Laboratory evaluation of the effect of nitric acid uptake on frost point hygrometer performance

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Abstract. Chilled mirror hygrometers (CMH) are widely used to measure water vapour in the troposphere and lower stratosphere from balloon-borne sondes. Systematic discrepancies among in situ water vapour instruments have been observed at low water vapour mixing ratios (<5 ppm) in the upper troposphere and lower stratosphere (UT/LS). Understanding the source of the measurement discrepancies is important for a more accurate and reliable determination of water vapour abundance in this region. We have conducted a laboratory study to investigate the potential interference of gas-phase nitric acid (HNO\textsubscript{3}) with the measurement of frost point temperature, and consequently the water vapour mixing ratio, determined by CMH under conditions representative of operation in the UT/LS. No detectable interference in the measured frost point temperature was found for HNO\textsubscript{3} mixing ratios of up to 4 ppb for exposure times up to 150 min. HNO\textsubscript{3} was observed to co-condense on the mirror frost, with the adsorbed mass increasing linearly with time at constant exposure levels. Over the duration of a typical balloon sonde ascent (90–120 min), the maximum accumulated HNO\textsubscript{3} amounts were comparable to monolayer coverage of the geometric mirror surface area, which corresponds to only a small fraction of the actual frost layer surface area. This small amount of co-condensed HNO\textsubscript{3} is consistent with the observed lack of HNO\textsubscript{3} interference in the frost point measurement because the CMH utilizes significant reductions (>10%) in surface reflectivity by the condensate to determine H\textsubscript{2}O.

1 Introduction

Water vapour is the most important greenhouse gas in the atmosphere. Changes in its atmospheric abundance are a principal feedback mechanism in the radiative forcing of climate (Held and Soden, 2000; Soden and Held, 2006; Trenberth et al., 2007; Dessler et al., 2008). Knowledge of the distribution of water vapour and how it is changing as the earth system changes is especially important in the upper troposphere and lower stratosphere (UT/LS) where water vapour plays critical roles in atmospheric radiative balance, cirrus cloud formation, and photochemical processes (Forster and Shine, 2002; Gettelman et al., 2004; Solomon et al., 2010). Dehydration processes reduce water vapour mixing ratios to part per million (ppm) values in the tropical UT/LS. The microphysics related to dehydration and cirrus cloud particle nucleation and growth are poorly understood at present (Jensen et al., 2005; Peter et al., 2006), limiting the ability to accurately model the dehydration process and, hence, our ability to estimate the influence of climate change on the UT/LS water vapour distribution. Part of this limitation derives from the uncertainties in available water vapour measurements in the 1 to 10 ppm range typical of this region of the atmosphere. Over the past several decades, there has been considerable disagreement (~25–100%) among in situ water vapour measurements by different instruments made at low mixing ratios in the UT/LS. The reasons for the observed differences in measured water vapour in the UT/LS are actively under investigation. A recent, chamber-based intercomparison of the principal in situ UT/LS water vapour instruments (AquaVIT, https://aquavit.icg.kfa-juelich.de/AquaVit/AquaVitWiki) revealed far less disagreement, indicating the differences observed in the atmosphere are likely related to the use of these
instruments on moving platforms or interferences from other atmospheric trace species.

