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Collection efficiency of the soot-particle aerosol mass spectrometer (SP-AMS) for internally mixed particulate black carbon

M. D. Willis¹, A. K. Y. Lee¹, T. B. Onasch², E. C. Fortner², L. R. Williams², A. T. Lambe², D. R. Worsnop², and J. P. D. Abbatt¹

¹Department of Chemistry, University of Toronto, Toronto, Ontario, Canada ²Aerodyne Research, Inc., Billerica, Massachusetts, USA

Correspondence to: M. D. Willis (megan.willis@mail.utoronto.ca) and T. B. Onasch (onasch@aerodyne.com)

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Abstract. The soot-particle aerosol mass spectrometer (SP-AMS) uses an intra-cavity infrared laser to vaporize refractory black carbon (rBC) containing particles, making the particle beam-laser beam overlap critical in determining the collection efficiency (CE) for rBC and associated non-refractory particulate matter (NR-PM). This work evaluates the ability of the SP-AMS to quantify rBC and NR-PM mass in internally mixed particles with different thicknesses of organic coating. Using apparent relative ionization efficiencies for uncoated and thickly coated rBC particles, we report measurements of SP-AMS sensitivity to NR-PM and rBC, for Regal Black, the recommended particulate calibration material. Beam width probe (BWP) measurements are used to illustrate an increase in sensitivity for highly coated particles due to narrowing of the particle beam, which enhances the CE of the SP-AMS by increasing the laser beam-particle beam overlap. Assuming complete overlap for thick coatings, we estimate CE for bare Regal Black particles of 0.6 ± 0.1 , which suggests that previously measured SP-AMS sensitivities to Regal Black were underestimated by up to a factor of 2. The efficacy of the BWP measurements is highlighted by studies at a busy road in downtown Toronto and at a nonroadside location, which show particle beam widths similar to, but greater than that of bare Regal Black and coated Regal Black, respectively. Further BWP measurements at field locations will help to constrain the range of CE for fresh and aged rBC-containing particles. The ability of the SP-AMS to quantitatively assess the composition of internally mixed particles is validated through measurements of laboratorygenerated organic coated particles, which demonstrate that the SP-AMS can quantify rBC and NR-PM over a wide range of particle compositions and rBC core sizes.

1 Introduction

Incomplete combustion of fossil fuels, bio-fuels and biomass emit teragram quantities of anthropogenic black carbon to the troposphere every year (Bond et al., 2007, 2013). Aerosol containing black carbon is an important absorber of short wave radiation in the atmosphere, with the ability to affect the energy balance at the surface and within the troposphere (Ramanathan and Carmichael, 2008). Black carboncontaining particles are estimated to be the third largest contributor to anthropogenic global warming, after carbon dioxide and methane (Bond et al., 2013). However, the magnitude of their direct effects on radiative balance may become even more significant when black carbon is internally mixed with other aerosol components such as sulphate, nitrate and organic species (Adachi et al., 2010; Jacobson, 2001), though the importance of this enhancement for ambient aerosol remains an open question (Cappa et al., 2012; Lack et al., 2012). The physical and chemical complexity of these particles compounds the uncertainty associated with the impacts of black carbon on global and regional climate change and human health (Seinfeld, 2008; Jansen et al., 2005).

A variety of measurement techniques have been used to quantify atmospheric black carbon. These include direct techniques, such as the single particle soot photometer (SP2) that uses laser induced incandescence to quantify refractory black carbon (rBC) and indirect methods using opticaltechniques, sample oxidation, or oxidation combined with optical methods (Bond et al., 2013; Lack et al., 2014). When aerosols contain a small black carbon mass fraction, reported mass concentrations from a range of measurement techniques can differ significantly (up to 80%) (Bond et al., 2013). This large range in black carbon mass loadings can be partially attributed to measurement bias, most often for optical methods, associated with the presence of other absorbing species in the aerosol. These measurement uncertainties can make it difficult to understand the spatial and temporal trends and the climate impacts of black carbon.

The soot-particle aerosol mass spectrometer (SP-AMS) is a recently developed instrument that combines a nearinfrared laser system for volatilization of rBC with the Aerodyne high-resolution time-of-flight AMS (DeCarlo et al., 2006) to quantify aerosol components (Onasch et al., 2012). Near-IR laser vaporization and electron impact ionization facilitate mass spectral detection of light-absorbing rBC and the simultaneous measurement of non-refractory species associated with rBC. Detection of rBC and NR-PM depends upon the ability of rBC to absorb the IR energy and produce a heating effect significant enough to volatilize the rBC coating material and, upon further heating, the rBC itself. The SP-AMS has been shown to respond linearly and reproducibly to laboratory generated rBC particles (Regal Black (Cabot R400) and flame generated soot) (Onasch et al., 2012). However, in contrast to the Regal Black particles generally used for SP-AMS calibration, freshly emitted particulate black carbon tends to be smaller and less compact (i.e. lower mass-mobility exponent, Cross et al., 2010). Over time these small particles grow and become internally mixed with organic and inorganic species through condensation and coagulation. With ageing, changes in chemical composition and physical properties of rBC particles affect their atmospheric lifetime, radiative properties, and their ability to act as cloud condensation nuclei and as ice nuclei (Liu et al., 2013; Bond et al., 2013). Therefore, it is crucial that the SP-AMS is able to quantify the extent of particle coating and the rBC mass loading. The ability to simultaneously detect rBC and species that condense on it with comparable accuracy is crucial for three reasons; first, to assess the rates at which black carbon processing occurs in the atmosphere; second, to understand the role that such internally mixed particles may play as cloud nuclei; and third, to assess direct effects on solar radiation.

