Size distribution and coating thickness of black carbon from the Canadian oil sands operations

Yuan Cheng1, Shao-Meng Li1, Mark Gordon2, and Peter Liu1

1 Air Quality Research Division, Environment and Climate Change Canada, 4905 Dufferin Street, Toronto, Ontario M3H 5T4, Canada
2 Department of Earth and Space Science and Engineering, York University, 4700 Keele Street, Toronto, Ontario M3J 1P3, Canada

Correspondence: Shao-Meng Li (shao-meng.li@canada.ca)

Received: 30 October 2017 – Discussion started: 22 November 2017
Revised: 25 January 2018 – Accepted: 9 February 2018 – Published: 23 February 2018

Abstract. Black carbon (BC) plays an important role in the Earth’s climate system. However, parameterizations of BC size and mixing state have not been well addressed in aerosol–climate models, introducing substantial uncertainties into the estimation of radiative forcing by BC. In this study, we focused on BC emissions from the oil sands (OS) surface mining activities in northern Alberta, based on an aircraft campaign conducted over the Athabasca OS region in 2013. A total of 14 flights were made over the OS source area, in which the aircraft was typically flown in a four- or five-sided polygon pattern along flight tracks encircling an OS facility. Another 3 flights were performed downwind of the OS source area, each of which involved at least three intercepting locations where the well-mixed OS plume was measured along flight tracks perpendicular to the wind direction. Comparable size distributions were observed for refractory black carbon (rBC) over and downwind of the OS facilities, with rBC mass median diameters (MMDs) between ∼135 and 145 nm that were characteristic of fresh urban emissions. This MMD range corresponded to rBC number median diameters (NMDs) of ∼60–70 nm, approximately 100% higher than the NMD settings in some aerosol–climate models. The typical in- and out-of-plume segments of a flight, which had different rBC concentrations and photochemical ages, showed consistent rBC size distributions in terms of MMD, NMD and the corresponding distribution widths. Moreover, rBC size distributions remained unchanged at different downwind distances from the source area, suggesting that atmospheric aging would not necessarily change rBC size distribution. However, aging indeed influenced rBC mixing state. Coating thickness for rBC cores in the diameter range of 130–160 nm was nearly doubled (from ∼20 to 40 nm) within 3 h when the OS plume was transported over a distance of 90 km from the source area.

1 Introduction

Oil sands (OS), a type of unconventional petroleum deposit, are naturally occurring mixtures of bitumen (a viscous form of crude oil), sand, water and small amounts of other contaminants. The OS deposit in Alberta, Canada, is estimated to contain about 1.7 trillion barrels of bitumen. This deposit is distributed in the Athabasca, Cold Lake and Peace River regions, covering a total area of ∼1.42 × 10⁵ km², of which about 10% can be recovered economically with existing technologies (Government of Alberta, 2009). Bitumen can be recovered in two ways, i.e., surface mining for the shallow reserves (e.g., less than 75 m below the surface) and using in situ technologies for the deeper deposits. Surface mining can be applied to an area of only 4800 km² area within the Athabasca region, and by 2013, about 19% of this surface minable area had been disturbed (Alberta Energy, 2017). As demand for crude oil fluctuated, oil production from the Alberta oil sands experienced periods of rapid expansion and stabilized production over the last decade, with total OS production doubling between 2004 (1.1 million barrels per day, with about 66% from surface mining) and 2014 (2.2 million barrels per day, with about 47% from surface mining) (Alberta Energy, 2016).
The OS industry in Alberta has raised concerns over environmental impacts. For example, measurement results from Kelly et al. (2009, 2010) and Kurek et al. (2013) showed that OS developments contributed organic (e.g., polycyclic aromatic hydrocarbons, PAHs) and inorganic (e.g., mercury, nickel, and thallium) pollutants to the Athabasca River watershed; these simulations by Parajulee and Wania (2014) indicated that the Canadian National Pollutant Release Inventory (NPRI) likely underestimated PAH emissions in the Athabasca OS region. Despite these studies, both the emissions and subsequent environmental impacts remain poorly understood for pollutants from the Alberta OS industry. To help address this lack of understanding, an aircraft campaign was conducted with measurements of an extensive set of air pollutants over the Athabasca OS region in the summer of 2013. Using results from the campaign, Shephard et al. (2015) validated profiles of ammonia, carbon monoxide, formic acid and methanol retrieved from the Tropospheric Emission Spectrometer (TES) satellite; Liggio et al. (2016, 2017) demonstrated the large OS surface mining facilities in Athabasca as a significant source of secondary organic aerosol (SOA) and gaseous organic acids, and Li et al. (2017) identified the surface mining facilities as a greater source of volatile organic compounds (VOCs) than previously realized.

In addition to gaseous pollutants and SOA, another focus of the 2013 aircraft campaign was black carbon (BC) emissions from the surface mining facilities and its transport downwind. BC is a distinct type of carbonaceous material formed during incomplete combustion of fossil and biomass fuels, which is strongly light-absorbing in the visible light spectral range, refractory, insoluble and typically appears as chain-like aggregates consisting of fewer than 10 to several hundred carbon spherules (Andreae and Gelencsér, 2006; Bond et al., 2013; Petzold et al., 2013; Buseck et al., 2014). BC plays a unique and important role in the Earth’s climate system as an effective absorber of solar radiation. It has relatively short atmospheric residence times but can exert a strong warming effect on global and regional climate (Ramanathan and Carmichael, 2008; Bond et al., 2013; Myhre et al., 2013). Therefore, BC emission reduction has long been considered as an important near-term climate mitigation target. However, each step along the way between the source and environmental effect of BC is complex. For example, anthropogenic BC emissions and the resulting temporal and spatial variations of BC, which can be simulated by chemical transport models, remain highly uncertain (Samset et al., 2014); parameterizations of BC size and mixing state have not been well addressed in state-of-the-art radiative transfer models (Morgenstern et al., 2017). Both factors are recognized as important sources of uncertainties in the estimate of climate forcing by BC (IPCC, 2013).

