

Energetic particle precipitation in ECHAM5/MESSy – Part 2: Solar proton events

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Abstract. The atmospheric chemistry general circulation model ECHAM5/MESSy (EMAC) has been extended by processes that parameterize particle precipitation. Several types of particle precipitation that directly affect NO_y and HO_x concentrations in the middle atmosphere are accounted for and discussed in a series of papers. In part 1, the EMAC parameterization for NO_x produced in the upper atmosphere by low-energy electrons is presented. Here, we discuss production of NO_y and HO_x associated with Solar Proton Events (SPEs). A submodel that parameterizes the effects of precipitating protons, based on flux measurements by instruments on the IMP or GOES satellites, was added to the EMAC model. Production and transport of NO_y and HO_x, as well as effects on other chemical species and dynamics during the 2003 Halloween SPEs are presented. Comparisons with MIPAS/ENVISAT measurements of a number of species affected by the SPE are shown and discussed. There is good agreement for NO₂, but a severe disagreement is found for N₂O similar to other studies. We discuss the effects of an altitude dependence of the N/NO production rate on the N₂O and NO_y changes during the SPE. This yields a modified parameterization that shows mostly good agreement between MIPAS and model results for NO₂, N₂O, O₃, and HOCl. With the ability of EMAC to relax the model meteorology to observations, accurate assessment of total column ozone loss is also possible, yielding a loss of approximately 10 DU at the end of November. Discrepancies remain for HNO₃, N₂O₅, and ClONO₂, which are likely a consequence from

the missing cluster ion chemistry and ion-ion recombination in the EMAC model as well as known issues with the model's NO_y partitioning.

1 Introduction

Solar flares and coronal mass ejections are eruptions on the sun's surface that lead to vastly increased fluxes of high-energy particles. Depending on the position of the Earth relative to the ejection, the particles can reach the earth's atmosphere. The phenomenon is then called a solar proton event (SPE). An SPE causes ionizations, dissociations, dissociative ionizations, and excitations in the atmosphere. This results in the production of HO_x and NO_y in the middle and upper atmosphere (Brasseur and Solomon, 1986; Jackman and McPeters, 2004). The altitude profile of the production is mainly determined by the type and number of precipitating particles and their energy distributions, which then also determines the SPE's significance in terms of associated ozone depletion through catalytic destruction by the HO_x and NO_x products. Such events are interesting natural experiments that allow us to test and improve our understanding of atmospheric chemistry, our observational capabilities, and numerical models.

The effects of SPEs on the middle atmosphere have been measured and modelled extensively for almost four decades, initiated by Weeks et al. (1972). Effects on NO_x chemistry were first presented by Heath et al. (1977). Recently, a review on observational and modeling efforts was presented by Jackman and McPeters (2004). The fourth largest period of SPEs in the past 40 years occurred in October/November



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2003 and has been termed Halloween events and received considerable attention. Jackman et al. (2005) investigated both short-term and long-term effects of the Halloween SPEs. 2-D model simulations were compared to NOAA16 SBUV/2 ozone and UARS HALOE NO_x measurements. In the lower and upper mesosphere, short-term ozone depletion up to 50% and greater than 70%, respectively, were found in the model results. Measurements showed a loss of approximately 40% in the lower mesosphere. Northern Hemisphere polar total ozone reduction greater than 0.5% was predicted to last for 8 months. SCIAMACHY/ENVISAT measurements of the ozone depletion were shown by Rohen et al. (2005) and Rohen et al. (2006). In addition, Rohen et al. (2005) compared the ozone depletion with results from a 2-D chemistry transport model based on SLIMCAT. In the five weeks following the SPEs the ozone depletion was captured fairly well by the model and agrees approximately with the model results of Jackman et al. (2005).

The MIPAS instrument on board ENVISAT has been used extensively to study effects of the Halloween SPE on the chemical composition of the middle atmosphere. For example, effects on NO_x and ozone are shown in López-Puertas et al. (2005a), von Clarmann et al. (2005) shows HOCl and ClO perturbations, HNO₃, N₂O₅, and ClONO₂ changes are discussed in López-Puertas et al. (2005b), and N₂O enhancements are discussed by Funke et al. (2008). Jackman et al. (2008) noted that MIPAS measurements of the HNO₃ enhancements were unexpectedly large; Verronen et al. (2008) attributed them to ion-ion recombination between NO₃⁻ and H⁺.

Jackman et al. (2007) discussed dynamical effects in the mesosphere using results from the TIME-GCM model. SPE-induced cooling up to 2.6 K in the lower mesosphere and heating up to 2 K at 90 km were found. Background wind velocities changed up to 25%.

The present study discusses the immediate effects of the Halloween SPEs using results from simulations with the ECHAM5/MESSy model system with an additional sub-model that parameterizes the effects of SPEs in the middle atmosphere. The model results are evaluated using data from the MIPAS instrument. Disagreements between model and satellite data are attempted to be reconciled using several different options.

The model and the satellite instrument are described in Sects. 2 and 3, respectively. The simulated time evolution of several trace gases affected by the SPE as well as comparisons with the satellite observations are described in Sect. 4. Options for improvements to the parameterization are presented, and the remaining discussion is based on the parameterization that yields the best results when compared to the MIPAS measurements. Finally, a summary and conclusions are presented in Sect. 5.

2 Model description

2.1 ECHAM5/MESSy

The ECHAM/MESSy Atmospheric Chemistry (EMAC) model is a numerical chemistry and climate simulation system that includes sub-models describing tropospheric and middle atmosphere processes and their interaction with oceans, land and human influences (Jöckel et al., 2006). It uses the Modular Earth Submodel System (MESSy, see Jöckel et al., 2005; Jöckel et al., 2010) to link multi-institutional computer codes. The core atmospheric model is the 5th generation European Centre Hamburg general circulation model (ECHAM5, Roeckner et al., 2006). The model has been shown to consistently simulate key atmospheric tracers such as ozone (Jöckel et al., 2006), water vapour (Lelieveld et al., 2007), and lower and middle stratospheric NO_y (Brühl et al., 2007). For the present study we applied EMAC (ECHAM5 version 5.3.02, MESSy version 1.8⁺) in the T42L90MA-resolution, i.e. with a spherical truncation of T42 (corresponding to a quadratic Gaussian grid of approximately 2.8 by 2.8 degrees in latitude and longitude) with 90 vertical hybrid pressure levels up to 0.01 hPa. This part of the setup matches the model evaluation study by Jöckel et al. (2006). Enabled submodels are also the same as in Jöckel et al. (2006) apart from the new submodels SPE and SPACENOX, and the sub-submodel FUBRad (Nissen et al., 2007), a high-resolution short-wave heating rate parameterization. The submodel SPACENOX, dealing with NO_x production from low-energy electrons in the upper atmosphere, is described in the companion paper (Baumgaertner et al., 2009), SPE is described here. The chosen chemistry scheme for the configuration of the submodel MECCA (Sander et al., 2005) is simpler compared to the configuration in Jöckel et al. (2006). For example, the NMHC (non-methane hydrocarbon) chemistry is not treated at the same level of detail. The complete mechanism is documented in the supplement.

The basic state of the model atmosphere is determined by a set of boundary conditions in order to make comparisons with observations more meaningful. Sea surface temperatures are taken from the Met Office Hadley Centre (HadISST), the equatorial zonal wind is weakly relaxed towards observed winds to yield the correct phase of the QBO. Long lived trace gases are relaxed towards observed concentrations at the surface, short lived species are emitted into the lowermost model layers (Baumgaertner and Brühl, 2008). The model can be nudged towards analyzed meteorology (ECMWF ERA-40), allowing direct comparisons with measurements. Results from setups that were nudged as well as from free-running setups will be shown here. Nudging was applied between levels 18 (~ 1.5 hPa) and 84 (~ 715 hPa) with relaxation e-folding times of 12 h for temperature, 6 h for vorticity, 48 h for divergence and 12 h for surface pressure. Below (down to level 13 or ~ 0.7 hPa) and above (up

to level 87 or ~ 920 hPa) these levels stepwise reduced coefficients were applied.

2.2 The submodel SPE

The new submodel with the name SPE parameterizes the effects of energetic protons by calculating ionization rates and, subsequently, production rates of NO_x and HO_x . The results are added to the model tendencies of NO_x and HO_x .

A column version of the submodel was also implemented which uses the same submodel core layer files (for a discussion of the MESSy structure see Jöckel et al., 2005) as the 3-D model, but a separate interface and a simple time control. It is not coupled to any other MESSy submodels.

For the calculation of ionization rates a method based on Vitt and Jackman (1996) was employed. The proton flux measurements by GOES (Geostationary Operational Environmental Satellite) or IMP (Interplanetary Monitoring Platform) are interpolated to a logarithmic energy spectrum. The energy deposited in any model layer is calculated directly using the thicknesses of the considered layer and an energy-range relationship. Here, we used the relationship by Bethe and Ashkin (1953),

$$R = (E/a)^b \quad (1)$$

where R is the range in meters (see Vitt and Jackman, 1996) and E is the particle energy measured in MeV. Bethe and Ashkin (1953) determined $a=9.3$ and $b=1.8$. In order to evaluate the ionization rate calculation, the SPE submodel was employed in column mode and the simulation results were compared to published ionization rates. For this, daily ionization rates provided by Jackman (2006) for the SOLARIS working group of SPARC (Stratospheric Processes and their role in climate, a project of the WCRP – World Climate Research Programme) were used as a reference. Resulting ozone depletion and other SPE effects can then be compared without large uncertainties in the underlying ionization rates. Figure 1 depicts the column model ionization rates (black lines) averaged over 28 October 2003 (dashed line) and 29 October (solid line) using hourly model output, and the data by Jackman (2006) (blue lines) for the same days. Although there is general agreement, the vertically integrated ionization rate averages for 29 October differ by 16%. In particular, the ionization rate altitude dependence is not satisfactory; at 45 km on 29 October for example an overestimation of approximately 50% is found. However, a correct altitude dependence is crucial due to the steep profiles of ozone and other constituents that the SPE affects. Therefore, a series of sensitivity studies was performed using different sets of a and b in Eq. 1. Values for a between 9 and 14, for b between 1 and 3 were tested. Using $a=12.3$ and $b=1.8$ yield the best agreement between the calculated and published ionization rates reducing the difference between the vertically integrated rates to 5%. The results are shown as the red lines in Fig. 1. Because of the better agreement

