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Backman, John

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On Aethalometer measurement uncertainties and an instrument correction factor for the Arctic

John Backman1, Lauren Schmeisser2,a, Aki Virkkula1,3,4, John A. Ogren2,5, Eija Asmi1, Sandra Starkweather2,5, Sangeeta Sharma6, Konstantinos Eleftheriadis7, Taneil Uttal5, Anne Jefferson2, Michael Bergin8, Alexander Makshtas9, Peter Tunved10, and Markus Fiebig11

1Finnish Meteorological Institute, Atmospheric Composition Research, Helsinki, Finland
2University of Colorado Boulder, Cooperative Institute for Research in Environmental Sciences, Boulder, USA
3University of Helsinki, Department of Physics, Helsinki, Finland
4Nanjing University, Joint International Research Laboratory of Atmospheric and Earth System Sciences, Nanjing, China
5National Oceanic and Atmospheric Administration, Earth System Research Laboratory, Boulder, USA
6Environment and Climate Change Canada, Climate Research Division, Downsview, Canada
7Institute of Nuclear and Radiological Science and Technology, Energy and Safety, Environmental Radioactivity Laboratory, NCSR “Demokritos”, Athens, Greece
8Duke University, Civil and Environmental Engineering, Durham, USA
9Russian Federal Service for Hydrometeorology and Environmental Monitoring, Arctic and Antarctic Research Institute, St. Petersburg, Russia
10Stockholm University, Department of Environmental Science and Analytical Chemistry, Stockholm, Sweden
11NILU – Norsk institutt for luftforskning, Dept. Atmospheric and Climate Research (ATMOS), Kjeller, Norway
a now at: University of Washington, Department of Atmospheric Sciences, Seattle, USA

Correspondence: John Backman (john.backman@fmi.fi)

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Abstract. Several types of filter-based instruments are used to estimate aerosol light absorption coefficients. Two significant results are presented based on Aethalometer measurements at six Arctic stations from 2012 to 2014. First, an alternative method of post-processing the Aethalometer data is presented, which reduces measurement noise and lowers the detection limit of the instrument more effectively than boxcar averaging. The biggest benefit of this approach can be achieved if instrument drift is minimised. Moreover, by using an attenuation threshold criterion for data post-processing, the relative uncertainty from the electronic noise of the instrument is kept constant. This approach results in a time series with a variable collection time (Δt) but with a constant relative uncertainty with regard to electronic noise in the instrument. An additional advantage of this method is that the detection limit of the instrument will be lowered at small aerosol concentrations at the expense of temporal resolution, whereas there is little to no loss in temporal resolution at high aerosol concentrations (> 2.1–6.7 Mm⁻¹ as measured by the Aethalometers). At high aerosol concentrations, minimising the detection limit of the instrument is less critical. Additionally, utilising co-located filter-based absorption photometers, a correction factor is presented for the Arctic that can be used in Aethalometer corrections available in literature. The correction factor of 3.45 was calculated for low-elevation Arctic stations. This correction factor harmonises Aethalometer attenuation coefficients with light absorption coefficients as measured by the co-located light absorption photometers. Using one correction factor for Arctic Aethalometers has the advantage that measurements between stations become more inter-comparable.
1 Introduction

Black carbon (BC) and soot, which originate from incomplete combustion, are particularly potent absorbers of solar radiation and comprise a complex part of the climate system (Bond et al., 2013). Light-absorbing particles, including BC and soot, influence the aerosol radiative forcing (ARF) by warming the atmosphere, changing the aerosol single-scattering albedo, and potentially altering cloud droplet evaporation and lifetime (Koch and Del Genio, 2010). In addition, trace amounts of absorbing particles deposited on snow can perturb snow grain size and thus lower the snow albedo (Hadley and Kirchstetter, 2012; Wiscombe and Warren, 1980a, b); a low albedo favours melting. Polar regions are particularly sensitive to changes in surface albedo, which subsequently impacts sea ice, snow cover, and ultimately surface temperature (Holland and Bitz, 2003; Serreze and Barry, 2011; Serreze et al., 2009). This polar amplification results in enhanced ice melt and more open water (Johannessen et al., 2004; Serreze et al., 2009). Brown carbon (BrC) absorbs sunlight primarily in the ultraviolet–visible region of the solar spectrum (Andreae and Gelencsér, 2006; Bergstrom et al., 2007), whereas the BC absorption efficiency is relatively uniform across the UV to near-infrared solar spectrum.

Given that BC is a particularly potent perturbing agent, in situ measurements of BC are important. A widely used technique to measure light absorption by aerosol particles is with filter-based absorption instruments such as the Aethalometer (e.g. Weingartner et al., 2003), the particle soot absorption photometer (PSAP; Bond et al., 1999; Virkkula et al., 2005), and the multi-angle absorption photometer (MAAP; Petzold and Schönlinner, 2004; Petzold et al., 2005). These instruments report either equivalent black carbon (eBC) mass concentrations or light absorption coefficients (Petzold et al., 2013).

The high variability of eBC, particularly in polar, high-altitude, and coastal regions, makes measurements with Aethalometers challenging. During clean periods, the eBC concentrations can easily be below the detection limit of the instrument. Data treatment methods such as boxcar averaging can improve the detection limit of the instrument.

An alternative method to reduce noise in Aethalometers has been proposed (Hagler et al., 2011). In this work, a criterion from Hagler et al. (2011) is used; an attenuation change ($\Delta$ATN) threshold needs to be exceeded for post-processing calculations to be invoked. Instead of using this one criterion for boxcar averaging intervals, $\Delta$ATN is used in the post-processing calculations using the Aethalometer equation. Here we explore this alternative method from a measurement uncertainty perspective and show that a constant relative uncertainty can be achieved using this one criterion for data post-processing. The result is a time series with a time resolution which is adapted to the measured aerosol concentration. The best performance of this method is achieved when drift in the Aethalometer is at a minimum.

While it is well known that Aethalometer measurements require some form of post-processing (Arnott et al., 2005; Collaud Coen et al., 2010; Schmid et al., 2006; Virkkula et al., 2007; Weingartner et al., 2003), the purpose of this paper is not to add a correction algorithm to the literature but to show how to reduce noise in Aethalometer measurements more effectively. This paper uses data from Arctic sites, regions with low signal and high susceptibility to ARF from eBC, to examine noise reduction in the Aethalometer signal. Aethalometer instruments have been used to make measurements in the Arctic since the 1980s (e.g. Bodhaine, 1995; Sharma et al., 2006, 2013).

Using the adaptive collection time method of data collection we present an Arctic correction factor ($C_f$) value to harmonise Aethalometer absorption measurements to other filter-based light absorption photometers. This correction factor can be used in existing Aethalometer correction schemes available in literature.

2 Measurements and instruments

The data used in this study comprise 3 years of measurements (2012–2014) at six Arctic stations. The actual eBC climatology of the stations will be presented in a following paper. Below we provide information about station location, operations, and environs, as well as the instruments deployed at each site. Each site has at least an Aethalometer and an additional filter-based absorption photometer. The instruments are summarised in Table 1.

2.1 Measurement sites

2.1.1 Barrow

The Barrow observatory is located on the northernmost coast of Alaska, just 5 km north-east of the town of Barrow, Alaska (population ~ 4200), and 2 km from the Arctic Ocean coast, at an elevation of 11 m above sea level (a.s.l.) and at coordinates 71.323°N and 156.612°W. The site is primarily influenced by regional air masses originating from the Beaufort Sea, though the station also measures pollution coming from the nearby town. All air masses originating from the direction of the town are marked as contaminated, and those data are not used in this analysis.

A 7-wavelength Magee AE31 Aethalometer has been operating at the station since 2010. The co-located light absorption instrument is the continuous light absorption photometer (CLAP; Ogren et al., 2017) that has been collecting aerosol absorption data since 2011 and was built by National Oceanic and Atmospheric Administration (NOAA). Previous descriptions of the aerosol optical property climatology at Barrow can be found in Bodhaine (1983, 1995) and Delene and Ogren (2002).
from 2003 to present. The co-located light absorption pho-
with a 1-wavelength (880 nm) Magee AE16 Aethalometer
Equivalent black carbon measurements here have been made
scientific operations at the Summit site that necessitate activ-
University in collaboration with the NOAA Earth Systems
est in elevation of the six sites. Measurements of equivalent
72.580
The monitoring station at Summit, Greenland, is located at
2.1.3 Summit
Table 1. Overview of instruments at the respective measurement stations. The flow rates are mean values in standard litres per minute (slpm). TOT means that it is a total aerosol inlet with no cutoff size. Filter change settings reported as either ATN or filter transmittance (Tr).