For large-scale industrial activities such as the OS surface mining operations in Athabasca, key concerns regarding BC include (but are not limited to) the quantities of BC emitted into the atmosphere, size distribution and mixing state of the freshly emitted BC particles, evolution of the BC particles including their size, mixing state and optical properties as the OS plumes are transported downwind, and BC deposition. In this study, a total of 17 flights conducted during the 2013 aircraft campaign were investigated to characterize BC emissions from six major OS surface mining facilities in the Athabasca region, with focuses on the evolution of BC size distribution and mixing state. Airborne BC measurements were performed by a single-particle soot photometer (SP2). BC mass and number size distributions were determined and compared not only for different facilities but also for different downwind distances. BC mixing state was estimated by coating thickness retrieved from the SP2, based on which the influences of photochemical aging were illustrated. Limitations of using this coating thickness to represent BC mixing state were also discussed. These results can provide insights into the evolution of BC aerosol in the real atmosphere.

2 Methods

2.1 Aircraft campaign

The aircraft campaign was conducted over the Athabasca OS region in northern Alberta between 13 August and 7 September 2013 in support of the Joint Canada-Alberta Implementation Plan for Oil Sands Monitoring (JOSM). Using instruments installed aboard the National Research Council Institute for Aerospace Research Convair-580 research aircraft, an extensive set of air pollutants (including both gaseous and particulate species) were determined with high time resolutions (Gordon et al., 2015; Liggio et al., 2016; Li et al., 2017). During this campaign, 22 flights were made for a total of about 84 h, without the influences of wet removal and cloud processing. These flights were designed (1) to quantify emissions of air pollutants from six major OS surface mining facilities including Syncrude Mildred Lake (SML), Suncor Energy OSG (SUN), Canadian Natural Resources Limited Horizon (CNRL), Shell Albian and Jackpine (SAJ), Syncrude Aurora (SAU) and Imperial’s Kearl oil sands mine (IKL), and (2) to determine atmospheric evolution of the primary pollutants. The details of the measurements, flight patterns and objectives of the flights were described in detail by Liggio et al. (2016) and Li et al. (2017). In 14 flights for emission quantitation, the aircraft was typically flown in a four- or five-sided polygon pattern encircling an OS surface mining facility, with level flight tracks at 8–10 altitudes increasing from 150 to 1370 m above ground and reaching above the mixed layer; these level flight tracks were stacked along the sides of the polygon to form a virtual box encasing the facility (Figs. 1a and S1a in the Supplement). Repeated emission flights were made over SML, SUN, CNRL and SAJ, whereas single flights were made over SAU and IKL.
of the OS source area. At least three screens were created were stacked vertically to create a virtual screen downwind direction at multiple altitudes; then these level flight tracks was flown along level flight tracks perpendicular to the wind direction measurements were used to guide the intercepting locations. The first intercepting locations were chosen at approximately 1 h downwind of the majority of the OS facilities so about 30 km. Composite Google Earth images showing flight tracks are presented in Fig. S1 for F_8/28 and F_9/4. Altitude shown here indicates the ellipsoid height.

Three flights were designed to study transformation of air pollutants emitted from the OS surface mining facilities. They were conducted in a Lagrangian pattern such that the same OS plume was sampled at different time intervals (approximately 1 h apart) as it was transported downwind from the source area (Figs. 1b and S1b). Real-time wind speed and direction measurements were used to guide the intercepting locations. The first intercepting locations were chosen at about 1 h downwind of the majority of the OS facilities so that the emitted air pollutants were well mixed and merged into large plumes. At each intercepting position, the aircraft was flown along level flight tracks perpendicular to the wind direction at multiple altitudes; then these level flight tracks were stacked vertically to create a virtual screen downwind of the OS source area. At least three screens were created for each transformation flight, without industrial emissions in between.

2.2 Aerosol sampling

Aerosols were sampled through an isokinetic, shrouded solid diffuser inlet (Droplet Measurement Technologies Inc., Boulder, CO, USA) with a NASA design as described in Huebert et al. (2004). The inlet was shared by all aerosol instruments inboard the aircraft, including a high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS; Aerodyne Research Inc., Billerica, MA, USA), a condensation particle counter (CPC, Model 3775; TSI Inc., Shoreview, MN, USA), an ultra-high-sensitivity aerosol spectrometer (UHSAS) and a single-particle soot photometer (Droplet Measurement Technologies Inc., Boulder, CO, USA). The UHSAS measures particle number size distribution in the 0.06 to 1.0 µm diameter range. Aerosol number size distributions were also measured using a forward scattering spectrometer probe (FSSP, Model 300; Particle Measuring Systems Inc., Boulder, CO, USA) housed in a pod and mounted under the right wing of the aircraft. The FSSP has a non-intrusive inletless design and measures particle number size distribution in the 0.3 to 20 µm diameter range. A comparison of the measurement results from the UHSAS and FSSP in the overlapping size range of 0.3 to 1.0 µm showed an agreement in terms of both particle numbers and their size distributions (Fig. S2). This comparison suggests that in the inlet and sampling line, both the particle loss and the evaporation loss from the particles were minimal for the < 1.0 µm size range.

2.3 BC measurements by the SP2

From the common inlet and sampling line, a SP2 was used to measure the refractory black carbon (rBC) cores on a particle-by-particle basis based on incandescent light emitted from heated rBC cores when they cross and absorb energy from a laser beam (Stephens et al., 2003; Baumgardner et al., 2004; Schwarz et al., 2006; Moteki and Kondo, 2010; Laborde et al., 2012a). The SP2 used in this study detected single-particle rBC cores in the mass range of ~0.3–16 fg, based on the calibration using regal black particles (Cappa et al., 2012). To account for the rBC cores outside this detection range, a lognormal fit was applied to the measured rBC size distribution and then extrapolated over 10–1000 nm (Schwarz et al., 2006). Here the rBC size refers to the mass equivalent diameter (D_MEV) calculated as \( [(6 \times m) / (\rho \times \pi)]^{1/3} \), where \( m \) and \( \rho \) are the mass and density of the rBC core, respectively. The value of \( \rho \) was assumed to be 1.8 g cm\(^{-3}\), which is the median \( \rho \) value recommended by Bond and Bergstrom (2006). Based on this \( \rho \) value, the SP2’s detection range for single-particle rBC core mass (~0.3–16 fg) corresponded to an rBC size detection range of ~70–260 nm in terms of \( D_{MEV} \). For either rBC mass or number concentration, a scaling factor \( F_{BC} \) was
calculated as $I_{\text{whole}}/I_{\text{detected}}$, where $I_{\text{whole}}$ indicates the integral of the lognormal fitting curve from 10 to 1000 nm, and $I_{\text{detected}}$ indicates the integral of the curve from 70 to 260 nm. Subsequently, the final rBC concentration could be determined as $F_{\text{BC}} \times C_{\text{detected}}$, where $C_{\text{detected}}$ is the detected rBC concentration (either mass or number) derived from the SP2.

