On the consistency of HNO$_3$ and NO$_2$ in the Aleutian High region from the Nimbus 7 LIMS Version 6 data set

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Abstract

This study uses photochemical calculations along kinematic trajectories in conjunction with Limb Infrared Monitor of the Stratosphere (LIMS) observations to examine the changes in HNO$_3$ and NO$_2$ near 30 hPa in the region of the Aleutian High (AH) during the minor warming event of January 1979. An earlier analysis of Version 5 (V5) LIMS data indicated increases in HNO$_3$ without a corresponding decrease in NO$_2$ in that region and a quasi-wave 2 signature in the zonal distribution of HNO$_3$, unlike the wave 1 signal in ozone and other tracers. Version 6 (V6) LIMS also shows an increase of HNO$_3$ in that region, but NO$_2$ is smaller than from V5. The focus here is to convey that V6 HNO$_3$ and NO$_2$ are of good quality, as shown by a re-examination of their mutual changes in the AH region. Photochemical model calculations initialized with LIMS V6 data show increases of about 2 ppbv in HNO$_3$ over 10 days along trajectories terminating in the AH region on 28 January. Those increases are mainly a result of the nighttime heterogeneous conversion of N$_2$O$_5$ on background stratospheric sulfuric acid aerosols. Changes in the composition of the air parcels depend on the extent of exposure to sunlight and, hence, on the dynamically controlled history of the trajectories. Trajectories that begin in low latitudes and traverse to across the North Pole in a short time lead to the low HNO$_3$ in the region separating the anticyclone from the polar vortex, both of which contain higher HNO$_3$. These findings help to explain the observed seasonal evolution and areal extent of both species. V6 HNO$_3$ and NO$_2$ are suitable, within their errors, for the validation of stratospheric chemistry–climate models.

1 Introduction and objectives

The Limb Infrared Monitor of the Stratosphere (LIMS) experiment operated on Nimbus 7 from 25 October 1978 through 28 May 1979. LIMS measurements were originally processed and archived to a Version 5 (V5) data set (see Gille and Russell, 1984, and references therein). Since then, the observed LIMS radiance profiles have been re-processed with improved, Version 6 (V6) algorithms to provide profiles of temperature, chemical...
species, and geopotential height (GPH), as a function of pressure–altitude from 65°S to 84°N latitude (Remsberg et al., 2004). There are improvements from the registration of the LIMS radiance profiles and from updated spectroscopic line parameters for retrievals of the V6 species profiles. Several studies already show that the V6 ozone is of better quality for scientific analysis (Natarajan et al., 2002; Stolarski et al., 2013; Remsberg et al., 2013; Shepherd et al., 2014). The two nitrogen species, nitric acid (HNO₃) and nitrogen dioxide (NO₂), are also of better quality, particularly NO₂ (Remsberg et al., 2010, 1994). Holt et al. (2012) quantified the exchange of V6 NO₂ from the mesosphere to the middle stratosphere within the polar vortex. Remsberg and Harvey (2016) also found good relationships on the 550 K potential temperature surface (near 30 hPa) for the highest values of V6 HNO₃, lowest values of ozone, and highest values of potential vorticity (PV) within the Arctic winter vortex. While there are residual effects from polar stratospheric clouds (PSCs) in the ozone and water vapor (H₂O), those effects are small in the HNO₃ and NO₂ profiles and occur only at temperatures < 194 K and from about 1 to 20 January. The V6 data set is part of the SPARC Data Initiative for chemistry–climate model comparison studies (Tegtmeier, et al., 2013; SPARC, 2017).

The northern hemispheric polar winter of 1978–1979 was dynamically active in the middle stratosphere, as determined from daily surface plots of GPH, PV, and ozone on the 850 K potential temperature surface (near 10 hPa) (McIntyre and Palmer, 1983; Leovy et al., 1985; Butchart and Remsberg, 1986). A major zonal wave 1 forcing brought about a rapid exchange of air between polar and middle latitudes from mid- to late January. Rood et al. (1993) (hereafter referred to as R93) analyzed the LIMS V5 data in the Aleutian High (AH) region for 14–27 January. They reported that the tracer-like species HNO₃ on the 30 hPa surface increased slowly during that time in the region of the relatively warm anticyclone. The absence in their analysis of a corresponding decrease in NO₂ accompanying the increase in HNO₃ led to concerns of an inconsistency in the LIMS data. The present study reconsiders that anomaly but using the V6 data set and a trajectory model that includes the chemistry of reactive nitrogen (NO₃), or the sum of NO, NO₂, 2N₂O₅, NO₃, HNO₃, HONO₂, and ClONO₂.

In their analysis of the V5 data, R93 also noted finding high values of HNO₃ in both the polar vortex and the AH and lower values around the periphery of both circulation systems, resulting in a quasi-wave 2 signature in the HNO₃ distribution. They suggested that, since dynamically controlled species like ozone showed a clear wave 1 signature similar to GPH and PV, the HNO₃ levels outside the polar vortex may have influences from a combination of chemistry and dynamics in order to account for its quasi-wave 2 variation. They explored the possible role of heterogeneous chemistry on background sulfate aerosols in converting NO₃ to HNO₃ and concluded based on a 2-D model study that those species changes from V5 are large in winter but not correct phenomenologically. Separate studies have shown that heterogeneous reactions involving sulfate aerosols do perturb the stratospheric photochemistry and HNO₃ levels (Austin et al., 1986; Rodriguez et al., 1991). With the availability of the improved V6 data set, we revisit the question of inconsistency for the LIMS HNO₃ and NO₂ observations in the AH region. We focus our attention on the same period, namely 14–27 January, when there was a minor stratospheric warming.
Section 2 describes briefly the improvements implemented in the retrievals of the V6 profiles. We also note that the residual effects from unscreened PSCs in the polar vortex are small for HNO$_3$ relative to those in ozone. Section 3 contains polar plots of ozone, HNO$_3$, NO$_2$, and GPH for 17 and 27 January 1979 and describes their changes, as the AH develops and the center of the vortex moves off the North Pole. Section 4 shows the changes in V6 HNO$_3$ and NO$_2$ at the center of the AH from 14 to 27 January. Section 5 describes an ensemble of trajectory calculations, including heterogeneous chemistry, for air parcels converging in the region of the AH on 27 January. Those calculations partition NO$_3$ into the observed HNO$_3$ and NO$_2$, plus the unobserved variations of nitric oxide (NO) and nitrogen pentoxide (N$_2$O$_5$). The calculated daily species values compare reasonably well with closest observed LIMS values, as shown from the results in Sect. 6. We then show in Sect. 7 time series of variations of PV and of V6 HNO$_3$, H$_2$O, and NO$_2$ on the 550 K potential temperature surface across most of the Northern Hemisphere for the entire 7.25 months of LIMS data. In this way, the variations for January are set in the context of the longer-term seasonal changes for those species. Section 8 is a summary of the findings.

