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**Liquid particle in the
Arctic stratosphere**

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Composition analysis of liquid particles in the Arctic stratosphere

C. Weisser¹, K. Mauersberger¹, J. Schreiner¹, N. Larsen², F. Cairo³, A. Adriani⁴, J. Ovarlez⁵, and T. Deshler⁶

¹Max-Planck-Institute for Nuclear Physics, Heidelberg, Germany

²Danish Meteorological Institute, Copenhagen, Denmark

³Institute for Atmospheric Science and Climate, Consiglio Nazionale delle Ricerche, Rome, Italy

⁴Institute for Interplanetary Space Physics, Consiglio Nazionale delle Ricerche, Rome, Italy

⁵Laboratoire de Meteorologie Dynamique, Ecole Polytechnique, Palaiseau, France

⁶University of Wyoming, Laramie, Wyoming, USA

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Correspondence to: C. Weisser (weisser@mpi-hd.mpg.de)

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Abstract

Synoptic scale polar stratospheric clouds (PSCs) that formed without the presence of mountain leewaves were observed in early December 2002 from Kiruna/Sweden using balloon-borne instruments. The physical, chemical, and optical properties of the particles were measured. Within the PSC solid particles existed whenever the temperature was below the equilibrium temperature for nitric acid trihydrate and liquid particles appeared when the temperature fell below an even lower threshold about 3 K above the frost point. The correlation of liquid supercooled ternary solution aerosols with local temperatures is a pronounced feature observed during this flight although the molar ratios $\text{H}_2\text{O}/\text{HNO}_3$ were about a factor of 2 higher than model predictions. In addition HCl has been measured for the first time in liquid aerosols. The chlorine isotope signature served as a unique tool to identify unambiguously HCl dissolved in STS particles. Within a narrow temperature range of about three degrees above the frost point, measured HCl molar ratios are below 1 weight%. There is only fair agreement with model predictions.

1. Introduction

Substantial progress has been made during the last few years to understand the formation and distribution of polar stratospheric cloud (PSC) particles, especially over Northern Europe. New instruments and well-organized combinations of experiments on balloon payloads or research aircrafts have advanced our understanding on formation processes, composition, size, phase, and temperature dependence of PSCs. A European-American campaign in 1999/2000 (SOLVE/THESEO 2000) was particularly successful, producing highlights such as the first stratospheric measurement of solid nitric acid trihydrate (NAT) particles (Voigt et al., 2000) and of unusually large PSC particles within the polar vortex (Fahey et al., 2001). A comprehensive description of many parameters that characterize stratospheric clouds and their environment is now

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available (Newman et al., 2002).

Today we know that polar stratospheric clouds (PSCs) can consist of solid and liquid particles and their existence is controlled by atmospheric temperatures and relevant gas phase mixing ratios. Apart from sulfuric acid tetrahydrate aerosol, nitric acid trihydrate (NAT) particles have the highest equilibrium temperature T_{NAT} . The critical temperature T_{STS} below which supercooled ternary solution (STS) droplets exist is well below T_{NAT} , around 3 K above the frost point T_{ice} . The formation of solid particles requires overcoming a nucleation barrier. Ice freezes out of STS only 3–4 K below the frost point (Koop et al., 2000). Ice particles can serve as a site for the heterogeneous nucleation of NAT particles (Koop et al., 1997). It has become clear that an additional formation process of NAT above T_{ice} is required to explain many PSC observations (Larsen et al., 2002; Drdla et al., 2003). In contrast to the formation of solid ice or NAT particles there is no nucleation barrier for the growth of liquid STS. Below a certain threshold they begin to swell due to the combined uptake of mainly water and nitric acid (Carslaw et al., 1994).

Reported in this paper are results from a balloon campaign performed from Kiruna, Sweden, in early December 2002. The campaign was part of the joint US and European SOLVE-2/VINTERSOL project. The measurements captured a very early period of PSC formation in the 2002/2003 arctic winter. On 4 December 2002 between 20 and 27 km several passes through a PSC were accomplished using controlled up-down motions of the balloon. The balloon flight was carried out exclusively in the cold polar vortex without mountain lee-wave activity. The coldest temperatures encountered during the flight were several degrees above T_{ice} . During the flight the majority of particles were liquid STS droplets, but solid particles were also present. In a separate paper Larsen et al. (2004) shed light on the formation process of the various particles encountered. In the following we address the close correlation of the observed liquid stratospheric particles with the ambient atmospheric temperature. We also compare model predictions with derived $\text{H}_2\text{O}/\text{HNO}_3$ molar ratios.

