The filter loading effect by ambient aerosols in filter absorption photometers depends on the mixing state of the sampled particles

Luka Drinovec\textsuperscript{1,2}, Asta Gregorič\textsuperscript{3}, Peter Zotter\textsuperscript{4}, Robert Wolf\textsuperscript{4}, Emily Anne Bruns\textsuperscript{5}, André S.H. Prévôt\textsuperscript{4}, Jean-Eudes Petit\textsuperscript{5,6,\$}, Olivier Favez\textsuperscript{5}, Jean Sciare\textsuperscript{6,7}, Ian J. Arnold\textsuperscript{8,\%}, Rajan K. Chakrabarty\textsuperscript{8,+}, Hans Moosmüller\textsuperscript{8}, Agnes Filep\textsuperscript{9}, Griša Močnik\textsuperscript{1,2}

\textsuperscript{[1]} Research and Development Department, Aerosol d.o.o., Ljubljana, Slovenia
\textsuperscript{[2]} Condensed Matter Physics Department, Jožef Stefan Institute, Ljubljana, Slovenia
\textsuperscript{[3]} Center for Atmospheric Research, University of Nova Gorica, Nova Gorica, Slovenia
\textsuperscript{[4]} Laboratory of Atmospheric Chemistry, Paul Scherrer Institute (PSI), 5232 Villigen PSI, Switzerland
\textsuperscript{[5]} Institut National de l’Environnement Industriel et des Risques, Verneuil-en-Halatte, France
\textsuperscript{[6]} Laboratoire des Sciences du Climat et de l’Environnement (CNRS-CEA-UVSQ), CEA Orme des Merisiers, Gif-sur-Yvette, France
\textsuperscript{[7]} Energy Environment and Water Research Center, The Cyprus Institute, Nicosia, Cyprus
\textsuperscript{[8]} Desert Research Institute, Nevada System of Higher Education, Reno, USA
\textsuperscript{[9]} MTA-SZTE Research Group on Photoacoustic Spectroscopy, Department of Optics and Quantum Electronics, University of Szeged, Szeged, Hungary
\textsuperscript{[#]} now at: Lucerne University of Applied Sciences and Arts, School of Engineering and Architecture, Bioenergy Research, Horw, Switzerland
\textsuperscript{[$\$$]} now at: Air Lorraine, Metz, France
\textsuperscript{[+] now at: Department of Energy, Environmental & Chemical Engineering, Washington University in St. Louis, St. Louis, MO, USA
\textsuperscript{[\%]} now at: College of Optical Sciences, University of Arizona, Tucson, AZ, USA

\textsuperscript{Correspondence to:} Luka Drinovec (luka.drinovec@aerosol.si) & Griša Močnik (grisa.mocnik@aerosol.si)
Abstract. Black carbon is a primary aerosol tracer for high-temperature combustion emissions and can be used to characterize the time evolution of its sources. It is correlated with a decrease in public health and contributes to atmospheric warming. Black carbon measurements are usually conducted with absorption filter photometers, which are prone to the filter-loading effect – a saturation of the instrumental response due to the accumulation of the sample in the filter matrix. In this paper, we investigate the hypothesis that this filter-loading effect depends on the optical properties of particles present in the filter matrix, especially on the black carbon particle mixing state. We conducted field campaigns in contrasting environments to determine the influence of source characteristics, particle age and coating on the magnitude of the filter-loading effect. High time resolution measurements of the filter-loading parameter in filter absorption photometers show daily and seasonal variations of the effect. The variation is most pronounced in the near-infrared region, where the black carbon mass concentration is determined. During winter, the filter-loading parameter value increases with the absorption Ångström exponent. It is suggested that this effect is related to the size of the black carbon particle core as the wood burning (with higher values of the absorption Ångström exponent) produces soot particles with larger diameters. A reduction of the filter-loading effect is correlated with the availability of the coating material. As the coating of ambient aerosols is reduced or removed, the filter-loading parameter increases. Coatings composed of ammonium sulfate and secondary organics seem to be responsible for the variation of the loading effect. The potential source contribution function analysis shows that high values of the filter-loading parameter in the infrared are indicative of local pollution, whereas low values of the filter-loading parameter result from ageing and coating during long range transport. Our results show that the filter-loading parameter can be used as proxy for determination of the particle mixing state, thus allowing for differentiation between local/fresh and transported/aged particles.

Keywords: black carbon, mixing state, Aethalometer, filter-loading effect
1 Introduction

Black carbon is a primary aerosol tracer for high-temperature combustion emissions and can be used to characterize the time evolution of its sources (Lack et al., 2014). Investigations performed on ambient black carbon mass equivalent concentrations ($BC$) have shown a stronger correlation with adverse public health effects than the ones observed for the total mass of particulate matter (Janssen et al., 2011; Janssen et al., 2012; Grahame et al., 2014; Olstrup et al., 2016). Black carbon absorbs sunlight very efficiently and heats the atmosphere with the top of the atmosphere forcing exceeding 1 W/m$^2$, and even though its lifetime is short compared to CO$_2$, it is still the second most important warming agent (Bond et al., 2013). Models show that its short lifetime opens up the possibilities for short-term mitigation of atmospheric warming (Shindell et al., 2012). While health effects are of interest in the context of local air pollution, climate change is investigated on regional and global scales and co-benefits from black carbon emission reductions are of great interest (Smith et al., 2009).

Measurements are conducted at locations suitable for all three different scales, with absorption filter photometers being the most common instrumentation employed to determine ambient black carbon mass concentrations.

Filter photometers measure the change of light transmitted by or reflected from the fiber filter on which particles are collected. These methods feature high sensitivity and high time resolution. The measurements are, however, prone to the filter-loading effect (FLE), a saturation of the instrumental response due to the accumulation of the sample in the filter matrix (Bond et al., 1999; Weingartner et al., 2003; Moosmüller et al., 2009). This non-linearity depends on the filter material and the type of instrument (Virkkula et al., 2007; Collaud Coen et al., 2010; Hyvärinen et al., 2013). It is important to compensate for FLE to obtain accurate measurements of black carbon concentration and for the determination of aerosol absorption spectra, which are used for source apportionment (Kirchstetter et al., 2004; Sandradewi et al., 2008).

