Reconciling aerosol light extinction measurements from spaceborne lidar observations and in-situ measurements in the Arctic

M. Tesche¹, N. Rastak¹, R. J. Charlson², P. Glantz¹, P. Zieger¹, and H.-C. Hansson¹

¹Department of Applied Environmental Science, Stockholm University, Stockholm, Sweden
²Department of Atmospheric Sciences, University of Washington, Seattle, WA 98195, USA

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Correspondence to: M. Tesche (matthias.tesche@itm.su.se)
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Abstract

In this study we investigate to what degree it is possible to reconcile continuously recorded particle light extinction coefficients derived from dry in-situ measurements at Zeppelin station (78.92° N, 11.85° E, 475 m a.s.l.) at Ny-Ålesund, Svalbard, that are recalculated to ambient relative humidity, and simultaneous ambient observations with the Cloud-Aerosol Lidar with Orthogonal Polarization (CALIOP) aboard the Cloud-Aerosol Lidar and Infrared Pathfinder Satellite Observations (CALIPSO) satellite. To our knowledge, this represents the first study that compares spaceborne lidar measurements to optical aerosol properties from short-term in-situ observations (averaged over 5 h) on a case-by-case basis. Finding suitable comparison cases requires an elaborate screening and matching of the CALIOP data with respect to the location of the Zeppelin station as well as in the selection of temporal and spatial averaging intervals for both the ground-based and spaceborne observations. Trustworthy reconciliation of these data cannot be achieved with the closest approach method that is often used in matching CALIOP observations to those taken at ground sites due to the transport pathways of the air parcels that were sampled. The use of trajectories allowed us to establish a connection between spaceborne and ground-based observations for 57 individual overpasses out of a total of 2018 that occurred in our region of interest around Svalbard (0 to 25° E; 75 to 82° N) in the considered year of 2008. Matches could only be established during winter and spring, since the low aerosol load during summer in connection with the strong solar background and the high occurrence rate of clouds strongly influences the performance and reliability of CALIOP observations. Extinction coefficients in the range from 1 to 100 Mm\(^{-1}\) were found for successful matches with an agreement of a factor of 1.85 (median value for a range from 0.38 to 17.9) between the findings of in-situ and spaceborne observations (the latter being generally larger than the former). The remaining difference is likely to be due to the natural variability in aerosol concentration and ambient relative humidity, an insufficient representation...
of aerosol particle growth in the used hygroscopicity model, or a misclassification of aerosol type (i.e., choice of lidar ratio) in the CALIPSO retrieval.

1 Introduction and motivation

Understanding and quantifying the climatic effects of natural and anthropogenic aerosols from direct observations requires a combination of data from a variety of instruments that usually apply very different measurement techniques. For example, ground-based in-situ measurements of aerosol optical, microphysical, and chemical properties (that are usually carried out with very high temporal resolution but only at a limited number of locations) can be combined with satellite observations or aircraft measurements (that generally provide us with better satellite observations or air- aircraft measurements (that generally provide us with better satellite observations or air- craft measurements (that generally provide us with better satellite observations or air-

1. Differences in measurement techniques: Different properties of the aerosols are sensed or observed by the various instruments. Satellite observations usually are based on optical properties, while in-situ measurements can be of optical properties as well as physical and chemical properties that can be transformed via theory or empirical data (i.e., parameterization) to optical properties (and vice versa).

2. Spatial resolution: Location and spatial resolution of the aerosol measurements are different. In-situ observations are often point measurements, while the swath width of passive satellite sensors can extend over up to a few thousand kilometers. In addition, active satellite sensors with narrow footprints often do not cover...
3. **Hygroscopicity:** The thermodynamic state of the air (especially the relative humidity, RH) has a strong effect on the aerosol optical properties (particularly in the lower marine troposphere) and is different for the different observations. Remote sensing of aerosols is normally performed at ambient condition (i.e., within the atmosphere), while most in-situ instruments dry the probed air during the sampling process before the aerosols are characterized (WMO, 2003).

4. **Temporal resolution:** The time periods over which the observations are averaged may be various. Short temporal averages (i.e., few hours) complicate a comparison since such an effort is only meaningful, when the different sensors actually observe the same air mass. Long-term averages (i.e., monthly means) on the other hand can generate arbitrary coherence of the data – especially when the considered data sets are of different size.

It is necessary to utilize these simultaneous but disparate data to be able to perform a closure study for the validation of remote-sensing data with independent in-situ measurements and vice versa. Such closure studies are not only important for validating the retrievals of aerosol optical thickness (AOT) or the aerosol extinction coefficient but also to investigate how the measured quantities are apportioned to different types of aerosol, e.g., how large the anthropogenic influence is on the optical properties of the atmosphere, and thus, the radiation balance. For this we have to be able to demonstrate that the measurement systems actually are sensing the same entity. The practical reality (i.e., it is not a simple matter to combine the in-situ and satellite data) is made into a doable but challenging task by the recognition at the outset that both the spaceborne and the in-situ instrument are well-tested devices that are operating correctly within the scope of their capabilities. Thus, the effort described here is not the usual ground truth
sort of activity done in order to constrain measurement uncertainties. We rather intend
to devise methods to bring the data sets into concordance.

Here, we consider in-situ measurements performed at the Arctic station at
Mt. Zeppelin (78.92° N, 11.85° E, 474 m a.s.l.), Svalbard, in comparison with data taken
simultaneously (or nearly so) with the Cloud-Aerosol Lidar with Orthogonal Polarization
(CALIOP) aboard the Cloud-Aerosol Lidar and Infrared Pathfinder Satellite Observa-
tions (CALIPSO, Winker et al., 2009) satellite. CALIPSO is operating in near-polar orbit
at an altitude of about 705 km.