One of the principal instruments currently used to measure atmospheric water vapour is the chilled mirror hygrometer (Oltmans et al., 2000a). A 30-yr record of balloon-borne CMH measurements provides important constraints on the evolving trend of stratospheric water vapour abundances, which have an important role in climate change (Oltmans et al., 2000b; Rosenlof et al., 2001; Solomon et al., 2010; Hurst et al., 2011). Observed decadal differences in the trend analysis of less than 1 ppm require high confidence in the calibration stability and the absence of measurement interferences in the CMH technique. The CMH technique utilizes the principle of condensation of water vapour as a liquid or solid on a chilled mirror. The temperature of the mirror is controlled in order to maintain a constant optical scattering from a thin dew or frost layer in contact with water vapour in an ambient sample flowing continuously past the mirror. An optical detector measures the specular reflectivity of the condensate-covered mirror, while a feedback controller regulates the mirror temperature such that the measured reflectivity of the mirror remains constant. Under this condition, the condensate layer on the mirror is maintained in approximate thermal equilibrium with the water vapour in the air passing over the mirror. The mirror temperature is assumed to be equal to the ambient dew point or frost point temperature. The water vapour partial pressure is calculated from this measurement using a formulation of the Clausius-Clapeyron equation for the saturation pressure over water or ice, and subsequently the water vapour mixing ratio is calculated by dividing by the measured ambient pressure. In this work we focus only on frost as the condensate, which is applicable to measurements at the low temperatures in the UT/LS.

Chilled mirror hygrometry provides an absolute measurement of water vapour with an accuracy that relies only on the calibration of the temperature sensor that measures the temperature of the mirror. This is true, however, only when two conditions are valid: (1) the surface reflectivity is inversely correlated with the amount of condensate, and (2) the saturation vapour pressure of the condensate layer remains that of pure hexagonal ice. A change in the composition of the mirror surface with a microscope camera through a sapphire window mounted directly opposite the mirror. The LED, IR

(Hanson and Mauersberger, 1988; Tolbert and Middlebrook, 1993). In the present study, we report a laboratory evaluation of the interference of HNO$_3$ in CMH frost point measurements under pressure, flow, and HNO$_3$ conditions representative of operation in the UT/LS.

2 Experimental

2.1 Chilled mirror hygrometer

The chilled mirror hygrometers evaluated in this study are a reconfiguration of the Cryogenic Frostpoint Hygrometer (CFH) sonde instrument. The CFH is a small (400 g) balloon-borne instrument that has been used extensively to measure water vapour profiles between the surface and the middle stratosphere at a number of locations around the globe. The stated accuracy in the determination of the frost point temperature is ±0.5 °C (Vömel et al., 2007), which corresponds to approximately ±0.2 ppm under UT/LS conditions, and the overall uncertainty near the tropopause is estimated to be 9%. The CFH units used in this laboratory evaluation (Fig. 1) were copies of the modified version previously employed in the AquaVIT chamber study. The modified housing is made of stainless steel and has a similar geometry to that of the balloon-borne instrument, but is hermetically sealed to allow sample flow at UT/LS pressures (<200 hPa). The optical path of the laboratory instrument has the light emitting diode (LED) source and infrared (IR) photodiode detector arranged at smaller (~40°) angles to the mirror surface. This arrangement allows for imaging of the mirror surface with a microscope camera through a sapphire window mounted directly opposite the mirror. The LED, IR
photodiode detector, and electronics were used unmodified from a standard CFH instrument.

The dimensions of the mirror, mirror stem and heater coil are identical to the balloon instrument as is the location of the mirror thermistor. However, a mirror mount was made of thin-walled (0.127 mm) stainless-steel tube 1.6 cm long, which was glued to the edge of the mirror disk and acted both as a seal and thermal separation from the housing, since the body of the housing remained at room temperature. The low thermal conductivity of the tube assures that the thermal properties of the mirror assembly are similar to those of the balloon instrument. A PCTFE (Kel-F) sleeve was placed around the mirror mount to insulate it from the sample flow. The cold finger, the part of the mirror assembly that is immersed in the cryogen to provide mirror cooling is longer than that of the balloon instrument, since the fittings for connection to the flow system necessitated a larger separation between the cryogen reservoir and the sensor housing. The cooling rate of the mirror, which is largely determined by the dimensions of the thin mirror stem, remained unchanged. All key operational parameters of the laboratory version, including proportional, integral, and differential values of the temperature control loop, are identical to those in the balloon instrument. The laboratory CFH was operated using liquid nitrogen as the cryogen instead of trifluoromethane (CHF$_3$) as used with the balloon version. A Teflon sleeve and foam collar were placed over the cold finger to reduce the heat transfer rate into the cryogen. This insulation largely compensated for the colder temperature of the liquid nitrogen as indicated by mirror heater currents and duty cycles that are comparable to those observed in flight instruments.

