Changes in aerosol properties during spring-summer period in the Arctic troposphere

A.-C. Engvall\textsuperscript{1}, R. Krejci\textsuperscript{1}, J. Ström\textsuperscript{2}, R. Treffeisen\textsuperscript{3}, R. Scheele\textsuperscript{4}, O. Hermansen\textsuperscript{5}, and J. Paatero\textsuperscript{6}

\textsuperscript{1}Department of Meteorology, Stockholm University, Stockholm, S 10691, Stockholm, Sweden
\textsuperscript{2}Department of Applied Environmental Science – Atmospheric Science Unit, University of Stockholm, Stockholm, S 106 91, Stockholm, Sweden
\textsuperscript{3}Alfred-Wegener-Institut für Polar- und Meeresforschung, Telegrafenberg A43, D-14473 Potsdam, Germany
\textsuperscript{4}Koninklijk Nederlands Meteorologisch Instituut, Postbus201, NL 3730, AE De Bilt, Netherlands
\textsuperscript{5}Norsk institutt for luftforskning, Postboks100, 2027 Kjeller, Norway
\textsuperscript{6}Finnish Meteorological Institute, P.O.B. 503, FI-00101 Helsinki, Finland

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Correspondence to: A.-C. Engvall (anki@misu.su.se)
Abstract

The change in aerosol properties during the transition from the more polluted spring to the clean summer Arctic troposphere was studied. A six year data set of observations from Ny-Ålesund, Svalbard, covering the months April through June serve as the basis for the characterisation of this time period. In addition four-day-back trajectories were used to describe air mass histories. The observed transition in aerosol properties from an accumulation mode dominated to an Aitken mode dominated size distribution is discussed with respect to long-range transport and influences from natural and anthropogenic sources of aerosols and pertinent trace gases. Our study shows that the air-mass transport is an important factor modulating the physical and chemical properties observed. However, the air-mass transport cannot alone explain the yearly repeatable systematic and rather rapid change in aerosol properties, occurring within a limited time window of approximately 10 days. With a simplified phenomenological model, which delivers the nucleation potential for new particle formation, we suggest that the rapid shift in aerosol microphysical properties between the Arctic spring and summer is mainly driven by the incoming solar radiation in concert with transport of precursor gases and changes in condensational sink.

1 Introduction

The Arctic environment has become a focus of the international scientific community during the last several decades due to its vulnerable ecosystem and climate system. Recent studies on the Arctic climate showed that due to its sensitivity to external perturbations, the Arctic could be seen as possible early warning of global climate change. Nowadays climate models predict the largest increase of the annual mean temperature for the Arctic (ACIA report, 2005).

Unlike for many other regions, there are only very few local human-derived sources of air pollution in the Arctic in addition to the metallurgical industry in the Russian Arc-
tic. Therefore, transport of pollution including aerosols and its gaseous precursors from industrialized mid-latitude regions in Europe, Asia, and North America are of great importance. It is suspected that interactions between solar radiation, high surface albedo, the aerosol particles and clouds magnify the radiative impact of atmospheric aerosols (Quinn et al. 2002) in the Arctic region. Thus, for a given aerosol distribution the specific optical impact is most likely increased in this high latitude region.

Previous experiments conducted over the last 40 years have mainly focused on the Arctic Haze phenomenon that occurs during late winter and spring. This was first noted in the literature by Mitchell (1957). Several years later scientist showed the seasonality of this phenomenon and that it was caused by anthropogenic sources at lower latitudes, especially from the Eurasia continent (Rahn, 1981; Barrie, 1986; Heintzenberg, 1989). Further studies show that the chemical composition appears to have a very strong annual variation with higher loadings of anthropogenic components during late winter and early spring compared to the summer months (Heintzenberg and Larssen, 1983; Beine, 1998; Sirois and Barrie, 1999; Quinn et al. 2002).

Apart from the air chemistry, ground-based measurements from the Zeppelin station, Svalbard, and Point Barrow, USA showed that the aerosol loading undergoes a systematic change from spring to summer (Quinn et al., 2001, Ström et. al., 2003). Results from these studies demonstrated that there is an increase of aged particles, i.e. accumulation mode particles larger than 100 nm diameter in winter and spring, which are associated with anthropogenic sources and long range transport. In summer, this mode is significantly smaller in terms of number density. Smaller sized aerosols, so called Aitken mode particles, which range between about 20 and 100 nm, dominate the size distribution. At the same time the total number density increases.

In this study we will put emphasis on the transition from the more anthropogenically influenced spring to clean summer conditions. It will be shown, that the transition from spring-type aerosol to summer-type aerosol occurs almost at the same time from year to year within a few weeks. We have selected different observational data in order to answer the question if this transition is mainly controlled by changes in the transport of
pollutants or if this is an effect of local processes in the Arctic. Transport patterns, trace gases and aerosol microphysics are investigated for the period April through June for the years 2000–2005.

2 Air mass trajectories and long-term measurements from the Zeppelin station

2.1 Air mass trajectories

Using the model TRAJKS (Stohl et al., 2001) three-dimensional four-day back trajectories were provided by the Royal Netherlands Meteorological Institute (KNMI). These were used to study the link between air mass origin and the aerosol properties. The trajectories were calculated on daily basis at 12:00 UTC in a regular grid centred at Ny-Ålesund (79° N, 11.9° E) for period April through June during years 2000–2005. The grid contains beside the point of Ny-Ålesund four surrounding additional points in an almost symmetric grid of more than 100 km sides. Two of the points are located over sea (78.5° N 14.5° E) and (79.5° N 14.5° E) and the other two are located over land (78.5° N 9.5° E) and (79.5° N 9.5° E). The objective to investigate this grid was to demonstrate how representative the trajectories to Ny-Ålesund are for a larger area of interest and to see possible difference between sea and land areas. Results from this study show that the flow pattern is essentially identical for all five chosen points. Thus for further calculations only the trajectories calculated for Ny-Ålesund (79° N, 11.9° E) were used.

In vertical, we investigate if there are any systematic differences in air transport between the boundary layer (BL) and free troposphere (FT) over the investigated period. The Micro Pulse Lidar (MPL) operated by the National Institute for Polar Research (NPIR) often show cloud tops around 2000 m altitude (Shiobara et. al., 2003). We thus define this altitude as the border between the BL and FT. The 1000 m and 5000 m altitudes are then taken to represent airflow in the BL and FT, respectively. Note that the Arctic atmosphere may present several stable layers and surface inversions are often observed (Tjernström 2004). However, we are interested in the layering that contains
“weather” (clouds and precipitation), which is why we use the altitude range indicated by the Lidar.

