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To cite this version:
G. Dufour, Sébastien Payen, Franck Lefèvre, M. Eremenko, A. Butz, et al.. 4-D comparison method to study the NOy partitioning in summer polar stratosphere – Influence of aerosol burden. Atmospheric Chemistry and Physics, European Geosciences Union, 2005, 5 (4), pp.926. hal-00328382

HAL Id: hal-00328382
https://hal.archives-ouvertes.fr/hal-00328382
Submitted on 21 Mar 2005

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4-D comparison method to study the NO\textsubscript{y} partitioning in summer polar stratosphere – Influence of aerosol burden

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Received: 12 May 2004 – Published in Atmos. Chem. Phys. Discuss.: 10 December 2004
Revised: 23 February 2005 – Accepted: 9 March 2005 – Published: 21 March 2005

Abstract. On 21–22 August 2001, NO, NO\textsubscript{2} and HNO\textsubscript{3} mixing ratio profiles were measured at high latitudes during sunset and sunrise using the Limb Profile Monitor of the Atmosphere (LPMA) and the DOAS experiments under stratospheric balloon. Photochemical simulations using the chemistry module of the Reprobus Chemistry Transport Model (CTM) that are constrained by ozone and total NO\textsubscript{y} balloon observations reproduce well the partitioning of NO\textsubscript{x} and NO\textsubscript{y} when model results are calculated at the exact time and location of the measurement for each tangent altitude. Taking the recently recommended reaction rate coefficients for the NO\textsubscript{x} partitioning (JPL-2003) and using realistic aerosol surface area in order to initialise the model leads to an agreement between calculations and measurements better than 10% all over the covered altitude range.

1 Introduction

Stratospheric ozone loss results mainly from catalytic cycles involving reactive nitrogen (NO\textsubscript{x}), hydrogen (H\textsubscript{2}O) and halogens (ClO\textsubscript{x} and BrO\textsubscript{x}) species. In the lower stratosphere, the nitrogen radicals (NO\textsubscript{x}=NO+NO\textsubscript{2}) play an important role by catalytically removing ozone and by moderating indirectly ozone loss through the coupling between the different radical families (Wennberg et al., 1994) and the formation of reservoir molecules. Many studies on nitrogen species, in particular addressing the polar winter stratosphere (Lary et al., 1997; Wetzel et al., 1997), demonstrated that for NO\textsubscript{x} the differences between model and measurement values are often larger than several ten percents. In particular the NO\textsubscript{2} volume mixing ratio, the NO\textsubscript{2}/HNO\textsubscript{3} ratio and the NO\textsubscript{x}/NO\textsubscript{2} (NO\textsubscript{y}=NO\textsubscript{x}+NO\textsubscript{3}+HNO\textsubscript{3}+2\times N\textsubscript{2}O\textsubscript{5}+HNO\textsubscript{2}+HNO\textsubscript{4}+ClONO\textsubscript{2}+ BrONO\textsubscript{2}) ratio are found to largely disagree in the lower stratosphere below 30 km. In order to improve the photochemical models, a better understanding of the partitioning of NO\textsubscript{x} and NO\textsubscript{y} is thus needed. The NO/NO\textsubscript{2} ratio is controlled by fast photochemistry which inter-converts NO and NO\textsubscript{2}, mainly through Reactions (R1) and (R2):

\begin{align*}
\text{NO}_2 + h\nu & \rightarrow \text{NO} + \text{O} \quad \text{(R1)} \\
\text{NO} + \text{O}_3 & \rightarrow \text{NO}_2 + \text{O}_2 \quad \text{(R2)}
\end{align*}