Previous research has applied the linear response of the SP-AMS to measure the size-resolved chemical composition of rBC-containing particles during roadside measurements of traffic related pollutants (Massoli et al., 2012), and to characterize the rBC mixing state for its relation to black carbon absorption enhancement (Cappa et al., 2012). However, the effect of rBC mixing with NR-PM on the collection efficiency of the SP-AMS laser vaporizer has not been explicitly addressed. In this work, we first examine the effect of rBC-particle coating on the SP-AMS sensitivity to the rec-



Figure 1. Laboratory schematic showing the basic particle generation, coating and characterization components. Experiments were conducted independently at the University of Toronto (UofT) and Aerodyne Research, Inc. (ARI) with similar set-ups, see text for details.

ommended calibration material, Regal Black. Secondly, we address differences in collection efficiency for bare and internally mixed rBC particles with beam width probe (BWP) measurements to demonstrate particle beam narrowing that occurs due to coating. Finally, using measured sensitivities for laboratory-generated particles, we demonstrate that the SP-AMS can quantify the fraction of rBC in coated particles over a wide range of internally mixed compositions.

2 Experimental

2.1 Particle generation and coating

Experiments were conducted at the University of Toronto (UofT) and at Aerodyne Research, Inc. (ARI) with similar experimental set-ups. A general experimental schematic is shown in Fig. 1. In both cases, rBC containing particles were generated from a water suspension of Regal Black (RB) (Regal 400R pigment, Cabot Corp.), with concentrations ranging from 10 to 1000 mg L⁻¹ in de-ionized water (18 M Ω – cm, TOC < 4 ppb), using a constant output atomizer (TSI Inc., model 3076). The polydisperse aerosol flow was dried with a diffusion drier containing silica gel (< 20 % RH) and directed to a differential mobility analyser (TSI Inc. model 3081) for size selection. The monodisperse aerosol was characterized "bare" (nascent) or after coating with *bis*(2-ethylhexyl)sebacate (BES) via condensation in a thermally controlled condenser.

The University of Toronto coating apparatus consisted of a 1.3 cm outer diameter, 30 cm long glass tube containing BES. The sample line was connected to the coating region via 0.63 cm diameter glass inlets. The temperature of the coating region was controlled using heating tape, monitored using a thermocouple and varied from room temperature up to 125 °C. Aerosol carrier flow through the coating region was 450 cm³ min⁻¹. The ARI coating apparatus consisted of a 60 cm long, 1.3 cm outer diameter glass tube wrapped in heating tape and temperature controlled using an Omega PID controller and a thermocouple. The tube has two temperature zones ($\sim 30 \text{ cm}$ each) and was operated with the downstream zone set to $\sim 20 \,^{\circ}\text{C}$ less than the upstream zone. The temperature of the upstream zone was varied from room temperature to 150 $\,^{\circ}\text{C}$. The flow through the coating tube was $750 \,\text{cm}^3 \,\text{min}^{-1}$. Experimental conditions are summarized in Table 1.

2.2 Particle characterization

The size selected bare and coated rBC particles were characterized for particle mass using a mass analyser (aerosol particle mass analyser (APM model 3600, Kanomax Inc.) or centrifugal particle mass analyser (CPMA, Cambustion), Olfert and Collings, 2005), number concentration using a condensation particle counter (CPC, TSI Inc., model 3776 and 3022a), and contributions from doubly charged particles using a scanning mobility particle sizer (SMPS. TSI Inc., model 3936). SMPS measurements were used to determine appropriate solution concentrations to use in the atomizer, such that doubly charged particles accounted for less than 10% of the total particle mass. Particle number concentration was monitored before and after coating, and used to monitor particle loss or homogeneous nucleation. The mass analyser was scanned using both the rotational speed and voltage to measure the size resolved mass distribution for coated and bare particles. From the mass analyser measurements of bare and coated Regal Black particles we define the mass ratio of organic coating to Regal Black ($R_{Org/RB} = mass_{BES}/mass_{RB}$) and the mass fraction of Regal Black ($f_{RB} = mass_{RB}/(mass_{BES} + mass_{RB})$), where $mass_{BES} = mass_{RB,coated} - mass_{RB}$.