All the rBC concentrations involved in this paper have been scaled by flight-specific $F_{\text{BC}}$.

In addition to emitting incandescent radiation, rBC-containing particles also scatter light when passing through the laser beam of the SP2. Coating thicknesses on rBC cores ($T_{\text{coating}}$, in nanometers, nm) can be retrieved from the scattering signals on a particle-by-particle basis, using Mie theory calculation with a series of assumptions (Schwarz et al., 2008a, b; Laborde et al., 2012b). To calculate $T_{\text{coating}}$ for an rBC-containing particle, the internally mixed particle needs to be idealized as a two-component sphere with a concentric core-shell morphology. In this study, the rBC core was assumed to have a complex refractive index of 2.26–1.26i, which was initially suggested by Moteki et al. (2010) and subsequently confirmed by Taylor et al. (2015). The coating material on a rBC core was assumed to have a complex refractive index of 1.5–0i, which is representative of the corresponding values determined for inorganic salts (e.g., ammonium sulfate) and secondary organic aerosol (Schnaiter et al., 2005; Lambe et al., 2013). The core size was held fixed at $D_{\text{MEV}}$ of the rBC core, whereas the diameter of the whole particle was varied in the Mie calculation until the modeled scattering cross section matched the measurement. The measured scattering cross section was determined by a leading-edge-only (LEO) fit to the recorded scattering signal (Gao et al., 2007). Finally, $T_{\text{coating}}$ was calculated as the difference between the radii of the whole particle and the rBC core.

A key step in retrieving $T_{\text{coating}}$ of an rBC-containing particle from its scattering signal (S) is the LEO fit, which requires at least S can be properly measured (Schwarz et al., 2008a, b; Laborde et al., 2012b; Liu et al., 2014). The LEO fit cannot be performed when $S$ is outside the SP2’s detection range of scattering intensity. Thus, $T_{\text{coating}}$ cannot be calculated for relatively small rBC cores with thin coatings (i.e., rBC-containing particles with $S$ below the lower detection limit of scattering intensity) or relatively large rBC cores with thick coatings (i.e., rBC-containing particles with $S$ above the upper detection limit of scattering intensity) (Metcalf et al., 2012; Dahlkötter et al., 2014). This limitation prohibits a direct comparison of $T_{\text{coating}}$ across all rBC cores with different sizes.

Moreover, the retrieved $T_{\text{coating}}$ could be considerably influenced by uncertainties introduced by the LEO fit. These uncertainties can be evaluated using non-rBC-containing particles. The scattering signals of non-rBC-containing particles always have the shape of a full Gaussian curve, since they will not evaporate or change in size when passing through the SP2’s laser beam. Thus, for non-rBC-containing particles, the LEO fit should in principle lead to the same scattering amplitude or the same optical size ($D_{\text{optical}}$) as that retrieved from a fit to the full scattering signal (i.e., the full-Gaussian fit) (Gao et al., 2007). In this study, the LEO and full-Gaussian fits agreed within approximately ±15% in determining $D_{\text{optical}}$ for non-rBC-containing particles (Fig. 2).

Here $D_{\text{optical}}$ were calculated from the fitted scattering amplitudes, by assuming a complex refractive index of 1.5–0i for non-rBC-containing particles. $D_{\text{optical}}$ was used in Fig. 2 to evaluate the agreement between the LEO and full-Gaussian fits because it was more directly related to $T_{\text{coating}}$ compared to the scattering amplitude.

### 2.4 Additional data sets used

Organic aerosol (OA) mass was measured with a time resolution of 10 s by the HR-ToF-AMS. Photochemical age was calculated as $-\log_{10}(\text{NO}_2 / \text{NO}_y)$, where $\text{NO}_y$ is the sum of nitrogen monoxide and nitrogen dioxide (i.e., $\text{NO} + \text{NO}_2$) and $\text{NO}_2$ refers to the total reactive oxidized nitrogen compounds (Kleinman et al., 2008). Measurements of OA, $\text{NO}_y$ and $\text{NO}_2$ during the aircraft campaign have been described elsewhere (Liggio et al., 2016).

### 3 Results and discussion

#### 3.1 rBC size distributions over the OS source region: facility-integrated results

For each flight, the measured masses of the individual rBC cores over the entire flight were first grouped into different size bins and then fitted by a lognormal curve:

$$\frac{dm}{d\log D_{\text{MEV}}} = A_{\text{mass}} \times \exp \left\{ \ln \left( \frac{D_{\text{MEV}}/X_{1,\text{mass}}}{X_{2,\text{mass}}} \right)^2 \right\}, \quad (1)$$

where $A_{\text{mass}}$, $X_{1,\text{mass}}$, and $X_{2,\text{mass}}$ are the fitting parameters. The fitting parameter $X_{1,\text{mass}}$ will be termed the mass median
of masses outside the SP2’s detection range, is independent based scaling factor (\(F_{\text{BC, mass}}\)), which accounts for the rBC masses outside the SP2’s detection range, is independent of \(A_{\text{mass}}\). Therefore, \(A_{\text{mass}}\) will not be further discussed in rBC size distribution. Similarly, rBC number size distribution could be expressed as follows:

\[
\frac{dN}{d\log D_{\text{MEV}}} = A_{\text{number}} \times \exp \left\{0 - \left[\frac{\ln(D_{\text{MEV}}/X_{1, \text{number}})}{X_{2, \text{number}}}\right]^2\right\}, \tag{2}
\]

where \(A_{\text{number}}\), \(X_{1, \text{number}}\) and \(X_{2, \text{number}}\) are the fitting parameters. \(X_{1, \text{number}}\) and \(X_{2, \text{number}}\) will be termed the number median diameter (NMD) and the number distribution width (\(\text{Width}_{\text{number}}\)), respectively. \(\text{Width}_{\text{number}}\) can be converted to the standard deviation of the rBC number size distribution (\(\sigma_{\text{number}}\)) by \(\sigma_{\text{number}} = \exp \left(\text{Width}_{\text{number}}/\sqrt{2}\right)\).