The partitioning between NO\textsubscript{2} and HNO\textsubscript{3}, and consequently between NO\textsubscript{x} and NO\textsubscript{y}, is dominated by slower reactions. In the lower stratosphere, the dominant sink of NO\textsubscript{x} is a two-step process involving the formation of N\textsubscript{2}O\textsubscript{5} during the night, followed by the heterogeneous hydrolysis of N\textsubscript{2}O\textsubscript{5} on sulphate aerosols, which converts N\textsubscript{2}O\textsubscript{5} to the more stable species HNO\textsubscript{3}. In summer at high latitudes, the extended daylight duration is implying efficient photolysis which inhibits the formation of the precursor molecule NO\textsubscript{3} and thus of N\textsubscript{2}O\textsubscript{5}. In effect, these conditions provide an opportunity to test our understanding of the NO\textsubscript{x} partitioning when gas-phase chemistry dominates the exchange between NO\textsubscript{x} and HNO\textsubscript{3}. Reactions (R3–R5) are the 3 main reactions that govern the NO\textsubscript{2}/HNO\textsubscript{3} ratio:

\begin{align*}
\text{HNO}_3 + h\nu & \rightarrow \text{NO}_2 + \text{OH} \quad \text{(R3)} \\
\text{HNO}_3 + \text{OH} & \rightarrow \text{NO}_3 + \text{H}_2\text{O} \quad \text{(R4)} \\
\text{NO}_2 + \text{OH} + \text{M} & \rightarrow \text{HNO}_3 + \text{M} \quad \text{(R5)}
\end{align*}

Several field studies performed during the 1997 Photochemistry of Ozone Loss in the Arctic Region In Summer (POLARIS) mission investigated the partitioning of NO\textsubscript{x} species (Osterman et al., 1999; Gao et al., 1999; Perkins et al., 2001). These studies showed that a significant disagreement between observed and modelled NO\textsubscript{x} is observed when using the reaction rates recommended by DeMore et al. (1997).

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Table 1. Time, latitude and longitude of LPMA sunset measurements for selected tangent altitudes $H_t$.

<table>
<thead>
<tr>
<th>Time (UT)</th>
<th>Latitude</th>
<th>Longitude</th>
<th>$H_t$ (km)</th>
</tr>
</thead>
<tbody>
<tr>
<td>18:45</td>
<td>68.07</td>
<td>21.01</td>
<td>38.68</td>
</tr>
<tr>
<td>18:50</td>
<td>68.32</td>
<td>20.21</td>
<td>38.29</td>
</tr>
<tr>
<td>18:55</td>
<td>68.59</td>
<td>19.44</td>
<td>37.69</td>
</tr>
<tr>
<td>19:00</td>
<td>68.86</td>
<td>18.71</td>
<td>37.77</td>
</tr>
<tr>
<td>19:05</td>
<td>69.14</td>
<td>18.01</td>
<td>35.51</td>
</tr>
<tr>
<td>19:10</td>
<td>69.43</td>
<td>17.34</td>
<td>33.96</td>
</tr>
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<td>19:25</td>
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<td>19:30</td>
<td>70.61</td>
<td>15.13</td>
<td>26.11</td>
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<td>19:35</td>
<td>70.91</td>
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<td>19:40</td>
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<td>71.75</td>
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<tr>
<td>19:55</td>
<td>72.04</td>
<td>13.70</td>
<td>14.75</td>
</tr>
<tr>
<td>19:59</td>
<td>72.26</td>
<td>13.58</td>
<td>13.06</td>
</tr>
</tbody>
</table>

In particular, Osterman et al. (1999) recommended a reduction in the rate coefficient of Reaction (R5) by about 35% to achieve a reasonable agreement (better than 10% for altitudes higher than 15 km and than 30% at lower altitudes) between the modelled and the measured NO$_2$/NO$_y$ ratios higher than 15 km and than 30% at lower altitudes) to achieve a reasonable agreement (better than 10% for all $H_t$ altitudes). For instance, Osterman et al. (1999) recommended a reduction in the rate coefficient of Reaction (R5) by about 35% to achieve a reasonable agreement (better than 10% for altitudes higher than 15 km and than 30% at lower altitudes) between the modelled and the measured NO$_2$/NO$_y$ ratio higher than 15 km and than 30% at lower altitudes.)