The filter loading effect, that is the reduction of the measurement sensitivity with increasing filter loading, in Aethalometers has been described by several models (Weingartner et al., 2003; Arnott et al., 2005; Schmid et al., 2006; Virkkula et al., 2007; Park et al., 2010; Collaud Coen et al., 2010; Drinovec et al., 2015). Offline methods for the quantification of FLE are based on jumps of reported $BC$ values just before and immediately after the filter change (Weingartner et al., 2003; Virkkula et al., 2007) or $BC$ vs. ATN method (Park et al., 2010; Segura et al., 2014; Drinovec et al., 2015). These low time resolution estimations of FLE showed big differences in the effect between seasons (Virkkula et al., 2007). Uncoated soot induces a higher loading effect compared to coated soot (Weingartner et al., 2003) and Virkkula et al. (2015) showed that the loading parameter depends on the backscatter fraction and the single scattering albedo (SSA).

Traditionally, Aethalometer data have been post-processed using fixed parameters characterizing the loading effect. With the introduction of the dual-spot Aethalometer it has become possible to measure FLE with high time resolution (Drinovec et al., 2015). This allows for a much better characterization of effects governing FLE. In this paper, we are testing the hypothesis that the filter loading effect depends on the optical properties of particles present in the filter matrix, especially on black carbon particle coating. Under laboratory conditions, coating can enhance optical absorption of soot particles by up to a factor of two (Weingartner et al., 2003; Sandradewi et al., 2008).
Schnaiter et al., 2003). Ambient experiments show a smaller effect from 6%-60% (Cappa et al., 2012; Moffet and Prather, 2009; Liu et al., 2015). Determination of the mixing state using FLE may also reduce uncertainties in radiative forcing estimates (Jacobson, 2001; Shiraiwa et al., 2008). We conducted field campaigns in contrasted environments to determine the influence of sources, particle age and coating on the magnitude of the FLE.

2 Materials and Methods

2.1 Instrumentation

BC was characterized with the dual-spot Aethalometer (Magee Scientific Aethalometer model AE33) with real-time loading compensation (Drinovec et al., 2015). The inlet air stream of the AE33 is split and the sample is collected on two filter spots concurrently within the instrument. The flow through each of the two spots is different so that loading rates on the respective sample spots are different. Different loading rates cause the accumulation of the sample to be different between the two spots, resulting in the different magnitude of FLE between the spots. Measurement of FLE enables the compensation of the data – using the parametrization described in Drinovec et al. (2015), the compensation parameter $k$ can be derived. This parameter describes the linear decrease of the instrumental sensitivity with loading of the spot with the absorbing sample for each wavelength of light at which the measurement is performed. The absorption coefficients are calculated at all wavelengths and $BC$ is determined from the measurement at the wavelength 880 nm. Absorption and $BC$ are compensated using the following equation:

$$BC_{\text{compensated}}^A = \frac{BC_{\text{non-compensated}}^A}{(1 - k_A \cdot ATN)}.$$  

where $ATN$ is optical attenuation of light passing through a sample loaded filter. $k_1, k_2, k_3, k_4, k_5, k_6, k_7$ are compensation parameters determined at the following wavelengths: 370, 470, 520, 590, 660, 880 and 950 nm. The absorption Ångström exponent (AAE) (Ångström, 1929) is calculated from absorption coefficient $b$ (Drinovec et al., 2015) measured at 470 nm and 950 nm: $AAE = \ln(b_{470\text{nm}}/b_{950\text{nm}})/\ln(950/470)$. Instruments were run at 5 lpm sample flow and 1 minute time resolution.

The measurement of non-refractory submicron (NR-PM$_1$) major chemical species (i.e., organic matter, nitrate, sulphate, ammonium and chloride) was performed using Aerodyne® Aerosol Mass Spectrometer (AMS) and/or Aerosol Chemical Speciation Monitor (ACSM), fully described elsewhere (Canagaratna et al., 2007; Ng et al., 2011; Fröhlich et al., 2013). Briefly, PM$_{2.5}$ aerosols are sampled at 3 L/min and then sub-sampled at about 85 mL/min through an aerodynamic lens, focusing submicron particles onto a 600°C-heated conical tungsten vaporizer where non-refractory material are flash-vaporized and quasi-instantaneously ionized by electron impact at 70 eV. AMS mainly differs from ACSM system by allowing for size-segregated measurements of the chemical composition by using a chopper in the AMS. Many AMS also use advanced mass spectrometers with higher resolution and better detection limits. However, for the measurements of NR-PM$_1$ species in ambient air,
a good agreement is commonly observed between data from these two instruments (Crenn et al., 2015; Fröhlich et al., 2015).

Individual aerosol particles were analyzed using field emission scanning electron microscope (FEG SEM, JEOL JSM 7100F TTLS) coupled with energy dispersive X-ray (EDX) system (X-Max Large Area Analytical Silicon Drift detector, Oxford Instrument). Aluminum foils used for sampling of aerosol particles (sampling described in section 2.2) were mounted directly on the sample holder by double-sided conductive carbon tape. SEM-EDX acquisitions were performed under high vacuum at 5 keV accelerating voltage. Micrographs were acquired by secondary electron detector at 4 – 6 mm working distance (WD). Samples were first scanned under low magnification to obtain overall distribution of particles and afterwards closely examined at magnifications from 10,000 to 50,000. Representative areas were chosen for further EDX analyses. The microanalysis was performed at 10 mm WD with acquisition time of 50 to 150 s.

2.2 Measurement campaigns

The Paris campaigns were conducted at the SIRTA observatory located about 20 km south of Paris, France (48º 42’ 47” N, 2º 12’ 29” E). The observatory sits on a plateau, about 160 m above sea level, in a semi-urban environment being divided between agricultural fields, wooded areas, housing and industrial developments. The prevailing winds are from the South-West, blowing air of maritime origin over the site (e.g., Crippa et al., 2013). North-easterly winds also occur quite frequently under high pressure systems. The latter conditions are associated with the advection of continental air masses as well as of polluted air from the Paris metropolitan area. Beside continuous AE33 and ACSM measurements since 2013, results from two intensive EMEP campaigns in summer 2012 (15 June 2012 – 10 July) and winter 2013 (24 January 2013 – 16 February) were available for the present study.

