Mass spectrometry of refractory black carbon particles from six sources: carbon-cluster and oxygenated ions


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Abstract

We discuss the major mass spectral features of different types of refractory carbonaceous particles, ionized after laser vapourization with an Aerodyne High-Resolution Soot-Particle Aerosol Mass Spectrometer (SP-AMS). The SP-AMS was operated with a switchable 1064 nm laser and a 600°C thermal vapourizer, yielding respective measurements of the refractory and non-refractory particle components. Six samples were investigated, all of which were composed primarily of refractory material: fuel-rich and fuel-lean propane/air diffusion-flame combustion particles; graphite-spark-generated particles; a commercial Fullerene-enriched Soot; Regal Black, a commercial carbon black; and nascent aircraft-turbine combustion particles.

All samples exhibited a spectrum of carbon-cluster ions $C_x^{n+}$ in their refractory mass spectrum. Smaller clusters ($x < 6$) were found to dominate the $C_x^{n+}$ distribution. For Fullerene Soot, fuel-rich-flame particles and spark-generated particles, significant $C_x^{n+}$ clusters at $x \gg 6$ were present, with significant contributions from multiply-charged ions ($n > 1$). In all six cases, the ions $C_1^+$ and $C_3^+$ contributed over 60% to the total $C_1^{<x<16}$ intensity. Furthermore, the ratio of these major ions $C_1^+/C_3^+$ could be used to predict whether significant $C_x^{n+}$ signals with $x > 5$ were present. When such signals were present, $C_1^+/C_3^+$ was close to 1. When absent, $C_1^+/C_3^+$ was < 0.8. This ratio may therefore serve as a proxy to distinguish between the two types of spectra in atmospheric SP-AMS measurements.

Significant refractory oxygenated ions such as $CO^+$ and $CO_2^+$ were also observed for all samples. We discuss these signals in detail for Regal Black, and describe their formation via decomposition of oxygenated moieties incorporated into the refractory carbon structure. These species may be of importance in atmospheric processes such as water uptake, aging and heterogeneous chemistry.
1 Introduction

Combustion-generated particles represent the most efficient light-absorbing particles in the atmosphere and are estimated to be the strongest anthropogenic climate warming agent after CO$_2$ (IPCC, 2007; Ramanathan and Carmichael, 2008; Bond et al., 2013). In addition to direct absorptive heating, such particles may enhance glacial melting (Flanner et al., 2007), alter convection and precipitation (Ramanathan and Carmichael, 2008), react with atmospheric trace gases (Monge et al., 2010), and serve as cloud condensation (Hitzenberger et al., 2003; Tritscher et al., 2011; Engelhart et al., 2012; Martin et al., 2012) or ice nuclei (DeMott et al., 1999; Koehler et al., 2009; Corbin et al., 2012).

Due in part to the short atmospheric lifetime (days to weeks, Cape et al., 2012) of combustion-generated particles, anthropogenic combustion sources have been singled out as ideal candidates for near-term climate mitigation. Such mitigation has been predicted to additionally improve both human health and crop yields (Shindell et al., 2012; Bond et al., 2013) when co-emitted species are also considered. Such co-emitted species may form non-absorbing particles, counteracting the warming potential of light-absorbing particles (Bond et al., 2013). Co-emitted species may also enhance the hygroscopicity of light-absorbing particles, thus altering their atmospheric lifetime (Zhang et al., 2008).

Techniques to measure and apportion combustion emissions to their sources include radiocarbon measurements (e.g. Zencak et al., 2007); X-ray, Raman or electron spectroscopy (Robertson, 2002); the combination of elemental carbon measurements with other tracers (e.g. Jeong et al., 2013); measurements of light-absorption Ångström exponents (e.g. Sandradewi et al., 2008); and single-particle mass spectrometry (e.g. Toner et al., 2008). The first three of these methods require offline filter analysis and do not provide high-time-resolution information. High-time-resolution measurements of light-absorbing carbonaceous particles are made by a number of mass- or light-absorption-based instruments (e.g. Petzold and Schönlinner, 2004; Laborde et al.,
2012), but mass spectrometry stands out for its ability to characterize chemical differences between combustion particles. Such chemical characterization may provide information on particle source, mixing state (Liu et al., 2013), and chemical evolution.

Commercial aerosol mass spectrometers typically employ either thermal-desorption or laser desorption/ionization (LDI) vapourizers. Thermal-desorption vapourization does not detect the refractory, light-absorbing carbon of combustion particles. On the other hand, LDI mass spectrometers may detect both refractory and non-refractory components. These mass spectrometers are often capable of detecting individual particles, which might thereafter be attributed to specific sources, such as different vehicle types (Toner et al., 2008). Multiple studies have correlated single-particle LDI signals with collocated mass measurements in order to estimate chemically-resolved mass loadings (Pratt and Prather, 2012; Healy et al., 2013). However, the sensitivity of LDI to matrix-influenced charge-transfer reactions (Reilly et al., 2000) and to the particle-to-particle variability of reactions within the ablation plume (Reinard and Johnston, 2008) hinders a robust quantification of atmospheric aerosols.

Although not able to directly measure refractory carbon species, the Aerodyne Aerosol Mass Spectrometer (AMS) (Jayne et al., 2000; Canagaratna et al., 2007) has in recent years been pivotal in quantifying source contributions to organic aerosol loadings (Zhang et al., 2011). The AMS quantifies Non-Refractory Particulate Matter (NR-PM) by thermal-desorption vapourization at 600 °C, followed by a separate electron-impact ionization step. This two-step vapourization-ionization process yields especially reproducible mass spectra, facilitating mass quantification (Jimenez et al., 2003). Yet, the AMS does not vapourize refractory particulate matter (R-PM) such as that in dust, sea salt, or refractory black carbon particles (rBC).