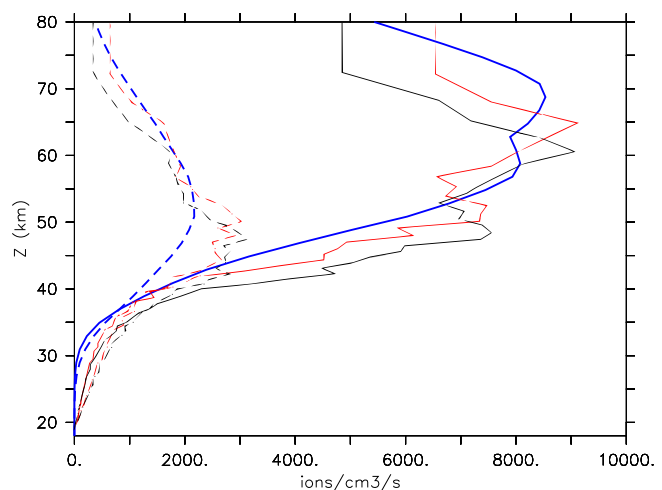


Fig. 1. Ionization rates for 28 and 29 October 2003 (dashed and solid lines, respectively) calculated by a column mode version of the SPE submodel using the original (black) and optimized (red) parameters for the energy-range relationship. Results provided by Jackman (2006) are shown in blue.

with the results by Jackman (2006) who used a more complex approach, this set of parameters was employed for the rest of this study. Note that the agreement above 50 km is affected by the different resolution of the data sets: the vertical resolution of EMAC in the L90 setup degrades to 8 km at the highest altitude.

Instead of using internally calculated ionization rates, externally calculated rates, 1-D or 3-D, generally with a lower time resolution compared to the online calculation, can also be applied. This option was however not used in the present study.

Since the magnetic field lines guide the charged particles toward the geomagnetic poles, full ionization was only applied poleward of 60° geomagnetic latitude. The production of N and NO was parameterized using commonly cited values of 0.55 N atoms and 0.7 NO molecules per ion pair (e.g., Jackman et al., 2005) because the model does not incorporate excited state chemistry of atomic nitrogen. However, in an attempt to improve the agreement between satellite and model results, we tested a height dependent N/NO production efficiency which is described in Sect. 4.1. The more complicated production of H and OH by ionization was parameterized using an approximation to data published in Solomon et al. (1981). HO_x production per ion pair depends on altitude and the baseline ionization rate. The altitude dependence was approximated as

$$f(h) = 2 - \exp((h - 83)/6), \quad (2)$$

where h is the altitude in km. Using thresholds $t(q)$ as in Solomon et al. (1981) for the baseline ionization rate dependence, the production rates H^{prod} and OH^{prod} are then

$$(\text{H}^{\text{prod}}, \text{OH}^{\text{prod}}) = 0.5 \cdot q \cdot f(h) \cdot t(q) \quad (3)$$

Table 1. Baseline versions for spectra version V3O of the employed MIPAS data.

O ₃	HNO ₃	N ₂ O ₅	NO ₂	N ₂ O	HOCl	ClONO ₂
7, 9	8, 9	9, 10	9, 11	12	3	11, 12

where q is the ionization rate. Note that $f(h)$ cannot become negative, therefore Eq. 2 cannot be used for altitudes greater than approximately 87.2 km.

2.3 The CAABA boxmodel

We also performed simulations with the atmospheric chemistry box model CAABA (Chemistry of the Atmosphere As a Box model Application, version 2.4d), described in Sander et al. (2010). CAABA contains the chemistry module MECCA (Sander et al., 2005) and the photolysis module JVAL (based on Landgraf and Crutzen, 1998), which are also part of EMAC, therefore the chemical and photochemical schemes are almost identical to the 3-D simulations. The box model does not contain heterogeneous chemistry, and the radiation environment is different in that a climatological ozone profile is assumed.

3 Satellite observations

The Halloween SPE effects on atmospheric chemistry have been observed by instruments on board the Environmental Satellite (ENVISAT). Here, we present measurements from the limb viewing spectrometer MIPAS (Michelson Interferometer for Passive Atmospheric Sounding) to assess if the SPE-induced production of trace gases and subsequent ozone depletion as seen in the model output are realistic. MIPAS observations of the Halloween SPE have previously been presented for example by López-Puertas et al. (2005a), López-Puertas et al. (2005b), von Clarmann et al. (2005) and Funke et al. (2008). Here, we use the retrievals performed with the IMK-IAA data processor and the Karlsruhe Optimized and Precise Radiative Transfer Algorithm (KOPRA), which were also used in the aforementioned MIPAS studies.

Only MIPAS spectra version V3O were employed. The employed baseline versions are listed in Table 1. The differences between the employed baseline versions are small and generally simply cover different days of the concerned period. When several versions were available for a specific day, the newest version was employed.

The MIPAS measurement errors and vertical resolution for a single profile are listed in Table 2 for 40 km and 60 km altitude. Note that if an ensemble of measurements is considered, the precision of the mean (also termed random error) is reduced by a factor equal to the square root of the number of measurements. The systematic error and the vertical resolution do not change with the number of measurements. Ad-

ditionally, the standard error of an ensemble mean needs to be considered, which accounts for natural variability as well as measurement precision. During highly disturbed conditions, such as for NO₂ during the initial phase of an SPE, the standard error can be much larger than the precision. The sampling “error”, i.e. the error related to the fact that only a limited number of measurement points in time and space of a “true distribution” exist, has not been quantified. Note, however, that due to the relatively homogeneous spread of MIPAS profiles over the polar area the difference between the measured and modelled distributions should be small. For the sampling in time, only species with a large diurnal cycle are likely to be adversely affected by this as discussed below.

To facilitate comparison with the model results, the MIPAS data were binned to create a daily time series. The data were also gridded onto the EMAC model grid used here (T42). No restrictions were made with respect to solar zenith angle limits and both day and night data were used. For species with a strong diurnal cycle, here especially NO₂, this is a potential error source if the fraction of measurement points made during sunlit conditions is different between model and measurements. For example, for NO₂ we found for 31 October in the region poleward of 70° N a daytime fraction of 10% for EMAC, and 13% for MIPAS, both calculated from the solar zenith angle. Differences of the same magnitude persist until mid-November when the area north of 70° N remains completely dark. Therefore, in the first half of November this introduces a negative bias for MIPAS NO₂ compared to EMAC, as more profiles were obtained in daylight when NO₂ is negligible.

In the vertical, the original geometric height grid was adopted as is. For comparison with the MIPAS data, the model output was transferred onto the same vertical grid using the simulated geopotential height.

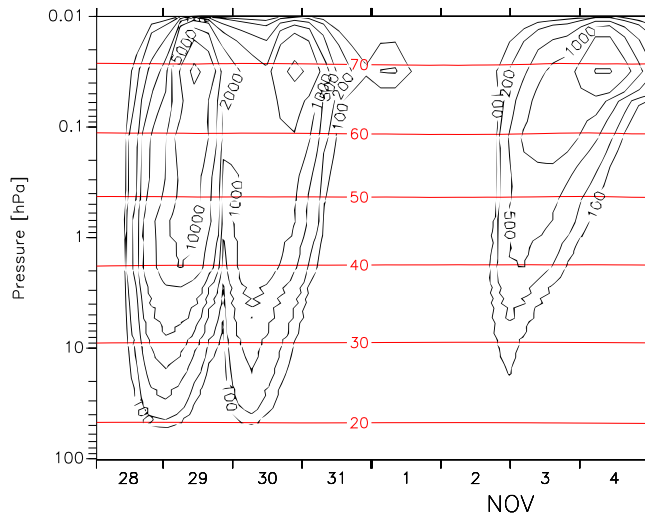
The vertical resolution of the MIPAS data is generally lower than the model vertical resolution. Therefore, the model results were convolved with the corresponding MIPAS averaging kernel (AK). The supplement contains figures with and without convolution with the MIPAS AK.

4 Results and discussion

The SPE submodel has been tested with EMAC for the time of the Halloween SPEs between 28 October and 4 November 2003. For the most intense period, 28–31 October 2003, the zonally averaged (70°–90° N) ionization rate profiles are depicted as a function of time in Fig. 2. The major episode of the SPE starts at approximately 12:00 UTC on 28 October 2003. Above 40 km altitude the ionization rate exceeds 10 000 ion pairs cm⁻³ s⁻¹ for several hours on 29 October 2003. Less intense periods follow on the 30 October and 2–4 November. There appears to be a very good agreement with the ionization rates calculated by Jackman et al. (2005), see their Fig. 3.

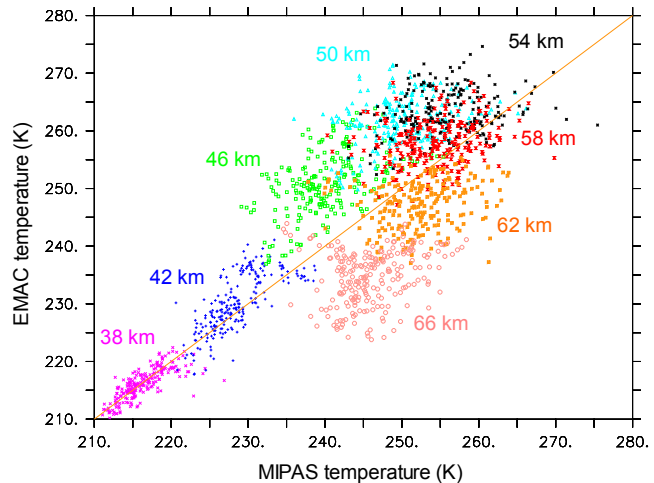
Table 2. MIPAS measurement error contributions and vertical resolution.

