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Simulation of stratospheric water vapor trends: impact on stratospheric ozone chemistry

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Abstract

A transient model simulation from 1960 to 2000 with the coupled climate-chemistry model (CCM) ECHAM4.L39(DLR)/CHEM shows a stratospheric water vapor trend during the last two decades of +0.7 ppmv and additionally a short-term increase during

- volcanic eruptions. At the same time this model simulation shows a long-term decrease in total ozone and a short-term tropical ozone decline after a volcanic eruption. In order to understand the resulting effects of the water vapor changes on stratospheric ozone chemistry, different perturbation simulations have been performed with the CCM ECHAM4.L39(DLR)/CHEM with the water vapor perturbations fed only to the chem-
- ¹⁰ istry part. Two different long-term perturbations of stratospheric water vapor, +1 ppmv and +5 ppmv, and a short-term perturbation of +2 ppmv with an e-folding time of two months have been simulated. Since water vapor acts as an in-situ source of odd hydrogen in the stratosphere, the water vapor perturbations affect the gas-phase chemistry of hydrogen oxides. An additional water vapor amount of +1 ppmv results in a 5–10%
- OH increase. Coupling processes between HO_x and NO_x/ClO_x also affect the ozone destruction by other catalytic reaction cycles. The HO_x cycle becomes 6.4% more effective, whereas the NO_x cycle is 1.6% less effective. A long-term water vapor increase does not only affect the gas-phase chemistry, but also the heterogeneous ozone chemistry in polar regions. The additional water vapor intensifies the strong denitrification
- of the Antarctic winter stratosphere caused by an enhanced formation of polar stratospheric clouds. Thus it further facilitates the catalytic ozone removal by the CIO_x cycle. The reduction of total column ozone during Antarctic spring peaks at -3%. In contrast, heterogeneous chemistry during Arctic winter is not affected by the water vapor increase. The short-term perturbation studies show similar patterns, but because of the
- short perturbation time, the chemical effect on ozone is almost negligible. Finally, this study shows that 10% of the simulated long-term ozone decline in the transient model simulation can be explained by the water vapor increase, but the simulated tropical ozone decrease after volcanic eruptions is caused dynamically rather than chemically.

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1. Introduction

Water vapor in the upper troposphere (UT) and lower stratosphere (LS) plays a key role in atmospheric chemistry. The oxidation of H_2O and CH_4 by excited oxygen $O(^1D)$ is the primary source of hydrogen oxides ($HO_x=OH+HO_2$) (Reaction R1–R2), which are involved in an important catalytic reaction cycle that controls the production and destruction of ozone in the lower stratosphere. The importance of the catalytic HO_x -cycles for the photochemistry of stratospheric O_3 has already been identified in 1950 by Bates and Nicolet. Additionally, OH is important for changing the partitioning of the nitrogen and the halogen family which are crucial for the O_3 removal in the stratosphere.

$$O(^{1}D) + H_{2}O \rightarrow 2OH$$
(R1)

 $O(^{1}D) + CH_{4} \rightarrow OH + CH_{3}$

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Several studies discussed an increase in stratospheric water vapor (Evans et al., 1998; Michelsen et al., 2000; Nedoluha et al., 1998; Rosenlof, 2002). For example, Rosenlof et al. (2001) combined ten different data sets between 1954 and 2000, and examined a trend of +1%/yr. Several reasons like enhanced methane oxidation, increased aircraft emission in the LS, a warming of the tropical tropopause, volcanic eruptions and large-scale changes in stratospheric circulation and tropospherestratosphere-exchange have been discussed, but the observed stratospheric water vapor increase is not yet understood (Evans et al., 1998, and references therein). In contrast, tropical tropopause temperatures have been found to decrease (Zhou et al., 2001).

Oltmans et al. (2000) analyzed 20 years of water vapor measurements with the NOAA Climate Monitoring and Diagnostics Laboratory (CMDL) frostpoint hygrometer over Boulder, CO, and reported a +1%/yr (+0.05 ppmv/yr) trend in the stratosphere. Figure 1 shows a comparison of this time series with a corresponding water vapor time series from a transient model simulation with ECHAM4.L39(DLR)/CHEM¹. The model

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(R2)





¹Dameris, M., Grewe, V., Ponater, M., et al.: Long-term changes and variability in a transient

results overestimate the observed water vapor mixing ratios more than 50%. Hein et al. (2001) stated that the water vapor minimum above the tropical tropopause is qualitatively reproduced, but stratospheric water vapor mixing ratios are systematically overestimated by ECHAM4.L39(DLR)/CHEM. Both time series show a positive linear trend.

