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Effects of long-range aerosol transport on the microphysical properties of low-level liquid clouds in the Arctic

Quentin Coopman^{1,2}, Timothy J. Garrett¹, Jérôme Riedi², Sabine Eckhardt³, and Andreas Stohl³

¹Department of Atmospheric Sciences, University of Utah, Salt Lake City, UT, USA

²Laboratoire d'Optique Atmosphérique, Université de Lille/CNRS, Lille, France

³Norwegian Institute for Air Research, Kjeller, Norway

Correspondence to: Quentin Coopman (quentin.coopman@ed.univ-lille1.fr)

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Abstract. The properties of low-level liquid clouds in the Arctic can be altered by long-range pollution transport to the region. Satellite, tracer transport model, and meteorological data sets are used here to determine a net aerosol–cloud interaction (ACI^{net}) parameter that expresses the ratio of relative changes in cloud microphysical properties to relative variations in pollution concentrations while accounting for dry or wet scavenging of aerosols en route to the Arctic. For a period between 2008 and 2010, ACI^{net} is calculated as a function of the cloud liquid water path, temperature, altitude, specific humidity, and lower tropospheric stability. For all data, ACI^{net} averages 0.12 ± 0.02 for cloud-droplet effective radius and 0.16 ± 0.02 for cloud optical depth. It increases with specific humidity and lower tropospheric stability and is highest when pollution concentrations are low. Carefully controlling for meteorological conditions we find that the liquid water path of arctic clouds does not respond strongly to aerosols within pollution plumes. Or, not stratifying the data according to meteorological state can lead to artificially exaggerated calculations of the magnitude of the impacts of pollution on arctic clouds.

1 Introduction

Due to growing concentrations of greenhouse gases and complex feedback processes, the Arctic region has warmed approximately 2 times faster than the global average (Serreze and Francis, 2006; Serreze et al., 2009; Richter-Menge and Jeffries, 2011), a trend that is anticipated to continue through this century (Yoshimori et al., 2013; Overland and Wang,

2013). Further, the Arctic is not pristine, even if it is remote from industrialized areas and major aerosol sources. Midlatitude aerosols can be transported to northern latitudes in relatively high concentrations when precipitation rates are low and there are strong temperature inversions that inhibit vertical mixing (Sirois and Barrie, 1999; Law and Stohl, 2007; Quinn et al., 2007; Law et al., 2014). The origins of arctic haze tend to be pollution from Eurasia (Shaw, 1995; Stohl, 2006; Shindell et al., 2008; Ancellet et al., 2014), and boreal forest fires in North America, Eastern Europe, and Siberia (Stohl, 2006; Stohl et al., 2007). Between spring and summer, the atmosphere becomes cleaner due to an increase in wet scavenging (Garrett et al., 2010).

Such aerosols have the potential to alter cloud properties in the Arctic (Garrett and Zhao, 2006; Lance et al., 2011). On one hand, thin low-level clouds with more numerous smaller droplets can radiate more longwave radiation thereby warming the surface (Garrett et al., 2002, 2004; Garrett and Zhao, 2006). On the other, polluted clouds can reflect more sunlight, leading to a cooling effect (Lubin and Vogelmann, 2007). Zhao and Garrett (2015) found that seasonal changes in surface radiation associated with haze pollution range from $+12.2 \text{ W m}^{-2}$ in February to -11.8 W m^{-2} in August. Annually averaged, the longwave warming and shortwave cooling nearly compensate, although the seasonal timing of the forcings may have implications for rates of sea ice melt (Belchansky et al., 2004; Markus et al., 2009).

The influence of aerosols on cloud microphysical properties is often quantified using an indirect effect (IE) or aerosol–cloud interaction (ACI) parameter that expresses the ratio of relative changes in cloud microphysical properties to

variations in pollution concentrations, most typically aerosol index, the aerosol optical depth, the aerosol concentration, or the cloud condensation nucleus (CCN) concentration (Feingold et al., 2001; Feingold; 2003b). Where the parameter is expected to decrease with increasing aerosols or cloud condensation nuclei (e.g., the effective radius), the ratio is multiplied by -1 so that the IE or ACI is positive.

Garrett et al. (2004) used ground-based retrievals of cloud-droplet effective radius and surface measurements of dried aerosol light scattering from Barrow (Alaska) to obtain a value for the cloud-droplet effective radius ACI that lies between 0.13 and 0.19. Satellite measurements show that ACI values for cloud-droplet effective radius range from 0.02 to 0.20 for midlatitude continental clouds (Nakajima et al., 2001; Feingold, 2003a; Lohmann and Feichter, 2005; Myhre et al., 2007) and from 0.03 to 0.15 for midlatitude oceanic clouds (Bréon et al., 2002; Sekiguchi, 2003; Kaufman et al., 2005; Myhre et al., 2007; Costantino and Bréon, 2013; Wang et al., 2014). Satellite instruments have the advantage of providing data over large spatial scales, however satellite retrievals of aerosol concentrations are normally obtained from air columns close to the analyzed cloud. The assumption is made that plumes are horizontally homogeneous both within and without the cloud, and that they are vertically collocated with the cloud top (Nakajima et al., 2001; Feingold et al., 2001; Sekiguchi, 2003). For large-scale cloud studies, this method potentially introduces bias since it is not obvious that pollution should be uniform for different meteorological regimes.

Colocating satellite cloud retrievals with pollution tracer output from a chemical transport model offers an alternative approach for assessing the effect of pollution on clouds (Berg et al., 2011; Lance et al., 2011; Tietze et al., 2011). Cloud microphysical properties and pollution concentrations can be estimated at the same time, location, and meteorological regime (Schwartz et al., 2002; Kawamoto et al., 2006; Avey et al., 2007). Active tracers experience both sources and sinks through wet scavenging, dry deposition, and chemical reactions that can be difficult to accurately model. Passive pollution tracers, on the other hand, are determined only by source emission strength and subsequent dilution. An example of a passive tracer is carbon monoxide (CO), which is a combustion byproduct that correlates with the anthropogenic CCN close to pollution sources (Longley et al., 2005). Since both are found in pollution plumes, CO can serve as a passive proxy for CCN that is relatively straightforward to model. A difference is that, unlike CCN, CO is unaffected by wet and dry scavenging. In the absence of scavenging, a linear relation exists between CCN and CO and it should be possible to see changes in clouds when CO concentrations are high. However, if CCN have been scavenged from pollution plumes, then the observed sensitivity of clouds to the pollution plumes should be expected to be low. Thus, passive tracers serve as an indicator of the net effect of pollution plumes on clouds that accounts for the effect

of scavenging during transport en route to the clouds themselves.