2 Improvements of the LIMS V6 species

The LIMS instrument obtained profiles of atmospheric limb radiance in six channels, a wide (W) and a narrow (N) band channel for CO$_2$ (CO$_2$ W and CO$_2$ N) and one each for ozone, H$_2$O, HNO$_3$, and NO$_2$ (Gille and Russell, 1984). Retrieved V6 profiles occur at a spacing of every 1.4° of latitude (~155 km) along the orbit, although their horizontal, tangent-path resolution remains no better than about 320 km. Bandpass filters (in cm$^{-1}$) for the LIMS instrument are CO$_2$ W (579–755), CO$_2$ N (637–673), ozone (926–1141), H$_2$O (1370–1560), HNO$_3$ (844–917), and NO$_2$ (1560–1630) in terms of their 5% relative response points. Both the H$_2$O and NO$_2$ channels have an instantaneous field-of-view vertical width at the horizon of 3.6 km, while the other four channels have half that width or 1.8 km.

Retrievals of the V6 temperature and associated species profiles were obtained by using all successive, up–down scan profile pairs along their observed orbital tangent-path locations and at 18 levels per decade of pressure–altitude, $p(z)$, or spaced about every 0.88 km. The effective vertical resolution is the same (~3.7 km) for the retrieved V6 temperature and for each of the species profiles (Remsberg et al., 2004), such that one can evaluate better the combined changes of HNO$_3$ and NO$_2$ at a given pressure level. Further, the spectral line parameters used for the retrieval of the NO$_2$ were updated for the production of V6, leading to values of nighttime NO$_2$ that are up to 20% smaller than those of V5 in the upper and middle stratosphere (e.g., Remsberg et al., 1994). An important addition to the V6 data set is the co-located GPH for each of the retrieved profiles.

LIMS-retrieved ozone has a nonlinear sensitivity to temperature and/or radiance biases and to the effects of PSCs; retrieved H$_2$O mixing ratio is even more sensitive (Remsberg et al., 2007, 2009). However, effects from PSCs are much less noticeable in HNO$_3$ and NO$_2$. As an example, Fig. 1 shows the relative effects for ozone and HNO$_3$ of the residual contamination or the unscreened effects of emission from PSCs, plus the associated temperature and GPH on the 31.6 hPa surface for 11 January. Grid-point data for the surface plots of Fig. 1 are from the V6 Level 3 sequential estimation (SE) algorithm product (Remsberg and
Lingenfelser, 2010). The ozone panel of Fig. 1 shows values in the cold vortex region that are of the order of 6 ppmv and not in keeping with the much lower surrounding values of 3 ppmv. White plus signs indicate where there was a screening of profile segments perturbed by emissions from PSCs in ozone, and the red dot indicates the presence of a PSC based on data from the Stratospheric Aerosol Monitor (SAM II) experiment for comparison purposes. While there can be descent of ozone in the vortex from higher altitudes, the excess ozone in the vortex region of Fig. 1 is due to the remaining, unscreened effects of the PSCs. Note also that the PSC features occur only where the environmental temperature is less than about 194 K. Thus, effects of PSCs are minimal for most of the Arctic lower stratosphere, and they are not present at all in the warmer AH region.

The lower right panel of Fig. 1 shows that observed HNO$_3$ is highest in the vortex and particularly in the presence of PSCs. As HNO$_3$ adsorbs onto PSCs, a decrease of gas-phase HNO$_3$ is also expected in the presence of and directly downwind of large PSCs (von Koenig et al., 2002; Lambert et al., 2012). However, those high LIMS values are nearly unaffected by PSCs because the relationship between the LIMS HNO$_3$ channel radiance and its retrieved mixing ratio is essentially linear. In fact, simulation studies indicate that a temperature bias error of 1 K has only a small, 3 % effect in the V6 HNO$_3$ mixing ratios from 10 to 50 hPa (Remsberg et al., 2010, their Table 1). Retrieved NO$_2$ also varies in a nearly linear way. Yet as with ozone, there are no perturbing effects from PSCs for H$_2$O, HNO$_3$, or NO$_2$ in the AH region that is the focus of the remainder of this study.

### 3 LIMS V6 GPH, O$_3$, HNO$_3$, and NO$_2$ during January 1979

R93 (and references therein) analyzed and presented results of GPH, ozone, and HNO$_3$ at 30 hPa from the V5 data set for 14, 17, 23, and 27 January. They noted that ozone behaves as a tracer in terms of its relation to changes in GPH and according to its associated transport fields. In contrast, they found that the HNO$_3$ distributions did not evolve in the same way, but developed a quasi-wave 2 rather than wave 1 character over a deep layer of the middle stratosphere (50 to 5 hPa). They also did not find clear anticorrelations between the distributions of V5 HNO$_3$ and NO$_2$ in the more isolated AH region, at least to the extent that the sum of those two gases do not change appreciably. We consider their variations again using the V6 data.

Initially, Rood et al. (1987) expressed some doubt about the accuracy of the Arctic GPH fields at 30 hPa from the operational meteorological analyses for January 1979. Figures 2b and 3b are polar plots of the zonal GPH anomalies for the equivalent level of 31.6 hPa and for two of the four days above, 17 and 27 January. Those GPH anomalies exhibit structures that are very similar to those found by R93 (their Fig. 1b and d). Large-scale flow is along isolines of the GPH anomalies or around the respective high and low centers. The AH strengthened significantly but remained at about the same location from 17 to 27 January.

Anomalies for the LIMS species in Figs. 2 and 3 are a result of subtracting the V6 zonal mean coefficient (or mixing ratio) from the observed values at each latitude–longitude grid point. However, the so-called “zonal anomalies” for ascending (local 13:00) and descending (local 23:00) NO$_2$ represent the result of subtracting the average of the ascending or
descending values for a latitude from the corresponding, observed values of NO\textsubscript{2} at each longitude. In other words, the separate analyses of the ascending or descending NO\textsubscript{2} in the LIMS Level 3 product do not account for likely diurnal variations in the NO\textsubscript{2}. Maximum ozone anomalies in Figs. 2a and 3a are nearly coincident with the center of the AH, again in close agreement with the findings of R93 (their Fig. 2b and d). There is strong meridional transport of air of relatively poor ozone along the western edge of the AH to across the North Pole from 17 to 27 January.

