Size resolved morphological properties of the high Arctic summer aerosol during ASCOS-2008

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10 Abstract

11 The representation of aerosol properties and processes in climate models is fraught 12 with large uncertainties. Especially at high northern latitudes a strong under-13 prediction of aerosol concentrations and nucleation events is observed and can only be 14 constrained by in situ observations based on the analysis of individual aerosol 15 particles. To further reduce the uncertainties surrounding aerosol properties and their 16 potential role as cloud condensation nuclei this study provides observational data 17 resolved over size on morphological and chemical properties of aerosol particles 18 collected in the summer high Arctic, north of 80° N.

19 Aerosol particles were imaged with scanning and transmission electron microscopy 20 and further evaluated with digital image analysis. In total, 3909 aerosol particles were 21 imaged and categorized according to morphological similarities into three gross morphological groups: single particles, gel particles and halo particles. Single 22 23 particles were observed between 15 and 800 nm in diameter and represent the 24 dominating type of particles (82%). The majority of particles appeared to be marine 25 gels with a broad Aitken mode peaking at 70 nm and accompanied by a minor 26 fraction of ammonium (bi)sulfate with a maximum at 170 nm in number 27 concentration. Gel particles (11% of all particles) were observed between 45 and 800 28 nm with a maximum at 154 nm in number concentration. Imaging with transmission 29 electron microscopy allowed further morphological discrimination of gel particles in 30 "aggregate" particles, "aggregate with film" particles and "mucus-like" particles.

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Halo particles were observed above 75 nm and appeared to be ammonium (bi)sulfate
(59% of halo particles), gel matter (19%) or decomposed gel matter (22%), which
were internally mixed with sulfuric acid, methane sulfonic acid or ammonium
(bi)sulfate with a maximum at 161 nm in diameter.

5 Elemental dispersive X-ray spectroscopy analysis of individual particles revealed 6 prevalence of the monovalent ions Na^+/K^+ for single particles and "aggregate" 7 particles and of the divalent ions Ca^{2+}/Mg^{2+} for "aggregate with film" particles and 8 "mucus-like" particles. According to these results and other model studies, we 9 propose a relationship between the availability of Na^+/K^+ and Ca^{2+}/Mg^{2+} and the 10 length of the biopolymer molecules participating in the formation of the 3D gel 11 networks.

12 **1** Introduction

13 Aerosol particles have major impacts on the climate of our planet. They alter the 14 planetary albedo both directly by absorbing and scattering sunlight and indirectly by 15 modifying the reflectivity, life-time and extent of clouds (Twomey, 1977; Albrecht, 16 1989; Solomon et al., 2007). Despite the known importance of the effects of aerosol 17 particles on clouds, they still give rise to large uncertainties in climate models 18 (Schimel et al., 1996; Penner et al., 2001; Forster et al., 2007). Detailed model 19 analyses have contributed to an enhanced understanding of the parametric 20 uncertainties in global aerosol models and point towards significant uncertainties 21 arising from an incomplete representation of aerosol processes and emissions in the 22 models (e.g., Lee et al., 2013; Carslaw et al., 2013; Mann et al., 2014). Especially for 23 high northern latitudes, a strong under-prediction of aerosol particle concentrations 24 and nucleation events in summer compared to measurements is recognized (Mann et 25 al., 2014). This can lead to an insufficient representation of cloud condensation nuclei 26 (CCN), which can activate and form cloud droplets, in models. Moreover, the 27 microphysical properties of the cloud droplets are strongly related to the size, 28 chemical composition, morphology and state of mixture of the activated CCN. Size 29 resolved data based on the analysis of individual particles are therefore indispensible 30 for an appropriate parameterization of aerosol particles within aerosol models.

The sources of aerosol particles in the Arctic are subjected to large regional and
seasonal differences. In late winter/spring, a pronounced anthropogenic influence on
the Arctic is observed, that is a phenomenon known as Arctic haze (Shaw, 1995).

1 During that time the Arctic air mass expands southwards towards Eurasia and North 2 America and anthropogenic emissions are transported into the Arctic where they 3 remain for prolonged times (Shaw, 1995; Douglas and Sturm, 2003). The aerosol 4 during periods of Arctic haze is characterised by relatively high concentrations of 5 aged-aerosol predominantly in the accumulation mode (Shaw, 1984; Heintzenberg 6 and Leck, 1994; Ström et al., 2003; Engvall et al., 2008; Korhonen et al., 2008). The 7 air masses arriving in summer, however, originate from sectors over the oceans with 8 limited man-made activities and the transport into the Arctic is slower compared to 9 winter conditions (Stohl, 2006). The summer conditions are thus much more pristine 10 and the aerosol shifts from a dominance of the accumulation mode to the Aitken mode 11 (Heintzenberg et al., 2006; 2015; Engvall et al., 2008).

12 Over the high Arctic pack ice north of 80° number concentrations of CCN show a 13 large temporal variability, ranging over 2 to 3 orders of magnitude but usually are below 100 cm⁻³ and occasionally less than 1 cm⁻³ (Lannefors et al., 1983; Bigg et al., 14 1996; Bigg and Leck, 2001a; Mauritsen et al., 2011; Leck and Svensson, 2015). These 15 16 relatively low CCN concentrations have a significant impact on the formation of low-17 level (stratiform) clouds prevalent in the high Arctic summer. Mauritsen et al. (2011) identified a regime with very low CCN concentrations ($< 10 \text{ cm}^{-3}$) where cloud 18 19 formation is limited mainly by the availability of CCN. Such low CCN concentrations 20 occur as a result of weak local aerosol sources and effective wet deposition (Nilsson 21 and Leck, 2002; Held et al., 2011a,b; Heintzenberg et al., 2006; Leck and Svensson, 22 2015) at the marginal ice zone and over the pack ice.

