Seasonal cycle and modal structure of particle number size distribution at Dome C, Antarctica

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Abstract. We studied new particle formation and modal behavior of ultrafine aerosol particles on the high East Antarctic plateau at the Concordia station, Dome C (75°06′S, 123°23′E). Aerosol particle number size distributions were measured in the size range 10–600 nm from 14 December 2007 to 7 November 2009. We used an automatic algorithm for fitting up to three modes to the size distribution data. The total particle number concentration was low with the median of 109 cm⁻³. There was a clear seasonal cycle in the total particle number and the volume concentrations. The concentrations were at their highest during the austral summer with the median values of 260 cm⁻³ and 0.086 µm³ cm⁻³, and at their lowest during the austral winter with corresponding values of 15 cm⁻³ and 0.009 µm³ cm⁻³. New particle formation events were determined from the size distribution data. During the measurement period, natural new particle formation was observed on 60 days and for 15 of these days the particle growth rates from 10 to 25 nm in size could be determined. The median particle growth rate during all these events was 2.5 nm h⁻¹ and the median formation rate of 10 nm particles was 0.023 cm⁻³ s⁻¹. Most of the events were similar to those observed at other continental locations, yet also some variability in event types was observed. Exceptional features in Dome C were the winter events that occurred during dark periods, as well as the events for which the growth could be followed during several consecutive days. We called these latter events slowly growing events. This paper is the first one to analyze long-term size distribution data from Dome C, and also the first paper to show that new particle formation events occur in central Antarctica.

1 Introduction

The climatic effects of atmospheric aerosol particles are tied strongly with their concentration, size distribution, chemical composition and dynamical behaviour in the atmosphere (Forster et al., 2007; Quaas et al., 2009; Ghan et al., 2012). A key process in this respect is atmospheric new particle formation, including nucleation from precursor gases and subsequent growth of nucleated clusters to larger sizes (Kulmala et al., 2004; Wang and Penner, 2009; Kazil et al., 2010; Makkonen et al., 2012). The formation rate of new aerosol particles is linked closely with the gaseous sulfuric acid concentration (e.g. Kulmala et al., 2006; Petäjä et al., 2009; Kerminen et al., 2010; Sipilä et al., 2010), which is related to sulfur dioxide originating mostly from anthropogenic sources.

The concentrations of anthropogenic aerosols have increased markedly since pre-industrial times, while at the same time the concentrations of natural aerosols have remained at roughly the same level (Charlson and Wigley, 1994). Antarctica is an ideal place for studying the natural aerosol processes, since it is the continent furthest away from anthropogenic pollution sources. There is practically no vegetation, and the oceans surrounding the continent are the main source of aerosol particles (e.g., Shaw, 1988; O’Dowd...
Aerosol number concentrations, size distributions and chemical composition have been studied at several stations around Antarctica, and long-term records of particle number concentrations have been reported, e.g. from Neumayer (Weller et al., 2011) and the South Pole (e.g., Samson, 1990). Particle number size distributions have been measured mainly during campaigns at coastal stations (e.g., Ito, 1993; Koponen et al., 2003; Virkkula et al., 2007; Asmi et al., 2010; Pant et al., 2011; Belosi et al., 2012) and on the upper plateau at the South Pole (e.g., Park et al., 2004). Hara et al. (2011) presented number size distributions and aerosol volatility measured at the Japanese Antarctic station Syowa, on the coast of Queen Maud Land in 2003–2005. Norwegian researchers recently started long-term size distribution measurements at the Troll station, more to the inland of Queen Maud Land (Hansen et al., 2009), but there are no long-term size distribution measurements from the upper plateau. The measurements presented here are the first step towards filling in this gap: particle number size distributions have been measured at the Concordia station at Dome C on the upper plateau of East Antarctica since December 2007.

New particle formation has been observed at several stations in coastal Antarctica (Ito, 1993; Koponen et al., 2003; Asmi et al., 2010). The motivation of this study was to observe and analyse new particle formation events in the inland Antarctica. In this work we will present seasonal variation of the particle number and volume concentration and modal structure of particle number size distributions, as well as analyses of new particle formation events during the first continuous period from December 2007 until November 2009.

2 Instrumentation and data analysis methods

2.1 Size distribution measurements

We measured particle number size distributions at the Italian–French Concordia station at Dome C (75°06′S, 123°23′E). The station is located on the upper plateau of East Antarctica at 3200 m above sea level and 1100 km away from the nearest coast. The measurement period was from 14 December 2007 to 7 November 2009.

The sampling site is the same as was used by Udisti et al. (2012) and Becagli et al. (2012) for taking filter samples for chemical analyses. This sampling site is located about 1 km southwest of the station main buildings, upwind in the direction of the prevailing wind. All motorized activity is forbidden south and within 300 m north of the sampling site (Udisti et al., 2012). The northeastern direction was declared as the contaminated sector (10°–90°) due to diesel generator and motor vehicle emissions at the station. Consequently, the data were omitted from further analysis when the measured winds were from the contaminated sector. Due to contamination, 6.6 percent of the measured data were removed from the analysis.

Snow mobiles and other traffic were active from early November to February and thus did not create major gaps to the winter measurements. However, there are minor gaps in the measured data due to power failures. Longer gaps in the data exist for the early spring 2008 and for winter 2009.

Particle number size distributions in the size range 10–600 nm were measured with a Differential Mobility Particle Sizer (DMPS) that consisted of a Hauke-type medium-size DMA (Winklmayr et al., 1991) in a closed-loop arrangement and a TSI Model 3010 condensation particle counter (CPC) that detects particles larger than 10 nm. The DMPS setup was similar to that used at Aboa by Virkkula et al. (2007). The time resolution of the raw data was 10 min. The size distribution data in this study are presented in the UTC time but the new particle formation plots are presented in local time (UTC + 8 h).