The Payerne station is a rural background air quality monitoring station located in Southwestern Switzerland (46º 48’ 47” N, 6º 56’ 40” E, 489 m a.s.l.), between the Jura and the Alps. It is situated about 1 km south-east of the small rural town of Payerne (Switzerland). The site is surrounded by agricultural land (grassland and crops), forests, and small villages. Intensive campaigns took place in summer (15 June 2012 – 10 July 2012) and winter (24 January 2013 – 16 February 2013). In addition to the Aethalometer AE33, a high resolution ToF-AMS (Aerodyne Research) was operated throughout the campaigns.

The Ljubljana campaign was held between 6 August 2014 and 12 September 2014 in a suburban background location, 10 km south-west of Ljubljana, Slovenia (46º 0’ 58” N, 14º 24’ 42” E), 293 a.s.l. The site is at the edge of a village surrounded by agricultural fields and at a distance of 500 m from a highway. It lies in the Ljubljana basin, which is surrounded by hills rising from 500 to 1700 m above the valley floor. The Ljubljana campaign used four AE33 Aethalometers measuring four different ambient air streams: non-treated, dried, thermodenuded, and thermodenuded and dried samples. Ambient, dried and thermodenuded air streams had separate inlets; the thermodenuded sample was separated in two streams, one connected directly to the Aethalometer, the second guided through a silica gel drier. The Dekati thermodenuder was set to a temperature of 300 °C with 10 l/min
sample flow, which was split for the two Aethalometers. The thermodenuder consists of a heated section and an air-cooled denuder where semi-volatile compounds are adsorbed onto activated charcoal. The estimated residence time for 10 l/min flow was 0.5 s. A diffusion membrane drier (Magee Scientific, Aerosol Inlet Dryer) and a silica gel drier with dew point reductions of 11.0 ± 1.0 °C and 11.5 ± 1.0 °C, respectively, were used. Scanning Mobility Particle sizer (SMPS) with a DMA (TSI, Model 3080) connected to a water-based Condensation Particle Counter (TSI, Model 3785 UW CPC) was used to acquire particle size distribution every 5 minutes. A Dekati DLPI low-pressure impactor was used to sample particles into 13 size fractions (from 30 nm to 10 μm) for scanning electron microscopy (SEM) and energy-dispersive X-ray (EDX) spectroscopy analysis. Impactor samples were collected on aluminum foils (DEKATI, CF-300) covered by a thin layer of APIEZON L grease. Impactor stage 4 with an expected particle size range between 170 – 260 nm was analyzed.

The K-puszta campaign took place at K-puszta observatory (Hungary, 46° 58′ 12″ N, 19° 33′ E, 125 m a.s.l.) during January and February 2013. During winter ambient aerosols are dominated by wood burning emissions from household heating. The laboratory biomass burning profile was obtained during an experiment performed at the Paul Scherrer Institute (Bruns et al., 2015). Laboratory diesel exhaust was characterized at Aerosol d.o.o. in Ljubljana from a EURO 3 diesel engine. Exhaust was transferred into the smog chamber, where it was diluted and analyzed. Ljubljana roadside measurements were conducted during November 2012 in the city center, close to a heavy traffic road. Paris and Payerne data were collected during summer intensive campaigns (see above).

The Reno laboratory experiment was conducted at Desert Research Institute (Reno, Nevada, USA). Pure mustard oil was burned in a simple ceramic bowl with a cotton wick. The diluted emissions were collected and injected into a distribution chamber from which the sample was routed to multiple instruments. Ponderosa pine needles (~50 grams) were combusted in a biomass combustion chamber, a close replica of which has been described by Tian et al. (2015). The emissions from the first 35 seconds of combustion were collected in the distribution chamber and analyzed by a SMPS (TSI, DMA model 2080 with CPC model 3775) and an Aethalometer (model AE33).

2.3 Trajectory analysis

To illustrate air mass origin during specific pollution episodes during the Paris and Payerne campaigns (please see section 2.2), 72 h backtrajectories were calculated every 3 h using the PC-based version (v.4.9) of HYSPLIT (Stein et al., 2015) with weekly Global Data Assimilation System (GDAS) meteorological field data. Backtrajectories were set to end at SIRTA and Payerne at 500 m above ground level (a.g.l.).

Potential Contribution Source Function (PSCF), was used to investigate the potential advection of particulate material over large geographical scales during the different measurement campaigns. In this methodology, the probability that an air parcel may be responsible for high concentrations measured at the receptor site is calculated based on back-trajectory analyses, as described in Polissar et al. (1999). Briefly, at each ij-th grid cells, probabilities are calculated as:
\[
PSCF_{ij} = \frac{m_{ij}}{n_{ij}},
\]  

where \( n_{ij} \) represents the total number of back-trjectories passing through each \( ij \)-th cell, and \( m_{ij} \) the number of back-trajectories passing through the same cell that are associated with measured concentrations higher than a user-defined threshold. A 0.5° x 0.5° grid cell was used, covering Western Europe. Similarly to Waked et al. (2014), wet deposition was roughly estimated with precipitation data along each trajectory, assuming that even low precipitation would clean up the air parcel. For graphical purpose, results were smoothed by a Gaussian filter, and eventually normalized (see fig. 9 with colors scales from 0 to 1).

3. Results

3.1 Variability of filter loading effect (FLE)

The magnitude of the FLE for aerosols collected on a filter is dynamically determined by the dual-spot Aethalometer (Drinovec et al., 2015). The FLE depends on the entirety of the sample accumulated in the filter matrix up to the time of the measurement. The value of the FLE compensation parameter \( k \) is calculated for each of the seven measurement wavelengths, yielding a specific spectral fingerprint, which differs substantially between different measurement campaigns (Figure 1). The campaign-averaged values of the loading parameter \( k \) for short wavelengths show limited variability with a range of 0.004 – 0.005, while the variability in the near-infrared is much greater with a range of 0.001 – 0.008 (Table 1 and Figure 1).

![Figure 1: Typical spectral fingerprints of the FLE compensation parameter \( k \) obtained during ambient and laboratory campaigns. Campaign/experiment average values are presented.](image)
Table 1: Typical wavelength dependence of the filter loading compensation parameter $k$ obtained for ambient and laboratory campaigns. Campaign average $k$ and absorption Ångström exponent $AAE$ are presented.