23 The physical and chemical properties which determine the ability of the summer high 24 Arctic aerosol particles to act as CCN are still not very well understood. Attempts to 25 theoretically predict concentrations of CCN in closure studies resulted in over- and 26 under-predictions of the observed CCN concentrations (Zhou et al., 2001; Bigg and 27 Leck, 2001a; Lohman and Leck, 2005; Martin et al., 2011; Leck and Svensson, 2015). 28 The most recent closure study by Leck and Svensson (2015) simulated the cloud 29 nucleation process by assuming Köhler theory together with a Lagrangian adiabatic 30 air parcel model that solves the kinetic formulation for condensation of water on size-31 resolved aerosol particles. The authors suggested a larger fraction of the 32 internally/externally mixed water-insoluble particles in the smaller aerosol size ranges 33 as well as and kinetically restricted growth of the activated particles. The non-water 34 soluble particle fraction was suggested to physically and chemically behave as

polymer gels¹ with a dichotomous behavior (low hygroscopic growth factor but a high 1 2 CCN activation efficiency) in cloud droplet activation as a result of the interaction of 3 the hydrophilic and hydrophobic entities on the structures of the high Arctic polymer 4 gels (Orellana et al., 2011). On average 32% of the Arctic surface ocean DOM 5 assembled as microgels (Orellana et al., 2011), a significantly higher percentage than reported for other ocean regions (10%; Chin et al., 1998; Verdugo et al., 2004). All 6 7 together these findings strongly supported the previously unverified hypothesis of a 8 link between cloud formation and polymer gels in the surface microlayer (SML, < 9 1000 µm thick at the air-sea interface) of the high Arctic open leads (Bigg et al., 2004; 10 Leck and Bigg, 1999; Leck et al., 2002; Leck and Bigg, 2005b; Bigg and Leck, 2008; 11 Leck and Bigg, 2010).

12 The transport of marine gels into the atmosphere is thought to happen via the burst of 13 air bubbles at the air-sea interface. Air bubbles rising within the water column 14 scavenge surface-active organic matter, especially from the surface microlayer to their 15 outer walls (Wotton and Preston, 2005). Bursting of the bubbles at the water surface 16 produces small jet and film drops containing organic surface-active compounds, 17 debris of phytoplankton, bacteria, viruses and sea salt (Blanchard and Woodcock, 18 1957; Blanchard, 1971, Blanchard and Syzdek, 1988; Gershey, 1983; O'Dowd et al., 19 1999), that are transported further up into the atmosphere through turbulent mixing 20 processes. However, studies of individual particles by Bigg and Leck (2001; 2008); 21 Leck et al., (2002); Leck and Bigg (2005a; 2005b) over the perennial ice have failed 22 to find evidence of sea salt particles of less than 200 nm in diameter. In the Arctic 23 breaking waves as a source of bubbles are rare due to low wind speeds and short 24 fetches between the ice floes (Tjernström et al., 2012). Even in the absence of wind-25 driven breaking waves a recent study has now confirmed both the presence and 26 temporal variability of a population of bubbles within the open leads (Norris et al., 27 2011). The considered mechanisms for bubble formation and mixing were induced by 28 changes in gas saturation. Other possible bubble formation mechanisms are 29 respiration from algae and phytoplankton (Medwin, 1970; Johnson and Wangersky, 30 1987) and the release of trapped air from melting ice (Wettlaufer, 1998).

¹ Phytoplankton and bacteria in surface seawater produce varying amounts of mucus- or gel-like matter comprised of biopolymers like proteins, polysaccharides or lipids that form 3-dimensional networks inter-bridged with divalent ions, preferably Ca^{2+} and Mg^{2+} . Embedded in the polymer network is a high content of water (99%) that prevents the network from collapsing (Chin et al., 1998). This type of supramolecular organisation is referred to as marine gels. (Verdugo, 2012 gives a review).

1 Due to the remoteness and the harsh conditions in the high Arctic, the number of 2 aerosol studies from this region is limited with data available only from four 3 expeditions onboard the Swedish icebreaker *Oden* during the summers of 1991, 1996, 4 2001 and 2008 (Leck et al., 1996; 2001, 2004; Tjernström et al., 2014). These 5 expeditions took advantage of the pristine conditions during the Arctic summer when 6 the Arctic is greatly separated from polluted mid-latitudinal sources, which provided a 7 unique opportunity to study aerosol particles from predominantly natural sources. All 8 other Arctic studies on aerosol chemical composition, morphology and state of 9 mixture were either performed during winter/spring when the transport of polluted 10 aerosol from lower latitudes into the high Arctic was strong (e.g. Hara et al., 2003; 11 Xie et al., 2007; Winiger et al., 2015) and/or they were located further south and 12 outside of the pack ice area. The latter event would allow missing potential aerosol 13 sources from the pack-ice area and at the same time include anthropogenic pollutions 14 (e.g. Geng et al., 2010; Chi et al., 2015).

15 An investigation of size, chemical composition, morphology and state of mixture on 16 the level of individual aerosol particles is required to further reduce the uncertainties 17 surrounding the CCN properties that promote or suppress cloud droplet formation 18 over the pack ice area. The present study will make use of aerosol particles collected 19 during the most recent icebreaker expedition under the name ASCOS (Arctic Summer 20 Cloud and Ocean Study) 2008. The Swedish icebreaker Oden departed from 21 Longyearbyen on Svalbard on 2 August and returned on 9 September 2008. After 22 traversing the pack-ice northward, the icebreaker was moored to an ice floe and drifted passively with it around 87° N between 12 August and 1 September 23 24 (Tjernström et al., 2014). We used electron microscopy (scanning electron 25 microscopy (SEM) and transmission electron microscopy (TEM)) to image aerosol 26 particles at high resolution, and subsequent digital image analysis to objectively 27 assess size and morphology of the particles on an individual basis. Earlier studies 28 north of 80° focused on a qualitative description of the aerosol in the high Arctic 29 (Leck and Bigg, 2005, 2008, 2010; Bigg and Leck, 2001b, 2008) complemented with 30 bulk chemical analyses (Leck et al., 2002; 2013; Leck and Svensson, 2015; Lohman 31 and Leck, 2005).

Firstly, by individually screening close to 4000 aerosol particles collected during the
ice-drift with SEM and subsequent digital mapping, we gained size resolved
information on the aerosol population as a whole. The obtained number size

1 distribution was compared with measurements from an independent method (Tandem 2 Differential Mobility Particle Sizer, TDMPS) to verify that a representative fraction of 3 the aerosol population was captured with SEM. Secondly, we sorted all mapped 4 particles according to morphological differences and a separate number size 5 distribution for each of the morphological types was obtained. Thirdly, to obtain 6 deeper insights into the morphological features of the collected particles and to 7 simultaneously assess their elemental composition with EDX spectroscopy 8 investigated a subpopulation of aerosol particles in TEM at very high resolution.