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

The soot-particle aerosol mass spectrometer (SP-AMS) is an Aerodyne high-resolution AMS equipped with an infrared laser vaporization module similar to that of the single particle soot photometer (Droplet Measurement Technologies) as described in detail by Onasch et al. (2012). SP-AMS instruments contain both the traditional tungsten vaporizer and the IR-laser vaporizer, and can be operated with both tungsten and IR-laser vaporizers on or with only the tungsten vaporizer only, the tungsten vaporizer must be removed because its lowest possible temperature is ~ 200 °C due to heating by the filament.

In brief, the SP-AMS laser vaporization module uses a continuous wave intra-cavity 1064 nm Nd:YAG laser, situated perpendicular to the particle beam, that allows vaporization and subsequent detection of rBC species (Onasch et al., 2012). The width of the laser beam (fundamental transverse mode, TEM00, with Gaussian profile) is dictated by the length of the lasing cavity and the curvature of the optics, which are fixed in SP-AMS instruments. Particles are focused with an aerodynamic lens into a narrow beam which crosses the IR laser beam. Particles containing rBC absorb IR energy and are heated, first vaporizing non-refractory (NR) coatings (estimated temperature < 600 °C) and then vaporizing the rBC (estimated temperature ~ 4000 °C Onasch et al., 2012). Neutral gas phase analytes are ionized using 70 eV electron impact ionization. Three SP-AMS instruments were used in this study (UofT SN 215-121, ARI SN 215-039 and 215-130), all of which are time-of-flight mass spectrometers operated in V-mode with both the tungsten vaporizer (~ 600 °C) and the laser vaporizer simultaneously, unless otherwise noted.

Particle beam alignment with the tungsten vaporizer and the IR laser beam was confirmed preceding these experiments and IR laser power was optimized for saturation of the rBC signal. The physical distance between the centre of the tungsten vaporizer and the vertical position of maximum sensitivity for Regal Black in the IR laser was < 0.5 mm. Data acquisition alternated between mass spectrum (MS) and particle time-of-flight (pToF) modes. Data were analysed using the AMS analysis software Squirrel version 1.51H and Pika version 1.10H. Signals of rBC from Regal Black were quantified by the sum of C_1^+ to C_9^+ peaks, using high resolution ($R \sim 2000$ at m/z 28) MS data. The average ratio of C_1^+ to C_3^+ (0.6), obtained from bare Regal Black, was used to correct the C_1^+ peak intensity for interference from nonrefractory species in coated particles. Signals from refractory oxygen containing species can arise during the vaporization of rBC and can contribute to signals at CO^+ and CO^+_2 (Corbin et al., 2013; Onasch et al., 2012). The signals arising from CO^+ and CO^+_2 were not included in the organic loading if it accounted for greater than 10% of the total organic signal, since the appropriate chemical fragmentation of these species arising from Regal Black was not quantified in this work.

For comparison to mass analyser derived mass fractions of Regal Black ($f_{\rm RB}$) in coated particles, chemical identification and quantification was carried out with each SP-AMS instrument. The SP-AMS derived $f_{\rm RB}$ (or $R_{\rm Org/RB}$) was determined through quantification of rBC and coating, taking into account NR-PM already present on Regal Black as follows:

$$f_{\rm RB} = \frac{\rm mass_{rBC} + \rm mass_{Org, RB}}{\rm mass_{rBC} + \rm mass_{Org, RB} + \rm mass_{BES}},$$
(1)

where $mass_{Org, RB}$ is the mass of organic species present on bare Regal Black.

2.4 Particle beam width measurements

Each SP-AMS was equipped with a beam width probe (BWP) consisting of a vertical wire of 0.46 or 0.41 mm in width (UofT and ARI, respectively). The BWP wire was oriented perpendicular to the laser beam axis and stepped across the particle beam parallel to the laser axis (horizontally).

	UofT apparatus	ARI apparatus
DMA	TSI model 3081	TSI model 3081
DMA sample flow (sccm)	450	300
DMA sheath flow (sccm)	4500	3000
Coating tube diameter (cm)	1.3	1.3
Coating tube length (cm)	30	60
Coating tube flow rate (sccm)	450	750
Coating tube temperature range (°C)	25-125	25–150, with two temperature zones ($\Delta T = 20$)

 Table 1. Summary of particle generation systems.