<table>
<thead>
<tr>
<th>Site</th>
<th>Instruments</th>
<th>Model</th>
<th>Wavelengths (nm)</th>
<th>Inlet</th>
<th>Flow rate (slpm)</th>
<th>Filter change at</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alert</td>
<td>Aethalometer</td>
<td>AE31</td>
<td>370, 470, 520, 590, 660, 880, 950</td>
<td>TOT</td>
<td>3.9</td>
<td>ATN = 75</td>
</tr>
<tr>
<td></td>
<td>PSAP</td>
<td>RR²</td>
<td>3λ</td>
<td>PM₁₀</td>
<td>1.1</td>
<td>Tr = 0.7</td>
</tr>
<tr>
<td>Summit</td>
<td>Aethalometer</td>
<td>AE16</td>
<td>880</td>
<td>PM₂₅</td>
<td>6.3</td>
<td>ATN = 75</td>
</tr>
<tr>
<td></td>
<td>CLAP</td>
<td>3λ</td>
<td>467, 528, 652</td>
<td>PM₂₅</td>
<td>0.6</td>
<td>Tr = 0.7</td>
</tr>
<tr>
<td>Barrow</td>
<td>Aethalometer</td>
<td>AE31</td>
<td>370, 470, 520, 590, 660, 880, 950</td>
<td>PM₁₀</td>
<td>4.8</td>
<td>ATN = 99</td>
</tr>
<tr>
<td></td>
<td>CLAP</td>
<td>3λ</td>
<td>467, 528, 652</td>
<td>PM₁₀</td>
<td>1.3</td>
<td>Tr = 0.7</td>
</tr>
<tr>
<td>Tiksi</td>
<td>Aethalometer</td>
<td>AE31</td>
<td>370, 470, 520, 590, 660, 880, 950</td>
<td>PM₁₀</td>
<td>4.9</td>
<td>ATN = 75</td>
</tr>
<tr>
<td></td>
<td>MAAP</td>
<td>5012</td>
<td>637°</td>
<td>PM₁₀</td>
<td>5.6</td>
<td>ATN = 20</td>
</tr>
<tr>
<td>Pallas</td>
<td>Aethalometer</td>
<td>AE31</td>
<td>370, 470, 520, 590, 660, 880, 950</td>
<td>TOT</td>
<td>2.9/4.0</td>
<td>8 h</td>
</tr>
<tr>
<td></td>
<td>MAAP</td>
<td>5012</td>
<td>637°</td>
<td>PM₂₅</td>
<td>8.7</td>
<td>ATN = 50</td>
</tr>
<tr>
<td>Zeppelin</td>
<td>Aethalometer</td>
<td>AE31</td>
<td>370, 470, 520, 590, 660, 880, 950</td>
<td>TOT</td>
<td>7.5</td>
<td>ATN = 75</td>
</tr>
<tr>
<td></td>
<td>PSAP</td>
<td>1λ</td>
<td>525</td>
<td>TOT</td>
<td>0.9</td>
<td>Tr = 0.7</td>
</tr>
</tbody>
</table>

* Radiance research; b Custom-built 1 λ PSAP; c Müller et al. (2011)

2.1.2 Alert

Alert is located in Nunavut, Canada, 12 km west of Cape Sheridan, at 82.492° N and 62.508° W and at an elevation of 8 m a.s.l. The monitoring station is operated by Environment and Climate Change Canada. Alert is the northernmost site of those analysed here, located just 817 km from the North Pole. Given the remote location, the aerosols there are not heavily influenced by human populations. The site is near the coast, which is ice covered in the winter but turns to open ocean during summer. A 7-wavelength Magee AE31 Aethalometer has been running at Alert from 2008 to present. Co-located light absorption measurements were made with a 3-wavelength PSAP from 2007 to present. More information on black carbon measurements at Alert can be found in Sharma et al. (2002).

2.1.3 Summit

The monitoring station at Summit, Greenland, is located at 72.580° N and 38.480° W, and at 3216 m a.s.l., is the highest in elevation of the six sites. Measurements of equivalent black carbon at Summit are supported and operated by Duke University in collaboration with the NOAA Earth Systems Research Laboratory. Although there are many established scientific operations at the Summit site that necessitate activities that produce anthropogenic aerosols, the site is generally very remote and measures very low aerosol concentrations. Equivalent black carbon measurements here have been made with a 1-wavelength (880 nm) Magee AE16 Aethalometer from 2003 to present. The co-located light absorption pho-
tometer at Summit is a multi-wavelength CLAP, running at the site from 2011 to present.

2.1.4 Zeppelin

The Zeppelin Mountain observatory is located at 475 m a.s.l. near the small research village of Ny-Ålesund on the island of Svalbard at 78.907° N and 11.889° E. The monitoring station is owned by the Norwegian Polar Institute and operated by the Norwegian Institute for Air Research, and the most recent version of the station building was established in the year 2000. The site is typically located above the inversion layer and thus measures air masses with minimal contamination. The observatory has long-term measurements of equivalent black carbon with Magee Aethalometers, namely AE9 from 1998 to 1999 and AE31 from 2001 to present (Eleftheriadis et al., 2009), and co-located light absorption measurements with a 1-wavelength PSAP.

2.1.5 Pallas

The Pallas measurement station is located in the Finnish Arctic in the Municipality of Muonio. The measurement station is operated by the Finnish Meteorological Institute. The main measurement building housing the instruments used in this study is located on top of the Sammaltunturi fell. The top of the fell is at an altitude of 565 m a.s.l. and above the tree line. The coordinates of the station are 67.973° N, 24.116° E. There are no major local sources close to the station, and the surrounding terrain is forested, consisting of pine, spruce, and birch trees in addition to barren fells.
The 7-wavelength Magee AE31 Aethalometer is connected to the total aerosol inlet which is heated in order to lower the relative humidity (RH) and causes cloud drops to evaporate. The co-located light absorption photometer is a MAAP (Thermo Scientific, model 5012). The MAAP is connected to a heated PM$_{2.5}$ inlet to lower the relative humidity. The different size cuts of the instruments could bias the Aethalometer towards higher absorption coefficients than the MAAP. A more thorough description of the site is provided by Hatakka et al. (2003).

2.1.6 Tiksi

The Tiksi measurement station is located in northern Siberia in Russia. The station is located 500 m from the coast of the Laptev Sea at an altitude of 30 m a.s.l. at 71.596° N, 128.889° E. The site is surrounded by tundra. The station is a cooperation between the Russian Federation’s Roshydromet, the US National Oceanic and Atmospheric Administration, the US National Science Foundation, and the Finnish Meteorological Institute. The station is located ≈ 4 km south of the town of Tiksi, which comprises the sole source of local air pollution. The data were screened using local wind direction and aerosol size distribution data to omit local pollution from the town (Asmi et al., 2016).

The measurement instruments used in this study consist of a 7-wavelength Aethalometer (Magee model AE31) and a MAAP (model 5012). The instruments are connected to a PM$_{10}$ inlet with self-regulating heating to avoid the build-up of ice on the inlet. By raising the temperature of the sample air to room temperature, the sample RH is kept below 30% (Asmi et al., 2015).

2.2 Data processing

2.2.1 The Aethalometer

The Aethalometer theory of operation relies on the measurement of light transmitted through a fibre filter as aerosol particles are collected on the filter. The filter is illuminated by a light source from one side with the detectors located on the other side of the filter. Light is transmitted through a pristine part of the filter with an intensity $I_0$. The light that traverses through the part where aerosol particle deposit is transmitted with an intensity $I$. The Aethalometer calculates, and reports, filter attenuation (ATN) as described in Eq. (1) (e.g. Weingartner et al., 2003).

$$\text{ATN} = -100\ln \left( \frac{I}{I_0} \right) \quad (1)$$

The term $I/I_0$ represents the transmission of light through the filter and is referred to as the filter transmittance. The factor of 100 in Eq. (1) is there for numerical convenience and will for this reason also be included throughout this work. The attenuation coefficient ($\sigma_0$) can be written in the form (e.g. Weingartner et al., 2003)

$$\sigma_0 = \frac{A}{Q \Delta I} \frac{\Delta \text{ATN}}{100} \quad (2)$$

In Eq. (2), $A$ is the filter spot size area, $Q$ is the sample flow rate, and $\Delta I$ is the time between the light intensity measurements. The term $\Delta \text{ATN}$ is the change in ATN over the time $\Delta t$, which is here called the collection time. When the fibre filter is loaded with aerosol, and the transmission of light through the filter has dropped too much, the filter spot needs to be changed. In the Aethalometer, filter changes can be set to occur automatically at an ATN value set by the operator. Alternatively, the filter can be set to change after a given time.

Although the Aethalometer actually measures $\sigma_0$, the instrument output is equivalent black carbon mass concentration (Petzold et al., 2013). The conversion from $\sigma_0$ to eBC is done using a wavelength-dependent mass attenuation cross section (MAC$_{AE}$) of 16.62 m$^2$g$^{-1}$ (at 880 nm), scaling inversely with wavelength (e.g. Arnott et al., 2005).

The firmware of the Aethalometer uses an internal collection time $\Delta t$ which is $2 \leq \Delta t \leq 5$ min. This is the inner data processing cycle of the AE31 Aethalometer. Any longer averaging times set by the operator will commence an outer cycle, which will average the readings obtained during the inner cycle. Therefore, the averaging time ($t_{\text{avg}}$) that can be set for the instrument by the operator is restricted to multiples of 5 min. In other words, the output of the outer cycle is an average of the inner cycle with a $\Delta t$ of 5 min. This is not always ideal since at very pristine sites a collection time of 5 min is not long enough, resulting in noisy data.

Choosing a longer averaging time (the so-called outer cycle) will reduce noise and, therefore, the detection limit of the instrument, at a rate of $t_{\text{avg}}^{-0.5}$. Increasing $t_{\text{avg}}$, however, results in a reduction of temporal resolution. Moreover, when $t_{\text{avg}} > \Delta t$ the instrument output can no longer be reproduced using Eq. (2) since the data that comprise the inner cycle are no longer reported by the instrument. Thus, the greatest versatility of post-processing can be achieved when $t_{\text{avg}}$ is equal to $\Delta t$; i.e. $t_{\text{avg}} \leq 5$ min for the Aethalometers in this study.

One can circumvent the outer cycle by data post-processing and achieve a lower detection limit. Included in the standard long-format output of the AE31 and AE16 Aethalometer models are the ATN values at the end of the averaging period, along with the aerosol flow rate. Thus, the standard output data can be used to post-process the data using Eq. (2) for an arbitrary value of $\Delta t$; i.e. an arbitrary collection time. The term $\Delta \text{ATN}$ is then simply the change in ATN from the time $t$ to $t + \Delta t$; i.e. $\Delta \text{ATN} = \text{ATN}_{t + \Delta t} - \text{ATN}_t$.

The benefit of this post-processing approach is that it reduces noise better than the boxcar averaging of the firmware. This is discussed and shown further on. This approach for reducing noise in Aethalometer measurements was originally suggested by Hagler et al. (2011). In their work, a $\Delta \text{ATN}$
change was used as a criterion for boxcar averaging, whereas, here, ΔATN is used in the calculations. Previous work using a PSAP has shown that the collection time approach can greatly reduce the noise of filter attenuation measurements (Springston and Sedlacek, 2007) to produce a time series with an adaptive collection time (Hagler et al., 2007). In this work, the method is elaborated on using uncertainty analysis, specifically for Aethalometers.