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2 LPMA observations

The balloon flight reported here took place from Kiruna (Sweden) on 21–22 August 2001. The measurements were performed during three flight phases: balloon ascent from 15:55 UT to 18:42 UT, sunset from 18:43 UT to 20:05 UT and sunrise from 1:11 UT to 2:29 UT. Infrared spectra have been recorded by the LPMA instrument, which is a remote sensing infrared Fourier transform interferometer operating in solar occultation (Camy-Peyret, 1995). The high spectral resolution and sensitivity of the LPMA instrument permits to retrieve vertical profiles of species with stratospheric mixing ratios of the order of a few tenths of ppbv, such as ClONO$_2$. On the same gondola, a UV-visible DOAS (Differential Optical Absorption Spectroscopy) instrument (Ferlemann et al., 2000) analysed the same sun light for O$_3$ and NO$_2$ profiles. Both instruments LPMA and DOAS use the same suntracker to lock onto the sun. Thus, their line of sight (LOS) is identical and the retrieved vertical profiles are directly comparable. The consistency of O$_3$ and NO$_2$ retrieved vertical profiles has been studied in detail (Butz et al., 2005). The LPMA instrument tracked the sun between 10 km up to the float altitude during balloon ascent and until loss of Sun, with a lowest tangent altitude of 12 km, during sunset. Spectra were recorded at sunrise from a tangent altitude of 15 km up to the float altitude of 39 km. Profiles at sunrise were retrieved down to 12 km. Tables 1 and 2 summarize the time (UT), the location and the altitude of the tangent points considered in the present study at sunset and sunrise, respectively.

The spectral retrieval of the target species relies on a multtifit algorithm that uses an efficient minimization algorithm based on the Levenberg-Marquardt algorithm (Press et al., 1992). It allows the simultaneous detection and retrieval of vertical profiles of CH$_4$, N$_2$O, NO, NO$_2$, HCl, ClONO$_2$ and O$_3$ in 7 micro-windows. HNO$_3$ profiles are retrieved using the same algorithm but on a larger spectral window.
(25 cm⁻¹). All the molecular parameters are extracted from the HITRAN2000 database (Rothmann et al., 2000), except for ClONO₂, for which we use the absorption cross-sections measured by Wagner and Birk (2003). The error bars of retrieved NO, NO₂, HNO₃ and ClONO₂ volume mixing ratio correspond to 2σ fitting error: they do not include, however, the uncertainties on spectroscopic parameters. These are later added to fitting errors in order to estimate the systematic errors: the total error bars are estimated to 10% for NO and NO₂, to 15% for HNO₃ and to 20% for ClONO₂. For tangent heights below 19 km, the spectral micro-window used for the retrieval of ozone is saturated and thus it is difficult to fit correctly the spectrum base line. As a result, the O₃ mixing ratio retrieved from LPMA is underestimated. Consequently, below 19 km, we have used the ozone vertical profiles (for sunset and sunrise) retrieved from the DOAS UV-visible measurements (Butz et al., 2005).

3 Comparison method between measurements and model results

The photochemical model used here is a one-dimensional version of the Reprobus chemical-transport model (CTM) (Lefèvre et al., 1994, 1998). The model provides a comprehensive description of the stratospheric chemistry by inclusion of 147 photolytic, gas-phase, and heterogeneous reactions. Most of the used absorption cross-sections, gas-phase reactions, and heterogeneous reaction probabilities are those recommended by the latest JPL compilation (Sander et al., 2002). The aerosol surface area used in the model is inferred from the SAGE-II satellite observations (Thomason et al., 1997). The model extends from the ground up to 0.1 hPa (about 65 km), on 42 vertical levels. We used a short chemical time step of 15 s in order to describe accurately the rapid variations of NOₓ species at sunrise and sunset.

Along the line of sight strong spatial variations of the radical species must be taken into account in the retrieval of vertical profiles. The NO volume mixing ratio (vmr) decreases rapidly during sunset and increases rapidly during sunrise. For example, the NO vmr varies by about 20% between 89° and 91° of solar zenith angle (SZA). In this case the assumption of a uniform mixing ratio along the line of sight would lead to significant errors. We use the Reprobus 1-D model to calculate the altitude-dependent diurnal variation of NO relative to a SZA reference value of 90°. Derived correcting factors are then used in the retrieval algorithm to correct the NO vmr on each side of the tangent point along the line of sight. The corrected NO profile is compared to the uncorrected one.
Fig. 3. Comparison between measured NO, NO$_2$ and HNO$_3$ profiles during sunset and corresponding calculated profiles with the Reprobus 1-D model initialised with Reprobus CTM.