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1 2 Methods for sampling of airborne aerosol particles during ASCOS

2 2.1 Collection of airborne particles

3 2.1.1 The sampling inlet

4 A PM₁₀-inlet (9 cm inner diameter) was deployed at ambient conditions (85–100 % 5 relative humidity (RH) and temperatures around 0 °C) to eliminate particles with 6 diameters > 10 micrometer in equivalent aerosol dynamic diameter (EAD) from the 7 sampled air. To optimise the distance from the sea-surface and the ship's 8 superstructure, the inlet was located forward 25 m above the sea-surface and 3 m above the roof of the laboratory container on the 4th deck of the icebreaker. Direct 9 10 contamination from the ship was excluded by using a pollution controller, located 11 directly after the inlet pipe that passed through the roof of the container. Provided that 12 the wind was within $\pm 70^{\circ}$ of the direction of the bow and stronger than 2 ms⁻¹, no 13 pollution reached the sample inlets (Leck et al., 1996). Directly downstream from the 14 pollution sensor the electrostatic precipitator and the TDMPS were connected to the 15 inlet with short stainless steel tubes (length ca. 1 m). To ensure that the sampling 16 conditions and losses were the same for both instruments, the inlet take-offs for the 17 two instruments were placed closely together. The temperature in the container was 18 kept at 20 °C which resulted in an RH of 20% in the secondary lines during sampling. 19 See Leck et al. (2001) for more details of the set-up for the sampling of aerosol 20 particles.

Volatile compounds on particle surfaces and weakly bound water molecules were probably lost during the sampling procedure. In the Arctic, the concentration of volatile compounds is generally lower than at lower latitudes (Bates et al., 1987), and losses due to evaporation can be considered very small.

25 2.1.2 The electrostatic precipitator

Using the electrostatic precipitator the aerosol particles were collected directly onto 3 mm copper 300 mesh Formvar-coated TEM grids (TED PELLA Inc.; Dixkens and Fissan, 1999; Leck and Bigg, 2008). Formvar-coated grids were chosen because of the hydrophilic and thus polar nature of the Formvar film (Rocha et al., 2005). The aerosol particles were charged at the inlet of the precipitator by a ⁶³Ni beta-emitting radioactive source and precipitated by a 12 kV cm⁻¹ electric field between the inlet and the collecting grid surface. The flow rate was kept very low (0.17 mL s⁻¹) in order to collect particles up to ~1 μ m in diameter. The collection efficiency of the electrostatic precipitator was intercompared with a TSI 1236 Nanometer Aerosol Sampler (⁶³Ni beta-emitting radioactive source and sample flow of 1 Lm⁻¹) mounted side-by-side with the electrostatic precipitator. Both collected a small but statistically significant number of particles < 25 nm in diameter. The precipitator took samples for 6 to 12 hrs. Before and after sampling the grids were placed within a grid holder box in a sealed plastic bag and stored in a desiccator at a constant temperature of 20 °C.

8 2.1.3 The TDMPS-sampling system

9 The TDMPS-sampling system to measure the number size distributions of dry (20% 10 RH) sub-micrometer particles used pairs of differential mobility analyzers (DMAs). 11 The TSI 3010 counters in the DMAs were size and concentration calibrated against an 12 electrometer and the TSI 3025 counters for particle sizes below 20 nm in diameter 13 according to Stolzenburg (1988). This set up yielded a complete number size 14 distribution from 3 nm to 800 nm diameter scanned over 45 size channels every 10 -15 20 min. Further details of the TDMPS system can be found in Heintzenberg and Leck 16 (2012). NIST (National Institute of Standards Technology) traceable calibration 17 standards of polystyrene latex spherical particles were used to determine error in 18 determination of the mobility diameter to ± 5 % (Wiedensohler et al., 2012).

In order to compare the number size distribution obtained from the precipitator
samples (section 2.1.2) with those simultaneously recorded by the TDMPS, we
assumed median particle number diameters for each of the 45 TDMPS size channels.
The particle diameters were then merged to form a complete set of diameters across
the TDMPS measuring interval.

24 2.2 Image recording and elemental analysis

25 2.2.1 Imaging with scanning electron microscopy (SEM)

The samples were investigated with a high-resolution SEM (JEOL JSM-7401) under high vacuum conditions, at less than 9.63 x 10-5 Pa (Stevens et al., 2009). A detailed description of the setup of the scanning electron microscope can be found in Hamacher-Barth et al. (2013). The Gentle Beam mode of the microscope was used to minimalize radiation damage of the aerosol particles, avoid surface charge-up and to demagnify the electron beam diameter (Michael et al., 2010). Correction for stigmatism and focusing of the electron beam was done every time before imaging an aerosol particle. The grey scale (contrast and brightness) was adjusted automatically
 before recording an image.

The imaging of the aerosol particles aimed to account for an uneven distribution of the particles on the TEM grid and to capture a representative fraction of the aerosol particles. In brief, particles were imaged at a magnification of 40.000 on the TEM grid squares along a diagonal from the center of the grid to the edge on 6 to 8 squares of the TEM grid. Each square was screened systematically to capture a representative fraction of the aerosol population. For a detailed description of the screening procedure see Hamacher-Barth et al. (2013).

10 **2.2.2** Imaging with transmission electron microscopy (TEM)

11 To image the samples with TEM they have to be coated by a thin metal layer. 12 Evaporation of a heavy metal thin coating at an oblique angle onto the sample 13 increases the mass contrast and accentuates the topography of the aerosol particle by 14 producing a shadow (William and Carter, 2006). Furthermore, shading has the 15 advantage that the metal cover protects the aerosol particles against heating by the 16 electron beam during examination, especially at high magnifications. It is also 17 advantageous that in case of any evaporation from the aerosol particle the metal 18 replica of the aerosol particle is still visible.

19 The aerosol particles were shaded with platinum (Pt) at an angle of $\arctan(0.5) = 26^{\circ}$ 20 (Okada, 1983) in a vacuum chamber at 10⁻⁶ mbar. Pt was evaporated from a Pt wire 21 (\emptyset 0.2 mm, 20 mm length). The Pt wire was drawn around a tungsten (W) wire and 22 evaporated clusters of Pt atoms when the W wire was heated up electrically by a 23 85 mA current for 30 sec. The shading procedure produces a layer of Pt particles of 1-24 2 nm in diameter on the TEM grid.

25 After shadowing the TEM grids were examined in TEM using a JEOL JEM-2100 26 high-resolution instrument, equipped with a LaB₆ filament and a Si/Li detector crystal. 27 The TEM grid containing the aerosol particles was mounted on a sample holder made 28 of Beryllium to avoid background signals from the sample holder material in the EDX 29 measurements (see chapter 2.2.3). A CCD camera (Gatan SC1000 Orius, 11 30 Megapixel) in bottom mount position was used to image the aerosol particles. Images were taken at high vacuum less than 35×10^{-5} Pa and at an accelerating voltage of 31 32 100 kV.

Particles were imaged on TEM grid squares along a diagonal from the center of the grid to the edge on 6 to 8 squares. To avoid imaging of particles that were damaged by prior imaging with SEM, an area of the grid was chosen, which was not exposed to any electron beam at high magnifications before. Screening each square for individual particles was done at a magnification of 30.000, and images were taken at magnifications between 25.000 and 80.000.