2.3 Arctic Aethalometer correction factor

The actual aerosol light absorption coefficient of the initially suspended particles is not \( \sigma_0 \). When aerosol particles deposit onto a filter, they will inevitably interact with the filter. The realisation of this has resulted in a variety of different data-processing correction schemes for the Aethalometer and the PSAP (Arnott et al., 2005; Bond et al., 1999; Collaud Coen et al., 2010; Schmid et al., 2006; Virkkula et al., 2005, 2007; Weingartner et al., 2003). The purpose of these corrections is to derive the actual light absorption coefficient (\( \sigma_{ap} \)) of the suspended particles devoid of filter-induced artefacts.

The Arctic measurement sites included in this study all have the same type of measurement instrument, namely the Aethalometer model AE31, except for Summit, which is an AE16 model Aethalometer. The benefit of having the same type and model of instrument is that measurement artefacts for the same type of instrument would be expected to be more similar than between different types of instruments. The comparison of aerosol properties between different sites should be more robust when all sites have the same type of instrument than if the instruments would differ from site to site. Both the AE31 and AE16 model use the same type of filter (Pallflex Q250F). However, it still has to be acknowledged that artefacts can differ between different stations depending on the difference in aerosol properties even though the same type of instrument is used.

For the sake of inter-comparability, a relative normalisation factor is introduced to harmonise the determination of the absorption coefficient at the Arctic stations. The normalisation factor is calculated as

\[
C_f = \frac{\sigma_0}{\sigma_{ap}},
\]

where \( \sigma_{ap} \) is the light absorption coefficient as measured by co-located filter-based absorption measurements. The co-located instruments are discussed in more detail in the next section. The interpretation of the correction factor is in essence how much greater the attenuation coefficient is in comparison to the light absorption coefficient of the co-located filter-based absorption photometers which have been corrected for loading and scattering artefacts. Thus, \( C_f \) will effectively be an inter-instrument comparison artefact.

This correction factor can be used in many of the available correction algorithms in the place of the multiple scattering correction factor (\( C_{ref} \)); see Collaud Coen et al. (2010) Table 2 for a list. However, the \( C_f \) values reported here are not multiple scattering correction factors because no true reference absorption measurements were available. The purpose of the \( C_f \) values here is to provide a general value that can be used in place of the \( C_{ref} \) value for the Arctic, in order to harmonise the determination of \( \sigma_{ap} \) from Aethalometers in the Arctic with other methods for determining aerosol light absorption coefficients.

2.4 Co-located filter-based absorption instruments

The Arctic sites in this study were chosen based on the criterion that they all have Aethalometers and an additional co-located filter-based photometer measuring aerosol light absorption coefficients. The additional instrument is either a MAAP, PSAP, or CLAP. These instruments will provide the \( \sigma_{ap} \) that is needed to calculate \( C_f \) for the sites, using Eq. (3). The MAAP is a filter-based absorption instrument that, in addition to transmittance measurements through the filter, also measures the back-scattered light at two angles (Petzold and Schönlinner, 2004). This allows for a radiative transfer scheme to be applied since the back-scattered light at multiple angles can be used to distinguish between diffusely scattered light and Gaussian scattered light. This information is then used to calculate the diffuse fraction of light scattered back by the filter in order to account for multiple scattering and apparent absorption effects by solving the radiative transfer equation from the measurements on the filter.

The PSAP and CLAP instruments measure transmission, and therefore are based on Eqs. (1) and (2). Both instruments use the same type of filter, and the optical design of the CLAP is very similar to the PSAP. The CLAP differs from the PSAP in that, instead of a single sample spot on a 10 mm filter, it has eight sample spots on a 47 mm filter. Solenoid valves are used to switch to the next sample spot once the filter transmittance reaches 0.7. Thus, the CLAP can run 8 times as long as the PSAP before requiring a filter change, which is ideal for remote sites that are not visited daily.

Both the PSAPs and CLAPs use the same type of Pallflex E70-2075W filters, with the only difference being their size. As the optical designs of the two instruments are very similar, both the PSAP and CLAP data used in this study were corrected using the Bond et al. (1999) correction along with the Ogren (2010) wavelength adjustment. It has been shown before that the same type of filter and a similar optical design yields very similar results (Miyazaki et al., 2008; Nakayama et al., 2010). The Bond et al. (1999) correction includes a multiple scattering correction, a filter loading correction, and an apparent absorption correction. The apparent absorption correction makes use of light scattering coefficients (e.g. from nephelometers). At all sites in this paper where light scattering coefficients were needed to correct the PSAP and CLAP, the light scattering was measured by TSI nephelometers (TSI Inc, model 3563; Anderson and Ogren, 1998).

Although the co-located instruments are based on collecting the sample aerosol on filters, there are differences. For
Table 2. Standard deviations of $\delta \Delta{\text{ATN}}_{\text{ND}}$ for the different Aethalometers and their measurement wavelengths. The filtered air noise measurements consist of at least 24 h of data, except for Alert where data was comprised of a few hours of measurements totalling 4 days. The standard deviation was calculated from subsequent reported ATN values as such and can therefore be used to reproduce Fig. 4. The $t_{\text{avg}}$ column shows the instrument setting for the outer cycle of the instrument during the time of the noise measurements.

<table>
<thead>
<tr>
<th></th>
<th>$N$</th>
<th>$t_{\text{avg}}$</th>
<th>370</th>
<th>470</th>
<th>520</th>
<th>590</th>
<th>660</th>
<th>880</th>
<th>950</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alert</td>
<td>1267</td>
<td>5</td>
<td>0.011</td>
<td>0.010</td>
<td>0.011</td>
<td>0.010</td>
<td>0.011</td>
<td>0.008</td>
<td>0.008</td>
</tr>
<tr>
<td>Summit</td>
<td>235</td>
<td>5</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.016</td>
<td></td>
</tr>
<tr>
<td>Barrow</td>
<td>745</td>
<td>5</td>
<td>0.016</td>
<td>0.016</td>
<td>0.015</td>
<td>0.015</td>
<td>0.015</td>
<td>0.015</td>
<td>0.015</td>
</tr>
<tr>
<td>Tiksi</td>
<td>316</td>
<td>5</td>
<td>0.007</td>
<td>0.008</td>
<td>0.006</td>
<td>0.007</td>
<td>0.007</td>
<td>0.074</td>
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<tr>
<td>Pallas</td>
<td>290</td>
<td>5</td>
<td>0.003</td>
<td>0.004</td>
<td>0.003</td>
<td>0.004</td>
<td>0.003</td>
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<td>0.004</td>
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<tr>
<td>Zeppelin</td>
<td>48</td>
<td>30</td>
<td>0.028</td>
<td>0.012</td>
<td>0.010</td>
<td>0.031</td>
<td>0.016</td>
<td>0.012</td>
<td>0.014</td>
</tr>
</tbody>
</table>

Table 3. Standard deviation of $\sigma_0 (\delta \sigma_0)$ in Mm$^{-1}$ when measuring particle-free air. These values can be used in Eq. (10) for an arbitrary value of $\sigma_0$ and $\Delta{t}$.

<table>
<thead>
<tr>
<th>$\Delta{t}$</th>
<th>370 nm</th>
<th>470 nm</th>
<th>520 nm</th>
<th>590 nm</th>
<th>660 nm</th>
<th>880 nm</th>
<th>950 nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alert</td>
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<td>0.251</td>
<td>0.286</td>
<td>0.253</td>
<td>0.282</td>
<td>0.215</td>
</tr>
<tr>
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<td>0.325</td>
<td>0.316</td>
<td>0.312</td>
<td>0.322</td>
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</tr>
<tr>
<td>Barrow</td>
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<td>0.155</td>
<td>0.117</td>
<td>0.129</td>
<td>0.151</td>
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</tr>
<tr>
<td>Tiksi</td>
<td>5</td>
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<td>0.144</td>
<td>0.149</td>
<td>0.111</td>
<td>0.114</td>
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<tr>
<td>Pallas</td>
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<td>0.026</td>
<td>0.021</td>
<td>0.065</td>
<td>0.032</td>
<td>0.024</td>
</tr>
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</table>

Aerosol particles that have a high single-scattering albedo ($\omega_0$), defined as the ratio of light scattering ($\sigma_{\text{sp}}$) to light extinction ($\sigma_{\text{ep}}$), the MAAP has been shown to be less prone to interpret light scattering as light absorption than a PSAP or an Aethalometer (Petzold et al., 2005). In the Arctic, this is an advantage because of the high $\omega_0$ of the aerosol. Research has also shown a good agreement between the MAAP and independent reference absorption ($\sigma_{\text{sp}}$-$\sigma_{\text{ep}}$) measurements for $\omega_0$ values in the range of 0.7–0.98 (Sheridan et al., 2005). The same study also showed that the PSAP, with the Bond correction applied, agrees better with independent reference absorption measurements for atmospherically relevant $\omega_0$ values (0.88) than for very dark aerosol ($\omega_0$ = 0.30). At a $\omega_0$ of 0.88, there was virtually no dependence of filter loading on $\sigma_{\text{ap}}$. MAAP-, PSAP-, and CLAP-derived $\sigma_{\text{ap}}$ will be cross sensitive to $\sigma_{\text{sp}}$ for a purely scattering aerosol, and the extent of the cross sensitivity depends on the loading of the filter (Müller et al., 2011).

The unit to unit variability of the MAAP is comparatively lower than for the other absorption photometer instruments used in this study (Müller et al., 2011). The same study also showed that the instrument noise of the PSAP and the MAAP are lower than for Aethalometers. The design of the PSAP and CLAP instruments should also make the measured flow through the instruments less uncertain than in instruments using a filter tape roll because the filters are sealed in place inside the instrument.