In-situ instruments usually measure aerosol properties under dry conditions with
a RH of 10–30 % in an indoor laboratory, while ambient conditions are usually asso-
ciated with much higher RH of up to 100 %. Hence, in-situ measurements need to be
transformed to ambient conditions by means of direct RH-dependent measurements
or a microphysical particle model to account for the loss in particle size due to drying
the aerosol particles (Tang and Munkelwitz, 1994; Tang, 1996; Zieger et al., 2013). On
the other hand, ambient aerosol extinction coefficients can be measured directly for
instance with active optical remote-sensing techniques such as lidar or differential op-
tical absorption spectroscopy (DOAS). Previous closure studies show that reasonable
agreement is found between results obtained from remote sensing of aerosols and find-
ings from in-situ observations when the effect of relative humidity has been accounted
for (Hoff et al., 1996; Masonis et al., 2002; Zieger et al., 2011, 2012; Hoffmann et al.,
2012; Ziemba et al., 2013; Skupin, 2013). However, studies in the literature mainly deal
with few single cases during intensive field campaigns rather than systematic compar-
isons of multi-year data sets.

The clean environment of the Arctic is very sensitive to anthropogenic impacts. Arctic
aerosol conditions are also strongly influenced by regional meteorology (Eneroth et al.,
2003), which controls the RH of the air. Changes in this parameter have a huge influ-
ence on aerosol particle size, and thus, on light scattering (Zieger et al., 2010, 2013)
and cloud formation (Mauritsen et al., 2011) in this region. Measurements with a hu-
midified nephelometer were carried out between 15 July 2008 and 12 October 2008
at Zeppelin station (Zieger et al., 2010). A comparison to Zeppelin’s dry nephelo-
ter (operating at RH < 20%) showed that the ambient 550 nm scattering coefficients at RH = 85 ± 2% were on average about three times higher (scattering enhancement factor $f(RH) = 3.24 ± 0.63$) than the ones of the dried aerosol sample (Zieger et al., 2013). Optical properties and concentrations of Arctic aerosols have been measured at Ny-
Ålesund, Svalbard, with in-situ instruments (Covert and Heintzenberg, 1993; Ström et al., 2003; Tunved et al., 2013) and by means of remote sensing (Herber et al., 2002; Hoffmann et al., 2009, 2012; Tomasi et al., 2007, 2012) for several years.

Hoffmann et al. (2012) performed a combined analysis of ground-based Raman lidar measurement at Ny-Ålesund and in-situ measurements at Zeppelin station. Instead of the aerosol extinction coefficient they compare the particle number concentration as obtained from a microphysical inversion of the lidar data and measured by the in-
situ instruments. As the ground-based lidar data cannot be used to derive aerosol optical properties below 750 m height accurately, measurements at Zeppelin station (474 m height) were instead compared to lidar findings obtained at a height of 850 m. Despite the elaborate comparison approach (e.g., different heights, assumptions in the inversion of lidar data) agreement of a factor of ca. two was found in the total aerosol number concentration for the investigated pollution event on 4 April 2009 with smaller lidar-derived values compared to the in-situ measurements.

The use of the spaceborne CALIPSO lidar has the potential to overcome the altitude limitations since its observations extend all the way down to the Earth’s surface. The high frequency of overpasses at high latitudes makes it attractive to consider the possibility of a combined analysis of ground-based in-situ and spaceborne lidar measurements in the Arctic. In principle, such an analysis connects information on the vertical and horizontal aerosol distribution from the CALIPSO satellite data to the more specific information about aerosol microphysical and chemical properties at the surface. In-situ measurements are quite limited to a few measurement locations, while satellites can (in principle) view the exact same volume of air that is being sampled at the surface. Satellite sensors also have vastly larger fields of view and allow for global
Di Pierro et al. (2013) used these advantages to perform a comprehensive study of the spatial and seasonal distribution of Arctic aerosols based on optical properties observed by CALIOP between 2006 and 2012. The authors introduce an empirical correction that accounts for the different measurement sensitivity during day and night – a crucial factor when it comes to summertime CALIOP observations in the Arctic. The authors found CALIOP aerosol extinction in the Arctic to be of the same order of magnitude as nephelometer observations at Barrow and Alert with the latter being transformed to ambient RH. However, in addition to using highly averaged data (i.e., monthly and seasonal mean values) the averaging methodology of Di Pierro et al. (2013) applies a detection frequency that is defined as the ratio of the number of height bins with detected aerosol layers to the total number of height bins in a given area and time period. This procedure is likely to decrease the magnitude of the obtained mean extinction profiles by introducing zero-values to the averaging. In fact, the authors show that the mean CALIOP extinction profile obtained for a comparison to measurements with a high spectral resolution lidar (HSRL) at Eureka yields much smaller values than the ground-based HSRL observations. Di Pierro et al. (2013) also provide the readers with the seasonal variation of CALIOP-derived mean extinction coefficients for different atmospheric layers. Their values for the layer from the surface to 2 km height are a maximum at around 10 Mm$^{-1}$ in March for the Atlantic sector that is most representative for the conditions at Svalbard. This relates to a maximum AOT of 0.02 for the polluted spring season if we assume that the majority of aerosols is present within this 2 km deep layer. Such a value is similar to what is observed in the Arctic troposphere around Svalbard during the clean summer season (Glantz et al., 2014). Note that it is more likely that the aerosol-containing planetary boundary layer at Svalbard is between 0.5 and 1.0 km deep – which would decrease the maximum AOT as derived from...
the values presented in Di Pierro et al. (2013) even further. This discrepancy calls for a more detailed investigation of the factors that influence the reconciliation of extinction coefficients from ground-based and spaceborne observations. We will return to this point in the conclusion.

A description of the instrumentation and the data processing used in this study is presented in Sect. 2. Section 3 describes the methodology for relating segments of individual CALIPSO overpasses to in-situ measurements at Zeppelin station. The findings of the comparison for the year 2008 are discussed in Sect. 4. The paper ends with a summary and conclusions in Sect. 5.