In an ideal setup, there would be an independent CPC for measuring total particle number concentrations and a DMPS from which the total number concentration can be calculated. The agreement of these two would increase the quality of the data. However, in the measurements discussed here there was no additional CPC available, so we cannot calculate the degree of closure between two independent measurement methods.

2.2 Data processing methods

2.2.1 Mode fitting

Log-normal modes were identified from the size distributions with an automatic algorithm (Hussein et al., 2005). This algorithm parameterizes aerosol particle number size distributions with a multi log-normal distribution function. The multi log-normal distribution function is widely in use to parameterize atmospheric aerosol particle size distributions. The algorithm used did not need a user decision for the initial input parameters, only the maximum number of fitted modes was set to be three, which is typically enough to represent atmospheric aerosol size distributions. The algorithm works by reducing the maximum number of possible modes with an overlapping test between adjacent modes. The quality of the log-normal fit is based on least-squares value between the fitted and measured size distribution. The modes found by the algorithm were numbered according to diameter from the smallest to the largest. The diameter of the fitted modes depended solely on the size distribution data. The term fitted
modes is used when referring to the modes obtained by the algorithm.

Later in this work the terms nucleation mode, Aitken mode and accumulation mode ranges refer to the measured data in pre-selected size ranges: nucleation mode range (< 25 nm), Aitken mode range (25–100 nm) and accumulation mode range (> 100 nm) (Dal Maso et al., 2005).

2.2.2 New particle formation event classification

New particle formation event days – simply event days below – were determined following the procedure introduced by Dal Maso et al. (2005). We counted as an event day those days when growth of the newly formed particles could be reliably followed as well the days when growth was clearly detected but could not be followed due to for example changes in the air masses. We divided the event days into class 1 events and into class 2 events. From a class 1 event, we could determine the growth rate in contrast to class 2 events. Furthermore we divided the class 1 events into normal events and into class 2 events. From a class 1 event, we could determine the growth rate in contrast to class 2 events. Furthermore we divided the class 1 events into normal events that remind events typically observed at continental sites (for example in Hyytiälä, Finland, e.g. Dal Maso et al., 2005) and into slowly growing events, when the growth could be followed for several days.

Event days were carefully checked to verify that the observed events were natural events and not due to contamination from the station. For this reason, the wind direction and speed were tracked two days before the event started. If the wind direction was from the polluted sector or the wind speed was lower than 2 m s\(^{-1}\) for more than one hour during the 48 h period after the event started, the event was excluded from the analysis. Altogether 21 new particle formation events were excluded. Examples of new particle formation events and the wind direction and speed during the events are shown in figures introduced in Sect. 3.

The lower limit of the instrument, 10 nm of the particle diameter, created a challenge of interpreting both the event starting time and the event duration. In addition, it was not certain whether new particle formation actually initiated on-site or whether we detected solely an appearance of a growing mode originating from particle formation that had occurred away from our station.

2.2.3 Growth rate calculations

Determining growth rates was not straightforward due to the unique nature of Antarctic events and we used several methods to determine growth rate depending on the type of event. For most of the normal events the method developed by Hirsikko et al. (2005) was used for determining growth rates. This method determines the particle growth by following the size class maximum. First, the times of the concentration maxima in each size class are defined. Then a line is fitted to the size class maximum times as function of size class diameter, and the slope of this line gives the growth rate. The method is limited for events in which the growth can be followed to larger sizes.

In case the of the normal events for which the method by Hirsikko et al. (2005) did not work, and in the case of slowly growing events, the growth rates were determined by mode-fitting method or by fitting a curve to the calculated geometric mean diameter. The mode-fitting method is based on the log-normal modes fitted to each number size distribution using the algorithm of Hussein et al. (2005). The method works by selecting the geometric mean diameter of the growing nucleation mode based on visual inspection of the daily contour plots of the particle size distributions. The growth rate is then obtained by a linear least-squares fit to these selected nucleation mode mean diameters as function of time. Further details of this method can be found in Dal Maso et al. (2005), Yli-Juuti et al. (2011) and Kulmala et al. (2012).

The method for calculating the growth rate from the geometric mean diameter was similar to the method described above but used calculated geometric mean from measured data instead of mode data. In some cases two different methods of determining growth rate could be used for the same day and we had to choose the method that was qualitatively the best. The methods used for calculating the growth rate on each event day are given in Table 3. The errors of the growth rates were determined as the standard error of the linear fit.

\[ J_{10} = \frac{dN}{dt} + \text{CoagS} \cdot N + \frac{\text{GR}}{\Delta d_p} \cdot N, \]

where \( N \) is the particle number concentration of 10–25 nm particles, \( \text{CoagS} \) is the coagulation sink due to pre-existing aerosol particles and \( \text{GR} \) is the particle growth rate over the size range of width \( \Delta d_p \). Coagulation sink for nucleation mode particles is calculated from the measured size distributions according to the method presented in Kulmala et al. (2012).

