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The scavenging processes controlling the seasonal cycle in Arctic sulphate and black carbon aerosol

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Abstract. The seasonal cycle in Arctic aerosol is typified by high concentrations of large aged anthropogenic particles transported from lower latitudes in the late Arctic winter and early spring followed by a sharp transition to low concentrations of locally sourced smaller particles in the summer. However, multi-model assessments show that many models fail to simulate a realistic cycle. Here, we use a global aerosol microphysics model (GLOMAP) and surface-level aerosol observations to understand how wet scavenging processes control the seasonal variation in Arctic black carbon (BC) and sulphate aerosol. We show that the transition from high wintertime concentrations to low concentrations in the summer is controlled by the transition from ice-phase cloud scavenging to the much more efficient warm cloud scavenging in the late spring troposphere. This seasonal cycle is amplified further by the appearance of warm drizzling cloud in the late spring and summer boundary layer. Implementing these processes in GLOMAP greatly improves the agreement between the model and observations at the three Arctic groundstations Alert, Barrow and Zeppelin Mountain on Svalbard. The SO_4 model-observation correlation coefficient (R) increases from: -0.33 to 0.71 at Alert (82.5° N), from -0.16to 0.70 at Point Barrow (71.0° N) and from -0.42 to 0.40 at Zeppelin Mountain (78° N). The BC model-observation correlation coefficient increases from -0.68 to 0.72 at Alert and from -0.42 to 0.44 at Barrow. Observations at three marginal Arctic sites (Janiskoski, Oulanka and Karasjok) indicate a far weaker aerosol seasonal cycle, which we show is consistent with the much smaller seasonal change in the frequency of ice clouds compared to higher latitude sites. Our results suggest that the seasonal cycle in Arctic aerosol is driven by temperature-dependent scavenging processes that may be susceptible to modification in a future climate.

1 Introduction

The effects of climate change are amplified in the Arctic, with projected temperature increases of 5 °C predicted before the year 2100 (IPCC, 2007). This amplified warming is manifested in the rapid decline in sea ice extent over the past few decades (Comiso, 2002). The retreat of Arctic sea ice could shift patterns of atmospheric circulation, increase coastal erosion and disrupt the ocean thermohaline cycle as well as devastate the fragile Arctic ecosystem (Serreze et al., 2007). Furthermore, the decrease in planetary albedo may lead to an acceleration of global and regional warming through the albedo feedback described in Curry et al. (1995).

The increase in Arctic surface temperatures since the mid-1970s is estimated to be between 1.2 and 1.8 °C (Shindell and Faluvegi, 2009). The modelling study of Shindell and Faluvegi (2009) suggests that less than one-third of this warming can be explained by changes in either greenhouse gas concentrations or the Arctic ozone budget. Changes in anthropogenic aerosol, particularly the decrease in global sulphate emissions since the late 1980s, may account for up to 1 °C of the observed warming trend between 1976 and 2008.

Anthropogenic aerosols, both in the atmosphere or deposited on ice and snow, have been observed in the Arctic since the 1880s (Garrett and Verzella, 2008). The high concentrations of aerosol observed in the Arctic spring, now referred to as Arctic haze, were first identified in the 1950s by

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Mitchell (1956). Analysis of the haze constituents by Rahn et al. (1977) and Shaw (1982) found a signature of Asian sources, and from the 1980s onwards numerous studies have confirmed the anthropogenic origin of Arctic haze events in the early spring (Gong and Barrie, 2005; Fisher et al., 2010; Hirdman et al., 2010; Sharma et al., 2006; Reddy and Boucher, 2007).

A strong seasonal cycle in Arctic sulphate and BC aerosol mass concentrations is evident in long term observations at Point Barrow in Alaska (Sharma et al., 2006; Quinn et al., 2009, 2002), Zeppelin Mountain on Svalbard (Eleftheriadis et al., 2009; Ström et al., 2003) and Alert in Canada (Gong et al., 2010). The Arctic haze forms due to the strong temperature inversion over the Arctic, which facilitates the ascent of transported pollutant plumes from lower latitudes over the pole. The cold dry conditions of the Arctic winter, which suppress removal via wet deposition, and the suppression of vertical mixing by the polar inversion are thought to provide suitable conditions for transported pollutants to accumulate over the winter and early spring (Shaw, 1995). This situation persists until the beginning of summer when changes to transport pathways (Stohl, 2006), the onset of vertical mixing due to solar heating (Kahl et al., 1996), and increases in wet scavenging remove existing aerosol and hinder further aerosol transport from low latitudes.

An analysis of observations at Barrow (Garrett et al., 2010, 2011) found that the onset of local wet scavenging may be the dominant process controlling low summertime aerosol, rather than changes in the efficiency of aerosol transport to Barrow. The inefficient long-range transport in the summer means that summertime aerosol is sourced from the marginal Arctic (sea-ice boundary and boreal forest regions) and probably consists of episodic soot from forest fires (Warneke et al., 2010), sulphate from dimethyl sulphide (DMS) emitted from the ocean surface (Ferek et al., 1995; Lunden et al., 2010), and sea-spray (Yang et al., 2008). Here, we suggest that the low aerosol concentrations observed in the Arctic boundary layer north of 70° N are the result of the cumulative effect of drizzling low clouds at high latitudes on these sources (see Sect. 7.2).

Analysis of decadal trends indicates that while summertime ground-station concentrations (with the exception of Barrow) have changed little since the 1970s, in winter and spring sulphate and equivalent black carbon (EBC, see Sect. 2) concentrations have declined over the last 20 yr (Quinn et al., 2007; Hirdman et al., 2010). This decline in both BC and SO₄ is also evident in central Greenland ice-cores (McConnell et al., 2007). The global decrease in aerosol concentrations resulting from stricter air quality controls in Europe and North America (IPCC, 2007) is a possible cause of this decline. However, the North Atlantic Oscillation (NAO) also strongly influences pollution transport to high latitudes with positive phases of the NAO shown to enhance winter surface pollutants by up to 70% (Eckhardt et al., 2003). Between 1970 and 2000 the NAO index

for December to February was on average strongly positive (\sim 2) compared to the average low negative index between 2000 and 2010 (seasonal NAO indices obtained from Hurrell, 2011). The recent decline in Arctic aerosol concentrations between winter and early spring is therefore most likely the result of a reduction in source emissions combined with a weakening of transport pathways post-2000.

Alternatively, it has also been shown (Vinogradova and Ponomareva, 2000) that scavenging efficiency, to which Arctic aerosol concentrations are particularly sensitive (Shindell et al., 2008), in liquid-phase clouds is far greater then that in ice-phase (see Sect. 5.1). Therefore it may be that the increase in global temperatures since the 1970s (IPCC, 2007) has increased the ratio of liquid to ice-phase cloud scavenging and so decreased the efficiency of long range-transport into the Arctic. Currently, little evidence exists to support or refute this hypothesis. Here, we suggest that wintertime Arctic scavenging rates are relatively insensitive to temperature increases of less than 5 °C (see Sect. 7.3.2).

At Barrow, in addition to the winter decline in aerosol concentrations, a summertime increase in non-sea-salt (nss) SO₄ of 8 to 12 % per year has been observed from 1976 to 2008. This increase in nss-SO₄ mass coincides with an equivalent increase in methane sulphonic acid (MSA) concentration, a by-product of DMS oxidation to SO2. Increases in MSA concentration suggest that climatic changes (i.e. increasing sea surface temperature) in the Arctic may be altering atmospheric DMS concentrations (Quinn et al., 2009), although this hypothesis is not explored here. In addition to increasing DMS concentrations it is likely that a decline in Arctic sea-ice extent (coupled with increases in wind speed) will increase sea-salt aerosol concentrations in the Arctic boundary layer (Jones et al., 2007; Struthers et al., 2011). However, to the knowledge of the authors no observational evidence exists to support this hypothesis.

In a multi-model assessment, Shindell et al. (2008) compared 13 models against measured EBC and sulphate mass concentrations at Alert, Barrow and Svalbard. They found that the majority of models significantly underestimated both BC and SO₄ concentrations through January to May while overestimating summertime SO₄. In almost half of the models the seasonal cycle in sulphate mass concentration was approximately 6 months out of phase with observations. Korhonen et al. (2008b) compared simulated aerosol size distributions in a particle size-resolving aerosol microphysics model (Spracklen et al., 2005) with observations at Zeppelin mountain on Svalbard (Ström et al., 2003) and again found them to be seasonally out of phase, with larger particles dominating the summertime size distribution. Both studies looked in detail at the sensitivity of the seasonal cycle to different deposition processes and found aerosol concentrations to be extremely sensitive to changes to modelled wet deposition

Here, two new deposition parametrisations, ice-cloud and low-cloud scavenging, have been added to the global aersol model GLOMAP-mode (Mann et al., 2010) to understand how changes to the local and global scavenging distribution affect the Arctic SO₄ and BC aerosol seasonal cycle. Through analysis of observed and modelled sulphate and BC aerosol surface concentrations it was demonstrated that the rapid shift in aerosol characteristics observed in the late springtime Arctic boundary layer is strongly controlled by the local onset of warm (liquid) cloud nucleation scavenging in the low Arctic troposphere.

2 Observations

Simulated surface SO₄ mass concentrations are evaluated using measured concentrations at three high latitude Arctic ground-stations: Zeppelin mountain on Svalbard (78.9° N, 11.9° E, 474 m a.s.l.), Alert station in the Canadian Arctic (82.5° N, 62.5° W, 210 m a.s.l.), and Point Barrow in northern Alaska (71° N, 156.6° W, 11 m a.s.l.). Below 70° N, SO₄ measurements were also available from three ground stations in the EMEP network (http://www.emep.int/): Karasjok in Norway (69° N, 24° E), Janiskoski in Russia (68.5° N, 28.5° E), and Oulanka in Finland (66° N, 29° E). The locations of all six sites are shown in Fig. 1. Observations from field campaigns at higher latitudes (above 85° N) were not used in this evaluation since they cover only short periods and the focus of this study is on the annual cycle.

Mass concentrations of non-sea-salt (nss) SO_4 at Zeppelin were sourced from the Norwegian institute for air research (NILU) database (www.nilu.no/projects/ccc/create/database. htm) as daily concentrations which were then converted to monthly means after the removal of statistical outliers (here assuming a normal distribution defined as data points greater then 1.5 standard deviations from the monthly mean). The particle size measured at Zeppelin was limited to below $10\,\mu m$. However, at this latitude this size limit is unlikely to have a significant impact on measured mass concentrations.

At Barrow weekly sampled nss-SO₄ mass concentrations for both submicron (diameter $\leq 0.1\,\mu m)$ and supermicron (0.1 $\mu m \leq$ diameter $\leq 10\,\mu m)$ particulates were sourced from the NOAA archive (ftp://ftp.cmdl.noaa.gov/aerosol/brw/chem/). SO₄ mass concentrations at Barrow were sampled over different time periods (ranging from 1–5 days) depending on the season and the size range. The two datasets were averaged over each month with statistical outliers removed within each. Both the submicron and supermicron monthly averaged SO₄ mass concentrations were then added to give the total monthly mean sulphate mass concentration for particulates below 10 μm in diameter.