- ⁵ The water vapor trend over Boulder (24–26 km) is +0.044±0.012 ppmv/yr (Oltmans et al., 2000). The simulated water vapor trend is 35% weaker than the observed trend. It amounts to +0.029±0.007 ppmv/yr. The uncertainties are the 95% confidence intervals using the t-statistic. At the same time the model results show a decrease in global mean total column ozone of approximately -1.7%/decade (-5.5 DU/decade).
- Several studies have shown that increasing stratospheric water vapor affects the ozone budget significantly (e.g. Evans et al., 1998; Dvortsov and Solomon, 2001). Figure 2a shows the trend in ozone in the transient model simulation, which is similar to observed values (WMO, 2002). One objective of this study is to assess the contribution of the simulated water vapor increase to the analyzed ozone decrease.
- ¹⁵ The model results show two pronounced water vapor peaks, associated with the eruption of the volcanos El Chichon (17.33° N, 93.20° W) in 1982 and Mount Pinatubo (15.13° N, 120.35° E) in 1991 (Fig. 1). The observations over Boulder also show enhanced water vapor mixing ratios in 1992 after the Pinatubo eruptions as indicated by the dashed line in Fig. 1. The volcanic eruptions lead to a short-term increase of strato-
- ²⁰ spheric water vapor. In the model simulation the water vapor mixing ratios return to prevolcanic levels 2 years after the maximum value. Since both volcanos are located in the tropics within the upwelling branch of the Brewer-Dobson circulation, it is expected that the gas-phase chemistry of the tropical lower stratosphere is primarily influenced by a short-term increased water vapor loading. Several effects of the volcanic eruption of
- ²⁵ Mount Pinatubo on the stratospheric water vapor have been discussed. Beside the direct injection of large amounts of water vapor into the stratosphere (Glaze et al., 1997), volcanic eruptions cause an enhanced sulphate aerosol loading which results in an important heating effect on the tropical tropopause (Stenchikov et al., 1998). A warmer

simulation with a chemistry-climate model employing realistic forcings, to be submitted in ACP.

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tropopause leads to an enhanced transport of water vapor from the troposphere to the stratosphere (Considine et al., 2001). Figure 2b shows the simulated tropical ozone decline after the last three major volcanic eruptions. Beside the ozone trend analysis a second major objective is to investigate whether these short-term ozone changes arise from a short-term water vapor increase.

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The effect of increasing stratospheric water vapor on ozone chemistry has also been discussed within the scope of supersonic aircraft emission studies. Gauss et al. (2003) analyzed the impact of water vapor emissions from aircraft on the atmosphere with a chemical transport model for the year 2015. They found that supersonic aircraft cause a pronounced water vapor perturbation in the lower stratosphere with values up to +400 ppbv (their Figure 8). Gauss et al. (2003) considered the stratospheric water vapor emissions as a passive tracer without chemical loss and applied a tropospheric lifetime of 8.75 days. Morris et al. (2003) applied a trajectory model to assess stratospheric water vapor changes caused by sub- and supersonic aircraft. They sim-

- ¹⁵ ulated maximum water vapor perturbations of 1.5 ppmv in the tropical stratosphere near 20 km. Since the only loss mechanism for water vapor considered in the simulations is transport to the troposphere, Morris et al. (2003) stated that the results should be viewed as upper limit. The impact of supersonic H₂O emissions on stratospheric ozone has been studied with various 2D- and 3D-CTMs in IPCC (1999). In contrast to the present study the IPCC model studies do not include detailed encluses of chamical
- the present study the IPCC model studies do not include detailed analyses of chemical processes and their changes caused by the H₂O perturbation.

In this paper we investigate the impact of increased stratospheric water vapor on stratospheric ozone chemistry by model simulations with the coupled climate-chemistry model ECHAM4.L39(DLR)/CHEM. We use a special method to prevent a feedback of

the simulated water vapor increase to the model dynamics in order to separate the chemical effect. A short model description is given in the next section. Section 3 describes the applied tracer approach. The results of our study are presented in Sec. 4.1 and 4.2. We differentiate between a short-term and a long-term water vapor increase. A discussion and summary is given in the last section.

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2. Model description

The coupled climate-chemistry model ECHAM4.L39(DLR)/CHEM (Hein et al., 2001, hereafter referred to as E39/C) consists of the dynamic part ECHAM4.L39(DLR) (E39) and the chemistry module CHEM. E39 is a spectral general circulation model, based

on the climate model ECHAM4 (Roeckner et al., 1996), and has a vertical resolution of 39 levels up to the top layer centered at 10 hPa (Land et al., 1999). A horizontal resolution of T30 (≈6° isotropic resolution) is used in this study. The tracer transport, parameterizations of physical processes and the chemistry are calculated on the corresponding Gaussian transform grid with a grid size of 3.75° ×3.75°. Water vapor, cloud water and chemical species are advected by a so-called semi-Lagrangian scheme.