2 Instrumentation and methods

2.1 In-situ measurements at Zeppelin station

The aerosol in-situ instruments at Zeppelin station include a differential mobility particle sizer for measuring the particle size distribution in the diameter range from 10 to 790 nm (time resolution of 20 min) and an integrating nephelometer (TSI model 3563) for measurements of particle light scattering coefficients at the wavelengths of 450, 550, and 700 nm (time resolution of 10 min) (Ström et al., 2003; Tunved et al., 2013). The nephelometer measurements were corrected for the truncation error and lamp non-idealities according to Anderson et al. (1996). All in-situ instruments are placed indoors and connected to an inlet without a particle size cut.

The location of the Zeppelin station at 79° N imposes a severe climatic situation with usually low outside temperature (from −25 to +15 °C) and correspondingly high RH, often near or at 100 %. On the other hand, the in-situ instruments in the laboratory are operated at ordinary room temperature of about 20 °C. Hence, sampled air is heated by as much as 40 K during its transit into the laboratory. Continuous aerosol in-situ observations are usually performed at dry conditions with RH < 30–40 % in order to avoid the influence of water uptake on the aerosol optical properties and to keep the
measurements at different ambient RH and at different sites comparable (WMO, 2003). However, the humidity effect on the scattering properties of the aerosol has to be accounted for if results are to represent actual atmospheric conditions.

Measurements with a humidified nephelometer operating at RH between 20 % and 95 % were carried out between 15 July 2008 and 12 October 2008 at Zeppelin station (Zieger et al., 2010). A comparison to Zeppelin’s dry nephelometer (operating at RH < 20 %) showed that the ambient scattering coefficients at RH = 85 % were on average about three times higher than the scattering coefficients of the dried aerosol sample. Direct measurements of the scattering enhancement factor were only available for 4 months in 2008.

A high-volume sampler with a PM$_{10}$ inlet was used to obtain the chemical composition of the Arctic aerosol with time resolutions of one day for sulfate and sea salt and one month for OC/EC during 2008.

### 2.2 Transferring dry measured optical parameters to ambient conditions

The aerosol extinction coefficient for ambient conditions is obtained with the help of a hygroscopicity model that uses measurements of outdoor humidity (hourly values), the dry particle number size distribution, and hygroscopicity information from measurements of aerosol chemistry. The model is in detail described in Rastak et al. (2014) and here only a brief description is being given.

The $\kappa$-Köhler theory (Kreidenweis et al., 2005; Petters and Kreidenweis, 2007) is used to account for hygroscopicity effects and to transform the dry particle size distribution to ambient conditions. The aerosol growth factor is derived by combining the individual aerosol volume fractions obtained from the analysis of chemical samples collected at Ny-Ålesund with the hygroscopicity parameter $\kappa$ of the respective components available in the literature. The components considered in this study are water-soluble and insoluble organics, sulfate, sea salt, and black carbon.

Aerosol scattering, absorption, and extinction coefficients are obtained from the humidified aerosol size distribution and refractive index by means of Mie-scattering theory.
Note that absorption contributes less than 7% to the ambient extinction coefficient of Arctic aerosols (Eleftheriadis et al., 2009; Zieger et al., 2010). Cases with ambient RH larger than 95% were considered to be measurements within clouds or fog, and thus, excluded from the procedure. All optical properties are calculated at a wavelength of 550 nm and with a temporal resolution of 1 h.

An extensive validation of the microphysical model also has been performed. A comparison of aerosol scattering coefficients measured by the dry nephelometer and calculated from the particle size distributions results in a slope close to unity and a squared correlation coefficient of $R^2 = 0.95$ (Rastak et al., 2014). A comparison between humidified scattering coefficients and measurements with the humidified nephelometer during the three months of parallel operation (Zieger et al., 2010) showed agreement with $R^2 = 0.64$, although with a slight tendency of the model to underestimate the measurements (Rastak et al., 2014). The enhancement factor $f(RH)$ is the ratio of ambient to dry extinction coefficients (Zieger et al., 2013). Values of $f(RH) = 4.30 \pm 2.26$ with a range from 1.5 to 12.5 were found for the year 2008. This is in agreement with the findings of Zieger et al. (2010) for Arctic aerosols at ambient RH at Zeppelin station.

The box plots in Fig. 1 visualize the importance of transforming dry optical properties to ambient conditions. About 75% of the hourly aerosol scattering coefficients at 550 nm measured with the dry nephelometer at Zeppelin station in 2008 are smaller than 5 Mm$^{-1}$. Humidity correction to ambient extinction coefficients increases the median value for 2008 from 2 to 7 Mm$^{-1}$ (numbers in the upper part of the figure). On average the ambient extinction coefficient is a factor of three to five larger than the dry one when resolved according to different seasons. The Arctic haze period in spring shows the highest median values of the ambient extinction coefficient (17 Mm$^{-1}$) followed by winter (8 Mm$^{-1}$). Summer and fall are associated with very low median values (3 and 4 Mm$^{-1}$, respectively). Summer is the slightly cleaner season and a larger variation is observed during fall. This is in agreement with previous observations at Zeppelin station (Ström et al., 2003; Zieger et al., 2010; Tunved et al., 2013).
The humidification of the particle number size distribution \( n(r) \) leads to an increase of the particle effective (surface-weighted) radius which is defined as

\[
    r_{\text{eff}} = \frac{\int r^3 n(r) dr}{\int r^2 n(r) dr}
\]

from 0.14 ± 0.02 to 0.23 ± 0.04 µm (yearly average, not shown). This moves much of the aerosol from an optically ineffective state to a size range in which they are very efficient in interacting with visible light.

### 2.3 CALIOP

The CALIOP is an elastic-backscatter lidar that emits linearly polarized laser light at 532 and 1064 nm wavelength and features three measurement channels. It has been operational since June 2006. An overview of the instrument as well as the data retrieval and interpretation algorithms can be found, i.e., in Winker et al. (2009); Young and Vaughan (2009), and Omar et al. (2009).