The condensational growth rate explained by certain vapor concentration \( C_v \) can be calculated using the formula (Nieminen et al., 2010)

\[ \text{GR} = \frac{2 \cdot K \cdot \beta}{3 \cdot \rho_v} \cdot \left( \frac{8kT}{\pi} \right) \cdot \left( 1 + \frac{d_v}{d_p} \right)^2 \cdot \left( \frac{1}{m_p} + \frac{1}{m_v} \right)^{1/2} \cdot m_v \cdot C_v, \]

where \( m_v, d_v \) and \( \rho_v \) are the vapor molecule mass, diameter and condensed phase density, \( d_p \) is the diameter of the grow-
ing particle and \( T \) the ambient temperature. \( Kn \) and \( \beta \) are the Knudsen number and the Fuchs–Sutugin transition regime correction factor for mass flux, respectively. Equation (2) can be used to calculate the vapor concentration required to explain the observed particle growth rates. Assuming molecular properties of sulfuric acid for the condensing vapor, the concentration of \( 10^7 \) cm\(^{-3} \) corresponds to the growth rate of 0.4 nm h\(^{-1} \) for nucleation mode (10–25 nm) particles.

The source rate \( Q_v \) for the condensing vapor can be calculated from the equation describing the evolution of vapor concentration:

\[
\frac{dC_v}{dt} = Q_v - CS \cdot C_v.
\]

(3)

In steady-state (\( dC_v/dt = 0 \)) the vapor source rate is

\[
Q_v = CS \cdot C_v.
\]

(4)

Here the condensation sink CS onto aerosol particles is calculated from the measured particle size distributions according to Kulmala et al. (2012).

2.2.5 Back-trajectory analysis

Air parcel 96 h back trajectories were calculated using the Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) Model (Draxler and Rolph, NOAA Air Res. Lab., Silver Spring, Maryland, http://www.arl.noaa.gov/ready/hysplit4.html). A GDAS meteorological dataset was used in the calculations. Back trajectories were calculated for arriving at three different heights: 100 m, 500 m and 1000 m above the ground level.

3 Results and discussion

3.1 General features and seasonal cycle

Over 81 000 size spectra were measured during the measurement period, from which over 76 000 size spectra were used in the data analysis. We calculated the statistics separately for the summer (December–February), autumn (March–May), winter (June–August) and spring (September–November) seasons. In summer, sunlight was present all the time, whereas the winter months were completely dark. The average temperature in winter was \(-63^\circ C\) and in summer it was \(-36^\circ C\).

A clear seasonal cycle of the particle number concentrations was seen in the number size distribution data (Figs. 1 and 2). The particle number concentrations were at their lowest in July and August and at the highest in January. In Fig. 1 the short peaks in the particle number size distribution are mainly contamination from the station (in non-cleaned dataset). However, we observed also natural peaks in the particle number concentration, which indicated new particle formation. Altogether during the measurement period, we observed 60 natural new particle formation days, from which we could analyze particle formation and growth rates on 15 days as explained in detail in Sect. 3.2.

After cleaning the size distribution data, the total concentration of measured particles larger than 10 nm varied from 4 to 1300 cm\(^{-3} \) and the median total concentration was 94 cm\(^{-3} \). The median particle concentrations were 20, 41 and 6 cm\(^{-3} \) in the nucleation, Aitken and accumulation modes, respectively. The geometric mean diameter of the measured particles varied from (5th and 95th percentile) 20 to 64 nm and the volume concentration varied from 0.004 to 0.140 \( \mu \)m\(^3 \) cm\(^{-3} \) (median value 0.033 \( \mu \)m\(^3 \) cm\(^{-3} \)). The particle number concentrations observed at Dome C are lower than those in coastal Antarctica. For instance at Neumayer, the median number concentration of 25 yr of CPC data was 258 cm\(^{-3} \) (Weller et al., 2011), which is 64 % higher than our measured median total concentration. Other coastal measurements in the summer have shown concentrations in the range of 300–1000 cm\(^{-3} \) (Ito, 1993; Gras, 1993). At the South Pole, reported aerosol number concentrations were about 100–300 cm\(^{-3} \) in summer and below 20 cm\(^{-3} \) in winter (Shaw, 1988; Park et al., 2004). These values are close to the mean and average values measured at Dome C (Table 1).

The seasonal cycle in the particle number concentration was similar to those observed at other Antarctic sites: high concentrations in summer and low in winter. The median number concentration was 260 cm\(^{-3} \) and 15 cm\(^{-3} \) in summer and in winter, respectively (Table 1). At Neumayer the annual maximum number concentration of 1000 cm\(^{-3} \) was reported in March and the minimum number concentration of \(< 100 \) cm\(^{-3} \) was reported in June/July.

Weller et al. (2011) also detected a diurnal cycle in particle number concentrations at Neumayer. For particles larger than 7 nm, this cycle was present for the months September through April but absent from May through August. The observations by Weller et al. (2011) do not differ much from our observations: at Dome C the diurnal cycle of particles larger than 10 nm was strongest in spring (September–November), not quite as strong in summer (December–February), and almost absent in both autumn and winter (Fig. 3). Weller et al. (2011) noted that the observed diurnal cycle is consistent with a photo-chemically induced process. The vicinity of the sea with higher aerosol and precursor gas concentrations is probably also affecting the diurnal cycles at Neumayer, whereas at Dome C diurnal cycles are most probably only of photochemical origin.