The chemistry module CHEM (Steil et al., 1998) is based on the family concept. It includes stratospheric homogeneous and heterogeneous ozone chemistry and the most relevant chemical processes for describing the tropospheric background NO_x -CH₄-CO-HO_x-O₃ chemistry with 107 photochemical reactions, 37 chemical species

- and 4 heterogeneous reactions on polar stratospheric clouds (PSCs) and on sulphate aerosols. CHEM does not yet consider bromine chemistry. Mixing ratios of methane (CH₄), nitrous oxide (N₂O) and carbon monoxide (CO) are prescribed at the surface following IPCC (2001) for the year 2000. Zonally averaged monthly mean concentrations of chlorofluorocarbons (CFCs) and upper boundary conditions for total chlorine
- and total nitrogen are taken from the 2-D model of Brühl and Crutzen (1993). Nitrogen oxide emissions at the surface (natural and anthropogenic sources), from lightning and aircraft are considered. The model set-up follows Hein et al. (2001) except for the total amounts of emissions, lightning NO_x, which is parameterized according to Grewe et al. (2001), and a more detailed sulphate aerosol chemistry.
- The model E39/C can be run in two different modes: with- and without-feedback. In the without-feedback mode the concentrations of the radiatively active gases H₂O, O₃, N₂O, and CFCs calculated by CHEM do not feed back to the radiative scheme of E39. Prescribed climatological mixing ratios of the radiatively active gases are used as input

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for the radiative scheme instead. A detailed model climatology is given in Hein et al. (2001).

The hydroxyl radical (OH) plays a crucial role for the present study. Since the model climatology given in Hein et al. (2001) does not consider OH, the OH distribution in E39/C will be discussed below. Figure 3 shows a vertical cross-section of the zonally 5 and monthly averaged OH distribution in E39/C. Because of the extreme variability of OH in time and space due to the high dependency on the solar zenith angle (Hanisco et al., 2001), it is inappropriate to compare in situ OH measurements (e.g. Wennberg et al., 1995) with model results which include night hours. Spivakovsky et al. (2000) calculated a global climatological distribution of tropospheric OH (referred to OH-S 10 hereafter) using observations of precursors for OH like O₃, H₂O, CO, hydrocarbons and NO_t (defined as NO₂ + NO + $2N_2O_5$ + NO₃ + HNO₂ + HNO₄, Spivakovsky et al., 2000) as well as observed temperatures and cloud optical depths as input for a photochemical box model. OH-S shows the highest zonal mean OH concentrations in the tropics at ca. 600 hPa, peaking at about 25×10^5 molecules/cm³. Within this altitude 15 range the tropical OH concentrations in E39/C are in good agreement with OH-S. The

OH distribution in E39/C has a maximum closer to the surface. This might be associated with the missing effects of non-methane-hydrocarbons (NMHCs) on OH in E39/C. OH is depleted over forested tropical continents by the influence of NMHCs (Lawrence

- et al., 2001). In both OH distributions the maximum OH concentrations are shifted to the respective summer hemisphere during January and July, following the position of the sun. During April and October both OH distributions are mostly hemispherically symmetric. The mid-latitudes exhibit a strong seasonal cycle that is associated with sunlight and water vapor variations. Both OH distributions show very low OH concen-
- trations at high latitudes during winter. OH-S shows a more pronounced vertical gradient between 500 and 100 hPa. Hence E39/C overestimates the OH concentrations in this altitude range. Generally, the comparison of both OH distributions shows that for the matter of this study the modeled OH distribution is realistically enough reproduced.

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3. Methodology

3.1. Water vapor perturbations

The intention of this study is to investigate the impact of an additional stratospheric water vapor loading on the stratospheric chemistry without changing atmospheric dynamics. Increasing stratospheric water vapor results in changing dynamics. This dynamical effect as well as the changing chemical composition leads to a modified ozone distribution. In turn, the changed ozone distribution affects the model dynamics again. For a better understanding of coupling processes between chemistry and dynamics the model has been run without feedback effects of the simulated water vapor increase on the model dynamics. For this purpose a special tracer approach was introduced in E39/C: Two water vapor tracers are differentiated - the background water vapor and a water vapor perturbation. The radiative calculations are performed with the background water vapor. The "chemical" water vapor field is defined as follows:

$H_2O_{Chemistry} = H_2O_{Background} + H_2O_{Perturbation}$

The water vapor perturbation is transported like an arbitrary tracer by the semi-Lagrangian advection scheme of E39/C. In order to treat the water vapor perturbation as realistic as possible it has to pass the same physical source and sink processes as the background water vapor. These processes include convection, condensation and vertical exchange by turbulence. This means, if cloud formation and precipitation cause a reduction of the background water vapor content by 10% within one model grid box, then the amount of the water vapor perturbation within this grid box is also reduced by

¹⁰ 10%. The water vapor perturbation is fixed to a constant value within the tropical lower stratosphere at each integration time step. Below 200 hPa the perturbation field is set to zero since this study concentrates on the stratospheric chemistry.