Weekly sampled sulphate mass concentrations at Alert are available from the Canadian National Chemistry (NatChem) database (http://www.msc.ec.gc.ca/natchem/particles/index_e.html) and were converted to monthly mean concentrations. At the EMEP sites SO₄ observations were taken directly from the EMEP database already converted to monthly mean

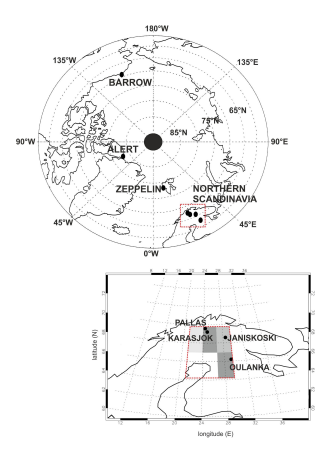


Fig. 1. Model projection of the Arctic. The Arctic ground-stations: Alert (62.5° W, 82.4° N), Barrow (156.6° W, 71.3° N) and Zeppelin mountain (11.9° E, 78.9° N) are labelled and the Northern Scandinavian sites: Oulanka (29.2° E, 66.2° N), Janiskoski (28.5° E, 68.6° N) and Karasjok (25.1° E, 69.3° N) are shown (red box). The bottom projection shows both the position of the latter three stations which are in close proximity to each other and the three model grid-boxes which represent this region (grey shading). The region outlined in red represents the model region used to calculate the Northern Scandinavian average used in Fig. 7d.

mass concentrations. Given the close proximity of the EMEP stations (in the context of the model resolution – see Fig. 1) and the consistency of observations in this region ($\sigma=0.08$), observations from all three stations were averaged to produce a single mass concentration.

Long-term BC mass concentration datasets were available only at Alert and Barrow, which limits model evaluation of BC concentrations to the Canadian and Alaskan Arctic. The majority of long-term BC measurements are derived from measurements of light absorption and so are termed equivalent black carbon (EBC). EBC is calculated using the total light absorption coefficient of an air sample so EBC concentrations are likely to overestimate BC mass due to the presence of other light absorbing aerosol species. This overestimation can be compensated for by the choice of a mass absorption efficiency (MAE) calibrated from

alternative (usually thermal) BC measurements (Sharma, 2002; Sharma et al., 2004). At Barrow hourly measured light absorption coefficients were available from the World Data Centre for Aerosols archive (http://wdca.jrc.ec.europa.eu/sites/brw/brw_labs.html) and converted to mass concentration using an MAE of $10 \, \mathrm{m}^2 \, \mathrm{g}^{-1}$ as recommended by the principal investigators and discussed in Sharma (2002).

Hourly EBC mass concentrations at Alert were obtained from the World Data Centre for Aerosols (WDCA – http:// wdca.jrc.ec.europa.eu/sites/alt/alt_ebc.html). Unlike Barrow, EBC measurements at Alert have been compared with alternative measurements from thermal filter sample analysis (Sharma et al., 2004). EBC measurements at Alert derived using a MAE of 19 m² g⁻¹ and thermally derived BC measurements were found to compare well with a R^2 correlation of 0.82 and a slope and intercept of 0.85 and 0.09 μ g m⁻³, respectively when fitted with a linear regression. In light of this comparison EBC measurements at Alert were derived from a seasonally varying specific absorption coefficient of $19.8 \,\mathrm{m^2\,g^{-1}}$ from November to April and $28.8 \,\mathrm{m^2\,g^{-1}}$ from May to October as described in Sharma et al. (2004). At both ground-stations EBC was converted from hourly measurements to monthly mean concentrations after the elimination of statistical outliers (as discussed above).

At the Alert and Barrow sites different instruments are used to measure equivalent black carbon concentrations so the two datasets are not directly comparable. The major uncertainty in these measurements is the presence of secondary absorbing species in the particles, which results in an overestimate of refractory black carbon mass. Due to the seasonality of aerosol mass concentration at Arctic sites the use of uniform MAE (such as at Barrow) is likely to overestimate summer BC mass while underestimating winter BC mass. The inter-annual variability of Arctic aerosol is also high so it is likely that from year to year the agreement of EBC mass with actual BC varies considerably. In this study we focus on reproducing the seasonal cycle (and in particular the late-spring transition) of BC and for this purpose these observations are applicable. However, quantitative comparison of these observations with the model (below) will be affected by this uncertainty.

3 Model description

GLOMAP-mode is a global microphysical aerosol model (Mann et al., 2010) developed as an extension of the off-line TOMCAT chemical transport model described in Stockwell and Chipperfield (1999). The model treats the aerosol size distribution using a two-moment modal scheme with 7 lognormal modes (4 soluble and 3 insoluble) and the 5 aerosol species dust, sea salt, BC, organic carbon (OC) and SO_4 . The 7 modes fall into four variable-size size categories: nucleation (~ 3 –15 nm), Aitken (~ 10 –50 nm), accumulation (~ 50 –200 nm) and coarse (~ 1000 nm). Concen-

trations of oxidants are specified offline every 6 h from a previous TOMCAT simulation (Arnold et al., 2005). Evaluation of GLOMAP-mode in the mid-latitudes and tropics suggest that the model performs well against observational datasets although black and organic carbon surface mass concentrations are underestimated in some regions (Mann et al., 2010).

The model was run at a horizontal resolution of $2.8^{\circ} \times 2.8^{\circ}$ (latitude \times longitude) with 31 pressure levels extending from the surface to 10 hPa (approximately 40 km) in the vertical. Meteorological fields of horizontal wind speed, temperature and humidity were taken from analyses of the European Centre for Medium-Range Weather Forecasts (ECMWF). The ECMWF analyses compare well with other analyses and describe the meteorology of the Arctic region reasonably well (Bromwich et al., 2007).

The model was run from January 2000 to December 2002 using annually-varying anthropogenic BC, organic carbon (OC) and SO_2 emission inventories from the AEROCOM hindcast database (http://dataipsl.ipsl.jussieu.fr/AEROCOM/emissions.html). AEROCOM inventories were also used for volcanic SO_2 while wildfire emissions were updated each month from the global fire emissions database version 2 (GFED) (Randerson et al., 2007). We assume that 2.5 % of the SO_2 from anthropogenic, volcanic and wildfire sources is emitted as SO_4 as recommended by Stier et al. (2005) to represent sub-grid nucleation and growth of new particles. The sub-grid nucleated SO_4 particles are partitioned equally between two lognormal size modes of geometric mean diameter 60 and 150 nm, respectively with a geometric standard deviation of 0.59 (Stier et al., 2005).

The microphysical processes in GLOMAP-mode include coagulation between and within modes, wet and dry deposition, ageing of BC/OC mixed particles from an initial insoluble state to water soluble, oxidation of SO_2 into SO_4 in cloud drops, condensation growth between modes (due to uptake of sulphuric acid and secondary organic vapours), and nucleation of SO_4 aerosol (Mann et al., 2010).

Wet deposition occurs in frontal and convective precipitating clouds by nucleation scavenging (due to activation of CCN into cloud drops and subsequent precipitation) and impaction scavenging of aerosol particles by falling raindrops. All water-soluble particles larger than 103 nm wet diameter are assumed to act as activated CCN in grid-boxes where precipitation has been diagnosed. The reason for using a diameter threshold to determine CCN populations is historic stemming from the use of a sectional size-resolving bin scheme in previous model versions. The aerosol wet diameter in GLOMAP is calculated from the dry diameter using the relative humidity diagnosed from the ECMWF temperature, pressure and specific humidity meteorological fields. Thus, at relative humidity between 70 and 80 % this process activates all aerosol with dry diameter greater than \sim 60 nm and at relative humidities greater than 80 % aerosol with diameter greater than ~ 50 nm are activated. The fraction of activated

aerosol removed in a time-step is assumed to be equal to the fraction of condensate precipitated.

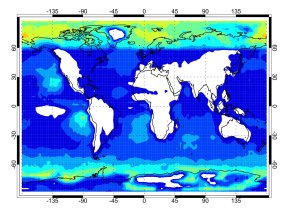
Occurrences of precipitating clouds are calculated from the ECMWF analyses used to drive the model (Pringle, 2006). In convective clouds the fraction of condensate is calculated from the precipitating water downward mass flux parameterised in the model via the methodology of Tiedtke (1989). Convective rain is assumed to be a sub-grid process that occurs in only 30 % of a grid-box (Giannakopoulos et al., 1999). In contrast, frontal rain scavenging is assumed to occur over an entire grid-box. Frontal rain is diagnosed from the ECMWF humidity profile (Mahowald et al., 1997; Giannakopoulos et al., 1999) and is assumed to remove 100 % of CCN during one model time step (Giannakopoulos et al., 1999).

In the baseline model, monthly resolved low cloud climatologies are taken from the International Satellite Cloud Climatology Project (ISCCP) database (Rossow and Duenas, 2004) which provides low-cloud fraction and cloud top pressure (converted in GLOMAP to vertical cloud fraction). Low (boundary layer) clouds are assumed not to precipitate, but only affect aerosol through aqueous phase oxidation of SO₂ to SO₄ by ozone and hydrogen peroxide (Mann et al., 2010). In several of the perturbed runs used in this study the IS-CCP fields are merged at 60° N with data from the MODerate resolution Imaging Spectrometer (MODIS) satellite (see Sect. 4). MODIS has been shown to measure Arctic cloud fraction with greater accuracy (compared to surface climatologies) than other instruments (Liu et al., 2004). This substitution is discussed further in Sect. 4.

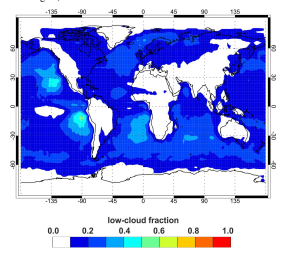
4 Low-level cloud

At high latitudes in the summer, the relatively high humidity allows the formation of low-level stratocumulus cloud decks in the boundary layer and lower troposphere. These low-level clouds and fogs produce frequent drizzle and can be present for as much as 90 % of the time (Curry et al., 1996; Lawson et al., 2001). It has been suggested that the low summertime aerosol concentrations can be explained by this increase in scavenging combined with a decrease in transport from outside the Arctic (Stohl, 2006).

In previous model versions monthly resolved low cloud fraction and cloud top pressure climatologies were sourced from ISCCP (Rossow and Schiffer, 1999). ISCCP cloud fractions north of 60° N have a median cloud fraction from 0 to 0.1 with generally higher fractions in summer and a maximum 95th percentile in April of 0.25 (Fig. 2). Although the temporal pattern in ISCCP cloud fraction is consistent with observations, the magnitude of cloud cover is significantly underestimated compared to observations from the Arctic surface, which rarely fall below 0.50 and can be as high as 0.90 (Curtis et al., 1998; Hahn et al., 1995). The satellite detection of polar clouds is complicated by the thermal and vis-



(a) JJA mean low-cloud fraction (MODIS/ISCCP merged climatologies)



(b) JJA mean low-cloud cover (ISCCP climatologies)

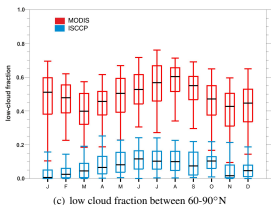


Fig. 2. Comparison of the merged MODIS/ISCCP (see section 4) and ISCCP low-cloud climatologies. (**a–b**) show the JJA average column maximum low cloud fraction in the MODIS/ISCCP (**a**) and ISCCP (**b**) climatologies. Note: white here indicates maximum cloud fraction below 10%. (**c**) compares the MODIS (red) and ISCCP (blue) median low cloud fraction between 60 and 90° N. Errors bar denote the 5th and 95th percentiles for each month, while the box plots show the interquartile range of the median.

ible similarity between clouds and ice-snow surfaces as well as the pervasive temperature and humidity inversions in the Arctic boundary layer (Liu et al., 2004). However, MODIS, which takes a greater frequency of high latitude observations, has been found to measure Arctic cloud fraction and cloud top pressure more accurately then other instruments (Liu et al., 2004).