<table>
<thead>
<tr>
<th>$AAE$</th>
<th>K-puszta winter</th>
<th>Laboratory biomass burning</th>
<th>Payerne winter</th>
<th>Laboratory diesel engine exhaust</th>
<th>Ljubljana roadside</th>
<th>Paris summer</th>
<th>Payerne summer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Filter loading parameter $k$ at the wavelength</td>
<td>370 nm</td>
<td>0.0047</td>
<td>0.0046</td>
<td>0.0044</td>
<td>0.0041</td>
<td>0.0045</td>
<td>0.0041</td>
</tr>
<tr>
<td></td>
<td>470 nm</td>
<td>0.0055</td>
<td>0.0052</td>
<td>0.0049</td>
<td>0.0045</td>
<td>0.0048</td>
<td>0.0036</td>
</tr>
<tr>
<td></td>
<td>520 nm</td>
<td>0.0059</td>
<td>0.0056</td>
<td>0.0051</td>
<td>0.0046</td>
<td>0.0050</td>
<td>0.0035</td>
</tr>
<tr>
<td></td>
<td>590 nm</td>
<td>0.0067</td>
<td>0.0060</td>
<td>0.0055</td>
<td>0.0048</td>
<td>0.0051</td>
<td>0.0034</td>
</tr>
<tr>
<td></td>
<td>660 nm</td>
<td>0.0072</td>
<td>0.0063</td>
<td>0.0059</td>
<td>0.0050</td>
<td>0.0052</td>
<td>0.0032</td>
</tr>
<tr>
<td></td>
<td>880 nm</td>
<td>0.0081</td>
<td>0.0072</td>
<td>0.0063</td>
<td>0.0055</td>
<td>0.0052</td>
<td>0.0024</td>
</tr>
<tr>
<td></td>
<td>950 nm</td>
<td>0.0084</td>
<td>0.0074</td>
<td>0.0064</td>
<td>0.0056</td>
<td>0.0053</td>
<td>0.0023</td>
</tr>
</tbody>
</table>

Table 1 also reports the average value of the absorption Ångström exponent obtained for each campaign. Occurrence of higher $k$ values appears to be linked to the latter parameter, which is commonly used as a tracer for biomass burning aerosols (Kirchstetter et al., 2004; Sandradewi et al., 2008). This is evident for both laboratory and ambient data, the most extreme case for field campaigns being K-Puszta winter data. For fresh diesel exhaust (laboratory and Ljubljana roadside measurements), obtained $k$ values fall in the middle of the parameter range, whereas lower $k$ values are obtained for summer ambient (Paris and Payerne) campaigns. This variability will be investigated in more detail in the following chapter.
3.2 Seasonal variation of filter loading effect

One-year-long black carbon mass concentrations at SIRTA show significant day-to-day variability with yearly average and standard deviation of 1068 ± 1123 ng/m$^3$ (Figure 2). Average concentrations are of 913 ± 932 ng/m$^3$ and 1263 ± 1299 ng/m$^3$ respectively for the hot (April - September) and cold (October - March) periods, respectively. The sources of black carbon impacting the measurement site vary during the year; during summer the absorption Ångström exponent ($AAE$) is close to 1, indicating the dominance of traffic emissions, while during winter its value is 1.4, suggesting the importance of biomass burning from residential heating (Kirchstetter et al., 2004; Sandradewi et al., 2008), which is consistent with previous $BC$ source apportionment studies performed in the Paris area during intensive campaigns (e.g., Favez et al., 2009; Petit et al., 2014) as well as long-term investigations (Petit et al., 2015).

![Figure 2: Black carbon mass concentration and absorption Ångström exponent $AAE$ measured at the Paris urban background site (SIRTA observatory) during 2013.](image)

Concomitantly with the absorption Ångström exponent, the parameter $k$ shows large seasonal variations (Figure 3). Our investigation is focused here on $k_6$ ($\lambda = 880$ nm), because of larger $k$ variability at longer wavelengths (Drinovec et al., 2015; Virkkula et al., 2015), and since Aethalometer measurements at 880 nm are traditionally used for the determination of black carbon mass concentrations. High values of $k_6$, between 0.006 and 0.012 are observed during winter, while in summer $k_6$ remains low, with still large oscillations (0 – 0.005), with episodes of low $k_6$ that can last for several days, indicating stronger influence of weather conditions than of daily cycles.

![Figure 3: Filter loading parameter $k$ measured at the Paris urban background site (SIRTA observatory) in 2013.](image)
Figure 4 shows monthly spectral fingerprints of the parameter $k$, which are in agreement with the offline analysis of the Aethalometer data (Virkkula et al., 2007; Virkkula et al., 2015): we measured the lowest values of the parameter $k$ during summer (red) and highest during winter (blue) months, with spring and autumn somewhere in between. Again, the variation of the parameter $k_6$ roughly correlates to the AAE, except for the months of April, May, and October, when the $k_6$ monthly mean varies between 0.0028 and 0.0041, while the AAE mean changes very little between these months (Table 2). This seems to indicate that two distinct factors - meteorology and sources – may significantly influence the value of the parameter $k$.

Table 2: Monthly averages and standard deviation of black carbon mass concentration $BC$, absorption Ångström exponent $AAE$ and compensation parameter $k_6$ for the Paris campaign (SIRTA observatory) in 2013.