More generally, the primary controls on cloud microphysical properties are not aerosols but rather meteorological conditions during cloud formation (Chang and Coakley, 2007; Brenguier and Wood, 2009; Kim et al., 2008; Painemal et al., 2014; Andersen and Cermak, 2015). For example, a reduced stability of the environmental temperature profile can allow for enhanced cloud-droplet growth through increasing convection (Klein and Hartmann, 1993). This would be expected to lead to greater mixing of the aerosols with the cloudy air and greater aerosol impacts on cloud microphysical properties (Chen et al., 2014; Andersen and Cermak, 2015). Also, in the Arctic during the winter, pollution plumes from Asia are often associated with higher values of potential temperature than pollution plumes from Europe (Stohl, 2006). Thus, the observed impact of pollution plumes on clouds may be correlated with a particular meteorological regime.

Using the approach of collocating a passive tracer from a tracer transport model with satellite observations, Tietze et al. (2011) presented an analysis of pollution–cloud interactions over the Arctic from March to July 2008. Anthropogenic and biomass burning aerosol pollution was represented using a CO passive tracer in the FLEXPART (FLEXible PARTicle dispersion model) tracer model (Stohl et al., 2005) as a proxy. CO was colocated with POLDER-3 (Polarization and Directionality of the Earth's Reflectance) and MODIS (MODerate-resolution Imaging Spectroradiometer) observations. Tietze et al. (2011) showed that the sensitivity of liquid-cloud-droplet effective radius (r_e) and optical depth (τ) to pollution has a maximum around the freezing point, and that the sensitivity decreases for both higher and lower temperatures. The optical depth was generally up to 4 times more sensitive than the effective radius. Their results also suggested that biomass burning pollution has a smaller yet significant impact on liquid cloud microphysical properties than anthropogenic pollution, and that the ACI parameters depend on altitude, LWP, and temperature.

Our study extends the Tietze et al. (2011) research by adding 2 years of data, 2009 and 2010, and in addition to temperature, stratifying the results by lower tropospheric stability (LTS) and atmospheric specific humidity (Matsui et al., 2006; Mauger and Norris, 2007). Our results highlight the importance of considering meteorological conditions when assessing the aerosol impact on cloud microphysical properties to show that r_e and τ have similar sensitivities to pollution.

2 Data

The analyses in this study are based on a collocation of satellite retrievals of cloud properties, tracer transport model simulations of pollution locations and concentrations, and re-analysis data sets for meteorological fields.

2.1 Satellite cloud property retrievals

In this study we used two instruments, both part of the A-train mission (Stephens et al., 2002). The MODIS instrument on board the Aqua satellite measures radiation in 36 different spectral bands with central wavelength from 400 to 14 400 nm in wavelength. For the effective radius, optical depth, and cloud top temperature we use Collection 5 Level-2 products (Platnick et al., 2003; King and Platnick, 2006). Regarding the technique applied for computation of the MODIS Level-2 products cloud top temperature is derived from the 11 μm infrared band. Cloud-droplet effective radius (r_e) and optical depth (τ) are retrieved from simultaneous cloud reflectance measurements in three water-absorbing bands (1.6, 2.1, 3.7 μm) and three nonabsorbent bands (0.65, 0.86, 1.2 μm) (Platnick et al., 2003). The pixel resolution of the retrievals at nadir is 1 km \times 1 km for cloud microphysics and 5 km \times 5 km for cloud top temperature.

The POLDER-3 camera on the PARASOL satellite platform (Polarization & Anisotropy of Reflectances for Atmospheric Sciences coupled with Observations from a Lidar) captures a wide field of view through spectral, directional, and polarized measurements of reflected sunlight (Fougnie et al., 2007). Multidirectional observations allow for a pixel to be observed from up to 16 different view angles. The instrument measures radiance in 9 spectral channels between 443 and 1020 nm, including three polarized channels at 490, 670, and 865 nm. POLDER-3 cloud microphysical properties retrievals have a 36 km \times 36 km spatial resolution. Cloud top pressure is derived from the cloud oxygen pressure (Bréon and Colzy, 1999). Cloud top pressure retrieved by POLDER-3 appears to be a better proxy for low-level cloud height than MODIS cloud top pressure derived using a thermal signature assuming a given temperature profile (Buriez et al., 1997; Weisz et al., 2007; Tietze et al., 2011; Desmons et al., 2013).

To determine the cloud thermodynamic phase we use a combination of MODIS and POLDER-3 measurements. The algorithm takes advantage of multiangle polarization data, shortwave, thermal infrared, and visible measurements to retrieve a thermodynamic phase index Φ between 0 for liquid clouds and 200 for ice clouds with varying degrees of confidence (Riedi et al., 2010). Figure 1 shows the distribution of the thermodynamic phase index for clouds between 200 and 1000 m and between 1000 and 2000 m altitude from 2008 to 2010 over a region with latitudes greater than 65°. We observe different modes in the phase index corresponding to liquid clouds with Φ lower than 70, clouds with undetermined phase, mixed phase, or multiple cloud layers, for which Φ lies between 70 and 140, and ice clouds with Φ greater than 140.

2.2 Anthropogenic pollution tracer fields

For determining anthropogenic pollution tracer fields, we use the Lagrangian particle dispersion model FLEXPART

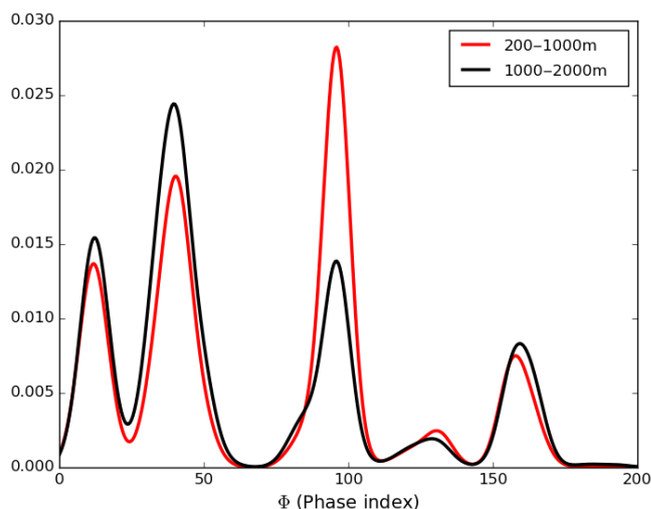


Figure 1. Normalized cloud thermodynamic phase index frequency distribution from the POLDER–MODIS algorithm, for pixels with the phase-index SD less than 10. Colors represent different cloud altitudes, between 200 and 1000 m in red and between 1000 and 2000 m in black.