The same seasonal cycle as for the particle number concentration was also observed for the particle volume concentration (Fig. 2). The mean volume concentration was the highest in summer, \(~ 0.1 \) \( \mu \)m\(^3 \) cm\(^{-3} \) and the lowest in winter, \(~ 0.02 \) \( \mu \)m\(^3 \) cm\(^{-3} \) (Table 1). These numbers can be compared with the mass concentrations obtained from filter and impactor samples taken from Dome C, even though not simultaneously with our measurements. Udisti et al. (2012) analyzed filter and impactor samples from December 2004 to December 2007, which is not the same period as in our
Table 1. Descriptive statistics of the total particle number concentration, geometric mean diameter and its geometric standard deviation, the particle volume concentration, modal mean diameters and their geometric standard deviation, and growth rates during the four seasons.

|        | Winter |        |        |        |        |        |        |
|--------|--------|--------|--------|--------|--------|--------|--------|
|        | Percentile | N  | Mean | Std | 95% | 50% | 5% |
| Total concentration | 18 313 | 20.2 | 37.4 | 40.5 | 15.4 | 4.27 |
| Geometric mean diameter [nm] | 19 005 | 39.4 | 19.3 | 71.1 | 34.2 | 19.6 |
| Geometric std of diameter | 19 005 | 2.27 | 0.30 | 2.71 | 2.25 | 1.92 |
| Volume concentration [µm³ cm⁻³] | 19 005 | 0.021 | 0.091 | 0.046 | 0.009 | 0.002 |
| Diameter of mode 1 [nm] | 16 595 | 29.1 | 30.2 | 85.5 | 18.8 | 9.00 |
| Diameter of mode 2 [nm] | 3740 | 34.7 | 40.9 | 98.7 | 23.3 | 9.00 |
| Diameter of mode 3 [nm] | 231 | 64.6 | 61.6 | 213 | 46.3 | 9.00 |
| Width of mode 1 | 16 595 | 1.62 | 1.33 | 4.08 | 1.19 | 9.00E-07 |
| Width of mode 2 | 3740 | 2.00 | 1.26 | 4.30 | 1.69 | 1.02 |
| Width of mode 3 | 231 | 1.77 | 0.97 | 4.22 | 1.38 | 1.07 |
| GR 10–25 nm [nm h⁻¹] | 0 | – | – | – | – | – |
| GR 25–600 nm [nm h⁻¹] | 1 | 0.500 | – | 0.500 | 0.500 | 0.500 |
| GR from growth limited events | 0 | – | – | – | – | – |

|        | Summer |        |        |        |        |        |        |
|--------|--------|--------|--------|--------|--------|--------|--------|
|        | Percentile | Mean | Std | 95% | 50% | 5% |
| Total concentration | 20 526 | 305 | 179 | 623 | 260 | 150 |
| Geometric mean diameter [nm] | 22 531 | 41.6 | 11.2 | 62.4 | 41.1 | 24.7 |
| Geometric std of diameter | 22 531 | 1.97 | 0.16 | 2.21 | 1.99 | 1.70 |
| Volume concentration [µm³ cm⁻³] | 22 531 | 0.103 | 0.148 | 0.184 | 0.086 | 0.051 |
| Diameter of mode 1 [nm] | 21 643 | 40.2 | 19.6 | 74.9 | 38.5 | 9.00 |
| Diameter of mode 2 [nm] | 11 603 | 61.0 | 38.1 | 124 | 58.7 | 9.00 |
| Diameter of mode 3 [nm] | 6507 | 57.9 | 46.2 | 142 | 41.7 | 9.00 |
| Width of mode 1 | 21 643 | 1.92 | 0.590 | 2.67 | 1.85 | 1.35 |
| Width of mode 2 | 11 603 | 1.63 | 0.460 | 2.43 | 1.52 | 1.25 |
| Width of mode 3 | 6507 | 1.58 | 0.480 | 2.35 | 1.47 | 1.19 |
| GR 10–25 nm [nm h⁻¹] | 9 | 3.37 | 2.34 | 9.01 | 2.79 | 0.792 |
| GR 25–600 nm [nm h⁻¹] | 7 | 0.640 | 0.410 | 1.13 | 0.780 | 0.210 |
| GR from growth limited events | 0 | – | – | – | – | – |

|        | Autumn |        |        |        |        |        |        |
|--------|--------|--------|--------|--------|--------|--------|--------|
|        | Percentile | Mean | Std | 95% | 50% | 5% |
| Total concentration | 16 957 | 131 | 145 | 421 | 87.1 | 24.0 |
| Geometric mean diameter [nm] | 18 467 | 35.4 | 11.3 | 55.5 | 33.7 | 20.5 |
| Geometric std of diameter | 18 467 | 2.03 | 0.140 | 2.67 | 2.03 | 1.80 |
| Volume concentration [µm³ cm⁻³] | 18 467 | 0.043 | 0.239 | 0.098 | 0.030 | 0.007 |
| Diameter of mode 1 [nm] | 17 843 | 32.4 | 21.3 | 76.5 | 29.7 | 9.00 |
| Diameter of mode 2 [nm] | 7791 | 48.5 | 31.9 | 104 | 40.3 | 9.00 |
| Diameter of mode 3 [nm] | 2772 | 46.3 | 37.4 | 110 | 35.0 | 9.00 |
| Width of mode 1 | 17 843 | 1.79 | 0.580 | 2.96 | 1.60 | 1.25 |
| Width of mode 2 | 7791 | 1.65 | 0.420 | 2.50 | 1.54 | 1.20 |
| Width of mode 3 | 2772 | 1.01 | – | 1.01 | 1.01 | 1.01 |
| GR 10–25 nm [nm h⁻¹] | 9 | 3.37 | 2.34 | 9.01 | 2.79 | 0.792 |
| GR 25–600 nm [nm h⁻¹] | 7 | 0.640 | 0.410 | 1.13 | 0.780 | 0.210 |
| GR from growth limited events | 2 | – | – | – | – | – |