A modified AMS, the Soot-Particle AMS (SP-AMS), addresses this limitation with the addition of a 1064 nm laser vapourizer (Onasch et al., 2012). Whereas the AMS vapourizes only NR-PM, the SP-AMS vapourizes 1064 nm Light-absorbing Refractory Particulate Matter (LR-PM) via radiative heating. LR-PM includes rBC as a subset. During radiative heating of LR-PM, any less-refractory PM internally mixed with LR-
PM may be vapourized by conduction. This commonly includes such species as sul-
fates, nitrates, and organic matter. These conductively-vapourized species distinguish
the SP-AMS from incandescence-based techniques such as the single-particle soot
photometer (SP2), which quantify rBC mass based only on its most-refractory, light-
absorbing component. The relationship between these PM subsets is further illustrated
in Sect. S1.

This paper presents a discussion of the SP-AMS mass spectra of six types of rBC
particles. The six samples comprise five laboratory aerosols as well as emissions from
an aircraft engine within a test cell at the Zurich Airport in Switzerland. The discussion
emphasizes refractory mass spectral features relevant to the interpretation of ambient
SP-AMS measurements. The observed refractory signals fall into two groups: carbon-
cluster ions and oxygenated-carbon ions. In the sections below, we begin with an up-
dated discussion of SP-AMS nomenclature before presenting experimental results.

2 Experimental

2.1 Laboratory setup

A series of laboratory experiments were performed in January 2012 as part of an inten-
sive laboratory campaign, “BC-Act”. This campaign was undertaken at ETH Zurich to
investigate the physical, chemical and optical properties of various rBC particle types
using a range of techniques. Oil-free stainless-steel tubing was used prior to the SP-
AMS. All gases used were high-purity (grade 5.6 synthetic air, grade 6.0 nitrogen, grade
6.0 argon).

The laboratory setup is shown in Fig. 1. rBC particles were generated by various
methods (Sect. 2.3), diluted when necessary, and introduced into a chamber with a resi-
dence time of approximately 30 min. Dilution was achieved using either an ejector
dilutor (VKL10, PALAS GmbH, Germany) with synthetic air or a filter system. From
the residence chamber, particles were passed through two bipolar chargers, split into
two equal sample flows, and size-selected by a Differential Mobility Analyzer (DMA) in each line before characterization by a suite of instruments. This suite included a Single-Particle Soot Photometer (SP2) and Aerosol Particle Mass analyzer (APM), which measure incandescent rBC mass and shape-independent single-particle mass, respectively. Details on the SP2 and APM are given in the supplement (Sect. S5). A previous publication has discussed selected Single-Particle Soot Photometer (SP2) results from this study (Gysel et al., 2012); the present work focusses on results obtained with the SP-AMS.

The two DMAs (DMA 1: custom-built (Wiedensohler et al., 2012); DMA 2: TSI model 3071, TSI Inc., Minnesota, USA) were each calibrated using polystyrene latex spheres of the diameters used in the experiments. The calibration particles were 125 nm (Duke Scientific Corp., Palo Alto, USA), 200 nm (Duke Corp.) and 305 nm (BS-Partikel GmbH, Wiesbaden, Germany) in diameter. Some experiments were also performed with 500 nm-selected particles and polydisperse aerosols. The sample flow rate through each DMA was consistent at 2 L min$^{-1}$, with a 5 L min$^{-1}$ sheath flow. A Condensation Particle Counter (CPC, model 3010, TSI Inc. USA) monitored particle number concentrations during each experiment.

### 2.2 Soot-Particle Aerosol Mass Spectrometer

The Aerodyne Soot-Particle Aerosol Mass Spectrometer (SP-AMS) is an evolution of the Aerodyne high-resolution time-of-flight Aerosol Mass Spectrometer (AMS), to include a continuous-wave 1064 nm laser vapourizer. In this section we describe the operation principles of the standard AMS, followed by a description of the SP-AMS and its sampling configuration.

The AMS (Aerodyne Research Inc., Massachusetts, USA) has been described in detail by Jayne et al. (2000) and Canagaratna et al. (2007). Briefly, aerosols are sampled through an aerodynamic lens to focus particles before they enter a vacuum chamber. Upon entering the chamber, particles are accelerated to their size-dependent terminal velocity and traverse a sizing region before reaching the vapourizer. The AMS vapour-
izer is a porous-tungsten cone held at \( \sim 600^\circ C \), which flash-vapourizes non-refractory aerosol components before 70 eV electron-impact ionization (EI) and high-resolution time-of-flight mass analysis. Interference by gases and internal background signals is accounted for by regularly blocking incoming particles with a mechanical disc, as well as periodically filtering the sampled aerosol. In an alternate mode of operation, the mechanical disc is continuously rotated, and positioned such that particles are pulsed through a mechanical slit on its edge. This so-called Particle Time-of-Flight (PToF) mode allows measurements to be taken as a function of particle vacuum aerodynamic diameter at the cost of signal-to-noise; the chopper has a duty cycle of only 1%. We refer to “MS mode” to distinguish normal measurements from PToF measurements.