2.2 Long-term measurements

Aerosol particle number density, aerosol size distribution, SO$_2$, CO and $^{210}$Pb observations at the Zeppelin station

The long-term measurements at the Zeppelin station were used to evaluate temporal variation of the spring-to-summer transition in aerosol properties on a multi-annual basis. The Zeppelin station is located on Mount Zeppelin 474 m above sea level in the community of Ny-Ålesund, Svalbard. Given the elevated location of the Zeppelin station the effect of local particle sources such as from the Ny-Ålesund community itself, sea spray and resuspension of dust are strongly reduced. However, occasionally the sea salt and dust can contribute significantly to the total mass of the particles. Wind fields over Svalbard are complex due to the topography and surface characteristics. Compared to ocean level observation the effects of local wind phenomenon, such as katabatic winds are reduced at the Zeppelin station. Measurements of chemical and physical properties at the Zeppelin station are included in the Cooperative Programme for Monitoring and Evaluation of the Long-range Transmission of Air Pollutants in Europe (EMEP) and the Global Atmosphere Watch (GAW) programme coordinated by the World Meteorological Organization (http://www.wmo.int). Further information about the instrumentation and database are summarized at the homepage http://www.emep.int.

In this study we use aerosol data obtained at the station including number density, size distributions and activity of the radioactive component lead-210 ($^{210}$Pb). In addition we also use sulphur dioxide (SO$_2$) and carbon monoxide (CO) measurements to analyse the history and origin of the air masses. Table 1 gives an overview of data availability for the time period of 2000 through 2005 of all data from Zeppelin station used in this study. In general the data availability is typically better than 90% each year, but some years are not covered by all variables. Year 2002 has low data coverage for aerosol microphysics and only less than 50% of $^{210}$Pb data are available for year 2004.
The physical properties of the aerosols such as the total particle number density and the size distribution are provided by the Department of Applied Environmental Science, Atmospheric science unit (ITM) at Stockholm University. Hourly data from the Zeppelin station includes total aerosol number density for particles larger than 10 nm ($N_{10}$) using a TSI 3010 Condensation Particle Counter (CPC) and size distribution that covers the size range from 20 nm to 630 nm for year 2000 through 2005. The size distribution observations are carried out using a custom build Differential Mobility Particle Sizer (DMPS) based on a Hauke type Differential Mobility Analyzer (DMA) (Knutson and Whitby, 1975) coupled to a CPC model TSI 3010. This system uses a closed loop sheath air circulation described in Jokinen and Mäkelä (1997). The aerosol sample flow is 1 L min$^{-1}$, while the sheath airflow is set to 5.5 L min$^{-1}$. This yields a rather broad transfer function, but improves counting statistics during periods of low aerosol loading. When the total number concentration is $\sim$100 cm$^{-3}$, the uncertainty arising from counting statistics, in the size classes close to mode of the size distribution is less than 5% (one standard deviation). The mobility distribution measured by the DMA is inverted to a number distribution assuming a Fuchs charge distribution (Wiedensohler, 1988). The final data is given as hourly average. As the number concentration range over several orders of magnitude the means are calculated as geometric means.

No dedicated cloud-detecting device is available at the Zeppelin station. However, typically when the station is in cloud, it is represented by low accumulation mode particle (diameter $>0.1 \mu m$) number density as the cloud drops scavenge aerosols from the air. Aerosol characteristics during cloud events are very much affected by the cloud itself and we make a rudimentary data reduction in an attempt to reduce the impact of cloudy episodes. From our experience with the data set we choose an accumulation mode (90–530 nm) number density of 35 cm$^{-3}$ as a threshold to indicate cloud-affected aerosols. Hence, these data were removed from the data set, which corresponds to a data reduction of 22%.

The Norwegian Institute for Air Research (NILU) performs long-term measurements of sulfur dioxide ($SO_2$) and carbon monoxide (CO). Daily $SO_2$ gas concentrations are
measured with KOH-impregnated Whatman 40 filter and further analysed with an ion chromatography. Gas chromatography with mercuric oxide reduction detection is used for CO measurements (Beine, 1998).

The Finnish Meteorological Institute (FMI) provides data of the activity concentration of lead-210 ($^{210}$Pb) on aerosols. $^{210}$Pb concentrations are measured by collection with a hi-volume aerosol particle sampler onto glass fibre filters (Munktell MGA). The sampler is made of stainless steel. The flow rate is ca. 120 m$^3$h$^{-1}$ and is measured with a pressure difference gauge over a throat. Three samples per week were collected with filter changes on Mondays, Wednesdays and Fridays. One out of 25 filters are left unexposed and is used as a field blank sample. The measurement of $^{210}$Pb is carried out by alpha counting of the in-grown polonium-210 (Mattsson et al., 1996; Paatero et al. (2003). We investigate data for five years 2001 to 2005 (see Table 1). Availability of data for each year is 100% except of 2004 when only 43% of data are available.

3 The spring to summer aerosol transition

3.1 Total number density

Due to a large range in the number densities, daily geometric means are calculated based on the hourly arithmetically averaged data. The temporal evolution of $N_{10}$ is presented in Fig. 1. To emphasis major trend in data, a running mean using a weekly window was applied.

The changing pattern from month to month is evident from the frequency distribution of $N_{10}$ number densities presented in Fig. 2. The months of April and May show a mode around 200 to 300 cm$^{-3}$ and only few occasions when aerosol number density exceeds 1000 cm$^{-3}$. June on the other hand shows a frequency distribution that is skewed towards higher aerosol number densities reaching several thousands per cubic centimetre.
3.2 Particle size distribution

To illustrate major changes in aerosol size distribution we present a multi year composite of monthly mean size distributions for each of the three months from April through June in Fig. 3. From Fig. 3 it is clear that a shift from an accumulation-dominated distribution to an Aitken-dominated size distribution occurs over the period from April through June. The integral number density for the different distributions do not change dramatically and range from 200, through 250, to 300 cm$^{-3}$ for the consecutive months. When including smaller particles ($N_{10}$) the change in the integral number density from 203 cm$^{-3}$ in April to 406 cm$^{-3}$ in June (see Table 2) is more pronounced. Hence, simple approximation can be drawn, that around one quarter of the $N_{10}$ aerosol is smaller than 20 nm in June.

Based on the size range of the measurements and data availability we term particles between 22 and 90 nm Aitken mode particles and between 90 and 560 nm Accumulation mode particles. The temporal evolutions of these two modes are presented in Fig. 4 as weekly running geometric mean and standard deviation over the six years period. The Accumulation mode number density shows a general decrease over the time period. However, the trend is not so clear within the given data variability. In contrast, the Aitken mode number density show more structure in the way that an increase from around day of the year (DOY) 150 is evident in both mean and in variability indicating increasing importance of the periods with very high aerosol number densities.

