Technical Note: Using DEG CPCs at upper tropospheric temperatures

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Abstract

Over the last few years, several Condensation Particle Counters (CPC) capable of measuring in the sub-3 nm size range have been developed. Here we study the performance of Diethylene glycol (DEG) based CPCs at different temperatures during Cosmics Leaving Outdoor Droplets (CLOUD) measurements at CERN. The data shown here is the first set of verification measurements for sub-3 nm CPCs under upper tropospheric temperatures using atmospherically relevant aerosol particles. To put the results in perspective we calibrated the DEG-CPC at room temperature, resulting in a cut-off diameter of 1.4 nm. All diameters refer to mobility equivalent diameters in this manuscript. At upper tropospheric temperatures between $-25^\circ$C and $-65^\circ$C, we found cut-off sizes in the range of 2.5 and 2.8 nm. Due to low number concentration after size classification, the cut-off diameters have a high uncertainty ($\pm 0.3$ nm) associated with them. Operating two laminar flow DEG CPCs with different cut-off sizes together with other aerosol instruments, we looked at the growth rates of aerosol population in the CLOUD chamber for particles smaller than 10 nm at different temperatures. A more consistent picture emerged when we normalized the growth rates to a fixed gas-phase sulfuric acid concentration. All of the instruments detected larger growth rates at lower temperatures, and the observed growth rates decreased as a function of temperature, converging with each other at temperatures over $0^\circ$C. The theoretical calculations had a much smaller temperature dependency.

1 Introduction

Condensation particle counters (CPCs) measure the total number concentration of particles by activating and growing the aerosol particles with a supersaturated vapour. Once the particles are grown they are counted optically with a laser (McMurry et al., 2000). They provide information about the total particle concentration above a certain size. The parameter used to describe the performance of a CPC is the cut-off diameter,
which is defined as the diameter of the particles that are successfully counted with at least 50% efficiency. By using several CPCs with different cut-off diameters, additional information such as the growth rates can be retrieved (Kulmala et al., 2007). In this paper all diameters refer to mobility diameters. CPCs capable of measuring reliably in the size range below 3 nm have been developed only recently (Kuang et al., 2011; Vanhanen et al., 2011; Jiang et al., 2011a; Sipilä et al., 2009). CPCs are widely used for ground-based measurements, but they are also deployed during airborne measurements (Crumeyrolle et al., 2010; O’Dowd et al., 1998; Schobesberger et al., 2013b), where ambient conditions, such as temperature and pressure, are changing. It has been shown (Kulmala et al., 2007; Wimmer et al., 2013; Kangasluoma et al., 2013) that the counting efficiency curves of the sub-3 nm CPCs change according to the particle composition. As the working principle of laminar flow CPCs is based on creating supersaturation and activation by temperature differences, their performance might depend also on the ambient temperature. One issue with CPCs measuring at low temperatures is whether the particles evaporate during the transition from the cold measurement environment to the warmer CPC. Another issue is whether the temperature profile (and thus supersaturation) inside the instrument changes when cold sample air is brought in, maybe resulting in a cut-off shift.

At CLOUD measurements were performed to study nucleation under upper tropospheric temperatures. Therefore it is highly interesting to determine whether the cut-off size of the DEG CPC that was used to directly measure the nucleation rate at 2 nm was changing due to the ambient temperature. As the growth rates at small sizes can be size dependent (Kulmala et al., 2013; Nieminen et al., 2010) it is also important to measure them reliably as a function of diameter. Growth rates were determined at 2 nm from different instruments and compared to each other. Here especially the comparison with an ion spectrometer gives important additional information as this method is temperature independent, which is not the case for CPCs.

The aim of this study is to investigate the possibility to use CPCs at very low temperatures, which can be encountered in the upper troposphere (e.g. during flight
measured efficiencies). Especially we studied how the counting efficiency curve of the DEG CPC changes due to temperature, and we also studied how the growth rates behave for different instruments at low temperatures.