Large-scale, zonal anomalies in HNO\textsubscript{3} (Fig. 2d and d) are opposite in sign to those of GPH and ozone. One exception is the relatively low values of HNO\textsubscript{3} near 80° N, 0° E on 17 January in Fig. 2d, which has been interpreted as due to an uptake of gas-phase HNO\textsubscript{3} onto PSC particles just upwind a day or so earlier (Remsberg and Harvey, 2016). However, by 27 January (Fig. 3d) the cyclonic circulation about the low GPH center indicates clearly that there must be net transport of low values of HNO\textsubscript{3} near 140° E, along the axis of the polar night jet, across the North Pole, and then to about 300° E. As a result, while both ozone and NO\textsubscript{2} display the same wave 1 structures as the GPH field on 27 January, the HNO\textsubscript{3} distribution exhibits quasi-wave 2 structure. Figures 2c and 3c represent NO\textsubscript{2} anomalies from profiles of just the LIMS V6 descending orbits or from its nighttime values near 23:00 local time (LT). Those anomalies for NO\textsubscript{2} are from four zonal waves minus the zonal mean coefficient, rather than from six zonal waves as for all the other LIMS Level 3 products.

Most of the NO\textsubscript{x} converts from NO to NO\textsubscript{2} at sunset, followed by a further, partial conversion of the NO\textsubscript{2} to N\textsubscript{2}O\textsubscript{5} up to the 23:00 observation time of LIMS (Brasseur and Solomon, 2005). The respective panels of Figs. 2 and 3 demonstrate that ozone and NO\textsubscript{2} have large-scale features of opposite sign in the AH region, while HNO\textsubscript{3} and NO\textsubscript{2} have anomaly patterns of the same sign on 17 January but not on 27 January. Clearly, it is important to consider the amount of NO\textsubscript{x} that is in the form N\textsubscript{2}O\textsubscript{5} in the AH region during that time span.

Figure 4 shows the zonal species variations from the V6 SE or Fourier coefficients at 31.6 hPa for 27 January at the latitude of 64° N and as calculated at grid points spaced every 5.625° of longitude. The LIMS Fourier coefficients for each of the species are the result of applying the SE algorithm to their Level 2 or profile data, as limited by the estimated data precisions. As a result, the V6 species variations (black curves) are relatively accurate and significant, at least to the scale of zonal wave 6 (or to wave 4 for NO\textsubscript{2}). Vertical error bars in Fig. 4 are the 2\sigma values from the error covariance matrix of the vector of the V6 Fourier coefficients (Remsberg and Lingenfelsler, 2010). Variations from the V5 data set at 30 hPa are included for comparison purposes (red curves).

HNO\textsubscript{3} within the polar vortex (0 to 90° E) is higher by nearly 2 ppbv compared to that in the AH region (180 to 240° E). Note from Fig. 3, however, that HNO\textsubscript{3} has a strong, positive Equator-to-North-Pole gradient in the 0 to 90° E sector, whereas that of ozone is weak and slightly negative. Ozone in Fig. 4 has a broad, wave 1 character, while HNO\textsubscript{3} exhibits two minima (at 130 and 290° E). At 10 hPa and lower pressures (or at higher z) a quasi-wave 2 structure is no longer apparent in HNO\textsubscript{3}. Variations of the descending and ascending (daytime or ~ 13:30 LT) NO\textsubscript{2} modes appear separately in Fig. 4. Diurnal differences for NO\textsubscript{2} are seen only in the longitude sector from 320 to 120° E or across the boundary of the cold
polar vortex and the warmer AH region. Relatively low ozone in this region results in only a slow conversion of NO\textsubscript{2} at sunset to NO\textsubscript{3} and finally to N\textsubscript{2}O\textsubscript{5}. Consequently, a larger fraction of NO\textsubscript{x} remains as NO\textsubscript{2} at the time of the LIMS descending mode observations, or ~23:00 LT. The warmer AH region from 150 to 270° E, in contrast, has higher ozone mixing ratio, and the decrease in NO\textsubscript{2} from its maximum at sunset is steeper, bringing NO\textsubscript{2} mixing ratios at the time of the LIMS descending mode observation close to the value of the ascending mode measurement. There is very little diurnal difference in NO\textsubscript{2} in that longitude region. Thus, air parcel history is important for interpreting observed changes in the distributions of these two reactive species even in the AH region. V5 NO\textsubscript{2} varies in a similar way, although its values are nearly twice those of V6.

4 Changing composition within the Aleutian High

Species variations with time are shown in Fig. 5, as determined from the V6 orbital or Level 2 profiles at 31.6 hPa and nearest to the center of the AH on each day and identified by the location of the maximum GPH anomaly. Note that Fig. 5, which is analogous to Fig. 7 of R93, also shows that the latitude of the AH moves poleward from about 60 to 68° N from 14 to 27 January. Its longitude moves slightly, too, from 186° E on 14 January to 230° E on 19 January and then retreats partially to 214° E by 27 January. Figure 6 illustrates those locations better as red boxes for each day. GPH of the 31.6 hPa level at the center of AH grows from 23.0 to 23.8 km during that same time.

Vertical bars in Fig. 5 represent root-sum-squared (RSS), single profile errors for 30 hPa (Remsberg et al., 2007, 2009), and the ozone and H\textsubscript{2}O variations are no greater than those error estimates of 15 %. Those two species remain rather steady in the AH and are in keeping with their small horizontal gradients on the 31.6 hPa surface and their relatively long chemical lifetimes. HNO\textsubscript{3} shows significant changes; it declines from 10 to 8.2 ppbv from 14 to 18 January, but then increases rather steadily again to 10 ppbv by 27 January. Single profile RSS uncertainty for HNO\textsubscript{3} is ~ 8 % at 30 hPa (Remsberg et al., 2010). Descending (or late evening) NO\textsubscript{2} declines from 1.1 to 0.5 ppbv from 15 to 19 January, increases to about 1.0 ppbv on 23 January, before declining again to 0.4 ppbv on 27 January. Ascending (early afternoon) NO\textsubscript{2} is smaller than descending NO\textsubscript{2} from 14 to 17 January, or when the AH center is at about 60° N. As the AH central latitude shifts northward after 17 January, both the ascending and descending NO\textsubscript{2} exhibit similar values and indicate that the daytime observations are from near to or within the polar night boundary. Generally, the RSS error for single NO\textsubscript{2} profiles is ~ 30 % at this pressure level, although profile registration uncertainties can lead to larger errors across the polar night boundary. Remsberg et al. (2010, their Fig. 3) showed an altitude–latitude plot of “zonal average NO\textsubscript{2}” from only the descending (23:00) orbital segments of 15 January, when the polar vortex is still nearly circular and centered on the North Pole. Its vertical distribution at 60° N declines from 2.5 ppbv at 20 hPa to a local minimum of < 1 ppbv at 35 hPa or about where the associated HNO\textsubscript{3} has its local maximum. In fact, the V6 retrieval algorithm sets NO\textsubscript{2} to zero, when the forward radiance for the tangent layer approaches the measurement noise for that channel (5.5 \times 10^{-4} \text{ w m}^{-2} \text{ sr}^{-1}). Thus, the effect of a finite (~ 3.7 km) vertical resolution is to smooth across that local NO\textsubscript{2} minimum, giving a high bias in its final retrieved value.
5 Trajectory model with photochemistry