7 2.2.3 Elemental X-ray spectroscopy

8 The elemental analyses were performed using an energy dispersive X-ray detector 9 JED-2300 attached to the JEM-2100 TEM. In order to avoid time consuming 10 realignment of the electron beam and focusing procedures, the EDX-analyses were 11 also performed at an accelerating voltage of 100 kV. The energy range measured was 0-20 keV, the counting rate was typically 1053 counts/sec⁻¹, the life time was 30 sec, 12 13 the real time was 33.00 sec and the dead time was 10 %. Generally EDX spectroscopy 14 allows the detection of elements \geq Be as their photon energies are above 100 eV and 15 thus lie within the X-ray region of the electromagnetic spectrum (Egerton, 2008). 16 Nevertheless, the detection of light elements like C, N, and O that are typical for 17 organic compounds can be difficult on a Formvar-coated copper grid since the signal 18 intensity can be biased by attenuation of the X-ray signal through absorption by the 19 adjacent copper grid. For this reason these elements were not reliably detectable and 20 are not part of this study.

Blank grids shadowed with platinum were used to identify the background noise and
signals from the TEM grid including copper from the grid and the Formvar film and
the Pt shadowing. The EDX spectra of blank grids showed only signals from Pt, the
supporting copper TEM grid and carbon and oxygen signals from the Formvar
substrate-film.

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27 2.2.4 Digital image analysis

Images taken with SEM at a magnification of 40.000 were evaluated using an optimized commercial image processing software (Aphelion[™] Dev 4.10). In brief, the maximal intensity of the neighboring background of each aerosol particle was determined. Using exactly the same image but including the aerosol particle allowed

the separation of the particle and measurement of the particle area in pixels
 (Hamacher-Barth et al., 2013).

3 The particle size was calculated according to Eq. (1)

$$4 D_{pa} = 2\sqrt{Area/\pi} (1)$$

with D_{pa} as the particle equivalent diameter, which is the diameter of a circle that
comprises the same area as the aerosol particle projected onto a two-dimensional
surface (Allen, 1997; Hinds, 1999). The value for the area is calculated from the
number of pixels counted for each particle. A number size distribution of the aerosol
sample was obtained using MATLAB 2011a and the freely available software
package EasyFit.

11

12 3 Results and discussion

13 Firstly, to verify that a representative fraction of the aerosol population has been 14 captured with SEM, we calculated a number size distribution of all aerosol particles 15 and compare it with measurements from TDMPS. Secondly, we sorted all particles 16 imaged according to morphological similarities into three gross groups, named: single 17 particles (SP), gel-like particles (GP) and halo particles (HP), shown in Fig. 2. Thirdly, 18 to obtain more subtle insights into the morphological features of the collected aerosol 19 particles and simultaneously assess their elemental composition we investigated a 20 subpopulation of the aerosol particles with TEM and EDX spectroscopy at very high 21 resolution.

22 3.1 Total number size distributions

23 In order to derive an overall number size distribution we imaged in total 3909 aerosol 24 particles at a magnification of 40,000 with SEM. The number size distribution of all 25 imaged aerosol particles exhibited an Aitken mode at 32 nm in diameter. In the 26 accumulation mode region maxima are observed at 89 nm and 147 nm with a shoulder 27 at 335 nm (see Fig. 3, red line). Hamacher-Barth et al. (2013) used the same image 28 mapping method as in this study and determined the error of sizing for polystyrene 29 latex spheres of several diameter sizes between 20 and 900 nm. The error values are 30 displayed in Fig. 3 as red arrows. For the TDMPS number size distribution we 31 assumed an error in determining the mobility diameter of 5% across the whole

1 measuring interval (Wiedensohler et al., 2012). The two approaches show an overall 2 good agreement between their number size distributions with a similar modal 3 structure, an Aitken mode below 80 nm and an accumulation mode at higher 4 diameters. The reduced particle number concentration in the Aitken mode seen by 5 SEM was probably caused by their partly weak contrast to the Formvar film, which 6 either resulted in an underestimation of the size or that the particles remained 7 undetected. The accumulation mode was separated into a double peak with particle 8 number maxima at 89 nm and 147 nm in SEM and 106 nm and 158 nm in diameter in 9 TDMPS. The aerosol particles at diameters > 100 nm often showed a patchy and 10 inhomogeneous appearance, that might have lead to an underestimation of their size 11 and the observed shift to smaller diameters in SEM with that of 147 nm compared to 12 173 nm in TDMPS and broadening of the maximum at 335 nm.

In general, the number size distributions determined for particles larger than 20 nm in
diameter showed modal features typical for aerosol particles collected in the high
Arctic summer boundary layer: an Aitken mode between 26 and 80 nm and a
multimodal accumulation mode between 80 and 1000 nm (Covert et al., 1996;
Heintzenberg et al., 2006; Heintzenberg and Leck, 2012) with the so called
Hoppelminimum around 80 nm in between (Hoppel, 1986).

19 3.2 Single particles

20 Single particles (SP) seen by SEM appeared as single entities that mostly contrasted 21 sharply and thus could be easily separated from their Formvar background for 22 diameters > 40 nm. At smaller diameters, however, the contrast to the backgroud was 23 often weak and probably resulted in an underestimation of particle size or non-24 detection of particles. Imaged examples of SP are shown in Fig. 4. Of the 3909 25 particles that were mapped, SP were the overall dominating type of particles with 26 82% of the total aerosol particles attributed to this group (Table 1). They were 27 observed over the whole size range between 15 nm and 800 nm in diameter with a 28 broad Aitken mode peaking at 64 nm accompanied by a less pronounced peak at 27 29 nm. The majority of SP (80%) appeared in the Aitken mode size region and below 80 30 nm in diameter (Table 1). In the accumulation mode size range, 18% of SP appeared 31 between 80 and 200 nm with a maximum at 106 nm in diameter and the remaining 32 2% of the SP were detected in diameter sizes (Table 1) > 200 nm (Fig. 5, upper panel). 33 We observed that 35% of the SP partly evaporated under the SEM electron beam but 34 retained their outer shape on the timescale of minutes. Also, at a higher magnification

1 using TEM, the same behavior was seen for 30% of the particles. We tentatively 2 assigned these particles to be ammonium (bi)sulfate particles. We were guided by the 3 results published by Heard and Wiffen (1969) and Bigg and Leck (2001b) where 4 particles with the same morphological features and instability under the electron 5 microscope were made up by ammonium sulfate, bisulfate or methane sulfonate 6 mixtures. The presence of ammonium sulfate or bisulfate particles would be 7 supported by the fact that ammonia has been reported to be the predominant base in 8 the remote marine troposphere (Söderlund, 1982) that undergoes primarily acid-base 9 reactions with non-seasalt H₂SO₄, an oxidation product of biogenic dimethyl sulfide, 10 DMS (Quinn et al., 1987). Leck and Persson confirmed the presence of ammonia 11 bisulfate particles both along the marginal ice edge and over the inner parts of the 12 pack ice. Over remote marine locations at lower latitudes, Meszaros and Vissy (1974) 13 observed ammonium bisulfate concentrations up to 38% and on average 24% 14 contained the highest particle concentrations between 100 nm and 1 um in diameter. 15 In the literature, chemical tests have also been used to identify ammonium and sulfate 16 in samples investigated by TEM (Bigg and Leck, 2001b). Such tests were not 17 implemented during this study since the use of chemicals would have added 18 additional mass onto the particles. This would have altered the size and the 19 morphology of the particles and hampered the investigation of the aerosol particles 20 with TEM and EDX spectroscopy.