Despite the progress in PSC research, one important minor constituent expected

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to be present in PSC particles – hydrochloric acid, HCl – has never been measured in liquid or solid particles in the polar stratosphere. HCl is a relatively inert reservoir molecule, which can be chemically changed (activated) through heterogeneous reactions in or on PSC particles when the temperature drops well below 200 K. Although the HCl content of PSCs is of substantial importance, it has only been estimated in atmospheric chemical or microphysical models. Here, we report first measurements of HCl in STS droplets.

The solubility of HCl in liquid aerosols has been treated in some detail in a review paper by Carslaw et al. (1997), which describes the fundamental principles of gas-liquid solution interactions and presents some experimental results besides model calculations. The authors derive a number of relationships between gas and liquid phases of participating species, based on Henry's law, to permit calculations of the amount of gases dissolved in liquid aerosols. It is important to recognize in experimental research that just a few degree change in temperature can alter considerably the H₂SO₄ concentration in liquid particles and as a consequence the solubility of species that include HCl and HBr by orders of magnitude. Thus, it is difficult to simulate stratospheric conditions in laboratory experiments because of the high sensitivity of trace gas solubility to temperature (Hanson and Ravishankara, 1993; Zhang et al., 1993). Here we use for comparison the results of the model by Carslaw et al. (1995), which are similar to the predictions of the Tabazadeh et al. (1994) model and to the work done by Luo et al. (1995). Those models predict at low temperatures just above the frost point the fraction of H₂O, HNO₃, H₂SO₄, and in addition a number of trace gases dissolved in liquid ternary aerosols of the lower polar stratosphere. Within a broad range of parameters that include pressure and HNO₃ mixing ratio, models show, for atmospheric temperatures of around 187 K, HCl mixing ratios in STS between 1 and 0.1 weight%.

2. Meteorological conditions

In contrast to observations with the same set of instruments during previous winters in mountain induced leewave conditions over northern Scandinavia (Deshler et al., 2003b; Schreiner et al., 2003; Voigt et al., 2000), the observations in early December 2002 were characterized by the presence of a large synoptic-scale region with low temperatures over the North Atlantic. The cooling started around mid November when small regions with temperatures below T_{NAT} grew to large areas by the end of November and persisted throughout December within the stable arctic vortex. On 4 December, when the measurements took place, the northern part of Scandinavia was clearly inside the cold pool on the 550 K potential temperature level. Due to low surface winds no orographic induced leewave activity occurred. The observed PSC particles most probably formed in air parcels that spent several days at temperatures between T_{ice} and T_{NAT} . Temperature histories (Larsen et al., 2004) show that the air did neither experience temperatures below T_{ice} nor strong cooling or heating rates, in contrast to the situation one year before in December 2001 (Deshler et al., 2003b). For the first time this kind of comprehensive in situ investigations of polar stratospheric clouds were performed exclusively inside the cold polar vortex without any leewave activity.

3. Observations

Over the last few years, an aerosol composition mass spectrometer (ACMS) has been developed that provided the first chemical analysis of polar stratospheric clouds (Schreiner et al., 1999b). More recent flights, with additional instruments on the gondola, extended the characterization of PSCs and resulted in the first identification of NAT particles (Voigt et al., 2000). Within the ACMS stratospheric particles are concentrated by an aerodynamic focussing lens into a narrow beam, which is directed through a differentially pumped vacuum system. The particle beam finally is stopped and evaporated within a small gold sphere. The evolved gases are analyzed by a

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mass spectrometer with the ion source directly located at the exit of the small evaporation sphere. Surrounding the beam and the mass spectrometer are liquid helium pumps that provide a sufficiently high pumping speed to perform an analysis of the major content of PSC particles such as H₂O and HNO₃. Atmospheric gases, which enter the analysis system with the particle beam, are present in considerably reduced form due to the high-speed pumping. They are occasionally measured to monitor the performance and stability of the experiment. Since the ACMS has a very high sensitivity for condensed-phase material, the detection of HCl became possible when, during the last two years, improvements were made in the reduction of background gases and in higher ion beam intensities. Comprehensive descriptions of the focussing lens and the analysis system are provided by [Schreiner et al. \(1999a, 2002\)](#).