#### 2.3.1 Data treatment

For the comparison presented here we use level 2 version 3.01 products with a vertical resolution of 60 m (below 20.2 km height) and a horizontal resolution of 5 km. To derive extinction coefficients for comparison we only considered CALIPSO profiles with Atmospheric_Volume_Description bits 1–3 equal to 3 (feature type = aerosol), a CAD_Score below −20 (screen artifacts from data), and an Extinction_QC_Flag_532 of either 0 or 1. A description of the CALIPSO lidar level 2 5 km cloud and aerosol profile and layer products can be found in the CALIPSO Users Guide (2012).

Retrieving extinction coefficients from CALIOP observations requires the assumption of an aerosol-type specific extinction-to-backscatter (lidar) ratio (Müller et al., 2007; Omar et al., 2009). The CALIPSO aerosol model separates between six aerosol types...
that are selected according to the location of the instrument (surface type) and the detected feature (aerosol layer close to surface or elevated), the intensity of the measured signal (integrated attenuated backscatter coefficient), and an approximated value of the aerosol depolarization ratio (Omar et al., 2009). The considered aerosol types are: clean marine, dust, polluted continental, clean continental, polluted dust, and smoke. The lowest 532 nm lidar ratio of 20 sr is that of clean marine aerosol, while the highest values of 65 and 70 sr are used for polluted mineral dust, polluted continental aerosol, and biomass-burning smoke. Background conditions are described by the clean continental type that features a lidar ration of 35 sr. Lidar ratios of 30–40 sr at 532 nm are reported by Hoffmann et al. (2012) and Stock (2012) for two cases at Ny-Ålesund during spring 2009 and 2008, respectively. Proper aerosol-type identification is crucial for accurate extinction-coefficient retrievals due to the wide range of available lidar ratios (Müller et al., 2007). Details regarding the CALIPSO lidar-ratio selection algorithm are presented in Omar et al. (2009).

2.3.2 Representativeness

To assess the representativeness of the CALIOP measurements in our region of interest around Svalbard it is worthwhile to first examine the availability of lidar profiles and the atmospheric conditions (i.e., the abundance of aerosols and clouds) encountered during these observations. Figure 2a shows the number of monthly available lidar profiles subdivided according to what has been detected in the individual profiles: no features (neither clouds nor aerosols), only aerosols (aerosol features but no cloud features in a profile), only clouds (cloud features but no aerosol features in a profile), or clouds and aerosols (both cloud and aerosol features in a profile). For the entire year 2008, only 5.8% of the considered 187 711 profiles show conditions of aerosols only (i.e., no disturbance by clouds) that are most favorable for the type of comparison that we pursue in this study. Best conditions are found during March (15.1% cloud-free profiles with aerosols features) while the summer month (May to September) and particularly July (0.6% cloud-free profiles with aerosol features) represent non-ideal
conditions for the comparison of surface measurements and spaceborne observations attempted in this study. About 10% of all CALIOP profiles contain neither aerosol nor cloud features with a maximum and minimum occurrence rate of 25% and 4% in July and January, respectively. This effect is due to the weaker signal-to-noise ratio (SNR) of CALIOP measurements during bright daytime conditions (i.e., polar summer) compared to the absence of sunlight during night and the correspondingly higher threshold value that has to be exceeded for feature detection (Winker et al., 2009; Young and Vaughan, 2009). Polar summer and winter can be recognized in the occurrence rate of no features (magenta bars) in Fig. 2a. Observation rates of 50% to 85% for clouds only (during March and August, respectively) illustrate that cloudiness is another main obstacle for deriving aerosol information from CALIOP measurements. Most of these clouds are optically thick and lead to significant or full attenuation of the laser light. As long as these clouds form the uppermost feature, no aerosol detection is possible even if cloud and aerosol layers are present at different height levels.

Figure 2b shows the occurrence rate of the number of height bins with aerosol information for profiles that fall into the categories aerosol only and clouds and aerosols (i.e., profiles identified to contain aerosol information). Note that the information given in Fig. 2a refers to the entire profile while Fig. 2b refers to the height-resolved observation provided by these profiles. Figure 2b shows that the detection rate of aerosol bins (i.e., the amount of aerosol-containing height bins per profile per month) is much higher during winter, when the background of sunlight is absent and clouds are also less frequent (Fig. 2a). During summer, almost no aerosol features are detected. This is probably due to the decreased SNR of the measurement during daytime, the generally cleaner conditions during this time of the year, or a combination of both. It is also apparent from Fig. 2b that most aerosol features are detected in combination with clouds in the same profile (red) rather than during cloud-free conditions (green). A view at the number of detected aerosol layers given in the CALIPSO products reveals that aerosols are restricted to a single layer during the majority of observations (not shown). Multiple
aerosol layers are restricted to polar night. The observation of two layers is already rare while the number of cases with four layers is negligible.

Summarizing Fig. 2, we can conclude that obtaining useful results from CALIOP measurements in the Arctic during summer is improbable and that only a very small fraction of all measurements will occur during cloud-free conditions that favor the kind of study we attempt to perform in this paper. Attempts to overcome the limitations of CALIOP observations during Arctic summer as of Di Pierro et al. (2013) who introduced a detection rate for correction are likely to produce incorrect data or will at the least overemphasize the few data available during summer. Nevertheless, it is worthwhile to proceed with our study for the limited number of available cases in order to assess the value of the combined data sets.

2.4 HYSPLIT trajectories

We use the HYSPLIT (Hybrid Single-Particle Lagrangian Integrated Trajectory, Draxler and Rolph, 2010) model of NOAA Air Resources Laboratory to study the advection of air parcels to and from the Zeppelin station. Forward and backward trajectories with time intervals of 1 h were calculated starting and arriving every 3 h at the height and location of the Zeppelin station, respectively.

Meteorological parameters are provided along the trajectories and used in this study to estimate RH at the location of the CALIPSO overpass.