In this study we merge cloud fractions from MODIS at high latitudes with ISCCP at low latitudes. Monthly resolved $1\times1^\circ$ cloud fraction and cloud top pressure data from the MODIS satellite (Hubanks et al., 2008) were interpolated to a $2.8^\circ\times2.8^\circ$ resolution and merged with interpolated ISCCP climatologies using a exponential function to calculate a weighted mean cloud fraction such that in grid-box $(x_{\text{lon}}, y_{\text{lat}})$ the low cloud fraction (CF $_{x_{\text{lon}}, y_{\text{lat}}}$) was calculated as:

$$CF_{x_{\text{lon}},y_{\text{lat}}} = \frac{M_{x_{\text{lon}},y_{\text{lat}}}e^{-\lambda(y_{\text{lat}}-y_{60})} + I_{x_{\text{lon}},y_{\text{lat}}}e^{-\lambda(y_{60}-y_{\text{lat}})}}{e^{-\lambda(y_{\text{lat}}-y_{60})} + e^{-\lambda(y_{60}-y_{\text{lat}})}} \quad (1)$$

where M and I are the MODIS and ISCCP cloud fractions, respectively and λ is equal to 0.5. Figure 2 shows the median low cloud fraction above 60° N for both the merged and ISCCP low cloud climatologies. In the merged climatologies cloud fraction has increased universally, with a 5th percentile greater than the ISCCP 95th percentile in the summer and greater than the upper quartile for the rest of the year. The use of $\lambda = 0.5$ results in MODIS climatologies replacing ISCCP over only two latitude grid-boxes. However, with this method, median summertime cloud fraction north of 70° N increases to 50–60% equivalent to that observed (Curtis et al., 1998).

We chose to merge the datasets rather than using MODIS climatologies universally as global comparisons of the two datasets show that in the ISCCP climatology areas of higher cloud fraction coincide with the stratocumulus regions described in Leon et al. (2008), while in MODIS maximum cloud fractions occur over the equatorial region and the southern ocean. This disparity would suggest that, despite underestimating the magnitude of cloud fraction at high latitudes, ISCCP performs better than MODIS at lower latitudes. This conclusion is consistent with Leon et al. (2008) who found that ISCCP climatologies compared well with observations from the CloudSat instrument on the CALIPSO satellite.

5 Model scavenging processes

5.1 Ice-cloud scavenging

The in-cloud lifetime of activated CCN with respect to wet scavenging in liquid-phase clouds is of the order 1 h (Wood, 2006; Curry et al., 1996). In addition, in-cloud CCN lifetimes in liquid-phase clouds are generally independent of the cloud precipitation rate as liquid water path scales to the precipitation rate, leading to cancellation in Eq. (2) (see below, Wood,

2006). However, scavenging processes in ice-phase clouds are much less efficient and in-cloud CCN lifetime can extend to 1 month if ice-nucleation is the predominant scavenging process (Davidson et al., 1987; Vinogradova and Ponomareva, 2000). The principal reason for this difference is that precipitation-sized drops in liquid phase clouds grow primarily by collection, a process that collects both cloud droplets and aerosol. Conversely ice crystals tend to form via the nucleation of ice onto individual particles and grow by vapour deposition rather than collision and coalescence. Thus precipitation in liquid-phase clouds can scavenge aerosol particles much more efficiently (Vinogradova and Ponomareva, 2000).

In the standard version of GLOMAP-mode (our control simulations) scavenging rates of soluble aerosol are independent of temperature (see Fig. 3) with all cloud-phases assumed to scavenge equal numbers of soluble aerosol. This assumption is likely to overestimate global scavenging rates particularly in regions where inefficient ice cloud scavenging is prevalent. For insoluble aerosol, a cloud temperature threshold is used to determine scavenging rates. Above -15 °C insoluble BC aerosol is not scavenged and below −15 °C they act as IN and are scavenged at the rate of CCN of the same size (Fan, 2004) (Fig. 3). It is possible that this process overestimates the deposition of insoluble aerosol from ice nucleation. However, the majority of insoluble particles are small and exist in the Aitken mode (freshly emitted BC and OC) and are therefore only weakly scavenged (0.1 % of the total global insoluble BC mass deposited) compared to impaction scavenging (22 %) and dry deposition (77.9 %). Thus, model assumptions about IN scavenging of insoluble BC particles have only a small effect on the model results.

To address the overestimation of soluble particle scavenging rates, we extended the cloud temperature threshold parameterisation also to soluble aerosol. In the absence of a well-defined temperature limit governing the onset of the ice-phase in clouds (Gayet et al., 2009b,a) the same limit of $-15\,^{\circ}\mathrm{C}$ was used to match the existing temperature threshold governing insoluble aerosol scavenging (Fan, 2004). Below this temperature, scavenging of soluble aerosol is prohibited, reflecting the absence of scavenging by coalescence and collision in ice-clouds, while insoluble particles continue (as in the standard model version) to act as ice nuclei (IN) (Baumgardner et al., 2008; Gorbunov et al., 2000).

This simple method of switching between efficient warm-cloud scavenging and effectively non-scavenging ice clouds is expedient in a chemical transport model that does not include the cloud microphysical processes required to predict hydrometeor phase. It is also consistent with the analysis of Garrett et al. (2010), who identified a scavenging temperature switch in Arctic clouds, and the observations of Gayet et al. (2009a) and Curry et al. (1996) which show that Arctic clouds are in general dominated by either ice or liquid phase droplets.

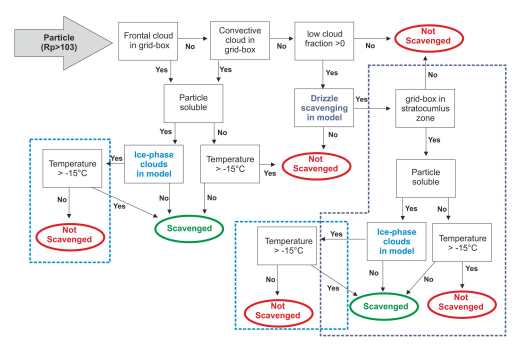


Fig. 3. Flowchart describing GLOMAP-mode nucleation scavenging scheme.

However, observations have shown that in spring mixedphase cloud is also present in the Arctic lower troposphere (Hobbs and Rangno, 1998). Mixed-phase cloud scavenging can occur via nucleation of ice on individual particles (as discussed above) and riming of ice particles with water droplets and is likely to have an efficiency between that of ice-nucleation scavenging and liquid-phase scavenging (Hegg et al., 2011) Therefore, our treatment of mixed-phase clouds as liquid-phase is likely to overestimate the scavenging rate in spring.

Mixed-phase cloud formation can occur over a wide temperature range. The scavenging efficiency of individual clouds is dependent on the ratio of liquid-to-total water, which is affected by physical processes such as the Wegener-Bergeron-Findeisen process and can be highly variable over limited temperature ranges (Smith et al., 2009). In models that include mixed-phase clouds (Bourgeois and Bey, 2011) assumptions still need to be made about scavenging rates that are not treated explicitly, and agreement with observations is not necessarily improved. Our switch from liquid to icephase at $-15\,^{\circ}\text{C}$ is consistent with what is seen in observations (Garrett et al., 2010), and is a reasonable assumption until the physical processes are resolved better. The implications of the exclusion of mixed-phase scavenging are discussed below (Sect. 7.3.2).

Figure 4 shows the percentage of total precipitation falling to the surface as snow in GLOMAP-mode as defined by this ice-cloud temperature threshold. The ratio of modelled snow to rain is highest in the Arctic winter and early spring, where 90% of precipitation falls as snow above 60° N in Asia and above 70° N in Canada and Alaska. Comparison

of the snow to rain ratio at different Arctic ground-stations indicates that, while in the winter Alert and Barrow are positioned in regions of negligible warm cloud scavenging, at Zeppelin, Oulanka, Karasjok and Janiskoski warm rain scavenging remains a significant removal mechanism throughout the winter and spring.

5.2 Low-cloud scavenging by drizzle

Local scavenging of aerosol in the Arctic can be caused by persistent light drizzle and sleet in the boundary layer as well as episodic dynamic rain fronts directed into the high latitudes by large scale meteorological systems. In GLOMAP, low-cloud precipitation is not treated in the standard model. Thus episodic large scale meteorological systems dominate scavenging north of 60° . More dynamic rain fronts reach the Arctic in winter than summer, although liquid precipitation (rain) is more likely in summer, with median values in summer of $\approx 1.5 \ \text{mm} \ \text{d}^{-1}$ compared to $0.5 \ \text{mm} \ \text{d}^{-1}$ in winter.

The reason for neglecting low cloud scavenging in GLOMAP-mode is historic: GLOMAP diagnoses precipitation in frontal clouds and convection (see Sect. 3), and wet deposition by these processes alone results in aerosol lifetimes of ~ 1 day that are well within the range of other models (Textor et al., 2006). In addition, given the uncertainties in marine aerosol and precursor emissions in regions where stratocumulus clouds are prevalent, it has not been possible to detect missing sinks when evaluating the model against observations (Spracklen et al., 2007; Korhonen et al., 2008a) (see also Sect. 7.5). Nevertheless, drizzle is a common occurrence in the summertime Arctic boundary

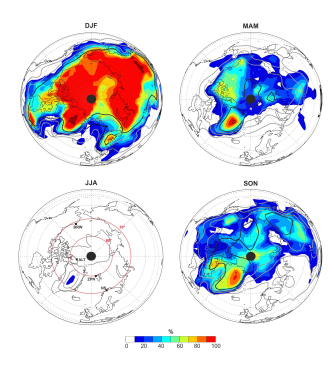


Fig. 4. Percentage of seasonal (DJF, MAM, JJA and SON) mean precipitation in an atmospheric column (kg m $^{-2}$ d $^{-1}$) formed at temperatures below $-15\,^{\circ}$ C and so identified in GLOMAP-mode as snow. Also shown are the surface $-15\,^{\circ}$ C (black solid line), $-10\,^{\circ}$ C, $-5\,^{\circ}$ C and $0\,^{\circ}$ C (grey solid lines as labelled) isotherms. In the bottom left-hand figure the positions of three Arctic ground-stations: Alert (ALT), Barrow (BRW) and Zeppelin mountain (ZPN) are shown as is the approximate position of three closely located Northern Scandinavian sites: Oulanka, Karasjok and Janiskoski (NS).

layer and, although it is not the dominant wet removal process on a global scale, it is likely to be an important scavenging process in regions such as the Arctic which have limited frontal and convective rainfall (Curry et al., 1996; Hobbs and Rangno, 1998; Hobbs et al., 2001; Lawson et al., 2001; Gayet et al., 2009a).