During the flight on 4 December 2002 from Kiruna, Sweden, clouds were present in a temperature range from as low as 184.5 K to about 187.5 K within the polar vortex. While the balloon moved up and down in a pressure range of 35 mbar to about 14 mbar, three layers of clouds were traversed. Figure 1 presents the measurements for the first ascent of the balloon through the cloud. Within the cloud, temperatures stayed below T_{NAT} , but remained above the frost point (panel A). If temperature had gone below T_{ice} and ice particles had formed large peaks would have appeared in the ACMS water signal. Backscatter ratios at 940 and 480 nm ([Rosen and Kjome, 1991](#)) are shown in panel B. The color index which is defined as the ratio between the backscatter ratio at 940 and 480 nm serves as an indicator of the particle phase, indicating solid particles below 500 K and above 580 K potential temperature. The phase can also be inferred from depolarization measurements at 532 nm using a laser backscattersonde ([Adriani et al., 1999](#)); however, these measurements are not shown. Particle size and number concentration were measured with four optical particle counters (OPCs) ([Deshler and Oltmans, 1998](#); [Deshler et al., 2003a](#)). One OPC measured condensation nuclei (CN), $r > 0.01 \mu\text{m}$. Three measured particles with $r > 0.15\text{--}10.0 \mu\text{m}$, whereas one had a heated inlet to evaporate PSC particles and thus measured the background stratospheric aerosol upon which PSCs condense (Eidhammer and Deshler, submit-

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ted, 2004¹). The cumulative size distribution from one OPC with unheated inlet is shown in Panel C. Finally, the particle composition, primarily water and nitric acid, was measured with the ACMS (Schreiner et al., 2002, 2003). In Fig. 1 only water signals are shown (panel D). The time constant of the system for detecting particle water is short and thus water measured on mass 18 is a very good indicator for the onset of particle detection as well as when leaving a cloud. The few spikes in the water signal show the evaporation of single, most probably solid particles in the sphere where the particle beam is stopped. Over the whole altitude range large particles, $>2.0\mu\text{m}$ existed, however, a broad middle layer from 500 to 580 K potential temperature was dominated by numerous liquid particles masking the embedded large particles. Above and below the middle layer were layers of low concentration solid particles as evidenced by the high color index and large particles.

The main constituents of PSC particles are water and nitric acid measured with the ACMS on mass 18 (H_2O^+) and on mass 63 (HNO_3^+) and 46 (NO_2^+). Of particular interest are the signals related to HCl (mass 36 and 38). When clouds were encountered, signals of those PSC particle related masses rose dramatically in the mass spectrometer. Figure 2 shows in panel C experimental evidence that indeed HCl contained in PSC particles was measured. Laboratory calibrations have shown that HCl undergoes little fractionation in the electron impact ion source and therefore masses 36 (H^{35}Cl) and 38 (H^{37}Cl) can be used to identify hydrochloric acid within the mass spectrum with the unique isotope ratio of 3.1. Because it is a trace gas, caution must be applied when the data gathered on those mass peaks are interpreted. ^{36}Ar and ^{38}Ar contributions, besides those from the instrument background, must be subtracted to derive a net HCl signal. From measurements of N_2 and ^{40}Ar , which are occasionally monitored, the potential abundance of the two lighter Ar isotopes can be estimated. In addition, their isotope ratio $^{36}\text{Ar}:^{38}\text{Ar}$ equals 5.5, while the ratio for $^{35}\text{Cl}:^{37}\text{Cl}$ is 3.1. This ratio, as an exclusive indicator for HCl, has been verified during laboratory calibrations of

¹Eidhammer, T. and Deshler, T.: Evaporation of polar stratospheric cloud particles, in situ, in a heated inlet, Atmos. Chem. Phys., submitted, 2004.

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the mass spectrometer. The general background correction was obtained when the gondola travelled outside the clouds. The average Cl isotope ratio, although with considerable scatter, is 3.2 ± 0.3 and thus identifies the isotopes of HCl. Both mass peaks are only present during those parts of the flight when the water signal increased during the encounter with stratospheric clouds (panel A, low mass 18 resolution).