Although primary aerosol sources such as sea spray exist in the Arctic, the Aitken mode particles are dominantly as a result of secondary particle formation. Through condensation and coagulation the newly formed particles grow into the Aitken mode. These processes will affect the growth of the particles and their residence time in a certain size interval, hence coagulation and condensation are major sources of variability for the Aitken mode particles (Williams et al., 2002).

In Fig. 4 the crossover between an accumulation-mode dominated to an Aitken dominated size distribution occurred around DOY 140 to 150, which corresponds to the last
To explore this transition further we generated an alternative figure where we make use of the ratio between the two modes. In Fig. 5a running mean of the ratio $N_{(22-90)}/N_{(90-560)}$ from all available data between 2000–2005 is presented. Low numbers below one representing that accumulation mode particle dominates the size distribution and vice versa for values above one. From Fig. 5 we see that the ratio around DOY 150 quickly exceeds and remains above 1.5. This indicates that once the change takes place the magnitude of the change in the distribution also increases.

The change in shape of the aerosol size distribution can be also viewed in terms of persistence. In Figs. 6a–b the ratio of days for which the Aitken mode number density is larger than the accumulation mode over a weekly window is presented. We term this the Aerosol Transition Index (ATI). The ATI appear to provide a more distinct measure of when the atmosphere has reached summer conditions i.e. dominance in Aitken mode particles. This can be defined at the point when the index reaching and remaining above 0.4 and last for at least 10 days. To study the year-to-year variability we calculated this index for each year (Fig. 6b). We note that all six years have reached summer conditions based on this index by day 145 using the threshold defined above. All years are rather similar with respect to the temporal evolution and to when the final transition takes place and differ only within 5–10 days. Taking into account that a weekly window is used this implies that the transition from spring to summer is completed within typically two weeks from year to year.

4 Coupling between aerosol properties, anthropogenic tracers and air mass transport and aerosol microphysics

4.1 Long-range transport

Eneroth et al. (2003) summarized the seasonal variation of the transport patterns to Ny-Ålesund (79° N, 11.9° E) in a form of air mass trajectory climatology for the pe-
period 1991–2001. Five-day back-trajectories were calculated two times a day (00:00 and 12:00 UTC) and divided into eight major transport patterns (clusters). Their study showed the variability of the atmospheric circulation, in a synoptic scale, month-to-month and year-to-year. They derived clear seasonal differences due to the difference in the strength of pressure gradients between the seasons. Longest trajectories (equal to a fast transport from lower latitudes of the Atlantic and Eurasia continent) are found during winter time and shortest (equal to a slow transport) during summer. Spring, defined as mid April to May, is associated with stable weather conditions and low cyclonic activity. The stable weather conditions favour persistent airflow from the Arctic, and slow transport from northern part of Russia, Scandinavia and the Atlantic. The fast transport associated with storm tracks from the Atlantic is shown to be infrequent during this period. Summer period i.e. June, July and August is associated with weak pressure gradients. During this period the transport from anthropogenic sources is not as efficient as in winter and spring.

In this section we will extend their study and emphasize on airflow patterns over a limited period of the year and furthermore link air-mass transport, anthropogenic tracers and aerosol properties. Our aim is to produce an overview of the temporal evolution of the air transport to Ny-Ålesund.

4.1.1 Vectorized trajectories

For this study we have calculated 546 4-day-back trajectories for two altitude levels, 1000 and 5000 m, from April through June for the years 2000 through 2005 (see details Sect. 2). Handling of this large amount of trajectories requires synthesising the data into a more hand-able data that can be coupled to other tracer data. Therefore a variable called “trajectory vector” was derived. For each trajectory the radius of gyration was calculated. This summarises all the intermediate time steps of the trajectory into one vector, with a direction and magnitude. The direction yields the main flow direction of the trajectory and the magnitude yields and indication on the speed with which the air travelled.
From this trajectory vector we further are able to calculate the north-south (N-S) and east-west (E-W) contribution of the air-mass transport to Ny-Ålesund. These open up the possibility to reduce the air mass trajectory information into single parameter and produce time series, which allow investigation of the air mass transport temporal evolution. Further we use these trajectory vector time series to evaluate the hypothesis that the air-mass transport controls the transition of the aerosol properties between spring and summer.

The coordinates are relative to Ny-Ålesund 79° N 11.9° E i.e. projection of the trajectory vector to the x-axis give us the contribution in the E-W direction relative to longitude 11.9° and projection to the y-axis the contribution in the N-S direction of latitude 79° N. A positive (negative) value of the x- and y-axis represents air from the east (west) and north (south).

4.1.2 Transport patterns

Combining the N-S and E-W components for each day gives us the horizontal information of the trajectory with respect to the direction and magnitude. Fig. 7a–b presents the calculated N-S contribution for each day trajectory arriving to level 1000 m (BL) and 5000 m (FT) for the time period from 2000 through 2005.

Comparing the different years for the fraction of the N-component an about a fifty-fifty contribution of the different wind components. The fraction of the N-component for each year (2000 through 2005) at level 1000 m (5000 m) is 66(65)%, 61(56)%, 49(44)%, 51(48)%, 41(40)%, and 55(43)% (cf. Fig.s 7a-b). To give some value how well the vectors between the two levels follow each other the correlation coefficients for the N-S-contribution have been calculated. For each year the result is $R_{2000}=0.75$, $R_{2001}=0.73$, $R_{2002}=0.62$, $R_{2003}=0.64$, $R_{2004}=0.70$, $R_{2005}=0.63$. The rather low correlation indicates differences in the wind patterns at the two altitudes. From Fig. 7 there are no apparent trends that could link to and explain the transition in aerosol properties.

The analysis above has essentially provided information about the horizontal transport. To investigate the vertical dimension we also calculated the time the trajectory
has spent in the boundary layer during the last 4 days of its transport to Ny-Ålesund. The different years are then averaged for each day of the investigated time period and are expressed as fraction time to the total time-length of the trajectory. In this context we define generally altitudes below 2000 m as BL. This altitude often represents the top of the low level clouds as seen by micro pulse lidar at Ny-Ålesund. The exceptions are cases when the air arrives within a sector that belongs to the Arctic basin i.e. representing the area within the two tangents that origins in the coordinates of Ny-Ålesund and tangents the north coast of Greenland and the northern coast of Siberia. Within this sector the BL height is defined as 1000 m. This difference in definition in BL is taking into account the stable stratification over the pack ice, especially during the summer months (Nilsson et al., 1999).