	Single profile precision for 70–90° N; given in vmr and percent under quiet and disturbed (in brackets, 30 October–5 November) conditions		Systematic error		Vertical resolution (km) for 70–90° N	
	40 km	60 km	40 km	60 km	40 km	60 km
O ₃	0.26 ppmv/6(6)%	0.11 ppmv/10(14)%	0.33 ppmv	0.0067 ppmv	4	9
HNO ₃	0.15 ppbv/9(7)%	0.37 ppbv/42(22)%	0.30 ppbv	0.32 ppbv	8	43
N ₂ O ₅	0.18 ppbv/18(40)%	0.041 ppbv/54(90)%	0.064 ppbv	0.0081 ppbv	7	52
N ₂ O	0.63 ppbv/45(67)%	0.46 ppbv/26(15)%	0.027 ppbv	0.45 ppbv	4	13
NO ₂	1.7 ppbv/11(13)%	4.6 ppbv/9(10)%	0.86 ppbv	5.1 ppbv	7	13
HOCl	0.12 ppbv/63(44)%	0.0019 ppbv/70(50)%	0.0013 ppbv	0.000049 ppbv	15	17
ClONO ₂	0.13 ppbv/17(14)%	0.02 ppbv/40(31)%	0.069 ppbv	0.0063 ppbv	13	16

**Fig. 2.** Calculated high-latitude (70°–90° N zonal average) ionization rates ($\# \text{cm}^{-3} \text{s}^{-1}$) during October and November 2003. The red lines denote the altitude in km.

In addition to the simulation with the SPE submodel switched on (denoted S-SPE in this paper), one simulation was performed without the submodel (denoted S-NOSPE). This allows to accurately assess the impact of the SPE by comparing the two simulations with each other. Both simulations were nudged towards the observed meteorology in the troposphere (see Sect. 2), so differences in the dynamics of the two simulations are small.

Before we present an analysis of trace gases that are affected by the Halloween SPEs, we discuss the simulated and observed temperatures before and during the events. Because many reactions involving NO_y species are temperature dependent, it is important to know how accurately the temperature is simulated by the model. In Fig. 3 the EMAC temperatures between 26 October and 30 November 2003 are directly compared to MIPAS measurements at altitudes between 38 and 66 km at 4 km intervals. All altitude bins can be distinguished by a different colour and symbol and are

**Fig. 3.** Comparison of zonal average temperatures of MIPAS and EMAC for 70–90° N for 26 October–30 November 2003.

labelled accordingly. In the upper stratosphere between 46 and 54 km a high bias of approximately 10 K can be noted (the model stratopause is situated at approximately 53 km, not shown). Higher up, the model shows a low bias increasing with altitude to 10 K at 66 km. These biases indicate that the model stratopause is lower than in the observations and are likely to affect the comparisons of simulated and measured trace gas concentrations, therefore sensitivity studies using a boxmodel with fixed temperatures are presented in Sect. 4.1.

Atomic nitrogen (N) and nitric oxide (NO) are among the direct products of the SPE, and the effects on NO_x are discussed and evaluated next. Figure 4 depicts changes in NO₂ with respect to 26 October for 70–90° N from MIPAS measurements (Fig. 4a) and the corresponding model results with the MIPAS NO₂ averaging kernel applied (Fig. 4b). Note that day and night data were used to create this figure and similar figures below.

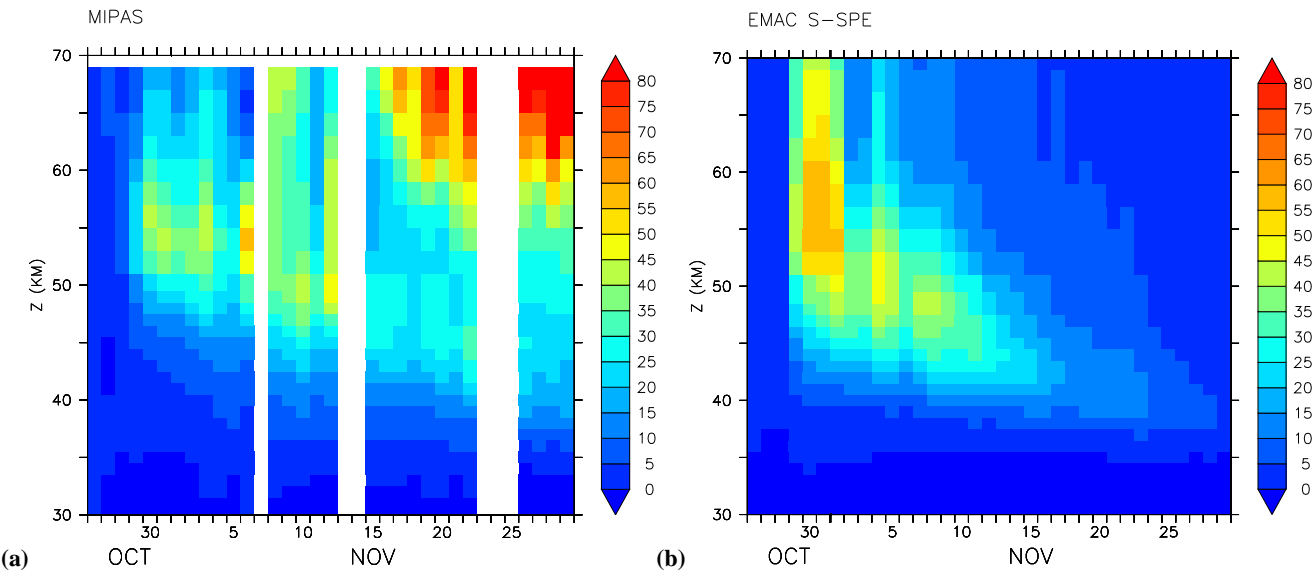


Fig. 4. NO₂ change (ppbv) with respect to 26 October for 70–90° N for (a) MIPAS, (b) EMAC simulation S-SPE.

During the SPE, MIPAS shows lower mesospheric enhancements up to 50 ppbv where the model reaches 60 ppbv. During the initial phase of the SPE the NO_x enhancements are centered around the geomagnetic pole, then gradually transported and mixed to other areas including lower latitudes (not shown). This leads to a very inhomogeneous spatial distribution of NO_x and thus large sampling errors (see Sect. 3). Also note that the systematic error of NO₂ at 60 km is 5.1 ppbv (see Table 2). In the light of these aspects, the agreement between EMAC and MIPAS is acceptable. Higher up the model overestimates the enhancements by up to 100%. In the following weeks downward transport of the enhancements can be seen in both the model results and the measurements. Note that the enhancements in the MIPAS data in the upper mesosphere during the second half of November are not related to in-situ production of NO_x but originate in the lower thermosphere and are transported downward. This effect is not subject of the work presented here, but is discussed in the companion paper by Baumgaertner et al. (2009). Also note that on 6 and 7 November only 5 profiles were obtained by MIPAS in this area, with an ensemble mean precision of 1.5 ppbv at 50 km. Similarly, on 12 and 13 November only 6 profiles were obtained, with an ensemble mean precision of 1.0 ppbv. Therefore the enhancements during both these periods are probably not representative for the full polar cap, given that flux measurements did not show any anomalies during these times.

The produced NO_x can partially react to form N₂O as has been shown e.g. by Funke et al. (2008). Figure 5 depicts N₂O changes in MIPAS and model data similar to the NO₂ changes shown above. It is evident that the model production of N₂O by far exceeds that of the enhancements observed

by MIPAS. The measurements show enhancements below 5 ppbv, while in the model 35 ppbv are reached at 60 km (not discernable from Fig. 5 due to the saturated scale which was chosen such that the MIPAS measurements can be analysed). This is a major deficiency of the model performance during this event and has been found in other model simulations as well (Funke et al., 2008). Before the effects on other species are discussed, possible options for improvements to the model or the SPE parameterization are explored in the next section.

4.1 Model improvements

The large discrepancy between simulated and observed N₂O could be due to various reasons:

1. reactions not accounted for or unknown, or inaccurate reaction rates,
2. incorrect external parameters that influence the reactions, such as temperature, or
3. inaccurate assumptions about SPE production of N/NO.

A similar discrepancy between MIPAS N₂O and the Canadian Middle Atmosphere Model (CMAM) during the Hal-loween storm period was reported by Funke et al. (2008). They attempted to resolve this problem by adding to the N₂O-forming reaction (reaction G3107 in the MECCA mechanism, see supplement.)



the branches



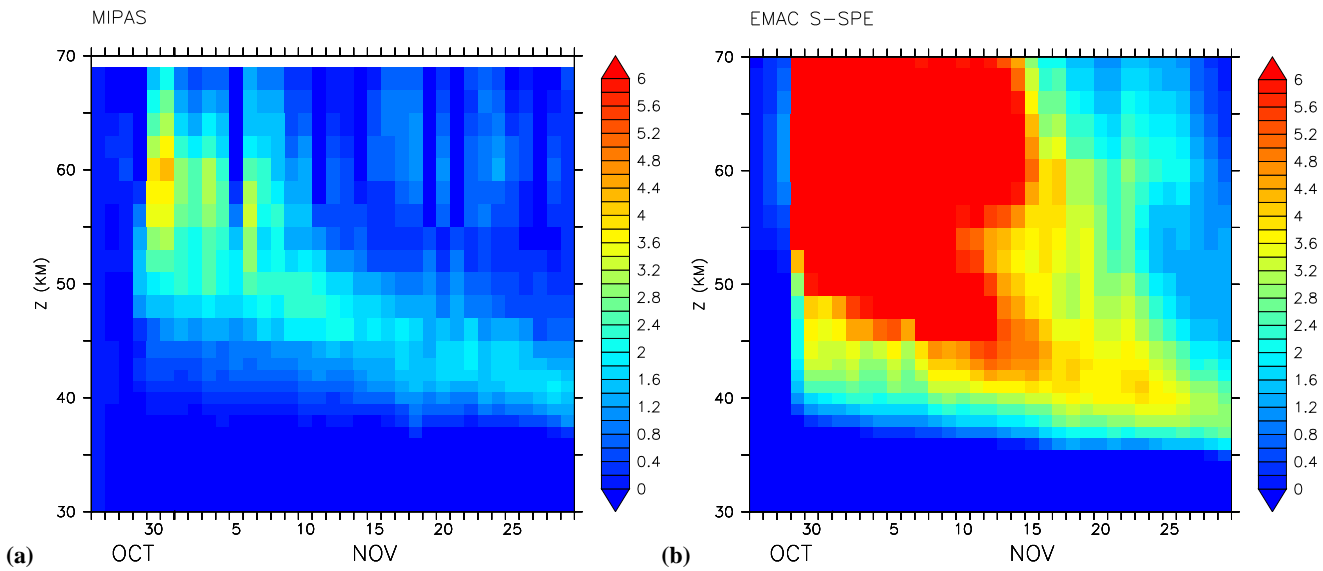


Fig. 5. Same as Fig. 4 but for N_2O (ppbv).



with 50% of the reaction rate going into the primary branch R1, and 25% each for the Reactions (R2) and (R3) chosen such that the model better reproduces the observations. The same modification was implemented in EMAC (see supplement) and the equivalent simulation to S-SPE was performed, termed S-SPE-FUNKE hereafter. The results for N_2O are shown in Fig. 6. There is generally less N_2O compared to the S-SPE simulation discussed above (Fig. 5), however, a large overestimation persists.