3 Comparison approach

Anderson et al. (2003) and Kovacs (2006) investigated the regional representativeness of local measurements of atmospheric aerosols by correlating these to the distance at which coincident satellite observations were performed. They concluded that the distance at which two measurements, both at ambient RH, along a trajectory show
acceptable correlation to establish a connection are 300 and 500 km for observations over land and sea sites, respectively.

In 2008, CALIPSO passed 2018 times over the area from 0 to 25° E and from 75 to 82° N that is considered in this study. The closest overpass occurs only 2 km away from the Zeppelin station, while the furthest one was at a distance of 360 km. Differences in exact location of the measurements pose a severe problem, since the humidity and aerosol content of air is highly variable in time and space (horizontally and vertically). Thus it is essential to select that part of the CALIPSO ground track for which it is most likely that both CALIOP and in-situ instrumentation actually sampled the same air mass. Following the approach presented in Tesche et al. (2013), air-mass trajectories are used to connect the in-situ station to the segment of the CALIPSO ground track that is most suitable for comparison. The length of the trajectories between Zeppelin station and the intersection with the CALIPSO ground track provide us with the time lag between fitting observations.

In a first step of screening the CALIPSO data, we selected only those overpasses that actually show extinction coefficients (i.e., signals above the CALIOP detection threshold) in a height range from 250 to 730 m that spans around the height of the Zeppelin station. This holds for 24% of all overpasses in the area of interest. For these cases, we investigated whether backward and forward trajectories starting every 3 h for 15 h after and before the CALIPSO overpass are actually crossing the ground track (second step). Cases with no such intersections were discarded from the investigation. This left 9% of all 2018 overpasses in 2008. Note that in contrast to the studies by Anderson et al. (2003) and Kovacs (2006) that referred to the lengths scale we use a time scale and restrict the comparison to a time delay of 15 h. This corresponds to a maximum distance of 360 km at a mean transport velocity of about 7 m s$^{-1}$. We believe that time rather than distance is a better parameter to assess changes in the aerosol properties in the atmosphere. The majority of the track segments for comparison were located either in the vicinity or to the north (beyond 81° N) of the ground site (not shown).
In the third and final step, we checked for the availability of (1) CALIOP extinction coefficients at the intersection of satellite ground track and air-mass trajectories and of (2) humidified extinction coefficients at Zeppelin station at the time of the CALIPSO overpass plus/minus the lag provided by the trajectories. That was the case for only 57 individual overpasses (3% of all 2018 overpasses) in 2008, which form the core of this study. The extinction coefficients from CALIOP were averaged in the vicinity of the crossing point of the ground track and the trajectory. The along-track averaging range was determined individually for each overpass according to the spread of the crossing trajectories. A change in the along-track average of the CALIOP extinction profile (i.e., from a range related to crossing trajectories with different starting time at the location of the ground site to a fixed interval) can result in large differences of the resulting mean extinction profile. Once an extinction profile could be obtained at the proper location for comparison, the values in the height range from 250 to 730 m (eight 60 m height bins) were averaged eight 60 m height bins. We chose this height range to account for vertical motions during the transport from the location of the CALIOP observation to Zeppelin station (backward trajectories) or the other way round (forward trajectories). Better agreement with the in-situ observation may be obtained for an average over a smaller height range. However, we chose a conservative range that is likely to be suitable for most cases. The average and the corresponding standard deviation (as a measure of vertical homogeneity) represent values used in the comparison to the findings of the measurements at Zeppelin station. To coarsely account for uncertainties in the trajectories, in-situ extinction coefficients were averaged over 5 h (five 1 h values) centered around the time during which the in-situ instruments sampled the same air parcels as CALIOP, i.e., time of a CALIPSO overpass plus the time lag determined from the length of the trajectories that connect this overpass to Zeppelin station.
4 Results and discussion

The time period from 22 to 28 January 2008 has been chosen to illustrate the analytical work and some of the results obtained. Figure 3 presents the dry scattering coefficient measured with the nephelometer at Zeppelin station and the ambient extinction coefficient calculated from the humidified size distribution during this period. The ambient RH given in the figure reflects the influence of hygroscopicity which causes the huge differences between dry scattering and ambient extinction values. The latter parameter has not been estimated when ambient RH exceeded values of 95%. The time period covered in Fig. 3 shows ten CALIPSO overpasses that were connected to the ground station with the help of trajectories (see colored triangles and corresponding numbers at the top of Fig. 3). Extinction coefficients extracted from the CALIPSO observations could be compared to ground-based measurements for six cases (overpasses 1, 2, 3, 6, 8, and 9). Four examples of how trajectories are used to connect the ground site with the proper segment of the CALIPSO track (overpasses 1, 6, 8, and 10) are given in the lower part of Fig. 3. Green triangles mark cases for which aerosol profiles were obtained during cloud-free conditions as indicated by a cloud optical thickness (COT) of zero. The examples of overpasses 1 and 8 show how the trajectories lead to a cloud-free part of the ground track. The different lengths and tracks of the trajectories indicate that time and distance should not be considered as synonyms. The satellite- and ground-based extinction coefficients agree within factors of 1.1–1.3 for the overpasses on 22 and 27 January 2008 with the shortest time delay of 6 h (201 km distance) and the longest time delay of 15 h (322 km distance). Note that ambient RH was above 90% on 22 January 2008 and that the difference between the dry scattering coefficient and the RH-corrected extinction coefficient is as much as a factor of 10. A much smaller ratio of ambient to dry extinction coefficients can be found for 26 and 27 January 2008, for which RH varies between 65% and 90%. The green cases in Fig. 3 illustrate the importance of accounting for the proper time delay between the
measurements of CALIOP and in-situ instrumentation. Using the in-situ measurements at the time of the satellite overpass decreases the agreement of the observations.

Using the trajectories as described above, a cloudy part of the CALIPSO ground track (COT > 0, AOT = 0) was identified for the overpasses 4, 5, 7, and 10. No comparisons could be performed since there is no aerosol information available for these cases. This kind of situation inhibited comparisons in 127 cases for the months January to April and October to December 2008. Typical scenarios are: no height bins are marked as containing aerosols at all, all aerosols are located above or below our height range of interest, or the obtained aerosols profile is of unreasonable shape and/or magnitude.