In this section, we examine the interplay between photochemistry and dynamics during the evolution of HNO$_3$ and NO$_2$ in the AH by making use of photochemical calculations along trajectories that terminate in the AH region on 27 January. In these calculations, we use the V6 level 2 profile data to initialize the air parcel composition, which allows a comparison between model results and co-located observations along the trajectory in the AH region. The trajectory model is driven by 3-D meteorological data from MERRA (Rienecker et al., 2011) corresponding to January 1979. This data set includes 3-hourly information on surface pressure, horizontal wind, vertical pressure velocity, and temperature on a 1.25° longitude by 1.25° latitude grid. A family of 70 back trajectories is generated, with trajectories beginning at 30 hPa from a grid of 2° latitude by longitude covering a domain defined by 210 and 218° E longitudes and 60 and 86° N latitudes. The starting time of the back trajectories is 09:00 GMT on 28 January, which corresponds to a local time of 23:00 on 27 January at 210° E. This is close to the local time of LIMS descending mode observations in this latitude region. The selected region overlaps the AH in the contour plot of V6 GPH anomalies for this day, as shown in Fig. 3b. The model uses a fourth-order Runge–Kutta advection scheme to generate 3-D kinematic back trajectories. We save trajectory parameters required for further calculations on an hourly basis.

Figure 6 shows the back trajectory beginning on 27 January at 214° E, 68° N and 30 hPa. The numbers in black along the trajectory represent the day numbers. Red colored squares with day numbers represent the history of the location of the AH center at 31.6 hPa based on the maximum V6 GPH anomaly. This V6 pressure level is the closest Level 2 value to 30 hPa. It is clear that between 22 and 27 January the trajectory and the AH center remain in a region north of 54° N latitude, and the AH region provides an isolated natural chemical laboratory for the constituents to evolve. Prior to 22 January the trajectory and AH center diverge with the trajectory going backwards to latitudes as far south as 27° N. As we will show later, the back trajectory beginning at 214° E and 60° N remains within the AH region or north of 45° N throughout the 10-day period. These differences among the trajectories affect their initial values and the changes in air parcel composition.

We conducted time-dependent photochemical calculations along the trajectories in the forward direction. Information used in these calculations include the vertical ozone column along the trajectories based on V6 Level 3 ozone data, background sulfate aerosol surface area densities for January 1979 adopted from the IGAC/SPARC CCMI recommendations for Reference Simulation 1 (Eyring et al., 2013), and solar zenith angle corresponding to the local time of day. Aerosol surface area in the model is $4 \times 10^{-9}$ cm$^{-1}$ at 31 hPa and 60° N for January (see Table 1 of Hofmann and Solomon, 1989). The starting location and mixing ratios of measured species are determined by identifying for each trajectory the spatially and temporally closest LIMS descending mode observation between 14 and 17 January. Longitude separation between the trajectory and the V6 data is within 15°, latitude separation within 7.5°, and time of measurement within an hour. We use an updated version of the stratospheric diurnal photochemical model (Natarajan and Callis, 1997), incorporating the chemical kinetics and photochemical data from the recent JPL evaluation (Burkholder et al., 2015), to calculate the changes in the composition of the air parcels until they reach the
AH region on 27 January. Results from a time-dependent, 2-D chemistry–transport model (Callis et al., 1997) simulation corresponding to January 1979 provide initialization estimates of other unmeasured HO\(_x\), Cl\(_x\), and NO\(_x\) species. The initialization procedure involves repeated diurnal calculations at the fixed starting latitude, altitude, and day. During each diurnal cycle, the mixing ratios of ozone, NO\(_2\), and HNO\(_3\) are set to the observed values at the local time of the LIMS descending mode measurement (Natarajan et al., 2002). Within five diurnal cycles, the short-lived chemical species reach near steady mixing ratios. Then, the photochemical model integration continues along each trajectory until reaching the endpoint of 09:00 GMT on 28 January.

6 Results and discussion

We show in this section the results of sample trajectories that behave similar to those of the ensemble and discuss the transformations occurring in the composition of the air parcels over their 10-day transit. All trajectories terminate at 30 hPa but, since we use 3-D kinematic trajectories based on MERRA, there are fluctuations in pressure and altitude as the parcel moves along its trajectory. The length of exposure to daylight also fluctuates and is not the same for all the trajectories. These differences certainly affect the photochemical changes that occur.

Figure 7a shows, on a polar stereographic projection map of the Northern Hemisphere, three different trajectories terminating at a longitude of 214° E. The latitude circles in the figure are 10° apart poleward of 20° N. The uppercase letters A, B, and C denote the starting location of the trajectories and the lowercase letters a, b, and c mark the endpoints at 60, 72, and 80° N latitude, respectively. The color scale ranging from 0 to 1 represents the accumulated hours of exposure to darkness along each trajectory expressed as a fraction of the total length of the trajectory in hours. The cumulative fraction of darkness at the endpoint is 0.70, 0.67, and 0.63 for the trajectories A–a, B–b, and C–c, respectively. Figure 7b shows the same three trajectories but now colorcoded to demonstrate the calculated variation of HNO\(_3\). Trajectory A–a starts at 222.1° E and 59.9° N on 17 January with an initial HNO\(_3\) mixing ratio of 9.05 ppbv, adopted from the nearby LIMS observation. Both the starting and termination points for this trajectory are very near the center of the AH as seen in the GPH anomaly contours (Fig. 3b). The mixing ratio of HNO\(_3\) increases to 10.8 ppbv, mostly due to the heterogeneous hydrolysis involving N\(_2\)O\(_5\) and sulfate aerosol. This conversion of NO\(_x\) predominates while the parcel is in darkness.