The calculated fraction ranges from 0 to 1, where 1 represents a trajectory spends all its time in the BL. The temporal evolution of this fraction is depicted in Fig. 8. The upper panel of Fig. 8 represents trajectories arriving to Ny-Ålesund at the altitude level 1000 m and the lower panel trajectories arriving to altitude level of 5000 m. Not surprisingly trajectories arriving at a lower altitude spend more time in the BL. Comparing the start and end of the period, we notice a decreasing trend in ratio of high-level trajectories spending its time in the BL (bottom Fig. 8). But these trends are not very strong and do not show the radical change between DOY 140 and 150 as the aerosol properties do.

4.2 Anthropogenic tracers

In previous section we have shown that transport alone does not explain the transition in aerosol properties observed on annual basis. Here we will explore possibility that the combination of various transport patterns and seasonal changes in the anthropogenic sources strength can shed a light on observed aerosol microphysics. To follow this hypothesis we used data of anthropogenic tracers namely SO$_2$, CO and $^{210}$Pb observed at the Zeppelin station and focused on their temporal variation. SO$_2$ is one of the major anthropogenic trace gases and highly relevant to aerosol properties. However, it also has a natural biogenic source contributing to SO$_2$ in the atmosphere through oxidation.
of DMS (Li and Barrie 1993, Li et al. 1993). Therefore, we also explored measurements of CO and $^{210}$Pb because in the Arctic the occurrence of these can be attributed exclusively to anthropogenic activities and continental regions, and we will use them as major tracers for anthropogenic source regions (Beine, 1998; Paatero, 2003).

Figure 9a and b shows the weekly running mean of SO$_2$ and CO. SO$_2$ shows a large inter-annual variability in data for the period April through June with episodes of high values in April and May and rather low concentration in June. Sulphur dioxide concentrations range between a minimum value of 0.01 µgSm$^{-3}$ (effective detection limit) to a maximum value of 2.13 µgSm$^{-3}$ and a median of 0.07 µgSm$^{-3}$. The monthly mean and standard deviation of SO$_2$ show their maximum to occur in May with 0.14(0.28) µgSm$^{-3}$ and a trend that decreases towards June with corresponding values of 0.07(0.04) µgSm$^{-3}$ (cf. Table 2).

The high variability of SO$_2$ in April and May reflects the frequent fast transport from anthropogenic sources that take place during springtime whereas the June is more associated with lower background concentrations perturbed by occasional higher values but not as high as April and May. For year 2005, SO$_2$ show a mean concentration of 0.13 µgSm$^{-3}$, which differ from other years where the mean concentration is much lower (0.07 µgSm$^{-3}$). Over all years we note that after day 140 SO$_2$ infrequently exceeds 0.07 µgSm$^{-3}$. The only characteristic that could relate to the aerosol transition is the fewer occurrences of higher SO$_2$ levels after DOY 140. However, this trend would tend to decrease the potential for new particle formation.

CO in Fig. 9 follows a clear decreasing trend from April through June. Carbon monoxide has a longer lifetime in the atmosphere than SO$_2$, about 90 days compared to about a week, which explains much of different features presented in Fig. 9. The total mean and standard deviation for all included years from 2002 to 2004 of CO concentration is 135(25) ppbv which is about 9% lower compared to yearly based mean of 146.7 ppbv (Beine, 1998). Beside some small deviation around DOY 155, the trend in mean values shows no special features and rather smoothly decreases with time. Variability is also rather constant over the time period.
The annual cycle of hydroxyl radical (OH) is of major importance for the seasonal evolution of CO as OH acts as a major sink for CO. OH is produced by photochemical reactions and therefore depends on the incoming solar radiation. Observations show that the CO accumulates in the Arctic troposphere during wintertime due to transport processes and the fact that sink processes by OH are not available (Dianov-Klokoc and Yarganov, 1989). During spring when the sun raises the sink processes for CO get active and the CO concentrations start to decrease and its minimum is reached in summer (Dianov-Klokoc and Yarganov 1989). The steady state decrease of CO mixing ratios only slightly perturbed by local maxima correlated with higher SO$_2$ also indicates that dilution and mixing with clean air masses and photochemical sink of CO dominates over the source in a form of long range transport during April–June period.

Lead-210 ($^{210}$Pb) originates almost exclusively from continents and as such it is tracer of air masses having rather recent contact with landmasses. With its long lifetime $^{210}$Pb (half-life 22 years) the atmospheric lifetime is mainly governed by the lifetime of aerosols (Paatero et al. 2003). Most of the atmospheric $^{210}$Pb is attached to accumulation mode aerosols. Based on the activity ratio of $^{210}$Pb and its progeny; mean aerosol residence times can be estimated. Within this study we only look at the temporal variation of $^{210}$Pb data itself. Paatero et al. (2003) presented the annual variation of the activity concentration of $^{210}$Pb at the Zeppelin station between 11 and 620 $\mu$Bqm$^{-3}$ for the year 2001. The maximum was observed in March–April and the minimum in summer. We calculated $^{210}$Pb activity concentration between 0 and 552 $\mu$Bqm$^{-3}$. The data presented in Fig. 10 agree well with Paatero et al. (2003) and his single year study, maximum values in April with a monthly mean and standard deviation of 165 and 134 $\mu$Bqm$^{-3}$, respectively and a much lower mean in June, 35(27) $\mu$Bqm$^{-3}$.

Lead-210 is one of the tracers investigated in this study that shows a more pronounced change around DOY 140. Together with a decrease in the mean activity, the variability also drops notably. It is worthy to note again that $^{210}$Pb is associated with accumulation mode particles and thus could be a proxy of relative increase or decrease of the aerosol surface area, which also are dominated by accumulation mode aerosol.
4.3 Coupling between the aerosols, anthropogenic tracers and air-mass transport

In Sect. 3.2 we have demonstrated that the Aitken mode particles show an increasing trend in number density from April to June and the accumulation mode particles show a slow decreasing trend. With the obtained vectorized trajectories (see Sect. 4.1) we will in this section couple the horizontal air-mass transport to the loading of the aerosols and the anthropogenic tracers loading for each day. Investigations have been performed for all years but we select one year, 2001, to illustrate our findings. For this year we have the most complete data set available (cf. Table 1).

Figure 11a (upper panel) shows the N-S component together with the daily average of the accumulation mode (solid line) and Aitken mode particles (dashed line) and the lower panel the corresponding E-W component. Combining the information of these two plots one can follow the aerosol loading and the main direction of the air-mass transport.

Peak of accumulation mode particles reach values around $300 \text{ cm}^{-3}$, which corresponds to the upper level of the variation of the number density (Fig. 4). The vectorized trajectories for these days, 125 and 132 (May), 165 and 172 (June), are characterised by north-east and south-east contributions.