The SP-AMS (Onasch et al., 2012) is identical to the AMS but for the addition of a switchable continuous-wave 1064 nm Nd:YAG laser (Droplet Measurement Technologies, CO, USA). The laser is similar to that used by the Single-Particle Soot Photometer (Schwarz et al., 2006; Laborde et al., 2012) but has higher fluence. The laser is installed orthogonally to both the particle beam and ion extraction path, intersecting the EI ionization chamber. Whereas the AMS detects only NR-PM, the SP-AMS vapourizes LR-PM by radiative heating (notably rBC and metals) as well as PM associated with LR-PM by conductive heating, as described in the introduction. After vapourization, EI ionization and mass analysis proceed in a similar fashion to the AMS. The different modes of vapourization in the AMS and SP-AMS may result in differences between the internal energies of the resulting vapours, which may influence consequent EI fragmentation patterns (Alfarra, 2004; Onasch et al., 2012).

In this study, the AMS and SP-AMS vapourizers were employed simultaneously, such that particles transited through the laser beam on their way to the AMS vapourizer. Laser-absorbing particles such as rBC particles are therefore expected to vapourize within the laser beam, without reaching the AMS vapourizer (Onasch et al., 2012). The SP-AMS laser was switched on and off during sampling, to obtain both “regular” AMS mass spectra (laser off) and SP-AMS mass spectra (laser on). The AMS vapourizer was always on. The ion time-of-flight chamber was operated in the shorter of two flight
modes, utilising a single pass through the reflectron. PToF mode was employed 25% of the time for all laboratory samples and measurements were averaged over 90 s. Data were analyzed using the community AMS analysis software SQUIRREL (version 1.51H) and PIKA (version 1.10H) as well as custom code written in Igor Pro (version 6, WaveMetrics Inc., USA).

2.3 Particle sources

Five types of rBC particles were produced in the laboratory, by combustion, spark-generation or nebulization of commercial samples. A sixth sample was measured directly behind an aircraft turbine at a commercial turbine servicing facility (SR Technics, Zurich Airport, Switzerland). Table 1 gives a summary of the measurement schedule.

2.3.1 CAST Black (CBK) and CAST Brown (CBW)

A Combustion Aerosol STandard burner (CAST, model 00-4, modified, Jing Ltd., Zollikofen, Switzerland) produced particles via a propane diffusion flame. The CAST burner was used to produce two types of particles. The first type, referred to as CAST “Black” (CBK), was produced at a fuel-equivalence ratio \( \phi = 0.85 \) (carbon-to-oxygen ratio \( \text{C/O} = 0.25 \)), and contained little organic matter. The second type, referred to as CAST “Brown” (CBW) after its visual appearance on a filter, was produced by a fuel-rich flame (\( \phi = 1.36; \text{C/O} = 0.41 \)) and contained significant amounts of OM (Fig. S2). Additional experiments were performed with CBW after thermodenuding at 250 °C. Denuding reduced volatile OC mass to near-detection-limit for the AMS.

2.3.2 GFG spark-generated particles (GFG)

A PALAS GFG 1000 Spark Generator (Helsper et al., 1993) produced carbonaceous particles by repeated electrical discharge across graphite electrodes. The PALAS GFG generator was sparked at 75 Hz in an argon atmosphere (grade 6.0), with no additional dilution. GFG particles are morphologically similar to diesel particles (Weingart-
ner et al., 1995; Burtscher, 2005; Schneider et al., 2006), although their microstructure and chemistry are significantly different (Saathoff et al., 2003; Schmid et al., 2011).

2.3.3 Regal Black (RB)

Regal Black (Cabot REGAL R400 pigment black), a carbon black distributed by Cabot Corp., USA, is the recommended calibration standard for the SP-AMS (Onasch et al., 2012). A sample of Regal Black from lot no. GP-3901 was received from Aerodyne Research Inc. to ensure consistency. The RB sample was suspended in Milli-Q water and nebulized with grade 5.6 synthetic air in a home-built Collison-type nebulizer, before drying within a silica gel diffusion dryer.

2.3.4 Fullerene-enriched particles (FS)

Fullerene-enriched carbonaceous particles were obtained from Sigma-Aldrich (“Fullerene soot as produced”, lot no. MKBB8240V). These particles are produced by resistive heating of graphite such that carbonaceous nanoparticles nucleate above the graphite surface (Krätschmer et al., 1990). When performed under \( \sim 10 \) kPa helium, this process results in nanoparticles containing \( \sim 7 \) wt\% fullerenes (\( \sim 6 \) wt\% \( \text{C}_{60} \), \( \sim 1 \) wt\% \( \text{C}_{70} \)) (Sigma-Aldrich internal report, personal communication, 2012). FS is commonly used to calibrate Single Particle Soot Photometers (SP2) instruments, as its incandescence-to-rBC-mass response is similar to that of rBC in ambient and diesel exhaust particles (Moteki and Kondo, 2010; Laborde et al., 2012).

2.3.5 Aircraft gas-turbine engine emissions (TU)

In a separate experiment to the laboratory campaign, emissions from a civil aviation gas turbine were sampled directly behind the engine. The turbine, a CFM56-5B4-2P engine, is commonly used by modern, short-haul civilian aircraft. Turbine exhaust was sampled through an 8 mm inner diameter, stainless steel, single-point sampling probe. The probe occupied a single, fixed position directly within the engine exhaust stream.
The measurements reported here were taken from an undiluted “ICAO Annex 16” industry-standard sampling line (Crayford, 2012) and represent fuel-rich combustion. Additional details on the sampling configuration and data analysis for TU are given in the supplement (Sect. S6.3).

3 Results and discussion

The dual vapourizer configuration of the SP-AMS (Sect. 2.2) allowed the laser vapourizer to be switched periodically on or off while the metal vapourizer remained continuously on. In laser-on mode, rBC particles are expected to vapourize within the laser, along with any internally-mixed non-absorbing material volatile below 4000 K. In laser-off mode, only NR-PM is observed, which in this study was largely organic matter (OM). To distinguish between these two modes, we will refer to “SP-AMS” (laser on) and “AMS” (laser off) data, along with the terms rBC and OM.