Figure 8a shows the variation with time of selected chemical constituents as the air parcel moves along trajectory A–a. Shown by the thick broken line at the top of the figure are the segments when the parcel along this trajectory is in darkness. It is clear that HNO\(_3\) increases during extended periods of darkness, which occur more often when the parcel traverses through higher latitudes. This increase in HNO\(_3\) comes at the expense of other reactive nitrogen species as can be seen in the decrease in the mean value and somewhat diminished amplitude of the diurnal variation of NO\(_2\). There is also a dampening of the diurnal variation of N\(_2\)O\(_5\) because of limited exposure to sunlight, although its mean value remains higher. The daytime peak mixing ratio of NO is low (not shown). When a trajectory is at lower latitudes (e.g., between 17 and 22 January), all NO\(_x\) species display larger diurnal variations.
with little change in mean value. HNO₃ also shows some diurnal fluctuations due to daytime photolysis followed by production due to heterogeneous chemistry in darkness and with a slight overall increase in mean mixing ratio. After 22 January, the air parcel moves to higher latitudes and experiences extended periods of darkness, leading to the steady increase in HNO₃. Thus, a combination of dynamics, which determines the trajectory that the parcel follows, and both gas-phase and heterogeneous photochemistry explains the higher levels of HNO₃ measured by LIMS near 30 hPa in the AH region on 27 January. Ozone, which has a longer chemical lifetime at this pressure level, shows almost no change. However, we also note that at lower pressures, e.g., at 5 hPa, there is a similar impact of dynamics and photochemistry in the formation of pockets of low ozone mixing ratio within the AH region during winter (Manney et al., 1995; Morris et al., 1998; Harvey et al., 2004). The photochemical time constant for ozone at those pressure levels is short enough that air parcels originating from lower latitudes and containing higher ozone go through a chemical transformation, when confined for an extended period of time within the AH region. This results in a destruction of ozone in reaching photochemical equilibrium under daylight conditions. Conversely, the increase in HNO₃ at 30 hPa is due mainly to the nighttime heterogeneous chemistry.

In order to highlight the role of the heterogeneous reactions involving background sulfate aerosols, we have repeated the photochemical calculations along the same trajectories but considering only gas-phase reactions. Figure 8b shows the mixing ratios in this case for trajectory A–a. HNO₃ decreases from the initial mixing ratio of 9.1 ppbv and settles to a value closer to 8.0 ppbv by 20 January. Because we initialized the parcel using LIMS HNO₃ data, that initial drop is indicative of the imbalance created by removing the source due to heterogeneous conversion. Small diurnal fluctuations are apparent during the passage through lower latitudes because of photolysis, but they are nearly absent when the parcel is in the high-latitude region. Without heterogeneous reactions, N₂O₅ remains the primary reservoir of NOₓ during the nighttime and reaches its peak values just when the parcel is about to emerge from darkness. The mean mixing ratio of N₂O₅ is about 2 ppbv between 23 and 25 January, when the parcel experiences that extended period of darkness at high latitude. The diurnal cycle for NO₂ exhibits larger amplitudes when using only gas-phase reactions, since the absence of additional production of HNO₃ keeps the NOₓ mixing ratio higher. The difference in the mixing ratio of HNO₃ in the AH region between the two simulations shows the impact of heterogeneous reactions in the partitioning of odd nitrogen. While the agreement for HNO₃ between model and the LIMS data of Fig. 5 is good with the inclusion of heterogeneous reactions, the comparison for NO₂ is worse; the model gives mixing ratios that are lower than the measurement. However, that is also when the V6 NO₂ in the AH region is near its local minimum of about 0.5 ppbv, and we noted in Sect. 4 that such small retrieved values are likely to have a high bias.

Trajectory B–b, shown in Fig. 7a, starts from 173.3° E and 23° N on 14 January. After spending a few days in the lower latitudes, the air parcel along this trajectory takes a nearly meridional path to the AH region. During the last 5 days the parcel remains confined in the AH region, similar to the parcel along trajectory A–a. The chemical evolution along the trajectory, shown in Fig. 9a, is also similar to that along trajectory A–a displayed in Fig. 8a. Large diurnal variations in NO₂ and N₂O₅ occur during the initial period between 14 and 20
January, when the trajectory is in the lower latitudes. Even HNO₃ displays noticeable variations with an increase during night caused by the heterogeneous conversion of N₂O₅ followed by a decrease due to photolysis during daytime. Amplitudes of the diurnal variations diminish in the high-latitude AH region due to reduced photolytic loss, especially for HNO₃, during the shorter daylight period and at higher zenith angles. The corresponding increase in the nighttime heterogeneous conversion leads to HNO₃ mixing ratios greater than 10 ppbv at the end of the trajectory. Between 22 and 25 January, NO₂ displays a steady diurnal cycle, while N₂O₅ shows a declining peak value. The increase in HNO₃ does not occur with a corresponding decrease in NO₂. During extended periods of darkness, NO₂ decreases to a negligible amount as shown at the end of the trajectory on 27 January. When there are only gas-phase reactions, the variation of HNO₃ is as shown by the red line. Again, initialization of the photochemical model uses the V6 data, and the absence of heterogeneous reactions introduces an imbalance leading to the negative tendency in HNO₃.