As detailed in Huffman et al. (2005), the BWP wire blocks a fraction of sampled particles from reaching the vaporizer at each step. By analysing the results from multiple wire steps across the particle beam, the particle beam divergence can be determined. In the horizontal orientation, the laser beam produces uniform vaporization of rBC, thus allowing direct measurement of rBC particle beam widths. Particle beam widths (σ) in this work are determined by fitting a onedimensional Gaussian profile to the measured transmission profile as a function of the distance the BWP wire travels along the horizontal axis of the laser beam. These widths are slightly different widths than those derived by Huffman et al. (2005), where a two-dimensional Gaussian profile was assumed for the particle beam and the tungsten vaporizer geometry was explicitly included in their modelling. BWP measurements were also carried out by rotating the BWP such that the wire was oriented parallel to the laser beam axis and could be stepped across the particle beam perpendicular to the laser beam (vertically). Measurements in this orientation can provide some insight into the effective laser beam width for rBC. Note that the UofT SP-AMS was operated without the tungsten vaporizer during these vertical BWP measurements. See Supplement Fig. S1 for an illustration of BWP orientation for particle and laser beam width measurements.

3 Results and discussion

3.1 The effect of rBC-particle coating on SP-AMS collection efficiency

3.1.1 SP-AMS calibration

The goal of this section is to examine the effect of rBC mixing state on SP-AMS sensitivity to rBC and associated NR-PM. To do so, we measure the apparent ionization efficiency of rBC and organic coating material relative to nitrate ($RIE_{rBC, app}$ and $RIE_{Org, app}$) for a variety of particle coating thicknesses and compare these observations to uncoated values. The relative ionization efficiency, defined in previous work (Jimenez et al., 2003; Onasch et al., 2012), is applied to quantify the mass concentration of a particu-

late species. Using the mass based ionization efficiency for nitrate (mIE_{NO3}, in units of ions per picogram) the mass concentration of species "s" can be determined as follows:

$$C_{\rm s} = \frac{1}{\rm CE_{\rm s} \cdot \rm RIE_{\rm s} \cdot \rm mIE_{\rm NO_3} \cdot Q} \sum_{i} I_{\rm s, i}, \qquad (2)$$

where C_s is the mass concentration of species "s," RIE_s = mIE_s/mIE_{NO3}, Q is the flow rate, CE_s is the species specific collection efficiency, and each $I_{s, i}$ is an ion signal associated with species "s." The apparent mass based ionization efficiency is dependent on the collection efficiency (i.e. RIE_{s, app} = RIE_s · CE_s) for the particular particle type, which is given by the product of terms associated with loss due to the aerodynamic lens (E_L), beam divergence (E_S), and particle bounce (E_B) off of the tungsten vaporizer (CE = $E_L \cdot E_S \cdot E_B$) (Huffman et al., 2005). For particles with unit lens transmission, the most important loss mechanism when considering the laser vaporization module of the SP-AMS is beam divergence, such that CE $\cong E_S$ (Onasch et al., 2012).

 $RIE_{rBC, app}$ and $RIE_{Org, app}$ values were determined from independent measurements of $mIE_{rBC, app}$, $mIE_{Org, app}$ and mIE_{NO_3} . Good agreement was observed between $RIE_{rBC, app}$ values obtained from bare Regal Black over the particle size range of 250–400 nm, which suggests minimal difference in collection efficiency for these uncoated particles.

3.1.2 The effect of rBC-particle coating

RIE_{rBC, app} and RIE_{Org, app} values were measured over a range of BES coating thicknesses, with the mass ratio of BES to Regal Black ($R_{Org/RB}$) ranging from 0 to 6 (corresponding to a mass fraction of Regal Black, f_{RB} , ranging from 1 to 0.15). Particles with 200–500 nm mobility diameters, characterized in terms of their per particle masses and number concentrations, were used to ensure unit lens transmission ($E_L = 1$). Figure 2a demonstrates that as $R_{Org/RB}$ increases, the apparent sensitivity to rBC increases by approximately a factor of 2, for two SP-AMS instruments, and within the uncertainties of our measurements appears to saturate at $R_{Org/RB} > 3$. A similar trend is shown in Fig. 2b for the BES coating from each SP-AMS instrument. A comparable increase in sensitivity with particle coating was also observed M. D. Willis et.al.: SP-AMS collection efficiency

0

UofT SP-AMS (dual vapourizers)

ARI SP-AMS (laser vapourizer)

a) 0.4

Apparent RIE_{rBC}

0.3

0 2

0.1

10

8

6

2

0

1

b) 12

Apparent RIE_{Organics}

Figure 2. (a) RIE_{rBC, app} measured as a function of BES coating thickness, demonstrating an increase in sensitivity that appears to reach saturation at approximately $R_{\text{Org/RB}} = 3$. RIE_{rBC, app} values at $R_{\text{Org/RB}} = 0$ were obtained using 200 and 300 nm (d_{m}) particles. Data from two instruments is included: UofT (circles) and ARI data with the laser vaporizer only (squares, SN 215-130). (b) RIE_{Org, app} measured as a function of BES coating thickness, demonstrating a similar increase with coating thickness. The blue solid lines represent a best fit to the UofT SP-AMS data used in subsequent analysis (Sect. 3.5).