The high accumulation mode number densities in May are connected to air masses arriving to Ny-Ålesund from northeast. A further look on the trajectories shows the air mass origin from the northerly part of Russia and Siberia and they are most likely affected by pollution sources, e.g. metal smelter at Norilsk (Virkkula et al., 1995). The occasions in June show southeasterly air-transport and the air masses can be traced to Scandinavia and further south. Aiken mode particles show high number densities when the air masses originate in the north-west or south-west i.e. in the clean regions of the Arctic basin and the Atlantic Ocean.

Additional insight into the origin and sources of the high accumulation mode aerosol episodes can be done with help of $\text{SO}_2$ and $^{210}\text{Pb}$ (Fig. 11b). $\text{SO}_2$ show two spe-
cific peaks around day 125 and 130 in May, same as the accumulation mode aerosol (Fig. 11a) supporting the continental origin of the air mass and influence of the anthropogenic sources.

$^{210}$Pb data has a lower data rate (one sample about every third day) compared to particle data and $SO_2$ data (reason for the non-continues line in Fig. 11c). This makes it hard to make accuracy comparisons of $^{210}$Pb and for individual values of accumulation mode densities and $SO_2$. As it is shown in Fig. 10a the averaged concentration of $^{210}$Pb decrease during the period with an order of more than 5 from April through June. Why we also find the highest peak in April, day 95, which can be coupled to a weak northwesterly air-transport to Ny-Ålesund. The concentrations reach a value of $\sim 450 \mu$Bqm$^{-3}$ compared to the median value of 133 $\mu$Bqm$^{-3}$ for the specific period of this year 2001. This high value cannot be associated to any transport from continental origin or to be coupled to any corresponding high values for $SO_2$ or accumulation mode particles. Further investigation of the trajectories shows that the air descending towards Ny-Ålesund originates from the American continent. In an earlier study the effect of the North American continent on the airborne $^{210}$Pb in northern Finland was observed (Paatero and Hatakka, 2000). As mentioned before the residence time for $^{210}$Pb can vary from a few days up to a month depending on how effective the scavenging of the particles from the atmosphere are. Less efficient scavenging of the particles might be the case of this observed max value. However, enhanced values later during the period days 130–135 are within same period for when enhanced values of the density of accumulation mode particles (Fig. 11a) and $SO_2$ (Fig. 11c) and are coupled to northeasterly flow from northern part of Eurasia.

The data presented show that the air mass origin and transport are important factors controlling the aerosol microphysical properties, but against the expectations air mass transport alone cannot explain at all the spring – summer transition in aerosol properties. Moreover, it is very likely that other processes are of the same or higher importance and this alternative scenario will be discussed in the following section.

The variable showing most of a spring to summer transition other than the aerosol is
210Pb. On the other hand the lifetime of 210Pb in the atmosphere is intimately linked to aerosols through being attached to accumulation mode particles (see Fig. 10b). The accumulation mode particles are typically where the largest surface area can be found and thus is the largest sink for condensable species. New particle formation is a non-linear process that involves the competition between the sink of condensable species and the source of the same. Therefore the alternative scenario can be formulated: Can annual systematic changes in the source strength and sink rate of the condensable species together give the type of transition that the aerosol properties show?

4.4 Nucleation potential

4.5 Method

To explore possible link between the observed spring-summer transition in aerosol size distribution with respect to aerosol nucleation potential we first assume that the new particle formation involves sulphuric acid (H$_2$SO$_4$). The equilibrium vapour concentration of sulphuric acid is calculated for each day based on daily average data, for which the source rate and sink rate of H$_2$SO$_4$ is balanced. Following Kulmala et al. (2004) the time dependent vapour concentration of H$_2$SO$_4$, C, can be written as

$$\frac{dC}{dt} = Q - CS \cdot C,$$

where $Q$ is the source rate and $CS$ the condensational sink. Equilibrium vapour concentration here refers to zero rate of change and is equal to $dC/dt = 0$ and thus a balance of the source and sink. The equilibrium vapour concentration of H$_2$SO$_4$, $C$, is then given by

$$C = \frac{Q}{CS}.$$
4.5.1 Source

The source rate $Q$, is calculated by

$$k \cdot \text{OH} \cdot \text{SO}_2,$$

where $k$ is the reaction rate $10^{-12}$ (s$^{-1}$) (Seinfeld and Pandis, 1998), and OH and SO$_2$ are observed daily averages expressed as molecules per cm$^{-3}$. The OH radical is not observed and was parameterized from the global radiation observations $S$ (Wm$^{-2}$) (http://www.awi-bremerhaven.de/MET/NyAlesund/fulltimeresquery.html). To calculate OH concentration, $S$ was simply scaled by a maximum value (387) of the total data set for observed diffuse solar radiation and multiplied by $5 \cdot 10^6$ according to Eq. (4).

$$S \cdot 5 \cdot 10^6 \cdot \frac{387}{\text{387}}.$$  

(4)

As actual OH radical concentrations are unknown to us this scaling is entirely arbitrary and only serves to reproduce typical values for OH concentrations frequently reported in the literature (Seinfeld and Pandis, 1998).

4.5.2 Sink

The condensational sink (CS) is calculated based on observed size distributions following Kulmala (2001), including the correction factor for the transition between the molecular and continuum regimes given by:

$$CS = 2\pi D \int_0^\infty d_p \beta_M(d_p)n(d_p) d_p d_p = 2\pi D \sum_i \beta_M d_p,i N_i$$

(5)

where $D$ is the diffusion coefficient, $n(d_p)$ is the particle size distribution function and $N_i$ is the concentration of particles in the size section $i$. For the transitional correction
factor for the mass flux $\beta_M$ we use the Fuchs-Sutugin expression (Fuchs and Sutugin, 1971).

To derive the condensational sink, CS ($s^{-1}$), the hourly averaged size distributions for particles larger than 90 nm (accumulation mode particles) were used. The reason for restricting the condensational sink only to the accumulation mode particles is that Aitken particles are a result of relatively recent particle production, whereas the accumulation mode particles most likely were already present and available as a sink at the time the new particles were formed.

CS was calculated from the hourly average of particle data. We exclude days with less than 10 data points (hourly averages). It should also be noted that CS is calculated based on dry aerosol due to the measurement setup (see Sect. 2.2). Increase in particles’ size due to ambient relative humidity is not considered. Typically, the Arctic summer boundary layer is often very humid. In situations when humidity exceeds 80% the CS may be underestimated by a factor of 2–3, therefore our values of CS are very likely to be on the lower end of the possible range.