(Stohl et al., 1998, 2005). The model is driven with 3-hourly operational analysis wind fields from the European Centre for Medium-Range Weather Forecasts (ECMWF) with 91 model levels and a horizontal resolution of 1° \times 1°. We use the same simulations as described by Stohl et al. (2013), which consider a black-carbon tracer undergoing removal processes, two fixed-lifetime black-carbon tracers, and a carbon monoxide (CO) tracer. The CO tracer used for this study is considered as passive in the atmosphere but is removed from the simulation 31 days after emission, thus focusing the simulation on “fresh” pollution. For the CO emissions, ECLIPSE (Evaluating the CLimate and Air Quality ImPacts of Short livEd pollutants) version 4.0 emission data (Klimont et al., 2016; Stohl et al., 2015) are used. For the anthropogenic emissions considered here, the ECLIPSE emissions are based on the GAINS (Greenhouse gas–Air pollution Interaction and Synergies) model (Amann et al., 2011). The emissions are determined separately for every year of this study and, notably, include gas flaring emissions, which have been shown to be important for black carbon in the Arctic (Stohl et al., 2013). Emissions from the residential sector are temporally disaggregated using a heating degree-day approach (Stohl et al., 2013).

Studies that have used FLEXPART CO concentration fields (χ_{CO}) have found satisfactory agreement between model output and measurements (Stohl, 2006; Paris et al., 2009; Hirdman et al., 2010; Sodemann et al., 2011; Stohl et al., 2013, 2015; Eckhardt et al., 2015). In the Alaskan Arctic for the day of 18 April 2008, Warneke et al. (2009) described a slope of 0.9 for a linear fit between FLEXPART model output of χ_{CO} and airborne measurements of CO with a least-squares correlation coefficient of 0.63.

Table 1. Cloud products, pollution tracer, atmospheric reanalysis used in this study with the corresponding spatial and temporal resolution.

Parameter(s)	From	Resolution(s)
Cloud parameters (T , r_e , τ)	MODIS, POLDER-3	Spatial resolution: 36 km ²
CO tracer concentration from anthropogenic sources	FLEXPART	Spatial resolution: 1° × 2° Temporal resolution: 3 h
Specific humidity, temperature profile	ERA-I (ECMWF)	Spatial resolution: 1.5° × 1.5° Temporal resolution: 6 h

The FLEXPART model outputs used here have a temporal resolution of 3 h and a spatial resolution of 1° × 2° (in latitude and longitude) divided into 9 different vertical levels. FLEXPART CO concentration (χ_{CO}) output is provided in units of mg m⁻³ but converted to units of ppbv (parts per billion by volume) to remove the atmospheric pressure dependence. Since the focus here is on the effect of anthropogenic pollution on clouds, only FLEXPART spatial bins where anthropogenic sources comprise more than 80 % of total CO concentrations are considered for comparison with cloud properties.

2.3 Meteorological data

ERA-Interim (ERA-I) reanalysis data from ECMWF (Berrisford et al., 2009) extend from 1989 to the present with an improved version released in 2011 (Dee et al., 2011). The temporal resolution is 6 h at 60 pressure levels. Reanalysis data from ERA-I show good agreement with satellite retrievals and aircraft data for cloud fraction and cloud radiative forcing in the Arctic (Zygmuntowska et al., 2012). Wesslén et al. (2014) analyzed ERA-I data with the Arctic Summer Cloud Ocean Study (ASCOS) campaign measurements in 2008 and calculated biases of about 1.3 °C, 1 %, and -1.5 hPa, respectively, for temperature, relative humidity, and surface pressure; root mean square errors of about 1.9 °C, 3.7 %, and 8.7 hPa, respectively; and correlation coefficients of approximately 0.85 for temperature and surface pressure and 0.31 for the relative humidity.

The goal of this study is to use satellite, tracer transport model, and meteorological data sets to determine the effects of long-range aerosol transport on cloud microphysics due only to the pollution itself and not to the meteorological state. The focus is on temperature, specific humidity, and LTS since these have been identified as basic meteorological quantities that correlate with cloud microphysical properties (Matsui et al., 2006; Mauger and Norris, 2007). Defining the potential temperature (θ) as

$$\theta = T \cdot \left(\frac{P_0}{P} \right)^{\frac{R}{c_p}}, \quad (1)$$

where T and P are the air temperature and pressure; P_0 equals 1000 hPa; R and c_p are, respectively, the gas constant for air and the isobaric heat capacity; and the LTS is

defined as the potential temperature difference between 700 and 1000 hPa (Klein and Hartmann, 1993).

$$\text{LTS} = \theta_{700} - \theta_{1000} \quad (2)$$

We also consider clouds with values of LWP greater than 40 g m⁻² separately from clouds with LWP values less than 40 g m⁻². This approach separates clouds according to their thermal radiative properties since a cloud with low LWP will tend to act as a graybody and potentially be radiatively susceptible to pollution at thermal wavelengths (Garrett and Zhao, 2006; Lubin and Vogelmann, 2006; Mauritsen et al., 2011). Thick clouds act as blackbodies, and their long-wave radiative properties are determined by temperature only (Garrett and Zhao, 2006; Garrett et al., 2009).

The different data sets used in this article are summarized in Table 1.

3 Methodology

This study examines data between 2008 and 2010 over the ocean at latitudes greater than 65°. Passive satellite sensors measure interactions of solar radiation with the atmosphere so as to retrieve cloud microphysical parameters of interest from visible wavelength measurements so analyses are restricted to the period between 1 March and 30 September.

3.1 Colocation of satellite retrieval and model pollution tracer fields

CO tracer concentrations from a FLEXPART grid cell are defined as the average between two temporal points, averaged over a spatial box. For example, model CO concentrations at 03:00 UTC and at the latitude–longitude coordinates of (70°, 80°), represent an average over a box between the latitudes of 70–71° and longitudes of 80–82° and between 00:00 and 03:00 UTC.

For an A-train satellite overpass time of 00:45 UTC, we match space-based retrievals to FLEXPART concentration output at 03:00 UTC representing the average concentration between 00:00 and 03:00 UTC and then linearly interpolate ECMWF meteorological fields to the LTS and specific humidity values for 01:30 UTC.

Here, as with many prior studies looking at aerosol–cloud interactions in the Arctic, we consider only low-level clouds

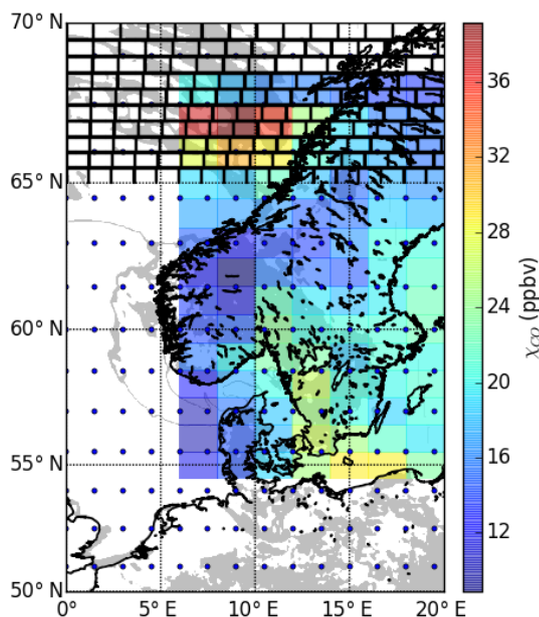


Figure 2. Illustration of the horizontal collocation method, showing satellite data corresponding to cloud top pressures below 1000 m altitude (gray shading), the average FLEXPART CO concentration between 1 and 2 km (colored shading), and the spatial resolution of temperature profiles and specific humidity in blue points. The black grid, at the top of the map, corresponds to the sinusoidal equal-area grid used in this study for collocating each data set.