Calibrations of the mass spectrometer before and – if possible – after the balloon flight are a critical component in the analysis of the data. As in past experiments the instrument was calibrated using a dynamic flow system (Kohlmann, 2000; Schreiner et al., 2002), which permits to admit well known amounts of gases that include N_2 , Ar, H_2O , HNO_3 , and HCl to the mass spectrometer ion source. Since the payload was recovered in good condition the instrument was also re-calibrated for the same gases of interest. Fortunately, the sensitivity of the instrument was found to be nearly the same as before the flight. We have only indirect checks on the instrument performance during the flight when atmospheric gases serve as monitors for its stability. Overall, the performance of the mass spectrometer during the balloon flight reported in this paper appears to be stable with no changes in sensitivity detected. Therefore the molar ratios H_2O/HNO_3 and the abundance of HCl in liquid particles will be based on those calibrations.

4. Results

The discussion of results will start with the appearance and molar ratios of STS particles followed by a description of HCl in liquid aerosols.

4.1. STS particles and atmospheric temperatures

On 4 December 2002 PSCs existed between 490 and 610 K potential temperature, approximately from 20 to 27 km altitude. Closely correlated PSC particle properties were detected with the different instruments, as shown in Fig. 1. The cloud was dominated

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by a broad layer of mostly liquid particles ranging from 500 to 580 K potential temperature. These relatively small particles appear in high number concentrations with nearly all background aerosols activated. The ACMS responded with continuous high count rates of water and nitric acid signals. Further characteristics are a high backscatter ratio, a low color index, and low depolarization. Low concentrations of considerably larger, probably solid particles were present over the whole altitude range wherever $T < T_{\text{NAT}}$ as can be seen from the optical particle counter measurements. This is also supported by the volume depolarization. Although the depolarization values are low, they are not as low as expected for a pure liquid cloud of that backscatter ratio. Due to the reduced transmission efficiency of the aerodynamic lens for such large particles only a few were measured with the ACMS. For a composition analysis, however, the counting statistic is too low. In the backscatter data these low concentrations of solid particles are masked by the high number density of liquid particles. They can only be identified at the top and the bottom of the liquid layer by low backscatter, but higher color index and depolarization, when the temperatures become too high for the existence of liquid particles. This gives rise to a “sandwich” structure, which has been previously observed by LIDAR measurements (Shibata et al., 1997; Stein et al., 1999; Biele et al., 2001). Here it seems clear that this structure is simply determined by the growth of STS as the temperature decreases in the middle of the cloud, rather than some features related to the development of the PSC.

One parameter of special interest in PSC research is the molar ratio of $\text{H}_2\text{O}/\text{HNO}_3$ of aerosol particles. The calibrations before and after the flight are intended to derive those ratios with sufficient accuracy. Figure 2, panel B shows the measured molar ratios in comparison with model predictions (Carslaw et al., 1995). All measured ratios are between a factor of 2 and 3 higher than model values, i.e. particles are either deficient in HNO_3 or water-rich. In past experiments (Schreiner et al., 2003; Deshler et al., 2003b), which carried the same ACMS instrument with the same calibrations, molar ratios of liquid particles agreed well with model predictions. Those flights were conducted under lee-wave conditions while here particles in synoptic PSCs were analyzed.

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Both ACMS and OPC data allow to derive the total aerosol volume. Panel E in Fig. 1 shows the measured aerosol volume for the first ascent. For the ACMS the fast and accurate water signal, the laboratory calibrations, and the measured molar ratios were used to assess the aerosol volume. Small corrections for the transmission efficiency of the lens were applied. The aerosol volumes based on measurements of the two instruments agree very well. For comparison predictions for STS particles from the model by Carslaw et al. (1995) are shown assuming 11 ppbv gas phase HNO_3 and using the measured water mixing ratio and the sulfuric acid mixing ratio derived from the background sulfate aerosol measurement. Gas phase HNO_3 was not monitored on the gondola. Measurements of the MIPAS-B instrument that was flown two days later reveal HNO_3 values of 10 ppbv (Oelhaf, personal communication, 2003) in areas with temperatures above T_{STS} , but between 5–10 ppbv due to the uptake in the STS particles. The high atmospheric mixing ratio is consistent with the early period of PSC observations prior to any denitrification.