21 The presence of biogenic nitrate as a counter ion to ammonium can be considered 22 rather unlikely since the formation of ammonium nitrate happens only after all sulfate 23 has been neutralized (Kuhn et al., 2010). Nitrate concentrations from impactor 24 measurements during ASCOS show nitrate values that are one order of magnitude lower than sulfate concentrations at the same time, usually below 0.1 nmolm^{-3} (C. 25 26 Leck pers. comm., 2015). Moreover, ammonium nitrate does not evaporate and is 27 stable under the electron beam (Rao et al., 1989). To minimize biases due to 28 evaporative losses and beam damage the ammonium sulfate particles were imaged as 29 quickly as possible. Fig. 6 (upper panel) shows the number size distribution of the 30 ammonium sulfate particles derived from the TEM images with a maximum at 172 31 nm in the accumulation mode.

The remaining 65% of the imaged particles (Fig. 5, upper panel) were stable under the heat of the electron beam and showed no sign of evaporation or changes in morphology. Some of those particles appeared as skeletal structures (Fig. 4B) that collapsed and merged to an unstructured flat appearance after exposure times to the electron beam significantly longer than the justified time for imaging of the particles.
 None of the SP particles showed an apparently crystalline appearance that could be
 attributed to sea salt or any other inorganic crystalline matter.

4

5 3.3 Gel-like particles

6 Aerosol particles classified as gel-like particles (GP) using SEM showed an 7 amorphous texture with an inhomogeneous distribution of pixel intensity. Their 8 diffuse structure and weak contrast to the Formvar-film suggested that these particles 9 predominantly contain light elements like C, H, N and O, which are typical 10 components of organic matter. The contrast between the particles and the Formvar-11 film provides (indirect) information about the elemental composition of the aerosol 12 particle since the number of the detected secondary electrons increases with 13 increasing atomic number of the elements present in the aerosol particle (Zhou et al., 14 2006) suggesting that the aerosol particles under investigation are built up by matter 15 of biological origin. The potential similarity in chemical composition between the GP 16 and the Formvar-film might have lead to an underestimation of the particle size which 17 resulted in the shift of the higher accumulation mode peak at 173 nm in TDMPS to 18 147 nm in the total number size distribution (Fig. 3).

GP appeared in the Aitken mode at diameters above 45 nm but were most frequently
observed in the accumulation mode with a maximum at 174 nm, covering all sizes up
to 800 nm (Fig. 5, middle panel). In total 11% of the 3909 particles that were imaged
were classified as GP of which 24% were observed in the Aitken mode > 45 nm,
while 49% appeared in the accumulation mode between 80 and 200 nm, and 27%
were observed > 200 nm (Table 1).

25 Particles classified as GP were further evaluated with TEM. The higher resolution of 26 the TEM images revealed better insight into the morphology of the particles and the 27 GP could be further divided into subgroups (see Fig. 2). Fourteen percent of the 28 particles consisted of a conglomeration of smaller spherical subunits that were welded 29 together and formed small chains or agglomerates (Fig. 7A, B). Those particles were 30 named "aggregate" particles. Twenty nine percent of the gel particles appeared as 31 "aggregate with film" particles where "aggregate" particles were covered with a 32 diffuse and nearly electron-transparent film that partly obscured the underlying 33 subunits and produced a more smooth appearance compared to the bare "aggregate"

particles (Fig. 7 C, D and E). However, for the majority of GP, 57% showed a *"mucus-like"* texture that was many times widely outspread on the Formvar-film
(Fig. 8 A, B), partly in long drawn-out structures (Fig. 8 C) or with small electron
dense inclusions (Fig. 8 D).

5 The individual subunits of "aggregate" particles and the dense spots in "mucus-like" 6 particles exhibit diameters between 11 nm and 109 nm with a maximum in number 7 size distribution at 39 nm and a smaller maximum at 28 nm. Fig. 9 compares the size 8 distributions of "aggregate" components from this study (red line) with those from 9 previous studies in the high Arctic and at lower latitudes (Leck and Bigg, 2005a; 10 2008; 2010). Similarity with previous studies outside and within the pack ice (Leck 11 and Bigg, 2005b; Orellana et al., 2011, same period as this study) strongly suggests 12 the presence of airborne marine gels. Entanglements, ionic or hydrophobic 13 interactions and/or hydrogen bonds stabilize the three-dimensional biopolymer networks of the marine polymer gels, with electrostatic bonds involving Ca^{2+} or Mg^{2+} 14 15 ions being the most dominating (Verdugo, 2012; Orellana and Leck, 2015). In 16 seawater the observed size range of gel particles ranges from solvated nanogels (100-17 200 nm; Bigg et al., 2004) that can further anneal into microgels (> 1000 nm) by 18 interpenetration and entanglement of neighboring nanogels or hydrophobic interaction. 19 Orellana and Verdugo (2003) have shown that changes in environmental factors like 20 enhanced UV-B radiation will avert gel particle formation or induce fragmentation of 21 existing gel particles in seawater (Orellana and Verdugo, 2003). Changes of pH and 22 temperature will lead to volume change of the gel polymer assemblies (Tanaka et al., 23 1980; Orellana et al., 2011). The transport of marine gels from the ocean water into 24 the atmosphere is likely to result in an enhanced exposure of the particles to solar UV-25 B radiation and might have lead to the fragmentation of airborne gel particles. In 26 cloud processing and the condensation of acidic gases onto the gel particles is 27 suggested to lead to a volume change of the gel particles similar to those observed in 28 seawater and produce gel particles with smaller diameters compared to the gel 29 particles observed in seawater (Leck and Bigg, 2005b; Orellana et al., 2011).

30 3.4 Halo particles

Besides SP and GP we observed particles with a halo-like appearance (halo particles,
HP) on the TEM grid where a relatively large central particle was surrounded by a
droplet ring structure of numerous smaller particles (for examples see Fig. 10).

