Continuous standalone controllable aerosol/cloud droplet dryer for atmospheric sampling

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Abstract

We describe a general-purpose dryer designed for continuous sampling of atmospheric aerosol, where a specified relative humidity (RH) of the sample flow (lower than the atmospheric humidity) is required. It is often prescribed to measure the properties of dried aerosol, for instance for monitoring networks. The specific purpose of our dryer is to dry highly charged cloud droplets (maximum diameter approximately 25 µm) with minimum losses from the droplet size distribution entering the dryer as well as on the residual dry particle size distribution exiting the dryer. This is achieved by using a straight vertical downwards path from the aerosol inlet mounted above the dryer, and removing humidity to a dry closed loop airflow on the other side of a semi-permeable GORE-TEX membrane (total area 0.134 m²).

The water vapour transfer coefficient, $k$, was measured to $4.6 \times 10^{-7}$ kg m$^{-2}$ s$^{-1}$ % RH$^{-1}$ in the laboratory and is used for design purposes. A net water vapour transfer rate of up to $1.2 \times 10^{-6}$ kg s$^{-1}$ was achieved in the field. This corresponds to drying a 5.7 L min$^{-1}$ (0.35 m$^3$ h$^{-1}$) aerosol sample flow from 100 % RH to 27 % RH at 293 K (with a drying air total flow of 8.7 L min$^{-1}$). The system was used outdoors from 9 May until 20 October 2010, on the mountain Brocken (51.80° N, 10.67° E, 1142 m a.s.l.) in the Harz region in central Germany. Sample air relative humidity of less than 30 % was obtained 72 % of the time period. The total availability of the measurement system was > 94 % during these five months.

1 Introduction

Atmospheric aerosol is important for climate. Large efforts are done to systematically investigate and monitor properties thereof. Many aerosol properties vary with relative humidity (RH). For measurements, and especially for data quality concerns for comparisons between stations/networks, a dry aerosol is important. Aerosol dryers and the drying process have been described earlier (e.g. Martinsson et al., 1992). One
automatically regenerating dryer has recently been presented by Tuch et al. (2009), see references therein for additional drying methods. That dryer used a three-way valve alternating between two silica gel dryers. Transmission for that system was well characterized in the diameter size range from 3 nm to 800 nm. The 50 % transmission for larger particles was calculated at 6 µm diameter. In our application we want to extend the aerosol measurement size range to drying of larger particles and cloud/fog droplets up to 25 µm diameter, which require a straight vertical path, in order to reduce impaction and sedimentation losses. This excludes the use of a standard valve or tee. The dryer should also work continuously. Therefore the design presented in this paper was used. In addition, the drying airflow rate can be varied to maintain a set RH (lower than ambient) in the sample flow, should that be required (in the future an additional humidifying setup can also be envisaged to maintain constant say 25 % RH). A closed loop for the drying air was designed for minimum variability due to flow changes and gas transfer into the aerosol sample flow.

2 Design of the dryer

The main principle of the dryer is to dry sample air, flowing vertically downwards, in counterflow with dry airflows, these flows being separated by membranes. Two concentric, cylindrical membranes were used, with the sample flow in the middle, annularly, between the dry airflows (see Fig. 1). The membrane is needed due to different flow velocities and flow directions of sample flow and drying flow, which would otherwise result in mixing between the two. The membrane allows water vapour exchange by diffusion, but no particle transfer. The two dry airflows used are arranged in closed loops, in order to prevent leakage of air between the flows (either through the membrane or at connections).