Fig. 4. (a) Comparison between measured NO$_x$/NO$_y$ profile during sunset and the corresponding calculated profile with the Reprobus 1-D model initialised with Reprobus CTM. (b) Comparison between measured NO/NO$_2$ profile during sunset and the corresponding calculated profile with the Reprobus 1-D model initialised with Reprobus CTM.

4 Results and discussion

In this study, the Reprobus 1-D model is initialised shortly before the LPMA measurements at 16:00 UT on 21 August 2001 (sunset) and at 1:00 UT on 22 August 2001 (sunrise). In a first analysis, all species are initialised from the result of a three-dimensional simulation of the Reprobus chemical-transport model on 21 August 2001. Figure 2 plots the ozone and NO$_2$ vertical profiles computed near the two extreme locations of the sunset measurements, for the same solar zenith angle. A large spatial variability is observed, with differences of 1.4 ppmv around the maximum of ozone and 0.7 ppbv for NO$_2$ (Fig. 2). In order not to introduce additional errors, it is very important to compare measurements and model results at the same time, altitude, longitude and latitude: one must perform a 4-D comparison.

During the LPMA and DOAS occultation measurements, the location of the tangent point varies by several degrees in latitude and longitude. Vertical profiles of stratospheric species may thus show substantial variations within the area sampled by the instruments. This is illustrated by the output of a three-dimensional simulation of the Reprobus chemical-transport model on 21 August 2001. Figure 2 plots the ozone and NO$_2$ vertical profiles computed near the two extreme locations of the sunset measurements, for the same solar zenith angle. A large spatial variability is observed, with differences of 1.4 ppmv around the maximum of ozone and 0.7 ppbv for NO$_2$ (Fig. 2). In order not to introduce additional errors, it is very important to compare measurements and model results at the same time, altitude, longitude and latitude: one must perform a 4-D comparison.
measured and simulated profiles of NO, NO$_2$ and HNO$_3$ are summarized on Fig. 3. Calculated volume mixing ratio of NO, NO$_2$ and HNO$_3$ are underestimated by the Reprobus 1-D model. The disagreement for HNO$_3$ is explained by the underestimation of the total amount of NO$_y$ in the model. The partitioning between NO$_x$ and NO$_y$ and between NO and NO$_2$ are shown on Fig. 4. For the NO$_x$/NO$_y$ ratio (Fig. 4a), the model is simulating well the ratio for the full altitude range. Thus, relative values of NO$_y$ species are well reproduced but not their absolute values. The NO/NO$_2$ ratio is correctly modelled by Reprobus 1-D for all the altitude range, except for altitudes lower than 19 km. Since during high latitude summer the partitioning between NO and NO$_2$ is mainly governed by Reactions (R1) and (R2) the NO/NO$_2$ ratio can be approximated by:

$$\frac{[\text{NO}]}{[\text{NO}_2]} \approx \frac{J_{\text{NO}_2}}{k_1[O_3]} \quad (1)$$

Accordingly if the ozone concentration is overestimated, this ratio becomes smaller and the NO and NO$_2$ concentrations calculated with a photochemical steady state model are both underestimated. Measured vertical profiles of ozone by LPMA and DOAS experiments are compared to the calculated one on Fig. 5. Ozone volume mixing ratios are overestimated by the model for all the altitude range, except for altitudes lower than 19 km. Since during high latitude summer the partitioning between NO and NO$_2$ is mainly governed by Reactions (R1) and (R2) the NO/NO$_2$ ratio can be approximated by:

$$\frac{[\text{NO}]}{[\text{NO}_2]} \approx \frac{J_{\text{NO}_2}}{k_1[O_3]} \quad (1)$$