4.5.3 Result

Following the above formulas the equilibrium $\text{H}_2\text{SO}_4$ concentration was calculated based on the average incoming radiation, SO$_2$ and condensational sink for each day. In Fig. 12 these values for the data set are plotted as function of day. The data ranges from close to zero to around $1\times10^8$ molecules cm$^{-3}$ with an increasing trend with time. We are interested to find out whether there is any particular value of C above which small particles are more likely to occur. To do this we analysed the observed integral aerosol number densities of various sizes versus C (Fig. 13a–c). There are different amount of data in each scatter plot presented in Figs. 13a–c due to data reduction of time resolution of the individual instruments. Note that data in 13a and 13b are entirely independent, whereas in Fig. 13c the accumulation mode part of the size distribution enters both variables.

All three scatter plots present a systematic trend with more frequent high number
densities as C increases. However, it is difficult to distinguish a very clear critical value for C that would indicate a nucleation threshold. Based on the three scatter plots it appears that a value of C above approximately $3 \cdot 10^6$ molecules cm$^{-3}$ separates C values with generally low number densities and C values associated with the highest number densities.

Note that we work with daily averages given by measured data from Ny-Ålesund. The time that new formed particles need to reach a size of 10 nm (detectable size for the instrument) may take several hours up to a day due to the low concentration of condensable material in the Arctic. Hence, the observed conditions when there are high number densities of aerosols in Ny-Ålesund are not necessarily identical to the conditions existing where the particle formation actually took place. To give the reader a feeling of the spatial scale of our approach we may think of how far an air parcel is transported in one day. Given a mean wind speed of 5 m s$^{-1}$ this distance is equivalent to more than 400 km transport during one day. This means that processes taking place 400 km from Ny-Ålesund affect our daily averages.

4.5.4 Trace gases influence on observed aerosol loading

Whether or not the critical value is below or above $3 \cdot 10^6$ molecules cm$^{-3}$ we can see the potential effect of such threshold on the size distribution over time. We may think of it as a nucleation potential. Moreover, it is not only of interest if the conditions reach the threshold, but also to what extent C may exceed above these values, as seen from Figs. 13a–c. We illustrate this by making weekly moving average of the ratio between the number of data points exceeding a certain threshold divided by the total number of data points for the time window over all six years. If the equilibrium threshold is reached often we expect that formation of particles occur readily during that time. The higher the value of this threshold is set the more vigorous we expect the particle formation to be.

We explore three different values of H$_2$SO$_4$ equilibrium concentration threshold $0.6 \cdot 10^7$, $0.9 \cdot 10^7$ and $1.1 \cdot 10^7$ molecules cm$^{-3}$ (Fig. 14a). From comparing the differ-
ent ratios it is clear that changing the threshold from $0.6 \cdot 10^7$ to $1.1 \cdot 10^7$ molecules cm$^{-3}$ have the largest impact on the evolution of the ratios in the middle of the time period where the reduction in the ratio is twice as large as in both end of the time period. Thus, the difference between the ratios in April and May compared with June are accentuated. This is most evident for the highest threshold value, where the ratio increases from low values to about 0.2 over the first two months and then quickly increases to above 0.4 in about a week’s time.

A small amount of data (about 6%) is clearly affected by rapid transport from anthropogenic sources evident in the very high SO$_2$ concentrations for the time period. These high SO$_2$ concentrations would yield to enhanced values of equilibrium H$_2$SO$_4$. As we only consider the aerosol size distribution up to 530 nm (due to the limitation of the measuring equipment) in calculating CS, it is possible that a significant contribution to CS from larger particles is missed during strong pollution events. This would result in an underestimation of C, whereas nucleation events are typically quenched in polluted air due to large aerosol surface area. To test the influence of these events on our results we repeated the ratio calculation after screening data where SO$_2$ exceed $0.6 \cdot 10^7$ molecules per cm$^{-3}$ (Fig. 14b). This limit was chosen by investigating data for when SO$_2$ concentration reached high values that probably were influenced by anthropogenic sources. Excluding the pollution events makes no large changes between Figs. 14a and 14b. But excluding these outliers makes the transition from low ratios in the beginning of the period to higher ratios in June somewhat clearer. The highest threshold for instance never occurs before DOY 115. Between DOY 115 and DOY 145 the fraction of time the threshold is exceeded increases to 20%. Only 10 days later this fraction is more than doubled and from about DOY 155 the fraction wiggles around 50%. Hence in late May and beginning of June the occurrence of the potential conditions for particle formation increases dramatically.
5 Discussion

Transport patterns and source strengths of anthropogenic tracers change over the year with the net effect that the Arctic atmosphere in late winter and springtime is the most polluted period and the summer is the cleanest period of the year. Over this time period the aerosol change from being an accumulation mode dominated aerosol to become an Aitken mode dominated aerosol. Our observations and analysis show that this transition is not typically a gradual change but occurs over a rather short time period each year.

Intrigued by the temporal persistence of the transition we investigated systematically pertinent tracers and transport patterns over the period April, May and June. Although seasonal changes in the flow pattern exist (Eneroth et al., 2003; Stohl, 2006) our analysis could not reveal any systematic pattern that could help explain the rapid transition observed for the aerosol properties.

The fact that the SO$_2$ values are at the lowest during the period with most active particle production is perhaps unexpected. Shaw (1989) discussed particle formation in clean areas and suggested that one viable path way is the cleaning of pre-existing aerosol area through precipitating clouds. The temporal evolution of the condensational sink, CS, depicted as a weekly running mean in Fig. 15a indeed shows persistently low values from about DOY 150 and onwards. The temporal characteristics of CS in many ways resemble those presented by $^{210}$Pb in Fig. 10a. As a matter of fact, the correlation between these two variables is better than 0.92. As the $^{210}$Pb is associated with accumulation mode particles this close relation can be understood.

At the same time as clouds remove pre-existing aerosol surface area and pave way for new particle nucleation, clouds also scavenge SO$_2$ through liquid oxidation and produce particulate sulphate on already existing particles. This removal of SO$_2$ will in turn reduce the potential for subsequent nucleation of new particles. The fact that the monthly mean SO$_2$ concentration is about half in June of what it is in April and May (c.f Table 2) is consistent with scavenging by the extensive clouds that occupy the Arctic in
We note that whereas $^{210}$Pb continues to decrease throughout our period of investigation, CS appears to recover at the end. As $^{210}$Pb is a tracer for continental air masses, this deviation between the two variables would be consistent with a rebuilding of the aerosol surface area that takes place within the Arctic basin or over ocean further south. Given the typically low wind speeds in the Arctic summer atmosphere (Stohl, 2006; Eneroth et al., 2003; Heintzenberg, 1991), this decrease in $^{210}$Pb and increase in CS is most likely the origin from secondary aerosol formation taking place within the Arctic.