Trajectory C–c, shown in Fig. 7a, starts from 79.9° E and 27.6° N on 17 January. Until 24 January, this parcel stays south of 40° N and then takes a meridional path directly northward reaching 80° N by 28 January. This is slightly north of the AH but still outside the polar vortex. Figure 9b shows variations of the species mixing ratios along the trajectory C–c for the simulation with heterogeneous chemistry. HNO₃ mixing ratio at the beginning of the trajectory C is about 7.8 ppbv, and it increases to about 8.7 ppbv at the endpoint c. Measurements in Fig. 3d indicate a quasi-wave 2 feature in HNO₃ at 31.6 hPa, with peak values in the polar vortex region and AH. The minimum in HNO₃ between these two regions is a result of the rapid transit of air parcels from lower latitude along trajectories similar to C–c. The air parcels remain in the low latitudes for a longer time, and photolysis during daylight hours keeps the net change in HNO₃ low. Only later is there a noticeable increase in HNO₃ for the high-latitude segment of the trajectory. NO₂ and N₂O₅ display diurnal variations as expected in the middle latitudes. After 21 January, there is a decrease occurring in the peak value of NO₂ at every successive sunset along the trajectory. The corresponding minimum in N₂O₅ shows a small increasing trend especially after 24 January, and this is due to the increasing lifetime against photolysis at winter high latitudes. The parcel is in complete darkness during the last 24 hours, when NO₂ continuously decreases to a very low mixing ratio. N₂O₅ declines slowly, and HNO₃ increases. Figure 10 shows HNO₃ mixing ratios calculated on a longitude–latitude grid for the end time of the ensemble of 70 trajectories, corresponding to 09:00 GMT on 28 January. The latitudinal variation in HNO₃ between 60° N in the AH region and the North Pole clearly shows a dip to lower values near 80° N in the model calculations. This spatial distribution occurs even when heterogeneous reactions are not included. However, calculations with heterogeneous chemistry simulate the magnitude of the LIMS HNO₃ observations better. Combined with the higher HNO₃ values in the polar vortex (not a focus of this paper), the differing trajectories explain the formation of a quadrupole structure at 30 hPa seen in the LIMS HNO₃ observations during the minor warming. As reported by R93, this feature is present in the LIMS V5 data also, except that V5 HNO₃ is nearly of the same magnitude in the AH and the polar vortex at 30 hPa on 27 January. For the same conditions, the V6 HNO₃ is larger in the polar vortex than in the AH by about 2 ppbv.
Even though we have shown and discussed the model results for only three of the trajectories, all of which end along 214° E longitude, the results for the entire ensemble provide a consistent picture. Figure 11a is a scatter plot of the calculated HNO\textsubscript{3} along the 70 trajectories versus the spatially and temporally closest LIMS observations. Differences between the model and LIMS data are of the order of the RSS error of 0.8 ppbv, except for mixing ratios greater than 10 ppbv where the model values are higher. Figure 11b is a similar scatter plot for NO\textsubscript{2}; model values are lower than the V6 observations that have an RSS error of 0.24 ppbv. Deviations from the diagonal dashed line for both HNO\textsubscript{3} and NO\textsubscript{2} could be due to a variety of factors, in addition to bias errors of the data. The criteria we used for selecting the closest observation are coarse, but tightening those criteria reduces the amount of data available for initializing the model and for comparison along the trajectory. While we used the closest LIMS observation to constrain the initialization of the model, other unmeasured species as well as total odd nitrogen are from two-dimensional model output that introduces some uncertainty. We have used the recommended kinetic rate constants, but any uncertainty in key reaction rates could affect the calculated variations in the composition. Another possible source of error is the background stratospheric aerosol, surface area density. We used the climatology from the IGAC/SPARC CCMI database, which is zonally averaged data. There were no major volcanic perturbations in late 1978 and early 1979, so large perturbations from this database are most unlikely. While at the lower latitudes photolysis during the daylight hours is important in limiting the impact of heterogeneous reactions, the aerosol data directly affects that rate of conversion of N\textsubscript{2}O\textsubscript{5} to HNO\textsubscript{3}. It may be that background aerosols in the vortex at high latitudes are less abundant than prescribed, which could explain the apparent high bias in model HNO\textsubscript{3} in Fig. 11a for values greater than 10 ppbv. Although LIMS did not measure N\textsubscript{2}O\textsubscript{5}, analyses involving ATMOS measurements show clearly the role of N\textsubscript{2}O\textsubscript{5} and heterogeneous reactions in the stratospheric odd nitrogen chemistry (Natarajan and Callis, 1991). A more comprehensive study of the changes in atmospheric composition in the AH region using data from more recent satellite experiments is beyond the scope of this study.

7 Seasonal evolution of PV, HNO\textsubscript{3}, H\textsubscript{2}O, and NO\textsubscript{2}

Anticyclone features usually develop in the Northern Pacific stratosphere and are present about 60 % of the time during winter (Harvey and Hitchman, 1996; Baldwin and Holton, 1988). Therefore, we place the V6 species variations of 14–27 January into the broader context of their seasonal variations. As before, one can ignore the effects of any remnants from PSCs for the species away from the polar winter vortex. First, Fig. 12 is a time series plot of the dynamical tracer, PV, on the isentropic surface of 550 K (near 31.6 hPa) for 25 October 1978 through 28 May 1979.

\begin{equation}
PV = (f + \zeta)/\sigma,
\end{equation}

where \( f = 2\Omega \sin \phi \) is the local vertical component of the planetary vorticity on a pressure surface and \( \zeta = \left( r \cos \phi \right)^{-1} \left( \partial v / \partial \lambda - \partial (u \cos \phi) / \partial \phi \right) \) is the relative vorticity in polar coordinates (longitude \( \lambda \) and latitude \( \phi \)). \( \sigma \) is isentropic density \( (\text{kg m}^{-2} \text{K}^{-1}) \) and \( 1/\sigma = -g \partial \theta / \partial \rho = (1/\rho) \partial \theta / \partial z \) is static stability. Geostrophic wind components, \( u \) and \( v \), are
calculated at grid points from the V6 GPH fields. Then, daily values of the vertical component of PV are computed at each grid point from the zonal and meridional components of the wind \((u \text{ and } v)\), plus the local vertical gradients of potential temperature versus pressure from V6, following Harvey et al. (2009). The ordinate of Fig. 12 is in terms of equivalent latitude \(\phi\) from the North Pole \((90^\circ)\) to \(15^\circ\) N and is from a monotonic ordering of the daily PV from high values inside the polar vortex to lower values outside (see, e.g., Butchart and Remsberg, 1986). Thus, equivalent latitude is a vortex-centered coordinate that assigns the highest PV values (located in the center of the vortex) to be at \(90^\circ\) N. Tic marks along the abscissa denote the middle of each month, and the PV time series have a seven point smoothing. The ordinate is linear in \(\phi\) to accentuate variations in the PV field at high equivalent latitudes. The effects of the AH on the displacement and erosion of the PV vortex during 14–27 January are in Fig. 12 from \(\phi\) of about 60 to \(90^\circ\) and following the tic mark labeled 01 on the abscissa.

The effect of the AH in peeling away material contours from the edge of the polar vortex has been described aptly (e.g., McIntyre, 1995; Juckes and McIntyre, 1987; Rose, 1986). Figure 12 indicates the continual erosion of highest PV values during late winter and early spring due to zonal planetary wave 1 (the AH) and wave 2 activity. The adjacent “surf zone” region of lower PV values expands and exhibits weakened gradients \((\phi \sim 30 \text{ to } 60^\circ)\) from the meridional mixing of PV across both the lower and higher latitudes (McIntyre and Palmer, 1983). It is also noted that the vortex was split (wave 2) at four separate times at the level of \(31.6\) hPa: late October, late November–early December, late February, and early April. Both scales of zonal forcing are indicative of the effects of planetary wave activity as it propagates from the troposphere to the \(31.6\) hPa level (Juckes and O’Neill, 1988) and/or of a poleward eddy heat flux (Colucci and Ehrmann, 2018). The meridional gradient of PV is quite weak equatorward of \(\phi \sim 60^\circ\) N in winter and then across all latitudes by mid-April, or after the polar vortex has undergone significant erosion. The large-scale anticyclones and associated zonal easterlies expand toward the middle latitudes by April.