As design criterion a sufficient transfer rate of water vapour is required to maintain the dried sample flow below a user specified maximum RH. This specified RH and the minimum expected temperature of the dried sample flow define the maximum allowed
specific humidity tolerated, after drying. The ambient corresponding parameters to consider for this criterion are the estimated maximum ambient RH and the maximum ambient temperature (i.e. the maximum ambient specific humidity), at times when drying is required. The difference between these two values of specific humidities gives the transfer rate of water vapour required, for a given sample flow. This transfer rate, in turn, determines the area of the membrane ($A_{\text{mem}}$) needed, and as well defines the drying airflow rate required, which takes up the humidity. We used GORE-TEX membrane due to performance, flexibility in fabrication options and dimensions, as well as having former experience with this type (Weingartner et al., 2002). The GORE-TEX membrane itself is thin and fragile and must be fixed on a holding surface, such as textile or plastic fiber mat, we chose plastic (see below for details). For our setup, tests we made (at 293 K) resulted in a practical water vapour transfer coefficient, $k$, of $4.6 \times 10^{-7} \text{ kg m}^{-2} \text{ s}^{-1} \% \text{ RH}^{-1}$ from the humid to the dry side, defined as the amount of water vapour passing one m$^2$ membrane per s per % RH difference between the two sides of the membrane. % RH is chosen in the unit here as that is most readily available during measurements. The transfer rate for a membrane can vary substantially with temperature, due to physical changes in the microstructure of the membrane material, influencing its porosity.

The actual dryer described here is used for an instrument measuring cloud droplet size distributions, the Droplet Aerosol Analyser (DAA) (Martinsson, 1996). The DAA requires typical cloud droplets to fall vertically into and inside the dryer, at least until dried to submicrometer-sized droplets or particles, in order to minimize droplet losses due to impaction. Such losses prevent us from using any ordinary type of valve before the drying process, and we have thus chosen a straight vertical path from the inlet through the dryer. The dryer considered consists of two concentric, cylindrical stainless steel mesh metal nets (wire diameter 0.35 mm, open area 2.25 mm$^2$) (see Fig. 1), with the sample flow in the middle between these. On the other sides of the metal nets are two cylindrical membranes (inner and outer membrane tube). The plastic fiber mats supporting these membranes are facing towards from the aerosol flow. Two separate
closed loop dry airflows circulate on each of the other sides of the membrane tubes, achieving the transfer of humidity from the sample flow due to the humidity gradient (pumps used are model 6025se/12vdc, Thomas, USA). The metal nets reduce electrostatic charging of the membranes and are fixed/grounded to the stainless steel main housing of the dryer, and also act to stabilise the membranes mechanically. The dryer has a total effective area of 0.134 m$^2$ (see Table 1).

Once the area of the membrane has been chosen, several design geometries can be proposed. One can envisage for instance a maximum distance from a sample air streamline to the dry airflow in order to avoid too large inhomogeneities in humidity in the sample flow, as well as a simpler geometry than ours, with only one layer of membrane used (one tube, with aerosol in the middle and dry air on the outside). Further design geometries considerations are outside the scope of this paper. However, there are three further design criteria to be considered once the area is determined, namely limitations of diffusion/impaction losses (e.g. von der Weiden et al., 2009), electrostatic losses (Fuchs, 1989) and residence time needed for drying. These will be briefly described in the following three sections. The criteria could be less stringent if only submicrometer aerosol is dried, or one criterion might require more attention, such as diffusion losses if nanoparticles should be studied, for instance.

### 2.1 Diffusion/impaction losses

The purpose of the DAA is to measure submicrometer aerosol down to a minimum dry size, $D_p$, of about 50 nm. For $D_p$ 50 nm particles, the fraction penetrating the dryer due solely to diffusion losses (from top to bottom, tubing afterwards not included) is larger than 99.7 % penetrating fraction according to theoretical calculations, which is acceptable. We used the equation describing diffusional losses under laminar conditions, i.e. the transport efficiency $\eta_{\text{diff}}$ (Willeke and Baron, 2005):

$$\eta_{\text{diff}} = e^{(-\xi S h)}$$

(1)
where $Sh$ is the Sherwood Number, $\xi = DL/Q$, $D$ is the particle diffusion coefficient, $L$ is the tube length and $Q$ is the flow rate. On the other hand for $D_p$ 3 nm particles, would that be the aim of a study, the fraction penetrating is theoretically only about 87%, where an improvement would require optimal flowrate (towards limit of turbulence) and shortest possible dryer length, for a specified required drying capacity. For particles larger than $D_p$ 50 nm diffusion losses are negligible for many applications, but for supermicron particles impaction and sedimentation losses become important. In our case, cloud droplets are aligned above the dryer to fall vertically into the cross-section between the two membranes, where they dry out (see below) and become residual particles. The residual particle size distribution is measured and is typically in submicrometer-size (depending on measurement site), with low losses as mentioned above. If supermicrometer-sized dry aerosol particles would be a topic of a study, the exit of a dryer and the following tubing used into relevant instrumentation would require careful design (design tactics: straight, vertical and few diameter contractions) see von der Weiden et al. (2009).