To address this model deficiency a low-cloud scavenging scheme was added to the model. Low-cloud drizzle is assumed to occur in grid-boxes without convective or frontal rainfall and with a low-cloud fraction greater than zero (see Fig. 3). The CCN lifetime is assumed to be equal to the condensate lifetime, which is calculated using the methodology of Wood (2005):

$$\tau_{\rm driz} = \frac{\rm LWP}{R_{\rm driz}} \tag{2}$$

where LWP is the low-cloud liquid water path (kg m⁻²) and $R_{\rm driz}$ is the drizzle rate at cloud base (kg m⁻² s⁻¹). This removal rate is applied to a fraction of the CCN in each grid box according to the low cloud fraction (Sect. 4). We assume that the drizzle does not evaporate, thus our CCN scavenging rates are an upper limit for the given condensate lifetime.

Low cloud drizzle rates and LWP over mid-latitude and tropical regions were specified in the model based on observations from satellite climatologies (Leon et al., 2008) and are described fully in Table 2. In the Arctic, while measurements of LWP are documented in the literature, observations of low-cloud precipitation rates are to the best knowledge of the authors not available. Arctic drizzle rates were therefore estimated using the empirical relationship between cloud thickness (z) and cloud droplet number concentration ($N_{\rm d}$) observed by van Zanten and Stevens (2005) and calculated using the methodology of Wood (2006), such that

$$R_{\text{driz}} = \frac{K_{\text{VZ}}z^3}{N_{\text{d}}},\tag{3}$$

where $K_{\rm VZ}=1.9\times10^{-5}\,{\rm kg\,m^{-8}\,s^{-1}}$ is an empirically derived constant. This empirical relationship between drizzle rates and stratocumulus cloud properties was based on observations of nocturnal marine stratocumulus clouds off the Californian coast (van Zanten et al., 2005) and evaluated against observations off the European coast (Wood, 2005) and over the North Atlantic (Pawlowska, 2003). Derived drizzle rates were found to correlate strongly with observations (R=0.97 and 0.79, respectively), although estimated N. Atlantic drizzle rates were biased low against observations ($-79\,\%$).

Analysis of observations over the Beaufort Sea (70–76° N, $124–127^\circ$ W) of summertime Arctic cloud thickness and drop concentration in precipitating low-cloud (Hobbs and Rangno, 1998) yield a summertime Arctic drizzle rate of $0.44~\rm kg~m^{-2}d^{-1}$. In spring no precipitating clouds were observed but an average cloud droplet number of $73~\rm cm^{-3}$ was calculated in low clouds with a non-trivial liquid water path. In conjunction with data from Hobbs et al. (2001), which suggest an average cloud depth of $150~\rm m$ in the Arctic springtime boundary layer, these observations yielded a low cloud springtime drizzle rate of $0.08~\rm kg~m^{-2}d^{-1}$ albeit with a high degree of uncertainty.

Arctic drizzle rates calculated using Eq. (3) (scaled with mean low cloud fraction of 80–90%) compare reasonably well to averaged precipitation measurements at Barter Island (70° N, 143° W) and Barrow of 0.6 kg m $^{-2}$ d $^{-1}$ in summer and 0.08 kg m $^{-2}$ d $^{-1}$ in spring (Curtis et al., 1998) assuming that the majority of measured precipitation is drizzle. However at northern European sites measured precipitation rates are higher than those calculated (0.3–1.0 kg m $^{-2}$ d $^{-1}$ in spring and 1.7–2.0 kg m $^{-2}$ d $^{-1}$ in summer). It is unlikely that all of the precipitation measured is drizzle. Thus, the spring and summer drizzle rate in the European Arctic (defined in Table 2) was changed to 0.42 kg m $^{-2}$ d $^{-1}$ and 0.8 kg m $^{-2}$ d $^{-1}$, respectively. The effect of this increase on CCN grid-box lifetime is shown in Fig. 5.

Table 2 summarizes the modelled and observed drizzle rates and LWP in the stratocumulus zones (used to calculate in-cloud CCN lifetime as in Eq. 2). Figure 5 shows the grid-box CCN lifetime with respect to drizzle scavenging (incloud CCN lifetime divided by low cloud fraction). Outside

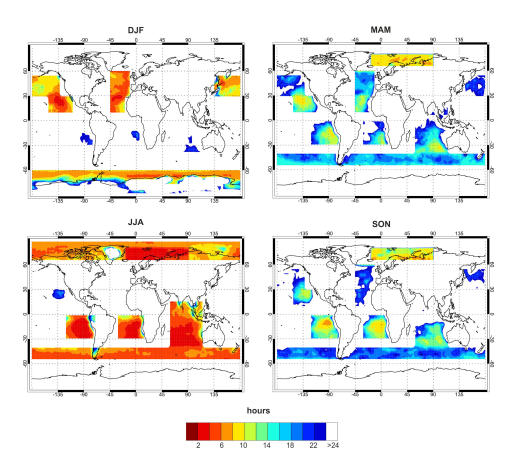


Fig. 5. Comparison of the CCN grid-box lifetime with respect to low cloud drizzle scavenging. Note that the colour scale is reversed from that in previous figures with red denoting a shorter CCN lifetime.

the main stratocumulus zones the minimum grid-box CCN lifetime is greater than 24 h and in most regions exceeds 4 days. Although Arctic drizzle rates are lower than in any other stratocumulus zones (Table 2), the higher low-cloud fraction introduced north of 60° N results in shorter average Arctic CCN grid-box lifetime from June–August (of 2–6 h) equivalent to that in the southern ocean stratocumulus zone.

The importance of the drizzle rate to the scavenging efficiency in stratocumulus zones is discussed below in several sensitivity experiment.

6 Description of model experiments

Five sets of model simulations were completed to determine the importance of ice and low cloud scavenging processes for Arctic aerosol concentrations. All five simulations use identical meteorological fields and emission inventories and were spun up for 3 months (October–December 1999) before the beginning of each 3-yr period (2000–2002). The simulations are described below and summarized in Table 1:

- CTRL-ISCCP: a control run using global ISCCP low cloud climatologies including convective and frontal rain scavenging but without low-cloud precipitation. Scavenging of soluble aerosol is permitted to occur in both warm-phase and ice-phase clouds (see Fig. 3). This run uses the same set-up as Mann et al. (2010).
- CTRL-MODIS: a control run using the merged MODIS/ISCCP low-cloud climatologies as described in Sect. 4. Convective and frontal rain scavenging processes are identical to CTRL-ISCCP and low-cloud precipitation is likewise excluded. The merged cloud fields will be used in the remainder of the simulations, so this simulation provides a new baseline in place of CTRL-ISCCP.
- DRIZZ: this is the same as CTRL-MODIS except that low-cloud drizzle scavenging in the Arctic and stratocumulus cloud zones is permitted (as described in Sect. 5.2) with the restrictions outlined in Fig. 3. Convective and frontal rain scavenging processes remain identical to both CTRL-ISCCP and CTRL-MODIS.

Run Name	low cloud climatology	low cloud	convective and frontal cloud	ice-cloud (frontal and convective)	ice-cloud (low)
CTRL-ISCCP	ISCCP	_	X	_	_
CTRL-MODIS	ISCCP/MODIS	_	X	_	_
DRIZZ	ISCCP/MODIS	X	X	_	_
ICE	ISCCP/MODIS	_	X	X	_
DRIZZICE	ISCCP/MODIS	X	X	X	X

Table 1. Description of core simulations. Included cloud scavenging processes in each simulation are marked with an X.

Table 2. Drizzle rate $(kg m^{-2}d^{-1})$ and cloud liquid water path $(g m^{-2})$ in 11 stratocumulus zones used in drizzle parametrisation sourced from Leon et al. (2008) as observations (obs) or calculated from cloud characteristics (estimated).

Region	Winter $kg m^{-2} d^{-1}$	$\begin{array}{c} \text{Spring} \\ \text{kg m}^{-2} \text{d}^{-1} \end{array}$	$\begin{array}{c} \text{Summer} \\ \text{kg m}^{-2} \text{d}^{-1} \end{array}$	$\begin{array}{c} \text{Autumn} \\ \text{kg m}^{-2} \text{d}^{-1} \end{array}$	$\begin{array}{c} LWP \\ gm^{-2} \end{array}$	source $R_{\rm driz}/{\rm LWP}$	Latitude	Longitude
N.Atlantic	2.8	0.8	0.29	0.8	144	obs /obs	30–60° N	10–45° W
N.pacific	2.3	0.68	0.29	0.68	131	obs/obs	30–55° N	130° W-105° E
Australian	2.3	0.57	0.26	0.57	90	obs/obs	40°S–15° N	60–115° E
Circumpolar	2.5	0.74	0.28	0.74	150	obs/obs	55–40°S	180° E−180° W
Nambian	1.7	0.55	0.24	0.55	84	obs/obs	0-30°S	30° W−15° E
Canarian	2.1	0.61	0.25	0.61	77	obs/obs	10–30° N	45–20° W
Peruvian	1.9	0.6	0.26	0.6	91	obs/obs	0-30°S	70–120° W
Californian	1.9	0.57	0.25	0.57	90	obs/obs	10-30° N	110–150° W
Arctic	0.0	0.08	0.42	0.08	68	estimated/obs	60–90° N	180° E–180° W
Euro Arctic	0.0	0.42	0.8	0.42	68	estimated/obs	60–80° N	15° E–150° W
Antarctic	0.0	0.08	0.42	0.08	68	Arctic values used	60-90°S	180° E-180° V

- ICE: this is the same as the CTRL-MODIS run except that ice-cloud scavenging of soluble aerosol is prohibited at temperatures less than -15 °C. As in CTRL-MODIS low clouds do not precipitate, the ice-cloud effect is applied to convective and frontal rain-clouds only.
- DRIZZICE: a combination of the DRIZZ and ICE runs, which includes merged MODIS and ISCCP low-cloud climatologies parameterisation of low-cloud drizzle in the Arctic and stratocumulus zones. To represent the slow removal rate in ice-phase clouds, scavenging of soluble aerosol is prohibited at $T < -15\,^{\circ}\mathrm{C}$ in low, frontal and convective clouds.

7 Results and discussion

7.1 Control runs

We first compare SO₄ and BC concentrations in the two control runs using different high latitude low-cloud climatologies, CTRL-ISCCP and CTRL-MODIS (Fig. 6). In these runs the only effect of the low-level (non-drizzling) clouds is on SO₄ production due to in-cloud oxidation and the growth of aerosol by aqueous phase reactions in non-precipitating clouds or cloud processing (Mann et al., 2010).

Predicted Arctic surface concentrations of BC mass are similar in CTRL-MODIS and CTRL-ISCCP throughout the year although the lower quartile from 60 to 90° N from December to February is typically lower in CTRL-MODIS (Fig. 6c). Likewise surface concentrations of SO₄ are typically lower north of 60° in CTRL-MODIS compared with CTRL-ISCCP (Fig. 6a). This mass decrease is the result of an increase in the rate of model cloud processing which counterintuitively decreases aerosol mass concentrations by decreasing SO₄ and BC aerosol lifetime with respect to wet deposition (through changes in the aerosol size distribution).

Compared with surface observations, both control runs predict the seasonal SO_4 mass concentration variation poorly. At Karasjok, Oulanka and Janiskoski (Fig. 7d), where observations show no seasonal variation in SO_4 mass concentrations, both control runs have a summertime maximum that is up to a factor 10 greater than observed. At all three stations SO_4 mass increases slightly with the use of MODIS cloud climatologies. This increase is small in the summer but is significant in the early winter (40–50%) and stems from a greater SO_2 oxidation to SO_4 flux (100–200%) in the vicinity of Finland and Northern Norway, which mitigates the regional wet deposition resulting from increases in cloud processing.