2.2 HNO$_3$ measurement

The NO$_y$ instrument used here to measure HNO$_3$ is described in detail elsewhere (Fahey et al., 1989). The NO$_x$ instrument has been used previously in the stratosphere during several field campaigns (e.g., Fahey et al., 1989; Kawa et al., 1992). HNO$_3$ and other NO$_y$ compounds are measured by conversion to NO in a 300 °C gold tube catalyst in the presence of CO (Fahey et al., 1985), followed by detection of the NO using NO/O$_3$ chemiluminescence. The sensitivity of the instrument was determined during each experiment by adding NO upstream of the catalytic converter from a gas standard with a known mixing ratio. Background signals in the chemiluminescence detector were subtracted prior to calculation of HNO$_3$ mixing ratios. The overall uncertainty in the HNO$_3$ measurement is 20%. There is no interference in the NO$_y$ measurement from water vapour mixing ratios in the range used here.

2.3 Experimental setup

The experimental setup used to evaluate the potential interference of gas-phase HNO$_3$ on the performance of chilled mirror hygrometers is shown in Fig. 2. The system consists of two identical hygrometers (described above) connected in series, sources of low mixing ratios of water vapour and HNO$_3$, the NO$_x$ instrument, and the pressure control and pumping unit. The flow system manifold consisted of 1.58 cm inner diameter (ID) stainless steel tube sections connected with ISO-KF16 fittings. A distance of 23.5 cm separated the two CFH mirrors. A mass flow controller (Tylan FC-261) was used to deliver a stable flow of synthetic air (Scott Specialty Gases, Zero Air) through the system. The water vapour content of the zero air was determined to be less than 0.3 ppm for all cylinders used. The total gas flow rate through the manifold in most experiments was ~2000 standard (0 °C, 1013.25 hPa) cm$^3$ min$^{-1}$ (sccm), but experiments were conducted with flows between 1200 and 6000 sccm. Pressures were measured using 100 and 1000 Torr capacitance manometers (MKS, Inc. 626B Baratron). The pressure sensor used to monitor the pressure of the gas flowing past the frost point mirrors was attached to the flow tube about 20 cm downstream of the second CFH instrument. The flow tube ID remained 1.58 cm from the frost point hygrometers to the pressure sensor and no significant pressure gradient is expected under this configuration.

Two methods were used to add water vapour to the manifold sample flow. The first was a zero air flow (20 sccm) past a temperature-controlled water vapour permeation tube (Kin-Tek Laboratories, HRT 010.00). The permeation source provided 1.9 to 5.7 µg min$^{-1}$ of water vapour for temperatures between 60 and 80 °C, respectively. When added to the main sample flow, water vapour mixing ratios were adjustable in the range of 1 to 4 ppm, depending on the sample flow rate. The second method was to add a small flow of
room air into the manifold through a needle valve. This allowed higher mixing ratios of water vapour (up to 20 ppm) in the flow system. Given the water vapour content of lab air (>6000 ppm), this flow was at most a few sccm, ≪1% of the total system flow. During experiments, the value of the water vapour mixing ratio in the flow system was determined using the measured frost point temperature from the upstream CFH instrument.