|        | Spring |        |        |        |        |        |        |
|--------|--------|--------|--------|--------|--------|--------|--------|
|        | Percentile | Mean | Std | 95% | 50% | 5% |
| Total concentration | 20 351 | 138 | 176 | 457 | 85.8 | 17.9 |
| Geometric mean diameter [nm] | 19 005 | 34.5 | 16.4 | 66.2 | 29.8 | 15.8 |
| Geometric std of diameter | 21 667 | 1.99 | 0.250 | 2.46 | 1.95 | 1.66 |
| Volume concentration [µm³ cm⁻³] | 21 667 | 0.057 | 0.220 | 0.098 | 0.030 | 0.007 |
| Diameter of mode 1 [nm] | 19 713 | 32.4 | 27.4 | 91.2 | 24.0 | 9.00 |
| Diameter of mode 2 [nm] | 11 084 | 38.3 | 28.1 | 93.3 | 30.0 | 9.00 |
| Diameter of mode 3 [nm] | 3649 | 40.4 | 36.4 | 113 | 34.5 | 9.00 |
| Width of mode 1 | 19 713 | 1.76 | 0.830 | 2.94 | 1.56 | 1.16 |
| Width of mode 2 | 11 084 | 1.75 | 0.690 | 2.49 | 1.56 | 1.20 |
| Width of mode 3 | 3649 | 1.75 | 0.650 | 2.93 | 1.56 | 1.20 |
| GR 10–25 nm [nm h⁻¹] | 2 | 13.3 | 1.13 | 14.1 | 13.3 | 12.5 |
| GR 25–600 nm [nm h⁻¹] | 2 | 3.30 | 3.17 | 5.32 | 3.30 | 1.29 |
| GR from growth limited events | 1 | 0.490 | – | 0.490 | 0.490 | 0.490 |
Table 2. Descriptive statistics for data that came from the polluted sector.

| Percentile               | N   | Mean | Std | 95 % | 50 % | 5 % |
|--------------------------|-----|------|-----|------|------|-----|
| Total concentration [cm$^{-3}$] | 5412 | 910  | 4250| 2620 | 245  | 18.2 |
| Geometric mean diameter [nm]     | 5411 | 37.1 | 16.2| 62.2 | 34.7 | 18.4 |
| Geometric std of diameter      | 5411 | 2.05 | 0.260| 2.04 | 1.68 |
| Volume concentration [µm$^3$ cm$^{-3}$] | 5413 | 0.210| 0.634| 0.941| 0.070| 0.008|

Fig. 1. Time series of (a) the particle number size distribution, (b) total particle number concentration, (c) total particle number concentrations in the nucleation, Aitken mode and accumulation mode size ranges, and (d) the total particle volume concentration.

work, yet the general level can be compared. They did not weigh the samples but analyzed them for the concentrations of major ionic constituents. In addition, they did not present the total concentrations but the concentrations of sea salt and its contribution to the sum of analyzed ions for particles with $D_p < 10 \mu$m. The 3 yr average concentration of sea salt was 10.7 ng m$^{-3}$ in summer and 58.8 ng m$^{-3}$ in winter. The respective average contributions were 11.2 % and 84.3 %, so it can be calculated that in the data of Udisti et al. (2012), the average mass concentrations were $\sim 96$ ng m$^{-3}$ in summer and $\sim 70$ ng m$^{-3}$ in winter. The average volume concentrations calculated from our DMPS data were 0.103 µm$^3$ cm$^{-3}$ and 0.021 µm$^3$ cm$^{-3}$ in summer and winter, respectively (Table 1). If it is assumed that the particle density is that of water, 1 g cm$^{-3}$, the mass concentrations were 103 ng m$^{-3}$ and 21 ng m$^{-3}$ in summer and winter, respectively. With the density of ammonium sulfate, 1.8 g cm$^{-3}$, the concentrations would be 185 ng m$^{-3}$ in summer and 38 ng m$^{-3}$ in winter. It has to be noted here that the samplers of Udisti et al. (2012) were at ambient relative humidity, whereas the DMPS sample air got warmer and thus drier when taken into the laboratory air. However, the difference in the relative humidity on the roof and in the DMPS does not change the conclusion. The higher relative humidity in the impactor on the roof just moves the size distribution towards somewhat larger sizes but does significantly not change the total PM10 mass concentration that was compared here with the DMPS-derived mass concentration. The above calculations show that aerosol mass concentrations derived from two very different types of measurements were of similar magnitude. However, a detailed comparison needs to be performed for a period when both number size distribution measurements and chemical sampling are being conducted simultaneously.

We compared particle concentrations in nucleation, Aitken and accumulation size ranges. The particle number concentration in the nucleation and Aitken mode ranges were typ-
Table 3. Particle growth rates (GR) and the standard error of the growth rate in the 10–25 nm size range, the derived formation rate ($J_{10}$) and vapor concentrations (CV) required to explain the observed growth (the vapor is assumed to have properties of sulfuric acid) and condensation sink (CS) values during the particle formation. The methods used to determine growth rate were to fit a curve to the geometric mean diameter (A), to use the method presented by Hirsikko et al. (2005) (B), and to fit a curve to calculated mode data (C).