At Alert, Barrow and Zeppelin the SO₄ mass concentration seasonal maximum occurs in early summer and is 3–4 months out of phase with the observations (Fig. 7a–c). In

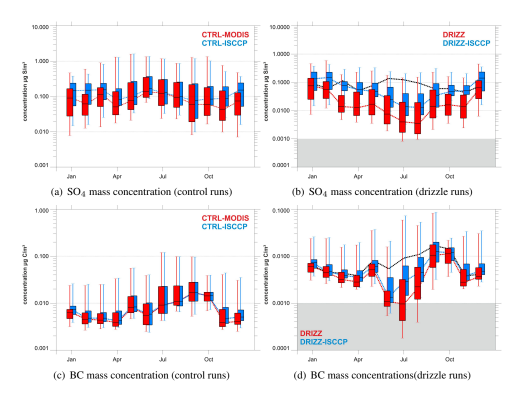


Fig. 6. Comparison of median and interquartile range SO_4 (**a, b**) and BC (**c, d**) surface mass concentration (in the year 2000) between 60° and 90° N for control runs CTRL-MODIS (red) and CTRL-ISCCP (blue) and drizzle runs DRIZZ (red) and DRIZZ-ISCCP (blue). The Errors bar denote the 5th and 95th percentiles in each month. Note: the ISCCP simulation results have been offset along the x-axis, this was done to maintain clarity and does not indicate any difference in temporal resolution. In (**c-d**) the median SO_4 and BC surface mass concentration in the CTRL-MODIS simulation are also shown (black dashed lines).

summer, SO₄ mass concentrations at Alert and Barrow fall within one standard deviation of the observed mean, but at Zeppelin simulated SO₄ concentrations from June to August are up to 4 times greater than measured. In winter modelled SO₄ concentrations at Alert and Barrow are more than 2 standard deviations from the observational mean while at Zeppelin the model is less than 0.1 standard deviations from the observation mean (see Table 4).

The model prediction of BC mass concentrations and EBC data at Alert and Barrow is worse than for SO₄ (Fig. 7e–f). The model is about 6 months out of phase, with a significant over-prediction in summer (a factor of 2.3σ from the observation JJA mean at Barrow) and under-prediction in winter. The simulated summer BC maximum at both sites is skewed by anomalously high concentrations in August 2001. This maximum is due to the monthly updated GFED biomass burning (BB) emission inventories used in GLOMAP-mode. The monthly emission inventories assume a constant emission rate over the month. This assumption emphasises the contribution of forest fires which occur when meteorological conditions (which are updated every 15 min) suppress the transport of BB emissions to ground-station sites. The overestimation of BB aerosol concentrations at these sites could be reduced by using daily resolved wildfire emission inventories in the model, however no such inventories were available to the authors pre-2008.

In summary, irrespective of the cloud climatology used, GLOMAP-mode fails to simulate the observed seasonal cycle, with insufficient BC and SO_4 mass in the winter and early spring but too much BC and SO_4 in the summer. The use of MODIS climatologies north of 60° has had a negligible effect on modelled SO_4 and BC surface mass concentration. However, the use of a different low-cloud climatology is likely to be of greater impact in simulations with low-cloud scavenging included, which we address below.

7.2 Simulation DRIZZ

7.2.1 Effect of drizzle scavenging at the Arctic sites

Scavenging due to drizzle reduces SO_4 aerosol concentrations at all Arctic sites between March and November, with the greatest effect occurring in summer when Arctic drizzle rates are at a maximum (Fig. 7). By disproportionately reducing summertime concentrations, drizzle scavenging goes some way to correcting the negative correlation across the seasonal cycle. At stations north of 70° N the R correlation coefficient increases from -0.39 (CTRL-MODIS) to 0.47

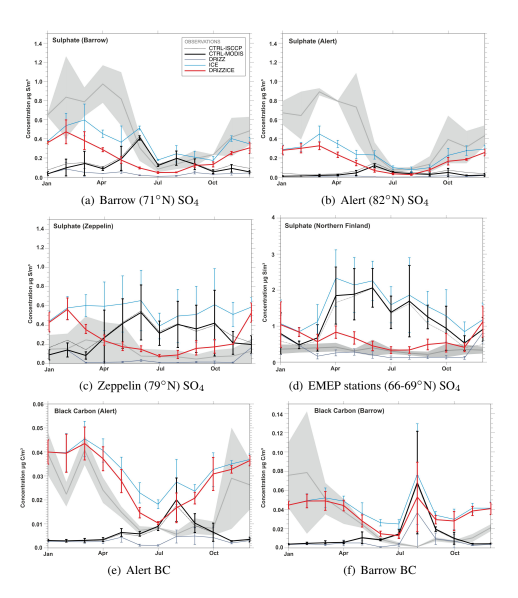


Fig. 7. Monthly mean SO₄ and BC surface mass concentration at three Arctic ground-stations averaged over the model run period 2000–2002. Mass observations are shown in grey with the monthly inter-annual standard deviation indicated by light grey shading. Observed SO₄ and BC mass concentrations are compared with both control runs (MODIS-black, ISCCP-black dashed), DRIZZ (dark blue), ICE (light blue) and DRIZZICE (red). Error bars represent the inter-annual standard deviation between yearly model runs. In (d) monthly SO₄ mass concentration observations at the three EMEP stations (Karasjok, Janiskoski and Oulanka from 2000–2002) have been averaged and compared with modelled surface concentrations over Northern Norway and Finland (see Fig. 1).

(DRIZZ) at Alert, from -0.23 to 0.4 at Barrow and from -0.43 to 0.11 at Zeppelin. However, year-round drizzle also exacerbates the low model bias in winter, spring and autumn, increasing the annual mean negative bias to -97% at Alert and to -89% at Barrow (Table 3). At Zeppelin (Fig. 7c) drizzle scavenging has significantly decreased model bias, with SO_4 concentrations annually underestimated by 60% in DRIZZ simulations.

At marginal Arctic sites (Karasjok, Janiskoski and Oulanka see Fig. 7d) modelled SO₄ mass in DRIZZ is now on average less than one inter-annual standard deviation from

observations (-0.4, -0.1 and -0.3, respectively) and the model-observation bias is about -10 to -20% (Table 3). Despite the consistent negative bias at all three stations the DRIZZ simulation is a considerable improvement on the average 200-300% positive bias in the control simulations. This analysis suggests that, while drizzle scavenging has very different effects on SO_4 at the five Arctic sites, in general the effect of drizzle in the Arctic is to push modelled SO_4 concentrations below observations.

Like SO₄, Arctic BC mass concentrations have decreased in the DRIZZ simulation compared to CTRL-MODIS. At

Barrow and Alert the correlation between observations and DRIZZ is still negative, as in CTRL-MODIS, but summer BC mass concentrations have decreased by a factor of ~ 3 in DRIZZ, reducing the average bias from 230 % to 80 %. However, at Alert the negative model bias has increased in DRIZZ from -40 % to -75 % and summertime BC mass concentrations are now -1.4 standard deviations from the observation mean (Tables 3 and 4).

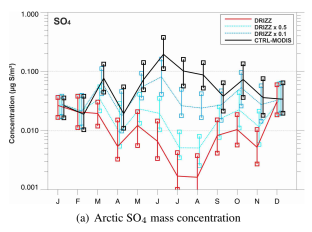
7.2.2 How sensitive is Arctic aerosol to the Arctic drizzle rate?

There are two significant uncertainties in our modelling of Arctic drizzle: the uncertainty in low-cloud cover (ISCCP versus MODIS) and the uncertainty in stratocumulus drizzle rates. The sensitivity of surface SO₄ and BC concentrations to global drizzle rates was tested using four additional simulations: DRIZZx0.1 where global drizzle rates were decreased by a factor of 10, DRIZZx0.5 where drizzle rates were decreased by a factor of 2, DRIZZx2 where drizzle rates were increased by a factor of 2 and DRIZZx10 where drizzle rates were increased by a factor of 10.

Changing global drizzle rates has a large effect on median surface SO_4 mass concentrations north of 70° (Fig. 8a). Halving drizzle rates increases Arctic surface SO_4 concentrations from March to November by a factor of 2 and 3 (March–April and June–August, respectively) compared to DRIZZ simulations, while dividing drizzle rates by 10 increases summertime SO_4 by a factor ~ 10 . However, even with the very low scavenging rates in the DRIZZx0.1 simulations, SO_4 concentrations are still a factor of 3 lower than in the CTRL-MODIS simulations without any drizzle. Thus, Arctic SO_4 surface concentrations are highly sensitive to even very low drizzle rates.

Arctic surface BC concentrations are less sensitive to the drizzle rate than SO_4 , with median summertime concentrations above 70° N in DRIZZx0.1 increasing by a factor of ~ 3 compared to DRIZZ (Fig. 8b). The effect on the bias at Alert and Barrow is also negligible. In August all three drizzle runs have similar median BC concentration to the control (Sect. 7.1). The negligible effect of drizzle scavenging is confined to August and stems from a high fraction of hydrophobic BC in the Arctic BC load. Therefore, we suggest that BC concentrations during this month are sourced from biomass burning emissions (see Sect. 7.1) transported rapidly to the Arctic and unaffected by ageing processes (see Sect. 7.4 for further discussion).

We note that at Barrow and Alert the DRIZZx0.1 simulation is in much better agreement with observed summertime SO_4 than the DRIZZ simulation (mean JJA bias reduced from -93% to -30% and from -96% to -56%, respectively), but results in a large positive bias at the other sites (as in CTRL-MODIS). However, we do not draw conclusions about the best model simulation until we introduce the effect of reduced scavenging in ice clouds (Sect. 5.1). The question



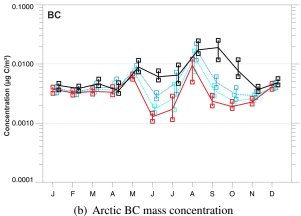


Fig. 8. Median SO₄ (a) and BC (b) mass concentration north of 70° for CTRL-MODIS (black line), DRIZZ (red) and two drizzle sensitivity runs where global drizzle rates have been multiplied by 0.5 and 0.1 (blue dashed lines). Error bars represent the lower and upper quartile or the interquartile range. Note: all datasets have been offset from DRIZZ results along the x-axis so that the figure remains clear but each represents the same time interval.

we now ask is from where this sensitivity to the drizzle rate arises: from changes to scavenging close to source regions in the extra-Arctic or local changes in the high-latitudes.

7.2.3 Where does drizzle affect Arctic aerosol?

The DRIZZ simulation includes drizzle scavenging globally, thus reductions in Arctic aerosol will be due to scavenging close to low-latitude sources, during transport from outside the Arctic, and at high latitudes. The relative effects of scavenging in these regions will likely vary with season and location because of seasonal variations in local and high latitude aerosol sources, cloud fractions, and precipitation rates. To determine the relative importance of drizzle north and south of 60° we performed an additional model experiment in which drizzle occurs only between 0 and 60°.