It should be noted that none of the filter changes for any of the instruments can be considered to be synchronised with each other; e.g. the PSAP filter is not changed at the same time as an Aethalometer tape advance. Thus, when comparing a reference instrument to an Aethalometer, using the whole time series, any remaining cross sensitivity to the state of the filter on a reference instrument will represent the mean or median bias.

Because the reference instruments operate at different wavelengths than the Aethalometers, Ångström exponents ($\alpha$) were used to interpolate or extrapolate data to a matching wavelength; $\alpha$ were also used to match nephelometer wavelengths to reference absorption wavelengths when using the correction schemes. The Ångström exponent was calculated as follows:

$$\alpha = -\frac{\log(\sigma_1) - \log(\sigma_2)}{\log(\lambda_1) - \log(\lambda_2)},$$

(4)

where $\sigma_1$ and $\sigma_2$ represent absorption or scattering coefficients at their respective wavelengths $\lambda_1$ and $\lambda_2$. Using $\alpha$, the absorption coefficient ($\sigma_x$) can be calculated for a desired wavelength $\lambda_x$ using

$$\sigma_x = \sigma_1 \left(\frac{\lambda_1}{\lambda_x}\right)^\alpha.$$
Using Eqs. (4) and (5), the Aethalometer data was interpolated to the wavelengths 467, 525, and 637 nm. The reference absorption instruments that did not already measure at these wavelengths were also interpolated to these three wavelengths. The 1-wavelength Aethalometer at Summit was interpolated from 880 to 637 nm using an $\alpha$ of 0.814 in Eq. (5).

3 Aethalometer uncertainty analysis

In order to investigate how the collection time approach can improve the Aethalometer measurements, the measurement uncertainties must be known. By applying the equation for the propagation of uncertainty for uncorrelated variables

$$\delta \sigma_0 = \sqrt{\sum_{i=1}^{n} \left( \frac{\partial \sigma_0}{\partial x_i} \right)^2 \delta x_i^2} \tag{6}$$

to Eq. (2), the relative uncertainty of the measurements can be solved. In Eq. (6), $x_i$ represents the independent variables $-\Delta \text{ATN}$, $A$, $Q$, and $\Delta t$ of Eq. (2) – and $\delta x_i$ represents their uncertainties.

However, the uncertainty in $\Delta \text{ATN}$ has more than one component. Therefore, prior to applying Eq. (6) to Eq. (2), the term $\Delta \text{ATN}$ is decomposed into two components. The first component is the true change in $\Delta \text{ATN}$ that contains no drift, here denoted as $\Delta \text{ATN}_{\text{ND}}$. The second component that contributes to the uncertainty in $\Delta \text{ATN}$ is drift, here denoted as $\Delta \text{ATN}_D$. Furthermore, drift can be expressed as a rate of change over the time $\Delta t$ as $k_D = \Delta \text{ATN}_D / \Delta t$. The influence of drift for an arbitrary $\Delta t$ then becomes $k_D \Delta t$. Thus, $\Delta \text{ATN}$ has been decomposed into $\Delta \text{ATN} = \Delta \text{ATN}_{\text{ND}} + k_D \Delta t$. Substituting the total change in $\Delta \text{ATN}$ with $\Delta \text{ATN}_{\text{ND}} + k_D \Delta t$ into Eq. (2) and applying uncertainty propagation (Eq. 6) yields after some rearrangements

$$\frac{\delta \sigma_0}{\sigma_0} = \sqrt{\frac{(\delta A)^2}{A^2} + \left( \frac{\delta Q}{Q} \right)^2 + \left( \frac{\delta \Delta \text{ATN}_{\text{ND}}}{\Delta \text{ATN}_{\text{ND}}} \right)^2 + \left( \frac{\delta k_D}{k_D} \right)^2}. \tag{7}$$

Note that the term $\delta \Delta t$ has been dropped here since any normal drift in the clock can be neglected.

The determination of both $\delta A$ and $\delta Q$ is to some extent dependent on the instrument operator. The term $\delta A$ can be estimated using a magnifier glass with a scale or digital image analysis to measure the area of the sample spot. Here we will assume that the filter size area can be determined with a 2% uncertainty using digital image analysis.

The value of $\delta Q$ comes from both the accuracy of the calibration and the performance of the flow controller of the instrument. The uncertainty of the flow meter (Sierra Instruments, model 824-RFQ-2430) is reported (by the manufacturer) to be 1.5%, which is what will be assumed here. The flow measured by the flow meter is not the exact flow that enters the instrument since there is also a lateral flow through the fibre filter. The lateral flow will bias the internal flow meter readings towards higher values than the actual flow entering the system. The lateral flow is likely to be a function of the pressure difference between the sampling line and the room air, which further adds to the uncertainty in the flow rate.

The drift term $(\delta k_D/k_D)^2$ of Eq. (7) is the most demanding to assess as it may vary greatly from station to station and instrument to instrument for a number of reasons. Drift can be expected to ensue from changes in temperature or relative humidity, changes in lateral flow due to pressure changes in the sampling line, changes in semi-volatile constituents that have deposited onto the filter, etc. The sources that contribute to drift, and the impact of drift on instrument performance, are best studied under controlled conditions in a laboratory. Therefore, drift will largely be omitted in the uncertainty analysis and discussed on the basis of observations.

By substituting $\delta Q$ in Eq. (7) with the flow rate uncertainty ($f_q$) as a fraction of the total flow $Q$, the term $\delta Q^2$ becomes $(f_q Q)^2$. Equivalently, if the uncertainty of the spot size area ($f_a$) is a fraction of the total area $A$, the term $\delta A^2$ becomes $(f_a A)^2$. Equation (7) then becomes

$$\frac{\delta \sigma_0}{\sigma_0} = \sqrt{f_a^2 + f_q^2 + \left( \frac{\delta \Delta \text{ATN}_{\text{ND}}}{\Delta \text{ATN}_{\text{ND}}} \right)^2}. \tag{8}$$

Because the drift term has been left out Eq. (8) describes the best case scenario without any drift taken into account. It should be noted that the term $\delta \Delta \text{ATN}_{\text{ND}}$ describes the random error that originates from the electronics in the instrument. The relative uncertainty of $\delta \Delta \text{ATN}_{\text{ND}}/\Delta \text{ATN}_{\text{ND}}$ can be expressed in terms of measurement-derived values using particle-free air as

$$\frac{\delta \Delta \text{ATN}_{\text{ND,air}}}{\Delta \text{ATN}_{\text{ND,air}}} = \frac{Q}{A} \frac{\delta \sigma_0}{\sigma_0}. \tag{9}$$

In Eq. (9), $\delta \sigma_0_{\text{air}}$ is the standard deviation of $\sigma_0$ at the time resolution of $\Delta t_{\text{air}}$. When determining $\delta \sigma_0_{\text{air}}$, $\Delta t_{\text{air}}$ should be short so that $\Delta t_{\text{air}}$ is the same as the inner cycle for the Aethalometer. Similarly, $\Delta \text{ATN}_{\text{ND}}$ can be written as a function of $\sigma_0$ and substituted into Eq. (9), which yields

$$\frac{\delta \sigma_0}{\sigma_0} = \sqrt{f_a^2 + f_q^2 + \left( \frac{\delta \sigma_0_{\text{air}} \Delta t_{\text{air}}}{\sigma_0 \Delta t} \right)^2}. \tag{10}$$

It is often desirable to know the absolute uncertainty $(\delta \sigma_0)$ of the measurement in units of the quantity measured. Equation (10) then becomes

$$\delta \sigma_0 = \sqrt{\sigma_0^2 \left( f_a^2 + f_q^2 + \frac{\delta \sigma_0_{\text{air}} \Delta t_{\text{air}}}{\Delta t} \right)^2}. \tag{11}$$

Equation (11) implies that the absolute uncertainty of the Aethalometer scales proportionally to $\Delta t^{-1}$ when post-processing using Eq. (2) for a fixed $\Delta \text{ATN}$ and no drift; note
that $\sigma_0$ inside the square root contains $\Delta t^{-1}$. Solving $\delta \sigma_0$ from Eq. (8) yields the same conclusion.

The $\Delta t^{-1}$ dependency can be verified by measuring particle-free air. First, a time series of measurements on particle-free air is needed. This was obtained by measuring particle-free laboratory air with an absolute filter on the inlet of an Aethalometer and logging the extended format of the Aethalometer. Then, the drift in ATN was removed by subtracting a running mean of three points from the reported ATN values, yielding a time series of ATN free of drift ($\text{ATN}_{\text{ND}}$).

Figure 1 depicts the decomposition of ATN of laboratory measurements when measuring particle-free air through an absolute filter. From the figure, it is clear that ATN increased even though no particles should have entered the instrument because of the absolute filter connected to the sample inlet of the instrument. This test implies that there can be instrumental drift that only becomes apparent in long time series. ATN and $\text{ATN}_{\text{ND}}$ shown in Fig. 1a and 1c constitute the data used to produce Fig. 2 in addition to the eBC data that was used for the boxcar average $t_{\text{avg}}$ in the figure. Figure 1c also strengthens the argument that the term $\delta \text{ATN}_{\text{ND}}$ is close to the random error from the electronics when using a running mean to derive $\text{ATN}_{\text{ND}}$ from ATN.

The origin of the drift is not evident and there can well be more than one source for the observed drift. The sampling line comprised a short tube connecting a low resistance absolute filter and a flow meter to the instrument which was open to laboratory air in the other end. The low resistance absolute filter, and the moderate flow rate, should only lower the pressure in the sampling line minutely. If this pressure drop were the reason that unfiltered air enters the sampling line after the filter causing drift, then the drift should be greater when the instrument is connected to a high-volume inlet at a measurement station.