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3.2. Experimental design

For the current study various model simulations with E39/C have been performed. A short overview of the performed simulations is given in Table 1. All model simulations were performed in the without-feedback mode. Thus, all experiments show the same meteorology. The model runs simulate atmospheric conditions of the year 2000. The adopted mixing ratios of greenhouse gases and NO_x emissions of different sources are given in Table 2 and 3. The simulations CNTL, H2O_1 and H2O_5 were integrated over 11 years in a quasi-stationary state (time slice simulations). The last five years of the integration period have been analyzed. The simulation CNTL was performed without a water vapor perturbation as reference simulation. As mentioned above, this paper deals with two different objectives. The first one is to investigate the effects of a short-term increase of stratospheric water vapor which e.g. occurs during a volcanic eruption. Therefore, an ensemble of five 1-year-simulations has been analyzed, representing the last five years of CNTL (hereafter referred to as VOLC). Each annual cycle starts in July,

- since the Mount Pinatubo eruption occurred June 15, 1991. During July and August additional water vapor is "emitted" in the tropical lower stratosphere by the above mentioned method. The water vapor perturbation in the tropical lower stratosphere is set to +2 ppmv. For the rest of the year the emission is stopped. Five different years have been chosen to cover a range of dynamical situations. Figure 4 shows the annual cycle
- of the water vapor perturbation for the five simulations at 89 hPa at 30° N/S and 60° N/S. Within the tropics the perturbation shows a rapid decrease after the peak in August. The e-folding time of the perturbation is 2 months. The variability within the ensemble of five 1-year-simulations is very small. At 30° the interhemispheric differences are just as little as the differences between the different annual cycles of VOLC. At 60° the
- interhemispheric differences are much more pronounced. In the northern hemisphere the perturbation peaks in September whereas the southern hemispheric peak occurs 3 month delayed in December. This pattern indicates a more intense northward transport in the model within the lowermost stratosphere during summer. This transport pattern

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has also been identified by Grewe et al. (2004) who studied the impact of horizontal transport caused by streamers on the chemical composition of the tropopause region.

The second part of this study deals with the impact of a long-term water vapor trend. For that purpose two long-term simulations including a stratospheric water vapor per-

- ⁵ turbation have been performed, H2O_1 and H2O_5. In H2O_5 the water vapor perturbation in the tropical lower stratosphere is set to ca. +5 ppmv. This value corresponds roughly to a doubling of water vapor within this atmospheric region. This model experiment was designed as a maximum impact scenario. Taking into account a stratospheric water vapor trend of +0.05 ppmv/yr the doubling of water vapor would be
- ¹⁰ reached in ca. 100 years. The simulation H2O_1 was performed like H2O_5, but the perturbation was set to +1 ppmv in the tropical lower stratosphere. Figure 5 shows the climatological mean distribution of the water vapor perturbation in H2O_5 for January and July. The dark shaded area indicates the "source region" where the perturbation is fixed. Approximately 70% of the tropical perturbation can be found around
- ¹⁵ 70° N (~3.5 ppmv), whereas the perturbation is much lower around 70°S (southern hemispheric values 15%–20% lower than northern hemispheric values). Again, these results indicate a more intense northward transport. The strong dehydration of the Antarctic winter stratosphere is clearly visible. The dehydration of the Arctic winter stratosphere is less intensive. The minimum values of the additional water vapor in
 the Antarctic winter stratosphere fall below 1 ppmv in H2O_5 (H2O_1: ~0.1 ppmv),
- whereas the minimum values in the Arctic winter stratosphere are around 3 ppmv in H2O_5 (H2O_1: ~0.45 ppmv). The results for simulation H2O_1 show the same pattern, just the absolute values are 5 times smaller.

4. Results

As mentioned above, the oxidation of H_2O and CH_4 by $O(^1D)$ (Reaction R1 and R2) is the primary source of OH in the stratosphere. Excited oxygen atoms, $O(^1D)$, are produced by photolysis of O_3 at wavelengths shorter than about 320 nm. Associated

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with strong actinic fluxes the photolysis rate of O_3 is highest near the equator. Since the tropical lowermost stratosphere is the region of the highest H_2O perturbation, the most pronounced changes of the OH distribution are expected for the tropical lowermost stratosphere within all perturbation scenarios. In turn, the production of H_2O by the methane oxidation is enhanced:

$$CH_4 + OH \xrightarrow{O_2} CH_3O_2 + H_2O$$
(R3)

The stratospheric OH increase causes a variety of changes concerning the different ozone production and destruction cycles. The results of our study will show that these changes yield to a net ozone destruction. At first a short overview concerning the role of OH within the stratospheric ozone chemistry is given. Different catalytic cycles take part in stratospheric ozone destruction. These cycles can be written in a general form:

 $\begin{array}{l} X+O_3 \rightarrow XO+O_2 \\ XO+O \rightarrow X+O_2 \end{array}$

 $\mathrm{O}_3 + \mathrm{O} \rightarrow \mathrm{O}_2 + \mathrm{O}_2$

where X can be one of the free radicals H, OH, NO, CI or Br. Beside this general form additional catalytic cycles exist, but only an additional HO_x cycle is mentioned here:

 $\begin{array}{l} \mathsf{OH} + \mathsf{O}_3 \rightarrow \mathsf{HO}_2 + \mathsf{O}_2 \\ \mathsf{HO}_2 + \mathsf{O}_3 \rightarrow \mathsf{OH} + \mathsf{O}_2 + \mathsf{O}_2 \end{array}$

 $O_3 + O_3 \rightarrow O_2 + O_2 + O_2$

Of special interest for our study is the coupling of the HO_x and NO_x/ClO_x cycles, since coupling reactions change the catalytic effectiveness of the separate catalytic cycles. The coupling of the HO_x and NO_x cycles occurs by the following reactions:

 $HO_2 + NO \rightarrow NO_2 + OH$

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(R4)

(R5)

(R6)

$$\begin{array}{c} \mathsf{OH} + \mathsf{NO}_2 + \mathsf{M} \to \mathsf{HNO}_3 + \mathsf{M} & (\mathsf{R7}) \\ \\ & \mathsf{HNO}_3 \xrightarrow{h\nu} \mathsf{OH} + \mathsf{NO}_2 & (\mathsf{R8}) \\ \\ & \mathsf{HNO}_3 + \mathsf{OH} \to \mathsf{H}_2\mathsf{O} + \mathsf{NO}_3 & (\mathsf{R9}) \end{array}$$

An increase of OH should result in an enhanced transformation of NO₂ into the reservoir species HNO₃. Since Reaction R9 is less effective in releasing NO_x which is tied in the reservoir species HNO₃, the coupling of the HO_x and NO_x cycle is expected to reduce the catalytic effectiveness of the NO_x cycle. Reaction R6 controls the partitioning between OH and HO₂ which is important for the effectiveness of the HO_x cycle. As OH increases, ozone destruction by the CIO_x cycle becomes more effective:

 $OH + HCI \rightarrow H_2O + CI$ $HO_2 + CIO \rightarrow HOCI + O_2$ (R10)
(R11)

The reservoir species HCl is converted back into active Cl by Reaction R10. Hypochlorous acid (HOCl) is photolyzed to OH and active Cl. Increasing stratospheric H_2O (and hence OH) decreases the catalytic effect of the NO_x cycle and enhances the effectiveness of the ClO_x cycle and the HO_x cycle on the O_3 destruction. The following sections present a quantification of the impact of increasing stratospheric water vapor on stratospheric chemistry.

4.1. Short-term increase of stratospheric water vapor

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This section deals with the impact of a short-term increase of stratospheric water vapor as associated with strong volcanic eruptions like the Mount Pinatubo eruption in 1991 (simulation VOLC). It should be pronounced that the volcanic impact on stratospheric aerosol is not considered in the simulations. Neither an enhanced ozone depletion caused by the increasing aerosol surface area nor the aerosol effect on the tropical tropopause temperature is taken into account. Figure 6 shows the percentage changes of the OH concentration at 50 hPa (upper panel) and 80 hPa (lower panel) in the VOLC

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simulation depending on latitude and season. The OH concentration increases in both hemispheres. The most pronounced changes occur within the tropics associated with the highest H₂O perturbation. The tropical OH increase peaks in July with up to +15% and more at 80 hPa and in August with up to 7% at 50 hPa. The 50 hPa level is above
the "source" region of the H₂O perturbation (≈96–70 hPa). Thus, the upward transport within the tropics takes about one month between 80 and 50 hPa. Of special interest are the two peaks at ca. ±30–40° at 80 hPa. This pattern is associated with the polwards and downwards transport of the maximum OH signal. In contrast, at 50 hPa the maximum is located symmetrically around the equator. The percentage changes of the monthly and zonal mean OH concentration depending on the altitude for August are

- displayed in Fig. 7. The downward transport at ca. $\pm 30-40^{\circ}$ is clearly visible in both hemispheres, but it is more pronounced in the southern hemisphere. The changes within the polar winter regions are not relevant due to the low OH background concentrations. As mentioned above changing stratospheric OH levels affect the different
- ¹⁵ ozone destruction reactions. Figure 8 shows the relative changes of the effectiveness of the HO_x, ClO_x and NO_x cycle as well as of the reaction H₂O + O(¹D) \rightarrow 2OH (removal of odd oxygen) at 50 hPa and different latitudes (upper panel: tropics, lower panel: mid-latitudes) caused by the stratospheric OH increase. The background contribution of the different processes to the ozone destruction in CNTL is given in the first
- ²⁰ row of Table 4. The dominant effect is the increase of ozone destruction through the HO_x cycle. Within the tropics the increase of the HO_x cycle peaks in September with approximately +7% compared to CNTL. Additionally, the CIO_x cycle and the reaction $H_2O + O(^1D)$ become more effective, but both ozone destroying reactions have only little importance in the tropics or mid-latitudes. The changes of the NO_x cycle are neg-
- ligible (≤-1%). Qualitatively, the mid-latitudes show the same results. The changes of the effectiveness are less pronounced than in the tropics since the OH increase is less. Additionally, the peaks are slightly shifted to November/December because of the transport times between tropics and mid-latitudes.