2 The CLOUD facility

The Cosmics Leaving OUtdoor Droplets (CLOUD) experiment was designed to study nucleation in a controlled environment under various conditions (Duplissy et al., 2010; Kirkby et al., 2011). In the design, great care was taken to ensure high cleanliness, as it has been shown that impurities of trace gases (already at concentrations as low as few pptv), such as NH₃ influence the nucleation rates (Ball et al., 1999; Kirkby et al., 2011; Schobesberger et al., 2013a). The chamber as well as all the gas lines and connections to the chamber are made of stainless steel. The CLOUD facility comprises a 26.1 cm³ cylindrical stainless steel chamber with a fibre optic UV system (Kupc et al., 2011). It includes a very stable temperature control unit (0.01 K) with a temperature range from −70°C up to +100°C. Two mixing fans are installed in the upper and lower part of the chamber to ensure homogenous internal mixing (Vogtländer et al., 2012). Measurements can be performed under high ion pair production rates (pion beam from CERN proton-synchrotron on, clearing field grounded), natural level of ion production rates (beam off, clearing field grounded) and completely neutral conditions (beam off, clearing field on). The clearing field is created by applying ±30 kV to two circular grids at the top and the bottom of the chamber, removing the small ions, which are continuously generated by the galactic cosmic rays in less than 1 s. The nucleation measurements in the CLOUD5 campaign, which took place in fall 2011, were mainly performed at three different ranges of temperatures: between [206–208] K, [221–223] K and [245–248] K. To compare the performance of the chamber and the instruments to previous experiments, measurements at +5°C were also performed. In CLOUD the instruments draw air from sampling lines with a length of 75 cm and an outer diameter of 1/2 inch. To guarantee the same temperature in the chamber and in the sampling lines, the
sampling lines were thermally coupled to the chamber, using copper. To avoid thermal exchange with the room air, the sampling lines were wrapped with thermal insulation material on top of the copper wires.

For the measurements analyzed in this study only sulphuric acid and water were introduced into the chamber. A Chemical Ionisation Mass Spectrometer (CIMS, Kürten et al., 2010) was used for measuring the gaseous sulphuric acid (SA) concentration. The sub-5 nm CPCs that were used for measuring the total particle concentrations and calculating nucleation rates were a Particle Size Magnifier (PSM; Airmodus A09), a butanol based laminar flow CPC (3776; TSI Inc.) and two DEG based laminar flow CPCs. Details about the setup and performance of the DEG CPCs can be found in Wimmer et al., 2013. It should be noted that the DMA that was used for size selecting the particles for the calibration measurements shown here was a standard nano-DMA, whereas for the calibration measurements at room temperature, a high resolution DMA was used, which means that the calibrations in the laboratory have a better size resolution. A comparison of the performance of the different DMAs can be found in Jiang et al. (2011b).

### 3 Calibrating CPCs using the CLOUD chamber as an aerosol source

A set of calibration measurements was performed using the equipment available in CLOUD. The calibration unit consisted of the CLOUD chamber as the particle source. Doing so, the calibration particle composition was exactly the same as during the nucleation experiments. The particle production was triggered by switching on the UV lamp after adding SO$_2$, O$_3$ and water vapour to the chamber. The particles were sampled from a core sampling probe, which was used to minimize the diffusion losses. The core sampling probe consisted of a coaxial tube of with a conical tip and an outer diameter of 6 mm. A fraction of the flow was sampled through that tube from the center of the flow, whereas the flow closer to the walls of the main tube was discarded. No external charger was used, only the naturally negatively charged fraction of particles.
which were size selected with a nano Differential Mobility Analyzer (nano-DMA; Grimm Aerosol Technik, S-DMA, 55-100) were used for the calibration. The laminar flow DEG CPC is the instrument that was calibrated. Figure 1 shows the setup for the calibration measurements. To measure the total concentration after the DMA, as a first test, an electrometer was used as a reference. The particle concentrations after the size selection were low due to the low charging probability and internal losses in the DMA. This resulted in an unacceptably high noise to signal ratio and therefore it was decided to use a PSM as the reference instrument instead of using an electrometer. The PSM has a significantly lower cut-off size than the laminar flow DEG CPC and a steep counting efficiency curve ($d_{50} = 1.1$ nm, $d_{90} = 1.5$ nm for ammonium sulphate Wimmer et al., 2013) and it is a single particle counter. We assume that the PSM behaves similarly to the laminar flow DEG CPC regarding the ambient temperature. With this setup, the problem with a too low signal to noise ratio could be reduced.