The second possibility listed above refers to external parameters that influence the production of N_2O . The Reaction (R1) that forms N_2O after the SPE is temperature dependent with a rate coefficient of

$$k = 5.8 \times 10^{-12} \cdot e^{220/T} \quad (4)$$

(see supplement), therefore the temperature discrepancy found above (see Fig. 3) could explain some of the overestimations of N_2O . This hypothesis is tested by performing sensitivity simulations of the chemistry only. For this, the boxmodel CAABA, described in Sect. 2.3, is used. CAABA was initialized with the temperature and tracer mixing ratios of the corresponding spatial and temporal location in the 3-D simulation. Note that the temperature is kept constant throughout the CAABA simulation. The solar proton event was simulated by applying the ionization rates that were calculated in the 3-D study. Temperature sensitivity of the reactions was explored by offsetting the temperature T_{caaba} with respect to the 3-D simulation $T_{3\text{-D}}$. As an example, the resulting N_2O mixing ratios for a CAABA simulation for 60 km altitude, 0° E, and 75° N are shown in Fig. 7 for the temperature $T_{3\text{-D}}$ (black), as well as for $T_{3\text{-D}} + 10\text{ K}$ (red) and

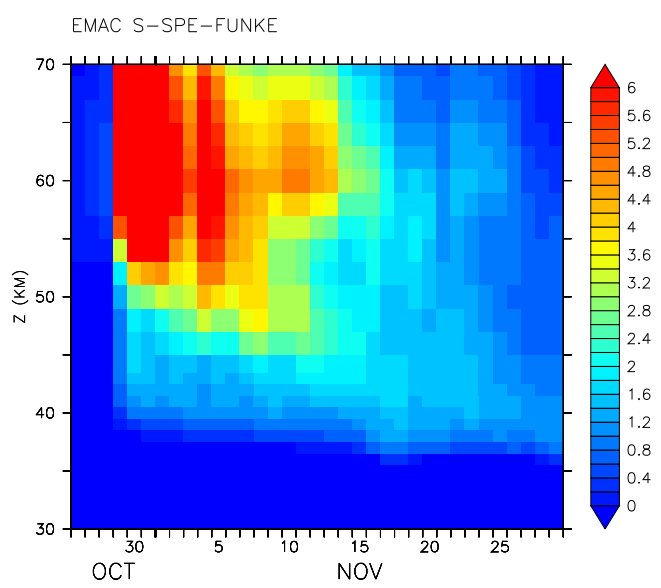


Fig. 6. N_2O change (ppbv) in EMAC simulation S-SPE-FUNKE with respect to 26 October for $70\text{--}90^\circ\text{N}$.

$T_{3\text{-D}} + 20\text{ K}$ (green). The crosses denote MIPAS measurements in the area $75^\circ\text{N} \pm 5^\circ$ latitude.

As expected from the 3-D simulation with the same chemical mechanism, there is a sharp increase of N_2O during the SPE, but no change in the following weeks since there is no N_2O photolysis and no O^1D present to deplete N_2O . A significant temperature dependence of the amount of N_2O produced is evident. However, mixing ratios are still strongly overestimated when compared to the MIPAS observations.

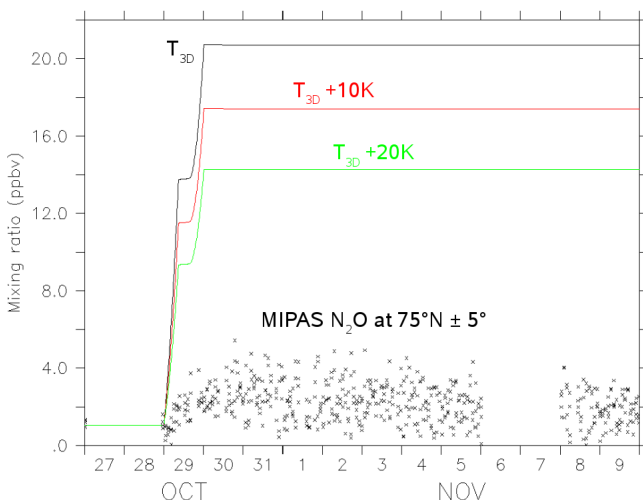


Fig. 7. Mixing ratios of N_2O at 60 km, 75°N from the CAABA box model at three different temperatures (lines), and MIPAS N_2O measurements for the same latitude region.

Finally, we discuss the third possibility, referring to inaccurate assumptions about the production efficiency of N and NO per ion pair.

We have chosen this branching ratio as tuning parameter because neither experiments nor theoretical studies have provided accurate values by now. Values ranging from 0.33 to 2.5 for the total production of N and NO have been reported in previous work. Thus, its adjustment by means of a fit to MIPAS data is appropriate as long as the resulting branching ratio does not significantly exceed the spread of values reported in the literature. It should be noted, however, that we do not aim at providing a scientific justification for modification of this parameter (which would be beyond the scope of this paper) but at identifying potential error sources responsible for the model/data mismatch and – in this sense – to motivate future experimental studies.

The method presented here, which is used to reconcile N_2O concentrations, appears to be a manifest approach because the $0.55 \text{N}^4\text{S}/0.7 \text{NO}$ branching ratio commonly used has only been derived by “best fit” between models and observations in the past. We support this statement with the following discussion of the involved processes and of previous studies on this subject.

Impacting protons with energies from 100 keV to 1 GeV that reach the mesosphere or stratosphere are too energetic to be efficient in dissociating nitrogen (Crutzen et al., 1975), but deposit their energy in the accessible states of N_2 and O_2 by secondary electrons of low energy ($< 100\text{eV}$), see Porter et al. (1976). These act on N_2 and O_2 to initially produce N , N^+ , N_2^+ , O , O^+ , and O_2^+ ; for details on these reactions see e.g. Rusch et al. (1981). Further reactions lead to the production of N^4S , N^2D , and N^+ (which via reaction with O_2 also reacts further to N^2D) as well as other electroni-

cally excited states of N and O probably of less importance (for nitrogen: N^2P , $\text{N}(3s^2\text{P})$, $\text{N}(3s^4\text{P})$, $\text{N}(3s^2\text{D})$, $\text{N}(4s^2\text{P})$). Note that several of the involved reaction rates and branching ratios have not been experimentally determined (Rusch et al., 1981). The following reactions then lead to the formation of NO:



Reaction (R4) is slow and highly temperature dependent, while Reaction (R5) is fast and thus the main source for NO.

Additionally, N^2D can react with O or NO:



Additional to the Reaction (R1), N^4S also takes part in the following reactions:



Many atmospheric chemistry models do not include ion or detailed excited state chemistry, so in order to parameterise solar proton events, the formation of ground state N and NO is required as a function of ionization rate. However, because of reactions (R5) and (R6) it is not possible to assume an NO production rate normalized to the ionization rate, as pointed out e.g. by Rusch et al. (1981). We would also like to point out that the branching ratio $\text{N}^4\text{S}/\text{N}^2\text{D}$ is not necessarily equal to the ratio $\text{N}^4\text{S}/\text{NO}$ because of the additional reactions listed above.

Crutzen et al. (1975) mentioned that from the experimental studies of Winters (1966), Rapp et al. (1965), and Rapp and Englander-Golden (1965) a rate of production of 1.5 N atoms per ion pair can be estimated. However, the ratio $\text{N}^4\text{S}/\text{N}^2\text{D}$ or $\text{N}^4\text{S}/\text{NO}$ was not determined. Porter et al. (1976) write that additional studies concerning atomic nitrogen production were limited to high energy secondary electrons ($> 100\text{ keV}$) and are thus not applicable for SPEs. The work of Porter et al. (1976) is the only theoretical study that used an atomic cross section approach to derive a production of 1.27 atoms per ion pair for proton impact at 1 MeV, distributed over 8 different states including N^+ (see their Table V). While the ratio of $\text{N}^4\text{S}/\text{N}^2\text{D}$ can be deduced from this, it is not possible to accurately derive the N/NO production ratio. It is also interesting to note that Porter et al. (1976) found that, “it is observed that the efficiency for production of atomic species changes with the particle energy”, which effectively means that such production at least for proton energies below 1 MeV is height dependent since the particle energy determines the height it reaches.

As discussed in the following, newer SPE studies generally base their branching ratios either on approximations to Porter et al. (1976), often assuming that the ratio N(⁴S)/NO ratio is equal or similar to the N(⁴S)/N(²D) ratio, or derive such branching ratios by fitting their model results to observations. A review article by Jackman and McPeters (2004) states “Estimates of the number of NO_y constituents created per ion pair range from 0.33 (Warneck, 1972) up to 2.5 (Fabian et al., 1979). Recent publications show only small differences and range from 1.25 (Jackman et al. 1990) up to 1.3 (Reid et al., 1991) NO_y constituents produced per ion pair.” The study by Fabian et al. (1979) based their assumption on rocket measurements of mesospheric and thermospheric NO concentrations as well as ionisation rates. The work by Warneck (1972) is a modeling study based on production rates of 0.64 for N₂⁺ and 0.16 for N⁺ as published by Dalgarno (1967). However, Dalgarno (1967) derived from the initial N₂⁺ and N⁺ creation a production of 0.8 N and 0.5 NO “with a possible doubling of [the production rate of] NO at night”, which is in disagreement with the 0.33 NO_y constituents derived by Warneck (1972).