1 Several authors (e.g., Farlow et al., 1977; Bigg, 1986; Bigg et Leck, 2001b) have 2 found that the formation of droplet ring structures from sulphuric acid containing 3 aerosol is a result of humidity, hydrophilicity of the collection surface and impact 4 velocity effects. Bigg and Leck (2001b) observed that a solution of sulfuric acid wets 5 out on a hydrophilic surface but retracts when humidity is reduced leaving behind 6 small droplets in a symmetrical ring. In our study, the sampling procedure led to a 7 drastic reduction in relative humidity from around 100% at ambient conditions to 20% 8 within the sampling manifold (see chapter 2.1.1). The aerosol was impacted onto a 9 surface with hydrophilic properties (TEM grid). We will assume that the HP 10 originally existed as one particle in the atmosphere that splashed out into the droplet 11 ring structure upon impaction onto the substrate.

HP comprised 7% of the total number of aerosol particles (Table 1) and were observed at diameters above 75 nm and in the accumulation mode with maxima at 161 and 293 nm (Fig. 5, lower panel, left). The very weak contrast of the satellite particles against the Formvar background (Fig. 3) probably shifted the particle number size distribution towards smaller sizes to some extent.

17 Imaging with TEM allowed a more detailed investigation of the HP and revealed 18 three morphologically different types of the central particle. We observed central 19 particles in HP that consisted predominantly of skeletal structures or 20 "aggregate"/"aggregate with film" particles. Another types of particles was unstable 21 under the electron beam (Fig. 2). Central particles of skeletal structures or 22 "aggregate"/"aggregate with film" accounted for 19% and 22% of the HP. Examples 23 of both particle types are shown in Figs. 10(A) and 10(B). The majority of central 24 particles (59%), however, was unstable and seemed to partly evaporate during the 25 imaging process leaving more transparent structures behind, and was similar to the SP 26 described in chapter 3.2. We sized the central particles individually in order to 27 compare them with particles of similar morphology within SP or GP. The combined number size distribution of the stable "aggregate", "aggregate with film" and 28 29 skeletal particles exhibited a maximum at 270 nm in diameter (Fig. 6(B) green line). 30 The number size distribution of unstable central particles showed a maximum at 171 31 nm (Fig. 6(B), red line). Unstable SP exhibited a maximum at nearly the same 32 diameter as unstable central particles, at 172 nm (Fig. 6(A)). The consistency in peak 33 position for unstable central particles and unstable SP suggested a similar chemical nature but different atmospheric processing that resulted in condensation of further
 chemical species and the formation of HP.

3 The satellites (particulates or droplets) exhibited varying morphologies. Sometimes 4 numerous small satellites surrounded the central particle in a symmetrical ring (Fig. 5 10(A) whereas in other cases the satellite droplets were larger but fewer (Fig. 10(B), 6 (C)). In the literature three types of compounds have been described to form satellites 7 when airborne aerosol particles impact on a collection substrate: sulfuric acid (Ayers, 8 1978; Ferek et al., 1983), ammonium sulfate and bisulfate (Bigg, 1980; Busek and 9 Pósfai, 1999), and methane sulfonic acid, MSA, (Bigg et al., 1974). Sulfuric acid 10 exhibits a distinctive morphology: a central particle surrounded by a droplet-halo of 11 numerous smaller satellites. Neutralization of sulfuric acid by ammonium or a high 12 content of methane sulfonic acid produces a halo of larger and fewer droplets (Bigg 13 and Leck, 2001a). The morphology of the droplet-halos we observed in this study 14 points towards the presence of sulfuric acid often in a mixture with ammonium sulfate 15 or bisulfate and/or methane sulfonic acid. As discussed in section 3.2, these sulfur-16 containing components have not only been reported to be present over the Arctic pack 17 ice area in summer (Bigg and Leck, 2001a) but also to frequently occur in the remote 18 marine atmosphere (Barnard et al., 1994; Capaldo and Pandis, 1997; Kettle et al., 19 1999). The observed number size distribution for all HP (Fig. 5, lower panel) is in 20 agreement with results from the high Arctic reported by Hillamo et al. (2001), which 21 observed the first maximum in sulfate containing aerosol particles at diameters > 8022 nm and in ammonium and MSA containing particles at diameters > 100 nm.

23 3.5 EDX measurements

24 To determine the elemental composition of the aerosol samples an EDX spectrometer 25 coupled to TEM was used. EDX spectra of 103 aerosol particles were recorded in 26 conjunction with the imaging process. Molecular dynamics studies on polysaccharides by Li et al. (2013) and Sun et al. (2014) have shown that not only the divalent ions 27 Ca^{2+} and Mg^{2+} but also the monovalent ions Na^{+} and K^{+} can stabilize the three-28 dimensional biopolymer gel networks. Inspired by these results, we focused on the 29 detection of the alkali ions Na^+ and K^+ and the divalent ions Ca^{2+} and Mg^{2+} in the gel-30 type particles. In the following, we will refer to Na^+ and K^+ as Na^+/K^+ and Ca^{2+} and 31 Mg^{2+} as Ca^{2+}/Mg^{2+} . 32

The analysis revealed the following characteristics: Na^+/K^+ was detected in 91% of 1 the SP. Thirteen percent of these particles contained exclusively Na^+/K^+ whereas 78% 2 contained both types of metal ions, Na^+/K^+ and Ca^{2+}/Mg^{2+} with the latter only in 3 4 minor quantities (Fig. 11(A)). Examples of EDX spectra for the different types of particles are shown in Fig. S1. The "aggregate" particles contained exclusively 5 Na^+/K^+ in 20% of the particles and predominantly Na^+/K^+ and minor contents of 6 Ca^{2+}/Mg^{2+} in 80% of the particles (Fig. 11(B)). For the particle types "aggregate with 7 *film*" and *"mucus-like"* particles, however, a clear dominance of only Ca²⁺/Mg²⁺ was 8 detected. Eighty-four percent of "aggregate with film" particles contained Ca²⁺/Mg²⁺ 9 as the dominating type of ions. Seventeen percent of the "aggregate with film" 10 particles contained exclusively Ca^{2+}/Mg^{2+} and 67% of the "aggregate with film" 11 particles contained Ca^{2+}/Mg^{2+} that were accompanied by minor contents of Na^{+}/K^{+} 12 (Fig. 11(C)). "Mucus-like" particles also showed a clear dominance of Ca²⁺/Mg²⁺. 13 Eleven percent of "mucus-like" particles contained exclusively Ca²⁺/Mg²⁺ and 86% 14 contained Ca^{2+}/Mg^{2+} that was accompanied by minor contents of Na^{+}/K^{+} (Fig. 11(D)). 15 16 In summary, we observed a gradual transition from particles with a dominating content of Na^+/K^+ to particles with a dominating content of Ca^{2+}/Mg^{2+} moving from 17 SP over "aggregate" particles and "aggregate with film" particles to "mucus-like" 18 19 particles. Therefore, we hypothesise a connection between the morphology of the 20 particles and the respective dominating crosslinking ion within the polymer 3-21 dimensional network of the marine gels. Li et al. (2013) compared polysaccharides 22 with 3 and 4 repetition units of molecular weights of 1.9 kDa and 2.5 kDa, 23 respectively as representations for organic matter in seawater (Verdugo, 2004). Their results showed that the assembly of the longer polysaccharide chains seems to be 24 accelerated in the presence of Ca^{2+} whereas the presence of Na^{+} has a positive effect 25 26 on the assembly of shorter polysaccharide chains. Considering the observed 27 morphology of the aerosol particles that are built up by marine gel matter, a high content of Ca²⁺/Mg²⁺ could facilitate the formation of fluffy and less compact 28 "*mucus-like*" gel matter. The presence of Na^+/K^+ , however, favored a more compact 29 30 structure of type "aggregate" and SP.