During the calibration measurements, the DMA was set to stepping mode, to obtain better statistics for each size bin, since the particle population in the chamber was growing constantly. Another experimental challenge during the calibration measurements was to not only produce enough particles, but also keep them growing slowly enough, as the size range of the DMA was limited to below 10 nm. Slow enough growth was achieved by switching on the UV lights at very low intensities for a few minutes, letting the particles grow for a couple of minutes. As soon as the particles grew to sizes bigger than 10 nm, the UV was switched on again to trigger new growth of small particles. To ensure stable temperature of the calibration system, the DMA was located in a temperature controlled rack, which was kept at $-30^\circ$C for the measurements at $-65.9^\circ$C and $-50.5^\circ$C, whereas the rack was at $+2^\circ$C for the measurements at $-26.9^\circ$C. Due to technical limitations of the temperature control of the racks and the heat load of the instruments inside the racks, it was impossible to reach the same temperature as in the chamber. For the inlet system to the DMA, a core sampling line was used to avoid high diffusion losses. The make-up flow for the DMA was set to 10 lpm, while the aerosol flow through the DMA was 2 lpm. The sheath flow of the DMA passed through a heat
exchanger. The dilution flow after the DMA was taken from a different sampling line to guarantee the same composition and temperature of the stream. By setting the sheath flow to 17 lpm, a good sheath air to sample flow rate ratio was achieved to ensure a narrow size distribution in the DMA (Jiang et al., 2011b).

4 Method for determining growth rates from the DEG CPCs

The low cut-off CPCs (DEG-CPCs and PSMs) have been used at CLOUD also to determine the initial growth rate (GR) of the particles (Lehtipalo, 2014). This is relevant e.g. for scaling the nucleation rates, measured at larger sizes (Kirkby et al., 2011; Alemeida et al., 2013; Kulmala et al., 2013) down to smaller sizes (e.g. 1.7 nm). To determine whether the shift in the cut-off curves influences the scientific outcome of the measurements with the laminar flow DEG CPCs significantly, the growth rates have been determined and compared to other instruments (NAIS, Mirme et al., 2013, and PSM). As the two laminar flow DEG CPCs used here were developed for CLOUD mainly and they have not been used before for atmospheric measurements the method for calculating the growth rates from these instruments had to be determined.

An example illustrating the chosen method is shown in Fig. 2. The total measured number concentration was normalized to the maximum for each run separately. Next the time steps when the concentrations reached half of the maximum concentrations were determined. Based on the rise times of the different CPCs used here (PSM $d_{50} = 1.1$ nm, TSI 3776 $d_{50} = 2.9$ nm), the difference in the cut-off diameter of the two DEG CPCs was estimated to be 0.3 nm. The time difference was then used for determining the growth rates. The times used for the growth rate analysis are marked with arrows in Fig. 2. The half maximum points here were reached at 14:10:45 and 14:12:50 UTC resulting in a time difference of 125 s and therefore in a GR of $9.3 \text{ nm h}^{-1}$. In addition, the GRs from the PSMs were determined in the same manner (Lehtipalo et al., 2014).

The temperature in the chamber in this case was $-50^\circ$C and the electrodes of the high voltage control were set to $\pm 30$ kV, so it was a neutral run. The experiments were done...
at different temperatures and sulphuric acid concentrations. To be able to compare the
determined GR at different temperatures the growth rates were normalized to the same
sulfuric acid concentration ($10^7 \text{ cm}^{-3}$). The normalization was done using the following
equation

\[ \text{GR}_{\text{norm}} = \frac{\text{GR}_{\text{calc}}(\text{SA}) \times 10^7}{\text{SA}_{\text{meas}}}, \]

\[ \text{GR}_{\text{calc}} \]

GR here means the GRs calculated from the CPC measurements. Growth rates
higher than 20 nm h$^{-1}$ have been neglected, as then the time difference between the
two laminar flow DEG CPCs becomes too small.