Reid et al. (1991) indeed use a value of 1.3 NO_y molecules per ion pair: “Following Rusch et al, 1981, we assume that 1.3 NO_y molecules are produced per ion pair, partitioned between the ground N(⁴S) state and exited states (mainly 2-D) at 20 and 80%.” The Rusch et al. (1981) paper is a theoretical study that includes a comparison of model results and observations, however, they base their results on several assumptions that were not experimentally verified, and their conclusions are also build on “best fit” to observations: “A production of 1.3 odd nitrogens per ion par, an 80% branching ratio for the production of N(²D) by the primary reactions dissociating N₂, and inclusion of the loss of NO_x through N+NO in a time-dependent calculation appears to produce an ozone destruction most compatible with the measurements.” The other study cited in Jackman and McPeters (2004) is Jackman et al. (1990). They state “This calculation assumed 1.25 nitrogen (N) atoms produced per ion pair which is similar to the value given by Porter et al. (1976), derived using a detailed theoretical energy degradation computation. The agreement between the predicted and measured NO increase following the July 1982 SPE has given us confidence in the reliability of the computations for NO_y species increase caused by SPEs. We therefore assume that 1.25 N atoms are produced per ion pair for all base model computations in this paper.”

The N/NO branching is discussed in a newer modeling study by Jackman et al. (2005) who state “In order to best represent the production of NO_y constituents by the protons and their associated secondary electrons, we assume that 45% of the N atoms produced per ion pair result in the production of N(⁴S) (\sim 0.55 per ion pair) and that 55% of the N atoms produced per ion pair result in the production of NO (\sim 0.7 per ion pair).” It appears that this choice is also based on a best fit to observations of NO_y.

We conclude, that established N(⁴S)/NO ratios are not firmly supported by theoretical or experimental work or are inconsistent.

However, the observed discrepancy of N₂O could possibly be resolved by changing the N/NO branching ratio since the amount of produced N₂O is mostly dependent on the amount of available N atoms. Reducing the number of N atoms produced by an ion pair would therefore yield better results for N₂O. Since the 3-D model is computationally too expensive to perform a large number of tests of other N/NO production efficiencies, the following approach was chosen. The box-model CAABA, described in Sect. 2.3, was used to simulate the SPE at MIPAS measurement locations for N and NO production per ion pair ranging from 0 to 1.2 using steps sizes of 0.05. There are approximately 200 MIPAS profiles of N₂O and NO₂ between 31 October and 2 November north of 70°N available, all of which were included in this simulation set. Such a short period was chosen in order to avoid strong contributions by vertical transport, which cannot be captured by CAABA. The simulations were performed at 4 km height intervals starting at 42 km.

For each simulation CAABA was initialized at 27 October, one day before the SPE, with the temperature and tracer mixing ratios of the corresponding spatial and temporal location in the 3-D simulation. As in the CAABA simulations above, the SPE was simulated by applying the ionization rates that were calculated in the 3-D study. The simulation was stopped at the time step closest to the MIPAS measurement. Then the percentage deviations from the MIPAS values of both species were calculated. From all deviations the median was calculated separately for NO₂ and N₂O. Finally, the two medians were averaged using equal weighting. The entire simulation and processing chain is depicted in Fig. 8. It also shows the average deviations for 58 km altitude. From this, the N and NO productions per ion pair minimizing the total deviation was determined. Figure 9, left, shows the resulting production per ion pair for N (filled circles) and NO (filled squares), also indicating the remaining total deviation in percent. On the right, Fig. 9 shows the ratio of N/NO production.

At 42 km, the remaining total deviation is still large (83%), but is below 10% at 46, 50, and 58 km. At 62 km, where the model vertical resolution and the agreement of the ionization rates (see Fig. 1) degrade, deviations rise again up to 40%. Generally, there appears to be a strong height dependence, with an almost negligible production of N atoms below 45 which is in disagreement with the generally assumed production parameterization of 0.55 N atoms. Above 55 km the production of NO is also in disagreement with the previously used value of 0.7. This is mainly due to the fact that MIPAS yields only small increases of NO₂ and N₂O at these altitudes, despite the large ionization rates present there. This discrepancy can only partially be explained by the MIPAS averaging kernels for NO₂ and N₂O and warrants further investigation. Note that the production of N and NO per ion pair, as well as the N/NO production ratio appear to

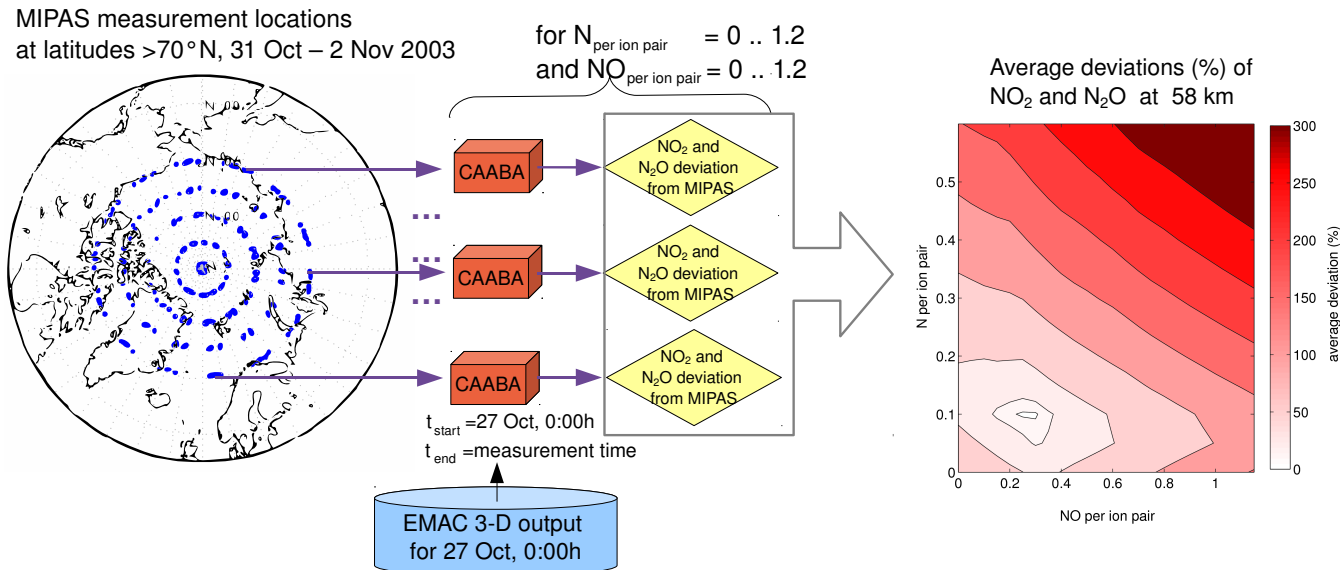


Fig. 8. Datasets, simulations, and postprocessing used to determine the N and NO production per ion pair that fits best to MIPAS observations. The depicted processing chain is performed for each altitude separately.

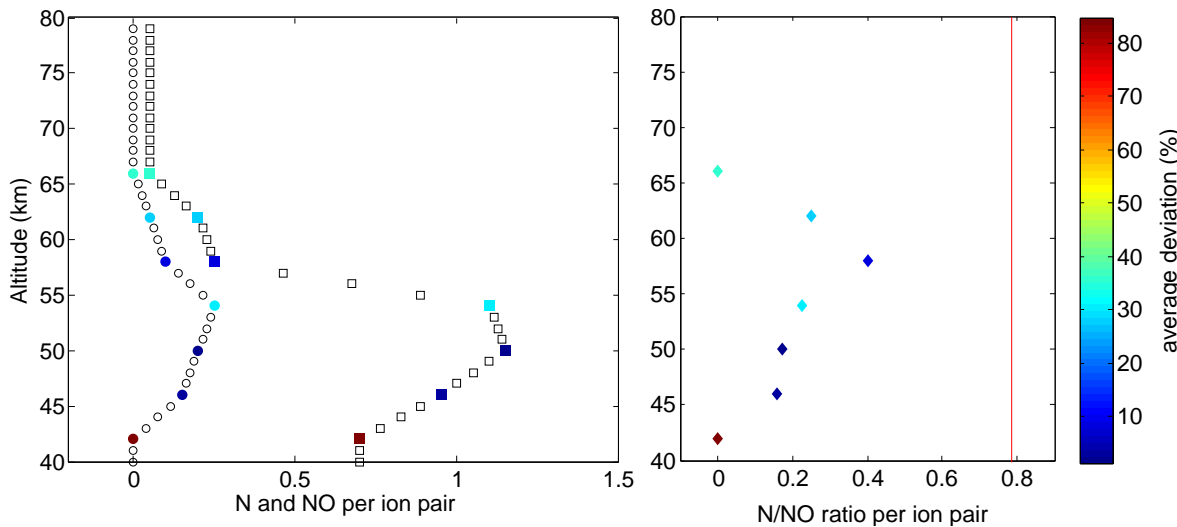


Fig. 9. Left: N (filled circles) and NO (filled squares) production efficiencies determined from CAABA simulations. The colour indicates the average deviation between model and MIPAS NO₂ and N₂O in percent. Empty circles and squares show the linearly interpolated and extrapolated production efficiencies used in the EMAC model simulations S-SPE-NNOEFF. Right: N/NO ratio per ion pair from CAABA simulations (diamonds) and commonly used value (red line).

maximize at approximately the height of the stratopause (see Fig. 3), possibly indicating a temperature dependent production mechanism.

A modified branching ratio of N^{4S} and NO as suggested by our analysis implies that an important amount of energy is lost by exothermic reactions involved in the reaction chain following the particle event, and hence would contribute to chemical heating.

A linear interpolation was performed to obtain N and NO production per ion pair with a resolution of 1 km, also shown in Fig. 9 (empty circles and squares). We use these values subsequently in a 3-D EMAC simulation, hereafter called S-SPE-NNOEFF. Except for these height dependent N/NO production coefficients, the simulation was identical to S-SPE.