Mass and number size distributions of rBC are summarized in Fig. 3 for the 14 emission flights. As shown in Fig. 3, the rBC MMD and NMD were typically in the range of 135–145 and 60–70 nm, respectively, while both the mass and number distribution widths were approximately 0.7 (the corresponding \(\sigma_{\text{mass}}\) and \(\sigma_{\text{number}}\) were about 1.6). Most of the rBC from the surface mining facilities were from the heavy diesel trucks used to transport the mined oil sands ores to centralized locations in each facility for bitumen separation from the sands. In most cases, rBC emissions from the six major OS surface mining facilities exhibited similar size distributions. These rBC size distributions are comparable with those observed for urban emissions and source (or near-source) samples representing different types of engine exhausts (Table 1). For example, (1) during an airborne measurement conducted as part of the CalNex 2010 campaign, rBC MMD was estimated to be 122 nm over the Los Angeles Basin (Metcalfe et al., 2012); (2) rBC MMD observed in urban outflows were typically in the range of 140–180 nm, as evidenced by ground-based measurement downwind of Tokyo (Shiraiwa et al., 2007), and by aircraft-based observations over Texas (Schwarz et al., 2008a), California (Sahu et al., 2012), and western and northern Europe (McMeeking et al., 2010); (3) when mainly impacted by traffic emissions, rBC MMDs were about 100 and 120 nm for a suburban site in Paris (Laborde et al., 2013) and an urban site in London (Liu et al., 2014), respectively; (4) rBC MMD measured at urban sites in Tokyo, Japan (Kondo et al., 2011b), and Sacramento, CA (Cappa et al., 2012), were between 140 and 150 nm; (5) a laboratory study showed that the MMD was about 125 nm for rBC cores emitted from a diesel car (Laborde et al., 2012b); (6) a MMD of 126 nm was observed for rBC at Cranfield Airport in the UK, attributable to aircraft engine exhausts (McMeeking et al., 2010). Although not all of these studies determined rBC MMD and NMD simultaneously, rBC NMD were typically in the range of ~60 to 80 nm for urban emissions dominated by contributions from fossil fuel combustion (e.g., Schwarz et al., 2008a; Kondo et al., 2011b; Metcalfe et al., 2012).

A comparison of rBC size distributions between this study and previous ones confirms the finding that rBC cores emitted from fossil fuel combustion were smaller in size compared to those from biomass burning (e.g., Schwarz et al.,
Table 1. A summary of rBC MMDs representative of different types of emission sources. The calibration material and assumed density of rBC are also shown. The corresponding rBC NMDs are presented in parentheses when available.