We found that concentrations of BC and SO₄ in this run differed negligibly from CTRL-MODIS at all six Arctic

site	species	CTRL-ISCCP		CTRL	-MODIS	DI	RIZZ		ICE	DRIZZICE	
		R	bias (%)	R	bias (%)	R	bias (%)	R	bias (%)	R	bias (%)
Alert (2000, 2002)	BC	-0.68	-43.7	-0.69	-39.8	-0.29	-74.58	0.65	92.7	0.72	58.95
Barrow	BC	-0.42	231.33	-0.43	723.46	-0.14	84.8	0.29	461.6	0.44	315.7
Zeppelin	SO_4	-0.42	306.8	-0.43	288.2	0.11	-61.9	0.03	530.2	0.4	149.7
Alert	SO_4	-0.33	-67.6	-0.39	-76.2	0.47	-97.1	0.72	-18.4	0.71	-52.1
Barrow	SO_4	-0.16	-36.6	-0.23	-40.9	0.40	-89.2	0.58	11.5	0.70	-46.9
Karasjok	SO_4	0.31	250.3	0.37	287.3	0.2	-17.7	0.48	397.9	0.46	98.9
Oulanka	SO_4	0.04	252.0	0.11	289.3	0.27	-17.9	0.2	377.5	0.50	82.8
Janiskoski	SO_4	0.43	188.4	0.47	222.9	0.12	-8.6	0.51	305.6	0.27	82.7

Table 3. Arctic Model-observation (ground-station) *R* correlation and mean bias for CTRL-ISCCP, CTRL-MODIS, ICE, DRIZZ and DRIZZICE simulations from 2000 to 2002 (with the exception of Alert where no BC observations from 2001 where available).

Table 4. Arctic model-observation error calculated as the mean difference between the model and observations divided by the standard deviation of the observed monthly mean at the ground-stations Alert, Barrow and Zeppelin. Data is shown averaged over the total time period 2000–2002 and for the DJF and JJA seasonal means.

simulation	Alert (BC)			Ва	Barrow (BC)			Alert (SO ₄)			Barrow (SO ₄)			Zeppelin (SO ₄)		
	tot	DJF	JJA	tot	DJF	JJA	tot	DJF	JJA	tot	DJF	JJA	tot	DJF	JJA	
CTRL-MODIS	-0.3	-1.0	0.005	0.03	-1.5	2.6	-2.8	-2.9	-0.8	-1.4	-2.5	0.6	4.0	-0.02	7.4	
DRIZZ	-1.0	-1.0	-1.4	-0.5	-1.6	1.0	-3.5	-2.9	-2.6	-2.2	-2.6	-1.2	-1.0	-0.2	-1.3	
ICE	1.8	0.02	0.9	1.8	-0.05	4.0	-0.7	-1.3	1.2	-0.07	-0.9	1.3	8.1	5.4	9.7	
DRIZZICE	1.1	0.02	0.9	1.3	-0.05	2.3	-1.9	-1.3	-1.3	-1.1	-1.0	-0.7	2.4	5.0	0.9	

ground stations. This result implies that drizzle at low latitudes ($<60^{\circ}$ N) is unimportant for Arctic aerosol. That is, reduced aerosol concentrations in the extra-Arctic source regions are not the explanation for reductions in Arctic aerosol when drizzle is added to the model. This conclusion is further supported by the negligible change in SO₄ and BC surface concentrations north of 60° from December to February (when Arctic drizzle rates are set to 0) shown in Fig. 9a–b even after global drizzle rates are raised by a factor of 10. Thus, any change in Arctic aerosol surface concentrations can be attributed to scavenging occurring at high latitudes.

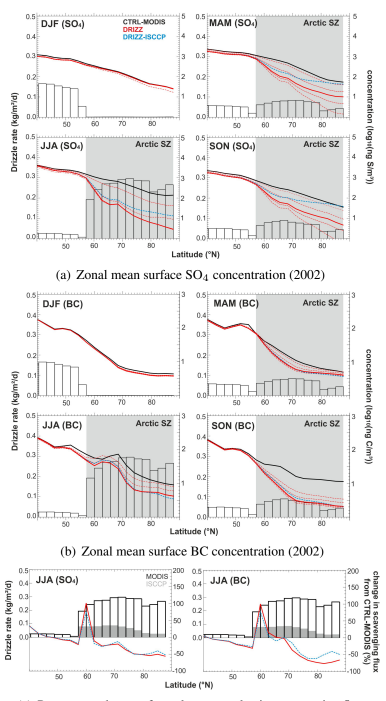
We now address the question of where low-cloud scavenging occurs as aerosol is transported north of 60° . Figure 9a–b shows the zonal mean surface SO₄ and BC concentration for the simulations CTRL-MODIS, DRIZZ and all four drizzle rate sensitivity runs in the year 2002. Also shown is the effect of replacing the MODIS cloud climatologies with ISCCP at high latitudes.

In the Arctic summer the cumulative effect of drizzle scavenging causes a decrease in surface SO_4 concentrations by an order of magnitude between the edge of the stratocumulus zone and 65° N (Fig. 9a). This decrease is in addition to the approximate factor 2 decrease due to other deposition processes (shown by the CTRL-MODIS black line). Thus drizzle is a dominant deposition process during this period. The effect on BC concentrations is less than for SO_4 , but mass concentrations are still reduced by an order of magnitude between 60 and 75° N.

Figure 9a–b shows that increases in the summertime drizzle rate beyond those used in the DRIZZ simulations have a negligible additional effect because the rate is already very high. Halving the drizzle rate hardly affects the latitude at which BC and SO₄ surface concentrations have fallen by an order of magnitude. However, the cumulative effect of the decreased drizzle rate is to approximately double SO₄ and BC mass concentrations north of 80°. In DRIZZx0.1 simulations, summertime zonal mass concentrations have increased by an order of magnitude compared to DRIZZ. Nevertheless, the average CCN boundary layer grid-box lifetime in these very low scavenging simulations is still $\sim\!2$ days, and SO₄ mass concentration still decrease by a factor of 3 below CTRL-MODIS by 80° N, and by a factor 10 taking into account all scavenging processes.

In addition Fig. 9c shows that substituting MODIS clouds with ISCCP reduces the scavenging and increases the SO₄ concentrations at 75° N by about a factor of 2. However, despite the large difference in cloud cover, the latitude by which aerosol concentrations have fallen by an order of magnitude remains approximately the same.

The overall effect of introducing drizzle is to improve the correlation of modelled and observed SO_4 and BC concentrations at all sites (by suppressing spring-to-autumn concentrations more than winter), but it results in large negative biases. The underestimation of SO_4 and BC mass concentrations north of 70° N may be caused by a too high Arctic low-cloud drizzle rates or result from our failure to account



(c) Percentage change of zonal mean nucleation scavenging flux

Fig. 9. Histogram of zonal drizzle rate (kg m $^{-2}$ d $^{-1}$)) compared with zonal SO₄ (**a**) and BC (**b**) surface mass concentration (log₁₀(ng *X* m $^{-3}$) in CTRL-MODIS (black) and DRIZZ (red). In addition, also shown are results from four sensitivity studies where global drizzle rates have been homogeneously multiplied by 0.1, 0.5, 2.0 and 10.0 (red dashed lines) and from the drizzle simulation DRIZZ-ISCCP which used ISCCP cloud climatologies north of 60° N (blue dashed line). Note: drizzle rates in (**a**) and (**b**) have been calculated using the merged MODIS and ISCCP cloud climatologies discussed in Sect. 4. (**c**) shows the JJA zonal drizzle rate calculated using both MODIS/ISCCP merged fields and ISCCP climatologies (gray) compared with the percentage change in the column integrated SO₄ (left) and BC (right) nucleation scavenging rate from CTRL-MODIS in DRIZZ (red) and DRIZZ-ISCCP (blue) simulations. Grey shading indicates the latitudinal extent of the Arctic stratocumulus zone.

for re-evaporation processes in drizzling clouds. However, sensitivity studies suggest that drizzle rates would need to decrease by a factor of 10 in order to agree with measurements at Alert and Barrow. This drastic reduction results in an overestimation of summertime SO₄ mass concentrations at Zeppelin, Oulanka, Janiskoski and Karasjok.

Our analysis suggests that Arctic drizzling low cloud acts as a filter, strongly suppressing the transport of aerosol within the lower troposphere between the marginal (60–70° N) and high (70° N) Arctic. In addition, it is likely that the cloud layer acts to filter air vertically mixed down from the upper troposphere, reducing the effect of high altitude aerosol plumes (Warneke et al., 2010) on boundary layer concentrations. The spatial extent and efficiency of this filter is dependent on the imposed drizzle rate and stratocumulus boundary in the model as well as the height and thickness of the simulated cloud layer.

Although Arctic drizzle is an additional deposition process in the model, it is sufficiently rapid at the edge of the Arctic that net deposition of SO₄ and BC is enhanced by drizzle only between about 60 and 70° N, but decreased at higher latitudes (Fig. 9). This result suggests that the majority of aerosol transported into the Arctic boundary layer enters horizontally through the lower troposphere/boundary layer and is quickly deposited to the surface after vertical mixing into the cloud layer. In the next section we introduce a temperature threshold for all nucleation scavenging in GLOMAP-mode.

7.3 Simulation ICE

7.3.1 Effect of reduced ice scavenging

The ICE run is based on CTRL-MODIS (i.e., no drizzle) but includes a complete suppression of scavenging of soluble aerosol below $-15\,^{\circ}\text{C}$ (see Fig. 3). At Alert and Barrow (Fig. 7) this ice-cloud scavenging scheme has raised wintertime aerosol concentrations by up to a factor of 6 resulting in a SO₄ and BC seasonal cycle that is in phase with observations. The R correlation between the model and SO₄ observations has increased to 0.72 at Alert and to 0.58 at Barrow. The EBC *R* correlation has increased to 0.65 at Alert and 0.29 at Barrow. In contrast, at Zeppelin ice-cloud scavenging, while increasing sulphate mass, has done little to correct the modelled seasonal cycle, reducing the *R* correlation to 0.03 and increasing model bias from 300 % to 530 % (see Table 3).

At Karasjok, Janiskoski and Oulanka the effect of suppressed ice-cloud scavenging is much smaller than at the other sites, increasing SO₄ by a few percent in summer and a maximum of a factor 2 in winter. These changes in SO₄ aerosol at the different sites are broadly consistent with the column-mean fraction of ice cloud in the vicinity and southward of the sites. Alert and Barrow are well within the region of high ice-cloud occurrence in winter and even in spring and autumn (Fig. 4) while the other sites are only marginally

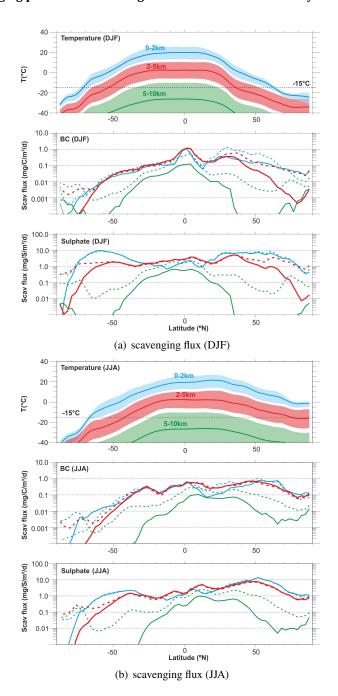


Fig. 10. Zonal mean temperature (top) between 0–2 km (blue), 2–5 km (red) and 5–10 km (green) compared with BC (middle) and SO_4 (bottom) zonal mean nucleation scavenging flux (mg X_s m⁻² d⁻¹) in CTRL-MODIS (dashed line) and ICE (solid line) integrated between equivalent atmospheric regions. The shading in the zonal mean temperature plots indicates the minimum and maximum temperature in each atmospheric region. (a) shows analysis of the December–February seasonal mean while (b) shows analysis of the June–August mean.

within this region in winter and well outside it in other sea-

We can analyse the vertical profile of temperature and the column-integrated nucleation scavenging flux (mg m $^{-2}$ d $^{-1}$) to determine where the effect of the ice clouds is greatest. Figure 10 shows the zonal mean temperature and SO₄ and BC integrated scavenging fluxes in CTRL-MODIS and ICE for the altitude levels 0–2 km (boundary/residual layer), 2–5 km (lower troposphere) and 5–10 km (mid-upper troposphere), the sum of which is equivalent to the deposition per m 2 per day. In the ICE simulation, the wintertime SO₄ scavenging flux decreases considerably above the boundary layer for latitudes greater than about 50° N, corresponding to locations with mean temperatures below -15 °C. In summer, the effect of the ice scavenging is evident only above \sim 5km altitude, and therefore has a minimal effect on surface level aerosol (Fig. 7).