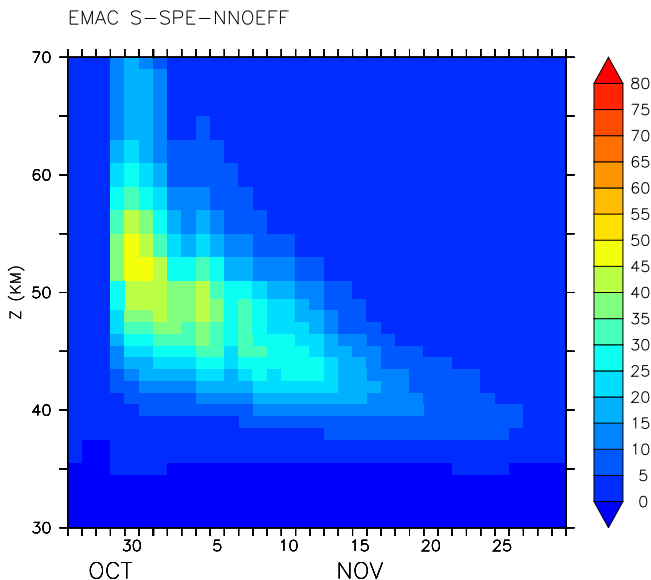


Fig. 10. NO₂ changes (ppbv) for 70–90° N relative to 26 October 2003 for simulation S-SPE-NNOEFF.

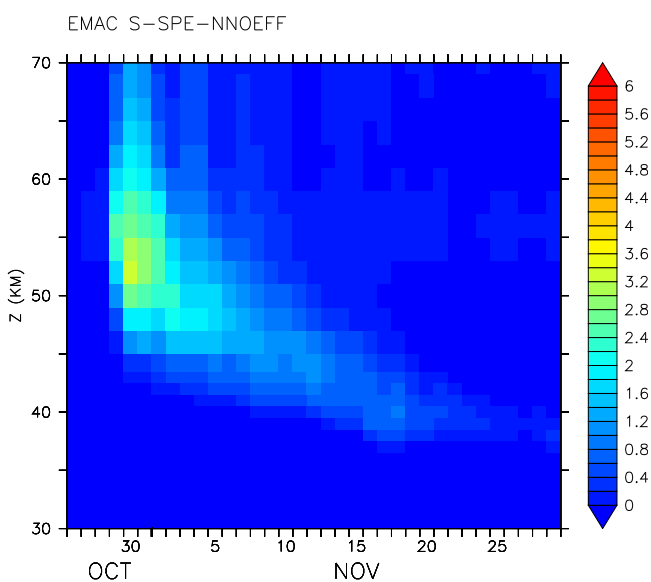


Fig. 11. Same as Fig. 10 but for N₂O changes (ppbv).

4.2 Evaluation of the improved parameterization

The SPE effects on the polar chemistry in the S-SPE-NNOEFF simulation are now evaluated. Firstly, simulated NO₂ (Fig. 10) and N₂O (Fig. 11) are compared to the MIPAS observations similar to the evaluation of the simulation S-SPE above. In the stratosphere and lower mesosphere the location of the peak values of NO₂ with respect to time and altitude is similar for EMAC and MIPAS (compare Figs. 10 and 4a). The magnitude of the first maximum at the end of October reaches approximately 50–60 ppbv. The agree-

ment with MIPAS is better compared to the simulation S-SPE, because the maximum between 55 and 60 km is similarly pronounced in the model and the observations, whereas NO₂ from simulation S-SPE showed a very broad maximum stretching from 50 to 70 km (Fig. 4b).

Note however that especially in the second half of November in the upper stratosphere the model appears to underestimate NO₂. Geomagnetic-activity related NO_x enhancements are at this time mainly found in the mesosphere, indicating that there might be problems concerning the model NO_y partitioning, see also Sect. 4.3.

The enhancements in N₂O shown in Fig. 11 are drastically reduced compared to the simulation S-SPE (Fig. 5b), and are now in satisfactory agreement with the MIPAS observations (Fig. 5a). The maximum during the SPE is located at around 55 km, 5 km below the maximum found by MIPAS. In the observations, N₂O enhancements reach 4.5 ppbv, which is overestimated by the model at approximately 6 ppbv. Note that the precision and standard error of the MIPAS profile ensemble amount to 0.08 ppbv and 0.14 ppbv at 50 km on 31 October, when the N₂O mixing ratio maximises. Assuming that the sampling error is smaller than these errors, there remains a bias between model and measurements.

4.3 Further SPE-induced composition changes

The enhancements of both NO_x and OH results in severe ozone depletion in the polar caps. Fig. 12 shows MIPAS (a) and EMAC S-SPE-NNOEFF (b) ozone changes in percent relative to 26 October 2003. Both MIPAS and EMAC S-SPE-NNOEFF results show an ozone depletion of up to 50% in the upper stratosphere. In the mesosphere, short-term ozone depletion exceeds 50%. In the sunlit Southern Hemisphere mesosphere ozone loss is less severe (not shown), which can be explained by differences in the ambient HO_x production (Rohen et al., 2005) and the shorter lifetime of NO_x. An additional estimate of ozone depletion is obtained by comparing the EMAC S-SPE-NNOEFF and S-NOSPE simulations, shown in Fig. 12c. It is evident that the SPE related ozone depletion is significant and not an artefact of the illustration technique. However, stratospheric ozone depletion due to the SPE appears to be overestimated by up to 20 percentage points, and contributions to ozone loss from intrinsic processes appear to play a non-negligible role.

Ozone depletion is restricted to altitudes above 35 km, thus the expected impact on total ozone content (TOC) is small. The effect of the SPE on TOC can be estimated by comparing EMAC S-SPE-NNOEFF TOC and S-NOSPE TOC for the area north of 70°N, shown as the solid and dashed line, respectively, in Fig. 13. Before the SPE, both simulations show almost identical behaviour of TOC. Deviations such as around 22 October are eliminated by the nudging on the relaxation e-folding timescales. From 28 October onwards, a reduction of approximately 5 DU is evident, growing to 10 DU at the end of November. Note that the dominating

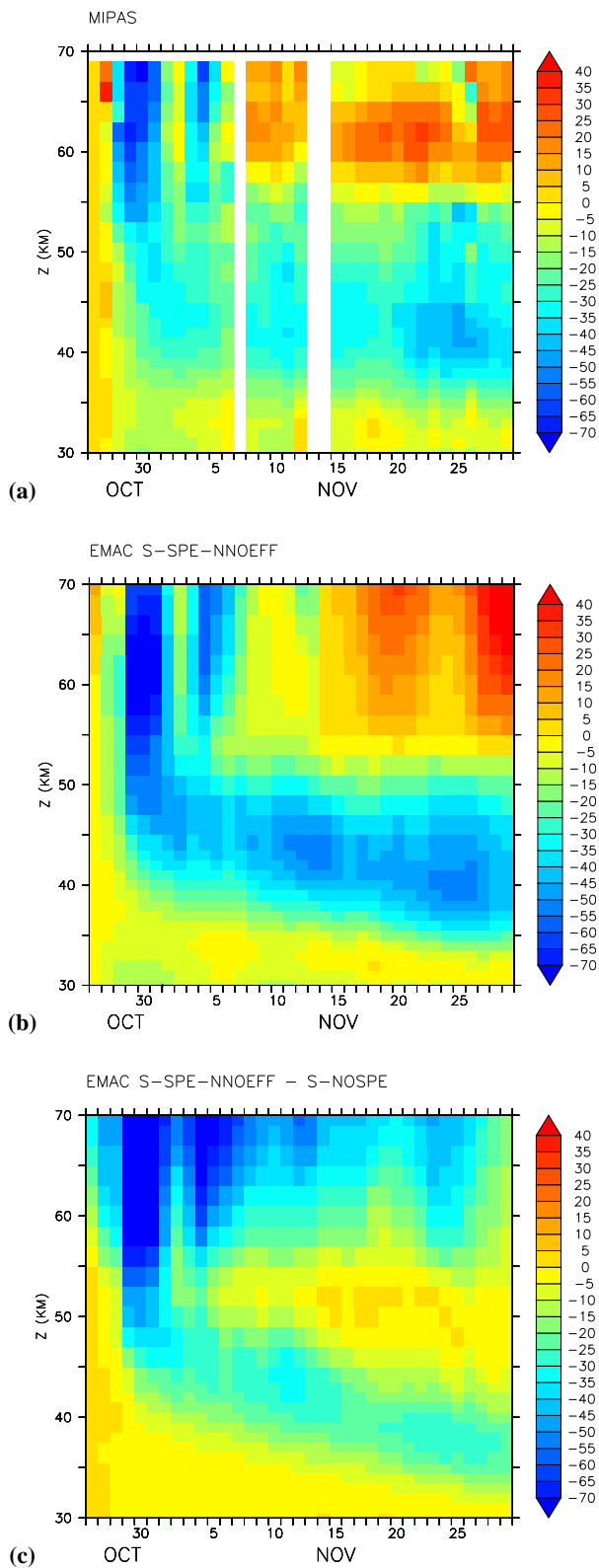


Fig. 12. Ozone mixing ratio percentage change; (a) MIPAS, relative to 26 October, (b) EMAC S-SPE-NNOEFF, relative to 26 October, (c) difference between EMAC S-SPE-NNOEFF and S-NOSPE.

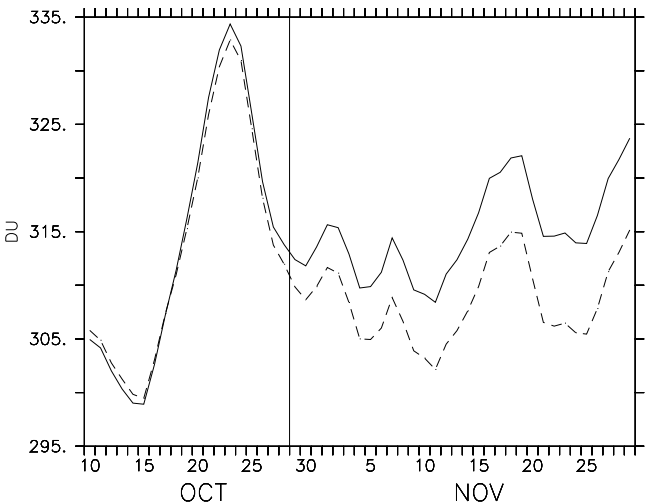


Fig. 13. Change of total ozone between the simulations S-SPE-NNOEFF (dashed line) and S-NOSPE (solid line) in the region 70–90° N.

variations on a timescale of a few days are synchronous because both simulations are nudged to the observed meteorology. The increase in TOC loss during the course of November is likely due to the downward transport of NO_x and associated ozone depletion (see above), becoming more and more important for TOC. This TOC loss is greater than simulated by Vogel et al. (2008), who reported an upper limit of 5.5 DU at the end of November. However, because a different geographical area was used and because of the large influence of natural variations, these results are unlikely to contradict each other.