| Date   | Method | GR (nm h$^{-1}$) | $J_{10}$ (cm$^{-3}$ s$^{-1}$) | CV ($10^7$ molec cm$^{-3}$) | CS ($10^{-4}$ s$^{-1}$) |
|--------|--------|------------------|-------------------------------|----------------------------|--------------------------|
| 9.1.2008 | B      | 4.6 ± 1.3        | 0.023                         | 10.5                       | 2.1                      |
| 10.1.2008 | B     | 2.4 ± 1.0        | 0.020                         | 5.5                        | 3.8                      |
| 26.2.2008 | B     | 3.4 ± 1.9        | 0.022                         | 7.9                        | 1.7                      |
| 3.3.2008  | B     | 10.1 ± 1.5       | 0.084                         | 23.3                       | 1.9                      |
| 9.5.2008  | A      | 1.3 ± 0.4        | 0.0012                        | 3.0                        | 0.3                      |
| 3.10.2008 | A     | 0.5 ± 0.02       | 0.00022                       | 1.1                        | 0.3                      |
| 29.11.2008 | B    | 12.5 ± 1.6       | 0.083                         | 28.8                       | 1.7                      |
| 29.11.2008 | B    | 14.1 ± 2.1       | 0.11                          | 32.5                       | 2.2                      |
| 11.12.2008 | C    | 4.6 ± 0.6        | 0.030                         | 10.6                       | 2.3                      |
| 20.12.2008 | C    | 0.8 ± 0.07       | 0.053                         | 1.8                        | 2.5                      |
| 23.12.2008 | C    | 3.1 ± 0.9        | 0.016                         | 7.1                        | 1.8                      |
| 30.1.2009  | B     | 1.6 ± 0.2        | 0.041                         | 3.6                        | 2.5                      |
| 22.2.2009  | B     | 1.7 ± 0.3        | 0.0043                        | 3.9                        | 1.0                      |
| 24.2.2009  | B     | 2.5 ± 0.2        | 0.017                         | 5.7                        | 1.7                      |
| 11.3.2009  | B     | 1.2 ± 0.3        | 0.071                         | 2.8                        | 1.5                      |
| Average     |        | 4.3              | 0.038                         | 9.9                        | 1.8                      |
| Median      |        | 2.5              | 0.023                         | 5.7                        | 1.8                      |
| Std         |        | 4.4              | 0.034                         | 10.0                       | 0.9                      |

Fig. 2. The annual cycle of (a) total number concentration and (b) volume concentration. The red bars present the median value of volume concentration, the blue box the 25th and 75th percentiles and the black bars the 5th and 95th percentiles.

Fig. 3. The diurnal cycle of total number concentration in (a) summer, (b) winter, (c) autumn and (d) spring. Red bars present the median value of total concentration, blue box the 25th and 75th percentile and black bars the 5th and 95th percentile.

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seasonal cycle with a summer maximum and winter minimum.

To further visualise the differences in the size distributions between the summer and winter, simple descriptive statistics were calculated, i.e., cumulative concentrations in each size channel of the data (Fig. 4). Figure 4 shows clearly that the modes of the size distributions were smaller in winter than in summer. In winter the mode of the median size distribution was at about 20 nm and in summer at about 40 nm. We also found that in summer particle number concentrations were higher in every size class.

3.2 Modal structure of measured particle size spectra

We studied the modal structure of the particle size distributions by using the automatic mode-fitting algorithm discussed above. We compared the modes calculated with the algorithm to measured particle size distribution data (Fig. 5). The cumulative sum of the fitted modes matched the measured size distribution well, both in the case where there are three modes present as well as when only one mode was observed.

We wanted to study the occurrence of the fitted modes and at which sizes the modes are fitted (median diameter of modes). If one mode was present, the median diameter of the fitted mode was 19 nm in the winter and 39 nm in the summer. When two or more modes were present, particles were found in smaller sizes. This indicates that the strongest new particle formation events occurred in summer and, as explained above, the growth did not reach sizes of the accumulation mode. Also in winter when the second and the third mode had the same median values, particles were found in smaller sizes when three modes were observed than when only one or two modes were observed.

Figure 6 shows the relative frequency of the modes calculated by the automatic algorithm and how they fit to the nucleation mode, Aitken mode and the accumulation mode size range. We calculated the relative frequency by summing up the occurrence of the modes in the nucleation, Aitken and accumulation size range in each month and dividing it with the number of modes in each month. When the particle number concentration was low (in winter), most of the time only one mode was present. When the new particle formation took place or particle number concentrations were high, usually three modes were found. The automatic algorithm fitted most of the modes in nucleation mode or Aitken size range. We found that most of the modes were in the nucleation size range from May to August when the total concentrations were the lowest, whereas other months most of the modes were in the Aitken size range. The occurrence of fitted modes in the accumulation size range was the highest during winter months when nucleation was most frequently observed and the particles were also able to grow to bigger sizes. This all indicates that the particles in the high Antarctic east-plateau air are small and that they are growing either slowly or not at all.

3.3 New particle formation and growth

We classified the days with respect to new particle formation into class 1 event days, class 2 event days, apple events (Vana
et al., 2008), undefined days and non-event days (Fig. 7). Before the analysis the contamination events were excluded from the event days (Fig. 8). We observed event days mainly in the Antarctic summer, and the event frequency was peaking in November and February. In 22 February percent of the days were event days. The month with the lowest occurrence of events was July when we observed events in 4 percent of the days. Most of the events belonged to the class 2, which means that the particle growth rate could not be determined. We observed class 1 events during all the summer months and also in March, May, October and November. The highest fraction of class 1 events was observed in November and December, roughly 7% of all the days were event days. The fraction of undefined days varied from 0 to 29 percent, the highest percentage being in November. The fraction of non-event days varied from 46 to 95 percent.

To further analyze the different events, we divided the class 1 events into “normal” events and “slowly growing” events. By normal event we mean similar kinds of events that have been observed practically all over the world in different environments (Kulmala et al., 2004), whereas slowly growing event are characterized by the continuous and slow particle growth of up to several days. Nine of the 13 normal events (for an example see Fig. 9) were observed during the Antarctic summer. The continuous growth of the nucleated particles during several hours suggests that the new particle formation occurs in an area that is from tens of kilometres up to hundreds of kilometres wide, depending on the wind conditions. Since the distance from Dome C to the nearest coastline is several hundred kilometres, this means that the newly formed particles were growing when the measured air masses were already over the plateau of Antarctica.