The changing ozone destruction yields a global reduction of total column ozone ex-

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cept in tropical and mid-latitude regions during July as presented in Fig. 9. The shortterm total ozone increase within the tropics during July is associated with an enhanced ozone production by the following reactions (enhanced NO₂ photolysis):

$$HO_2 + NO \rightarrow OH + NO_2$$
 (R12)

$$NO_2 \xrightarrow{\mu\nu} NO + O \tag{R13}$$

$$O_2 + O \rightarrow O_3 \tag{R14}$$

$$O_2 + O \rightarrow O_3$$

This increased ozone production is restricted to the altitude range 100-80 hPa and compensates the enhanced effectiveness of the HO_x cycle. One month later the water vapor perturbation has also reached higher altitudes, where the ozone depletion by the HO_v cycle becomes more important. Then, the enhanced ozone destruction by the ⁵ HO_x cycle dominates the increased ozone production at lower altitudes. Whether a tropical water vapor perturbation leads to ozone loss or ozone production depends on the altitude.

No results are shown for southern high latitudes during Antarctic winter and spring in Fig. 9. Numerical effects lead to an unrealistic increase of CIONO₂ in the Antarctic stratosphere, resulting in a reduced ozone removal by the CIO_x cycle. The CIONO₂ increase occurs right after the start of the experiments. Dynamical reasons for this CIONO₂ increase are not possible, since the experiments CNTL and VOLC simulate identical meteorology. Chemical conversion between the individual members of the CIX-family caused by the OH increase is also very unlikely, since the changes of other chemical species like HCl or ClO_x do not show an adequate behavior. Additionally, the 15 transport times between the tropical "source" region of the water vapor perturbation and the southern polar stratosphere are inappropriate to explain these rapid changes. Interestingly, the most pronounced total column ozone reduction takes place at northern high latitudes during spring/early summer. Since the chemical ozone destruction is only slightly enhanced in VOLC during Arctic spring this result seems to be associ-20



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port of tropical air masses to higher latitudes at pressure levels between 100 hPa and 30 hPa with the climate-chemistry model E39/C. Low ozone air masses are transported from the tropics to higher latitudes by wave breaking events, so-called streamers. They showed that streamers cause a 30% (50%) decrease of ozone at the extra-tropical tropopause of the summer (winter) hemisphere.

The changes displayed in Fig. 9 are statistically significant although the changes are less than 1% of the total column ozone. For this study only the variability of the chemical signal and not the variability of the dynamical signal is used for the *t*-test since all simulations have an identical meteorology.

- ¹⁰ Unfortunately, no observational data are available for the period right after the Mount Pinatubo eruption which would be suitable for a comparison with the model results. Following the volcanic eruption SAGE II (Stratospheric Aerosol and Gas Experiment) ozone values were being systematically overestimated caused by the Pinatubo aerosol. SAGE II ozone values below 22 km altitude are affected by aerosol for approximately 2
 ¹⁵ years after the Pinatubo eruption (SPARC, 1998). HALOE (Halogen Occultation Exper-
- iment) and MLS (Microwave Limb Sounder) satellite data are available since October 1991, i.d. no pre-Pinatubo observations are available for comparison.
 - 4.2. Long-term increase of stratospheric water vapor

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This section presents the impact of a long-term increase of stratospheric water vapor on the stratospheric chemistry as simulated in the model experiments H2O_1 and H2O_5. Since the results of both experiments show the same patterns the following figures refer to the simulation H2O_5. Figure 10 shows the relative changes of OH in H2O_5 compared to the reference simulation CNTL. The zonal mean changes, averaged over 5 years, for July and January are displayed. As expected the stratospheric OH content increases. The highest OH increase occurs in the tropical lower stratosphere. An additional stratospheric water vapor content of +5 ppmv (+1 ppmv) results in a 50% (10%) increase in OH (≈+20×10⁵-+25×10⁵ molecules/cm³). Thus, the OH response to increasing stratospheric

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water vapor is almost linear. Shaded areas indicate the statistic significance (*t*-test) of the changes. Since the existence of OH is coupled to the sunlit atmosphere, OH background concentrations are very low during the permanent polar night. Thus the high percentage changes in the winter polar hemispheres are negligible. In turn, the water vapor production by the oxidation of methane (Reaction R3) is up to 75% (10%) higher

in H2O_5 (H2O_1) than in CNTL (not shown). The regions of maximum increase in water vapor production are directly connected to the regions of highest OH increase.