The OH radical is one of the direct products of SPEs as seen above. The production parameterization leads to short-term enhancements of OH in the upper stratosphere and mesosphere of about 5 ppbv as shown in Fig. 14. MIPAS does not measure OH, but several reactions (e.g. outlined in Jackman et al., 2008) involving HO_x source gases also lead to the buildup of chlorine containing trace gases. For example, von Clarmann et al. (2005) and López-Puertas et al. (2005b) have presented evidence for buildup of HOCl and ClONO_2 in the stratosphere using MIPAS observations. Figure 15 depicts measured (left) and simulated (right) HOCl enhancements. As reported by von Clarmann et al. (2005), the enhancement commences at the start of the SPE and decreases exponentially afterwards, such that there is no evidence for HOCl enhancements after 5 November. The model reproduces this behaviour well, but overestimates the enhancement by about 0.05 ppbv. For ClONO_2 , the start of the enhancement is delayed by 1–2 days with respect to the SPE and lasts for several weeks as shown in Fig. 16a. López-Puertas et al. (2005b) attributed this behaviour to the relatively slow reaction of the enhanced NO_2 to ClONO_2 , providing a reservoir

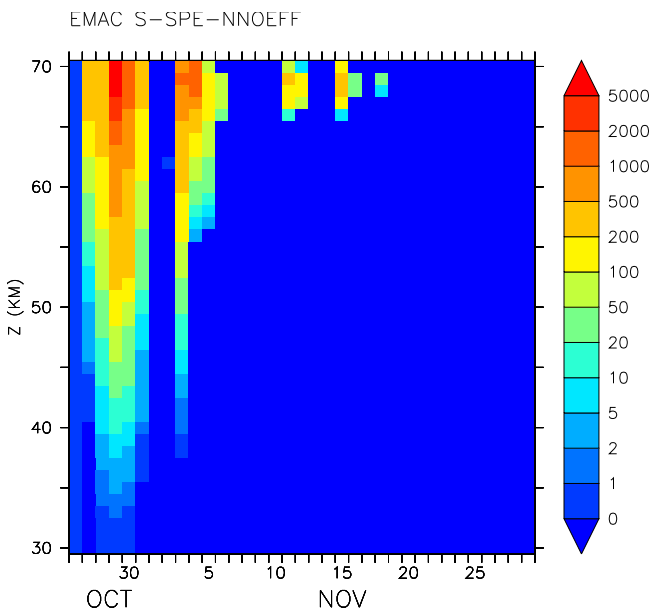


Fig. 14. Simulated change (pptv) of OH for 70–90° N with respect to 26 October 2003.

for the chlorine containing gases. The model (Fig. 16b) qualitatively reproduces the ClONO₂ maximum between 32 and 42 km in the weeks after the SPE, but underestimates the buildup by about 50%.

Figure 17 depicts changes of N₂O₅ mixing ratios in the same manner. Qualitatively the model reproduces the slow buildup of N₂O₅ centred around an altitude of 40 km. Clearly, the measured enhancements of up to 2 ppbv are overestimated by up to a factor of 5. This discrepancy was present in all simulations (S-SPE, S-SPE-FUNKE and S-SPE-NNOEFF). While the reactions that convert NO_x to N₂O₅ are sensitive to temperature, the agreement between MIPAS and EMAC temperature at 40 km (see Fig. 3) implies that the disagreement is not related to temperature effects. Instead, the lack of cluster ion chemistry in EMAC is probably causing this disagreement. Stiller et al. (2005) found that after the conversion of NO_x to N₂O₅ the latter reacts with cluster ions to form HNO₃. Since this reaction is not considered in EMAC, the N₂O₅ enhancements are much stronger than observed.

Indeed, EMAC simulations predict only an HNO₃ mixing ratio increase of 0.6 ppbv (see Fig. 7 in the supplement) in late November, compared to enhancements of 1–5 ppbv reported also by López-Puertas et al. (2005b) using MIPAS observations. Note that EMAC also does not reproduce the short-term increase of HNO₃ during and in the days after the SPE, similar to other models (see e.g. Jackman et al., 2008). As discussed by Verronen et al. (2008), the observed short-term increase is probably related to ion-ion recombination reactions missing in EMAC.

If MIPAS and model NO_y (here: NO₂ + 2×N₂O₅ + HNO₃ + ClONO₂) are compared (see supplement Fig. 11), a good agreement is found. The model overestimations of N₂O₅ and ClONO₂ especially toward the end of November compensate the more rapid loss of NO₂ in the model. This indicates problems concerning the partitioning within the NO_y family and has already been reported by Brühl et al. (2007).

4.4 Dynamical effects

In sunlit areas, the SPE induced perturbation of ozone can potentially lead to temperature changes because of its radiative importance. In order to examine SPE effects on temperature and winds, two further simulations were carried out where the nudging of tropospheric meteorology was turned off. The two simulations are otherwise equivalent to the simulations S-NOSPE and S-SPE-NNOEFF, respectively. The simulations were started on 1 October 2003 from the same initial conditions as all other simulations presented here. For the Halloween events, effects on dynamics are only expected to play a role in the sunlit southern polar area. There, the model predicts an ozone loss of 0.5–1 ppmv, equivalent to 30%–70%, in the lower mesosphere (not shown), but prevalent only during the SPE period. The associated temperature changes in the area 70° S–90° S relative to the simulation without the SPE are depicted in Fig. 18a. An average cooling of up to 2.6 K in the lower mesosphere is evident during the first week after the SPE. The cooling is strongest at 60 km. Jackman et al. (2007) found a cooling up to 2.6 K at 65 km altitude and 90° S from their model simulations. If the EMAC analysis is restricted to 90° S as in Jackman et al. (2007), the maximum cooling amounts to 3.0 K at 60 km. Therefore, the magnitude of the cooling is comparable, although there is a five kilometer altitude offset between the areas of the largest response. Using a mechanistic model, Becker and von Savigny (2010) also found a similar response (−3 K). For comparison, a model study by Krivolutsky et al. (2006) of the stronger July 2000 SPE found a cooling of up to 8 K centered around 60 km. Around the stratopause the cooling lasts longest and is still discernible at the end of November. The observed cooling is likely due to the changes in UV absorption. As a consequence, the model predicts a change of 3 m/s in the zonal mean wind (Fig. 18b), maximizing at 70 km. Changes in the meridional wind were not significant (not shown). These dynamical effects will be subject to further analyses in the future.

5 Summary and conclusions

A new submodel for the parameterization of solar proton events has been implemented in the atmospheric chemistry general circulation model EMAC and tested for the 2003 Halloween storm period. The chemistry submodels of EMAC are also contained in the box model CAABA,

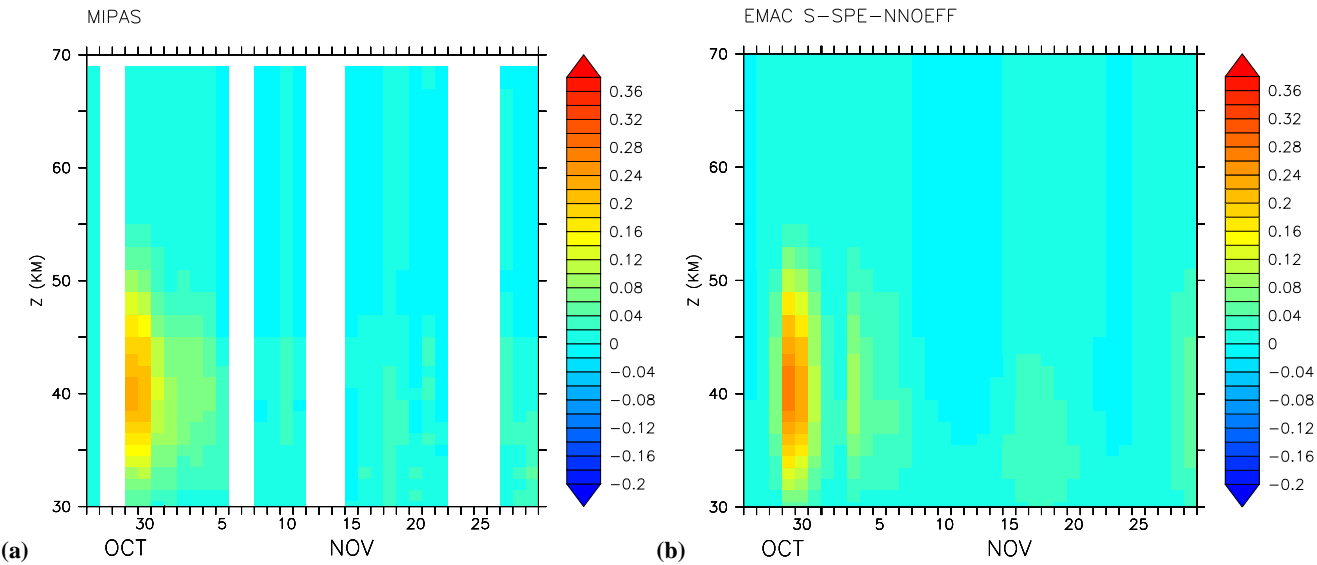


Fig. 15. HOCl changes (ppbv) for 70–90° N relative to 26 October 2003; **(a)** MIPAS, **(b)** EMAC simulation S-SPE-NNOEFF.