<table>
<thead>
<tr>
<th>Campaign information</th>
<th>Calibration material</th>
<th>Density (g cm(^{-3}))</th>
<th>MMD (nm)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Urban emissions dominated by contributions from fossil fuel combustion</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Airborne measurement over California, USA, in May 2010</td>
<td>Aquadag</td>
<td>1.8</td>
<td>122 over the Los Angeles basin (NMD ≈ 60 nm) between 145 and 150</td>
<td>Metcalf et al. (2012)</td>
</tr>
<tr>
<td>Ground-based measurement downwind of Tokyo, Japan, in the summer of 2004</td>
<td>thermally denuded ambient soot</td>
<td>1.77</td>
<td>170 for urban emissions (NMD ≈ 70 nm) averaging 175 for urban emissions</td>
<td>Shiraiwa et al. (2007)</td>
</tr>
<tr>
<td>Airborne measurement over Texas, USA, in September 2006</td>
<td>glassy carbon spheres</td>
<td>2.0</td>
<td>173 and 178 for urban outflows from Liverpool, UK, and Cabauw, Netherlands, respectively (NMD ≈ 80 nm for both cases)</td>
<td>Sahu et al. (2012)</td>
</tr>
<tr>
<td>Airborne measurement over California, USA, in June 2008</td>
<td>thermally denuded ambient soot</td>
<td>2.0</td>
<td>173 and 178 for urban outflows from Liverpool, UK, and Cabauw, Netherlands, respectively (NMD ≈ 80 nm for both cases)</td>
<td>Sahu et al. (2012)</td>
</tr>
<tr>
<td>Airborne measurement over western and northern Europe in April and May 2008</td>
<td>Aquadag</td>
<td>1.8</td>
<td>146, typically in the range of 130–170 (NMD averaging 64 nm)</td>
<td>Kondo et al. (2011b)</td>
</tr>
<tr>
<td>Ground-based measurement in Paris, France, during the winter of 2010</td>
<td>fullerene soot</td>
<td>1.8</td>
<td>~ 100 when impacted by fresh traffic emissions</td>
<td>Laborde et al. (2013)</td>
</tr>
<tr>
<td>Ground-based measurement in London, UK, during the winter and summer of 2008</td>
<td>Aquadag</td>
<td>1.8</td>
<td>146, typically in the range of 130–170 (NMD averaging 64 nm)</td>
<td>Liu et al. (2014)</td>
</tr>
<tr>
<td>Ground-based measurement in Tokyo, Japan, from late August to early September 2009</td>
<td>thermally denuded ambient soot</td>
<td>1.72</td>
<td>146, typically in the range of 130–170 (NMD averaging 64 nm)</td>
<td>Kondo et al. (2011b)</td>
</tr>
<tr>
<td>Ground-based measurement in Sacramento, USA, in June 2010</td>
<td>Aquadag</td>
<td>1.8</td>
<td>~ 145</td>
<td>Cappa et al. (2012)</td>
</tr>
<tr>
<td>Near-source measurement at Cranfield Airport, UK, in September 2008</td>
<td>Aquadag</td>
<td>1.8</td>
<td>126</td>
<td>McMeeking et al. (2010)</td>
</tr>
<tr>
<td>Laboratory study for source emissions from a diesel car</td>
<td>fullerene soot</td>
<td>1.8</td>
<td>~ 125</td>
<td>Laborde et al. (2012b)</td>
</tr>
<tr>
<td><strong>Biomass burning emissions</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Airborne measurement over Texas, USA, in September 2006</td>
<td>glassy carbon spheres</td>
<td>2.0</td>
<td>~ 210 for biomass burning plumes (NMD ≈ 140 nm) averaging 193 for biomass burning plumes (NMD averaging 141 nm)</td>
<td>Schwarz et al. (2008a)</td>
</tr>
<tr>
<td>Airborne measurement over California, USA, in June 2008</td>
<td>thermally denuded ambient soot</td>
<td>2.0</td>
<td>187 for fresh biomass burning plumes in Canada (NMD = 136 nm); 207 for aged biomass burning plumes transported from Asia to the Arctic (NMD = 141 nm)</td>
<td>Kondo et al. (2011a)</td>
</tr>
<tr>
<td>Airborne measurements over Canada between June and July 2008, and over the Arctic in April 2008</td>
<td>thermally denuded ambient soot</td>
<td>2.0</td>
<td>194 and 196 for two biomass burning plumes not impacted by wet deposition (NMD = 137 and 128 nm, respectively); 152 for a biomass burning plume impacted by wet deposition (NMD = 100 nm)</td>
<td>Taylor et al. (2014)</td>
</tr>
<tr>
<td>Airborne measurement over eastern Canada in July 2011</td>
<td>Aquadag</td>
<td>1.8</td>
<td>200 and 220 (NMD between 120 and 140 nm)</td>
<td>Schwarz et al. (2010)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Aged air masses in remote areas</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ground-based measurement at a remote island in Japan during the spring of 2007</td>
<td>thermally denuded ambient soot</td>
<td>1.77</td>
<td>between 200 and 220 (NMD between 120 and 140 nm)</td>
<td>Shiraiwa et al. (2008)</td>
</tr>
<tr>
<td>Ground-based measurement at a tropospheric site in Switzerland from February to March 2007</td>
<td>glassy carbon spheres</td>
<td>1.9</td>
<td>~ 200</td>
<td>Liu et al. (2010)</td>
</tr>
<tr>
<td>Airborne measurement over western and northern Europe in April and May 2008</td>
<td>Aquadag</td>
<td>1.8</td>
<td>199 over the Atlantic Ocean (NMD ≈ 90 nm)</td>
<td>McMeeking et al. (2010)</td>
</tr>
<tr>
<td>Airborne measurement over the remote Pacific in January 2009</td>
<td>fullerene soot</td>
<td>2.0</td>
<td>~ 180 for remote atmosphere and ~ 225 for the Arctic</td>
<td>Schwarz et al. (2010)</td>
</tr>
</tbody>
</table>
The rBC MMDs and NMDs measured in biomass burning plumes were typically around 200 and 140 nm (Table 1), respectively, as supported by airborne measurements over Texas (Schwarz et al., 2008a), California (Sahu et al., 2012), Canada (Kondo et al., 2011a; Taylor et al., 2014) and the Arctic (Kondo et al., 2011a). However, wet deposition could lead to a large decrease (e.g., as much as 50 nm) in the MMD of rBC cores in biomass burning plumes (Taylor et al., 2014), suggesting that an rBC MMD substantially smaller than 200 nm does not exclude the possibility of biomass burning contributions.

Different assumptions have been made by aerosol–climate models for the size distribution of black carbon. For example, the NMD of black carbon emitted by fossil fuel combustion were assumed to be 30, 40 and 60 nm by Dentener et al. (2006; for AeroCom Phase I models), Heald et al. (2014; for a radiative transfer model coupled with GEOS-Chem) and Stier et al. (2005; for the aerosol–climate modeling system ECHAM5-HAM), respectively. According to the SP2 measurement results on rBC, including results from the present study, a NMD of 60 nm would be a more appropriate input parameter in the models for black carbon emissions from fossil fuel combustion. However, there is also a need to evaluate the unimodal assumption for black carbon size distribution (Liggio et al., 2012; Buffaloe et al., 2014), given the SP2’s limited detection range of rBC core size.

3.2 rBC size distributions over the OS source region: time-resolved results

In addition to the facility-integrated results (Fig. 3), lognormal fits were also applied to 2 min intervals of rBC data derived from the SP2. Figures 4 and 5 show results from the emission flights conducted for CNRL on 26 August 2013 (i.e., F_8/26) and for SUN on 28 August 2013 (i.e., F_8/28), respectively. In both cases, the rBC mass and number size distributions did not exhibit major temporal variations, despite the minor fluctuations observed during F_8/28. The stable rBC size distribution within a flight can be more readily seen from Fig. 6a, which indicates that the rBC MMD, mass distribution width and therefore the mass-based scaling factor (\(F_{\text{rBC, mass}}\)) were independent of rBC concentration. As shown in Fig. 6a and Table 2, the variations of rBC MMD, mass distribution width and \(F_{\text{rBC, mass}}\) were within 5% for F_8/26. Larger variations in rBC size distribution were observed for F_8/28, but the variations in these three parameters were still within 10%. The variations of rBC NMD, number distribution width and number-based scaling factor (\(F_{\text{rBC, number}}\)) were also within 10% for both F_8/26 and F_8/28 (Table 2).

![Figure 4](image_url). Time-resolved rBC (a) mass size distribution, (b) number size distribution and (c) concentrations observed over the CNRL facility during F_8/26. Solid lines in (a) and (c) indicate MMD and NMD, respectively. The horizontal axis shows UTC time.