The results presented here will focus on those mass-spectral signals present only with the SP-AMS laser on, i.e., the refractory component of the particles. All SP-AMS mass spectra contained carbon-ion clusters $C_{x}^{n+}$ as the major signals. In general, these signals were negligible in AMS mass spectra, as shown by Fig. 2. The figure compares the AMS and SP-AMS signals at $C_{1}^{+}$ and $C_{3}^{+}$ on the lower ordinate. Two sequential monodisperse experiments are shown, at 200 and 305 nm. The upper ordinate shows that CPC number concentrations were stable throughout the period. While clear $C_{1}^{+}$ and $C_{3}^{+}$ signals were observed by the SP-AMS (red symbols), the AMS data (green) were at or below detection limit. These refractory $C_{x}^{n+}$ signals are discussed in detail in Sect. 3.1.

In addition to $C_{x}^{n+}$, significant SP-AMS signals were detected from refractory $CO^{+}$ and $CO_{2}^{+}$. Figure 2 additionally shows the time series of $CO_{2}^{+}$ in circle symbols on the lower ordinate. The signal clearly increases during SP-AMS periods. Relative to $C_{x}^{+}$, a significant $CO_{2}^{+}$ signal remains in AMS periods. This is discussed in Sect. 3.2.
3.1 Carbon-cluster fragmentation patterns

The following subsections present a general description of each of the $C_x^n^+$ fragmentation patterns (Sect. 3.1.1) followed by an intercomparison of the $C_x^n^+$ patterns for the six samples (Sect. 3.1.2).

3.1.1 $C_x^n^+$ ions

For all six samples, carbon clusters containing 1 to 3 atoms ($C_x^1^–3^+$) dominated the SP-AMS mass spectrum, as illustrated by Fig. 3. The figure shows the percentage contribution of $C_x^1^–3^+$ to the total $C_x^n^+$ signal for $n = 1$ and $1 < x < 16$ ($C_x^1^–16^+$). In total, $C_x^1^–3^+$ made up 60–85% of the $C_x^n^+$ signal depending on the sample. The two ions $C_x^1^+$ and $C_x^3^+$ each contributed at least 30% of $C_x^n^+$ for all samples. For $C_x^2^+$, the contributions ranged from 3–18%. This is consistent with the expected higher stability of odd-numbered, positively-charged carbon clusters (Raghavachari and Binkley, 1987).

Carbon-cluster ions with $x > 16$ and $n > 1$ were observed for three samples: GFG, CBW, and, especially, FS. The especially-high presence of these signals in the FS mass spectrum suggests that they originated from fullerenes, although ring structures may also form when clusters contain between 7 and 36 atoms (von Helden et al., 1993; Onasch et al., 2012). Peaks at $C_x^n^+$ were generally less intense than peaks at $C_x^1^–6^+$. For example, for FS the most intense peak above the mass-to-charge ratio of $C_{16}^+$ ($m/z$ 192) was at $m/z$ 216, and represented only 2.6% of the peak intensity at $m/z$ 36. Moreover, this $m/z$ 216 peak included signals from not only $C_{18}^+$, but also from $C_{36}^{2^+}$ and $C_{54}^{3^+}$. The presence of these multiply-charged ions was evidenced by the isotope peaks $^{13}C_1^{12}C_3^{2^+} (m/z$ 216.50), $^{13}C_1^{12}C_{34}^{3^+} (216.33)$ and $^{13}C_2^{12}C_{35}^{3^+} (216.66)$. No other multiply-charged peaks were present at $m/z$ 216.00, as confirmed by the absence of significant $^{13}C_1^{12}C_{71}^{4^+}$ signal at $m/z$ 216.25. The peak areas of $^{13}C_1^{12}C_{35}^{2^+}$ and $^{13}C_1^{12}C_{54}^{3^+}$ could therefore be combined with the isotopic abundance of $^{13}C_1$ to estimate the contributions of $C_{18}^+$, $C_{36}^{2^+}$, and $C_{54}^{3^+}$ to $m/z$ 216.00 (Fig. S3). We found
that $C_{36}^{2+}$ contributed the majority of the signal at $m/z$ 216.00, representing 62.8 ± 0.8 % of the total. Of the remainder, the majority was due to $C_{18}^+$ (33.1 ± 0.6 %). $C_{54}^{3+}$ contributed only 4.1 ± 0.6 %. Extrapolating this result to all $C_x^+ > 16$ peaks suggests that fullerene ions dominated the high-$m/z$ range, although ring structures were present in significant amounts.

A high intensity of multiply-charged $C_n^{n+}$ ions is expected for fullerenes – which exist for $C_x > 30$ (von Helden et al., 1993) – due to their unique electronic and geometric structure (Scheier et al., 1994; Matt et al., 1996). However, ring- or linear-shaped carbon clusters at $C_x < 36$ may also acquire multiple charges. In the FS mass spectrum discussed above, we also observed multiply-charged $C_3$ via the isotope ion $^{13}C_1^{12}C_2^{2+}$ at $m/z$ 18.050. This ion indicates that $C_3^{2+}$ was present at only 5 % of the $C_3^+$ intensity. This is consistent with the much higher energy required to form smaller doubly-charged $C_x^{n+}$ ions (Díaz-Tendero et al., 2002; Zimmerman et al., 1991). The relative intensity of multiply-charged ions is therefore low for smaller $C_x^+$ ions, but high for fullerenes. Because of the potential for misinterpreting these multiply-charged fullerene ions, and potential differences in their production between instruments, their abundance should be checked in future SP-AMS studies.