(Garrett et al., 2004; Garrett and Zhao, 2006; Lubin and Vogelmann, 2006; Mauritsen et al., 2011), with POLDER cloud top altitudes between 200 and 1000 m, and between 1000 and 2000 m. The cloud top pressure is translating to cloud top altitude by a pressure profile specific to the Arctic region. These two layers correspond to the FLEXPART vertical bin resolution. We average POLDER and MODIS data that fall within the height bins so that they are collocated with the corresponding FLEXPART CO concentrations.

Regarding horizontal collocation, Fig. 2 shows how the data sets are combined. We project data from satellite, model, and reanalysis data sets onto an equal-area sinusoidal grid such that the grid cell resolution is $0.5^\circ \times 0.5^\circ$ at the equator corresponding to an area of $54 \text{ km} \times 54 \text{ km}$. The sinusoidal projection conserves the grid cell area independently of longitude and latitude. One grid cell can include up to 81 POLDER-MODIS pixels. Satellite and tracer transport model data are averaged over each grid cell.

One consequence of the averaging is that each grid cell can include up to 81 different pixel-level values of Φ . We place a limit on the SD of the averaged phase index within each grid cell such that $\sigma_\Phi < 10$ in order to satisfy an a priori requirement of there being a nearly homogeneous phase within each grid cell.

3.2 The net aerosol–cloud interactions parameter

Assuming a constant LWC and a monodisperse size distribution of cloud droplets, the droplet effective radius (r_e) decreases as the CCN number concentration N_c increases following the relation (Feingold et al., 2001)

$$\left. \frac{d \ln r_e}{d \ln N_c} \right|_{\text{LWP}} = -\frac{1}{3}. \quad (3)$$

Here, we take a different approach, which is to examine how cloud properties change in response to changes in a CO tracer under the presumption that the CO tracer serves as a proxy for the potential of long-range pollution transport, of which CCN may be a part. Of course, N_c and CO tracer concentrations (χ_{CO}) do not represent the same quantity. However, cloud condensation nuclei and CO are both byproducts of combustion. The two quantities are expected to be highly correlated close to pollution sources where relative changes in one quantity can serve as a proxy for relative changes in the other (Avey et al., 2007; Tietze et al., 2011). The reason for using CO is 2-fold. First, CO is passive and therefore easier to represent in a dynamic model. Second, the analysis here is less focused on the local physics of aerosol–cloud interactions and more focused on the actual impact of anthropogenic activities on clouds far from combustion sources. These are similar but not identical questions. The aerosol–cloud interactions (or the ACI) parameter addresses the precise physics of the extent to which aerosols can modify clouds. However, interactions are a two-way street: when aerosols have been scavenged en route to distant clouds, there is potential for a pollution plume to be present but its impact on cloud properties to be weak. To account for scavenging, we employ the term $\text{ACI}_\tau^{\text{net}}$ or the net aerosol–cloud interaction parameter. $\text{ACI}_\tau^{\text{net}}$ is the same as the ACI while additionally accounting for any reduction of the ACI due to dry or wet scavenging of aerosols during transport. $\text{ACI}_\tau^{\text{net}}$ can be interpreted as a measure of the sensitivity of a cloud at any given location to pollution plumes from distant sources. It allows for the passive components of a plume (e.g., CO) to remain while aerosols have been removed:

$$\text{ACI}_\tau^{\text{net}} = \frac{d \ln \tau}{d \ln \chi_{\text{CO}}} \quad (4)$$

$$\text{ACI}_{r_e}^{\text{net}} = -\frac{d \ln r_e}{d \ln \chi_{\text{CO}}}. \quad (5)$$

For example, in the absence of scavenging, N_c from Eq. (3) is linearly related with χ_{CO} and Eq. (5) will be bound by a theoretical maximum value of $\text{ACI}_{r_e}^{\text{net}} = 1/3$ (Twomey, 1977; Feingold et al., 2001). Further from source regions, the correlation of CO concentration and aerosols is invariant to dilution but it may be affected by wet and dry scavenging (Garrett et al., 2010, 2011). If scavenging rates are low, CO and CCN tend to covary, but when precipitation is high along transport

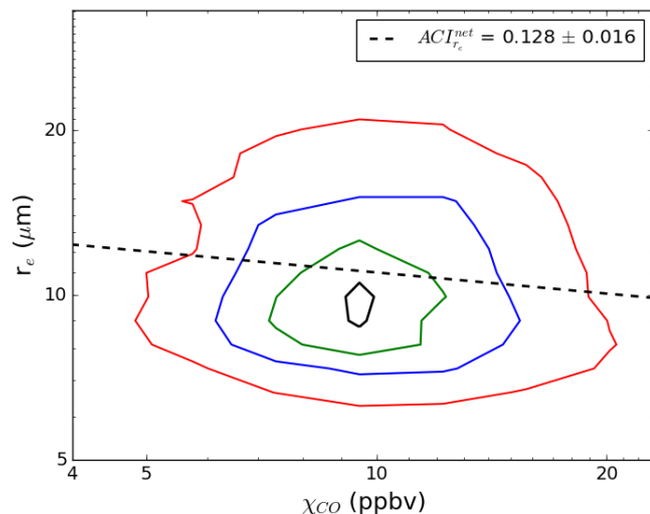


Figure 3. Calculation of the $ACI_{r_e}^{net}$ parameter from a probability distribution of values in the effective radius and CO tracer concentration for liquid clouds with cloud top altitudes between 1000 m and 2000 m, and cloud top temperatures between -12 and 6°C . The color scale indicates higher density of values in linear intervals. The $ACI_{r_e}^{net}$ number indicates the negative slope of the linear fit (dashed line).

pathways then aerosols are removed and values of ACI_{τ}^{net} will tend to be lower. Garrett et al. (2010) found that in Barrow, Alaska, when the temperature exceeds 4°C at the surface, wet scavenging efficiently removes CCN from the atmosphere. In this event, cloud microphysical properties will not be affected by pollution plumes.