<table>
<thead>
<tr>
<th>Month</th>
<th>$BC$ (ng/m$^3$)</th>
<th>$AAE$</th>
<th>$k_6$</th>
</tr>
</thead>
<tbody>
<tr>
<td>February</td>
<td>1390 ± 1320</td>
<td>1.28 ± 0.47</td>
<td>0.0059 ± 0.0023</td>
</tr>
<tr>
<td>March</td>
<td>1370 ± 1070</td>
<td>1.29 ± 0.23</td>
<td>0.0054 ± 0.0024</td>
</tr>
<tr>
<td>April</td>
<td>570 ± 640</td>
<td>1.17 ± 0.43</td>
<td>0.0041 ± 0.0016</td>
</tr>
<tr>
<td>May</td>
<td>640 ± 660</td>
<td>1.16 ± 0.46</td>
<td>0.0028 ± 0.0011</td>
</tr>
<tr>
<td>June</td>
<td>750 ± 670</td>
<td>1.10 ± 0.37</td>
<td>0.0023 ± 0.0010</td>
</tr>
<tr>
<td>July</td>
<td>1250 ± 1000</td>
<td>1.04 ± 0.31</td>
<td>0.0023 ± 0.0010</td>
</tr>
<tr>
<td>August</td>
<td>870 ± 690</td>
<td>1.09 ± 0.32</td>
<td>0.0018 ± 0.0012</td>
</tr>
<tr>
<td>September</td>
<td>1310 ± 1380</td>
<td>1.10 ± 0.29</td>
<td>0.0027 ± 0.0010</td>
</tr>
<tr>
<td>October</td>
<td>960 ± 910</td>
<td>1.18 ± 0.33</td>
<td>0.0035 ± 0.0015</td>
</tr>
<tr>
<td>November</td>
<td>1180 ± 950</td>
<td>1.25 ± 0.25</td>
<td>0.0052 ± 0.0009</td>
</tr>
<tr>
<td>December</td>
<td>1580 ± 1930</td>
<td>1.41 ± 0.26</td>
<td>0.0063 ± 0.0010</td>
</tr>
</tbody>
</table>
To analyze the influence of black carbon sources, we investigated in more detail the dependence of $k_6$ on the AAE during the heating season. The November – December period was analyzed because of stable weather conditions. The results confirm that $k_6$ increases with the AAE (Figure 5), indicating higher values for biomass burning emissions compared to traffic emissions. This is in agreement with the laboratory biomass burning experiment, where high $k_6$ values were also obtained (Figure 1). Figure 5 shows a value for $k_6$ close to 0.0055 at an absorption Ångström exponent of 1, as expected for diesel exhaust from source and ambient measurements (Table 1). We hypothesize that higher values of $k_6$ for biomass burning compared to diesel exhaust are related to the difference in black carbon agglomerate size which is investigated in more detail in the following section.

Indeed, diesel engine exhaust soot mobility diameter was reported in 30 – 100 nm range (Harris and Maricq, 2001; Keskinen and Rönkkö, 2010; Ning et al., 2013) in contrast to the 130 – 160 nm range for wood-stove emissions (Laborde et al., 2013) and 260 – 590 nm for wildfires (China et al., 2013). In special cases of large-scale sooty and turbulent fires with long flame residence time it is possible to obtain super-aggregates with geometric diameter larger than 1 $\mu$m (Chakrabarty et al., 2014).

Figure 5: The correlation between the FLE compensation parameter $k_6$ and the absorption Ångström exponent AAE for November and December 2013 in Paris (SIRTA observatory).
3.3 Particle size and the filter loading effect

To investigate the influence of black carbon particle size on $k$, we characterized emissions from a mustard oil lamp and from flaming ponderosa pine ($Pinus ponderosa$) needles combustion under laboratory conditions. Mustard oil burning produces particles with a mean mobility diameter of 210 nm (Figure 6A), which compares fairly well with the median aggregate mobility diameter of 296.8 ± 73.9 nm reported by Chakrabarty et al. (2013) for mustard oil lamp emissions. These particles are dominated by black carbon as indicated by an absorption Ångström exponent of 1 and low single scattering albedo of 0.23 (Table 3). Our Ångström exponent of 1 (470 nm, 950 nm) is somewhat lower than that determined by Chakrabarty et al. (2013, Fig. 2a) as 1.331 ± 0.004 (405 nm, 781 nm) with a three-wavelength photoacoustic instrument. Our single scattering albedo is in agreement with the lower values of 532-nm SSA measured by Chakrabarty et al. (2013, Fig. 4a) with an integrated photoacoustic-nephelometer. Chakrabarty et al. (2013) argue that higher values (up to ~0.8) of single scattering albedo shown in their Fig. 4 were caused by flickering of the flame increasing organic matter emissions; which may also have caused their increased absorption Ångström exponents. Flaming pine needle combustion produces considerably smaller particles with a mean diameter of 50 nm. We sampled only the flaming phase of Ponderosa pine needle combustion, where the emissions feature low values of the AAE (i.e., 1.25) and SSA (i.e., 0.49) as shown in Table 3. This is in agreement with the low SSA of 0.41 ± 0.14 at 532 nm that can be derived from the ratio of fuel-based emission factors of absorption and extinction cross sections measured by Chen et al. (2007, Table 1) with a photoacoustic absorption instrument and a CRD/CED extinction instrument, respectively, for flaming combustion of Ponderosa pine needles. We observed a larger filter loading effect for mustard oil lamp emissions (Figure 6a), which we believe is related to the larger particle size. This might explain the correlation between $k$ and AAE in Figure 5. The values of the loading parameter for both sources and their wavelength dependence are similar to the ambient observations in winter (Figures 1 and 4). The loading parameter values for fresh combustion products seem to be influenced by the size of the combustion products with an approximately 15% increase in the IR wavelength region for an aerosol mobility diameter increase of about 4 times. The increase of the loading parameter in the IR with size is consistent with the reported soot particle diameters for different ambient sources (Ning et al., 2013; China et al., 2013). The difference in size is accompanied by the difference in the absorption Ångström exponent and single scattering albedo. In addition to the size dependence of $k_s$, the dependence of the loading parameter on SSA we report here agrees with the one reported for Nanjing (Virkkula et al., 2015) – the decrease in SSA increases the loading parameter.
Figure 6: The filter loading parameter $k$ for laboratory measurements of emissions from mustard oil lamp and flaming Ponderosa pine needles (A). Aerosol size distributions for both sources (B).

Table 3. Filter loading parameter $k_6$, mean aerosol mobility diameter, single scattering albedo $SSA$ at 532 nm and absorption Ångström exponent $AAE$ of soot generated by mustard oil lamp and flaming Ponderosa pine needles.