In order to present carbon-cluster mass spectra for higher $x$, we restricted the present analysis to nominally singly-charged $C_x^+$ ions with $1 < x < 16$ ($m/z$ 12–200) and their corresponding $^{13}C_1C_{x-1}^+$ daughters. This may include interferences from multiply-charged $C_x^{n+}$ with $m/z = 12n$, along with $^{13}C_2^{12}C_{x-2}^{2n+}$ isotope ions. The $C_x^+$ mass spectra for all samples are given in Figs. S4–S9. Constraining the present discussion to $C_{1–16}^+$ allows a representation of the highest-intensity ions while avoiding ambiguity for CBW, where significant PAH signals at high $m/z$ (> 200) were difficult to distinguish from $C_x^{n+}$ signals in our mass spectra. We will below refer to the set of considered ions as $C_x^+$.
3.1.2 $C_1^+ / C_3^+$ to distinguish rBC types

Given that $C_1^+$ and $C_3^+$ represented the majority of the $C_x^+$ signal from all six samples, and that $C_x^+$ represented the majority of SP-AMS signals, this section explores the possibility of using these two ions as markers to distinguish between different sources of rBC based on their carbon-cluster fragments.

The ratio of signal intensities at $C_1^+$ and $C_3^+$ are plotted in Fig. 4 for all samples and mobility diameters listed in Table 1. The figure shows the 25th to 75th percentile as a box, with whiskers extending to the 10th and 90th percentile. The horizontal line across each box represents the median. The data include a large range of signal intensities: ion rates for GFG were 0.05 times lower than CBW, while CBK ion rates were 3 times higher. The samples fall broadly into two categories, with FS, GFG, CBW and CBWTD all having $C_1^+ / C_3^+$ close to 1, while TU, RB and CBK each have $C_1^+ / C_3^+ < 0.8$. The range of values for TU are especially narrow due to the fact that one stable engine condition is presented, as described in Sect. S6.3 of the supplement. In order to verify that the $C_1^+ / C_3^+$ ratio is reproducible, we compared our RB $C_1^+ / C_3^+$ ratio to that measured by Onasch et al. (personal communication, 2012) using the same RB sample. Onasch et al. observed a $C_1^+ / C_3^+$ ratio of 0.63 ($\circ$ in Fig. 4). The present data have a mean $C_1^+ / C_3^+$ ratio of 0.67 ($\times$ in Fig. 4), indicating that the $C_1^+ / C_3^+$ ratio may be a reproducible characteristic of this sample.

We next hypothesized that the $C_1^+ / C_3^+$ ratio may depend on the chemical structure of the rBC, and compared the $C_x^+$ mass spectra of each sample relative to FS. FS was chosen as a reference since it is commercially available, commercially traceable, and gave high-intensity signals at a large $m/z$ range. Figure 5 shows the $C_x^+$ spectral intensity for each sample plotted against FS on a log-log scale. Each spectrum is normalized to unity and excludes isotope ions for clarity. Since all spectra were dominated by $C_{1-3}^+$ (Sect. 3.1.1), these ions are found near the top-right corner, as labelled in white for RB. The white labels also apply to all data points vertically above or below the RB data points. Ions $C_{4-5}^+$ lie near to each other. None of the signals for $C_{6}^+$ were greater
than 0.1 % \(10^{-3}\) of the \(C_x^+\) signal for RB, TU, or CBK. The normalized spectra fall near the 1 : 1 line when the fractional contribution of \(C_x^+\) is similar to the respective fractional contribution of \(C_x^+\) for FS, and below it when the contribution is lower.

Figure 5 shows that the six mass spectra are grouped into the same two categories as the \(C_1^+/C_3^+\) ratios in Fig. 4. The mass spectra of GFG and CBW fall near the 1 : 1 line and are therefore similar to FS. In contrast, TU, RB, and CBK are generally shifted downwards, due to the lower relative intensities of heavier ions. The \(C_x^+\) spectra of FS, GFG, and CBW all contain significant signals up to and beyond \(C_{16}^+\), while TU, RB, and CBK were dominated by the smaller \(C_{1-5}^+\) clusters. For TU, RB and CBK, the contribution of \(C_x^+\) ions with \(x \geq 6\) was always less than 0.1 % of the total \(C_x^+\). This absence was not due to a lack of sufficient signal: the CBK mass spectrum was obtained at signal intensities 60 times higher than for GFG.

Minor quantities of \(C_1^+\) and \(C_3^+\) can be produced by organic molecules in the AMS and SP-AMS. These signals are typically negligible when compared to the overall AMS mass spectrum (e.g. Alfarra, 2004), but may be sufficient to interfere with \(C_x^+\) signals from rBC when organic mass fractions are high. We tested the possibility of organic interference in our analysis by removing OM from CBW with a thermodenuder at 250 °C prior to size selection. After thermodenuding, the AMS organic signal for 305 nm CBW particles was reduced to near detection limit (Fig. S2) whereas the \(C_1^+/C_3^+\) ratio did not change appreciably (Fig. 4).