However, for overpass 6 (blue triangle in Fig. 3) aerosol information was obtained in cloudy environment (COT > 0, AOT > 0). Even though this overpass occurred only 21 km from the ground site, the CALIPSO observation is in poor agreement with the result of the in-situ measurement. This emphasized that using a closest approach for comparison of ground-based measurements and CALIPSO observations might not always be the best choice. The case also illustrates that even few clouds can disturb aerosol measurements with spaceborne lidar. Note also that trajectories might actually lead to a track segment that is not closest to the ground site, as is the case for overpass 8.

Finally, 57 cases of the 2018 overpasses in 2008 were suitable for comparing extinction coefficients from CALIOP observations and humidified ground-based measurements (Fig. 4). Even though CALIOP extinction coefficients are generally larger than the ones derived from the in-situ measurements, most comparisons agree to a factor of one to five with a majority not exceeding a factor of two. This is a surprisingly good finding considering the data processing that is necessary to come up with comparable quantities. There is no indication that a closer distance between satellite ground track and in-situ ground site (or a smaller time lag, not shown) would give a better agreement. Suitable agreement actually occurs for many cases associated with overpasses at larger distances from the ground site. These cases would not have been included in
this study if we had chosen a distance in range rather than time for comparison. In fact, Fig. 4 shows that most of the cases exceeding a factor of five in agreement correspond to comparisons at distances of 120 km or less (blue, light blue, and light green dots in the upper left part of the plot). This suggests that the method of comparing local point or column-integrated measurements to the closest-approach observation of CALIPSO is likely to yield misleading results.

We performed a deeper analysis of the factors that could explain why a difference of as large as a factor of five occurs for some of the cases included here. Besides the spatial distance and temporal delay between the observations we considered the relative humidity at Zeppelin station and at the crossing point of the satellite ground track and trajectories, the occurrence of clouds and rain along the trajectory, and the wind direction at the ground site. However, only the latter parameter could be linked to the outliers in Fig. 4. Figure 5a shows that the largest absolute difference in the ambient extinction coefficients from CALIOP and in-situ measurements occur during westerly flow. It could be that aerosol conditions are more stable for air masses approaching Zeppelin station from the north and via ice-covered ocean compared to the open water to the west. In addition, we investigated the dominant aerosol type selected in the CALIPSO data retrieval for the individual comparisons. It was found that the most characteristic outliers in Figs. 4 and 5a occur for cases that were identified predominantly as polluted dust, polluted continental, and dust in the CALIPSO retrieval. These aerosol types are rather uncommon at 78° N and suggest misclassification in the CALIPSO retrieval. Misclassification can occur as a result of signal noise or due to surface effects.

Clean continental aerosol (i.e., background conditions) was classified for most comparison cases (see color coding in Fig. 5a) and seems to be the most appropriate choice of aerosol together with clean marine. It remains unclear, why half of the clean marine cases are within the set of outliers. In addition, classifying aerosol features as polluted dust or smoke (lidar ratio of 65–70 sr) instead of clean continental aerosol (lidar ratio of 35 sr) will only result in a factor of two difference, while the disagreement we obtain in our comparison for cases classified as something other than clean continental
shows factors in the range from 6 to 18. The range is 0.38 to 5 for cases classified as clean continental.

Strong variation in RH between the location of the CALIPSO ground track and Zeppelin station could also cause the scatter of values presented in Fig. 4. Such RH differences have a direct effect on the scattering enhancement factor $f(\text{RH})$, and thus, on the difference between dry and ambient extinction coefficients. The scattering enhancement factor was found to be much higher for Arctic aerosol compared to observations at continental, background, or marine sites (Zieger et al., 2013). Consequently, we should expect that even small differences in RH between the measurements at Zeppelin and along the satellite track can lead to high differences in the ambient extinction coefficient. This holds especially for high RH $> 85\%$. We investigated if we can find a connection between the difference in RH ($\Delta \text{RH}$) at the two measurement locations (i.e., the CALIOP ground-track segment and Zeppelin station) and the agreement in the comparison of ambient extinction coefficients at those sites. The RH at the location of the CALIOP observation is taken from the meteorological data provided with the trajectory analysis and thus highly uncertain. For the considered 57 cases, the $\Delta \text{RH}$ showed a mean value of $12 \pm 10 \%$ (mean RH of $80 \pm 12 \%$ at Zeppelin station) with a maximum value of around 30 % (not shown). Though $\Delta \text{RH}$ was considerable for several cases, we could not establish that this factor or the resulting difference in $f(\text{RH})$ can fully explain the disagreement found in the ambient extinction coefficients. Figure 5b shows the connection between the relative difference in $f(\text{RH})$ at the locations of CALIOP and in-situ observations and the relative difference in the ambient extinction coefficients obtained from these observations. Values should align along the 1 : 1 line, if hygroscopic growth was the only factor we would have to consider in our comparison. Deviations are likely to be related to the observation of different air masses at the two locations (i.e., different aerosol size distributions or chemical composition) or the improper representation of meteorological parameters (i.e., RH) in the trajectory model.

Table 1 gives a detailed overview of the results obtained from the comparison of spaceborne and ground-based observations subdivided according to the months of
2008 and to whether cloud-free or cloudy CALIOP aerosol profiles were used in the comparison. For the 57 considered cases Table 1 shows that time delay is rather evenly distributed between 0 and 15 h with a median of 8 h. 39 of the 57 suitable cases occurred during most favorable cloud-free conditions (AOT > 0, COT = 0), while the remaining 18 cases represent cloudy comparisons (AOT > 0, COT > 0). Resolving the comparison according to cloudiness in the CALIPSO observations (not shown) leads to ambiguous results: for 7 of the 18 cloudy cases (39 %) a difference larger than a factor of two is found between the extinction coefficients from CALIOP and Zeppelin station, while for the cloud-free cases 17 out of 39 (44 %) exceed this difference. The average time delay is $9.2 \pm 3.8$ h for cloud-free cases, while it is only $6.2 \pm 3.9$ h for cloudy cases. Accordingly, cloud-free cases show a mean distance of $228 \pm 100$ km and cloudy ones $178 \pm 116$ km. Extinction coefficients from CALIPSO vary between 4.6 and 127.0 Mm$^{-1}$ for cloud-free cases, while the range of values for cloudy profiles is much narrower and only spans from 14.3 to 91.6 Mm$^{-1}$.