HNO₃ was added to the sample flow manifold using a 10 sccm flow of zero air past a HNO₃ permeation source (Kin-Tek Laboratories, SRT-002.00-2022/40) operated at either room temperature or 40 °C. HNO₃ permeation sources are known to emit a small amount of NO₂ or other NOₓ species along with the HNO₃. The non-HNO₃ emission was determined by measuring the NOₓ signal after passing the permeation source flow through a plug of nylon wool, which completely scrubs HNO₃ from the flow. The residual (non-HNO₃) signal, which was ~5% of the NOₓ signal without the nylon wool in line, was subtracted from the measured NOₓ signal to determine the HNO₃ mixing ratio during experiments. The HNO₃ addition point was located between the two CFH units in order to expose only the downstream unit (Fig. 2). The sampling point for the NOₓ instrument was located 15 cm downstream of the second CFH mirror. The inner surfaces of the flow tubes were coated from the HNO₃ addition point between the hygrometers to a distance past the NOₓ sampling point with a thin film of FluoroPel (Cytotix Inc., PFC 504A-FS) to minimize HNO₃ adsorption on the walls. HNO₃ transmission through the flow system was determined by comparing the NOₓ signal with HNO₃ added to the flow system with that observed when the HNO₃ flow was added to the NOₓ instrument sample flow immediately in front of the catalyst. After 10–20 min, typical transmission values were ≥85%. HNO₃ addition to the flow system was controlled using a vacuum line connected to the HNO₃ flow line immediately upstream of the manifold. When the valve in this auxiliary line was closed, flow past the HNO₃ permeation tube was added to the manifold flow. When the valve in this auxiliary line was opened, the HNO₃ flow was diverted away from the flow system to the vacuum pump along with a small amount of air from the flow system. Opening and closing this valve resulted in prompt addition or removal of HNO₃ in the flow system.

One difference between a CFH instrument operating in the atmosphere on a balloon sonde and in our laboratory experiments is that in the former the temperatures of the sample flow and housing walls are near ambient as the altitude changes while in the latter they remain at room temperature. This difference is not expected to significantly affect the results of our experiments because it is the mirror surface that controls the temperature of absorbing and desorbing molecules. This result follows from a consideration that the molecular mean free path (<1 µm) is much smaller than the thickness of the laminar-flow thermal boundary layer (~1 mm) over the mirror surface and much smaller than the characteristic mirror dimension (1 cm).

3 Results and discussion

3.1 Frost point interference

HNO₃ exposure tests were conducted first by allowing both hygrometers to achieve a stable frost layer at low water...
vapour conditions (1–5 ppm) in the absence of HNO₃ and determining the NOᵧ instrument background. HNO₃ was then added to the manifold and the behaviour of the two CFH instruments was compared to evaluate the possible interference. A typical experimental run lasting ~4 h is shown in Fig. 3. In this experiment, the permeation source was used to produce a constant background water vapour mixing ratio of 2.3 ± 0.2 ppm in the sample flow as measured consistently by both CFH units. The absolute value determined from the CFH measurements agreed to within 10% of the value calculated from the manufacturer’s stated permeation rate. The higher water vapour values shown in Fig. 3, up to 4.3 and 11 ppm, were produced by periodically opening a valve (shown in Fig. 2) to admit a small flow of air from the room. The two CFH units exhibited slightly different transient responses to the rapid increases and decreases in water vapour, but reported the same value (within 0.3 ppm) upon reaching steady state after a few minutes.

Just before 2 h in the time series in Fig. 3, the HNO₃ diversion valve was closed, thereby exposing the downstream CFH unit, #2, to HNO₃. The HNO₃ mixing ratio in the sample flow rapidly reached 3.5 ppb and then continued to increase gradually over the next 2 h. The long rise time for HNO₃ is caused by the strong adsorption of HNO₃ to all exposed surfaces, particularly those not covered by the Fluoropel coating. The variations observed in the HNO₃ mixing ratio are due to slight changes in HNO₃ transmission through the flow system that arise from changes in surface wetness and mirror temperature, and mirror frost evaporation and condensation related to rapid changes in water vapour. The frost point temperature reported by CFH#2 never deviated significantly from that of CFH#1, even after a 2.5-h exposure of up to 4 ppb of HNO₃. Furthermore, the transient and steady state responses of CFH#2 to the periods of increased water vapour were unchanged following HNO₃ exposure. A typical CFH balloon flight to 30 km altitude typically requires less than 2 h and HNO₃ is typically <0.5 ppb in the middle and upper troposphere and gradually increases to 4 ppb within a few kilometres above the tropopause (Popp et al., 2009). Thus, based on several experimental runs similar to that shown in Fig. 3, over a range of UT/LS conditions, there is no interference (<0.1 °C frost point change) due to ambient HNO₃ in CMH balloon sonde measurements.