Since r_e and the optical depth (τ) are linked through $\tau = \frac{3}{2} \frac{LWP}{\rho_w r_e}$ for homogeneous clouds, it follows that

$$\frac{d \ln \tau}{d \ln \chi_{CO}} = -\frac{d \ln r_e}{d \ln \chi_{CO}} + \frac{d \ln LWP}{d \ln \chi_{CO}} \quad (6)$$

or,

$$ACI_{\tau}^{net} = ACI_{r_e}^{net} + ACI_{LWP}^{net}. \quad (7)$$

Figure 3 shows an example of how ACI_{τ}^{net} is calculated for temperatures between -12 and 6°C and altitudes between 1000 and 2000 m, for all LWP values. We first calculate $ACI_{r_e}^{net}$ as the linear fit of the natural logarithm of the effective radius to the natural logarithm of CO concentrations. The fit used in this study is based on the robust linear method (RLM) (Huber, 1973, 1981; Venables and Ripley, 2013). RLM uses an iterative least-squares algorithm: every measurement has initially the same weight; the weights of each point are updated, giving a lower weight to points that appear as outliers with respect to the entire data set. The process iterates several times and stops when the convergence tolerance of the estimated fitting coefficients lies below 10^{-8} . The slope is therefore less sensitive to outlier points. In Fig. 3, points indicated by the red line are weighted similarly to those indicated by

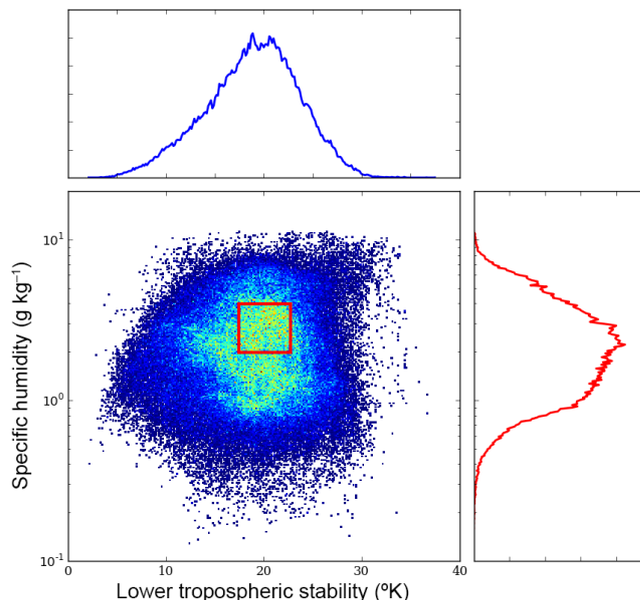


Figure 4. A 2-D histogram of the specific humidity and the LTS retrieved by ECMWF reanalysis from 2008 to 2010. The red rectangle corresponds to the range where there is a maximum of measurements within a bin corresponding to 15 % of the total range length of the corresponding parameter.

Table 2. Summary of the different ranges of the logarithm of the specific humidity and the LTS over the region of interest, detailing the method used to determine the final range of parameters considered. The Δ defines the difference between the maximum and the minimum of the total range. The considered range is chosen to keep the maximum number of measurements within a fixed interval of 15 % of the range, corresponding to the red square on Fig. 4.

	\log_{10} (specific humidity)	LTS ($^\circ\text{K}$)
Total range [min, max]	[$-0.89, 1.0$]	[2.1, 38]
Δ total range	1.9	36
15 % interval	0.28	5.4
Stratified range [min, max]	[0.30, 0.60]	[17, 22]
	= [2.0, 4.0] (g kg^{-1})	

black and blue lines. The slope retrieved by the linear fit, in Fig. 3, is -0.13 ± 0.016 . Referring to Eq. (5), $ACI_{r_e}^{net}$ equals $+0.13 \pm 0.016$.

3.3 Stratifying the data for specific humidity and lower tropospheric stability

Figure 4 presents a 2-D histogram of frequency distribution of the specific humidity and the LTS. The LTS ranges from 2.1 to 37 K and the specific humidity from 0.13 to 11 g kg^{-1} . The median values for specific humidity and LTS are 2.0 g kg^{-1} and 19 K, respectively.

Table 2 describes the method used here to stratify the data set according to meteorological conditions. We identify

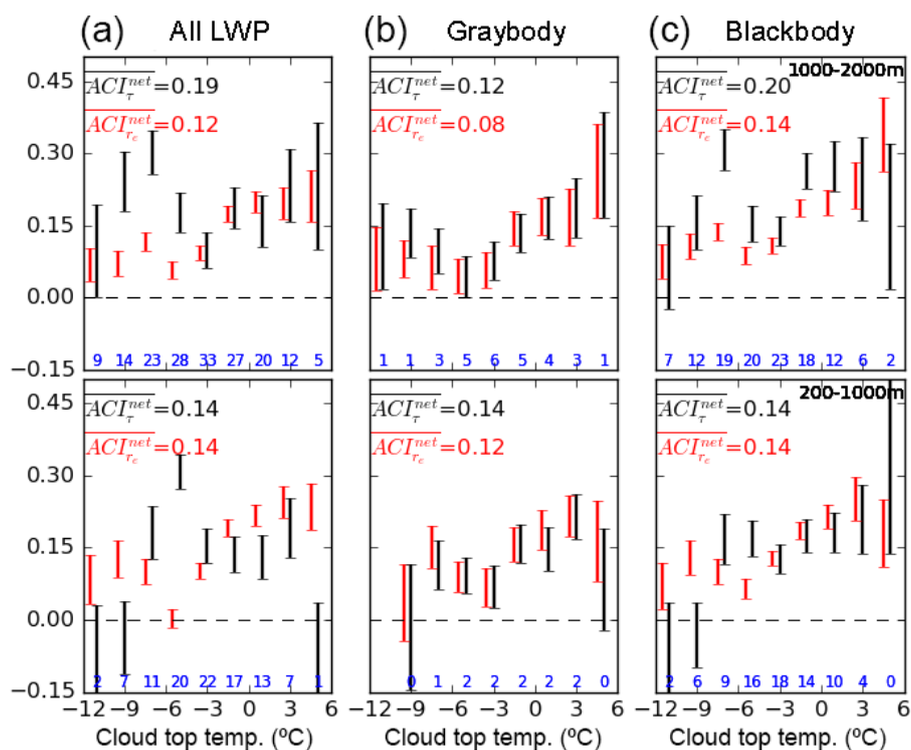


Figure 5. ACI^{net} parameter of the effective radius (r_e) (red) and optical depth (τ) (black), as a function of temperature calculated for liquid clouds between 200–1000 m (lower row) and 1000–2000 m (upper row). The bars indicate the 95 % confidence limit in the calculation of the mean ACI^{net} value. Each column corresponds to different thresholds for LWP (blackbody: $LWP > 40 \text{ g m}^{-2}$, graybody: $LWP < 40 \text{ g m}^{-2}$). Blue numbers indicate the number of grid cells, in hundreds, that are used to calculate each ACI^{net} value. In each figure the ACI^{net} value averaged over the temperature and weighted according to the inverse of the uncertainty is indicated.

a range in LTS and specific humidity that occupies 15 % of the total space of observed values but that is centered at the mode of the respective distributions. The total LTS range is 2.1–37 K, so the interval size is 5.3 K. The specific humidity is distributed over several orders of magnitude. To better represent the distribution, we use a logarithmic scale for this parameter. The logarithm base 10 of specific humidity has an interval of 0.28. The most common values of meteorological state, defined here as the maximum number of measurements, is delimited by the red rectangle in Fig. 4. The rectangle corresponds to a range of 2.0 to 4.0 g kg^{-1} for specific humidity, and a range of 16.5 to 21.8 K for LTS. It is these ranges that are focused on for calculation of the ACI^{net} parameter. We assume these intervals are sufficiently narrow that the variability within the interval has limited impact on cloud microphysics.