However, the flow was a constant $3.87 \pm 0.02$ litres per minute ($L \text{min}^{-1}$), and the room temperature was a constant $23.2 \pm 0.3 ^\circ C$ (mean $\pm$ standard deviation). Fluctuations in the flow rate and room temperature did not coincide with clear changes in ATN, thus not supporting a pressure difference nor a temperature drift hypothesis, at least not directly. A time series of all wavelengths of ATN, flow rate, and room temperature is shown in Fig. A1 in Appendix A. The times when the most visible jumps occurred were close to midnight on 26 December and 1 January when there was no activity in the lab. The timing of the abrupt changes suggests that unintentional human interference is not likely. Involuntary movement of the filter could well cause an ATN change; but that it occurs by itself seems very unlikely but not impossible.

A hypothesis that could contribute to the observed drift is the adsorption of semi-volatile organic compounds onto the filter. The possible adsorption of organics with an absolute filter in front would likely be severely hampered in comparison to what the effect would be without the filter because of adsorption in the absolute filter. The absorption Ångström exponent ($\alpha_{\text{ap}}$, see Eq. 4) of $\sigma_0$ during the measurements was 1.29. As the origin of the ATN drift is unclear, so is the meaning of $\alpha_{\text{ap}}$. Furthermore, it is likely that changes in relative
humidity will cause ATN to fluctuate, but this is unlikely to cause the steady increase in ATN as seen in Fig. 1.

The ATNNd of Fig. 1 was used to calculate ΔATNNd (and ΔATN) for a range of Δt values (2, 8, 16, 32 . . . 1024 min) to produce new time series of σ0 using Eq. (2). From these time series, the standard deviation of σ0 was calculated and plotted as a function of Δt as shown in Fig. 2. The time series used comprised 13 days of measurements with a Δt of 2 min. Consequently, the values used to calculate δσ0 in the figure decreased with increasing Δt.

Figure 2 shows that when the drift is removed the absolute uncertainty δσ0 follows the predicted Δt⁻¹ relationship. The curve fit for the drift-free δσ0 as a function of Δt gives a slope of −1.026; see Fig. 2. When the drift is not removed, using the running mean method described before, δσ0 is not reduced nearly as rapidly as for the non-drift situation. The difference is arguably due to drift. Also shown in the figure is δσ0 of boxcar-averaged σ0 converted from the eBC output of the instrument as σ0 = MACAE·eBC. The same time interval was used for boxcar averaging (tavg) as was used for Δt.

4 Measurement results

4.1 Measured uncertainties

Aethalometers that are deployed in clean environments can appear at times to just be reporting noise. By simple data post-processing, the signal can be extracted with a greater accuracy, albeit at the expense of temporal resolution (Hagler et al., 2011). This can be done by allowing for a temporal resolution that matches the concentration of species that creates the instrument response, namely the change in ATN, by choosing a constant relative uncertainty (Eq. 8). Equation (8) states that when fQ and fA are constant, the relative uncertainty depends on the change in filter attenuation (ΔATNNd). This fact can be used to produce a time series with a constant relative uncertainty.

However, it should be acknowledged that there is an additional uncertainty due to instrument drift, but in principle a constant uncertainty could also be achieved using Eq. (7). That would require a thorough investigation into the sources for the drift and how they vary between stations, which is not feasible in this study given the remote locations of the stations. However, based on the laboratory measurements the drift can be significant on a timescale from hours to days. When the aim is to determine the drift at a station, the absolute filter should be attached to the sampling line to capture the pressure changes in the sampling line relative to ambient pressure, and changes in relative humidity, on a pristine filter. This could possibly be extended to include loaded filters for different aerosol types and filter loadings.

The measurement uncertainties for the six Aethalometers at the respective stations were determined by measuring particle-free air. For all stations except Alert, particle-free air was sampled for at least 24 h with an absolute filter connected to the instrument inlet. These measurements are shown in Fig. 3 at a wavelength of 590 nm (Summit 880 nm). For Alert, the particle-free air was sampled for a few hours per week comprising 4 days of data in total.

Figure 3 depicts the ATN drift in the different Aethalometers during the particle-free air measurements. The figure shows that all the tested Aethalometers experienced drift during the particle-free air measurements. Also evident from the figure is that the drift of the different Aethalometers (and different sites) can differ. Based on the figure, it is not enough to conduct measurements on particle-free air for a few hours in order to assess the instrument performance at the site. Filtered air measurements should instead be performed over a period of 24 h or more. These measurements should be conducted on a pristine filter to minimise the influence of semi-volatile constituents that could have been deposited onto the filter (Cappa et al., 2008; Lack et al., 2008).

For reference, a linear drift of ΔATN0 = 1.0 in 24 h is shown in Fig. 3, which corresponds to a σ0 value of 0.07 Mm⁻¹ when using Q = 5 L min⁻¹, A = 0.5 cm², and Δt = 24 h. The consequence of a linear drift of 1.0 in 24 h would also set the lowest value achievable. As can be seen from the inserts of Fig. 1, ATN0 does need not be increasing all the time, and thus lower values of σ0 are achievable during periods with little drift. It should be mentioned that this drift will also affect the eBC concentrations reported by the instrument. For the five instruments evaluated here, the drift uncertainty is shown in Fig. 3 to be roughly 0.01–0.1 Mm⁻¹.
The particle-free air measurements were not all conducted in the same manner, which is both fortunate and unfortunate. Figure A2 in Appendix A shows the zero air measurements for ATN in greater detail. The Aethalometers at Alert, Barrow, Tiksi, and Pallas were not connected to the common aerosol inlet at the sites but drew air from inside, through a low resistance absolute filter. Again, this setup should only result in a negligible lower pressure in the sampling line after the filter. The most intriguing of the zero air measurements is Tiksi. Similar to Fig. 1, Tiksi shows abrupt changes in ATN that cannot be related to pressure changes in the sampling line. One of the abrupt changes occurred at night when there was no personnel at the station (see Fig. A2). The Tiksi Aethalometer also shows uneven drift, when compared to other channels, for the 370 and 950 nm channels that cannot be explained with aerosol deposition onto the filter. The hypothesis is therefore that this is electronic drift. Changes in filter morphology or position should affect all wavelengths. The Tiksi Aethalometer also has a noisy 880 nm channel which is clearly visible in Fig. A2.

Pallas, Zeppelin, and Summit all experienced drift immediately following the change to a pristine filter although the drifts were quite different. For Pallas, the drift was observed as a gradual and increasing ATN. The drift was largest for 370 nm at both Pallas and Zeppelin but not in the same very clear trend breaking way that was observed at Tiksi; although at Zeppelin the 370 nm channel behaved somewhat differently than the other channels. At Summit, the ATN changes were faster although smaller in magnitude. Again we hypothesise that these changes in ATN during the particle-free air measurements can be due to deposition or evaporation, or both, of semi-volatile organic compounds or changes in sample air relative humidity. The decreasing ATN at Zeppelin could be due to evaporation of adsorbed water or organics from the filter. At Barrow, the zero air measurements were conducted on a loaded filter. After an absolute filter was placed before the instrument, ATN started to drop. Possible causes can be the evaporation of organics or water vapour. It is possible that the absolute filter changes the partial pressure of one or more gas-phase constituents that subsequently affect the ATN. The rapid changes in the Barrow ATN values correlate well with rapid fluctuations in the sampling line temperature which is exposed to room air and therefore also to the air-conditioning unit. The ATN values increased when room temperature was dropped and vice versa.

The standard deviations of the \( \Delta \text{ATN}_{\text{ND}} \) measurements made with an absolute filter in line are shown in Table 2. Because the only wavelength-dependent variable in Eq. (8) is \( \delta \Delta \text{ATN}_{\text{ND}} \), the change in the ATN measurements with respect to wavelength will also be the sole source of the difference in the relative uncertainty between different wavelengths. The values that describe the relative uncertainty in terms of \( \Delta t \) and \( \sigma_0 \) (Eq. 10) are presented in Table 3.

Figure 4 shows the relative uncertainty (\( \delta \sigma_0 / \sigma_0 \)) of Eq. (8) as a function of \( \Delta \text{ATN}_{\text{ND}} \) based on measurements conducted with an absolute filter upstream of the instrument. For clarity, the figure was produced using a mean of all wavelengths to represent the typical relative uncertainty of the instrument. The mean values were calculated from Table 2. Figure 4 shows how the relative uncertainty decreases when \( \Delta \text{ATN} \) increases. The upper x-axis scale of \( \sigma_0 \) in the figure was calculated for reference using \( A = 5 \text{ L min}^{-1}, Q = 0.5 \text{ cm}^{-2} \), and \( \Delta t = 60 \text{ min} \).

Implicit from both Fig. 4 and Eq. (8) is that the relative uncertainty of the instrument changes with the aerosol concentration when using a fixed \( \Delta t \); for a fixed \( \Delta t \), \( \Delta \text{ATN} \) will change according to the concentration. The equation for the relative uncertainty (Eq. 8) can be used as a criterion to achieve a more constant level of uncertainty which was not captured when the method was introduced by Hagler et al. (2011). This can either be determined from Fig. 4 directly or calculated from Eq. (8) after the term \( \delta \Delta \text{ATN}_{\text{ND}} \) has been determined.

One way to characterise the performance of an Aethalometer is to calculate the \( \Delta \text{ATN} \) value at which the flow (\( f_0 \)) and spot size (\( f_a \)) uncertainties together are equally important as the \( \Delta \text{ATN} \) uncertainties. This is shown in Table 4. The crossover was calculated by solving \( \Delta \text{ATN} \) from the terms under the square root of Eq. (8), namely \( \Delta \text{ATN}_{\text{ND}} = (\delta \Delta \text{ATN}_{\text{ND}}^2 / (f_a^2 + f_q^2))^{1/2} \). The uncertainty in the flow rate, relative to the uncertainty in the ATN measurements, diminishes exponentially when \( \Delta \text{ATN} \) decreases (Fig. 4). Here, a criterion of \( \Delta \text{ATN} \geq 2 \) was used in the post-processing of the data to also allow for a lower detection limit in the boxcar-averaged reference data that is discussed in the next section.
end up with seven time series for each instrument, with different timestamps. That could make further data analysis unnecessarily convoluted. It is worth pointing out that if the data set being analysed is going to be averaged, then the ATN values included with the averaged data set should not be averaged – an averaged ATN would make $\Delta t$ less well defined. Instead, either the first or the last ATN value during the averaging period should be incorporated into the averaged data set.