5 Results

5.1 Upper tropospheric temperature calibration results

Figure 3 shows the calibration curves at different temperatures ($-65.9^\circ \text{C}$, $-50.5^\circ \text{C}$,
$-26.9^\circ \text{C}$, $+25^\circ \text{C}$). Due to the huge scatter and large error bars a number of the red
data points seem to indicate a small cut-off at $-65^\circ \text{C}$, but the majority of the data is
at zero and the fit shows that the cut-off is shifted. The data points Fig. 3 are taken
from the experiments with the highest particle concentrations after the DMA. Three-
parameter sigmoidal fits are applied to the measured data points

\[ \text{fit}(d_p) = a + \frac{b}{1 + \exp(-(d_p - b)/c)), \]

and are weighted based on the error bars of the measured data points. To fit the
$-65.9^\circ \text{C}$ data, the parameter $c$ had to be fixed to 0.3 and also the base value (para-
meter $a$) was fixed to 0, otherwise the fit would not have converged due to the large
error bars in the smaller sizes. The values at half maximum (= parameter $b$) given by
the fits are $2.5 \pm 0.05$ for the data points at $-65.9^\circ \text{C}$, $2.6 \pm 0.1$ for $-26.9^\circ \text{C}$ and $2.8 \pm 0.05$
for the data taken at $-50.5^\circ \text{C}$. As the overall particle concentration after the DMA was
typically 10 cm$^{-3}$, the relative errors of the measurements are large. This means that the results are qualitative. Time and technical limitations prevented the acquisition of more data. Nevertheless there is a clear shift in the cut-off diameter towards larger sizes at lower temperatures. This could indicate that there is some evaporation of the smallest particles, preventing detection in the CPC. The reason for that might also be due to the fact that the rack with the calibration setup was at −30 °C for all the measurements, because the temperature control for the rack was not capable of going lower, as the heat load in the rack was bigger than expected. The trend is not very clear on the other hand as the −26.9 °C curve does not fall in line with the other results. The shift in the cut-off size could also be caused by shifts in the temperature-dependent diffusion losses, but the calculation (Baron and Willeke, 2001) shows that this difference is negligible. The diffusion loss corrected $d_{50}$ is at 2.7 nm, whereas it is 2.8 nm for the uncorrected data points for the −26.9 °C data points. Another explanation for the unexpected behaviour for the calibration results at −26.9 °C might be that for those measurements the temperature inside the temperature controlled rack was at +2 °C whereas it was −30 °C for the other low temperature calibration measurements.

5.2 Comparison of growth rates

Figure 4 shows the growth rates in nm h$^{-1}$ on the x-axis as a function of temperature for the laminar flow DEG CPCs, the PSMs and the NAIS. The growth rates are normalized to a sulfuric acid concentration of $10^7$ cm$^{-3}$. As the growth rates were calculated from different nucleation experiments performed at different sulfuric acid concentrations, the normalization was done as described in Eq. (1). All the GRs are determined at about 2 nm. The GRs are averaged for the different temperature ranges [206–208] K, [221–223] K and [245–248] K and also the temperatures are averaged for each range. The error bars show the standard deviation of the averages of the growth rates. When comparing GR from NAIS, PSM and DEG CPCs, all instruments show the same trend. The growth rates are higher for lower temperatures and the data points converge for
the +5°C and +25°C data. The GR calculated from the DEG CPCs and the PSM agree within a factor of 1.2, whereas the GR from the NAIS are about a factor of 2.2 higher compared to the DEG CPCs at −65°C. The higher GR from the NAIS might be due to the fact that it was measuring negatively charged particles only which are expected to be higher than neutral GRs, also they were calculated following the mode of the distribution (i.e. the maximum concentration in the respective size bin). The calculated collision limited growth rates show the opposite trend, which is very likely due to the effect that it does not take evaporation into account.