The temporal variations of rBC concentration shown in Figs. 4 and 5 were mainly driven by the in- vs. out-of-plume differences. There was a sharp increase in rBC concentration when the aircraft flew into a plume, whereas the rBC concentration deceased rapidly when the aircraft left the plume. Therefore, the stable rBC size distributions observed for the emission flights, which were clearly independent of rBC concentration (e.g., Fig. 6a), signify negligible difference in the rBC size distribution between the in- and out-of-plume conditions over the OS source region. The size distribution consistency for rBC is observed regardless of the threshold rBC concentration used to distinguish the in- and out-of-plume conditions, which is flight-dependent (e.g., \(\sim 0.1 \mu g m^{-3}\) in terms of 2 min averaged rBC mass concentration for F_8/26 as shown in Fig. S3). The implications of consistent size distributions for rBC near the sources are further discussed in Sect. 3.3 together with results from the transformation flights.

In addition to rBC concentration, the in- and out-of-plume air masses had different photochemical ages as indicated by their values of \(-\log_{10}(NO_x / NO_y)\), determined from concurrent measurements of NOx and NOy. As shown in Fig. 6b, there was a robust negative correlation between the rBC mass
Table 2. Variations of the parameters derived from time-resolved lognormal fits to single-particle rBC data measured during F_8/26 and F_8/28. Variations are determined as relative standard deviations (RSDs, in percentage).

<table>
<thead>
<tr>
<th></th>
<th>MMD Width</th>
<th>$F_{rBC, mass}$</th>
<th>NMD Width</th>
<th>$F_{rBC, number}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>F_8/26</td>
<td>1.46</td>
<td>4.42</td>
<td>2.82</td>
<td>4.48</td>
</tr>
<tr>
<td>F_8/28</td>
<td>6.85</td>
<td>8.46</td>
<td>9.47</td>
<td>7.94</td>
</tr>
</tbody>
</table>

Figure 5. Time-resolved rBC (a) mass size distribution, (b) number size distribution and (c) concentrations observed over the SUN facility during F_8/28. Solid lines in (a) and (c) indicate MMD and NMD, respectively. The horizontal axis shows UTC time. The flight track of F_8/28 is shown in Fig. 1a.

Figure 6. Dependences of (a) rBC MMD, mass distribution width ($Width_{mass}$) and mass-based scaling factor ($F_{rBC, mass}$), and (b) photochemical age on rBC concentration during F_8/26. Time resolution is 2 min for all the parameters shown here. Based on the temporal variation of 2 min averaged rBC mass concentration (Fig. S3), rBC $\leq 0.1 \mu g m^{-3}$ can be used as an indicator for typical out-of-plume conditions during F_8/26. Uncertainties introduced by lognormal fitting are within $\pm 5$ nm and $\pm 0.06$ for the time-resolved MMD and $Width_{mass}$, respectively.

3.3 rBC size distributions downwind of the OS source region

Mass and number size distributions of rBC are shown in Figs. 7 and 8, respectively, for the transformation flight conducted on 4 September 2013 (i.e., F_9/4) which reached a downwind distance of approximately 120 km (relative to the downwind edge of the OS source area; Fig. 1b). As can be seen from the time-resolved lognormal fitting results (Figs. 7a and 8a), both the rBC mass and number size distributions were fairly stable during F_9/4, without major temporal change patterns. For the typical in- and out-of-plume conditions of F_9/4, the rBC MMDs were 143 and 142 nm with mass distribution widths of 0.72 and 0.71, respectively (Fig. 7b); the rBC NMDs were 71 and 69 nm with number distribution widths of 0.68 and 0.69, respectively (Fig. 8b). These rBC size distributions (Figs. 7b and 8b) were derived from the SP2 measurements performed on the various virtual screens, where the aircraft was flown along level flight tracks (primarily at $\sim 450$ and 600 m) perpendicular to the wind direction. For the level flight tracks, the typical in- and out-of-plume conditions (i.e., segments) were distinguished by rBC concentration (Fig. 9), i.e., the typical out-of-plume conditions were identified by relatively low and constant rBC concentrations whereas the typical in-plume conditions were
Figure 7. (a) Time-resolved rBC mass size distribution observed during the transformation flight F_9/4, (b) comparison of rBC mass size distributions between typical in- and out-of-plume conditions, (c) comparison of in-plume rBC mass size distributions among successive flight screens, and evolutions of (d) average rBC mass concentration and (e) photochemical age from screen 1 (S1) to screen 4 (S4). Scaling of out-of-plume rBC size distribution in (b), scaling of rBC size distributions for screens 2 to 4 in (c), and reasons for excluding results from screen 5 in (d) to (e) are explained in the text.

The rBC mass size distribution was characterized by sharp increases in rBC concentration above the out-of-plume level. In Fig. 7b, the rBC mass size distribution was scaled for the out-of-plume conditions to reveal their lower rBC concentrations compared to the in-plume conditions (Fig. 7d). When performing the scaling, the in-plume rBC size distribution was used as a reference (i.e., kept unchanged). The out-of-plume rBC size distribution was scaled to make the $I_{\text{out-of-plume}}$ to $I_{\text{in-plume}}$ ratio equal to the rBC out-of-plume to rBC in-plume ratio, where the individual terms, in sequence, represent the integral of the scaled out-of-plume rBC size distribution curve, the integral of the reference in-plume rBC size distribution curve, the average out-of-plume rBC mass concentration (54 ng m$^{-3}$, derived from Fig. 7d) and the average in-plume rBC concentration (208 ng m$^{-3}$, derived from Fig. 7d). In Fig. 8b, the out-of-plume rBC number size distribution was scaled similarly. As can be seen from Figs. 7b and 8b, the in- vs. out-of-plume difference was negligible for rBC size distribution downwind of the OS region.