The time series of the 1 h boxcar-averaged Aethalometer data is shown in Fig. 5. The time series of the adaptive collection time is shown in Fig. 6. When using the adaptive collection time, it is clear that when the absorption coefficient is low, the time resolution is low. At higher absorption coefficients, the time resolution is better. This is desirable since it means that at high concentrations of light absorbing aerosol particles there is no loss in temporal resolution, whereas at low concentrations this adaptive method is capable of reaching lower detection limits quicker than boxcar averaging when drift is minimal.

Comparing Figs. 5 and 6, it is clear that the adaptive collection time approach is to be favoured when $\sigma_{ap} < 0.1 \text{ Mm}^{-1}$ because of instrument noise. In Fig. 5, it is shown that at low $\sigma_{ap}$ the 1 h averages of the data set are clearly more scattered than when using the adaptive collection time method when $\sigma_{ap}$ is low (Fig. 6). Since the $y$-scale of Fig. 5 is logarithmic, negative values are not shown, although they are still present in the 1 h averaged time series. By definition, the adaptive collection time approach will not produce negative $\sigma_0$ values since $\Delta \text{ATN}$ is always positive.

In fact, for the measurements studied here, when $\sigma_0$ is above 2.1–6.7 $\text{Mm}^{-1}$, there is no loss in the temporal resolution in the 1 h averaged data of Fig. 5. The range in $\sigma_0$ is due to the fact that the different Aethalometers at these six Arctic sites are operated at different flow rates. Figure 7 shows histograms of $\Delta t$ for the different stations using the adaptive collection time approach.

Figure 6 shows values that are lower than the example drift in $\Delta \text{ATN} = 1.0$ in 24 h (Fig. 3), which implies that there are periods where the drift can be substantially lower. In Figs. 5 and 6, the $\sigma_{ap}$ values come from $\sigma_0$ values that have been corrected using a $C_f$ value of 3.45 as discussed in the next section. Thus, the drift uncertainty seen in Fig. 3 becomes 0.003–0.03 $\text{Mm}^{-1}$ after $C_f$ is applied.

### 4.2 Aethalometer correction factor for the Arctic

The determination of the Aethalometer correction factor for the Arctic was done according to Eq. (3). For the calculations, $\sigma_0$ values were obtained using the collection time approach because a constant relative uncertainty is desirable. The collection time approach applied to the Aethalometer data impacts the co-located absorption photometer data as well. The collection times for the Aethalometers were adapted to $\Delta \text{ATN}$. Subsequently, data from the co-located instruments were boxcar-averaged to the same time intervals as the Aethalometer data in order to calculate $C_f$. The co-located instruments will therefore have a longer averaging during times when $\sigma_{ap}$ is low, given that $\sigma_{ap}$ and $\Delta \text{ATN}$ correlate. For standard boxcar averaging, random measurement noise reduces proportionally to $t_{\text{avg}}^{-0.5}$ (e.g. Springston and Sedlacek, 2007). Using the collection time approach, $\delta \sigma_0$ can at best be reduced at a rate of $\Delta t^{-1.0}$ (Fig. 2).

Applying uncertainty propagation on Eq. (3) yields

$$\frac{\delta C_f}{C_f} = \sqrt{\left(\frac{\delta \sigma_0}{\sigma_0}\right)^2 + \left(\frac{\delta \sigma_{ap}}{\sigma_{ap}}\right)^2}.$$  \hspace{1cm} (12)

The term $\delta \sigma_0/\sigma_0$ is discussed in this work and cannot be lower than the spot sizes and flow rate uncertainties. This also holds true for the co-located absorption photometers. Equation (12) also implies that the relative uncertainty of $C_f$ increases when $\sigma_{ap}$ and $\sigma_0$ are low if the absolute uncertainties $\delta \sigma_{ap}$ and $\delta \sigma_0$ remain constant. Using $\Delta \text{ATN}$ to determine the averaging time for $\sigma_{ap}$ will lower $\delta \sigma_{ap}$ proportionally to $t_{\text{avg}}^{-0.5}$. For non-drift situations, the adaptive collection time approach will reduce $\delta \sigma_0$ at a rate of $\Delta t^{-1.0}$, which is faster than boxcar averaging and desirable because Aethalometers are generally more noisy than PSAPs and MAAPs. For reference, the PSAP and MAAP absolute uncertainties are typically 0.02 and 0.06 $\text{Mm}^{-1}$ for 5 min averages (Müller et al., 2011). The 5 min PSAP absolute uncertainty was calculated from 0.05 $\text{Mm}^{-1}$ for 1 min averages using a $t_{\text{avg}}^{-0.5}$ dependency. These absolute uncertainties of the PSAP and MAAP
Figure 5. Time series of 1 h averaged absorption data for $\lambda = 520$ nm. In the figure, the attenuation coefficients have been corrected using the Arctic correction factor $C_f$ of 3.45.

Figure 6. Time series of absorption coefficients using the adaptive collection time approach at a wavelength of 520 nm. The attenuation coefficients have been corrected using the Arctic correction factor $C_f$ of 3.45.

are much lower than the Aethalometer uncertainties shown in Table 3.

The uncertainty of $C_f$ is greater than the uncertainty that originates in measurement noise. Filter-based absorption photometers are generally considered to be accurate to within 20–30 % of the true $\sigma_{ap}$ value (Bond et al., 2013). The accuracy is a combination of electronic noise, instrument variability, and calibration uncertainty (e.g. Sherman et al., 2015). The adaptive averaging time approach will only lower measurement uncertainties from electronic noise and not the uncertainties associated with the measurement technique itself.

The $C_f$ values presented here can be used in many of the existing Aethalometer correction algorithms in the place of the multiple scattering enhancement factor. However, $C_f$ values calculated here should be viewed as a harmonising correction factor for the Arctic Aethalometers to the co-located filter-based absorption photometers and not as a literal multiple scattering enhancement factor.

There are several possible issues with the derivation of $C_f$ values presented here. First, in this study, the co-located absorption photometers also rely on measurements using filter-based absorption techniques – it remains unclear to which extent this will affect the absolute values of $C_f$ because no absorption standard measurements were available at the sites. However, since the filter changes of the different instruments are not synchronised, and because the data sets cover 3 years...
at each site, it can be assumed that there is very little coincidence with respect to filter loading effects. Thus, the $C_f$ values presented here should represent typical values for the different sites. This argumentation should especially hold true for a moderately loaded filter, e.g. ATN < 10. Second, the flow rates of the different instruments differ, which can affect the $C_f$ values due to different penetration depths (Lack et al., 2009; Nakayama et al., 2010). Third, it has to be acknowledged that there can be a bias in the absolute $C_f$ values because of imperfect corrections of filter artefacts in the reference instruments (Backman et al., 2014; Müller et al., 2011). However, this bias should not fundamentally alter the ATN dependency because filter changes were not performed in sync. The reasoning is elaborated on in Appendix B. As the filter gets loaded with aerosol particles, loading effects come into play. These loading effects change between the filter spots depending on the optical properties of the aerosol that is being deposited on that particular spot (Virkkula et al., 2015) and even during sampling on the same spot (Drinovec et al., 2015).

Such detailed analysis of filter loading effects is not feasible with this data set since it would require data with a high temporal resolution and preferably concurrent non-filter-based light absorption measurements. In general, the goodness of evaluation for all filter-based light absorption measurements should be the continuous light absorption coefficients over filter spot changes so that a filter spot change would go unnoticed; this should hold true for all aerosol types and loadings. This means that there would not be an ATN dependency when compared to non-filter-based light absorption measurements.

It has been shown that published Aethalometer correction algorithms, which aim to compensate for filter loading and multiple scattering effects, do not necessarily remove the ATN dependence when applied on data from different stations (Fig. 4 in Collaud Coen et al., 2010). Again, the aim is not to add another correction algorithm to literature. Instead, the $C_f$ values presented here should be interpreted as a means to make Aethalometers in the Arctic more inter-comparable by introducing a $C_f$ value for the Arctic using the co-located absorption photometers.

Figure 8 shows the calculated $C_f$ values as a function of ATN for the six Arctic sites. Since the data depicted in Fig. 8 were produced using a concentration-adapted collection time, the statistics in the figure were calculated using a collection-time-weighted percentile (Hyndman and Fan, 1996). Without this weighting, the statistics would have effectively been concentration weighted. Figure 8 is equivalent to Fig. 4 of Collaud Coen et al. (2010) for the values labelled “AE manufacturer” in their figure.

In general, Tiksi and Pallas show the highest $C_f$ values, whereas Summit shows the lowest. Summit stands out as an outlier in Fig. 8; it is the station at the highest elevation and uses a 1-wavelength Aethalometer (880 nm). The Summit Aethalometer data were interpolated to a wavelength of 637 nm using an $\alpha$ of 0.814 obtained from the co-located 3-wavelength CLAP. A summary of the different $C_f$ values calculated for the stations is presented in Table 5.