4 Results

4.1 Net aerosol–cloud interactions

Figure 5 summarizes ACI^{net} values calculated using combined POLDER-3, MODIS, and FLEXPART data for the period between 2008 and 2010 for latitudes greater than 65°

over the ocean, stratifying the data set by cloud top temperature in bins of 2° between −12 and 6 °C. The results are categorized according to bins in temperature, altitude and LWP, and LTS and specific humidity stratified. The number of grid cells used to calculate each ACI^{net} parameter per bin ranges from 100 to 3300. The ACI^{net} parameter is almost always positive but sometimes close to zero. $ACI_{r_e}^{net}$ ranges from 0 for graybody clouds between 1000 and 2000 m altitude with a cloud top temperature between −6 and −4 °C, to 0.34 for blackbody clouds between 1000 and 2000 m altitude with a cloud top temperature between 4 and 6 °C. ACI_{τ}^{net} ranges from −0.10 for all clouds between 200 and 1000 m altitude with a cloud top temperature of −11 °C, to 0.35 at 3 °C for blackbody clouds between 1000 and 2000 m altitude. In general, ACI_{τ}^{net} and $ACI_{r_e}^{net}$ are of the same order of magnitude and the maximum values of ACI^{net} are found for clouds with temperatures above the freezing temperature.

We define the uncertainty in ACI^{net} as the 95 % confidence limit in the calculation of the slope of the linear fit. The uncertainty in the calculated values of $ACI_{r_e}^{net}$ is generally less than 0.1, except for clouds with temperatures between 4 and 6 °C and between −12 and −10 °C, where the uncertainty bar is approximately 0.2. For the optical depth, the uncer-

Table 3. ACI_{τ}^{net} parameter calculated for the optical depth and the effective radius considering all clouds, graybody clouds, and blackbody clouds, averaged from values presented in Fig. 5 and weighted considering the inverse of the uncertainty in the mean.

	All LWP	Graybody	Blackbody
$ACI_{r_e}^{\text{net}}$	0.12	0.10	0.14
ACI_{τ}^{net}	0.16	0.13	0.17

tainty is typically approximately 0.1, although larger values are observed for high and low cloud top temperatures.

For blackbody clouds between 1000 and 2000 m altitude, the average values of ACI_{τ}^{net} and $ACI_{r_e}^{\text{net}}$ equal 0.20 and 0.14, respectively. For cloud tops between 200 and 1000 m altitude, ACI_{τ}^{net} and $ACI_{r_e}^{\text{net}}$ equal 0.14. For graybody clouds between 1000 and 2000 m, ACI_{τ}^{net} and $ACI_{r_e}^{\text{net}}$ equal 0.12 and 0.08, respectively. For cloud tops between 200 and 1000 m altitude, ACI_{τ}^{net} and $ACI_{r_e}^{\text{net}}$ equals 0.14 and 0.12, respectively. Thus, the value of ACI^{net} appears to be fairly robust to altitude and cloud thickness and to whether r_e or τ is considered. Table 3 presents the average ACI_{τ}^{net} and $ACI_{r_e}^{\text{net}}$. For all cases, ACI^{net} values are near 0.13 ± 0.03 .

4.2 Dependence of ACI^{net} on pollution concentration, specific humidity, and lower tropospheric stability

In what follows, we examine the influence of LTS, specific humidity, and pollution concentration on ACI^{net} . Table 4 shows values of $ACI_{r_e}^{\text{net}}$ and ACI_{τ}^{net} for graybody and blackbody clouds, and for $\chi_{\text{CO}} < 5.5$ ppbv and $\chi_{\text{CO}} > 10.0$ ppbv, corresponding respectively to the lower and upper quartile of CO tracer concentration, and for a range in LTS and specific humidity. For graybody and blackbody clouds, ACI_{τ}^{net} and $ACI_{r_e}^{\text{net}}$ are highest for low values of χ_{CO} . The difference in ACI^{net} values between low and high polluted environments is slightly greater for $ACI_{r_e}^{\text{net}}$ than for ACI_{τ}^{net} . Table 4 suggests that cloud-droplet effective radius and cloud optical depth are most sensitive to pollution when pollution concentrations are low. Previous studies have hypothesized that the effect of CCN on cloud microphysical properties saturates when cloud-droplet concentrations are high (Bréon et al., 2002; Andersen and Cermak, 2015). This effect does not explain the differences presented in Table 4 because Eqs. (4) and (5) already take into account the potential for linear saturation by considering the logarithms of χ_{CO} and cloud parameters.

We now present the sensitivity of the ACI^{net} parameter to five different ranges of meteorological parameters delimited by the percentiles values presented in Table 5. We do not stratify the data according to LWP. Figures 6 and 7 show the influence of pollution loading on the cloud-droplet effective radius and cloud optical depth for each of the different specific humidity and LTS regimes. Figure 6 presents the ACI^{net} parameter with respect to the cloud optical depth and cloud-

Table 4. ACI^{net} parameter calculated for the optical depth and the effective radius considering all clouds, graybody clouds, and blackbody clouds, for two different regimes of CO concentration representing lower and upper quartiles of CO concentration.

	All LWP		Graybody		Blackbody	
	$ACI_{r_e}^{\text{net}}$	ACI_{τ}^{net}	$ACI_{r_e}^{\text{net}}$	ACI_{τ}^{net}	$ACI_{r_e}^{\text{net}}$	ACI_{τ}^{net}
$\chi_{\text{CO}} < 5.5$ ppbv	0.23	0.36	0.31	0.26	0.28	0.24
$\chi_{\text{CO}} > 10$ ppbv	0.09	0.35	0.09	0.16	0.14	0.16

Table 5. Percentile values of specific humidity and LTS used to defined different regimes of the meteorological parameters.