3.2 HNO₃ surface coverage

HNO₃ is known to deposit readily to ice surfaces at temperatures similar to those of the CFH mirrors in these experiments in both the laboratory (Abbatt, 2003) and atmosphere (Popp et al., 2004; Voigt et al., 2006). The lack of any observed interference of HNO₃ on the measured frost point temperature raises the question of the amount of HNO₃ that accumulates on the frost during exposure. By rapidly heating the CFH mirror and vaporizing the frost layer after a certain HNO₃ exposure time, any adsorbed or co-condensed HNO₃ on the mirror is released into the sample flow and detected by the NOᵧ instrument. Separate tests showed that mirror heating did not desorb appreciable HNO₃ from surfaces of the insulated mirror mount (Fig. 1). A similar rapid mirror evaporation is typically programmed to occur during balloon ascent of the CFH instrument to ensure the condensate layer is frost and not supercooled liquid (Vömel et al., 2007). The resulting pulses of HNO₃ measured by the NOᵧ instrument can be clearly seen in the plot in Fig. 4. Examination of the NOᵧ trace in Fig. 4 reveals that the magnitude of the desorbed HNO₃ pulses is a function of the exposure time. The mass of adsorbed HNO₃ was determined by integrating the pulse measured by the NOᵧ instrument.
**Fig. 4.** Time series of CFH mirror temperatures and HNO$_3$ mixing ratios. The stable mirror temperature corresponds to a water vapour mixing ratio of 2.5 ppm. The pulses visible in the CFH#2 mirror temperature are rapid increases used to evaporate the frost layer. Corresponding pulses were observed in HNO$_3$ as measured by the NO$_y$ instrument when the frost layer had been exposed to HNO$_3$. The negative excursions in the CFH#2 temperature occur when the frost layer is being reformed following removal.

**Fig. 5.** Mass of HNO$_3$ measured during mirror frost layer evaporation events plotted against the frost layer exposure to HNO$_3$ expressed as the time integrated mixing ratio. The legend shows the experimental conditions of pressure, sample flow ($Q$), and water vapour and HNO$_3$ mixing ratios, along with the derived growth rate from the best fit line with a zero intercept.

Figure 5 shows the measured mass of desorbed HNO$_3$ plotted against the exposure expressed as the product of the HNO$_3$ mixing ratio and exposure time. HNO$_3$ accumulated approximately linearly with exposure at mixing ratios of up to 4 ppb over time periods up to 3 h. Laboratory studies have shown HNO$_3$ monolayer densities to be approximately $1–2 \times 10^{14}$ molecules cm$^{-2}$ (Abbatt, 2003), so the highest observed HNO$_3$ mass loadings ($\sim 8$ ng) in our experiments represent roughly monolayer coverage of the geometric surface area of the mirror ($0.37$ cm$^2$). There is no information on the specific surface area (cm$^2$ cm$^{-2}$) of the frost, but based on electron micrographs recently published by Leu and Keyser (2009) of frost layers formed by water vapour deposition at similar temperatures, it is likely to be $\gg 1$. This implies that only a small fraction of the surface area of the frost is affected by HNO$_3$ adsorption and, therefore, no significant interference in the CFH measurement would be expected.

