rBC-containing particles. This calculation focuses only on determining the thickness of HOA coating derived from traffic emissions; however, OOA, BBOA, and secondary inorganic species were mixed with rBC in more aged accumulation mode particles in these urban studies and a similar calculation could be carried out for these particle classes (Lee et al., 2015a, b).

We assume a core-shell structure in order to determine the thickness of HOA coating on fresh rBC-containing particles. In addition, we assume a spherical rBC core, a uniform thickness of coating, and an HOA density of 0.9 g cm\(^{-3}\). To simulate the effect of the fractal structure of ambient rBC, the effective density of rBC (\( \rho_{\text{eff,rBC}} \)) was varied between 0.3 and 1.3 g cm\(^{-3}\), values that have been observed for laboratory soot standards and engine exhaust (Maricq and Ning, 2004; Gysel et al., 2011). Figure 4 presents the two-dimensional histograms (\( m_{f_{BC}} \) vs. particle aerodynamic diameter, \( d_{\text{va}} \)) of rBC-rich and HOA-rich particles identified by cluster analysis in the two urban studies, with calculated coating thickness curves overlaid for comparison (dashed lines). The coating thickness is relatively insensitive to variations in \( \rho_{\text{eff,rBC}} \) (Fig. 4, dashed lines).

These observations demonstrate that rBC-rich particles are only very thinly coated with HOA material, as compared to HOA-rich particles that have only very small rBC inclusions (Fig. 4). Here we use two specific examples to illustrate that the coating thickness on HOA-rich and rBC-rich particles can vary significantly. First, a 200 nm (\( d_{\text{va}} \)) rBC-rich particle (i.e., physical particle diameter ~ 245 nm, assuming \( \rho_{\text{eff,rBC}} = 0.8 \text{ g cm}^{-3} \)) with 72–86 % rBC by mass (the average \( m_{f_{BC}} \) of rBC-rich particles) would be covered by a 5–11 nm HOA coating. Second, a 300 nm (\( d_{\text{va}} \)) HOA-rich
Figure 4. Two-dimensional histograms (mf_{rBC} vs. particle aerodynamic diameter) of rBC-rich (a) and HOA-rich (b) particle classes. The colour scale represents the number of particles in each particle class. The dashed lines represent the physical thickness (nm) of organic coating on rBC-containing particles determined by modelling the particles as a core-shell structure. Note that the d_{va} of model outputs are calculated by the physical diameter (d_p) and the density (\rho_p) of particles (i.e., d_{va} = d_p \cdot \rho_p, where \rho_p is the linear combination of mass-weighted HOA and rBC density). To examine the potential effect of fractal structure of ambient rBC particles on predicted coating thickness, the effective density of rBC was varied (green: 0.3 g cm\(^{-3}\); white: 0.8 g cm\(^{-3}\); red: 1.3 g cm\(^{-3}\)).

Figure 5. Time series of (a) ensemble organic aerosol and (b) refractory black carbon mass loadings during the roadside study. Positive matrix factorization (PMF) results for ensemble data (c–f): (c) regionally sourced rBC mixed with oxygenated organic aerosol, (d) biomass-burning organic aerosol mixed with rBC, and traffic-related rBC in an rBC-rich and HOA-rich factor (e) and (f). Mass spectra of all PMF factors are shown in Fig. S1.

3.2 Black carbon mass is dominated by BC-rich particles

Single particle measurements provide direct insight into mixing state at an individual particle level; however, single particle detection has an inherent bias towards large particle sizes because light scattering triggers particle detection (Lee et al., 2015a; Cross et al., 2009; Freutel et al., 2013; S. Liu et al., 2013). Ensemble size distributions of rBC peaked at \sim 100 nm in both roadside and non-roadside studies indicating that fresh rBC-containing particles emitted from vehicle exhaust are small (Fig. 2a and b). Smaller rBC-containing particles fall below the lower size limit of single particle detection (Fig. 2c and d). Therefore, single particle observations do not necessarily quantify mixing state at the population level. To provide a complementary view of rBC mixing state we turn to ensemble measurements from the roadside study where only rBC-containing particles were detected in the SP-AMS. Ensemble measurements better represent mixing state on a population basis compared to single particle observations, because they cover a wider range of particle size (i.e., 80–1000 nm vacuum aerodynamic diameter, d_{va}).