2

3

 $R_{Org/RB} = m_{Org}/m_{RB}$

4

5

6

with a third SP-AMS instrument (see Fig. S2); however, particle mass analyser measurements were not available during this experiment so the data were not included in Fig. 2. The difference in sensitivity enhancement evident in Fig. 2a between the two instruments could be due to differences in the final coated particle size or to differences in the uniformity of particle coating provided by each coating apparatus, both of which could result in differences in particle focusing in the aerodynamic lens and the resulting particle beam–laser beam overlap. The possibility of homogeneously nucleated organic particles, which cannot be neglected in these experiments, could also lead to differences in the overall RIE enhancement.

The relatively large values of RIE_{Org, app} observed in this work for BES were higher than the recommended value of 1.4 for all sampled organics from ambient sources (Al-

farra et al., 2004). As noted in Jimenez et al. (2003) individual organic molecules may exhibit different RIE values, including values greater than 1.4. Utilizing the results published in Jimenez et al. (2003) (Fig. 6), organic hydrocarbon compounds have an apparently higher RIE than highly oxygenated organic compounds (RIE ~ 2 for hydrocarbons vs. ~ 1.3 for oxygenated compounds). BES has a relatively low O: C ratio (4 : $26 \sim 0.15$), and thus may exhibit an RIE similar to hydrocarbons up to ~ 2 . For these data, the apparent RIE (defined as SP-AMS measured ions divided by the difference of the coated to uncoated per particle mass from the mass analyser) measured for BES under thin coating conditions are highly uncertain due to the difference term in the denominator, which goes to zero with thin coatings. Under thicker coating conditions, the observed RIEOrg, app values are significantly higher than 2 and increase with coating thickness. Reasons for this may be due to differences between the laser and tungsten vaporizers and could include; (1) an enhanced ionization efficiency for gas phase molecules generated in the laser vaporizer due to their position relative to the filament; (2) an improved ion extraction efficiency of ions originating from neutral gas-phase molecules vaporized at the laser vaporizer, due to their position in the ion chamber; or (3) a collection efficiency on the tungsten vaporizer of less than one $(E_b < 1)$ for organics on rBC particles. The fact that an $RIE_{Org, app} > 2$ for coated particles is observed not only with dual vaporizers, but also with the laser vaporizer alone may suggest an enhanced sensitivity to species volatilized from the laser vaporizer. In addition, the fractional increase in RIE_{Org, app} is larger than that of RIE_{rBC, app}, which may indicate the relevance of possibility (3), above, since organic species are volatilized from both vaporizers in this instrumental configuration. Further work is required to fully understand the differing responses to rBC and NR-PM in the SP-AMS.

The observed dependence of SP-AMS sensitivity on particle coating thickness (Fig. 2) is likely caused by decreased particle beam divergence with coating, resulting in greater particle beam-laser beam overlap. The saturation behaviour shown in Fig. 2 can be used to estimate collection efficiency (E_S) for bare Regal Black particles and for the organic species on thinly coated Regal Black particles. With the assumption that the sensitivity measured at saturation (i.e. highly coated particles) corresponds to complete particle beam-laser beam overlap ($E_{\rm S} = 1$ and $\rm RIE_{\rm rBC, app} =$ RIE_{rBC}), then E_S can be inferred by dividing $RIE_{rBC, app}$ (RIE_{Org, app}) values for the bare (nascent) Regal Black by the average maximum value observed in each instrument for coated particles (RIE_{rBC, app} and RIE_{Org, app}). From the data sets presented in Fig. 2, for bare (nascent) Regal Black, an $E_{\rm S}$ value of 0.6 ± 0.1 was determined. This result indicates that Regal Black calibrations used in the past, with mIE_{rBC} of $\sim 150\pm25\,\text{ions}\,\text{pg}^{-1}$ and corresponding RIE_{rBC} of 0.2 ± 0.1 (Onasch et al., 2012), are underestimated by approximately a factor of 2.



3.2 rBC particle beam divergence

In this section, the impact of particle coating on SP-AMS collection efficiency is explored using particle beam width measurements. Non-spherical particles produce more divergent particle beams compared to spherical particles, owing to lift and drag on irregularly shaped particles. As non-spherical rBC-containing particles are coated with a liquid organic species, they become more spherical in shape (e.g. Cross et al., 2010). Therefore, a change in rBC-containing particle beam divergence enough to alter the extent of particle beam–laser beam overlap, thus altering the collection efficiency (E_s). To investigate this difference in sensitivity with particle coating, the change in particle beam width was measured as a function of coating thickness with a beam width probe (BWP).