### 2.2 Electrostatic losses

The GORE-TEX membrane is made of highly resistive PTFE (teflon) and as such can acquire electrostatic charging locally inside the dryer. Static charging would result in an electric field, which would influence charged aerosol particles with a force, causing them to move. Assuming the charge is situated at a point on the membrane wall, the particle would move sideways, and if arriving at the side of dryer would mostly be deposited there, i.e. this process would result in a lower fraction of aerosol particles penetrating. Collisions with other (neutral, or larger, with lower mobility) aerosol particles would during this transport be negligible, due to the often low total concentrations in the DAA. Furthermore, such radial movements of aerosol particles could cause inhomogenities in the laminar flow exiting the dryer, and care must then be taken while sampling from this flow, as to avoid a streamline with artificially higher or lower concentrations. In order to reduce electrostatic charging and thus to reduce
the resulting electric field in the dryer, grounded metal nets are fitted on both sides facing the sample flow (see above). Furthermore, if the particles are charged, they might themselves create an electric field (see Fuchs, 1989) resulting in particle movements, called electrostatic scattering. Measurements done in laboratory confirmed that no significant (< 5%) losses of accumulation mode sized particles occur for up to 1000 charges on each single particle and relevant concentrations in the dryer, which was also noted by Martinsson (1996).

2.3 Residence time

Residence time needed to dry the largest ambient droplet that contain relevant aerosol for the study in question must be guaranteed. Otherwise the losses, especially for the larger aerosol particles, will increase by impaction and sedimentation. For instance for the DAA system, the inlet can sample up to approximately 25 µm diameter droplets. These droplets should have enough residence time to be dried before entering various tubing and instrumentation after the dryer in order to reduce transport losses. Droplet residuals can however be supermicron, and such large particles are difficult to sample, increasingly with increasing size. Assuming a 25 µm water droplet, the total droplet evaporation time needed can be calculated to 1.09 s at 293 K, see Table 2. The average residence time in the dryer is 5.5 s, and the RH is below 50% more than half that time (see Fig. 2). The drying time necessary increases significantly if the bulk RH surrounding the aerosol particle is higher. Further, at lower temperature, for instance at 273 K the corresponding total drying time required is 3.0 s, approaching the limit of the system. Furthermore the sample flow is laminar, thus the center streamline flow velocity is approximately twice the average, consequently the center streamline has half the average residence time. If more drying time is required, the dryer must be larger, or the flow smaller. We use an additional modular section with the same flange for increased residence time in this case. The two above mentioned requirements of minimum diffusion losses (increases with time) vs. residence time for drying (increasing time is
advantageous) need to be balanced. The two can in many situations be characterised and corrected for, to some extent.

3 Generation of dry air

The dry air came from circulation through aluminium oxide (Al₂O₃) pellets (spherical diameter 2–4 mm) (Martinsson et al., 1992). Each container had 1 kg Al₂O₃. The Al₂O₃ pellets were dried, regenerated, with low-pressure (about 4 hPa) evaporation drying, alternating between two containers (see left side of Fig. 1). A membrane vacuum pump was in constant use to generate the low pressure (Vacuubrand model MV 2NT, Germany). Two 0.002 m³ stainless steel containers were used for each drying flow used (i.e. 4 in total, in two pairs), each pair alternating in either regenerating mode or supplying dry air after the airflow had passed the Al₂O₃ pellets (at ambient pressure). The switching was done with three-way impulse solenoid valves. At the moment the valves switch, initially the low pressure from the container just having been dried is connected to the inlet volume. To avoid a large pressure change, two vacuum vent valves, that open if pressure is less than ca. 500 hPa, were fitted, one on each of the drying airflow circuits (see Fig. 1). The switching of the main dryer valves can be set when different conditions are met, such as: if the aerosol RH is too high, or if the RH in the drying branches are too high. We used as condition that the dryer (both branches simultaneously) should switch if the RH of the outer drying airstream was larger than 17 %, measured at its approximately ambient temperature (in an outdoor metal box) before entering the drier at the lower port. Further we used a minimum time between switching, which was set to 20 000 s (about 5.5 h), and in addition the switch was done only at end of a measurement cycle. During ambient conditions close to 100 % RH the system switched constantly at this time interval. Following a switch the system waited 30 s before next measurement started again, in order to minimise any disturbances.
4 Method

Below the method for particle transmission characterisation is detailed, as well as the method to measure the water vapour removal of the dryer.