By measuring very close to a busy road, ensemble SP-AMS observations of rBC and organic aerosol enabled the analysis of more than 100 vehicle exhaust plumes over the course of the roadside study (Fig. 5a and b). PMF (Zhang et al., 2011; Ulbrich et al., 2009) indicates three major sources of rBC-containing aerosol in the roadside environment: transported BBOA, regional background (OOA), and traffic emissions comprised of two PMF factors (HOA-rich and rBC-rich factors; Fig. 5c–f). Inorganic species evident in the bulk aerosol size distributions (Fig. 2a and b) are largely associated with OOA and BBOA factors, while the traffic-related factors contain the majority of rBC and HOA (Lee et al., 2015a, b). Mass spectra of all four PMF factors and a discussion of selection of the number of factors are presented in Supplement Sect. 2. Not previously observed with an AMS, the HOA-rich and rBC-rich factors represent two types of vehicle exhaust plumes with different amounts of rBC and HOA (Fig. 1e and f). The mf_{rBC} of HOA-rich and rBC-rich factors are 0.16 and 0.75, respectively. High-time-resolution SP-AMS measurements (Fig. 6) also demonstrated varying organic and rBC levels in vehicle plumes (Fig. 1g and h),
Mass spectra shown in Fig. 1g and h correspond to periods used for the subtraction of gas-phase contributions from soot spectrometer (PASS-3). Gaps in SP-AMS data correspond to single scattering albedo (SSA) at 405 nm from a photoacoustic high-time-resolution (1 Hz) sampling, with measurements of air signals in unit mass resolution spectra (Lee et al., 2015a).

Examples of rBC-dominated (a) and organic-dominated (b) plumes observed using the SP-AMS with high-time-resolution (1 Hz) sampling, with measurements of single scattering albedo (SSA) at 405 nm from a photoacoustic soot spectrometer (PASS-3). Gaps in SP-AMS data correspond to periods used for the subtraction of gas-phase contributions from particle signals. Mass spectra shown in Fig. 1g and h correspond to plumes in (b) and (a), respectively.

Figure 6. Examples of rBC-dominated (a) and organic-dominated (b) plumes observed using the SP-AMS with high-time-resolution (1 Hz) sampling, with measurements of single scattering albedo (SSA) at 405 nm from a photoacoustic soot spectrometer (PASS-3). Gaps in SP-AMS data correspond to periods used for the subtraction of gas-phase contributions from particle signals. Mass spectra shown in Fig. 1g and h correspond to plumes in (b) and (a), respectively.

with corresponding differences in optical properties (SSA at 405 nm) indicating that rBC-rich plumes were more highly absorbing. The origin of different plume types at this site has been characterized through long-term measurements and is related to a variety of engine types, operating conditions, and pollution control features (Wang et al., 2015).

Mass spectra of HOA-rich and rBC-rich PMF factors (Fig. 1e and f) are strikingly similar to the mass spectra of HOA-rich and rBC-rich particle classes identified in the single particle measurements (Fig. 1c and d), validating the division of traffic emissions into the two factors observed here. The single particle and ensemble mass spectra show some differences. First, the HOA-rich factor has a larger rBC content compared to the HOA-rich particle class; second, the rBC-rich factor contains higher CO\(^+\) and CO\(_2\)\(^+\) signals compared to the rBC-rich particle class (inserts of Figs. 1d, e, and f). The latter difference arises because highly oxygenated organic fragments (m/z 28 (CO\(^+\)) and 44 (CO\(_2\)\(^+\))) were excluded in the single particle analysis due to interference of air signals in unit mass resolution spectra (Lee et al., 2015a).

Differences in mf\(_{rBC}\) between single particle and ensemble measurements, especially for HOA-rich particles that contain a small amount of rBC, could be due to the probability of rBC detection in a single particle such that single particle mf\(_{rBC}\) may be underestimated relative to ensemble mf\(_{rBC}\) Lee et al. (2015a).

With support of direct measurements of single particle mixing state, this work presents the first interpretation of ensemble AMS results in terms of rBC and HOA mixing state. Using the intensity of the two traffic-related factors in plumes (Fig. 5e and f), we estimate that rBC-rich particles account for \(~90\%\) of the observed total traffic-related rBC mass (Fig. 3d). In a similar manner, we estimate that \(~60\%\) of the total HOA mass is due to HOA-rich particles (Fig. 3b). Purple dashed lines shown in Fig. 3b and d represent the inclusion of all CO\(^+\) fragments (due to surface functionality of ambient rBC, Corbin et al., 2014) in the calculation of rBC mass (see Supplement Sect. S1), providing similar results. Dashed lines in Fig. 3a, b, and c represent the effect of a 50 % overestimation of HOA mass (see Sect. S1), again providing similar results.