A complicating aspect with respect to the discussion above about scavenging of $\text{SO}_2$ by clouds is that, if one studies the temporal evolution of $\text{SO}_2$ over the last part of the time period it appears as if $\text{SO}_2$ increases (Fig. 9a). It is not obvious how to fit this behaviour with the observation of $^{210}$Pb and CS above. On one hand it may be indicative of an increasing marine source of $\text{SO}_2$ associated with the summer-time biological activity in the ocean through oxidation of dimethyl sulphide (DMS) and subsequent formation to $\text{SO}_2$. On the other hand this increase may be a greater influence from anthropogenic sources during parts of the summer as reported by (Heinzenberg, 1989). In the later case the scavenging of accumulation mode particles must be efficient while letting some fraction of $\text{SO}_2$ survive the cloud passage or passages. Nevertheless, the question about the origin of $\text{SO}_2$ in the summer Arctic, anthropogenic versus natural is an interesting subject and related to aerosol-cloud interactions and climate forcing and needs further investigations.

The third leg in this simple method to determine the nucleation potential is solar radiation, which in turn is a proxy for OH in the atmosphere. Ström et al. (2003) presented data that showed a close relation between the yearly cycle of particle number density and incoming solar radiation at Ny-Ålesund, which supports the importance of photochemical reactions. At about mid-March the Sun returns to the Svalbard region and the summer solstice occurs about three weeks into June. Hence, the spring to summer period presents a dramatic increase in the amount of radiation that reaches Ny-Ålesund.
Therefore, the decrease in the precursor gas \( \text{SO}_2 \) due to changes in transport patterns is more than compensated by the combined effect of an increased radiation flux and a decreased condensational sink.

This is illustrated by using some simple numbers. Sulphur dioxide decreases by about a factor of two between the beginning and end of the time period (c.f. Table 2). The condensational sink decreases over the same time by about a factor of 2 to 3 (cf. Fig. 15). However, the weekly median radiation increases by about a factor of 5 (Fig. 15b). Hence, the temporal evolution of CS and \( \text{SO}_2 \) largely compensate each other and the nucleation potential is mainly driven by the radiation. The net effect is that the nucleation potential is driven by planetary motions and will therefore reach some critical value at about the same time each year.

Particle nucleation is highly non-linear and some critical super-saturation of condensable vapours must be reached before new particles will form. Below this value nucleation does not readily occur. The more this critical value is exceeded the larger the chance is that the newly formed particles will grow to sizes that can be detected by the instruments at the Zeppelin station for instance.

Based on the temporal evolution of the Aitken mode (Fig. 4) and the aerosol transition index (Fig. 6a and b) we believe that this critical value is typically met every year around DOY 145 plus minus a week. Hence we believe that photochemical reactions govern much of the aerosol dynamics in the Arctic.

6 Summary

The main motivation for this paper was to have a closer look on the spring – summer period (April, May and June), with emphasis on the transition in aerosol properties observed in the Arctic troposphere on annual basis. In the study we have used four-day-back trajectories and long-term observations of the aerosols and trace gases from the Zeppelin station at Ny-Ålesund, Svalbard. We have described the differences between periods, spring and summer, the transition between them and we tried to link the
observations to the large-scale circulation and to influences from natural and anthropogenic sources of aerosols and gases.

We have investigated the hypothesis that the air transport to the Arctic is controlling the systematic change in the physical aerosol properties that are observed at the Zeppelin station. To summarise this work, we ended up with four main conclusions:

1. Using air mass back trajectories we have shown that the transport alone cannot explain the repeating transition from spring-type to summer-type in aerosol observed in the Arctic troposphere.

2. Blocking the advection of the polluted air masses from south into the Arctic is important contribution of the transport, but it should be seen more like a necessary prerequisite instead of the main process controlling the spring-summer transition in aerosol properties.

3. With a simplified model, which delivers the nucleation potential for new particle formation in a form of vapour equilibrium concentration of $H_2SO_4$ we suggest that the aerosol microphysical properties are a result of a delicate balance between incoming solar radiation, transport and condensational sink processes.

4. The temporal evolution of the condensational sink and $SO_2$ concentrations indicate that at large degree they compensate each other and the nucleation potential is mainly driven by the solar radiation. The net effect is that the nucleation potential is driven by planetary motions and will therefore reach some critical value about the same time each year. This is consistent with a repeating pattern of the aerosol transition between spring and summer.

The whole picture is further complicated by the fact that remote sensing data (Treffisen et al., 2006) show the whole troposphere to be involved in a similar transition. Remote sensing within the Stratospheric Aerosol and Gas Experiment SAGE II and SAGE III suggest a change between spring and summer for the optical properties of the Arctic.
aerosol in the upper free troposphere above approximately 4 km altitude (Treffeisen et al., 2006). At this time it is not clear how and to what extent these processes in the boundary layer and free troposphere are interlinked. In order to get better insight into this phenomenon, airborne in-situ measurements covering the whole tropospheric column are necessary.

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References


Shiobara, M., Yabuki, M., and Kobayashi, H.: A polar cloud analysis based on Micor-pulse


Table 1. Available trace gas- and aerosol data used in this study.

<table>
<thead>
<tr>
<th>Trace gas</th>
<th>Unit</th>
<th>Method used to measure</th>
<th>Time resolution of the measured data</th>
<th>Source of compound in atmosphere</th>
<th>Residence time in atmosphere</th>
<th>Year (available data) [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sulphur dioxide SO$_2$</td>
<td>µgS m$^{-3}$</td>
<td>Filter samples</td>
<td>1 day</td>
<td>Anthropogenic, fossil fuels</td>
<td>∼ 4 days</td>
<td>2000(100) 2001(100) 2002(100) 2003(100) 2004(96) 2005(99)</td>
</tr>
<tr>
<td>Carbon monoxide CO</td>
<td>Ppb(v)</td>
<td>Gas chromatography</td>
<td>1 day</td>
<td>Biomass burning, CH$_4$ oxidation, oxidation natural HC, anthropogenic</td>
<td>∼ 1.5 month</td>
<td>2002(100) 2003(95) 2004(98)</td>
</tr>
<tr>
<td>Lead-210 $^{210}$Pb</td>
<td>Bq µm$^{-3}$</td>
<td>High volume sampler</td>
<td>Every third day</td>
<td>Earths crust</td>
<td>From a few days up to two months</td>
<td>2001(100) 2002(100) 2003(100) 2004(43) 2005(100)</td>
</tr>
<tr>
<td>Total particle number density sizes &gt; 0.01µm N$_{10}$</td>
<td>cm$^{-3}$</td>
<td>CPC TSI3010</td>
<td>1 h</td>
<td>–</td>
<td>–</td>
<td>2000(96) 2001(91) 2002(12) 2003(88) 2004(99) 2005(76)</td>
</tr>
<tr>
<td>Size distribution sizes 0.02-0.630 µm DMPS</td>
<td>cm$^{-3}$</td>
<td>DMA and CPC TS3760</td>
<td>1 h</td>
<td>–</td>
<td>–</td>
<td>2000(95) 2001(93) 2002(17) 2003(85) 2004(97) 2005(99)</td>
</tr>
</tbody>
</table>
Table 2. Monthly mean and standard deviation (std) for atmospheric trace gas concentration and aerosol number density.