<table>
<thead>
<tr>
<th></th>
<th>$k_6$</th>
<th>Particle size (nm)</th>
<th>$SSA_{532}$</th>
<th>$AAE$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mustard oil lamp</td>
<td>0.0074 ± 0.0003</td>
<td>210</td>
<td>0.222 ± 0.003</td>
<td>1.02 ± 0.02</td>
</tr>
<tr>
<td>Flaming needles</td>
<td>0.0064 ± 0.0014</td>
<td>50</td>
<td>0.43 ± 0.05</td>
<td>1.25 ± 0.03</td>
</tr>
</tbody>
</table>
3.4 Influence of the mixing state on the filter loading effect: material available for coating

The second factor influencing $k$ was assumed to be related to the meteorological conditions. To test this assumption, high time resolution variation of $k$ was investigated during EMEP campaigns taking place during summer and winter in Paris and Payerne (Figure S1). During summer, we observed periods with $k_6$ close to zero, which lasted for several days with short intervals with $k_6$ close to 0.005. During the winter campaigns, we observed higher values of $k_6$ (up to 0.01) at both locations, whereas in Paris there were also two periods with low $k_6$ values. To investigate the dependence of the loading parameter $k$ on the available coating material we combined Aethalometer data with Payerne and Paris AMS/ACSM data. Here we define the coating factor ($CF$) as a mass of the potential material available for coating (a sum of sulfate, ammonia, nitrate, and organic mass) normalized to black carbon mass:

$$CF = (SO_4^{2-} + NO_3^- + NH_4^+ + ORG)/BC.$$  

The time trend of $CF$ and $k_6$ from the Paris summer campaign shows a similar time variation pattern (Figure 7). High $CF$ coincides with low value of parameter $k_6$ and vice versa. However, there is a time lag between both measurements which is related to the way the loading parameter $k$ is determined (Drinovec et al., 2015): the parameter $k$ is a cumulative property of all particles on the filter; hence all particles on the filter, collected from the last time the tape has been advanced, contribute to the loading effect. This means that the time-lag varies with and depends on ambient black carbon concentration. The second reason is geometric smoothing employed in the algorithm, which takes the last $k$ value determined from the previous filter spot into account. The time lag impairs the $k_6$ vs $CF$ correlation resulting in higher standard deviation of $k_6$ averaged in each $CF$ bin (Figure 8).

![Figure 7: Coating factor and filter loading parameter $k_6$ time series during the Paris summer campaign. Note the inverted scale for $k_6$.](image-url)
Figure 8: Correlation between the filter loading parameter $k_6$ and the coating factor $CF$ during winter and summer campaigns in Paris and Payerne. Hourly data is averaged in $CF$ bins 2 units wide. The error bars represent standard deviation for data points inside each bin.

Table 4: Fitting parameters of the filter loading parameter $k_6$ versus the coating factor and absorption Ångström exponent $AAE$ during winter and summer campaigns in Paris and Payerne.

<table>
<thead>
<tr>
<th></th>
<th>slope ($10^{-4}$)</th>
<th>intercept at $k=0$</th>
<th>$AAE$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paris summer</td>
<td>-1.2 ± 0.1</td>
<td>27 ± 3</td>
<td>1.06</td>
</tr>
<tr>
<td>Payerne summer</td>
<td>-0.8 ± 0.5</td>
<td>35 ± 20</td>
<td>1.16</td>
</tr>
<tr>
<td>Paris winter</td>
<td>-0.8 ± 0.3</td>
<td>57 ± 22</td>
<td>1.34</td>
</tr>
<tr>
<td>Payerne winter</td>
<td>-0.2 ± 0.1</td>
<td>340 ± 180</td>
<td>1.35</td>
</tr>
</tbody>
</table>

We have observed a decrease of $k_6$ with increasing coating factor for all four campaigns. The results are most pronounced for the Paris summer campaign with the largest absolute slope of the linear regression line. In Payerne, the slope is slightly lower. In summer, $k_6$ reaches zero at coating factor values of approximately 30 for both locations (Table 4). These results show that during summer the coating material causes a reduction of the filter loading effect. In Paris, $CF$ is much larger compared to Payerne. Because most of the non-refractory material in Paris during the summer is of secondary origin (Freutel et al. 2013), it is possible to form internally mixed aerosols in the atmosphere. The correlation of particle size with coating factor in Paris supports this assertion (Figure S2).
The slope obtained for the Paris winter campaign is lower compared to summer and $k_6$ reaches zero at $CF=57$, with the dependence being more complicated than the linear one observed in summer. There is almost no correlation between $k_6$ and $CF$ during winter in Payerne. This might be a consequence of biomass burning emissions which are dominated by organics usually externally mixed with black carbon – as confirmed for Paris region during the MEGAPOLI campaign (Healy et al., 2013). The amount of black carbon and organics from biomass burning depends strongly on the combustion conditions, with black carbon being created at high temperature flaming combustion and organics at low temperature smoldering combustion (Chakrabarty et al., 2013; Pagels et al., 2013). Especially during summer, elevated amounts of secondary non-refractory material are available for coating and we observe a good correlation of $k_6$ with $CF$, which indicates that coating is responsible for reduction of filter loading effect. The decrease of $k_6$ with increasing $CF$ is at first counterintuitive if it is compared with the influence of the measured aerosol mobility diameter on $k_6$. It was observed that the increase in the mobility diameter may induce a $k_6$ increase (Figure 6), and one could speculate that the availability of material for coating black carbon cores, and hence a high value of $CF$, would increase the aerosol size and thereby increase the $k_6$ value. In fact, the contrary happens, increasing $CF$ decreases $k_6$. However, increase in $CF$ should increase SSA as well, and it has been shown for ambient aerosols in Nanjing (Virkkula et al., 2015) that the increase in SSA decreases $k_6$. To resolve this question in the following sections, we will investigate first the influence of aerosol age, and continue with removing or reducing aerosol coating.
3.5 Aerosol age and origin influence the filter loading effect

Influence of the aerosol age was indirectly investigated by using potential source contribution function (PSCF). We divided the dataset into two subsets distinguished by $k_b$ being bigger or smaller than the campaign average of 0.002. The analysis was carried out for the summer period, when the planetary boundary layer is thicker compared to the winter with stronger influence of long-range transport than in winter, and good vertical mixing to the ground, where the measurement station is positioned.