Figure 3 shows the change in aerosol volume for the transit from inside the layer with liquid particles to outside at the top of the cloud around 590 K potential temperature as a function of temperature difference $\Delta T = T - T_{\text{ice}}$. This is an impressive demonstration, how the existence of STS particles is driven by ambient temperature. As soon as temperature drops below a certain threshold the sulfate background aerosols begin to swell due to the uptake of water and nitric acid and become STS droplets. If they have enough time and if there is enough gas phase nitric acid available they reach equilibrium, which here seems to be the case. It should be noted, however, that the observations do not strictly follow the temporal evolution of the growth or evaporation of the droplets, but show the transition from warmer air masses to a layer with lower temperature where STS can exist. Figure 3 also compares the measurements with model predictions of the total aerosol volume for STS and NAT particles for a range of 5 to 11 ppbv HNO_3 calculated from the model by Carslaw et al. (1995).

There are also data that do not fit the STS nor NAT model predictions. At the bottom and top of the PSC (500 and 610 K potential temperature) the aerosol volumes

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lie above the theoretical calculations for STS, but below NAT. These data points result from larger particles, that are present over the whole altitude range of the PSC as can be seen clearly in the size distribution measured by the OPC, Fig. 1, panel C. The depolarization measurements let us conclude that these particles are solid. In the ACMS measurements and the backscattersonde data they are masked by the high concentrations of small liquid droplets when temperature is below the STS threshold. The large solid particles could be of NAT composition, but their volume did not reach equilibrium expected for NAT, while, as shown above, the volume calculations suggest that the measured STS droplets are in equilibrium with their environment. The deviation of the large solid particles from NAT equilibrium are also reproduced by model calculations presented in the accompanying publication by [Larsen et al. \(2004\)](#).

4.2. First measurements of HCl in STS particles

Model calculations by [Carslaw et al. \(1995\)](#) show a very high sensitivity of dissolved HCl in particles to atmospheric temperature and pressure. Over a small temperature range the HCl solubility will increase dramatically: Between 195 and 187 K, according to the Carslaw model, the HCl content will change by at least a factor of 100. Similarly, substantial changes are calculated over changes in altitude, all related in the end to the change in H_2SO_4 and the water activity of the aerosol. Molar ratios from our measurements, displayed in Fig. 2 panel D, fall within 1 and 0.1 weight% and thus are in the range of model values for the flight conditions (~ 22 mbar, measured 6 ppmv H_2O and estimated 11 ppbv HNO_3 and 1 ppbv HCl in the gas phase). The molar ratios were derived by integrating the measured signals over the time period shown. Adsorption and desorption processes for HCl on the walls of the small particle evaporation sphere are the main limitation on accuracy.

To conclude, HCl in STS has been measured for the first time in a PSC with aerosols present in a temperature range of about 3 K above T_{ice} . The derived molar ratios are clearly higher, even when the range of uncertainty is included, than predictions of PSC models ([Carslaw et al., 1995](#); [Tabazadeh et al., 1994](#)). This difference may have impli-

cations for the main reaction ($\text{HCl} + \text{ClONO}_2 \rightarrow \text{Cl}_2 + \text{HNO}_3$) for chlorine activation involving liquid particles. Clearly it will proceed, as the analysis of [Hanson and Ravishankara \(1994\)](#) suggested, and thus will be a major factor in the subsequent magnitude of ozone reductions.

5. Conclusions

Synoptic scale polar stratospheric clouds that formed without the influence of mountain leewaves were observed in early winter 2002/2003. Measurements described here as well as observations from previous campaigns ([Deshler et al., 2003b](#)) show clearly that over wide areas solid and liquid particles coexist in the Arctic stratosphere. Relatively large solid aerosol particles with low number density, most likely composed of NAT, existed over a broad altitude range when $T < T_{\text{NAT}}$. When temperatures fell below T_{STS} liquid STS particles immediately appeared. The high number density of the small droplets masked the solid particles for identification with backscattersondes, creating the appearance of a sandwich structure of the PSC with a broad layer of liquid droplets surrounded by solid particles at the bottom and on top. The observations here and in December 2002 suggest that the presence of solid particles is almost independent of the temperature at the time of the observation as long as $T < T_{\text{NAT}}$, but it may strongly depend on the temperature history of the air parcels. In contrast, the measurements show a close correlation of the appearance of liquid STS droplets with local ambient temperature. We confirm, particularly because of the accurate temperature measurements, that the predictions made by current STS models ([Carslaw et al., 1995](#); [Tabazadeh et al., 1994](#)) are accurate in describing the growth of liquid STS particles as a function of temperature in the polar stratosphere.