5.3 Theoretical calculation of growth rates

The GR can be described as,

\[ \text{GR} = k_c \Delta d_p, \]  

(3)

\( \Delta d_p \) is the change of diameter due to the condensing sulphuric acid and \( k_c \) is the condensation rate. The condensation rate and the change in diameter was determined for a particle of 2 nm with the SAWNUC model (Lovejoy et al., 2004). Equation (3) is only valid for situations without evaporation. Very small particles have higher evaporation rates than bigger particles, due to the Kelvin effect (McKenzie et al., 1984). In fact particles below the critical diameter, if such a diameter exists, will mainly evaporate while particles bigger than the critical diameter will grow. In these calculations the evaporation effect on the growth rate is given by,

\[ \text{GR} = (k_c - k_e) \Delta d_p. \]  

(4)

With \( k_e \) being the evaporation rate as described by Lovejoy et al., 2004. This equation allows the evaporation of particles if the vapour pressure of condensing vapour is low, i.e. evaporation dominates over condensation. At higher temperatures the growth rates were calculated for a sulphuric acid concentration of \( 7 \times 10^8 \text{ cm}^{-3} \), as at this temperature a significant background contamination of NH₃ was present in the CLOUD 12806
chamber (Kirkby et al., 2011). NH$_3$ can lower the saturation vapour pressure of sulfuric acid over these small aerosols, though the magnitude of this effect depends on the composition in gas and aerosol phase (Clegg et al., 2008). SAWNUC includes only aerosols containing H$_2$SO$_4$ and water in the calculation, the calculations give nevertheless a good indication about the growth rate behaviour when evaporation can not be neglected. The results from those calculations compared to the experimental data are shown as the purple triangles in Fig. 4. The comparison with the data shows that the SAWNUC predictions show a similar temperature trend as the experimental data (i.e. smaller GR for higher temperatures). This drop off at higher temperatures remains unclear, nevertheless the higher GR retrieved from the NAIS data compared to the SAWNUC model sand to the GRs from the CPCs at the lower temperatures might be due to a charge enhancement (Laakso et al., 2003).

6 Conclusions

We investigated the performance of the Diethylene Glycol (DEG) based CPC under temperatures typical for the upper troposphere, by performing a set of calibration measurements at temperatures between $-25^\circ$C and $-65^\circ$C. Here the CLOUD chamber was used as aerosol particle source and a nano-DMA to size select the particles. As the total particle concentration was low, a single particle counter (Particle Size Magnifier; PSM) was used as reference instrument for the calibration measurements. The results show that there is a shift in the cut-off diameter of the CPC towards bigger sizes compared to the laboratory calibrations at room temperature. The cut-off sizes were 2.5 nm for $-65.9^\circ$C, 2.8 nm for $-50.5^\circ$C, 2.5 nm for $-26.9^\circ$C and 1.4 nm for $+25^\circ$C (room temperature) for a DEG CPC. The calibrations at $-26.9^\circ$C do not fall in line with the other temperatures as the cut-off curve is between the $-50^\circ$C and $-65.9^\circ$C measurements. However as the maximum concentration of particles after the DMA was about 10/cc, the uncertainty in the cut-off diameters retrieved from those measurements are large. This means that the results show a clear shift in the cut-off diameters.
towards the bigger sizes, but the exact numbers for the cut-off diameters need to be treated carefully. Nevertheless for the first time, a sub-3 nm CPC has been calibrated under upper tropospheric relevant temperatures. In addition the calibration measurements were done using the same particle composition as for the nucleation measurements. That means that the shift in the cut-off diameters is only due to the different temperatures and not influenced by the chemical composition of the particles.