	Specific humidity (g kg^{-1})	LTS (K)
Minimum	0.13	2.1
20th percentile	1.2	14
40th percentile	1.7	18
60th percentile	2.4	20
80th percentile	3.6	23
Maximum	11	37

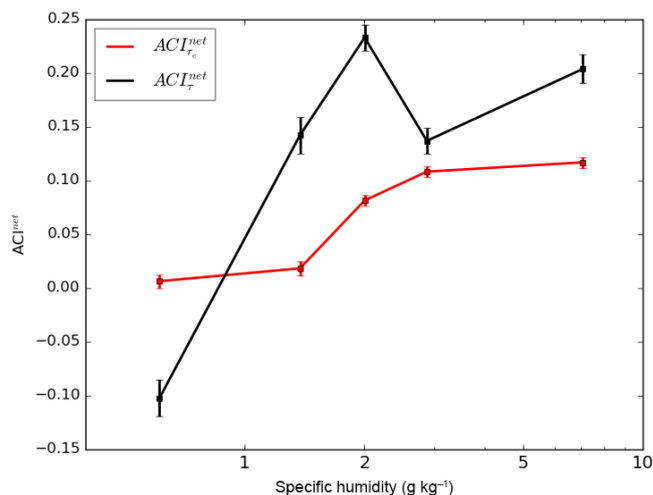


Figure 6. $ACI_{r_e}^{\text{net}}$ (red) and ACI_{τ}^{net} (black) for different bins of the specific humidity, stratified by LTS between 17 and 22 K. Each marker is placed in the middle of the corresponding bin.

droplet effective radius as a function of the specific humidity, stratifying the data by LTS according to the method described in Sect. 3.3. Figure 7 is the same as Fig. 6 except that it shows ACI^{net} as a function of LTS for a range of specific humidities.

Figure 6 shows that $ACI_{r_e}^{\text{net}}$ and ACI_{τ}^{net} tend to increase with the specific humidity independent of LTS. The ACI^{net} parameter is close to zero, or negative, for low values of specific humidity. It increases rapidly with specific humidity, saturating at a maximum value of about 2.5 g kg^{-1} . We note that cloud top temperature and specific humidity are weakly correlated. The correlation coefficient (r^2) of the linear re-

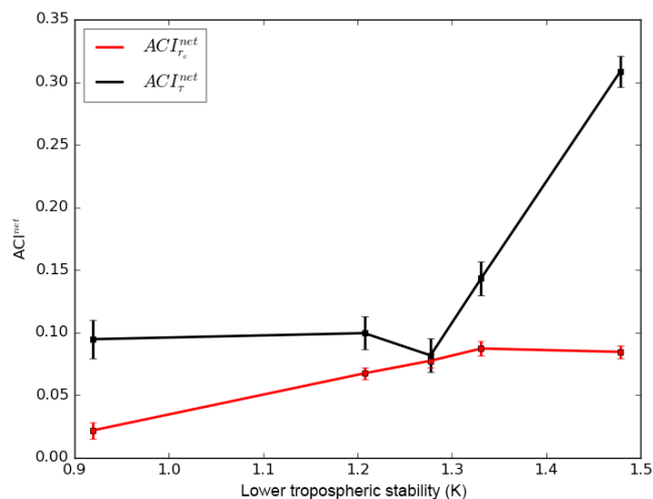


Figure 7. $ACI_{r_e}^{net}$ (red) and ACI_{τ}^{net} (black) and ACI_{LWP}^{net} as a function of the lower tropospheric stability, stratified by specific humidity between 2.0 and 4.0 $g\ kg^{-1}$.

gression of the two parameters is 0.20. The same applies for the specific humidity and the LWP. The correlation coefficient of the two parameters is 0.05.

$ACI_{r_e}^{net}$ increases with LTS from 0.02 for values of LTS ranging between 2.1 and 14 K to 0.09 for values of LTS between 23 and 38 K. The ACI_{τ}^{net} dependence on LTS is larger: ACI_{τ}^{net} equals 0.10 for LTS values between 2.1 and 14 K and it equals 0.32 for LTS values between 21 and 38 K.

5 Discussion

The results presented here show values of the ACI^{net} parameter with respect to the cloud-droplet effective radius and optical depth, for clouds over oceans north of 65° lying between 200 and 2000 m, and for the years between 2008 and 2010. We find ACI^{net} values that range from 0.00 to 0.34 for the cloud-droplet effective radius, and from -0.10 to 0.35 for the optical depth.

Prior studies examining the Arctic region have retrieved ACI values ranging from -0.10 to 0.40 (Garrett et al., 2004; Lihavainen et al., 2009; Zhao et al., 2012; Sporre et al., 2012). Tietze et al. (2011) calculated ACI^{net} values ranging from 0.00 to 0.17 using a similar satellite–FLEXPART collocation method. What differs in this study is that we examine solely anthropogenic pollution, and that we extend the data set from 1 year to 3 years, stratifying the data set according to specific humidity and LTS. The larger ACI^{net} values we find in this study suggest a higher sensitivity of cloud microphysical properties to pollution plumes from distant sources than was found by Tietze et al. (2011).

However, Tietze et al. (2011) also found values of ACI_{τ}^{net} that were greater than $ACI_{r_e}^{net}$ by a factor of 4, and they attributed this difference to unknown dynamic or precipitation

Table 6. Difference between ACI_{τ}^{net} and $ACI_{r_e}^{net}$ (i.e., ACI_{LWP}^{net}) for graybody, blackbody, and all clouds when lower tropospheric stability and specific humidity are stratified and when they are not stratified. The averaged ACI^{net} values are shown in Table 3.

	All LWP	Graybody	Blackbody
Stratified	0.04	0.03	0.04
Not stratified	0.12	0.04	0.08

feedbacks that make ACI_{LWP}^{net} greater than zero (Eq. 7). In contrast, our results show that the $ACI_{r_e}^{net}$ and ACI_{τ}^{net} parameters are more similar, suggesting no such feedback. Table 6 compares the differences between ACI_{τ}^{net} and $ACI_{r_e}^{net}$ that are presented in Table 3, along with their corresponding values when no control is made for specific humidity and LTS. The difference between $ACI_{r_e}^{net}$ and ACI_{τ}^{net} is largest when meteorological parameters are not controlled for. For all clouds considered, the maximum difference increases from 0.04 when the data are considered within narrow meteorological bands to 0.12 when the data are not. This result is important since it suggests that the hypothesized feedbacks discussed by Tietze et al. (2011) may have in fact been due to the natural sensitivity of clouds to local meteorology. Not controlling sufficiently for meteorology may lead to artifacts that exaggerate the magnitude of the aerosol indirect effect.