In the wintertime boundary layer the SO₄ scavenging rate does not significantly change between CTRL-MODIS and ICE, but falls substantially at higher altitudes. In contrast, boundary layer BC scavenging decreases by a factor ~4 between 30 and 60° N in the ICE simulation and increases by up to a factor 3 above 70° N. This change in BC scavenging at temperatures where the ICE parameterisation ought to have a negligible effect is due to the response of the mixing state of the BC particles to changes in SO₄. In CTRL-MODIS the majority of BC mass in the Arctic is insoluble (\sim 80 %) and within the Aitken mode (\sim 90 %) while in the ICE simulation BC mass is predominantly soluble (70-90 %) and split evenly between the Aitken and accumulation modes. Therefore, annually a greater proportion of BC mass is scavengable in ICE then in CTRL-MODIS, resulting in an increase in BC scavenging rates above 70° N. The ratio between Aitken mode and accumulation mode SO₄ mass has also changed, with a greater proportion of boundary layer SO_4 mass within the Aitken mode ($\sim 10\%$). However, this small decrease in accumulation mode mass has done little to alter the SO₄ scavenging flux between 0 and 2 km.

The shift in BC characteristics between CTRL-MODIS and ICE suggests that in ICE a greater proportion of Arctic BC aerosol is composed of aged particles transported from latitudes below 60° N. While the increase in SO₄ mass concentration at all stations coupled with a shift in SO₄ particle size from the Aitken to the accumulation mode at $\sim 500\,\mathrm{hPa}$ suggests an equivalent increase in SO₄ transport from the extra-Arctic.

7.3.2 Sensitivity to the ice threshold temperature

The ice scavenging temperature threshold is very uncertain and is controlled by many processes that are not simulated in GLOMAP. To test the sensitivity of Arctic aerosol to this threshold, we performed three additional experiments in which the threshold temperature was changed to -10, -5 and $0\,^{\circ}$ C. Changing the ice-phase temperature threshold to

 $-10\,^{\circ}\mathrm{C}$ (ICE-10) and $-5\,^{\circ}\mathrm{C}$ (ICE-5) increased wintertime median BC concentrations north of 70 $^{\circ}$ N by 8 % and 25 %, respectively and SO₄ concentrations by 35 % and 66 %, respectively.

The relatively low sensitivity of winter surface aerosol concentrations to the ice-phase threshold is explained by the sharp meridional temperature gradient (Fig. 4). In late spring when the zonal temperature gradient is weaker, changing the ice-phase temperature threshold to $-10\,^{\circ}\text{C}$ (ICE-10) and $-5\,^{\circ}\text{C}$ (ICE-5) increases SO₄ surface concentrations north of 70° N by 85 % and 250 %, respectively and BC concentrations by 22 % and 70 %, respectively. Comparison of the ICE-10 simulation with surface observations suggests that increasing the ice-phase temperature threshold has decreased model SO₄ bias at Alert (from $-18\,\%$ to $-2\,\%$) and Zeppelin (530 % to 463 %) but increased model SO₄ bias at Barrow (11 % to 37 %). At all other sites model bias has changed negligibly.

The non-uniform springtime response at different sites to increases in the ice-phase temperature threshold might change if mixed-phase scavenging processes were included. The scavenging efficiency of mixed-phase clouds is likely to be spatially variable due to variations in liquid-to-ice cloud ratios between regions (Smith et al., 2009). Thus, better treatment of mixed-phase cloud processes has the potential to improve the simulated spatial and temporal patterns of aerosol concentrations.

It is possible that by suppressing (or overestimating) scavenging rates in mixed-phase clouds during the spring we delay (or accelerate) the simulated transition from the Arctic late-spring haze period to the "clean" summer boundary layer. However, this effect is not evident in the majority of our ground-station comparisons (Fig. 7).

An unrealistic ice-phase threshold of $0\,^{\circ}\text{C}$ does result in a significant change in winter–early spring BC and SO_4 mass, increasing concentrations by 50 and 100%, respectively. However, the greatest effect on surface aerosol concentrations for this simulation occurs in summer where SO_4 and BC concentrations increase by 550 and 120%, respectively. This increase results in a modelled maximum at the majority of Arctic sites that is two months out of phase with observations.

In summary the suppression of soluble aerosol scavenging in ice-clouds increases concentrations of aged sulphate and BC aerosol in the Arctic boundary layer. Including this suppressed scavenging in the model corrects the negative bias of the model versus the observations in the winter and spring. However, in the summer the positive bias seen in CTRL-MODIS has been exaggerated by the ice-cloud scavenging scheme. This is particularly evident at marginal Arctic sites. Sensitivity studies suggest that, in winter surface concentrations of SO₄ and particularly BC are relatively insensitive to increases in the ice-phase temperature threshold of less then 5 °C. In the next section we show how a combination of suppressed wintertime ice-phase scavenging and enhanced

summertime drizzle scavenging results in the correct seasonal cycle at both high latitude and marginal Arctic sites.

7.4 Simulation DRIZZICE

Simulation DRIZZICE includes the low-level cloud drizzle scavenging processes of DRIZZ and the suppressed ice-cloud scavenging of the ICE simulation (now applied to all clouds, including drizzling clouds). Section 7.2 showed that drizzle suppresses Arctic aerosol in all seasons (with a greater effect in summer because of the higher cloud coverage) but exacerbates the model low bias in winter. The ice scavenging process acts in the other direction by suppressing scavenging in winter. We now study the combined effect of DRIZZ and ICE.

A comparison of DRIZZICE and CTRL-MODIS with ground-station SO₄ mass observations shows that the model-observation correlation at Alert, Barrow and Zeppelin has increased from a negative coefficient at all stations in CTRL-MODIS to 0.71, 0.7 and 0.4, respectively in DRIZZICE (Table 3). Thus, the combined effect of drizzle and suppressed scavenging in ice clouds has had the desired effect of decreasing summertime SO₄ and increasing wintertime SO₄ compared to CTRL-MODIS. The model captures the large seasonal cycle at the high latitude Arctic sites (Alert, Barrow and Zeppelin) and the weaker cycle at the marginal Arctic sites Karasjok, Oulanka and Janiskoski (Fig. 7).

For BC, the correlation coefficient has changed sign increasing from -0.7 (CTRL-MODIS) to 0.72 at Alert and from -0.43 to 0.44 at Barrow. The wintertime (DJF) bias in BC has reduced from -94 % to 2 % at Barrow and -92 % to -11 % at Alert, and the model is within 0.1 standard deviations of the DJF observation mean at both sites. However, the annual mean model bias at both stations has increased from 70 % in CTRL-MODIS to 200 % in DRIZZICE at Barrow and from -53 % to 59 % at Alert. This increase in model bias is due to the effect of ice-cloud scavenging in winter and spring (where the model negative bias has been reduced) combined with the inefficiency of drizzle scavenging on fresh Aitken mode BC which typically dominates the modelled summertime aerosol size distribution.

The dominance of insoluble Aitken mode BC in the late summer and early autumn over the Canadian and Siberian Arctic (70–90% of the total BC mass) is the result of the ageing scheme used in GLOMAP. Aerosol in GLOMAP is aged by condensation of SO₂. BC (and OC) particles become hydrophilic when they have a sufficient coating of SO₂, assumed here to be 10 monolayers of H₂SO₄. This is significantly higher than the previous threshold of 1 monolayer, and was altered to improve the model agreement with remote observations of BC (Mann et al., 2010). A similar decision was made by Liu et al. (2011) in their global model. However, in Liu et al. (2011) the required thickness of H₂SO₄ was explicitly calculated from the Köhler equations.

The effect of assuming that BC rapidly becomes soluble can be estimated from the dependence of SO_4 on drizzle rates. In August, high latitude SO_4 concentrations fall by a factor of 600 between runs CTRL-MODIS and DRIZZ and by a factor of 200 between runs CTRL-MODIS and DRIZZx0.5, while BC falls by a factor of 1.6 and 1.3 respectively (Fig. 8). Thus, assuming highly soluble BC would reduce BC by a factor of \sim 200, simulated August BC concentrations would be in better agreement with observations at Alert and Barrow (Fig. 7). However, it is not known whether the overprediction of August high latitude BC is the result of inaccurate emissions or incorrect scavenging.

In conclusion, the use of both drizzle and ice-cloud scavenging schemes in GLOMAP-mode has significantly improved the Arctic model evaluation compared with both CTRL-ISCCP and CTRL-MODIS (Fig. 7 and Table 3). This analysis is clear evidence for two processes affecting surfacelevel aerosol: scavenging in the free troposphere before the aerosol is entrained into the boundary layer (which is controlled here by the change in ice scavenging), and scavenging within the boundary layer itself caused by low-level warm clouds. In the winter, Arctic aerosol is controlled mostly by the free tropospheric scavenging, which is reduced when ice forms, and in the summer is controlled less by scavenging in the free troposphere but by scavenging in low-level drizzling clouds which filter air transported north of 60°. These adjustments to the model have also affected aerosol concentrations outside the Arctic, which we briefly review below.

7.5 Global impact of scavenging processes

Previous comparisons of GLOMAP-mode with numerous aerosol datasets outside of the Arctic (Mann et al., 2010) show a good agreement between the control model and observations. We now show that inclusion of drizzle and ice-cloud scavenging effects leads to small improvements in the model elsewhere.

The DRIZZICE, ICE and DRIZZ simulations were compared with annual mean SO₄ surface concentrations from the EMEP (26 stations), IMPROVE (12 stations), GAW (1 station) and the University of Miami ground-station networks (28 stations) and annual mean EBC surface concentrations from the IMPROVE network (12 stations) (Stier et al., 2005). The IMPROVE and EMEP networks are situated in N. America and Europe, respectively while stations in the University of Miami network range from urban (Miami at 80.2° W and 25.8° N) to isolated sites (Palmer in Antarctic at 64° W and 64.8° S) and include numerous island and coastal stations situated in or near modelled stratocumulus zones (Cook island, Reunion Island).

Comparison of the CTRL-MODIS simulation with all SO₄ datasets concurs with the findings of Mann et al. (2010) that GLOMAP-mode predicts surface concentrations of SO₄ well at all stations, with an average (over all networks) correlation coefficient of 0.88 and positive mean model bias of

74 %. Likewise, in both DRIZZ and DRIZZICE simulations the model-observation correlation is \sim 0.8, but the bias decreases to 32 % in DRIZZ and 65 % in DRIZZICE. The ICE simulation shows no significant change from the control.