We used a method to analyze growth rates based on the difference in the cut-off sizes between two DEG CPCs, using the results from the calibration measurements and compared the calculated growth rates with those derived from other instruments. To evaluate the validity of the results, the GR have been compared to the growth rates determined from mixing type CPCs (PSMs) and an ion spectrometer (NAIS) at different temperatures. All the instruments show higher GRs for lower temperatures. The average value of the growth rates from the NAIS show somewhat higher values (factor of 2.2) compared to the GRs determined from the DEG CPCs. The growth rates between PSM and DEG CPCs agree within a factor of 1.2. The growth rates converge for all instruments for temperatures above 0 °C. This indicates that growth rates at low temperatures might be under-estimated. However the CPCs detect both charged and neutral particles, whereas the NAIS measured only negatively charged particles. The GRs calculated from kinetically limited collision rates of sulfuric acid and water show the opposite temperature trend, but here evaporation is not taken into account. The SAWNUC (Lovejoy et al., 2004) model where evaporation is taken into account, shows a similar temperature trend. We have shown that when using CPCs at upper tropospheric temperatures, it has to be taken into account that cut-off diameters might be shifted towards bigger sizes. This effect might be due to evaporation of the smallest particles or to a change in the saturator temperature (i.e. supersaturation) profile. This means that one should be careful when using CPCs for measuring nucleation mode particles and when calculating nucleation and growth rates. Taking the cut-off shift into account CPCs can still be reliably used as total particle counters, depending on the size distribution.
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References


Figure 1. Sketch of the calibration setup. The calibration unit was in a temperature controlled rack, while the CPCs were at room temperature. The temperature inside the rack was $-30\, ^\circ C$ for the measurements at lower temperatures, whereas the rack temperature was $+2\, ^\circ C$ for the measurements at $-26.9\, ^\circ C$ in the chamber. To minimize diffusion losses, a core sampling probe was used from the chamber to the calibration system. The CLOUD chamber was used to produce naturally charged particles. A nano-DMA was used to size select the particles and as reference a PSM, with a significantly lower cut-off (1.1 nm) than the calibrated CPC. The sheath air of the DMA and the dilution flow after the DMA went through a heat exchanger, consisting of copper tubing to ensure the same temperature for all flows.
Figure 2. Example of a nucleation experiment during CLOUD5. The temperature inside the chamber in the shown example was $-50{\degree}C$ and the clearing field was at high voltage, meaning it was a neutral run. The red, blue, green and the black curves show the normalized concentrations for the DEG CPCs, PSM and the TSI 3776 CPC respectively. The grey line shows the normalized sulfuric acid concentration measured by the CIMS. The growth rates are calculated by finding time difference at half maximum between the two DEG CPCs, assuming a difference in diameter of 0.3 nm.
Figure 3. Results from calibration measurements. The blue, green and red curves show fits to the results from calibration measurements done at CLOUD in fall 2011 and the symbols show the measured data points, at three different temperatures (−26.9 °C, −50.5 °C, −65 °C). Black curve shows laboratory calibrations (Wimmer et al., 2013), using negatively charged ammonium sulfate clusters. The $d_{50}$ is shifted from about 1.4 nm at room temperature to 2.5 nm in the case of −65.9 °C about 2.6 nm for −26.9 °C and for the −50.5 °C, the cut-off is shifted to 2.8 nm. To illustrate the values for the cut-offs, dashed lines pointing to the according sizes are added to the figure. Due to low statistics, the error bars are rather large.
Figure 4. The growth rates normalized to a sulfuric acid concentration of $10^7 \text{ cm}^{-3}$ as a function of temperature. The GRs have been calculated using the laminar flow DEG CPCs, the PSMs and the NAIS. Red symbols show the average growth rates at the given temperature determined from the DEG CPCs, from the NAIS in blue and from the PSMs in green. The error bars show the standard deviation of the average. The lines are fits to the averages and are weighted for the error bars. The black line shows the expected growth rates based on kinetic collisions of sulfuric acid and water molecules (Nieminen et al., 2010). The purple triangles show calculations using the SAWNUC model. The modeled results show a similar temperature trend, as here evaporation is taken into account.