In this context, signal transmission is defined as the signal intensity at a given BWP position relative to that of an unblocked beam, normalized to particle number concentration. With a BWP centred in the particle beam, a lower signal transmission will indicate a narrower beam since a larger fraction of particles are blocked by the BWP wire. Figure 3 compares signal transmission versus BWP position curves for 200 nm uncoated Regal Black, BES coated Regal Black particles ($R_{\text{Org/RB}} \sim 3$) and 300 nm ammonium nitrate (as an example of a well focused beam), each fit to a 1-D Gaussian distribution. The standard deviation (σ) of this distribution is used to describe the particle beam width (i.e. 4σ , or $\pm 2\sigma$, defines 95.4 % of the particle beam width). As shown in Table 2 BWP measurements indicate that the particle beam narrows from $\sigma \sim 0.4$ mm for bare, 200 and 300 nm Regal Black particles to approximately 0.1-0.2 mm (ARI and UofT measurements, respectively) when the particles are coated with BES $(R_{\rm Org/RB} \sim 3)$. The differences in beam width measured in the two experiments likely arise from differences in the final coated particle size (\sim 375 nm and $d_{\rm va}$ \sim 280 nm, see Table 2). The beam width for these coated Regal Black particles approaches that of pure 230 nm BES particles ($\sigma = 0.12$ mm) (Huffman et al., 2005). However, it should be noted that 1-D Gaussian fits to BWP curves at low σ (i.e. a well focused beam) may overestimate the particle beam width (Huffman et al., 2005).

Figure 4a demonstrates the relationship between sensitivity to rBC, particle beam width and coating thickness. With a BWP centred in the Regal Black particle beam, signal transmission decreases as $R_{\text{Org/RB}}$ increases (i.e. the particle beam width decreases as coating thickness increases). Furthermore, with the BWP positioned outside of the particle beam, mIE_{rBC, app} increases as $R_{\text{Org/RB}}$ increases. These two observations suggest that as rBC particles become more coated the particle beam–laser beam overlap becomes more complete, giving rise to enhanced signal intensity for rBC and the associated NR-PM. RIE_{app} vs. SP-AMS derived $R_{\text{Org/RB}}$ plots for this data set, similar to those in Fig. 2, are presented in Fig. S2. Figure 4b demonstrates that rBC



Figure 3. Comparison of BWP signal transmission (signal intensity relative to that of an unblocked beam, normalized to particle number concentration) traces for 200 nm bare ($f_{RB} = 1$), BES coated ($R_{Org/RB} = 3$) Regal Black and 300 nm ammonium nitrate particles vaporized using the tungsten vaporizer as an example of a well focused beam. Data from the UofT SP-AMS.

and organic signal transmission both decrease due to particle coating, with an approximately one-to-one relationship. This concept is discussed further in Sect. 3.5.

3.3 Measure of effective laser width for rBC

To aid in assessing the importance of incomplete laser beam– particle beam overlap for SP-AMS quantification of rBC, an upper limit for the effective laser beam width was estimated and compared to ambient particle beam width measurements.

When the SP-AMS is operated without the tungsten vaporizer and the BWP is positioned such that the wire could be moved vertically (perpendicular to the laser beam axis), some information can be gained about the effective width of the laser beam in much the same manner as is done for particle beam widths. Since the resulting BWP position versus signal transmission curve is affected not only by the true laser beam effective width but also by the particle beam and the BWP wire width, these measurements only allow for estimation of an upper limit for the effective SP-laser width for rBC. Figure 5 demonstrates that with a 0.46 mm BWP centred in front of the IR laser an average of $97 \pm 2\%$ of the rBC signal was attenuated. From this information and the projected BWP wire width (see Fig. 5) the effective laser beam width (4σ) can be estimated as less than 0.73 mm. Therefore, the effective laser beam width (σ) is at most 0.18 mm. The narrowness of the effective laser beam width highlights the importance of determining $E_{\rm S}$ for accurate quantification of rBC mass loadings. Further measurements are required to unambiguously determine the effective IR laser beam width.

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Table 2. Summary of particle beam width measurements.

Particle type	Beam width σ (mm)	Reference
Flame soot	0.77	Huffman et al. (2005)
Ambient rBC (roadside, Toronto)	0.46 ± 0.03	this work
300 nm Regal Black	0.40 ± 0.08	this work
200 nm Regal Black	0.39 ± 0.04	this work
Ambient rBC (non-roadside, Toronto)	0.26 ± 0.04	this work
300 nm Nitrate (tungsten vaporizer)	0.22 ± 0.05	this work
Coated Regal Black (UofT, 150 nm core ^a)	0.23 ± 0.08	this work
Coated Regal Black (ARI, 300 nm core ^b)	0.11 ± 0.03	this work
230 nm DOS	0.12	Huffman et al. (2005)
Laser vaporizer (upper limit)	$\leq 0.18^{c}$	this work

^a $R_{\text{Org/RB}} \sim 3$, coated $d_{\text{va}} = 280$ nm.

^b $R_{\text{Org/RB}} \sim 3$, coated $d_{\text{va}} \ge 375$ nm.

^c Derived from BWP attenuation with the probe parallel to laser beam axis, all other measurements were made with

a perpendicular BWP orientation.