4.1 Aerosol particle transmission characterisation

In order to characterise the aerosol transmission through the dryer, measurements were done comparing the aerosol concentration at the exit of the dryer, $C_{\text{exit}}$, with that at the entry, $C_{\text{in}}$. Concentrations were measured with condensation particle counters (CPCs, Model 7610, TSI, USA). The transmission was defined as $C_{\text{exit}}/C_{\text{in}}$. Particles in the range $D_p$ 0.03–4.9 µm were evaluated. The set-up can be seen in Fig. 3.

Firstly a vibrating orifice aerosol generator (VOAG, model 3450, TSI, USA) was used to produce dry (RH 30 %) ammonium sulphate (AS, p.a.) particles, from a solution in MilliQ water, ranging from geometric diameter ($D_{\text{geo}}$) 0.84 to 4.9 µm. The aerodynamic diameter, $D_{\text{aero}}$, was measured with an aerosol particle sizer (APS, TSI model 3321), and was converted to $D_{\text{geo}}$ under assumption of spherical particles according to $D_{\text{geo}} = D_{\text{aero}}/\sqrt{\rho}$, where $\rho$ is particle density (1770 kg m$^{-3}$ for AS). The aerosol was neutralised with a $^{63}$Ni source (case design at Lund University, serial number KF11, source model NB14, 555 MBq, 2008, QSA Global, USA). The particles were fed directly into the dryer, with a tee to a first CPC, a second CPC measured the concentration after the dryer.

Secondly, smaller sized AS particles were produced with a TSI atomizer (TSI 3076 model), ranging from dry mobility diameter 0.03 to 0.4 µm. The aerosol was neutralised, and a monomodal size was selected with a Vienna type DMA (Differential Mobility Analyzer) and then neutralised again. The particles were fed directly into the dryer, with a tee to a first CPC, a second CPC measured the concentration after the dryer.
4.2 Ambient measurements

The dryer was used for continuous long-term measurements at the summit of the mountain Brocken (51.80° N, 10.67° E, 1142 m a.s.l.) in the Harz region in central Germany. The RH at ambient temperature ($T_{amb}$) directly after the exit of the dryer ($RH_{dried}$) was evaluated by using a Rotronic capacity RH sensor. However, this sensor was measuring the sample aerosol RH ($RH_{station}$), inside the measurement station (at its temperature, $T_{station}$ and in a small side-flow from a tee in the main sample flow). This reading was converted to $RH_{dried}$ as follows. The vapour pressure ($p_{H_2O}^{station}$, Pa) in the sample flow is defined as:

$$p_{H_2O} = \frac{RH_i}{100} p_{sat}$$

(2)

where $p_{sat}$ is the saturation vapour pressure of water, calculated from the following empirical formula (Weingartner et al., 1997):

$$p_{sat} = 610.8e^{(\frac{5350}{273.15 - \frac{1}{T_i}})}$$

(3)

with $T_i$ in K and $i$ = station. The above vapour pressure equation was confirmed with the Magnus equation (e.g. Murray, 1967, with a deviation of ≤ 0.1 % in the range 273–293 K. That same vapour pressure is then used in the above equations, rearranged, to calculate the $RH_{dried}$, at $T_{amb}$ (assuming isobaric cooling without condensation).