The first analysis shows that the NO$_x$ gas-phase chemistry and the temporal evolution of NO$_x$ are well represented in the model. The disagreements observed for individual profiles are mainly due to the model initialisation. Thus a better initialisation, in particular for ozone and total NO$_y$ amounts, is necessary. One way to initialise NO$_y$ in the model would be to use the N$_2$O profile measured by LPMA and the observed correlation between N$_2$O and NO$_y$ in the stratosphere. However, LPMA measurements of N$_2$O are not sufficiently accurate for a firm inter-comparison. Therefore in a second model run, we chose to constrain the model initialisation by the LPMA measurements of total NO$_y$ and ozone. For NO$_y$, the three dominant species (NO, NO$_2$ and HNO$_3$) are inferred from LPMA retrievals, and they represent between 90 and 95% of total NO$_y$ depending on the considered altitude. The other NO$_y$ species (N$_2$O$_5$, ClONO$_2$, ...) are derived from the output of the 3-D simulation. The time (hh mm UT), the latitude $\lambda$ and the longitude $\mu$ corresponding to each sounded tangent altitude during sunset and sunrise are reported in Tables 1 and 2. The simplified notation ($\lambda$, $\mu$) hh mm is used in the following. As no measurement was performed exactly
at initialisation time, \(hh_i \text{ mm}_i\) UT, (16:00 UT for sunset and 1:00 UT for sunrise) in the different locations \((\lambda, \mu)\), the \(O_3\) and \(NO_y\) profiles retrieved from LPMA during sunset and sunrise have to be used to initialise the model. They are scaled to their estimated value at 16:00 UT and 1:00 UT. This value is computed from the actual measurements, scaled by the variation predicted by the CTM between the initialisation time and the measurement time. We apply Eq. (2) to the volume mixing ratio of species \(X\) for each relevant tangent altitude reported in Tables 1 and 2, i.e. for each corresponding time \((hh \text{ mm} \text{ UT})\) and location \((\lambda, \mu)\).

\[
X_{LPMA}^{(\lambda, \mu)hh_i \text{ mm}_i} = \frac{X_{Reprobus}^{(\lambda, \mu)hh_i \text{ mm}_i}}{X_{LPMA}^{(\lambda, \mu)hh_i \text{ mm}_i}}
\]

Note that constraining the model with LPMA measurements somewhat disturbs the balance within the \(NO_y\) family, as minor \(NO_y\) species are not measured. Thus, before analysing the results, the 1-D model is run for about 4 to 50 days (depending of the considered altitude) until a satisfactory balance is reached.

Using this model initialisation, a very good agreement between measured and modelled vertical profiles of \(NO, NO_2\) and \(HNO_3\) is observed at sunset and sunrise (cf. Fig. 6). Except for a slight overestimation of \(NO\) between 20 and 24 km, the volume mixing ratio values of \(NO, NO_2\) and \(HNO_3\) are well reproduced by the model. A reasonably good agreement is also obtained for \(ClONO_2\), as shown on Fig. 7. The measured \(NO_x/NO_y\) profile is then well reproduced by the model, as for the first model initialisation with the CTM Reprobus (Fig. 8a). The percentage difference between measured \(NO_x/NO_y\) profiles and corresponding calculated values for sunset and sunrise is lower than 5% for altitudes higher than 30 km and of the order of 15% for altitudes lower than 30 km and the difference never exceeds 25%. The model reproduces very well the partitioning between \(NO\) and \(NO_2\). In particular, this second model run shows a better agreement with LPMA below 19 km, as a result of the more realistic ozone initialisation in this altitude range (Fig. 8b). Between 25 and 40 km the \(NO/NO_2\) percentage difference is smaller than 10% and 5% for sunset and sunrise, respectively. For altitudes lower than 25 km, this difference changes more with the observed conditions and it is of the order of 30% at sunrise. For sunset the discrepancy is around 20% for the three lowest altitudes, whereas the simulated ratios do not fall within the range given by the error bars at 22 km. We believe that this discrepancy can be explained to a large extent by an incomplete coincidence of measurement and model times.

Fig. 7. Comparison between measured \(ClONO_2\) profile during sunset and the corresponding calculated profiles with the Reprobus 1-D model constrained by measured \(O_3\) and \(NO_y\) amounts.