3.3 Black carbon mixing state at emission impacts

modelled optical properties

These novel single particle and ensemble observations of rBC mixing state were used to initialize the particle-resolved aerosol box model PartMC-MOSAIC (see Sect. 2.3) (Zaveri et al., 2008; Riemer et al., 2009), which simulates the evolution of aerosol due to condensation and coagulation in an idealized urban setting. As described in Sect. 2.3, we simulate two cases to isolate the impact of mixing state of rBC emissions on aerosol properties: first, a uniform mixing state case in which all particles are assigned identical composition at emission (measured average mf\(_{rBC}\)); second, a measurement-constrained mixing state case where mf\(_{rBC}\) is prescribed directly from our mixing state observations. The evolution of dry mf\(_{rBC}\) over a 24 h period is illustrated in Fig. 7a and b for the two cases. As ageing proceeds mf\(_{rBC}\) shifts to lower values, but in the uniform initial composition case no particles ever have mf\(_{rBC}\) larger than \(~45\%\). Note that the bulk aerosol composition and number size distributions are identical in the two cases (Fig. S10), and any differences in optical properties arise only due to the distribution of aerosol components amongst the particles.

Optical properties are determined for each particle in the population using Mie calculations and assuming a core-shell structure. We acknowledge that the underlying assumptions of Mie calculations may not be appropriate for rBC-containing aerosol in the real atmosphere (e.g., Adachi et al., 2011; Scarnato et al., 2013), which is often not spherical and may not exhibit a core-shell configuration (e.g., China et al., 2015). In particular, the enhancement of rBC absorption due to non-absorbing coatings may be overestimated (Cappa et al., 2012; Healy et al., 2015), except in situations
where rBC sources and ageing promote extensive coating (Liu et al., 2015; China et al., 2015). However, Mie calculations are commonly applied in regional and global models (Chung and Seinfeld, 2005; Zhao et al., 2013; Fast et al., 2006) to estimate optical properties, and we therefore include this comparison to illustrate the sensitivity to mixing state. 

Consistent with previous studies (e.g., Zaveri et al., 2010; Matsui et al., 2013), we observe a difference in volume absorption and scattering coefficients ($B_{abs}$ and $B_{scat}$, Fig. 7c) and a resulting increase of 0.1 in SSA from the uniform mixing state to the measurement-constrained case that is present during the emission period in the simulation and, notably, persists throughout ageing (Fig. 7d). Note that a direct comparison between measured and modelled SSA is not valid since we measure SSA of the bulk aerosol population and model it only for a subset of this population (i.e., the rBC-containing particles). No significant differences in calculated cloud condensation nuclei activity were observed in these simulations (Fig. S11), owing to the very similar hygroscopicity of rBC and HOA species. Calculations of aerosol DRF are very sensitive to changes in SSA (McComiskey et al., 2008) (e.g., uncertainties in SSA on the order of only 0.02 can result in a DRF uncertainty of ∼0.02 for a particular particle type; Chin et al., 2009), illustrating the importance of accurately measuring and simulating mixing state for calculating climatologically relevant aerosol properties.

4 Conclusions

We present mass-based measurements of the mixing state of BC-containing aerosol, using a soot-particle aerosol mass spectrometer, from traffic emissions in an urban environment. Observations from single particle mass spectrometry indicate that rBC co-exists with HOA in two distinct particle types: those containing a larger mass fraction of rBC, and those containing a larger mass fraction of HOA. Source apportionment of ensemble mass spectral observations using positive matrix factorization also indicates two types of rBC-containing aerosol related to traffic: rBC-rich and HOA-rich aerosol, validated by single particle observations. Ensemble measurements expand the particle size range over which mixing state can be investigated, providing a better insight into the mixing state of the particle population and indicating that approximately 90% of rBC mass resides in rBC-rich particles. These measurements were used to drive the particle-resolved aerosol box model, PartMC-MOSAIC. Our results indicate an increase in SSA of ∼0.1 when mixing state at the point of emission is treated accurately in the model compared to the assumption of uniform mixing state.
The approach described here for quantitative assessment of black carbon mixing state from traffic can also be used to assess mixing state from other sources and to explore the evolution of mixing state during atmospheric processing. Such measurements will be crucial to drive accurate model assessment of black carbon climate impacts on a broader scale.

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References