CBW had the highest OM content (based on the relative signal of AMS organics to \(C_x^+\) ions) of the laboratory samples discussed herein. Nevertheless, \(C_1^+\) signals were a factor of 250 lower in the AMS than the SP-AMS for 305 nm non-thermodenuded CBW particles, and fell below detection limit for thermodenuded 305 nm CBW particles. A factor of 250 is large, but not insignificant: the corresponding spherical equivalent diameters required for a 1 : 1 AMS:SP-AMS signal ratio at \(C_1^+\) are approximately 50 nm for a CBW “core” and 440 nm for CBW organic matter (assuming respective material densities of 1.8 g cm\(^{-3}\) (Park et al., 2004) and 1.3 g cm\(^{-3}\), Slowik et al., 2007). The CBW OM was dominated by PAHs, which may have a \(C_1^+\) yield different to that of
typical ambient OM. However, this estimation indicates that OM mixed with rBC in ambient studies may significantly interfere with the determination of $C_1^+ / C_3^+$ from ambient rBC. To avoid such interference, either organic material may be thermodenuded prior to sampling, or a different marker using higher-mass $C_x^+$ may be useful. All samples in the present study yielded carbon clusters $C_{1-5}^+$. Considering all possible combinations of $C_{1-5}^+$, we found that $C_4^+ / C_3^+$ is probably the best alternative ratio when organic interferences are significant (Fig. S10). Like $C_1^+ / C_3^+$, this ratio also distinguishes between particle sources according to their $C_x^+$ mass spectra. The disadvantage of this ratio is its relatively lower limit of quantification; since fewer $C_4^+$ ions than $C_1^+$ ions are produced per unit mass of rBC vapourized, a higher rBC mass would be required to determine $C_4^+ / C_3^+$ rather than $C_1^+ / C_3^+$. Additional studies using a variety of organic coatings should be performed to investigate whether such coatings influence these ratios.

3.2 Refractory CO$_x^+$ signals (rCO$_x$)

3.2.1 Identification of rCO$_x$

In addition to $C_n^+$, SP-AMS signals included oxygen-containing ions such ions as CO$_2^+$, H$_2$O$_2^+$, OH$_2^+$ and O$_2^+$, which we refer to as rCO$_x$. Because similar ions may also form from OM (both refractory and non-refractory), the identification and understanding of these signals is important for the interpretation of SP-AMS measurements from unknown samples. In this section, we discuss rCO$_x$ signals from RB. The focus is on CO$_2^+$ and CO$_+$, since these ions could not have resulted from the ionization of residual water on the particles. We focus on RB as a model sample because its $C_n^+$ mass spectrum was dominated by the singly-charged ions $C_{1-5}^+$, simplifying its interpretation (Sect. 3.1.1). Additionally, RB is the recommended calibration standard for the SP-AMS (Onasch et al., 2012).

The refractory nature of rCO$_x$ is illustrated by Fig. 6. The figure plots the PToF spectrum at $m/z$ 28 (N$_2^+$ or CO$_2^+$) for both AMS (green) and SP-AMS (red) operating modes while sampling polydisperse RB particles. The SP-AMS PToF spectrum of $m/z$ 36 ($C_3^+$,
black symbols) is shown for reference. Symbols represent raw data, while curves represent smoothed data (using a 4th-order, 25-point Savitsky-Golay filter to preserve peak shape). Both the red SP-AMS and green AMS m/z 28 curves show a strong peak from gas-phase N\(_2\)(g), at approximately 0.6 ms. However, only the SP-AMS shows a second peak at 3 ms. This peak is clearly within the particle phase, and corresponds to a mode vacuum-aerodynamic diameter of 386 ± 1 nm. This second peak is coincident with the arrival of the RB particles, as demonstrated by the coincident peak at m/z 36. We infer that the second peak at m/z 28 most likely originated from CO\(^+\) formed when the rBC was vapourized.

The peak at m/z 36 appears bimodal, whereas the peak at m/z 28 appears to decrease faster than at m/z 36. This suggests that the refractory CO\(^+\) content of RB is a function of particle size. A size dependence is also indicated by a decreasing ratio of CO\(_2\)^+ to C\(_3\)^+ with increasing mobility diameter for size-selected experiments. In these experiments, the CO\(_2\)^+ / C\(_3\)^+ ratio decreased from 1.6 (125 nm) to 0.8 (500 nm). The PToF signal at m/z 44 (CO\(_2\)^+) also peaked simultaneously with m/z 36 (C\(_3\)^+) (Fig. S11), demonstrating its particulate origin as previously observed by Onasch et al. (2012).

The relative significance of rCO\(_x\) was estimated by comparing the areas of lognormal curves fitted to the polydisperse PToF spectra. Bimodal fits were used for particulate m/z 36 (C\(_3\)^+) and m/z 44 (CO\(_2\)^+). The estimated CO\(_2\)^+ / C\(_3\)^+ ratio was 1.0, consistent with the ratio found from the size-selected measurements. The corresponding CO\(^+\) / C\(_3\)^+ ratio was estimated as 20. This ratio may be positively biased by interference from N\(_2\)^+.

Furthermore, this ratio cannot be interpreted as a direct mass ratio before accounting for the ionization efficiencies of both C\(_x\)^+ and rCO\(_x\)^+. When accounting for C\(_x\)^+ and not rCO\(_x\), the ionization efficiency of RB is 0.1 (Onasch et al., 2012), suggesting that a mass-weighted CO\(^+\) / C\(_3\)^+ ratio may be lower. For a mass-weighted ratio, the ionization efficiencies of both C\(_x\)^+ and rCO\(_x\) must be known. Nevertheless, such a high ratio indicates clearly that rCO\(_x\) species are an important component of RB.

Significantly higher PToF signals in the SP-AMS than the AMS were also observed at other m/z such as 18 (H\(_2\)O\(^+\)), 17 (OH\(^+\)) and 16 (O\(^+\)) for RB. Furthermore, significant
rCO_x signals were also observed for every sample discussed above. The potential origin and significance of these signals are discussed in the next section.