Table 4 summarizes the changes of different ozone destroying cycles and reactions at 50 hPa and different latitudes caused by the H_2O perturbation. The first row displays

- the contribution of the different ozone destroying reactions to the net ozone destruction in the control simulation CNTL. The following two rows display the percentage changes in the perturbation experiments H2O_5 and H2O_1 compared to CNTL. The dominating effect is the enhanced effectiveness of the catalytic ozone removal by the HO_x cycle (Reactions R4–R5). Within the tropics the effectiveness of the HO_x cycle is 29.0%
- (6.4%) higher in H2O_5 (H2O_1) than in CNTL. The effectiveness of the HO_x cycle increases almost linear with rising OH concentrations. The changes of the NO_x cycle and the removal of excited oxygen by the reaction with water vapor also show a linear behavior with increasing OH. The last mentioned process plays only a minor role in the ozone destruction. In contrast, the ClO_x cycle shows a strong non-linear behavior.
- ²⁰ The ClO_x cycle is 7.8% more effective in H2O₁, but above 100% more effective in H2O₅. However, the contribution of the ClO_x cycle in the tropics is very small. Within mid-latitudes the different ozone destruction cycles show qualitatively the same pattern except the ClO_x cycle which increases only slightly with increasing OH.

The increase in OH does not only affect ozone destruction, it also results in an enhanced ozone production. For example, as stratospheric ozone declines, ultraviolet radiation penetrates deeper into the stratosphere which leads to an enhanced ozone production by the Chapman mechanism:

$$O_2 + h\nu \xrightarrow{\lambda < 242} nm O + O$$

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(R15)

 $O_2 + O + M \rightarrow O_3 + M.$

This process affects mainly the tropical lower stratosphere, but the ozone production by the Chapman mechanism is only slightly enhanced (less than 5% in H2O_5). Ozone production increases as well caused by the increased NO₂ concentration, $HO_2 + NO \rightarrow OH + NO_2$, which is photolyzed. Additionally, the methyl peroxy radical CH_3O_2 , a product of the methane oxidation (Reaction R3), reacts with NO to NO_2 , which again leads to an increased ozone production, but this process is two orders of magnitude less important than the other mentioned reactions. However, the increased ozone production can not compensate the enhanced ozone destruction. This effect was already mentioned in the previous section.

The above mentioned results deal with the gas-phase chemistry and do not consider possible effects on the heterogeneous reactions on polar stratospheric clouds (PSCs). PSCs occur between 10 and 25 km at temperatures below \approx 190 K. Such low temperatures are more frequent in the Antarctic polar vortex. The release of active chlorine from the reservoir species HCl and ClONO₂ is very slow in the gas-phase. PSCs promote the activation of chlorine. At sunrise, activated chlorine compounds are easily photolyzed and catalytic ozone destruction starts. The chemistry module CHEM considers the following heterogeneous reactions on PSCs and sulfate aerosols:

$HCI + CIONO_2 \rightarrow CI_2 + HNO_3$	(R17)
$H_2O + CIONO_2 \rightarrow HOCI + HNO_3$	(R18)
$HOCI + HCI \rightarrow CI_2 + H_2O$	(R19)
$N_2O_5 + H_2O \rightarrow 2HNO_3$	(R20)

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Table 4 displays the percentage changes of the different ozone destroying reactions at 50 hPa in polar regions during spring. There is a remarkable interhemispheric difference of the importance of the NO_x and ClO_x cycle. At the North Pole, the NO_x cycle dominates the ozone destruction, whereas the ClO_x cycle contributes nearly 90% to the ozone destruction at the South Pole which is associated with the strong denitrification of the Antarctic strategy during winter. The denitrification removes NO_x from

 $_{\rm 15}$ tion of the Antarctic stratosphere during winter. The denitrification removes $\rm NO_x$ from

(R16)

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the system which prevents the re-formation of $CIONO_2$ and further supports catalytic ozone destruction by the CIO_x cycle. The impact of increasing water vapor is more pronounced at the South Pole where ozone destruction in general (sum of all ozone destroying reactions/cycles) increases (Table 4). The most important changes during Antarctic spring concern the NO_x and ClO_x cycle. The effectiveness of the ClO_x cycle is \approx 50% higher in H2O_1 and 2.5 times higher in H2O_5. The additional water vapor leads to an increase of PSCs Type II (ice) within the Antarctic polar vortex which is obvious in the strong dehydration of the Antarctic winter stratosphere (Fig. 5). In turn, the activation of chlorine from the reservoir species HCl and ClONO₂ is enhanced. In contrast, the NO_x cycle becomes \approx 50% less effective in H2O_1 and is almost vanished in H2O_5. The additional stratospheric water vapor seems to have no impact on the betrageneous chamistry in parthern polar entropy. The azona destruction rate in gen

heterogeneous chemistry in northern polar spring. The ozone destruction rate in general remains nearly unchanged, but the contribution of the different ozone destroying reactions is shifted (Table 4).