Apple events took place in most of the months, excluding winter months and we observed them during 0 to 7 percent of the days depending on the month. In apple events nucleation occurs in a more localized area than in the class 1 events, and these locally formed particles arrive at the measurement station after they have grown a certain time, which depends on the air mass properties. This event type has been previously shown to be typical for new particle formation in coastal regions, where this phenomenon is connected with coastal emissions following the low tide (Ehn et al., 2010). Apple events have also been observed at the near-coastal Antarctica site Aboa (Asmi et al., 2010; Kyrö et al., 2013). In our measurements, the classification of apple events was somewhat uncertain because we could not be sure from which sizes the new-particle formation started and how the event was shaped at sizes below 10 nm.
In the size ranges of 10–25 nm and 25–100 nm and for slowly growing events in the size range of 10–25 nm (Fig. 11). We were unable to determine the growth rate in the size range of 25–100 nm for either slowly growing events or winter events, since the growth could not be followed above 25 nm in those cases. The statistics were calculated season-wise.

The growth rate of all the class 1 events varied from 0.5 to 14 nm h$^{-1}$ in the size range 10–25 nm, and the median growth rate was 2.5 nm h$^{-1}$ based on 15 events (Table 3). These values are comparable to those usually observed in continental rural and clean sites (e.g. Kulmala et al., 2004; Manninen et al., 2010; Asmi et al., 2011). Most of the class 1 events were normal events. We were able to determine the growth rate statistics of normal events only for summer when the growth rate varied from 0.8 to 4.6 nm h$^{-1}$ with the median of 2.5 nm h$^{-1}$. In spring we were able to derive only two growth rates and in autumn only one growth rate from the total of 25 events observed during those seasons, but all being of unusually high values in Antarctica. The median growth rate of normal events in the size range of 25 to 100 nm was 0.8 nm h$^{-1}$. For the slowly growing events the median growth rate was 1.0 nm h$^{-1}$, indicating low concentrations of condensable gases. The mean standard error of the derived growth rates was 21%, which means that exact quantification of growth rates is difficult, but nevertheless we are able to prove that faster growth of the early nucleation mode is present.

The formation rates of 10 nm particles, $J_{10}$, of the class 1 events varied from 0.0043 to 0.11 cm$^{-3}$ s$^{-1}$ with the median of 0.023 cm$^{-3}$ s$^{-1}$ (Table 3). The median formation rate of 10 nm particles in Dome C is about an order of magnitude smaller than that measured at a coastal Antarctic station.
Fig. 11. Seasonal growth rate statistics for (a) the growth rates of normal events in the size range of 10–25 nm, (b) the growth rates of normal events in the size range of 25–600 nm, and (c) the growth rates determined from the slowly growing events. The seasons were winter (June, July, August), spring (September, October, November), summer (December, January, February) and autumn (March, April, May). Concerning the normal events in the size class of 10 to 25 nm, the growth rate statistics for spring were determined from two cases, for summer from 10 cases and for autumn from one case. Concerning the normal events in the size class of > 25 nm, the growth rate statistics for winter were determined from one case, for spring from two cases, for summer from seven cases and for autumn from one case. Concerning the slowly growing events, the growth rate for spring was determined from one case and for autumn from two cases.

Aboa or at a boreal forest site in Hyytiälä, Finland (Table 4). The value of the condensation sink varied from 0.00003 to 0.00038 s$^{-1}$ with the median of 0.00018 s$^{-1}$ in Dome C. These values are similar to those observed in Aboa, but about a magnitude lower than those observed in Hyytiälä (Table 4). Figure 12 shows the relation between the average values of $J_{10}$ and CS during the class 1 events. There seems to an overall tendency of more intense new particle formation at higher values of the condensation sink. Since the condensation sink is closely related to the aerosol surface area and submicron aerosol mass concentration, this connection might suggest that the air masses having more particulate matter have also more vapors participating in new particle formation.

It is unknown which vapors condenses onto newly formed aerosol particles and make them grow (e.g. Riipinen et al., 2011). Since sulfuric acid is very likely involved in new particle formation and early growth, we used Eq. (2) here to estimate how high sulfuric acid concentration would be needed for explaining the observed growth. The calculated vapor concentrations in the 15 class 1 events varied from 1.1 to $32 \times 10^7$ cm$^{-3}$ with the median of $5.7 \times 10^7$ cm$^{-3}$. Mauldin et al. (2001, 2004) measured sulfuric acid concentration at the South Pole in the Antarctic summer. The median concentration was $2.7 \times 10^7$ cm$^{-3}$. If sulfuric acid concentrations at Dome C are similar to those in the South Pole, this compound can explain only a small fraction of the particle growth rate in there.

Fig. 12. The formation rate of 10 nm particles as a function of the condensation sink for the class 1 events.

3.4 Origins and atmospheric pathways of particles reaching Dome C during nucleation events

In order to study the origin and possible transport pathways of particles reaching the Dome C station during class 1 and class 2 nucleation events, we calculated 4-day back trajectories that arrived at the station at the starting times of all the class 1 and class 2 events. The trajectories were calculated for three heights: 100 m, 500 m and 1000 m above the ground level. All of the calculated back trajectories for the class 1 event days came from inland, as well as the majority of all the calculated class 2 back trajectories. Most of the inland trajectories travelled close to the South Pole. The trajectories travelled a median distance of 1400 km during last four days before arriving at the measurement station.