We found, however, that the molar ratios of $\text{H}_2\text{O}/\text{HNO}_3$ are about a factor of 2 higher than predicted by models. The ACMS was carefully calibrated before and after the balloon flight and underwent the same preparations as instruments in previous flights when in mountain leewave conditions the molar ratios of STS particles agreed well

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with model predictions (Schreiner et al., 2003; Deshler et al., 2003b). Finally HCl was measured in liquid particles for the first time. Its volume abundance is higher than models predict by about a factor of 5 or more. Recalibrations after the flight confirmed that the instrument was stable and that indeed HCl was detected.

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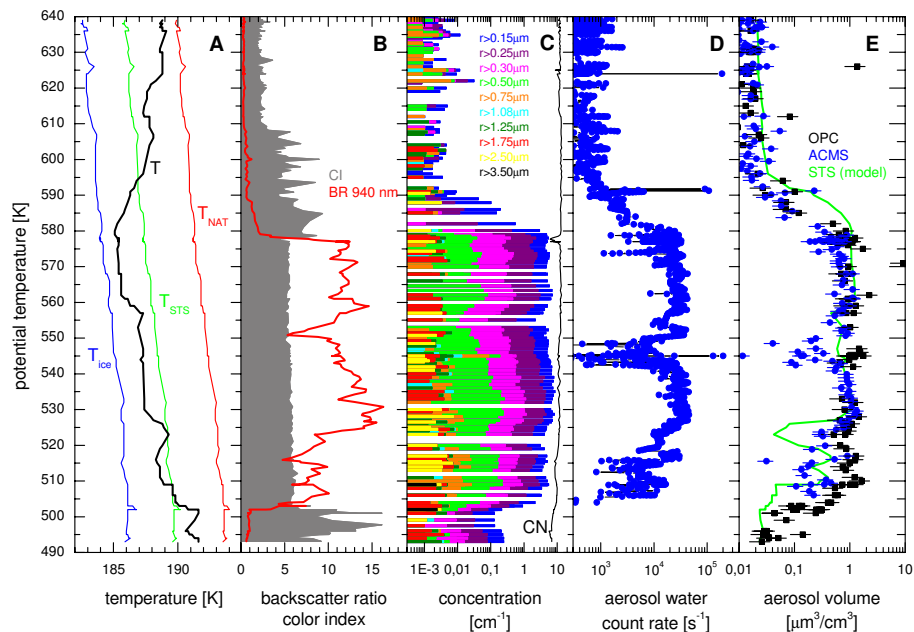


Fig. 1. In-situ measurements during the first ascent through a PSC on 4 December 2002. **(A)** Measured air temperature, accuracy ± 0.5 K, compared to measured T_{ice} and estimates of T_{NAT} and T_{STS} . T_{ice} is measured with a frost point hygrometer (Ovarlez and Ovarlez, 1994). T_{NAT} is calculated from (Hanson and Mauersberger, 1988) assuming 5 and 11 ppbv nitric acid and using the measured water vapor concentrations. T_{STS} is defined as the temperature at which the liquid particle volume changes by 30% for a temperature step of 0.1 K due to the condensation of nitric acid and water, calculated from Carslaw et al. (1995). **(B)** Backscatter ratio at 940 nm and color index. **(C)** Aerosol size and number concentration measured with optical particle counters. **(D)** Aerosol water (mass 18) measured by the aerosol composition mass spectrometer. **(E)** Aerosol volume derived from ACMS and OPC measurements in comparison with model predictions for STS using the measured water vapor and assuming 11 ppbv nitric acid.