Photochemical ages were older for the out-of-plume conditions compared to the in-plume ones, by $\sim$0.3–0.5 in terms of $-\log_{10} (\text{NO}_x / \text{NO}_y)$ for different screens of F_9/4 (Fig. 7e). Therefore, the consistent rBC size distributions between the in- and out-of-plume conditions indicated that photochemical age had little influence on rBC size distribution downwind of the OS region. This conclusion was also strongly supported by the comparison of in-plume rBC size distributions among different downwind distances. As the OS plume was transported downwind, the in-plume rBC concentration decreased due to dilution (Fig. 7d), from $\sim$310 ng m$^{-3}$ for the first screen (screen 1) to $\sim$110 ng m$^{-3}$ for the fourth screen (screen 4); meanwhile, the in-plume photochemical age $-\log_{10} (\text{NO}_x / \text{NO}_y)$ increased (Fig. 7e), from $\sim$0.1 for screen 1 to $\sim$0.5 for screen 4. The last screen (screen 5) did not differ largely from screen 4 with respect to either in-plume rBC concentration or photochemical age, appearing to indicate that the dilution and aging processes had slowed down or even stopped since screen 4. However, it should be noted that, unlike the first four screens, screen 5 did not capture the full OS plume, i.e., the plume edges were missed. Compared to the central portion of the plume, the plume edges had lower rBC concentrations and older photochemical ages. Therefore, the average rBC concentration and $-\log_{10} (\text{NO}_x / \text{NO}_y)$ could not be compared directly between screen 5 and the first four screens, and consequently, results...
Figure 9. Identification of typical in- and out-of-plume conditions for two level flight tracks at ∼450 and 600 m (in terms of ellipsoid height, equivalent to ∼150 and 300 m above ground) on the first virtual screen of the transformation flight F_9/4.

from screen 5 were not involved in Fig. 7d and e. Nonetheless, for all successive screens of F_9/4, the in-plume rBC MMDs and NMDs were found to fall into a narrow range of 140–145 and 69–72 nm, respectively, while both the mass and number distribution widths were about 0.7 (Figs. 7c, 8c and 10). In Figs. 7c and 8c, rBC size distributions derived from successive screens were scaled to reveal the decrease in rBC concentration caused by dilution, using the same approach as that described in detail for Fig. 7b. The scaling requires rBC concentration representative for the full plume and thus was not performed for screen 5. A direct comparison of rBC size distributions between screen 5 and the first four screens is provided by Fig. 10. Figure 10 also demonstrates consistent in-plume rBC size distributions among successive screens for the other two transformation flights that were conducted on 19 August and 5 September 2013, respectively (i.e., F_8/19 and F_9/5), providing further solid evidence for the negligible influence of atmospheric aging on rBC size distribution downwind of the OS source region.

As shown in Table 1, previous studies conducted in remote areas (either ground- or aircraft-based) typically showed rBC MMD between 200 and 220 nm (Shiraiwa et al., 2008; Liu et al., 2010; McMeeking et al., 2010; Schwarz et al., 2010), substantially higher than those observed over urban areas (e.g., 122 nm over the Los Angeles basin; Metcalf et al., 2012) or at urban locations (e.g., 146 nm in Tokyo, Japan; Kondo et al., 2011b). Moreover, the rBC MMD was found to be 20 nm larger for aged urban plumes from Nagoya, Japan, compared to fresh emissions from the same urban area (Moteki et al., 2007). Therefore, it has been argued that rBC size distribution tends to shift toward larger sizes during aging (e.g., McMeeking et al., 2010). Results from the present study, especially the comparison of rBC size distributions among successive flight screens (Fig. 10), indicate that this is not necessarily the case. It is inferred that not all aging processes will change rBC size distribution and instead, influences of aging on rBC size distribution may partially depend on the presence of atmospheric processes that can lead to increased rBC core mass and size in a single particle (e.g., rBC coagulation and evaporation of cloud droplets containing multiple rBC particles). In this study, it appears that no such processes were at play, and within the photochemical ages encountered, rBC core masses and sizes did not change.

In addition to the evolution of in-plume rBC concentration, Fig. 7d shows that the out-of-plume rBC concentration decreased until screen 3. This decrease was associated with an increase in $-\log_{10}(NO_x/NO_y)$ for the out-of-plume conditions (Fig. 7e). For screen 4, both the out-of-plume rBC concentration and photochemical age were nearly the same as the respective values observed for screen 3. Therefore, the out-of-plume conditions identified for screens 3 and 4 should be more representative of the background. For screens 3 and 4, rBC size distributions agreed well between the in- and out-of-plume conditions, within ±3 nm in terms of MMD or NMD, indicating that the background did not differ significantly from the OS emissions with respect to rBC size distribution. Consistent in- and out-of-plume rBC size distributions observed at smaller downwind distances (i.e., for screens 1 and 2) and over the OS source area (i.e., for the emission flights) pointed to the same conclusion, although the out-of-plume conditions in these cases were less representative of the background. The rBC cores in the background could be from the OS emissions and/or long-range transported urban emissions that had not been influenced by atmospheric processes that can change single-particle rBC core size. These two kinds of emissions did not differ largely in rBC size distribution (as discussed in Sect. 3.1) and therefore they were difficult to further distinguish by rBC size only.
3.4 Evolution of rBC mixing state

Coating thickness ($T_{\text{coating}}$) was found to exhibit a decreasing trend with the increase in rBC $D_{\text{MEV}}$ for both the transformation (Fig. 11) and emission flights (Fig. S4). This trend was primarily attributed to the limitation that the detection range of $T_{\text{coating}}$ is rBC $D_{\text{MEV}}$ dependent (as explained in Sect. 2.3), rather than indicating that relatively small rBC cores were more thickly coated than larger ones. Besides $T_{\text{coating}}$, the fraction of rBC cores that can be assigned a coating thickness ($F_{\text{assigned}}$, in percentage) was also rBC $D_{\text{MEV}}$ dependent such that $F_{\text{assigned}}$ was found to be the highest (between ~35–45 %) for rBC cores in the $D_{\text{MEV}}$ range of 130–160 nm (Figs. 11 and S4). The rBC-containing particles in this $D_{\text{MEV}}$ range were selected for further discussions on $T_{\text{coating}}$ (their $T_{\text{coating}}$ will be specified as $T^*$), with a focus on the evolution of rBC mixing state as the OS plumes were transported downwind.