In Paris, the compensation parameter $k$ is very well correlated with aerosol age: high values indicate local sources while values close to zero indicate a regional contribution (Figure 9). For low $k$ values it seems that the sources are distributed over France, part of Great Britain, and the Atlantic Ocean. This fact may be explained by influx of potential coating material that is brought to Paris by mid- and long-range transport and mixed with more local sources of black carbon (Bressi et al., 2013; Petit et al., 2015). In Payerne, the meteorological situation is different having a smaller source area. This can be explained by the station being located between the Jura mountains and the Alps where the air is in general more stagnant compared to Paris, and is channeled from southwest to east-northwest due to the mountain ranges. Still we can observe that low $k$ measurements are related to distributed sources further from the station. At both locations, the distribution of the coating factor resembles that of the low $k_b$. This is in agreement with the correlation analysis in the chapter above, where $k_b$ decreases with $CF$ (Figure 8, A, B). A low value of $k_b$ is indicative of aged particles, and a high value of fresh, local aerosols. This is in agreement with the spectral fingerprint of fresh diesel engine emissions (Figure 1). Measurements in Paris during the MEGAPOLI campaign showed that fresh traffic BC particles are non-coated, whereas BC particles from long-range transport exhibit substantial coatings of non-refractory material (Laborde et al., 2013). On average, regional transport contained 7% of EC mass compared to 37% from local emissions (Healy et al., 2013), in accordance with our PSCF analysis of the coating factor. Theoretical studies show that coating consisting of secondary organic and inorganic species is mostly transparent in the visible part of the spectrum and can affect the absorption of light by the particles (Fuller et al., 1999; Bond et al., 2006). The fact that coating can increase absorption was confirmed in ambient studies (Wei et al., 2013b; Liu et al., 2015) and the coating may also be responsible for the decrease of parameter $k$. 
Figure 9: The potential source contribution function analysis in Paris and Payerne for measurement with low/high values of the filter loading compensation parameter $k_6$, for the coating factor ($CF$) and black carbon ($BC$) being higher than 75th percentile.
3.6 Influence of the mixing state on the filter loading effect: removing the coating of ambient aerosols

Our coating hypothesis was put to the definite test during the Ljubljana campaign; we investigated the effect of the ambient aerosol coating thickness reduction on the loading effect. The campaign was carried out in summer, when the atmospheric mixing is stronger, so we expected aged and internally mixed aerosols to be ubiquitous. We have used two approaches to reduce the coating thickness: drying and thermo-denuding the inlet stream. Drying is expected to influence particles coated with hygroscopic material; thermo-denuding was used to remove certain organic compounds (Huffman et al., 2009; Cappa et al., 2012; Liu et al., 2015), but it also removes nitrates and sulfates which decompose above 280°C (CRC, 1920). The thermo-denuder in our experiment was set to an operation temperature of 300°C. At this temperature part of low volatility oxygenated organic aerosols (LV-OOA) is not removed from the sample (Poulain et al., 2014).

Figure 10: SEM images of ambient (A) and thermo-denuded (B) impactor samples (size range 170 – 260 nm) taken during the Ljubljana campaign. The energy-dispersive X-ray spectra (D) were obtained at the selected areas on image (C). Red and yellow arrows mark soot agglomerates and secondary ammonium sulfate residue, respectively.
The de-coating efficiency of the thermo-denuder was tested using SEM analysis of the impactor samples taken during mid-day, the time when a well-mixed atmosphere is expected. Coating residues were most pronounced for above-average size particles (Figure S4); thus the impactor stage with 170 – 260 nm particle size range was selected for the analysis (Figure 10). We can observe soot agglomerates (white, red arrows), organic residue (dark patches around agglomerates), and brain-shaped dendritic residues (yellow arrows). Dark patches around soot agglomerates can be also be found on the thermo-denuded sample, indicating low volatility of these compounds. The main difference between the non-perturbed and thermo-denuded sample is the removal of the dendrite-shaped residue; similarly shaped residue was reported in different studies (Buseck and Posfai, 1999; Adachi et al., 2014) as secondary ammonium sulfate particles. To confirm the presence of secondary inorganic particles and analyze their chemical composition, we utilized energy-dispersive X-ray spectroscopy (EDX). In the background area (Figure 10 C, spectrum 1), we identified spectral peaks specific for carbon, oxygen, and aluminum, as expected for impactor aluminum foil covered with thin layer of grease. In the residue area (spectrum 2) we observed carbon and oxygen peaks with higher amplitude compared to background, and additional sulfur and nitrogen peaks. This indicates the presence of oxidized organic compounds, ammonium sulfate, and possibly nitrates, further confirming the presence of coatings and in line with findings of previous studies (Adachi et al., 2014).
The de-coating experiment consisted of four instruments sampling ambient, dried, thermo-denuded, and thermo-denuded and dried air streams simultaneously. Diurnal profiles of $k_6$ show qualitative and quantitative differences between the four treatments (Figure 11): the loading parameter $k_6$ for ambient aerosols reaches its highest value during the night and decreases during the afternoon, when the atmosphere is completely mixed. Drying increases the average value of parameter $k_6$ and amplifies the night/day variation. For the thermo-denuded samples, $k_6$ is almost constant during the day with higher average values for the thermo-denuded and dried sample stream. The average campaign value of parameter $k_6$ shows a statistically significant difference between all treatments ($p = 0.05$). The $k_6$ for thermo-denuded samples is similar to that of fresh diesel exhaust (Table 1) as expected for ambient aerosol after removal of inorganic coating by the thermo-denuder. These results are consistent with the findings of the coating factor analysis above, where coating causes reduction of $k_6$.

![Figure 11](image)

**Figure 11**: Influence of dried and/or thermo-denuded ambient aerosols on the filter loading parameter $k_6$: diurnal plot (A) and campaign averages (B). Average values of $k_6$ for different treatments are significantly different with $p = 0.05$.

Freshly emitted BC particles are hydrophobic (Weingartner et al., 1997; Zuberi et al., 2005) and show low relative humidity induced particle growth (Laborde et al., 2013). Immediately after sunrise, soot particles in the atmosphere begin to age by developing a coating of secondary species including ammonium nitrate, sulfate, and organics (Moffet and Prather, 2009). This coating makes particles hydrophilic and sensitive to relative humidity induced growth (Hu et al., 2010; Zhang et al., 2015). Drying the inlet stream during the Ljubljana campaign reduced the thickness of the hydrophilic coating and caused a large increase of $k_6$. An even greater effect was observed for the thermo-denuded sample, where most of the inorganic coatings were removed. Interestingly, we observed the effect of drying also for thermo-denuded particles. This is attributed to the presence of oxygenated organic aerosols that are less efficiently removed because of their low volatility (Huffman et al., 2009; Cappa and Jimenez, 2010; Poulain et al., 2014). For the urban environment, it was shown that most of secondary (oxygenated) organic aerosols are water-soluble (Kondo et al., 2007) and could be prone to relative humidity induced growth. This experiment further supports the hypothesis that coating is responsible for the reduction of the filter loading effect in Aethalometer.
4 Conclusions