In addition to the different $C_f$ values observed over the ATN range in Fig. 8, there are other differences among the stations. Some of the $C_f$ values decrease as a function of ATN. In the ATN range of 0–10, the median $C_f$ values for Alert and Tiksi are greater than at the other stations, but at higher ATN the Alert and Tiksi $C_f$ values decrease. This is an expected behaviour and is due to the filter loading effect causing a decrease in Aethalometer sensitivity. However, a decrease in $C_f$ with ATN is barely noticeable for the Barrow and Zeppelin data sets, although the variation in $C_f$ at Zeppelin makes the trend — or lack thereof — less clear.
Figure 8. Correction factor ($C_f$) as a function of filter attenuation (ATN) calculated using Eq. (3). The grey dashed line and the right hand y axis show the number of data points that each ATN range comprise. The blue boxes represent the 25th to 75th percentile range, whereas the red circles represent the median values. $C_f$ values in the figure are for all available wavelengths. The figure also shows the median single-scattering albedo ($\omega_0$), back-scatter fraction ($b$), and scattering Ångström exponents ($\alpha_{sp}$). The $\omega_0$ of the aerosol was calculated using the absorption coefficients from Fig. 6. The slope of the ATN dependence is shown as the value $k$ so that $C_f^{-1} = \sigma_{ap}/\sigma_0 = 1 + k \cdot ATN$.

Table 5. Aethalometer correction factors ($C_f$) for the different stations. The values were calculated using averaging-time-weighted percentiles because of the adaptive average time used to derive them. The top portion of the table reports the $C_f$ values for all available wavelengths of the co-located absorption instruments. Aethalometer wavelengths were interpolated to these co-located absorption photometer wavelengths using absorption Ångström exponents. The Summit AE-16 data were extrapolated to a wavelength of 637 nm using $\alpha = 0.814$. The bottom portion of the table reports the statistics of $C_f$ using all available wavelengths. The last row in the table shows the number of data points ($N$) used for the statistics. The overall statistics comprise all stations except the high-altitude station of Summit.

<table>
<thead>
<tr>
<th></th>
<th>Alert</th>
<th>Summit</th>
<th>Barrow</th>
<th>Tiksi</th>
<th>Pallas</th>
<th>Zeppelin</th>
<th>Overall</th>
</tr>
</thead>
<tbody>
<tr>
<td>467 nm</td>
<td>3.43</td>
<td>–</td>
<td>3.17</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>525 nm</td>
<td>3.43</td>
<td>–</td>
<td>3.09</td>
<td>–</td>
<td>–</td>
<td>3.25</td>
<td>–</td>
</tr>
<tr>
<td>637 nm</td>
<td>3.43</td>
<td>1.61</td>
<td>3.12</td>
<td>4.01</td>
<td>4.22</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>25th</th>
<th>50th</th>
<th>75th</th>
<th>$N$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alert</td>
<td>2.70</td>
<td>3.43</td>
<td>4.37</td>
<td>3455</td>
</tr>
<tr>
<td>Summit</td>
<td>0.73</td>
<td>1.61</td>
<td>2.44</td>
<td>1055</td>
</tr>
<tr>
<td>Barrow</td>
<td>2.56</td>
<td>3.12</td>
<td>3.64</td>
<td>3590</td>
</tr>
<tr>
<td>Tiksi</td>
<td>3.34</td>
<td>4.01</td>
<td>4.77</td>
<td>2348</td>
</tr>
<tr>
<td>Pallas</td>
<td>3.36</td>
<td>4.22</td>
<td>5.85</td>
<td>3226</td>
</tr>
<tr>
<td>Zeppelin</td>
<td>2.28</td>
<td>3.25</td>
<td>6.91</td>
<td>2836</td>
</tr>
<tr>
<td>Overall</td>
<td>2.93</td>
<td>3.45</td>
<td>4.15</td>
<td>16510</td>
</tr>
</tbody>
</table>

Again, Summit shows a different behaviour altogether. As the filter ATN increases, so do the $C_f$ values. This is contrary to the expected behaviour of the filter loading effect in which loading generally decreases the sensitivity of a filter-based absorption measurement technique (Arnott et al., 2005; Virkkula et al., 2007). The filter loading effect is most pronounced for an aerosol with a low $\omega_0$ (Sheridan et al., 2005). The fact that Summit does not follow this trend suggests that the aerosol optical properties of Summit are different in relation to the other stations. The different behaviour, however, does not seem to be related to $\omega_0$ as the $\omega_0$ of Summit does not stand out.

The scattering Ångström exponent ($\alpha_{sp}$, Eq. 4) is shown in Fig. 8. The $\alpha_{sp}$ of an aerosol is indicative of aerosol particle size with values below unity indicating super micron aerosol, and values close to 4 indicate a predominantly fine-mode aerosol. Only the nephelometers at Pallas and Summit are connected to PM$_{2.5}$ inlets, with the rest connected to PM$_{10}$ inlets. Thus, both Pallas and Summit have smaller aerosol particles than the rest of the stations according to $\alpha_{sp}$ and very similar ones when compared with each other. If the
PM$_{2.5}$ inlet were to be the reason for a lower $C_f$ at Summit, then Pallas should also have to be influenced since the MAAP at Pallas is behind a PM$_{2.5}$ inlet. Also shown in Fig. 8 are aerosol back-scatter fractions ($b$) which are related to the size of the aerosol and have been shown to affect Aethalometers (Virkkula et al., 2015). The site that resembles Summit the most, considering $\alpha_{sp}$, $b$, and $\omega_0$ is Pallas. The Pallas Aethalometer was operated with a filter change interval of 8 h for most of the time, and therefore there are very few data points with an ATN above 10. Hence, it is questionable whether there are enough data to be able to draw conclusions about a trend in the $C_f$ and ATN relationship for Pallas.

Also shown in Fig. 8 are the slopes ($k$) of the ATN dependence. The slopes were calculated by linear regression to match the equation $\sigma_{ap}/\sigma_0 = 1 + k \cdot$ ATN that follows the notation of previous work (Virkkula et al., 2007, 2015). It is well known that light scattering by particles affects filter-based absorption photometers, which wrongly gets interpreted as light absorption, termed apparent absorption. For the co-located PSAP and CLAP instruments, this has been compensated for by subtracting a fraction of the light scattering from the light absorption (Bond et al., 1999; Ogren, 2010). For the co-located MAAP instruments, scattering correction is applied by the firmware (Petzold and Schönlinner, 2004). These corrections need not be enough for all types of aerosols, and a cross sensitivity to light scattering can remain (Müller et al., 2011). For $\sigma_0$, no scattering correction has been applied.

Figure 9 shows $C_f$ as a function of $\sigma_{sp}$. Since no scattering correction has been applied to $\sigma_0$ the expected behaviour is that $\sigma_0$ increases with $\sigma_{sp}$. This seems to be the case for Alert, Summit, Barrow, and Zeppelin, with a clearer trend for Alert and Barrow. All stations that show an increasing trend of $C_f$.
with $\sigma_{sp}$ have a co-located PSAP or a CLAP. It is possible that the non-compensation of $\sigma_{sp}$ on $\sigma_0$ is further influenced by a too-large or too-small compensation of $\sigma_{sp}$ on $\sigma_{ap}$. Since $\sigma_{ap}$ is the denominator of Eq. (3), an overcompensation of apparent absorption would also cause $C_1$ to increase with $\sigma_{sp}$. Both stations that have a MAAP as the co-located instrument show a decreasing $C_1$ as $\sigma_{sp}$ increases.

Another possible explanation for a changing $C_1$ with $\sigma_{sp}$ can be that different levels of $\sigma_{sp}$ are associated with different aerosol types and therefore also $C_1$. Different levels of pollution can be associated with different sources of pollutants. These different sources could have different chemical and physical properties that could impact the $C_1$ ratio. The right hand y axis in Fig. 9 shows the median $\omega_0$ of the aerosol for the range of $\sigma_{sp}$. The $C_1$ value seems to follow the general trend of $\omega_0$ so that when $\omega_0$ increases, so does $C_1$. It is not clear whether this is due to measurement artefacts or a difference in aerosol properties. The fact that the same behaviour is observed in Tiksi and Pallas indicates that the cross sensitivity to $\omega_0$ is not restricted to the PSAP and CLAP instruments.

In the upper part of Table 5, it can be seen that there is not much variation in the $C_1$ values at different wavelengths. In the table, the Aethalometer data were interpolated using Ångström exponents to match the wavelength of the reference instruments using Eqs. (4) and (5). It cannot be ruled out that the low variability in $C_1$ with wavelength is a consequence of the co-located instruments used. No multi-wavelength long-term measurements of aerosol absorption coefficients from non-filter-based measurement instruments were available. The lower part of Table 5 provides statistics of $C_1$ for each site. Because there does not seem to be a great wavelength dependence on the $C_1$ value at Alert and Barrow, the $C_1$ value for all wavelengths was calculated using all available wavelengths. The overall value of $C_1$ for sites across the Arctic was determined to be 3.45. The value of 3.45 was calculated using average-time weighted median as discussed earlier, and the weighted 25th and 75th percentiles for the all wavelength $C_{ref}$ values are 2.93 and 4.15, respectively. Because Summit appears to be significantly different from the other Arctic stations, the Summit $C_1$ was omitted from the grand median $C_1$ calculation.

Figure 10 depicts the relationship between the reference absorption instrument ($\sigma_{ap}$) and $\sigma_0$ which yields $C_1$; see Eq. (3). The figure is provided as an overview comprising all wavelengths and all stations except for Summit, because of the reasoning mentioned before. In the figure, in addition to the weighted median value of 3.45, the slope of the bivariate fit is also shown. The fit was performed using bivariate regression with the averaging time as weights (Cantrell, 2008). The slope of the bivariate regression becomes 3.59 (standard error 0.01). Both the regression method and the weighted median method yield values that are within 3 % of each other. The root-mean-square error (RMSE) was calculated from the predicted $\sigma_0$ using the results from the regression, which becomes 1.14 Mm$^{-1}$.