<table>
<thead>
<tr>
<th>Month</th>
<th>SO$_2$ mean(std) [$\mu$g Sm$^{-3}$]</th>
<th>CO mean(std) [ppbv]</th>
<th>$^{210}$Pb mean(std) [$\mu$Bq m$^{-3}$]</th>
<th>CPC geometric mean (std range) [cm$^{-3}$]</th>
</tr>
</thead>
<tbody>
<tr>
<td>April</td>
<td>0.12 (0.16)</td>
<td>159 (15)</td>
<td>165 (134)</td>
<td>203 (99-416)</td>
</tr>
<tr>
<td>May</td>
<td>0.14 (0.28)</td>
<td>137 (14)</td>
<td>119 (72)</td>
<td>287 (122-672)</td>
</tr>
<tr>
<td>June</td>
<td>0.07 (0.04)</td>
<td>109 (16)</td>
<td>35 (27)</td>
<td>406 (126-1386)</td>
</tr>
<tr>
<td>Total</td>
<td>0.11 (0.19)</td>
<td>135 (25)</td>
<td>113 (107)</td>
<td>274 (108-698)</td>
</tr>
</tbody>
</table>
Fig. 1. The total number density, $N_{10}$, for April through June for the years 2000–2005. The bold solid line is the running geometric mean and the thin solid lines represent plus minus one standard deviation.
Fig. 2. The relative frequency of the total number density \(N_{10}\) for the years 2000–2005 for the months (a) April, (b) May and (c) June.
Fig. 3. Monthly geometric mean and standard deviation (thin lines) of the size distributions for the years 2000–2005; Solid=April, dashed=May and dashed dotted=June.
Fig. 4. Weekly moving average of the Aitken mode particles (22–90 nm) (bold solid line) and the accumulation mode particles (90–560 nm) (bold dashed line) for the years 2000–2005.
Fig. 5. Weekly running mean of the ratio of particle concentration between Aitken mode and accumulation mode, $R_{\text{Ait/acc}} = \frac{N_{20-79}}{N_{79-630}}$ for the years 2000–2005.
Fig. 6. (a) (left hand figure) Weekly running mean of the Aerosol Transition Index (ATI), defined by the ratio between the frequencies of ratios $R_{\text{Ait/acc}} > 1$ divided by the frequency of ratios $R_{\text{Ait/acc}} < 1$ for a period of one week (for the years 2000–2005). (b) (right hand figure) Weekly running mean for each year of the Aerosol Transition Index (ATI), defined by the ratio between the frequencies of ratios $R_{\text{Ait/acc}} > 1$ divided by the total data points for a period of one week for the years 2000–2005.
Fig. 7a. The vectorized trajectories arriving to altitude-level 1000 m plotted with time (day of the year) for each year 2000–2005 (a–f). Y-axis shows the normalized North–South (N–S) contribution for the air-mass transport.
Fig. 7b. The vectorized trajectories arriving to altitude-level 5000 m plotted with time (day of the year) for each year 2000–2005 (a–f). Y-axis shows the normalized North–South (N–S) contribution for the air-mass transport.
Fig. 8. The mean relative fraction time each day trajectory (for the years 2000–2005) has spent in the boundary layer (BL), defined as altitudes below 2000 m (1000 m for the Arctic region). The blue line represent trajectory that arrives to the altitude level of 1000 m and red represents trajectories that arrive to the altitude of 5000 m. Linear regression is plotted for each data set, which present the trend of each line.
Fig. 9. Weekly running mean (bold lines) and plus minus one standard deviation (thin lines) of the trace gases SO$_2$ (a) and CO (b) measured at the Zeppelin station between for the years 2000–2005 and 2002–2004, respectively.
**Fig. 10.** (a) (Left plot) The activity concentration of lead-210 on aerosols measured at the Zeppelin station for the years 2001–2005. Weekly moving average and error bars representing one standard deviation. (b) (Right plot) The averaged lead-210 concentration plotted together with the average accumulation mode (90–630 nm) particles surface.
**Fig. 11.** (a) Vectorized trajectory for each day of 2001. Upper figure shows the N-S vs. the particle densities of the Aitken mode particles (solid line) and accumulation mode particles (dashed line). Lower figure shows the corresponding E-W contribution for each day trajectory. (b) Each day vectorized trajectory for 2001. Upper figure shows the N-S vs. SO\textsubscript{2} concentration. Lower figure show the corresponding E-W contribution for each day trajectory. (c) Each day vectorized trajectory for 2001. Upper figure shows the N-S vs. \textsuperscript{210}Pb concentration. Lower figure shows the corresponding E-W contribution for each day trajectory.
Fig. 12. Equilibrium vapour concentrations of H$_2$SO$_4$ April through June 2000–2005. Based on incoming solar radiation data, SO$_2$ and aerosols measurements from Ny-Ålesund, Svalbard.
Fig. 13. Number densities for three different ranges of particles sizes; (a) Total number densities $N_{10}$ based on measurements from CPC, (b) Aitken based on measurements from the DMPS system and (c) smaller sizes 10–20 nm using the $N_{10}$ minus integral total number from the DMPS, plotted as a function of the calculated equilibrium vapour concentration of H$_2$SO$_4$ April through June for the years 2000–2005.
Fig. 14. Fraction for when a certain criteria, $0.6 \times 10^7$ (solid) $0.9 \times 10^7$ (dashed) and $1.1 \times 10^7$ (dotted) of the equilibrium vapour concentrations of H$_2$SO$_4$ occur. (b) Same as a) but excluded for polluted events.
Fig. 15. Running weekly mean and standard deviation of the derived condensations sink (CS) derived from Eq. (5), based on DMPS data from the Zeppelin station for the years 2000–2005. (b) Average and plus minus one standard deviation Ny-Ålesund for the years 2000–2005.