### 3.3 Thermodynamic considerations

H$_2$O adsorption and desorption on the frost layer are close to steady state when the water vapour concentration is constant above the mirror surface and the CFH mirror temperature is at a nearly constant value; under these conditions the frost layer is neither growing nor shrinking significantly. In contrast, HNO$_3$ adsorption and desorption are clearly not
in steady state, as evidenced by the time dependence of the HNO$_3$ mass loading shown in Fig. 5. The state of the adsorbed HNO$_3$ molecules is not known for the frost layer in our experiments. However, the mirror temperature and H$_2$O and HNO$_3$ partial pressures present in these experiments fall thermodynamically within the region of stable NAT formation (Hanson and Mauersberger, 1988; Tolbert and Middlebrook, 1990; Worsnop et al., 1993; Zondlo et al., 1997). The estimated small fractional HNO$_3$ coverage of the frost surface area suggests that while NAT may exist locally on the frost surface or within the frost layer, the total HNO$_3$/H$_2$O molar ratio on the mirror is much less than 0.33, which would be required for complete conversion of the frost to NAT. The HNO$_3$ mixing ratios in our experiments represent a large supersaturation with respect to NAT at the mirror frost point temperatures (e.g. the ratio of the partial pressure of HNO$_3$ to the saturation vapour pressure over NAT is \(\sim 60\) for the conditions in the experiments shown in Fig. 5). The observed linear increase in HNO$_3$ adsorbed mass in the first hours of exposure indicates that the molar ratio may continue to increase with time until it reaches 0.33 in the frost, or perhaps in a layer at the frost surface, which would bring HNO$_3$ adsorption and desorption into steady state. If the NAT formation on the surface of the frost were to inhibit further formation of water ice, the CFH would then attempt to control at the NAT saturation temperature, several degrees higher than the frost point temperature under pure H$_2$O conditions (Hanson and Mauersberger, 1988; Tolbert and Middlebrook, 1990; Worsnop et al., 1993) and the mirror temperature would then lead to an overestimate of the water vapour partial pressure. The ppb-min exposure of HNO$_3$ required to reach NAT stoichiometry in the mirror frost appears to be much larger than that realized in this laboratory evaluation and, hence, is much larger than is relevant for CFH balloon-borne measurements. The interpretation of the interference in the reading of a commercial frost point instrument observed in the AIDA chamber was that HNO$_3$ had accumulated in sufficient amounts to convert the frost to NAT (Szakáll et al., 2001). The results presented here cannot confirm or disprove their interpretation since their experiment was conducted with much higher HNO$_3$ levels.

4 Conclusions

Two CFH frost point hygrometers and an NO$_y$ instrument were used to investigate the potential interference of gas-phase nitric acid (HNO$_3$) at atmospherically-relevant mixing ratios in measurement of low ppm water vapour mixing ratios by chilled mirror hygrometers under pressure and flow conditions representative of operation in the UT/LS. No detectable change (\(< \pm 0.1 ^\circ\)C) in the measured CFH frost point temperature was found with additions of up to 4 ppb HNO$_3$ for periods exceeding 2 h. HNO$_3$ that had adsorbed or co-condensed on the mirror with H$_2$O was detected during rapid frost layer evaporation using the NO$_y$ instrument. Our measured HNO$_3$ loadings of the CFH mirror surface indicate that over a typical balloon-borne CFH ascent period (90–120 min), condensed HNO$_3$ mass increased linearly at a rate of 0.033 ng min$^{-1}$ ppb$^{-1}$. The observed mass loading of \(\sim 8\) ng following 120 min exposure at 2 ppb HNO$_3$ corresponds to approximately monolayer coverage of the geometric mirror surface area, but likely much less than monolayer coverage of the frost surface area. We conclude that for typical atmospheric conditions, it is unlikely that HNO$_3$ alone could cause a significant error in water vapour measured by a balloon-sonde chilled-mirror hygrometer such as the CFH. The potential for HNO$_3$ to interfere with CMH water vapour measurement during extended sampling in the lower stratosphere from aircraft or other balloon platforms is likely small, but would require further study to critically assess.

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