We have tested the hypothesis that the filter loading effect present in filter-based photometers used for BC determination depends on the optical properties of particles present in the filter matrix, especially on coatings of black carbon particles. High time resolution measurements of the filter loading parameter in Aethalometer show daily and seasonal variations on the effect, which is most pronounced in the infrared region used for the determination of black carbon mass concentrations. The filter loading parameter value increases with the absorption Ångström exponent. It is suggested that this effect is related to the size of the black carbon agglomerates. The loading parameter for fresh combustion products seems to be influenced by the size of the combustion particles with an approximately 15% increase in the infrared wavelength region for an aerosol mobility diameter increase of about 4 times. On the other hand, we observed a reduction of the filter loading effect correlated with the availability of the coating material. High coating factor coincides with low value of loading parameter and vice versa. These results show that during summer the coating material causes reduction of filter loading effect. There is less or no correlation in winter, which may be a consequence of biomass burning organics being externally mixed with black carbon. To further test the coating hypothesis, a de-coating experiment using an aerosol drier and thermo-denuder was performed. De-coating the ambient aerosols increases the filter loading effect, as the coating is reduced or removed. The coating composed of ammonium sulfate and secondary organics seems to be responsible for the variation of the loading effect. The potential source contribution function analysis shows that high values of the filter loading parameter in the infrared are indicative of local pollution, whereas low values of the filter loading parameter result from ageing and coating during long range transport. Our results show that the filter loading parameter can be used as a proxy for determination of the particle mixing state, thus allowing to differentiate between local/fresh and transported/aged particles. Filter loading parameter is thus not only important for compensation of the Aethalometer absorption data but also provides additional information on the physical properties of aerosols.
Acknowledgements

The work described herein was supported by:
- EUROSTARS grant E!4825 FC Aeth
- JR-KROP grant 3211-11-000519
- JR-RK grant 3330-14-509063
- ACTRIS EU-FP7 grant n°262254
- 'PREQUALIF' project (ADEME contract n°1132C0020), DIM R2DS (AAP 2010)
- 'PARTICUL' AIR' project, and institutions CNRS, CEA and FEON.
- the National Science Foundation under Cooperative Support Agreement No. EPS-0814372
- NASA EPSCoR under Cooperative Agreement No. NNX14AN24A
- NASA ROSES under Grant No. NNX15AI48G
- National Science Foundation (NSF) under Grant No. AGS-1544425
- We thank the Slovenian Environmental Agency and AMES d.o.o. for the use of their measurement sites.

Conflict of interest declaration

L. Drinovec and G. Močnik are employed by the company Aerosol d.o.o. which develops and manufactures aerosol instrumentation including the Aethalometer AE33 and the drier used in the campaigns. H. Moosmüller is a co-inventor of the photoacoustic instruments (PAS S and PAX) manufactured by Droplet Measurement Technologies and receives licensing fees from their sale.
References


Favez, O., Cachier, H., Sciare, J., Sarda-Estève, R., and Martinon, L.: Evidence for a significant contribution of
wood burning aerosols to PM2.5 during the winter season in Paris, France, Atmos. Environ., 43, 3640–3644,

Freutel, F., Schneider, J., Drewnick, F., von der Weiden-Reinmüller, S.-L., Crippa, M., Prévôt, A. S. H.,
Stohl, A., Gros, V., Colomb, A., Michoud, V., Doussin, J. F., Borbon, A., Haefelin, M., Morille, Y.,
Beekmann, M., and Borrmann, S.: Aerosol particle measurements at three stationary sites in the megacity of
Paris during summer 2009: meteorology and air mass origin dominate aerosol particle composition and size

Fröhlich, R., Cubison, M. J., Slowik, J. G., Bukowiecki, N., Prévôt, A. S. H., Baltensperger, U., Schneider, J.,
chemical speciation monitor with TOFMS detection, Atmos. Meas. Tech. Discuss., 6, 6767–6814,

Fröhlich, R., Crenn, V., Setyan, A., Belis, C. A., Canonaco, F., Favez, O., Riffault, V., Slowik, J. G., Aas, W.,
Aijala, M., Alastuey, A., Artinano, B., Bonnaire, N., Bozzetti, C., Bressi, M., Carbone, C., Coz, E., Croteau,
P. L., Cubison, M. J., Esser-Gietl, J. K., Green, D. C., Gros, V., Heikkinen, L., Herrmann, H., Jayne, J. T.,
Lunder, C. R., Mingguillon, M. C., Mocnik, G., O'Dowd, C. D., Ovadnevaite, J., Petralia, E., Poulain, L.,
results from 15 individual, co-located aerosol mass spectrometers, Atmospheric Measurement Techniques,

Fuller, K. A., Malm, W. C., and Kreidenweis, S. M.: Effects of mixing on extinction by carbonaceous particles,

Grahame, T. J., Klemm, R., and Schlesinger, R. B.: Public health and components of particulate matter: The
changing assessment of black carbon, J. Air Waste Manage., 64, 620–660,

Harris S. J. and Maricq M. M.: Signature size distributions for diesel and gasoline engine exhaust particulate

Estève, R., McGuire, M. L., Jeong, C.-H., McGillicuddy, E., O’Connor, I. P., Sodeau, J. R., Evans, G. J., and
mass spectrometer and aerosol mass spectrometer measurements, Atmos. Chem. Phys., 13, 9479-9496,

Hu, D., Qiao, L., Chen, J., Ye, X., Yang, X., Cheng, T., and Fang, W.: Hygroscopicity of inorganic aerosols:

Huffman, J. A., Docherty, K. S., Aiken, A.C., Cubison, M. J., Ulbrich, I. M., DeCarlo, P. F., Sueper, D., Jayne,
J. T., Worsnop, D. R., Zieann, P. J., and Jimenez, J. L.: Chemically-resolved aerosol volatility measurements


Schmid, O., Artaxo, P., Arnott, W. P., Chand, D., Gatti, L. V., Frank, G. P., Hoffer, A., Schnaiter, M., and Andreae, M. O.: Spectral light absorption by ambient aerosols influenced by biomass burning in the Amazon...