### 3.2.2 Origin of rCO_x^+

The observation of increased CO_x^+ within the SP-AMS relative to the AMS mass spectrum, shown in Sect. 3.2.1, could be due to (1) oxygenated moieties incorporated into the refractory structure of the rBC, (2) fragmentation of refractory OM, (3) CO or CO_2 adsorbed to the particle surface, (4) gaseous CO or CO_2 trapped within internal voids, or (5) reaction of carbon vapour with gas-phase O_2. This section will argue that the first possibility is most likely, and discuss the implications.

The second possibility is unlikely, since the SP-AMS mass spectrum of RB did not contain significant signals from hydrocarbon ions (C_xH_y^+, C_xH_yO^+, etc.) even though the signal intensity of rCO_2^+ was close to that of C_3^+ in the RB mass spectrum. Thus, any refractory organic molecules would have to include a negligible contribution of carbon-hydrogen bonds, while simultaneously being large enough to be non-volatile at 600 °C. Moreover, for RB, these molecules would have had to form during its industrial production from hydrocarbon oils at 1500 °C. This combination of properties is physically unreasonable.

The third and fourth possibilities are not likely to be the major source of rCO_x in RB. Desorption of adsorbed CO_2 is expected to occur within minutes (Sevilla and Fuertes, 2011). The 30 min residence chamber used in the current experiments should have provided sufficient time for desorption after nebulizing in CO_2-free gas (Sect. 2.1). Voids within RB particles containing CO_2(g), would contain very little CO_2 mass per particle, since the density of CO_2(g) is 1000 times lower than rBC, yet the observed CO_2^+/C_3^+ ratio was 0.8–1.6. Finally, the fifth possibility would yield C_xO^+ and not CO_2^+ as the major product (von Helden et al., 1993). Such ions were not observed. Therefore, only the first possibility can be the major source of the observed rCO_x.

Oxygenated moieties incorporated into the refractory carbon structure of RB are expected based on its production method. RB is produced by atomizing a proprietary oil
within a furnace at 1800 K, resulting in a “carbon black” that contains only 0.5 – 1 wt% of material volatile at 1223 K. This carbon black is then treated with an HNO₃-like acid (Cabot Corp., personal communication, 2013), likely at elevated temperatures (Otake and Jenkins, 1993), in order to increase surface functionalization. This increased functionalization is indicated by an increase in the percentage of mass volatile at 1223 K, from 0.5–1 wt% to 3.5 wt%.

The acid treatment of rBC yields oxygenated moieties including hydroxyl, carbonyl, carboxyl, lactones, and acid anhydrides, depending on the reaction time (Sellitti et al., 1990; Vinke et al., 1994; Figueiredo et al., 1999, 2007). These groups thermally decompose into CO or CO₂ when heated slowly (∼5 Kmin⁻¹) under inert atmospheres, typically at 500–1200 K depending on the functional group and heating rate (Zielke et al., 1996; Figueiredo et al., 1999, 2007). The thermal decomposition of such rBC surface groups should therefore be expected in the SP-AMS. The decomposition temperature of oxygenated moieties also relates to possibility (3) above: if such moieties decompose at 500–1200 K, then no distinction between refractory, adsorbed CO₂(g) and chemically-bonded moieties can be made.

Although the acid treatment of RB is not atmospherically relevant, oxygenated surface groups may also form via reaction with such oxidants as O₂ (Otake and Jenkins, 1993; Figueiredo et al., 1999) or ozone (McCabe and Abbatt, 2008; Zielke et al., 1996) as well as other atmospheric oxidants (Saathoff et al., 2001). Reaction may occur within the combustion region (Frenklach, 2002) or afterwards (McCabe and Abbatt, 2008; Saathoff et al., 2001). In the present study, the presence of rCOₓ in the propane-flame samples, CBK and CBW, indicates the potential real-world significance of these SP-AMS species, even in the absence of further aging. The presence or formation of oxygenated groups at the rBC surface might be expected to enhance cloud-condensation-nucleus activities (Kuwata et al., 2013). Other real-world sources of rCOₓ, such as metal-oxide fuel additives (Bladt et al., 2012), may also be significant to SP-AMS measurements.
In addition to its atmospheric significance, rCO\textsubscript{x} may be significant for SP-AMS calibration. Neglecting rCO\textsubscript{x} during calibration would not only give an inaccurate sensitivity factor, but would lead to the misquantification of rBC with different degrees of oxidation. The degree of inaccuracy might be on the order of 10\%, considering the elemental analysis of acid-treated rBC by Figuieredo et al. (1999). Based on the estimated signal from CO\textsuperscript{+} in Sect. 3.2.1, this inaccuracy may be much higher.

4 Summary

For six rBC samples generated across a range of conditions, from aircraft-turbine to propane flame to spark vapourization, SP-AMS mass spectra indicate just two different overall carbon fragmentation patterns. One group of samples yielded carbon-cluster ions ranging up to a few hundred \( m/z \). The other group gave carbon-cluster mass spectra which did not extend appreciably beyond C\textsubscript{5}\textsuperscript{+}. The two groups could be distinguished according to the ratio C\textsubscript{1}\textsuperscript{+}/C\textsubscript{3}\textsuperscript{+}, which was approximately 1 for the former group and less than 0.8 for the latter.