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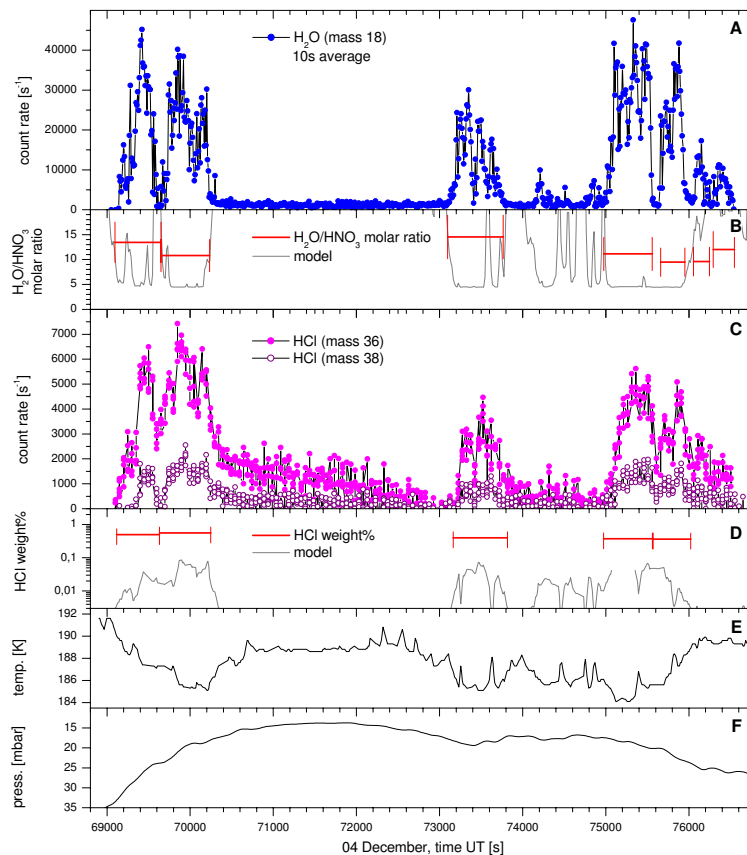


Fig. 2. (A) Water contained in PSC particles measured on mass 18 (H_2O), averaged over 10 s. (B) $\text{H}_2\text{O}/\text{HNO}_3$ molar ratios derived from measurements and model (Carslaw et al., 1995). (C) HCl contained in liquid PSC particles measured on masses 36 (H^{35}Cl) and 38 (H^{37}Cl). (D) HCl weight% derived from measurements and model (Carslaw et al., 1995). (E) Measured atmospheric temperature. (F) Measured atmospheric pressure.

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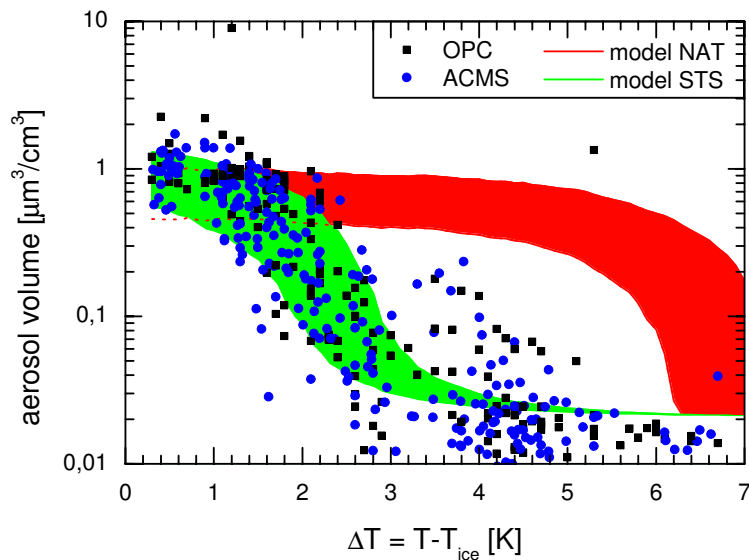


Fig. 3. Aerosol volume measured by the optical particle counters and derived from the aerosol composition mass spectrometer as a function of temperature difference $T - T_{\text{ice}}$. Measurements are compared with aerosol volumes for STS and NAT predicted by model calculations, using Carslaw et al. (1995). The measured water vapor mixing ratio was used, and a range of 5 to 11 ppbv gas phase HNO_3 was assumed.

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