5 Summary and conclusions

This study presents a comparison of extinction coefficients as determined from space-borne lidar (CALIOP) measurements and from ground-based in-situ measurements at Zeppelin station, Ny-Ålesund, Svalbard, during the year 2008. To obtain meaningful comparison, we had to consider several issues:

1. Neither in-situ instruments nor spaceborne lidar (CALIOP) provide us with direct measurements of the ambient aerosol extinction coefficient.

2. Approved methods were used to obtain ambient extinction coefficients from dry in-situ measurements of the particle size distributions in combination with a multi-component growth factor based on $\kappa$-Köhler theory and measured chemical aerosol properties.
3. Extinction coefficients from the spaceborne sensor were taken from operational CALIPSO products that underwent elaborate calibration and quality assurance.

4. Air-mass trajectories were used to ensure that all measurements compared were performed within the same air mass. This is necessary to establish a connection between the satellite’s ground track and Zeppelin station and to adapt the averaging intervals along the CALIPSO ground track according to the spatial spread of the crossing trajectories. The averaging height range of 510 m was centered at the elevation of the ground site and chosen to account for vertical displacement during travel along the trajectories. Temporal averaging of ground-based data of 5 h was introduced to further mitigate imprecision in the trajectory output.

The detailed matching procedure used in this study reduced the number of comparison cases from over 2000 overpasses in 2008 to 57 overpasses during 42 days of that year. Even though it is a costly and elaborate case-by-case comparison it is likely to yield more significant results than what is obtained by comparing monthly means of surface measurements with monthly regional means of CALIOP observations. However, since averaging times of only a few hours were applied in this study, we cannot draw conclusions about what will happen if the length of the temporal averaging window is increased. The median ambient extinction coefficient for the 57 comparison cases was 27.8 Mm$^{-1}$ for the CALIOP data compared to a value of 14.3 Mm$^{-1}$ for in-situ measurements that were corrected to ambient conditions. The different humidity during the measurement in the atmosphere and within a laboratory is an omnipresent limitation for studies like the one presented here. The thermodynamic state (e.g., RH) of the samples and the assumptions on the hygroscopic properties for the in-situ measurements are therefore vital factors for a successful comparison of aerosol extinction coefficients. In the case of our study, results are also influenced by the CALIPSO aerosol model that is required for the extinction-coefficient retrieval, the CALIOP feature detection limit, and the criteria that are used to match satellite observations to the measurement at the ground site.
Detailed knowledge of the humidity field is of vital importance when relating in-situ measurements to observations with spaceborne sensors. The effect of relative humidity on the light scattering properties of aerosol particles in the atmosphere is the dominating obstacle for a systematic reconciliation of measurements of the two platforms. Additional disturbing factors in the allocation procedure applied in this study were unfavorable wind direction (no intersection between trajectories and ground track), presence of clouds (RH > 95% at Zeppelin station and/or no aerosol information from CALIOP), no data from Zeppelin station or CALIOP, and the CALIOP detection threshold that prevents reliable aerosol detection in the presence of sunlight. CALIOP detects almost no aerosol features in the Svalbard region during Arctic summer even though the tropospheric median AOT is generally larger than 0.05 at visible wavelengths during May and June (Tomasi et al., 2007, 2012; Glantz et al., 2014). This is in agreement with a study by Di Pierro et al. (2013) that investigated the distribution of aerosols in the Arctic from CALIOP measurements. Consequently, CALIOP data have to be treated with great caution when they are used for studies of aerosol occurrence rate, transport patterns, radiative effects, and interactions with clouds under background conditions during polar day.

Based on the study presented here we also conclude that consolidating data sets that are averaged over large areas and/or long time periods can lure us into a feeling of arbitrary confidence, while there may actually be weak or no connection between individual observations. Using highly averaged parameters in the deduction of scientific findings is of particular importance for the validation of model simulations. Consequently, special emphasis should be placed on a proper selection of temporal and spatial averaging intervals when attempting to use spaceborne lidar observations in connection to ground-based measurements and model outputs.

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**Table 1.** Results of the comparison of CALIPSO observations and in-situ measurements at Zeppelin station subdivided according to months of the year 2008 and to cloud-free and cloudy conditions in the CALIPSO aerosol profiles. The first line (columns 3–7) refers to mean values and standard deviation, while the second line refers to median and range of values.