Since the absolute C\textsubscript{n}\textsuperscript{+} signal intensity is a function of rBC mass loading, using the most intense signals, C\textsubscript{1} and C\textsubscript{3}, to distinguish between samples would minimize the mass loading needed for sufficient signal. The C\textsubscript{1}\textsuperscript{+}/C\textsubscript{3}\textsuperscript{+} ratio may therefore be useful in distinguishing sources in ambient studies. However, because organic molecules produce some C\textsubscript{1}\textsuperscript{+} and C\textsubscript{3}\textsuperscript{+} ions in the SP-AMS, samples containing a high mass fraction of OM should be thermodenuded prior to characterization, or the alternate ratio C\textsubscript{4}\textsuperscript{+}/C\textsubscript{3}\textsuperscript{+} should be used.

In addition to C\textsubscript{n}\textsuperscript{+} ions, the SP-AMS mass spectrum of all six samples included oxygenated ions. These ions most likely originated from the thermal decomposition of oxygenated moieties within the refractory carbon structure. Their quantification was hindered by high signals from gas-phase molecules such as N\textsubscript{2}\textsuperscript{+}. Future work should aim to quantify these ions for a number of samples, to provide a basis for the interpretation of ambient SP-AMS spectra in which gas-phase interferences are unavoidable.
Further studies might also investigate whether the SP-AMS is capable of providing enough information to distinguish between different refractory functional groups, as has been done using well-established offline techniques.

Supplementary material related to this article is available online at http://www.atmos-chem-phys-discuss.net/13/27561/2013/acpd-13-27561-2013-supplement.pdf.

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References


### Table 1. Summary of samples, sizes measured, and experimental programme.

<table>
<thead>
<tr>
<th>Aerosol</th>
<th>Abbreviation</th>
<th>Mobility Diameter [nm]</th>
<th>Date</th>
</tr>
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<tbody>
<tr>
<td>CAST Black</td>
<td>CBK</td>
<td>125, 200, 305</td>
<td>Jan 26</td>
</tr>
<tr>
<td>CAST Brown</td>
<td>CBW</td>
<td>125, 200, 305</td>
<td>Jan 27</td>
</tr>
<tr>
<td>Spark-generated particles</td>
<td>GFG</td>
<td>125, 200, 305, 500</td>
<td>Jan 30</td>
</tr>
<tr>
<td>Regal Black</td>
<td>RB</td>
<td>125, 200, 305, 500</td>
<td>Jan 31</td>
</tr>
<tr>
<td>Fullerene-Enriched Soot</td>
<td>FS</td>
<td>125, 200, 305, 500</td>
<td>Feb 1</td>
</tr>
<tr>
<td>Thermodenuded CBW</td>
<td>CBWTM</td>
<td>125, 200, 305</td>
<td>Feb 2</td>
</tr>
<tr>
<td>Aircraft gas-turbine particles</td>
<td>TU</td>
<td>Polydisperse (mode 25)</td>
<td>Apr 30</td>
</tr>
</tbody>
</table>
Fig. 1. Laboratory setup used for all samples except turbine particles. Aerosols were diluted before passing through a 1 m$^3$ residence chamber with residence time $\sim 30$ min. Particles then flowed through two bipolar chargers at 4 L min$^{-1}$ before being split equally between two DMAs.
Fig. 2. Lower ordinate: time series of $C_1^+$ (×), $C_3^+$, and $CO_2^+$ (○) for Regal Black during two size-selected experiments for AMS (green symbols) and SP-AMS (red symbols) measurements. Upper ordinate: CPC particle number concentration.
Fig. 3. Percentage contribution of $C_{1-3}^+$ ions to the sum of $C_x^+$ signals (1 < x < 16) for CAST “Brown” (CBW), spark-generated particles (GFG), Fullerene-enriched Soot (FS), aircraft-turbine particles (TU), Regal Black (RB), and CAST “Black” (CBK), ordered by the summed $C_{1-3}^+$ intensity. Error bars for $C_1^+$ to $C_3^+$ are propagated from Poisson ion counting statistics. The data represent an average over all selected sizes.
**Fig. 4.** SP-AMS $C_1^+/C_3^+$ ratios for CAST “Brown” particles (CBW), thermodenuded CBW (CBWTD), spark-generated particles (GFG), Fullerene-enriched Soot (FS), aircraft-turbine particles (TU), Regal Black (RB), and CAST “Black” particles (CBK). All samples were monodisperse particles at multiple sizes, except TU. Each box spans the 25th to 75th percentile. Vertical lines extend to the 10th and 90th percentiles. The horizontal line within each box shows the median. The mean $C_1^+/C_3^+$ ratio for RB is shown for the present study (×) and for independent measurements by Onasch et al. (○).
Fig. 5. Scatterplot of mass spectral intensities at each $C_x^+$ up to $x = 16$. All spectra are normalized to unity and plotted against the FS $C_x^+$ spectrum. The white numbers labelling RB (red squares) indicate the number of carbon atoms $x$ in each $C_x^+$ for both RB and FS. For clarity, some labels have been omitted, namely $C_x^+$ with $x = 6$ (next to 8), 12 (next to 14), and 10 (next to 15). Error bars indicate ion counting uncertainties, and are not shown when they are smaller than the data symbols.
Fig. 6. Signal at \( m/z \) 28 (\( \text{N}_2^+ \) or \( \text{CO}^+ \)) and \( m/z \) 36 (\( \text{C}_3^+ \)) as a function of particle time-of-flight for polydisperse Regal Black particles. SP-AMS raw data are shown by the \( \times \) and \( \bullet \) symbols, AMS data by the + symbols. The lower curves (solid, SP-AMS; dashed, AMS) show the same data after smoothing; the upper curve (dash-dotted line) is the difference of the two lower curves. The first peak (0.6 ms) corresponds to gas-phase \( \text{N}_2 \), the second (3 ms) to \( \text{CO}^+ \) originating from the RB particles. Arrows indicate the axis on which the data are plotted.