Figure 13 shows the corresponding HNO\(_3\) distribution, as determined by averaging HNO\(_3\) values around the daily PV contours and then ordering them according to the \(\phi\) of Fig. 12. Those averages represent approximate, modified Lagrangian mean (MLM) values for HNO\(_3\) or its average values around the PV contours, and they enable one to identify differences in behavior (e.g., chemical changes) for trace constituents versus PV (McIntyre, 1980; Butchart and Remsberg, 1986). HNO\(_3\) varies nearly monotonically with latitude at this level, and values as high as 13 ppbv are found near the center of the vortex \((\phi = 90^\circ)\) by late November and v during the polar night. Such high values indicate a nearly complete chemical conversion of the available NO\(_y\) to its reservoir component HNO\(_3\). Poleward of about \(\phi \sim 70^\circ\) the HNO\(_3\) contours are aligned well with those of PV, indicating that HNO\(_3\) is an excellent tracer at \(550\) K, particularly in winter polar night when further chemical changes are inefficient. However, there should be some HNO\(_3\) uptake into PSCs, when temperatures become cold enough.

A sequence of polar orthographic plots (not shown) indicates that there is a buildup of HNO\(_3\) inside the polar vortex in November, punctuated by meridional transport during the zonal wave 1 events of early December and in January, and then followed by a splitting of
the vortex in mid- to late February. There is significant transport of HNO\textsubscript{3} from the polar region to middle equivalent latitudes ($\varphi = 45$ to $20^\circ$) during those events. Meridional gradients of HNO\textsubscript{3} are larger in winter at both $\varphi$ of 70 and near 20$^\circ$, marking the polar and subtropical edges of the region of efficient meridional mixing. The subtropical boundary of the so-called “tropical pipe” region shifts from $\varphi = 25$ to about $15^\circ$ from early December to late January and then remains at that location into springtime (Remsberg and Bhatt, 1996).

There is erosion of the high HNO\textsubscript{3} values of the polar vortex by early March. Thereafter, HNO\textsubscript{3} decreases at all equivalent latitudes, due to the daily effects of the chemical re-partitioning of NO\textsubscript{y} away from HNO\textsubscript{3} and toward NO and NO\textsubscript{2} under sunlit conditions.

Figure 14 displays the time series plot of the MLM for V6 H\textsubscript{2}O, a better tracer of stratospheric transport. Relatively large values of 5.5 to 6.0 ppmv occur at high latitudes from mid-November to early January and indicate the effects of the slow descent of higher values within the vortex. H\textsubscript{2}O is higher in the upper stratosphere from the oxidation of methane. Elevated H\textsubscript{2}O values at $\varphi >75^\circ$ on 13–15 and 19 January are due to residual emissions from PSCs. However, values of ~ 7 ppmv also appear in early February to mid-March, when temperatures are much too warm for the existence of PSCs. Although those higher values are within the uncertainties for retrieved H\textsubscript{2}O, they compare well with times when there is descent of higher ozone values in response to the major stratospheric warmings (see Fig. 18 in Leovy et al., 1985).

Finally, Fig. 15 is the MLM plot of V6 NO\textsubscript{2}, based on only its profiles along descending (nighttime) orbital segments. Smallest values of NO\textsubscript{2} occur in the polar vortex in late November and early December, when the HNO\textsubscript{3} values of Fig. 13 reach 13 ppbv. As with H\textsubscript{2}O, there are several minor increases in NO\textsubscript{2} at the highest latitudes in mid-January, and they occur at locations of residual effects from PSCs. The excess values of NO\textsubscript{2} poleward of $\varphi 75^\circ$ in February occur where the variations of PV in Fig. 12 also indicate the effects of transport and where there may have been descent of higher NO\textsubscript{2} values within the vortex (Holt et al., 2012). The distribution of NO\textsubscript{2} away from the vortex varies more slowly and smoothly. Figure 12 also indicates that there is considerable mixing for PV at $\varphi \sim 60$ to $75^\circ$ during February, and Fig. 15 shows that NO\textsubscript{2} is increasing along the PV contours. The vortex split into two sectors in the middle stratosphere from mid- to late February, when there was transport and descent of air with higher NO\textsubscript{2} values at the high latitudes. NO\textsubscript{2} increases steadily from March to May, due to the conversion of HNO\textsubscript{3} to NO\textsubscript{2} upon the return of sunlight.

There are significant seasonal variations of NO\textsubscript{2} displayed in Fig. 15 at middle equivalent latitudes. The re-partitioning of the NO\textsubscript{y} species to HNO\textsubscript{3} occurs in the presence of background aerosols from late autumn to winter, followed by photochemical conversion of the HNO\textsubscript{3} vapor back to NO\textsubscript{2} in springtime (e.g., Austin et al., 1986). Figure 15 shows that the very low values of NO\textsubscript{2} extend from near the North Pole to at least $\varphi = 30^\circ$ in early December and then retreat toward higher latitudes by late February. This variation of the NO\textsubscript{2} time series is an indicator of the so-called “Noxon cliff” feature of stratospheric column NO\textsubscript{2} during winter (e.g., Noxon, 1979).
8 Conclusions

A significant improvement of the LIMS data set from V5 to V6 is the better accuracy of the retrieved V6 NO\textsubscript{2} profiles. Both the V6 HNO\textsubscript{3} and NO\textsubscript{2} are of good quality, at least to within their respective error estimates and away from very cold regions of the vortex and their PSC remnants. The V6 species data are evaluated further in terms of the consistency of the HNO\textsubscript{3} and NO\textsubscript{2} distributions in the AH region during the minor warming event that took place in January 1979. In an earlier analysis of the LIMS V5 data, R93 highlighted an increase for HNO\textsubscript{3} at 30 hPa within the AH region but with little change in NO\textsubscript{2}, and they suggested the need for some unknown process leading to the production of HNO\textsubscript{3} and to the development of the quasi-wave 2 signature in its zonal distribution. The present study considers photochemical model calculations along kinematic trajectories over a 10-day period that terminate in the AH region on 28 January. The results indicate that there was an increase of about 2 ppbv in HNO\textsubscript{3} and a decrease of the order of 0.5 ppbv in NO\textsubscript{2}, mainly as a result of heterogeneous reactions converting N\textsubscript{2}O\textsubscript{5} on surfaces of background stratospheric sulfuric acid aerosols. R93 and Considine et al. (1992) alluded to this mechanism but reported that their three- and two-dimensional model studies with heterogeneous chemistry gave results that still did not agree well with the LIMS V5 species. On the contrary, the latitudinal variations of HNO\textsubscript{3} at the end of our Lagrangian trajectory calculations agree reasonably with the LIMS V6 data. Those variations depend on the initial conditions and the extent of exposure of air parcels to sunlit and dark conditions, and hence on the dynamically controlled histories of the separate trajectories. Our model calculations also reveal the formation of a dip in observed HNO\textsubscript{3} mixing ratios north of the AH due to meridional transport of low-latitude air across the North Pole. Therefore, we conclude that the study approach of R93 was valid and should have led them to better comparisons, if the V6 data set had been available earlier.