Comparison with the few EBC surface mass concentrations in the IMPROVE network (12 sites) shows the model-observation correlation in DRIZZICE has improved negligibly (from 0.32 to 0.35) and model bias has improved from $-37\,\%$ (CTRL-MODIS) to $-21\,\%$ (DRIZZICE). Similar analysis of ICE and DRIZZ with observations suggest that the decrease in BC model bias in DRIZZICE stems from the effect of ice-cloud scavenging rather than drizzle processes.

Global BC mass concentrations have increased in DRIZZICE particularly in the remote marine boundary layer (not shown here), despite the strong BC scavenging rates associated with the stratocumulus zones. In remote regions the increase in boundary layer BC concentrations in the DRIZZICE simulations stems from a number of contributing factors. Firstly, there is a significant decrease in BC particle size (from accumulation to Aitken mode) in the boundary layer in BC source regions such as Eastern Asia due to decreases in cloud processing of smaller to larger particles) which results in a decrease in BC scavenging in the lower troposphere. Secondly, the decrease in boundary layer SO₄ in DRIZZICE results in a increase in the model BC ageing lifetime, which mitigates the increase in BC scavenging in DRIZZ simulations.

Although the percentage increase in BC concentrations in the southern hemisphere is large ($\sim 100\,\%$) the absolute increase is small due to the low concentrations of BC in this region (BC concentrations remain less than 0.01 µg m⁻³). However, in the Northern Hemisphere the increase in BC mass from CTRL-MODIS to DRIZZICE is significant, with BC concentrations in interior Russia rising from $\sim 0.01\,\mu g\,m^{-3}$ to $0.1\mu g\,m^{-3}$ in DRIZZICE and North Atlantic BC mass concentrations rising from $< 0.02\mu g\,m^{-3}$ in CTRL-MODIS to $0.05-0.07\mu g\,m^{-3}$ in DRIZZICE.

BC mass concentration observations from a two-leg cruise in October 1992 from Nova Scotia to Morocco are available for comparison with modelled North Atlantic surface BC concentrations. The JGOFS 92-037 cruise measured a mean BC concentration of $0.024\,\mu g\,m^{-3}$ over the central North Atlantic (25–50° W) (Van Dingenen et al., 1995). The CTRL-MODIS simulation predicts $0.015\,\mu g\,m^{-3}$ in this region and $0.033\,\mu g\,m^{-3}$ in the DRIZZICE simulation (correlation of 0.82 and bias of 55 % versus the observations). However, conclusions about the validity of this global increase in BC mass concentrations remain tentative without more diverse observations.

In conclusion the simulation DRIZZICE compares as well (or better) with available SO₄ and BC mass concentration observations outside of the Arctic.

8 Conclusions

We have used a global model of aerosol processes to investigate how scavenging processes control the seasonal variation of SO₄ and BC aerosol in the Arctic. The effect of these scavenging processes was tested by comparing five model sensitivity simulations (Sect. 6) with surface observations of aerosol mass concentration at three independent Arctic ground-stations: Alert (82.5° N), Zeppelin Mountain on Svalbard (78.9° N) and Barrow (71° N), and averaged observations from the closely situated Scandinavian/Russian stations Karasjok, Oulanka and Janiskoski which lie between 66–69° N.

Inclusion of a new drizzle scavenging process and suppression of soluble aerosol scavenging in all clouds below $-15\,^{\circ}\mathrm{C}$ leads to a significant improvement in the agreement between the model and observations. In particular, the inverse seasonal cycle of aerosol concentration in the control run (with peak aerosol in the spring-summer) is corrected through a combination of these scavenging processes. The suppression of scavenging in ice clouds below $-15\,^{\circ}\mathrm{C}$ has the most pronounced effect, leading to a large increase in wintertime aerosol and reversing the modeled seasonal cycle.

The ice effect is most pronounced at the high latitude sites of Alert, Barrow and Zeppelin, leading to an average factor 6 increase in DJF SO₄, with the effect extending from autumn to spring. The effect is much weaker (factor 2) at the Scandinavian/Russian sites and is restricted to the winter months. This difference in the effect of ice clouds is consistent with the location of the sites relative to the $-15\,^{\circ}\text{C}$ isotherm, which lies asymmetrically over the pole (being shifted towards N. America and Asia, Fig. 4). The analysis suggests that the magnitude of the aerosol seasonal cycle is determined partly by the location of the site relative to the zone of lowest temperatures, with Barrow and Alert having the largest amplitude cycle, followed by Zeppelin, and the Finnish sites having a weak seasonal cycle. The seasonal cycle is further amplified by the occurrence of drizzle in lowlevel stratus clouds.

Our global 3-D model analysis is consistent with the observational analysis of Garrett et al. (2010), who identified a seasonal temperature "switch" controlling the distribution of Arctic aerosol mass. Here we have shown that the likelihood of a site exhibiting a characteristic Arctic aerosol seasonal cycle is dependent on its position relative to the winterspring $-15\,^{\circ}\mathrm{C}$ surface isotherm (see Fig. 4). Our approach of simply switching off scavenging below $-15\,^{\circ}\mathrm{C}$ is expedient in a model that does not simulate cloud microphysics. This threshold temperature may vary spatially and depend on the cloud and aerosol types, but the existence of some kind of threshold temperature is borne out by the Garrett et al. (2010) analysis and our model results, which also suggest that the precise temperature may not be very important.

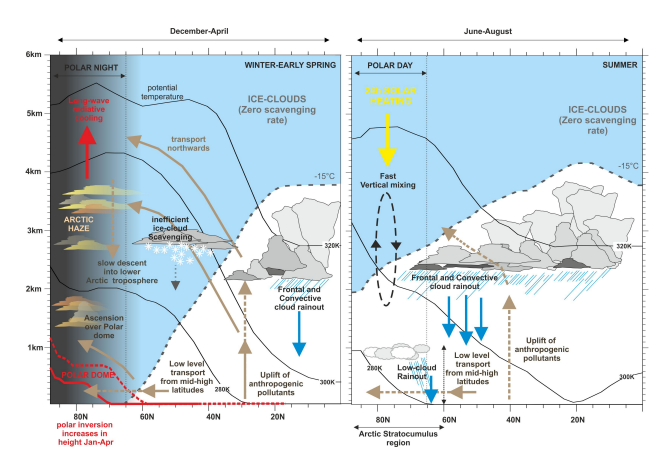


Fig. 11. Schematic showing major transport pathways and removal processes affecting the seasonal cycle in Arctic aerosol concentrations from December–April (left) and June–August (right). Transport pathways are labelled in brown with dashed lines indicating weak transport paths. Wet deposition processes are labelled in blue. Meteorological processes are labelled as required with zonal potential temperature isotherms (used in GLOMAP) shown as black solid lines. Long-wave radiative processes are shown in red while short-wave are in yellow. The $-15\,^{\circ}$ C zonal isotherm (ice-phase cloud temperature threshold) is shown as a dashed line and calculated from atmospheric temperatures diagnosed in GLOMAP. Light blue shading indicates atmospheric region where scavenging is deemed ice-phase and so suppressed.

Our results are also consistent with the coupled-climate model studies of Bourgeois and Bey (2011) and Liu et al. (2011), which found that reducing the scavenging efficiency of ice and mixed-phase clouds decreased model bias compared to observations. In addition, reducing ice and mixedphase cloud scavenging rates in both studies reversed the modeled seasonal trend in aerosol concentrations resulting in a winter to early spring BC and SO₄ maximum similar to that of the DRIZZICE simulations discussed here. Likewise, the CTM study by Huang et al. (2010) noted that the seasonal cycle in BC aerosol was driven by the seasonal cycle in local Arctic scavenging rates. However, here we suggest that the increase in winter-spring BC and SO₄ aerosol also stems from a reduction in scavenging during long-range high-level transport (as well as local suppression), a conclusion also noted by Bourgeois and Bey (2011).

Figure 11 summarises the effects of ice-cloud and drizzle scavenging processes in the context of the major transport pathways to the Arctic identified in the Lagrangian study Stohl (2006). These transport routes into the Arctic are defined by Stohl (2006) as: (1) low-level transport from the mid-latitudes followed by ascent over the polar inversion, (2) low-level transport directly into the Arctic boundary layer, and (3) uplift at lower latitudes followed by adiabatic ascent and transport into the Arctic free troposphere. In the summer, low-level transport through paths [1] and [2] is frequent from European anthropogenic sources and boreal forest fires, while transport from lower latitudes through path [3] is considerably weakened. In winter, transport path [2] occurred almost exclusively through Siberia (where surface temperatures are sufficiently low that air-mass entry into the polar dome is thermodynamically possible) but high-level transport through pathway [3] was found to be frequent from North American, European and Asian sources (Stohl, 2006).

From December to February, high-altitude deposition rates decrease substantially in DRIZZICE simulations due to the suppression of ice-cloud scavenging in the free troposphere (Fig. 10). The increase in winter concentrations of aged SO₄

and BC in the Arctic boundary layer in DRIZZICE suggests that transport (via pathways [1] and [3]) is a significant source of Arctic aerosol in GLOMAP during this period. The absence of low-level drizzling clouds in winter makes drizzle scavenging unimportant as a controlling mechanism. It also means that aerosol transport into the free troposphere through path [3] is not scavenged at low levels after it is mixed down in the Arctic.

In summer, analysis of the latitude dependence of the integrated atmospheric column scavenging flux in the drizzle simulations shows that scavenging increases sharply at the edge of the Arctic stratocumulus zone. The onset of scavenging removes low-level aerosol mass entering the Arctic. Thus inclusion of drizzle actually results in a reduced all-cloud scavenging flux above 70° N because aerosol concentrations are lower (Fig. 9c). From this result we can infer that in summer transport through paths [1] and [2] is both the dominant transport mechanism in GLOMAP and that this pathway is very strongly suppressed by drizzle scavenging. However, the predominance of low-level transport and the generally higher temperatures means that the suppression of scavenging in ice-clouds becomes much less important.

Drizzle scavenging in the Arctic effectively "filters" boundary layer air transported north of 60°. Summertime surface aerosol concentrations between 60° and 75° N are reduced by a factor 30 below the level due to other scavenging processes (dry deposition and rain-out in frontal clouds). But the effect of the drizzle is very dependent on the assumed cloud coverage and drizzle rates, neither of which are well defined in the Arctic. ISCCP cloud cover is approximately a factor 5 lower than that retrieved from MODIS, which leads to a factor 2 less removal of SO₄ but only 5 % less removal of BC by 75° N. The absolute differences in SO₄ and BC concentrations in the high Arctic caused by this uncertainty are only a few nanograms per cubic metre. Further simulations are needed to test whether these differences could be climatically significant in a region with low aerosol.

The possible ramifications of an Arctic drizzle "filter" and cloud scavenging temperature threshold are worthy of further investigation in the context of climate change. In winter, when weak ice-phase scavenging permits significant aerosol transport, Arctic concentrations of SO₄ and BC are relatively insensitive to temperature increases of less than 5 °C due to the sharp meridional temperature gradient. However, the weaker zonal temperature gradient in spring means that higher springtime temperatures are likely to increase scavenging, potentially reducing the lifetime of the Arctic haze. In summer, increased temperatures, reduced sea ice and higher humidity will cause more drizzle and reduced aerosol concentrations and deposition rates in the high Arctic. Thus, aerosol concentrations and deposition rates in the high Arctic are likely to fall as temperatures rise.

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