Furthermore the ambient RH ($RH_{amb}$) was measured (sensor model *GHTU.SGT010*, Greisinger, Germany). The difference between the ambient water vapour flow into the system and the water vapour flow after the dryer, is the water vapour removed from the sample flow (denoted $\dot{m}_{dried}$, kg s$^{-1}$). This was calculated according to:

$$\dot{m}_{dried} = x_{ambient} \dot{m}_{ambient} - x_{station} \dot{m}_{station}$$

(4)

where $\dot{m}_i$ and $x_i$ is the mass flow of dry air (kg s$^{-1}$) and specific humidity (kg kg$^{-1}$) for the ambient sample flow and the sample flow conditioned inside the station,
respectively. $\dot{m}_{\text{ambient}}$ and $\dot{m}_{\text{station}}$ are identical and represent the dry airflow through the system. The density changes up to 10% at occasions between these conditions.

5 Results

Here we present tests from the laboratory, aiming to describe the dryer under controlled conditions, as well as longterm results from the field, showing the overall capacity of the dryer.

5.1 Laboratory measurements

The transmission through the dryer from entry to exit is shown in Fig. 4. An average transmission for accumulation mode particles ($D_p 0.1–1 \mu m$) of 99.9% was observed. The transmission decreases with decreasing particle size below 0.1 µm. The Particle Loss Calculator (von der Weiden et al., 2009) was used to model losses in the dryer (thick line in figure). This software is described in the open access publication, and “employs relevant empirical and theoretical relationships found in established literature”. For supermicron-sized particles from the VOAG the transmission decreases due to two additional 90° bends in the experimental setup. These can be accounted for (modelled with the thin grey line) and indicate the importance of designing a proper exit from the dryer, should large particles be studied.

The uncertainty in transmission as shown in Fig. 4 is composed of counting statistics (Poisson error), flow settings (pressure changes) and dilution (leaks). The Poisson error was ≤ 1% for the accumulation mode sizes measured, the error in flows was estimated to ≤ 3% and dilution ≤ 3%, giving a total error of 4.4%. The uncertainty in transmission increases with increasing particle size, due to mainly lower particle concentrations generated. The uncertainty in size is 3% for the VOAG setup (standard deviation between size deduced from VOAG settings and measured size from APS) and estimated
to be $\leq 10\%$ for the atomizer-DMA setup (from uncertainty in dimensions, flows and high voltage supply for the DMA).

The water vapour transfer of the dryer was tested in the laboratory, geometry of dryer as described above. Humidity was produced by bubbling air through MilliQ-water in an impinger, heated to lab temperature (as it cools due to evaporation). 10.6\% RH at the dryer exit was achieved in the air sample flow with $5.5 \text{ L min}^{-1}$ ($0.33 \text{ m}^3 \text{ h}^{-1}$). The sample air entered with 87\% RH, measured with a capacity RH sensor (Rotronic). In Fig. 2 measured RHs are shown. The lines are a “guide to the eye” of what the profiles are inside the dryer, assuming a logarithmic function (Bierwerth, 2001). The setup is with one membrane outside of the sample flow (with an outside dry closed loop $10 \text{ L min}^{-1}$ flow), and one membrane on the inside of sample flow (with an inner dry flow of $4.7 \text{ L min}^{-1}$), thus in total just below three times as large drying airflow as sample flow. During these laboratory tests the flow ratio of the drying air (outside/inside) was 2.1, which was set too high. The a priori most favorable flow ratio, for a symmetrical drying effect, should be the ratio of the areas of the membranes, which is 1.3. During the operation in the field the flows where 5.0 and 3.7 L min$^{-1}$, outer and inner, and the ratio was 1.5 (the inner flow should have been set higher).

In order to estimate the vapour transfer property of the membrane the water vapour transfer coefficient $k$ was used (Bierwerth, 2001)

$$k = \frac{\dot{m}_{\text{H}_2\text{O}}}{(A_{\text{mem}} \Delta \text{RH}_{\text{log}})} \quad (5)$$

where $\dot{m}_{\text{H}_2\text{O}}$ is the transfer rate of water vapour ($\text{kg s}^{-1}$, using 0.015 kg kg$^{-1}$ H$_2$O in the dry air at $10^5 \text{ Pa}$), $A_{\text{mem}}$ is the membrane surface and $\Delta \text{RH}_{\text{log}}$ is the logarithmic mean difference in RH, defined as