When the effects of heterogeneous chemistry are included in the model calculations, the variations of HNO\textsubscript{3} and NO\textsubscript{2} along trajectories agree more reasonably with the LIMS V6 observations in the relatively isolated AH region. However, the model still underestimates NO\textsubscript{2} compared to the V6 values along some trajectory paths. A part of those differences may be due to an inability to retrieve a local minimum in the V6 NO\textsubscript{2} profile with good accuracy, at least based on the finite, vertical field of view of the NO\textsubscript{2} channel radiances and their associated LIMS temperature profiles. Still, the present study demonstrates that a combination of dynamical and photochemical changes can explain the maximum mixing ratios of HNO\textsubscript{3} in both the AH region and at the winter polar vortex, plus its lower values around the periphery of both circulation systems. HNO\textsubscript{3} re-partitions by photochemistry toward NO\textsubscript{2} from winter to springtime, when the anticyclone regions extend to middle latitudes. We also present examples of the seasonal evolution of HNO\textsubscript{3} and NO\textsubscript{2} during 1978–1979 as a separate aspect of the V6 data set for the validation of chemistry–climate models of the middle to lower stratosphere.

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Figure 1.
Polar orthographic projections of Northern Hemisphere ozone (a), geopotential height (GPH, b), temperature (c), and gas-phase nitric acid (d) for 11 January 1979 at 31.6 hPa; successive latitude circles are at every 10. The Greenwich meridian extends horizontally to the right. Contour intervals are every 0.75 ppmv for ozone, 0.25 km for GPH, 4 K for temperature, and 1 ppbv for nitric acid. White plus signs denote orbital profile segments that are missing; red dot denotes location of a SAM II PSC observation.
Figure 2.
Zonal anomalies of ozone, GPH, nitrogen dioxide, and nitric acid for 31.6 hPa on 17 January 1979.
Figure 3.
As in Fig. 2, but for 27 January 1979.
Figure 4.
Zonal variations of LIMS V6 (black) and V5 (red) species at 64° N on 27 January 1979. Vortex is between 0 and 90° E, and the AH region is from 180 to 240° E. Vertical bars on curves near 311° E are 2σ estimates of the error. Ozone has units of ppmv, while HNO₃ and NO₂ (its separate descending and ascending curves at bottom) have units of ppbv.
Figure 5.
Time series of observed LIMS V6 species at 31.6 hPa and the Aleutian High (AH) anomaly; latitude of AH center is the solid curve. The species curves have vertical bars near 24–26 January indicating their ±RSS errors. Ozone and water have units of ppmv, while NO$_2$ and HNO$_3$ have units of ppbv.
Figure 6.
History of the location of maximum GPH anomaly at 31.6 hPa, representing the AH center and displayed by red squares with day numbers. Latitude spacing is 10° beginning at 20° N and longitude spacing is 15° beginning at 120° E. Note that between 21 and 27 January there is some repetition of the AH center locations, and the corresponding red squares overlap. The location on 27 January is 214° E and 68° N. Black line with day numbers describes the back trajectory beginning at 214° E, 68° N, and 30 hPa.
Figure 7.
(a) Back trajectories beginning at 23:00 on 27 January (09:00 GMT on January 28) from 30 hPa, 214° E, and three different latitudes of (a) 60° N, (b) 72° N, and (c) 80° N. The corresponding endpoints A, B, and C are spatially and temporally closest to LIMS descending mode measurement locations between 14 and 17 January. The latitude grids are 10° apart, starting from 20° N and the prime meridian extends horizontally to the right. The color scale refers to the accumulated hours of darkness expressed as a fraction of the total length of the trajectory in hours as the air parcel moves in the forward direction starting from locations A, B, and C and ending at a, b, or c along 214° E longitude. (b) Evolution of HNO$_3$ along the three trajectories that were shown in panel (a).
Figure 8.
Mixing ratio of selected species as a function of time along the trajectory A–a shown in Fig. 7a. Air parcel terminates at 214° E and 60° N. HNO$_3$ (solid), NO$_2$ (dash), and N$_2$O$_5$ (dash-dot-dot) are in ppbv, and ozone (dash-dot) is in ppmv. The tick marks on the abscissa correspond to 09:00 GMT on the dates shown. The thick line at the top represents the periods of darkness along the trajectory with intermittent gaps corresponding to sunlit segments. (a) Results from the case including heterogeneous reactions, but with the red curve showing HNO$_3$ from only the gas-phase chemistry for comparison. (b) Results for all species from only the gas-phase reactions.
Figure 9.
Mixing ratio of selected species as a function of time for the case with heterogeneous reactions as shown in Fig. 8a but for trajectory B–b (a) and trajectory C–c (b). HNO$_3$ (solid), NO$_2$ (dash), and N$_2$O$_5$ (dash-dot-dot) are in ppbv, and ozone (dash-dot) is in ppmv. The red curve in both panels represent the HNO$_3$ variation for the case with only gas-phase reactions.
Figure 10.
Calculated HNO$_3$ at 09:00 GMT on 28 January, corresponding to terminal location of all 70 trajectories.
Figure 11.
Scatter plot of calculated species mixing ratios along the 70 trajectories and of the corresponding spatially and temporally closest LIMS V6 observation. (a) HNO$_3$; (b) NO$_2$. 
Figure 12.
Time series of LIMS isentropic PV vs. equivalent latitude at 550 K and with smoothing over 7 days. PV contour interval (CI) is 10 units.
Figure 13.
As in Fig. 12, but the averages of HNO$_3$ along PV isolines (CI is 0.5 ppbv).
Figure 14.
As in Fig. 13, but as averages of H$_2$O (CI = 0.2 ppmv).
Figure 15.
As in Fig. 13, but as averages of descending orbital (nighttime) NO$_2$ (CI = 0.2 ppbv).