The mass absorption cross section (MAC) describes the relationship between eBC mass concentrations and $\sigma_{ap}$. Similarly, $\text{MAC}_{AE}$ describes the relationship between $\sigma_0$ and eBC as given by the manufacturer; both MAC and $\text{MAC}_{AE}$ have units of m$^2$ g$^{-1}$. A simple evaluation can be performed to investigate whether the $C_1$ value is reasonable, assuming that the difference between $\text{MAC}_{AE}$ and MAC is $C_1$ (Arnott et al., 2005). If the Aethalometer measured at a wavelength of 550 nm, then $\text{MAC}_{AE}$ would be 26.59 m$^2$ g$^{-1}$. Compensating $\text{MAC}_{AE}$ with $C_1 = 3.45$ would yield a MAC of 7.7 m$^2$ g$^{-1}$, i.e. a MAC which is 3.45 times lower than $\text{MAC}_{AE}$. This MAC is within the range suggested by Bond and Bergstrom (2006), namely 7.5 ± 1.2 m$^2$ g$^{-1}$ at 550 nm, which implies that the $C_1$ value determined here is reasonable. This simple evaluation, however, does not take into account any apparent absorption or coating effects.

Multiple scattering correction factors ($C_{ref}$) have been reported for a range of different sites around the world. Although the $C_1$ values reported here cannot be considered to be equivalent to multiple scattering corrections, it is still useful to compare $C_1$ with $C_{ref}$. $C_{ref}$ values reported in the literature are summarised in Table 6.

Some of the variations in the reported $C_{ref}$ values of Table 6 can be attributed to the different ways in which they were calculated. Some $C_{ref}$ values were calculated with a filter loading correction applied and some without. Moreover, some of the $C_{ref}$ values were calculated with both a scattering and a filter loading correction applied. For the sake of inter-comparability, a $C_{ref}$ value calculated without any of the available correction algorithms would be preferable.

Comparing the grand median $C_1$ value of 3.45 conceived here shows that it is well within the range of studies reporting $C_{ref}$ values. The values that are the closest are those by Collaud-Coen et al. (2010), which is not surprising since that study had light absorption coefficients from a MAAP as reference and not true reference absorption measurements. The $C_1$ value of 3.45 conceived here is also close to 3.5, which is what the Global Atmospheric Watch’s World Calibrations Centre for Aerosol Physics recommends using globally (GAW Report No. 227; http://wmo-gaw-wcc-aerosol-physics.org/wmo-gaw-reports.html).

5 Conclusions

In clean environments, such as in the Arctic during summer months, measurements of aerosol light absorption coefficients can be below the detection limit of the instrument. Symptomatically, it is not uncommon to encounter measurements reporting negative equivalent black carbon concentrations or light absorption coefficients. These values are without physical meaning and originate from instrument noise and uncertainties.
Here a post-processing method for Aethalometer data based on collection time is elaborated on. This post-processing approach allows for an arbitrary collection time $\Delta t$, which lowers the electronic noise of the Aethalometer proportionally to $\Delta t^{-1}$. In comparison, boxcar averaging lowers the noise proportional to $t^{-0.5}$. The greatest benefit of this approach can be achieved when drift in the Aethalometer ATN measurements is minimised.

The noise characteristics of Aethalometers are best estimated using measurements of particle-free air. Based on these measurements, it is recommended that particle-free air measurements should be conducted for at least 24 h or more. Furthermore, the absolute filter used for particle-free air measurements should be connected between the instrument and the sampling line from which the sample is drawn during normal operation. From these data, the electronic noise and drift can be evaluated.

The uncertainty analysis showed that the collection time approach can be used with a simple criterion that keeps the signal-to-noise ratio constant, namely that the post-processing calculations are invoked once the filter attenuation of the instrument has changed by more than $\Delta \text{ATN}$. This criterion will cause the collection time to vary according to the concentration of absorbing aerosol particles. The collection time approach was applied to Aethalometer data from six Arctic monitoring sites using $\Delta \text{ATN} \geq 2$.

In addition, using co-located absorption photometer measurements at each site, an Arctic-specific Aethalometer correction factor ($C_f$) was calculated using the collection time approach as described above. This correction factor harmonises Aethalometer attenuation coefficients with light absorption coefficients as measured by the co-located light absorption photometers. For all wavelengths, and all low-altitude Arctic stations (i.e. all stations except Summit), the median $C_f$ value was calculated to be 3.45. The 25th to 75th percentile range of $C_f$ was 2.93–4.15. The $C_f$ value for Summit was calculated to be 1.61. The reason for the low $C_f$ value at Summit remains unresolved.

**Data availability.** The source data used for this article are stored in the EBAS database operated by the Norwegian Institute for Air Research http://ebas.nilu.no. EBAS is a database for surface in-situ measurements stations around the globe and is used as data repository by several frameworks. One of these is the World Meteorological Organisation Global Atmosphere Watch World Data Centre for Aerosol that also provided the data management service. The final data product resulting from the work presented here constitutes a secondary dataset targeted at one single analysis, as opposed to data stored in EBAS, which is not filtered for a specific purpose. The secondary data set is archived at the respective repository provided by the Aerosols, Clouds, and Trace gases Research Infrastructure (ACTRIS). ACTRIS also provided quality assurance by instrument comparison workshops for some of the instruments used here. The final data set is available at https://doi.org/10.21336/gen.1 (Backman et al., 2017).
Figure A1. Time series of (a) ATN for all Aethalometer wavelengths and (b) measured Aethalometer flow rate and room temperature with an external flow meter (TSI Inc 4000 series).

Figure A1 shows the time series of the ATN for all wavelengths during the laboratory measurements with an absolute filter. The flow rate at the inlet of the Aethalometer was measured with a TSI 4000 series flow meter in front of the instrument and after the absolute filter. The temperature that was measured by the flow meter is also shown in the figure.

Figure A2 shows all wavelengths of ATN for the five Arctic stations where measurements were conducted for over 24 h with an absolute filter connected to the instruments sample inlet.
Figure A2. ATN for all wavelengths from the measurements with absolute filters connected to the sample inlet. Alert is not shown because the measurements were not conducted for more than a few hours although they were conducted regularly.
Appendix B

The examination of the ATN dependence of an Aethalometer using a co-located absorption photometer requires the assumption that any remaining ATN dependence of the co-located instrument does not impact the ATN dependence of the Aethalometer substantially. An ATN dependence can be expressed as

$$\sigma_{ap} = (1 + k \cdot \text{ATN}) \sigma_0,$$  \hspace{1cm} (B1)

where \(\sigma_{ap}\) is the light absorption coefficient that has no ATN dependence and \(\sigma_0\) is an ATN-dependent light absorption coefficient. The term \(k\) describes the magnitude of the ATN dependence (Virkkula et al., 2015).

Consider a time series with a constant light absorption coefficient of 2 Mm\(^{-1}\). Solving \(\sigma_0\) from Eq. (B1) yields the equation that can be used to produce ATN-dependent time series of both a hypothetical Aethalometer and PSAP. The Aethalometer was given a \(k\) value of 0.002. The value of 0.002 is for a high single-scattering albedo aerosol (Virkkula et al., 2015). A \(k\) value of 0.001 was assigned to the hypothetical PSAP representing a non-ideal compensation of filter loading effects during data post-processing that would correspond to half the ATN dependence of the Aethalometer.

In order to achieve asynchronous filter changes, the ratio spot size area \((A)\) to flow rate \((Q)\) was chosen so that they differ. The Aethalometer was given a flow rate of 5 L min\(^{-1}\) and a spot size of 0.5 cm\(^2\). The PSAP was given a flow rate of 1.1 L min\(^{-1}\) and a spot size of 20 mm\(^2\). The time base for these calculations was 60 min. The change in ATN was calculated as

$$\Delta \text{ATN} = \frac{\sigma_0 Q \Delta t}{A}.$$  \hspace{1cm} (B2)

From Eq. (B2), a time series was constructed from increments of ATN so that ATN was set to begin from 0 after a threshold of 85 was reached, thus simulating a filter change. The ATN time series was then applied to Eq. (B1) to produce two time series of \(\sigma_0\) that both depend on ATN. Both time series are shown in the left panel of Fig. B1.

The ratio of \(\sigma_{ap}/\sigma_0\) was divided into ATN bins and plotted as a function of ATN (right panel of Fig. B1). Curve fitting to the median of each bin gives the \(\sigma_{ap}/\sigma_0\) ratio as a function of ATN, which is the equation \(\sigma_{ap}/\sigma_0 = 1 + k \cdot \text{ATN}\). The curve fit gives a \(k\) value of the simulated Aethalometer of 0.00190 which is 5% less than expected (it should be 0.002). Varying the \(k\) value of the co-located instrument between 0 and 0.004 yields \(k\) values of the simulated Aethalometer that are between −2 and −27% of from the expected \(k\) values.

The same exercise was repeated for a randomly varying concentration that is shown in Fig. B2, which should represent a real world situation. The time series was generated using random numbers. Changing the \(k\) value of the co-located instrument between 0 and 0.004 gives an ATN dependency that is between −2 and −29% of the expected. It should be noted that more data points make the slope closer to the expected value of 0.002. The ratio \(\sigma_{ap}/\sigma_0\) is equivalent to \(C_f^{-1}\) from the paper and should therefore support the interpretation of the ATN dependency of \(C_f\).

Figure B1. (a) shows the time series of an ATN-dependent \(\sigma_0\) for a hypothetical Aethalometer and co-located PSAP so that the filter changes were not performed in synchronisation. The “No ATN dependency” in the figure legend denotes \(\sigma_{ap}\) in Eq. (B1), from which both the hypothetical instrument ATN dependencies were calculated. (b) shows the ATN dependence of the Aethalometer when compared to the ATN-dependent co-located PSAP.
Figure B2. Same as Fig. B1 but with a variable concentration. Panel (a) shows the time series of a ATN-dependent $\sigma_0$ for a hypothetical Aethalometer and co-located PSAP so that the filter changes were not performed in synchronization. The “No ATN dependency” in the figure legend denotes $\sigma_{ap}$ in Eq. (B1), from which both the hypothetical instrument ATN dependencies were calculated. Panel (b) shows the ATN dependence of the Aethalometer when compared to the ATN-dependent co-located PSAP.
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