3.4 Ambient particle beam width measurements

Beam width probe measurements of ambient rBC-containing particles provide insight into the morphology, and collection efficiency, of these particles. Ambient rBC particles sampled at 5 m height and 16 m from a roadside in downtown Toronto gave wider particle beams compared to those sampled away from large roadways, suggesting that fresh rBC emissions produce more divergent beams, and that particle processing reduces particle beam divergence (see Table 2). Both ambient BWP measurements indicated beam widths narrower than that of flame soot ($\sigma = 0.77$ mm) (Huffman et al., 2005), and fall within the range of ambient particle beam widths observed by Salcedo et al. (2007).

The similarity in beam width for the "fresh" rBC particles measured here and the beam width for bare Regal Black could suggest similar collection efficiencies for these types of particles. However, previous comparisons of SP-AMS rBC mass loadings with those derived from other instruments, such as the multiangle absorption photometer (MAAP), have suggested lower overall collection efficiencies for ambient rBC-containing particles (i.e. 0.1 to 0.3) (Massoli et al., 2012; Dallmann et al., 2014). Future ambient BWP measurements, in conjunction with comparison to other instruments, will further constrain the SP-AMS collection efficiency for ambient rBC and NR-PM.

Comparison of the estimated upper limit of laser beam width ($\sigma \leq 0.18$ mm) with ambient and laboratory particle beam width measurements highlights the relevance of incomplete beam overlap for SP-AMS quantification. It is important to note that this overlap mismatch is only relevant in the dimension perpendicular to the IR laser, such that a simple comparison of particle and laser beam widths is insufficient to provide a quantitative estimate of collection efficiency.

3.5 Quantification of rBC mixing state in organic coated particles

In order to quantify mixing state in terms of the mass fraction of rBC present in ambient aerosol the SP-AMS must be operated without the tungsten vaporizer to avoid interference from externally mixed NR-PM; however, in laboratory experiments when all particles are known to contain rBC such a quantification is possible. In this section, the ability of the SP-AMS to quantify mixing state for laboratory-generated internally mixed rBC particles is examined. It is important to note that these experiments focused on liquid organic coatings, which are similar to hydrocarbon-like organic coatings but may not resemble more oxygenated organic coatings present in ambient aerosol.

To accurately quantify rBC and NR-PM simultaneously from the same particles, it is important that the effective IR laser beam width for vaporizing all particle phase species is similar. If, for example, NR-PM could be vaporized in lower energy regions of the IR laser (i.e. at the edges of the beam) that do not effectively vaporize rBC, NR-PM would have a wider effective laser beam width. In such a case, the contribution of organic species to the overall particulate mass loading could be overestimated relative to rBC for $E_S < 1$. Figure 4b demonstrates that the rBC and the organic signal transmission decrease with coating thickness in a similar manner when a BWP is placed in the centre of the particle beam. Furthermore, Fig. 2 demonstrates similar increases in RIE_{rBC} and RIE_{Org} with coating. Overall, these observations suggest that the effective IR laser beam width, in terms of its ability to vaporize different particle phase species, is similar for rBC and BES.

To further assess the ability of the SP-AMS to quantify the mixing state of rBC-containing particles, the mass fraction of Regal Black ($f_{RB} = mass_{RB}/(mass_{BES} + mass_{RB})$) in BES coated particles of various rBC core sizes (40 to



Figure 4. (a) mIE_{rBC, app} as a function of signal transmission with the beam width probe wire at the centre of the particle beam. The two sets of data were collected sequentially for each coating thickness by positioning the BWP wire outside the particle beam to measure mIE_{rBC, app} and then moving the wire into the centre of the beam to measure the associated signal transmission. Colours indicate organic coating thickness estimated from SP-AMS measurements of rBC and NR-PM (300 nm RB core). (b) Organic versus rBC signal transmission with a 0.41 nm BWP wire at the centre of the particle beam, as a function of BES coating thickness ($R_{Org/RB}$). The solid line represents a 1 : 1 relationship. Data from the ARI SP-AMS (SN 215-039).

300 nm mobility diameter) was measured with the SP-AMS and compared to f_{RB} obtained from particle mass measurements. Figure 6 demonstrates that f_{RB} values derived from the SP-AMS agree well with f_{RB} values from particle mass measurements when SP-AMS values are corrected for CE. The measured RIE_{app} in this case is the product of the RIE_{rBC} and CE, where RIE_{rBC} is the maximum ratio of the ions detected for rBC material sampled (mIE_{rBC}), assuming all rBC particles vaporize in the laser beam, to the ions detected for ammonium nitrate (mIE_{NO3}). The CE factor, which is unity or less, is a measure of the missed rBC ion signal due to poor overlap between the particle beam and the laser vaporizer; that is, it accounts for rBC particles that miss the laser