As shown in Fig. 12a for the transformation flight F_9/4, the in-plume $T^*$ exhibited an increasing trend with the increase in downwind distance or transport time, e.g., from ~22 nm for screen 1 to ~41 nm for screen 4. This trend is not surprising given the continuous formation of SOA during transport of the OS plumes (Liggio et al., 2016). For rBC near the sources, $T^*$ was close to zero as observed from the emission flights over the OS facilities. For example, $T^*$ was derived at ~3 nm for F_9/3 (Fig. S4). These freshly emitted rBC cores grew a coating of ~20 nm thickness in the first hour after emission, when the OS plume was transported from the sources in the OS facilities to the downwind edge of the OS region. $T^*$ were found to be comparable between the in- and out-of-plume conditions for screen 1, which were ~22 and 23 nm, respectively (Fig. 12a). It is unlikely that the out-of-plume $T^*$ could be as low as ~23 nm, if the majority of the out-of-plume rBC cores were from long-range transport. Therefore, the rBC cores observed in the out-of-plume conditions should also be influenced by emissions in the oil sands region, albeit at much lower air concentrations compared to the plumes, such as from on-road traffic that was not part of any oil sands surface mining facility.

Compared to in plumes, the increase in $T^*$ was smaller for the out-of-plume conditions as the OS plume was further transported from screen 1 (Fig. 12a) and moreover, the out-of-plume $T^*$ stopped increasing after screen 3 such that it was ~32 nm for both screens 3 and 4. One explanation for the different evolution patterns of the in- and out-of-plume $T^*$, which had comparable initial values (i.e., those for screen 1), is the less effective formation of coating materials (e.g., SOA and sulfate) for the out-of-plume conditions than in plumes. Coating precursors (volatile organic compounds and sulfur dioxide) were much more abundant in the plumes, from which fast formation of SOA was observed (Liggio et al., 2016). As shown in Fig. 12b, the in-plume OA-to-rBC mass ratio exhibited a robust increasing trend with the in-
increase in downwind distance (e.g., by ~150 % for screen 4 relative to screen 1), whereas the increase in OA-to-rBC ratio was less significant for the out-of-plume conditions (e.g., by only ~45 % for screen 4 compared to screen 1) which was negligible between screens 3 and 4.

We did not compare $T_{coating}$ measured in this study with results from previous ones due to the following reasons. (1) The detection range of $T_{coating}$ and therefore the estimated $T_{coating}$ depend on the SP2’s detection range of scattering intensity, which could differ among different SP2 instruments. This dependency indicates that different SP2 instruments might lead to different $T_{coating}$ estimates even for the same ensemble of rBC-containing particles. (2) The detection range of $T_{coating}$ and therefore the estimated $T_{coating}$ also depend on the rBC core size (i.e., $D_{MEV}$). Quite different $D_{MEV}$ ranges have been used by previous studies to estimate $T_{coating}$, e.g., 190–210 nm by Schwarz et al. (2008a, b) vs. 162–185 nm by Langridge et al. (2012), indicating that these $T_{coating}$ estimates are not directly comparable. (3) Comparison of the LEO and full-Gaussian fits for the determination of $D_{optical}$ or scattering amplitude, which should be done using non-rBC-containing particles, was not presented in many previous publications reporting $T_{coating}$. This is a concern because the LEO fit has been considered reliable as long as the LEO-to-full-Gaussian ratios are relatively constant (not necessarily around 1.0) for the fitted scattering amplitudes (e.g., Metcalf et al., 2012). Since an agreement between the LEO and full-Gaussian fits was not always required, previously reported $T_{coating}$ might be biased by the LEO-induced uncertainty to different extents, adding to the difficulties in comparing $T_{coating}$ across studies.

4 Conclusions

An aircraft campaign was conducted over the Athabasca oil sands region in the summer of 2013, during which the size distribution and coating thickness of refractive black carbon cores were studied as they were emitted from the sources and as they were transported downwind. The rBC size distributions were found to be comparable at the six major OS surface mining facilities, typically with mass median diameters of 135–145 nm and number median diameters of 60–70 nm that were characteristic of fresh urban emissions dominated by contributions from fossil fuel combustion. The observed rBC size distributions were consistent (in terms of MMD, NMD and the corresponding distribution widths) not only for the typical in- and out-of-plume segments of a flight but also for different downwind distances from the OS source area, indicating little dependence of the rBC size distribution on atmospheric aging within 4 to 5 h from the point of emission.

The coating thicknesses ($T_{coating}$) were retrieved for rBC-containing particles from their scattering signals, on a particle-by-particle basis. The uncertainty of the LEO fit, a key step to compute $T_{coating}$, was evaluated using non-rBC-containing particles. The LEO fit and the reference full-Gaussian fit were found to agree within approximately ±15 % in determining the optical size of non-rBC-containing particles. Mainly due to the SP2’s limited detection range of scattering intensity, however, $T_{coating}$ could not be calculated for all the detected rBC cores. The fraction of rBC cores that can be assigned a coating thickness was found to be the highest but still lower than 50 % for those in the diameter range of 130–160 nm. It is not surprising that $T_{coating}$ increased as the OS plumes were transported downwind, resulting from the formation mainly of secondary organic aerosols but also of sulfate. Such a coating increase with aging can significantly change the optical properties of the rBC-containing particles and hence their potentials for radiative forcing. Therefore, to resolve the coating impact on rBC radiative forcing, secondary organic aerosol and sulfate on the global basis need to be well quantified along with accurately determined black carbon emission rates.

Data availability. The rBC size distribution and coating thickness data involved in this paper can be found in the Supplement.

Supplement. The supplement related to this article is available online at: https://doi.org/10.5194/acp-18-2653-2018-supplement.

Competing interests. The authors declare that they have no conflict of interest.

Special issue statement. This article is part of the special issue “Atmospheric emissions from oil sands development and their transport, transformation and deposition (ACP/AMT inter-journal SI)”. It is not associated with a conference.

Acknowledgements. We would like to thank the National Research Council of Canada flight crew of the Convair-580, the technical support staff of the Air Quality Research Division and Stewart Cober for the management of the study. This project was supported by Environment and Climate Change Canada’s Climate and Clean Air Programme (CCAP) and the Canada-Alberta Oil Sands Monitoring program.

Edited by: John Liggio
Reviewed by: Darrel Baumgardner and two anonymous referees

References


Y. Cheng et al.: Black carbon over the oil sands region of Alberta

Atmos. Chem. Phys., 18, 2653–2667, 2018


www.atmos-chem-phys.net/18/2653/2018/