<table>
<thead>
<tr>
<th>month of cases</th>
<th>distance (km)</th>
<th>delay (h)</th>
<th>Extinction Coefficient (Mm$^{-1}$)</th>
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<tr>
<td></td>
<td></td>
<td>in-situ</td>
<td>CALIPSO</td>
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<tr>
<td>Jan</td>
<td>223 ± 112</td>
<td>8.9 ± 4.5</td>
<td>15.6 ± 7.5</td>
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<td></td>
<td>271, 21–343</td>
<td>9.5, 1.0–15.0</td>
<td>17.6, 1.2–27.8</td>
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<tr>
<td></td>
<td>288, 2–357</td>
<td>9.0, 0.0–15.0</td>
<td>12.8, 3.0–18.0</td>
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<tr>
<td>Feb</td>
<td>251 ± 110</td>
<td>7.8 ± 4.3</td>
<td>12.0 ± 4.4</td>
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<td>252, 44–360</td>
<td>9.0, 3.0–15.0</td>
<td>17.4, 4.0–43.0</td>
</tr>
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<td>Mar</td>
<td>223 ± 111</td>
<td>9.8 ± 4.1</td>
<td>21.2 ± 12.1</td>
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<tr>
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<td>203, 69–352</td>
<td>9.0, 3.0–13.0</td>
<td>27.7, 13.8–95.9</td>
</tr>
<tr>
<td>Apr</td>
<td>216 ± 104</td>
<td>8.5 ± 3.6</td>
<td>35.9 ± 27.5</td>
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<tr>
<td></td>
<td>203, 69–352</td>
<td>9.0, 3.0–13.0</td>
<td>27.7, 13.8–95.9</td>
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<tr>
<td>Oct</td>
<td>292 ± 40</td>
<td>7.5 ± 0.7</td>
<td>6.0 ± 1.7</td>
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<td></td>
<td>292, 263–320</td>
<td>7.5, 0.0–8.0</td>
<td>6.0, 4.8–7.2</td>
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<td>Nov</td>
<td>128 ± 66</td>
<td>5.9 ± 3.2</td>
<td>16.3 ± 24.6</td>
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<td>107, 23–226</td>
<td>5.5, 1.0–12.0</td>
<td>7.1, 1.6–75.7</td>
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<tr>
<td>Dec</td>
<td>106 ± 33</td>
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<td>106, 82–129</td>
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<td>all year</td>
<td>212 ± 107</td>
<td>8.2 ± 4.0</td>
<td>18.8 ± 17.8</td>
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<td>242, 2–360</td>
<td>8.0, 0.0–15.0</td>
<td>14.3, 1.2–95.9</td>
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<td>cloudy</td>
<td>178 ± 116</td>
<td>6.2 ± 3.9</td>
<td>23.0 ± 25.2</td>
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<td>169, 2–323</td>
<td>5.5, 0.0–13.0</td>
<td>14.7, 1.2–95.9</td>
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<tr>
<td>cloudfree</td>
<td>228 ± 100</td>
<td>9.2 ± 3.7</td>
<td>16.9 ± 13.0</td>
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<td>247, 23–360</td>
<td>9.0, 1.0–15.0</td>
<td>14.3, 1.6–74.9</td>
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Fig. 1. Statistical overview of the dry scattering (red) and ambient extinction (green) coefficients at 550 nm based on hourly measurements at Zeppelin station in 2008 according to the entire year and the different seasons winter (DJF), spring (MAM), summer (JJA), and autumn (SON). The numbers in the top of the figure mark the number of available hourly measurements, mean, standard deviation, and median values for an arithmetic mean of the data, as well as the mean and standard deviation for a geometric mean of the data (gray circles) for the different time periods. The geometric mean has a much lower standard deviation than the arithmetic mean and is similar to the arithmetic median value. The difference in data availability for dry scattering and ambient extinction coefficients is the consequence of cloud screening and an absence of input data required for humidity correction.
Fig. 2. Histograms of the monthly abundance of (a) CALIOP level 2 5 km aerosol profiles and (b) 60 m height-bins with aerosol observations as detected during 2018 CALIPSO overpasses in the region of interest during 2008. The color coding in (a) refers to the observed occurrence of atmospheric features (aerosols and/or clouds). The number of detected aerosol-containing height bins per month in (b) is subdivided according to whether clouds were absent (green) or present (red) in individual lidar profiles marked as aerosol only or clouds and aerosols in (a).
Fig. 3. Upper panel: CALIPSO extinction coefficient (532 nm, green circles) compared to in-situ measurements of the dry scattering coefficient (550 nm, red line and dots) and the ambient extinction coefficient (550 nm, green line and dots) for the time period from 22 to 27 January 2008. Red circles mark 5 h averages of the ambient extinction coefficient. Arrows show which values are compared. Ambient RH is given as blue line and squares. Values above RH > 95% were disregarded (dashed blue line). Colored triangles and corresponding numbers mark CALIPSO overpasses that could be connected to the ground site for the considered time period. The color refers to the availability of aerosol information in CALIOP profiles in the track segment chosen for comparison: only aerosol features (green), aerosol and cloud features (blue), and no or only cloud features (red). Lower panel: presentation of the use of trajectories to connect the in-situ site to the spaceborne measurements for four selected cases (marked as 1, 6, 8, and 10 in the upper plot). The CALIPSO ground track is marked by gray (no aerosol data available) and green (aerosol data available) circles that refer to individual 5 km aerosol profiles. Colored dots and lines mark backward trajectories starting close to the CALIPSO overpass (red) as well as 3 h (green), 6 h (blue), 9 h (magenta), and 12 h (orange) after the overpass. The time of overpass is given in the respective plots. The red star marks the location of the Zeppelin station.
Fig. 4. Comparison of the humidified 550-nm extinction coefficient from the in-situ measurements vs. the ambient 532-nm extinction coefficient extracted from CALIPSO overpasses for 57 suitable cases. The color coding describes the distance of the CALIPSO observation from the ground site. Error bars represent the standard deviation from averaging over five values of hourly humidified in-situ measurements and nine 60 m CALIPSO height bins between 250 and 730 m, respectively. Ratios of 1:1, 1:2, and 1:5 are marked by solid and dashed lines and the shaded area.
Fig. 5. Detailed view of (a) the effect of wind direction on the absolute difference in the ambient extinction coefficients derived from observations at Zeppelin and by CALIOP and (b) the connection between the relative difference $\Delta f(RH)$ of the scattering enhancement factors at Zeppelin station and at the intersection of trajectories and CALIPSO ground track and the relative difference in the ambient extinction coefficients observed at the two locations. The color coding refers to the dominant aerosol type identified in the CALIOP observations (cm – clean marine; d – dust; pc – polluted continental; cc – clean continental; pd – polluted dust; s – smoke, not observed) and the difference of RH observed at Zeppelin station and taken from the trajectory calculations at the location of the CALIPSO overpass, respectively. Relative values are normalized to the observation at Zeppelin station. The dashed line marks the 1 : 1 line.