It has been shown that a strong inversion is present at Dome C station throughout the year, except in December and January (Busetto et al., 2013; Genthon et al., 2010; Hudson and Brandt, 2005). In order to determine whether the measured air masses came from the inversion layer, we studied the vertical profiles of the back trajectories calculated to arrive at height 100 m (Fig. 13). The majority of the class 1 event back trajectories (Fig. 13a) and class 2 event back trajectories from the sea (Fig. 13c) originated at altitudes higher than 400 m that is above the inversion layer (Busetto et al., 2013). The shapes of the vertical profiles of the back trajectories are explainable with the circulation induced by the Antarctic drainage flow (James, 1989), resulting in that the origin of the particles measured during the class 1 events and some of the class 2 events were from the upper troposphere. In such circulation, air from upper troposphere is transported down to boundary layer in the central parts of Antarctica and then transported towards the coast by katabatic winds.

In contrast, the majority of the class 2 events that originated from land originated at altitudes below 400 m (Fig. 13b), which indicates that the measured particles came
from the inversion layer. Since the air mass travelled a distance of $\sim 1000$ km, we can rule out that the events were caused by contamination from the station that would have stayed in the inversion layer. However it is possible that the some of the class 2 events occurred locally somewhere further, and that we observed at Dome C those parts of the events that were preserved in the inversion layer. A further study of the origin of the particles observed at nucleation event days is not possible in the framework of this study. Based on older studies, we can speculate that the majority of the compounds reaching Dome C originate from the sea. Cosme et al. (2005) reported that oceans contribute more than 90% of the sulphate measured at Vostok station, close to Dome C. They also reported that volcanoes and anthropogenic sources contribute relatively more to the sulphate in inland than at coast. This might also be the case at Dome C due to the Antarctic circulation, supported by Fig. 13a and c.

4 Conclusions

We observed a clear seasonal cycle in the particle number concentrations, similar to that reported for other Antarctic stations. However, this study was the first one that presents the seasonal cycle of the frequency of the main three modes of the aerosol submicron number size distributions: the nucleation, Aitken and accumulation mode. An automatic algorithm was used to calculate modes from the particle size distribution data. The automatically fitted modes were located mainly in the nucleation and Aitken size ranges. Overall, the great majority of the particles were found in sizes below 100 nm of particle diameter, which is probably due to small amount of condensable vapours.

New particle formation events were observed in every month, and this phenomenon was most frequent during the summer months. A new finding was the winter events, which was not expected as there is no sunlight during winter. The apparent growth of particles during the winter events was limited, so we were not able to determine particle growth rates at sizes below 25 nm. Another interesting finding was the presence of slowly growing events, for which a continuous particle growth was evident during several consecutive days.

Analyzing the new particle formation events and calculating growth rates was not straight forward. Our size-distribution measurements started from 10 nm, which means that the beginning of the nucleation was not detected. The median growth rate we determined for the class 1 events in the size range of 10 to 25 nm, as well as the frequency of event days during the summer, was similar to that measured at coastal station Aboa (Asmi et al., 2010). This indicates that there is not a very significant difference in the growth rates between these two sites, which is somewhat surprising, considering that Aboa is close to the ocean that is the source of most condensable compounds in the region, whereas Dome C is high above the ocean and far away from the coast. In future, to better classify new particle formation events and calculate growth rates, it is crucial to be able to measure particle properties at sizes smaller than 10 nm.

It is of interest to contrast the observed seasonal behavior of the aerosol number size distribution in Dome C to that reported for a high-latitude Arctic site, Ny-Ålesund in Svalbard (Tunved et al., 2013). At both sites, there is a relatively stable background particle population from late au-

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Table 4. The comparison of growth rate (GR), formation rate ($J_{10}$) and condensation sink (CS) between Dome C, Aboa and Hyytiälä. The Aboa values represent the medians and ranges during the new particle formation events in January, 2010 (Kyrö et al., 2013). The GR values from Hyytiälä are the median and 10th to 90th percentile range of 809 new particle formation events, the value of CS value is the geometric mean of CS during these events, and $J_{10}$ represents the median 10 nm particle formation rate (Dal Maso et al., 2007).

|                | Dome C   | Aboa     | Hyytiälä |
|----------------|----------|----------|----------|
| **GR (nm h$^{-1}$)** | 2.5      | 5.5 (1.8–8.8) | 2.5 (1.1–5.3) |
| **$J_{10}$ (cm$^{-3}$ s$^{-1}$)** | 0.023 | 0.2 (0.003–0.3) | 0.4 |
| **CS (s$^{-1}$)** | 0.00018 | 0.00040 | 0.0017 |

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Fig. 13. Calculated 4-day back trajectories for class 1 events (a), class 2 events arriving from land (b) and class 2 events arriving from the sea (c). Time is hours to the past from the beginning of the event and height is in a.g.l. The red horizontal lines mark the median height, the edges of the blue boxes the 25th and 75th percentile and the black pillars the 5th and 95th percentile. The number $N$ indicates the number of trajectories used for statistics calculations. All the trajectories in (a), (b) and (c) are calculated to arrive at height 100 m a.g.l.
tumn until spring. In Ny-Ålesund this background is centered between about 100 and 200 nm of particle diameter, consisting mainly of long-range transported primary particles from anthropogenic sources. In Dome C the background is centered below 30 nm, suggesting that these particles have been formed in the atmosphere before entering the continental Antarctica. At both sites, late spring and summer are characterized by active new particle formation with subsequent particle growth up to the sizes of cloud condensation nuclei (> 50–100 nm). In Ny-Ålesund this process starts when the anthropogenic background is still present, eventually replacing the background particles altogether during the summer. In air masses entering Dome C, new particle formation and growth simply strengthen toward the summer, taking place more frequently close to the measurement site, and producing both more numerous and larger particles than in winter.

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