$$\Delta \text{RH}_{\text{log}} = \frac{(\Delta \text{RH}_{\text{Max}} - \Delta \text{RH}_{\text{Min}})}{\ln(\frac{\Delta \text{RH}_{\text{Max}}}{\Delta \text{RH}_{\text{Min}}})} \quad (6)$$

where $\Delta \text{RH}_{\text{Max}}$ and $\Delta \text{RH}_{\text{Min}}$ are the differences of RH at each side of the dryer, maximum and minimum of these, respectively. $\Delta \text{RH}_{\text{log}}$ was 20.6 and 19.9\% RH for the outer
and inner dry airflow, respectively. \( k \) was calculated to \( 4.6 \times 10^{-7} \) kg m\(^{-2}\) s\(^{-1}\) % RH\(^{-1}\). \( k \) can be compared to the manufacturer’s specification and literature values of approximately \( 8–13 \times 10^{-7} \) kg m\(^{-2}\) s\(^{-1}\) % RH\(^{-1}\) (Huang and Qian, 2007), for the membrane, and it is probable that we obtain a significantly lower value due to membrane support structure and metal nets fitted, which reduce the water vapour transfer.

### 5.2 Longterm ambient measurements

The dryer was used for continuous long-term measurements at the summit of the mountain Brocken (51.80° N, 10.67° E, 1142 m a.s.l.) in the Harz region in central Germany from May to October 2010. The project is in collaboration with the Air chemistry group of the Technical University of Brandenburg (BTU Cottbus), who has a cloud measurement site at mount Brocken since many years.

The average RH after the dryer during this time period (the measurements at Brocken) was \( 26.6 \pm 7.3 \) % RH (one standard deviation). The RH\(_{\text{dried}}\) is the quality criterion to ensure a sufficient dry sample flow for the drying of droplets. Furthermore, a second criterion for the highest allowable RH inside instrumentation in the station can be defined to guarantee indoor dryness (in the case of colder indoor temperatures than outdoors, for instance). In our case we considered data with RH\(_{\text{dried}}\) < 30 % RH to be used as is.

The specific humidity can be calculated as:

\[
x_i = \frac{M_{\text{H}_2\text{O}}}{M_{\text{Air}}} \left( \frac{p_{\text{H}_2\text{O}}}{p_{\text{amb}} - p_{\text{H}_2\text{O}}} \right)
\]  

(7)

where \( M_{\text{H}_2\text{O}} \) and \( M_{\text{Air}} \) are the molecular masses of water and air, respectively. \( p_{\text{amb}} \) is the ambient pressure and \( p_{\text{H}_2\text{O}} \) is calculated from the measured RH and temperature (station) with Eqs. (2) and (3). The sample mass flow of dry air relates to the volume flow (continuously monitored in the instrument) as follows:

\[
\dot{m}_{\text{station}} = \rho_{\text{air}} Q_{\text{samplestn}}
\]  

(8)
where \( \rho_{\text{air}} \) is the density of air (at \( P_{\text{amb}} \) and at \( T_{\text{station}} \)) and \( Q_{\text{sample-station}} \) is the dry aerosol sample flow. The above assumes the dried volume sample flow to be similar to the dry airflow, which are within 0.5% up to 293 K and 30% RH. The maximum \( \dot{m}_{\text{dried}} \) achieved was \( 1.2 \times 10^{-6} \text{ kg s}^{-1} \). Furthermore the fog/cloud droplet water is also removed, which is of lesser quantity, and if included would slightly increase the calculated water vapour removed from the system. The fog/cloud droplet entering the dryer can be estimated to rarely exceed \( 0.002 \text{ kg m}^{-3} \), (with an example cloud of 500 cloud droplets \( \text{cm}^{-3} \), with diameter 20 \( \mu \text{m} \)). Generally the specific humidity at the site was in the range \( 0.005–0.02 \text{ kg m}^{-3} \).

During these measurements the RHs of the dry airflows were not measured, thus the logarithmic mean RH was not known, so we could not relate the water vapour transfer to the performance of the membrane measured in the laboratory. The drying capacity, \( \dot{m}_{\text{dried}} \), represents what was achieved in the field and corresponds to drying a 5.7 L min\(^{-1} \) (0.35 m\(^3\) h\(^{-1}\)) aerosol sample flow from 100 to 27% RH at 293 K (with a drying air total flow of 8.7 L min\(^{-1}\)).