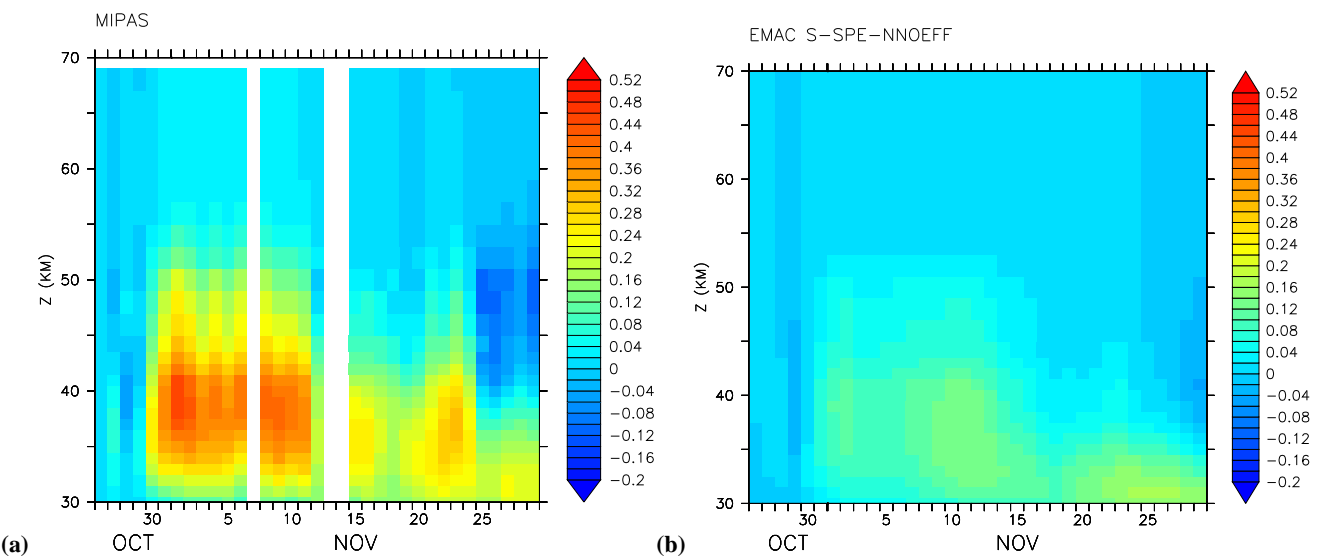


Fig. 16. ClONO₂ changes (ppbv) for 70–90° N relative to 26 October 2003; **(a)** MIPAS, **(b)** EMAC simulation S-SPE-NNOEFF.

allowing to assess the influence of temperature on SPE-induced changes of chemical species through temperature dependent reaction rates. In addition to this feature, EMAC is also well suited for this type of study because of the possibility to relax the model meteorology to reanalysis datasets and because of its interactive chemistry.

The internal calculation of ionization rates, based on particle flux measurements, was evaluated against published ionization rates. One of the parameters of the employed Bethe energy-range relationship was modified by 30% in order to reach better agreement. MIPAS observations were used to

evaluate NO₂, N₂O, N₂O₅, HOCl, ClONO₂, and O₃ which have all been shown to be affected by SPEs. For N₂O the model overestimates SPE related production grossly. Therefore, the production of N and NO per ion pair was modified using results from box model calculations: CAABA simulations were carried out for all individual MIPAS measurements of NO₂ and N₂O during and shortly after the SPEs, employing different N/NO production efficiencies. Using the optimized height dependent production efficiencies in the 3-D model yielded good agreement between EMAC and MIPAS for NO₂, N₂O and O₃. However, due to problems

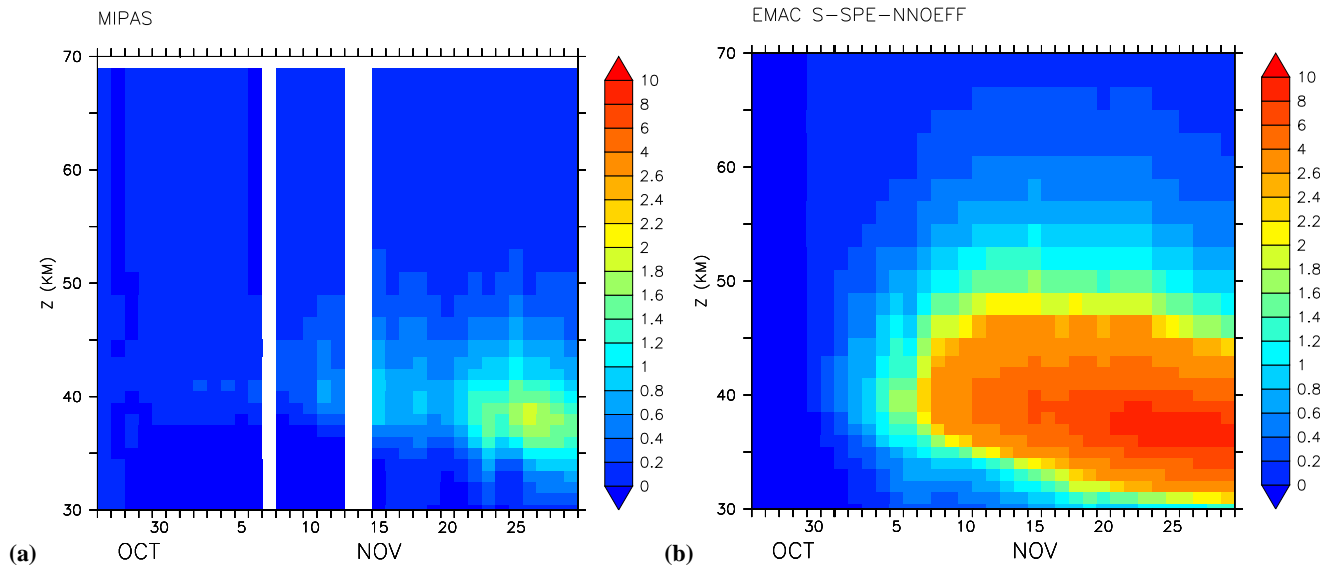


Fig. 17. N₂O₅ changes (ppbv) for 70–90° N relative to 26 October 2003; (a) MIPAS, (b) EMAC simulation S-SPE-NNOEFF.

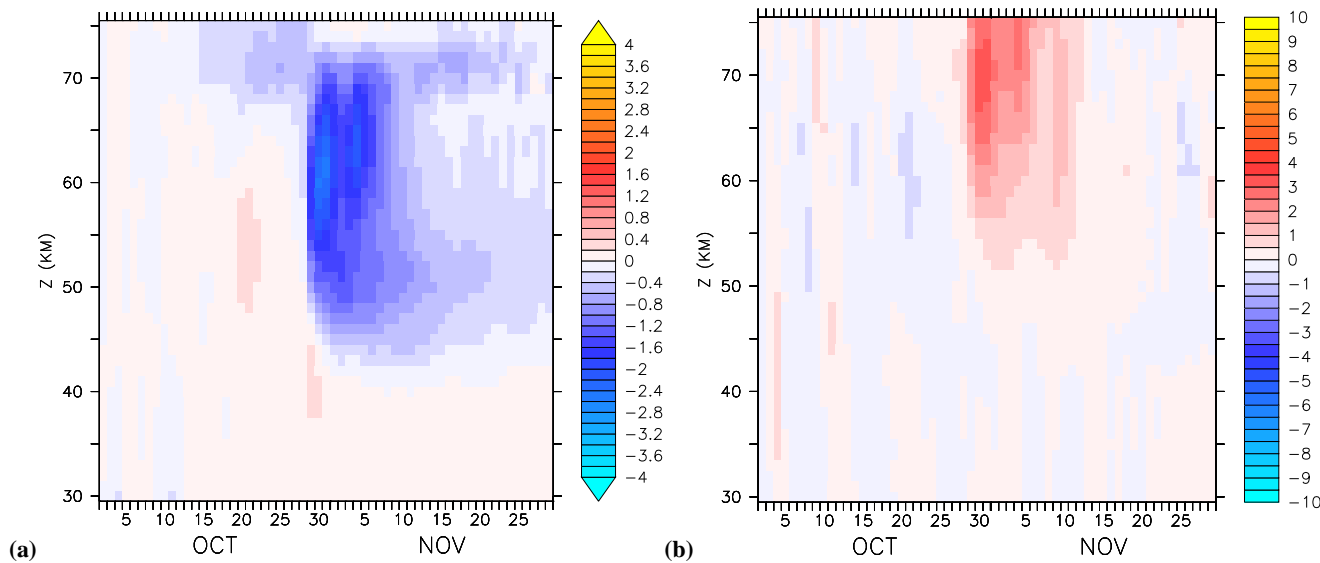


Fig. 18. (a) Change of temperature (K) between two free running simulations with and without SPE submodel turned on. (b) Change of zonal mean wind (m/s).

concerning the NO_y partitioning in the model as well as missing cluster ion chemistry, disagreements persist for HNO₃, ClONO₂ and N₂O₅. Despite these discrepancies the SPE induced ozone depletion in the polar Northern Hemisphere was shown to be in very good agreement with MIPAS throughout the middle atmosphere. Good agreement of HOCl as an indicator for perturbed HO_x was found. Both the modification of the energy-range relationship as well as the modification of the N/NO production efficiency will need to be tested further in EMAC as well as other general circulation mod-

els that include chemistry in order to examine their validity. In case further evidence mounts that such modifications are necessary to reconcile models and observations, further laboratory experiments are also conceivable.

Further results of this study include:

1. Significant changes in chemistry and dynamics already arise due to seasonal variation only, as can be shown by comparing SPE-simulations to an additional simulation without SPE. For example, stratospheric ozone loss

- reaches 50% at the end of November when compared to ozone before the SPEs, but only 35% when compared to a simulation where the SPE submodel was turned off.
2. The loss of total column ozone amounts to approximately 5 DU immediately after the SPEs. At the end of November, when NO_x and associated ozone loss reach lower altitudes, total column loss approaches 10 DU, an amount similar to natural variability and slightly larger than reported by previous studies.
 3. In the Southern Hemisphere, EMAC shows an SPE-induced cooling of about 2.5 K in agreement with another study which included more complex feedbacks that can play a role in the mesosphere-lower-thermosphere.

Supplementary material related to this article is available online at:

<http://www.atmos-chem-phys.net/10/7285/2010/acp-10-7285-2010-supplement.zip>.

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