Halo particles with a center of gel or fraction of a gel showed a high content of
Na⁺/K⁺: Fifty percent of the particles contained those metals exclusively whereas
another 25% contained mainly Na⁺/K⁺ with smaller amounts of Ca²⁺/Mg²⁺ (Fig. 12).
The high content of alkali metal ions in those particles suggested that they originated

from SP or fragmented *"aggregate"* particles, which were exposed to processes that
 lead to condensational growth of the original gel particles.

3

4 4 Summary and conclusions

5 Aerosol particles collected in the summer high Arctic north of 80° were individually 6 and objectively investigated with electron microscopy and subsequent image mapping. 7 This enabled a division of the aerosol particles into three size resolved gross 8 morphological groups: single particles (SP), gel particles (GP) and halo particles (HP). 9 Single particles (SP) dominated the aerosol population in terms of numbers and made 10 up 82% of all particles; they were observed over the whole sub-micrometer size range 11 and clearly dominated the Aitken mode. The majority of SP (65%) was stable under 12 the electron microscope and showed no signs of evaporation or morphological 13 changes during imaging. These particles with refractory properties appeared over the 14 whole size range of particles observed whereas the remaining 35% of SP appeared to 15 be heat instable, evaporated partly and were predominantly observed in the 16 accumulation mode. GP were observed at diameters > 45 nm predominantly in the 17 accumulation mode with a maximum at 154 nm and contributed 11% to the total 18 particle number. The GP exhibited various morphological features and appeared as 19 "aggregate" particles (14%) and as "aggregate with film" particles (29%), but the 20 majority (59%) was made up by "mucus-like" particles. Seventy percent of the GP in 21 our study appeared to be smaller than 100 nm in diameter, and 90% were smaller than 22 200 nm.

HP appeared mainly in the accumulation mode at diameters > 60 nm with a maximum
at 161 nm, that contributed up to 7% to the total particle number mapped. The
majority of HP consisted of heat instable particles, probably ammonium bisulfate
(59% of all particles), that were internally mixed with sulfur containing compounds
(sulfuric acid, ammonium bisulfate, methane sulfonic acid). The remaining fraction
was made up by "aggregate" particles (19%) and decomposed or fragmented gel
matter (22%) internally mixed with sulfur containing compounds.

Electron dispersive X-ray (EDX) spectroscopy revealed a gradual transition in the
 content of Na⁺/K and Ca²⁺/Mg²⁺ between different particle morphologies. SP and
 "aggregate" particles preferentially contained Na⁺/K⁺ whereas *"aggregate with film"* particles and *"mucus-like"* particles contained mainly Ca²⁺/Mg²⁺ with minor contents

of Na⁺/K⁺. Supported by model studies (e.g. Li et al., 2013; Sun et al., 2014), we
hypothesise a correlation between particle morphology and the presence of the ions
Na⁺/K⁺ and/or Ca²⁺/Mg²⁺. The prevalence of Ca²⁺/Mg²⁺ facilitates the formation of
large organic assemblies of GP type whereas a lack of Ca²⁺/Mg²⁺ and a prevalence of
Na⁺/K⁺ prohibit the formation of large assemblies and result in smaller entities of SP
type.

7 Insufficient knolwedge of the size resolved aerosol composition and especially the 8 role of organic compounds including their morphology and state of mixture, had 9 hampered a detailed understanding of the processes that lead to the activation of the 10 high Arctic aerosol particles and their role in the formation of cloud droplets (Leck 11 and Svensson, 2015; Martin et al., 2011; Zhou et al., 2001; Leck et al., 2002). One 12 evident outcome from this study is that the aerosol particles that would be activated 13 into cloud droplets over the Arctic pack ice areas cannot be seen as simply inorganic 14 salts. None of the aerosol particles showed an apparent cubic or otherwise crystalline 15 appearance that could be attributed to sea salt particles. The results from this study 16 clearly show that organic marine gel matter significantly contributes to the particle 17 number concentration over the whole sub-micrometer size range, especially at 18 diameters below 60 nm.

A parallel study conducted during the ASCOS campaign (Orellana et al., 2011) demonstrated that airborne aerosol particles contain hydrophobic moieties on their surface that play an important role for gel formation (Maitra et al., 2001) and increase the rate of gel assembly (Ding et al., 2008). The interaction of the hydrophilic and hydrophobic entities on the structure of the polymer gels likely will influence the water vapor pressure and decrease the surface tension of the cloud droplets to be formed (Leck and Svensson, 2015; Ovadnevaite et al., 2011).

26 Water-soluble particles like ammonium sulfate were present mainly in the 27 accumulation mode at diameters above 100 nm. Growth of the sub-Aitken particles 28 probably resulted from deposition of acids/organic vapors on polymer gel particles 29 and produced HP or sulfur containing particles with hygroscopic properties typical for 30 a gel nucleus covered by a sulfate-methane sulfonate-ammonium complex. At the 31 same time the fragmentation of larger particles is capable of adding numbers into the 32 Aitken mode (Leck and Bigg, 1999; 2010; Karl et al., 2013). Orellana and Verdugo 33 (2003) and Orellana et al. (2011) observed the sensitivity of marine gels to changes in 34 the physicochemical environment (pH and T) and the fragmentation of gel matter into

1 smaller entities as a result to UV radiation exposure. Acidic compounds typically 2 found in the marine atmosphere like sulfuric acid and dimethyl sulfide (DMS) 3 induced volume collapse of the swollen hydrated polymer gel network into a 4 condensed and more compact form (Tanaka et al., 1980; Leck and Bigg, 2010; 5 Orellana et al., 2011). Condensation of sulfur acidic compounds and in-cloud 6 processing of the marine gels in the atmosphere during their passage over the pack-ice 7 and continuous exposure to UV radiation due to the length of the polar day in summer 8 could produce smaller sized fragments of marine gels similar to the spherical subunits 9 observed in "aggregate"/"aggregate with film" particles and the dense inclusions in 10 "mucus-like" particles. Since SP showed a maximum in number concentration in the 11 same size range (i.e., 27 nm), it cannot be excluded that fragmentation of gel matter or 12 pH induced collapse of the gels lead to the formation of smaller entities that provided 13 a mechanism to produce SP and add particle numbers to the Aitken mode.