The percentage of the time the system operated was 94.2%. Some of the downtime was due to planned maintenance of the system, thus the reliability of the dryer was higher than this. During October the inlet/dryer also automatically shut down a few occasions due to freezing and icing conditions. The time fraction the dryer resulted in a sample RH below 30% was 72.2%, from 9 May until 20 October 2010, when data was available (see Fig. 5). We investigated possible reasons for the spread of RH\(_{\text{dried}}\), such as variations in ambient vapour pressure, \( T_{\text{amb}} \), diurnal variations and longer time series, however no single factor influenced the results. Specifically, the RH after the dryer does virtually not depend on neither ambient nor laboratory temperature. The ambient RH is not a good indicator for performance, because the water removed reduces at colder temperatures. However, the water vapour removed from the sample flow increases linearly with the ambient water vapour pressure, indicating that the system can dry air up to at least the humidities measured during these field tests.
6 Discussion

Although the dryer worked without failures during the field measurement period, at occasions the RH$_{\text{dried}}$ was above 30\%, which should be improved. No single environmental factor has been found to limit the system, rather a combination of factors contribute to time periods with RH$_{\text{dried}}$ above 30\%. Thus a general improvement is required. The two major refinements to do with the drying system presented is firstly to supply a higher dry airflow on the outside of the GORE-TEX membranes. We were presently limited to pumps that were slightly too close to design criteria (regarding flow, durability and ambient cold conditions). That improvement should reduce the sample RH. Secondly the drying circuits (the Al$_2$O$_3$ containers specifically) were also situated outdoors at cold conditions, and as these are regenerated, they cool further. If these circuits were to be situated indoors, or heated, the regeneration process would be more efficient. Indoor placement of the pumps would also be beneficial for durability. However, this would require adjustments depending on ambient and indoor temperature differences, in order to achieve a constant flow. The dry airflow can vary to some extent, without hampering the drying process.

Currently the regulation of the switching of dryers was when the RH in the drying airflow increased above a threshold value, as mentioned above. This would be improved by using RH$_{\text{dried}}$ of the sample at $T_{\text{amb}}$. The threshold for switching could then be set directly at a defined sample RH for instance 25\%. In addition a criterion for the indoor RH in the instrument should be used, which when exceeded should also switch dryers.

7 Conclusions

A dryer constructed for continuous sampling of atmospheric aerosol, where a specified relative humidity of the sample flow (lower than the atmospheric humidity) is required, has been designed and tested. The specific purpose of the dryer presented here is to dry fog/cloud droplets (maximum diameter approximately 25 µm), with minimum losses
of droplets and particles from entrance to exit of the dryer. The accumulation mode particle transmission was measured to be independent of particle size and 99.9 %, corresponding well with theory.

The water vapour transfer coefficient $k$ was measured to $4.6 \times 10^{-7} \text{ kg m}^{-2} \text{ s}^{-1} \% \text{RH}^{-1}$ in the laboratory and is used for design purposes.

The system was tested from 9 May until 20 October 2010, on the mountain Brocken ($51.80^\circ \text{N}, 10.67^\circ \text{E}, 1142 \text{ m a.s.l}$) in the Harz region in central Germany. A maximum water vapour transfer rate of $1.2 \times 10^{-6} \text{ kg s}^{-1}$ was achieved, corresponding to drying from 100 to 27 % RH at 293 K for a $5.7 \text{ L min}^{-1}$ ($0.35 \text{ m}^3 \text{ h}^{-1}$) aerosol sample flow. This can be improved with a closed loop higher dry airflow and warming of the dry air generation system, in order to achieve a more effective dry air regeneration, in our case. The sample flow was below 30 % relative humidity 72 % of that time period. Measurements were done > 94 % during this time period, and the dryer functioned all that time.