Fig. 8. (a) Comparison between measured \(NO_x/NO_y\) profiles during sunset and the corresponding calculated profile with the Reprobus 1-D model constrained by measured \(O_3\) and \(NO_y\) amounts. (b) Comparison between measured \(NO/NO_2\) profiles during sunset and the corresponding calculated profile with the Reprobus 1-D model constrained by measured \(O_3\) and \(NO_y\) amounts.
since the instrument assigns two consecutive spectra to the same line of sight and in consequence to the same tangent altitude. These two spectra correspond to a forward (when the moving mirror of the Michelson interferometer moves away from zero path difference or ZPD. Zero Path Difference) and a reverse (fly back of the mirror to ZPD) interferograms. The duration of this round-trip is around 100 s. A given LOS (Line Of Sight) and time are assigned to a reverse and forward scan sharing almost the same ZPD in the middle of this 100 s interval. The tangent altitude varies of several hundred meters and of the order of 0.1° in latitude and longitude during the recording time. Times and locations indicated in Table 1 and 2 are consequently mean times and locations and this is true for the tangent altitude of two consecutive spectra. The uncertainty on the time and location of the tangent point could explain a part of the discrepancy observed between calculated and measured values of NO/NO₂.

We further test the influence of the aerosol burden on simulated NOₓ. The previous results were obtained with the aerosol burden usually chosen in the Reprobus CTM seasonal calculations. Fig. 9 shows the comparison between the aerosol surface area profile assumed in the Reprobus calculations and the same profile deduced from balloon-borne aerosol measurements performed in 2002 (Deshler, 2003). The accuracy of the measured aerosol surface area is only 40% and the variability of aerosol surface area values between two consecutive years is quite large for altitudes lower than 18 km. However, the difference between modelled and measured surface area below 20 km is larger than the uncertainties on the measured surface area. The calculated aerosol distribution is then not completely realistic in particular for altitudes lower than 20 km. In another model sensitivity test, we thus use the aerosol surface area profile (below 30 km) deduced from balloon-borne measurements. The comparison of calculated and measured NOₓ/NOᵧ profiles is presented on Fig. 10. The partitioning of NOₓ and NOᵧ is governed by both gas-phase and heterogeneous chemistry. Although heterogeneous reactions involving N₂O₅ are not important in the polar summer stratosphere, the impact of aerosol burden initialisation is not negligible for NO, NO₂ and HNO₃ (from 10% to 25% on average, not shown). Taking into account a more realistic measured stratospheric aerosol burden is largely improving the agreement between calculated and measured NOₓ/NOᵧ profiles (cf. Fig. 10). This is especially true below 20 km, where CTM strongly underestimates aerosol. With the new surface area profile, model-observation percentage differences are then smaller than 10% over the covered altitude range, except for the lowest altitude, where it reaches 13%. Very similar results are observed for the sunrise comparison (not shown).

5 Conclusions

The LPMA/DOAS balloon-borne solar occultation measurements performed over Kiruna (Sweden) on 21–22 August 2001 allowed us to check our understanding of the NOᵧ and NOₓ partitioning during polar summer. In order to test the measured versus simulated NOᵧ partitioning, we used a state-of-art chemical transport model that includes the most recent gas-phase rate coefficients, absorption cross-sections and heterogeneous reactions as recommended by the NASA JPL2003 compilations. Vertical profiles of NO, NO₂,
HNO$_3$, NO/NO$_2$ and NO$_x$/NO$_y$, deduced from LPMA measurements are compared to the corresponding profiles calculated with the Reprobus 1-D model. Different model initialisations are tested and we show that measurements and calculations are in good agreement (both absolute concentrations and mixing ratios) when the photochemical model is constrained by measured ozone and total NO$_y$. This confirms earlier results obtained by Osterman et al. (1999) and Gao et al. (1999) that using the 2003 updated JPL reaction rate coefficients for the partitioning of NO$_y$, much of the earlier disagreement between measured and modelled stratospheric NO$_x$ and NO$_y$ disappears for the polar summer stratosphere. Moreover, taking realistic stratospheric aerosol burdens, cf. from the Deshler 2002 measurements, leads to a significantly better agreement.

Edited by: J. P. Burrows

References