Figure 5. rBC signal transmission as a function of BWP position parallel to the laser beam axis of the SP-AMS. When the 0.46 mm BWP wire was placed in the centre of the particle beam, an average of 3 ± 2 % signal transmission was observed indicating that the effective beam width (4σ) is less than the projected width of the wire on the laser vaporizer (shaded region). This corresponds to a beam width (σ) \leq 0.18 mm. Data from the UofT SP-AMS using uncoated 300 nm ($d_{\rm m}$) Regal Black particles.

beam and are not vaporized. To obtain the corrected values, the upward trend in RIE_{app} measured for rBC and NR-PM with increasing coating thickness (Fig. 2) was fit to a sigmoid relationship to obtain the appropriate values for a given coating thickness (determined by mass analyser measurements for different rBC-core sizes).

SP-AMS values derived using the RIE_{rBC} for uncoated Regal Black and RIE_{Org} typically used for organics (i.e. 0.2 and 1.4, respectively) can underestimate $f_{\rm RB}$ by up to ~ 50 % and data in Fig. 6 exhibit a pronounced non-linear relationship. For the specific case of BES, if the RIE_{Org} value is higher than the recommended value of 1.4, the underestimation would be less than that depicted in Fig. 6. When RIE_{app} values are corrected as described above the relationship in Fig. 6 shows excellent linearity ($R^2 = 0.98$, slope = 1.01 ± 0.05).

These results indicate that the SP-AMS can accurately measure the fraction of Regal Black in laboratory generated organic coated particles, down to $f_{\rm RB}$ of ~ 0.05 for small (i.e. 40 nm) inclusions, when instrumental sensitivity and CE are well known. The ability of the SP-AMS to quantitatively characterize rBC mixing state depends strongly upon the RIE_{rBC, app} and RIE_{Org, app} values used to calculate particulate mass loadings. In addition, knowledge of the value of $E_{\rm S}$, or particle loss owing to beam divergence, is essential for quantitative SP-AMS measurements of mixing state. Future ambient and laboratory beam width probe measurements may aid in determining CE for particles with unknown morphology and coating state.



Figure 6. The fraction of Regal Black in BES coated particles (f_{RB}) , obtained from SP-AMS mass loadings using uncorrected RIE_{rBC} (0.2) and RIE_{Org} (1.4) values (triangles) and RIE values corrected using the relationship observed in Fig. 2, plotted against f_{RB} obtained from particle mass measurements. Square points refer to RIE data shown in Fig. 2 that were used to determine the appropriate RIE_{app} values to apply to data from separate experiments at various rBC core sizes (circles). A best-fit line to the data using corrected RIE values yields a slope of 1.01 ± 0.05 with $R^2 = 0.98$. The solid line represents a one-to-one relationship. Data from the UofT SP-AMS. Analogous plots for two ARI SP-AMS instruments are shown in Fig. S3.

4 Summary

Our results demonstrate that particle morphology affects the SP-AMS particle beam width, that in turn affects the collection efficiency through the overlap of the particle beam and the laser beam. Observations from laboratorygenerated internally mixed rBC-containing particles in three SP-AMS instruments indicate that particle coating influences CE through beam divergence (E_S) . We estimate CE for bare Regal Black particles, the recommended calibration material, to be 0.6 ± 0.1 suggesting that previous rBC calibrations of the SP-AMS with Regal Black were underestimated by up to a factor of 2. CE for ambient rBC-containing particles will depend on their morphology and the resulting particle-beam width, which is largely unconstrained. Future work will aim to empirically relate measurements of particle beam width to CE using a number of SP-AMS instruments, to provide a method to determine the CE of particles with unknown coating state.

Ambient BWP measurements indicate that fresh rBC containing particles sampled at the roadside in downtown Toronto have particle beam widths similar to, but greater than, bare Regal Black, and thus likely a collection efficiency

less than bare Regal Black. More processed rBC particles have a smaller beam width and therefore a larger value of CE. The importance of measuring E_S is further highlighted by the upper limit estimate of the effective laser beam width for rBC, from BWP measurements, of $\sigma \leq 0.18$ mm. Further work is required to provide an unambiguous measurement of the IR laser beam width, which could be applied in conjunction with ambient particle beam width measurements to provide a more specific estimate of CE in different sampling environments.

This work demonstrates for the first time the ability of the SP-AMS to quantitatively characterize the levels of internally mixed particulate rBC and NR-PM in laboratorygenerated particles over a wide range of compositions, when instrumental sensitivity is well known. In particular, the SP-AMS can quantify the rBC mixing state even for highly organic-rich particles ($f_{\rm RB} < 0.05$) with atmospherically relevant rBC core sizes (i.e. ≥ 40 nm mobility diameter). The quantitative capability of the SP-AMS, combined with its ability to chemically characterize NR-PM coated on rBCparticles, can provide unique insight into the characteristics of ambient black carbon.

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