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References

**Table 1.** Characteristics of the two tubes used in the DAA dryer, and resulting characteristics for the aerosol flow space in between, at sample flow 5.5 L min\(^{-1}\) (0.33 m\(^3\) h\(^{-1}\)). For that sample flow a dry airflow of at least 5 and 6.5 L min\(^{-1}\) for the inner and outer side is recommended.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Inner tube</th>
<th>Outer tube</th>
<th>Sample air space</th>
</tr>
</thead>
<tbody>
<tr>
<td>GORE-TEX order number (tubes, welded at seam)</td>
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<td>Industrial dry filtration bag 4327 4327T67-0415</td>
<td>–</td>
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<td>Diameter [mm]</td>
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<td>(D_{\text{Inner}}: 67.2)</td>
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<td>Length [mm]</td>
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<td>415 (±3)</td>
<td>405</td>
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<td>Area, effective [m(^2)]</td>
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<td>0.076</td>
<td>0.134</td>
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<td>Outside dry air flow [L min(^{-1})]</td>
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<td>6.5</td>
<td>5.5</td>
</tr>
<tr>
<td>Width [mm]</td>
<td>–</td>
<td>–</td>
<td>12</td>
</tr>
<tr>
<td>Top cross-section [m(^2)]</td>
<td>–</td>
<td>–</td>
<td>0.0011</td>
</tr>
<tr>
<td>Air flow velocity [m s(^{-1})]</td>
<td>0.13</td>
<td>0.12</td>
<td>0.084</td>
</tr>
<tr>
<td>Residence time [s]</td>
<td>3</td>
<td>3.3</td>
<td>5.5</td>
</tr>
<tr>
<td>Reynolds number [–]</td>
<td>71</td>
<td>64</td>
<td>78 (laminar)</td>
</tr>
</tbody>
</table>
Table 2. Example calculation of droplet evaporation time for pure water at lab conditions and 50 % RH (from www.aerosols.wustl.edu/AAARworkshop08/software/AEROCALC-11-3-03.xls, the AeroCalc by Paul Baron).

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature [K]</td>
<td>293.15</td>
</tr>
<tr>
<td>Vapour diffusion coefficient [m² s⁻¹]</td>
<td>0.000024</td>
</tr>
<tr>
<td>Vapour molecular weight [kg mol⁻¹]</td>
<td>0.018</td>
</tr>
<tr>
<td>Saturation ratio [-]</td>
<td>0.5</td>
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<tr>
<td>Initial particle diameter [µm]</td>
<td>25</td>
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<tr>
<td>Particle density [kg m⁻³]</td>
<td>1000</td>
</tr>
<tr>
<td>Saturation vapour pressure [hPa]</td>
<td>23.4</td>
</tr>
<tr>
<td>Temperature at drop surface [K]</td>
<td>286.5</td>
</tr>
<tr>
<td>Vapour press. at drop surface [hPa]</td>
<td>15.4</td>
</tr>
<tr>
<td>Time to droplet evaporation [s]</td>
<td>1.09</td>
</tr>
</tbody>
</table>
Fig. 1. Schematic drawing of the inlet part of the DAA with dryer and dry air regeneration system of the outer drying air (left) and photo of the entry to the dryer only (right). The aerosol sample flows downwards into the three slots, the metal nets are visible on the inside of the slots. The central circular hole is for the exit flow of the inner drying airflow.
Fig. 2. RH-profiles inside the dryer. Points are measurements, lines a guide to the eye, assuming a logarithmic function describing the water vapour removal.
**Fig. 3.** Experimental set-ups for transmission measurements. In (A) the vibrating orifice aerosol generator (VOAG) is shown, aerosol particle sizer (APS), neutralizer (Ni 63). 2 CPCs were used, each with a flow of 1 L min$^{-1}$, similar in (B), showing the setup with an atomizer, for submicron-sized particles. A larger cone was used on top of the dryer, from where the sample flow to CPC 1 was drawn. The flow through the dryer was 5.7 L min$^{-1}$ for both setups.
Fig. 4. Transmission through dryer at 5.7 L min\(^{-1}\) sample flow. Particles were neutralised (i.e. negligible electrostatic scattering). Concentrations tested varied between 1–3500 cm\(^{-3}\). Thick line is model of transmission through dryer. The measurements showing low transmission above 1 µm are due to 90° bends in the experimental setup.
Fig. 5. Histogram of $\text{RH}_{\text{dried}}$, i.e. air sample RH after drying. 72% of the time the RH was below 30%. Approximately 140 days of data.