In hope of enhancing our understanding on CCN properties promoting/suppressing cloud droplet formation over the pack ice area in summer, and at the same time meet the demand for observational data for the evaluation of climate models, this study has presented critical size resolved data on particle morphology, chemical composition and state of mixture based on the analysis of individual particles.

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	15-80 nm	80-200 nm	> 200 nm	Sum (% of total)
Single particles	2609	573	57	3239
(% of total SP)	(80%)	(18%)	(2%)	(82 %)
Gel particles	97	198	108	403
(% of total GP)	(24%)	(49%)	(27%)	(11 %)
Halo particles	9	95	163	267 (7 %)
(% of total HP)	(3%)	(36%)	(61%)	
Total number of particles			3909 (100%)	

2 Table 1. Numbers and percentage of total for single particles (SP), gel particles (GP)

3 and halo particles (HP) imaged with SEM and used for calculating the number size

4 distributions.

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	No Na ⁺ /K ⁺ , no Ca ²⁺ /Mg ²⁺	Ca ²⁺ /Mg ²⁺	Na ⁺ /K ⁺	Ca^{2+}/Mg^{2+} and Na^{+}/K^{+}
Single particles	4 %	4 %	13 %	78 % (Na ⁺ /K ⁺)
"Aggregate" particles	-	-	20 %	80 % (Na ⁺ /K ⁺)
<i>"Aggregate with film"</i> particles	-	17 %	17 %	67% (Ca ²⁺ /Mg ²⁺)
"Mucus-like" particles	-	11 %	3 %	86 % (Ca ²⁺ /Mg ²⁺)
Halo particles	25 %	-	50 %	25 % (Na ⁺ /K ⁺)

Table 2. Fraction of particles containing the ions Na^+/K^+ or Ca^{2+}/Mg^{2+} , or both Na^+/K^+ and Ca^{2+}/Mg^{2+} , or neither Na^+/K^+ nor Ca^{2+}/Mg^{2+} in single particles, *"aggregate"* particles, *"aggregate with film"* particles, *"mucus-like"* particles and halo particles. The ions written in brackets in the last column indicate the prevalent type of ion in the respective type of particle.



Figure 1. Track of the icebreaker *Oden* in the Arctic (pink). The path during the icedrift is shown in the insert (red line); the circle indicates the start of the ice-drift, the ice edge (thin blue line) was passed on 12 August 2008.



Figure 2. Scheme of the aerosol particle types collected on Formvar grid; the particles observed with SEM are shaded in light blue, and particles observed in TEM are shaded in light green.



Figure 3. Number size distribution of the total aerosol collected for this study with the red line representing the SEM derived particle number distribution, and the red error bars representing the error in size determination retrieved from calibration measurements described in Hamacher-Barth et al. (2013). The blue line depicts the number size distribution from simultaneous Tandem Differential Mobility Particle Sizer measurements, and the blue error bars depict a 5% uncertainty of the data (Wiedensohler et al., 2012).



Figure 4. Examples for single particles (SP) observed with TEM: (A) A particle stable under the beam of the electron microscope; (B) a particle with a skeletal structure; (C) a particle stable under the electron beam; and (D) an example for a particle that is unstable under the electron beam.



Figure 5. SEM number size distribution of the particle types (single particles, gel-like particles and halo particles) evaluated for this study (on the left) plus the fraction of the different subgroups of particles derived from TEM (on the right). The dashed line in each figure marks the lowest diameter at which the respective particle type appears.



Figure 6. Number size distributions obtained from TEM images. Upper panel shows unstable SP while the lower panel exhibits central particles of HP. The red line represents the unstable central particles, while the green line represents the stable central particles comprising *"aggregate"*, *"aggregate with film"* and skeletal particles.



Figure 7. Examples for particles built up by aggregated subunits, observed with TEM: (A), (B) pure *"aggregate"* particles; (C), (D) and (E) aggregate particles covered with a thin film of gel, *"aggregate with film"* particles.



Figure 8. Examples for "*mucus-like*" particles observed with TEM: (A) mucus matter with small dense inclusions, partly outdrawn on the Formvar film; (B) mucus matter without dense inclusions; (C) mucus matter, extensively outdrawn on the formvar film; (D) mucus matter with numerous small dense inclusions.



Figure 9. Number size distributions of airborne aggregate particles and their building blocks at different locations: Mace Head (53°N, 10°W), Lizard Island (14.6°S, 145.5 °E), American Samoa (14 °S, 172 °W), Chichijima (27 °N, 142 °E), High Arctic (AOE-2001, between 88.9 °N and 88.2 °N; light blue line), and ASCOS 2008 (between 87 °N, 1°W and 87 °N, 11 °W, red line). All particles were assumed to be spherical in shape (from Bigg and Leck (2008), modified).



Figure 10. Examples for Halo Particles observed with TEM. (A) a central "aggregate" particle surrounded by satellite particles of sulfuric acid and a smaller amount of methane sulfonic acid. (B) a central particle of "aggregate with film", surrounded by satellite particles of sulfuric acid and methane sulfonic acid. (C) a central particle of ammonium sulfate, satellite particles formed by methane sulfonic acid, probably mixed with sulfuric acid. (D) a central particle of degenerated gel, surrounded by methane sulfonic acid mixed with sulfuric acid.



Figure 11. Fraction of particles containing the following ions: Na^+/K^+ (blue), Ca^{2+}/Mg^{2+} (yellow), Na^+/K^+ and minor contents of Ca^{2+}/Mg^{2+} (red), and neither Na^+/K^+ nor Ca^{2+}/Mg^{2+} (green). (A) Single particles comprised of gel matter; (B) *"mucus-like"* particles; (C) *"aggregate"* particles; (D) *"aggregate with film"* particles.



Figure 12. Fraction of HP containing the following ions: Na^+/K^+ (blue), Na^+/K^+ and Ca^{2+}/Mg^{2+} (red), and neither Na^+/K^+ nor Ca^{2+}/Mg^{2+} (green).