In contrast to most prior efforts, satellite-retrieved cloud properties are not compared to CCN or aerosol concentrations but rather to pollution concentrations – specifically CO simulated from a tracer transport model. CO serves as a proxy for CCN close to pollution sources although far from sources CCN and CO can become decoupled due to scavenging en route to distant clouds. The comparison we present through the ACI^{net} parameter implicitly accounts for this possibility. For temperatures below $-6^{\circ}C$, low values of ACI^{net} are observed. Tietze et al. (2011) hypothesized that at such temperatures, cloud supersaturations may be too small to activate aerosols as CCN or that clouds with lower temperatures have followed longer transport pathways nearer to the surface (Stohl, 2006) and therefore had greater exposure to dry deposition.

Table 4 suggests that ACI^{net} values are lowest when pollution concentrations are high. Figure 8 presents the normalized distribution of potential temperature for polluted and pristine clouds, defined as the upper and lower quartile for graybody clouds. We present results for graybody clouds because the ACI^{net} differences between polluted and clean cases are largest; results for blackbody and all clouds are not shown here, but have similar results regarding the potential temperature distribution.

Highly polluted air parcels are associated with potential temperatures around 280 K whereas pristine air parcels have a lower potential temperature – around 272 K. We hypothesize that higher values of potential temperature suggest pol-

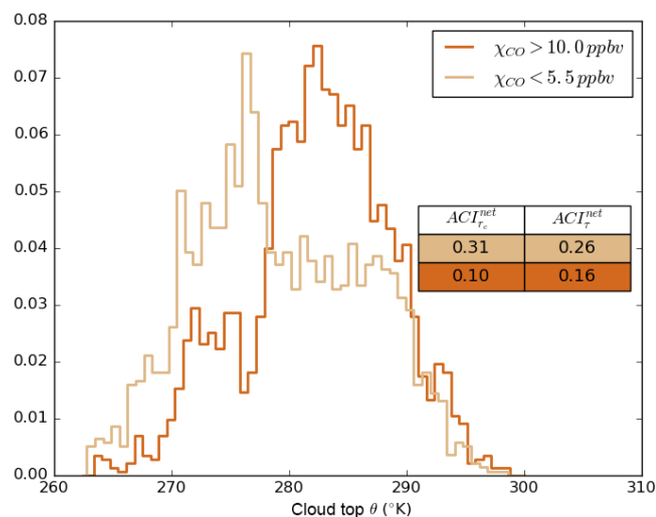


Figure 8. Normalized distribution of the cloud top potential temperature when clouds are associated with CO tracer concentrations (χ_{CO}) greater than 10 ppbv and less than 5 ppbv. The values of $ACI_{\tau_e}^{net}$ and ACI_{τ}^{net} associated with each histogram are presented also in Table 3.

lution sources from further south, so wet scavenging is more likely to occur during transport; this decreases the correlation between a CO tracer and CCN, therefore lowering the ACI^{net} parameter. Also, polluted air parcel and aerosols do not necessarily have the same physical and chemical properties at lower and higher latitudes, and this difference may impact the influence of aerosols on cloud microphysics and aerosols (Bilde and Svenningsson, 2004; Dusek et al., 2006; Ervens et al., 2007; Andreae and Rosenfeld, 2008).

In general, we observe that when moisture increases, the cloud sensitivity to pollution increases. From model simulations of stratocumulus, Ackerman et al. (2004) found that when the relative humidity (RH) above cloud top is high, cloud LWP increases with N_c consistent with theoretical arguments (Albrecht, 1989; Pincus and Baker, 1994), but that when the RH is low, the LWP decreases when N_c increases, as supported by some observations (Coakley and Walsh, 2002). The difference was attributed to the consequence of dry air into a cloud layer. Humidity inversions are common above low-level cloud tops in the Arctic (Nygård et al., 2014), so similar phenomena may be playing a role.

Studies of the indirect effect at midlatitudes suggest that values of ACI are highest under unstable conditions (Chen et al., 2014; Andersen and Cermak, 2015). Our results from the Arctic show the reverse, that conditions of high LTS are associated with higher values of ACI^{net} . Klein and Hartmann (1993) showed that, in general, higher values of LTS lead to greater stratiform cloudiness, except in the Arctic where radiative cooling prevails over convection as the driving mechanism for cloud formation. This result is similar to results found by Kim et al. (2008) who found that aerosol–cloud in-

teractions are strongest in clouds with adiabatic liquid water content profiles. Such clouds might be expected more frequently when LTS is high and there is reduced vertical mixing.

Finally, we find ACI_{τ}^{net} is more sensitive to changes in LTS than $ACI_{\tau_e}^{net}$. A consequence is that for values of LTS greater than 23 K, ACI_{τ}^{net} and $ACI_{\tau_e}^{net}$ differ by about 0.20. In a stable atmosphere with high LTS it appears that ACI_{LWP}^{net} increases more strongly in response to aerosols than in unstable environments (Klein and Hartmann, 1993; Qiu et al., 2015).

6 Conclusions

Satellite, numerical model, and meteorological reanalysis data sets from 2008 to 2010 are used here to calculate the sensitivity of cloud-droplet effective radius and optical depth in the Arctic to anthropogenic pollution transported from mid-latitudes. We focused on latitudes north of 65° for a period between March 2008 and October 2010. Using ECMWF reanalysis data, we stratified the data set according to temperature, LTS, specific humidity, altitude, and LWP. We find that the sensitivity of cloud properties to pollution, as quantified by values of ACI^{net} , lies close to a theoretical maximum value of $1/3$, assuming that a simulated CO tracer correlates well with CCN. Further, $ACI_{\tau_e}^{net}$ and ACI_{τ}^{net} seem to increase with specific humidity and LTS, highlighting that meteorological parameters have an important impact on aerosol–cloud interactions.

Globally, Klimont et al. (2013) have estimated that there was a drop of about 9281 Gg in anthropogenic sulphur dioxide emissions between 2005 and 2010 due to a reduction in European and American emissions and a flue gas desulfurization program on power plants in China. This reduction in emissions has led to a decrease of sulfate concentrations at the Arctic surface station (Hirdman et al., 2010). In the Arctic, the effect of a decrease in midlatitude pollution emissions may someday be offset by greater levels of Arctic industrialization (Lindholt and Glomsrød, 2012) and shipping (Pizzolato et al., 2014; Miller and Ruiz, 2014) that introduce new local aerosol sources. Further, an increase in the extent of open ocean due to sea-ice retreat may be expected to lead to an increase in the atmospheric humidity (Boisvert and Stroeve, 2015) and from the results presented here, a higher sensitivity of clouds to aerosols. However, this study also suggests that any associated decrease in LTS could partially counter-act this effect.

Climate warming is thought to stimulate boreal forest fires (Westerling et al., 2006). The impact of pollution from biomass burning has not been included in the present research. Given biomass burning aerosol can act as efficient ice nuclei (Markus et al., 2009), the analyses presented here might be extended to explore aerosol-induced changes in cloud thermodynamic phase.

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