- As a result of the above mentioned processes the ozone concentration decreases almost everywhere. Figure 11 displays the ozone changes determined by the differences of the zonal mean climatological ozone volume mixing ratio between the simulations H2O_5 and CNTL for January and July. The strongest ozone reductions occur in the polar regions of both hemispheres. The maximum in the winter hemisphere is slightly
- shifted downwards compared to the summer hemisphere. Tropospheric changes do not exceed more than −1%. The ozone decline in H2O_5 is ≈5 times higher than in H2O_1. The most pronounced ozone changes are computed in the southern hemisphere with values up to −10% (−3%) in the simulation H2O_5 (H2O_1). The change of total column ozone between the simulations H2O_5 and CNTL is presented in Fig. 12.
- In the tropics and the northern the pattern is very uniform without any seasonal variability. As expected, the most pronounced changes of total column ozone occur during southern polar spring with values up to -15% (H2O_5) and -3% (H2O_1), respectively. The southern polar ozone decline is not statistically significant during September/October and December/January (see also Fig. 11). This feature is associated with

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the seasonal variability of the water vapor perturbation in polar regions. This seasonal cycle is especially pronounced in the southern hemisphere. Throughout the southern winter and springtime the content of the water vapor perturbation decreases rapidly to values less than 1 ppmv (H2O_5) and 0.1 ppmv (H2O_1), respectively. This dehydration of the Antarctic winter lower stratosphere is associated with extremely low temperatures. The duration as well as the beginning of the dehydration of the Antarctic

winter lower stratosphere varies from year to year, resulting in a non-uniform ozone depletion pattern.

So far, this section dealt with the stratospheric changes caused by a water vapor increase. But as shown in Fig. 10 tropospheric OH concentrations increase slightly as well. As stratospheric ozone decreases the absorption of short-wave radiation by ozone is reduced and the amount of short-wave radiation reaching the ground increases. In the troposphere the photolysis of ozone at wavelengths < 319 nm is crucial for the formation of OH (Reactions R1–R2). Figure 13 shows percentage changes of the surface dose at 309 nm in the simulation H2O_5 compared to CNTL. The amount

- the surface dose at 309 nm in the simulation H2O_5 compared to CNTL. The amount of short-wave radiation reaching the ground increases. The pattern shown in Fig. 13 is directly correlated to the ozone changes displayed in Fig. 12: The percentage increase in surface dose is highest at southern high latitudes during southern spring when the total column ozone reduction is highest as well. In this region the surface dose inarcase at 200 nm in 450%. Within the trapica the surface dose is ophonead by 42%.
- ²⁰ crease at 309 nm is +50%. Within the tropics the surface dose is enhanced by +2%. The surface dose changes in simulation H2O_1 show the same pattern (not shown), but the percentage changes are 5 times smaller than in H2O_5.

5. Conclusions

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The impact of different idealized stratospheric water vapor perturbations on the stratospheric ozone chemistry without any feedback on atmospheric dynamics was studied with the coupled climate-chemistry model ECHAM4.L39(DLR)/CHEM. A long-term increase (+1 ppmv and +5 ppmv) as well as a short-term increase ("volcanic eruption",

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+2 ppmv, 2 month e-folding time) of stratospheric water vapor were simulated. The impact of water vapor perturbations on the effectiveness of different ozone destroying reactions as well as coupling effects between different chemical processes were detailed analyzed.

A stratospheric water vapor increase leads to an increased OH concentration which results primarily in an enhanced ozone depletion by the HO_x-cycle. The coupling of the HO_x-cycle with the NO_x- and ClO_x-cycle leads to a less effective NO_x-cycle and a slightly enhanced effectiveness of the ClO_x-cycle. A long-term increase in stratospheric water vapor affects the heterogeneous ozone destruction (ClO_x-cycle) within
 the Antarctic polar vortex caused by an enhanced formation of PSCs Type II (ice). Heterogeneous ozone chemistry in the Arctic polar vortex is not affected. The simulated short-term perturbation does not affect heterogeneous ozone chemistry.

A comparison between the short-term (VOLC) and long-term water vapor perturbations (H2O_1 and H2O_5) reveals some differences. VOLC shows an increase in total column ozone in the tropics during July and August. During this period of the simulation

- VOLC the water vapor perturbation is not yet evenly distributed and the OH increase is concentrated on the lowermost stratosphere. The ozone chemistry is dominated by the ozone production caused by an enhanced NO₂ photolysis (Reactions R12–R14). The HO_x cycle becomes more important in higher altitudes. The simulations H2O₁ and
- H2O_5 do not show a similar effect since the water vapor perturbation has reached a steady distribution. Simulation VOLC as well as simulation H2O_1 show a 7% increase in OH at 50 hPa within the tropics during August. Despite similar water vapor and OH perturbations VOLC and H2O_1 show different ozone changes within this region. The ozone reduction in H2O_1 amounts -1%, but only -0.07% in VOLC. At this alti-
- ²⁵ tude the ozone concentration is dynamically controlled (WMO, 1986). Thus, the effect of a short-term change in chemistry is less important than the impact of a long-term change resulting in different ozone transport.

The almost negligible impact of the short-term water vapor changes on tropical total ozone columns can therefore not explain the ozone decline in the transient model

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simulation of approximately 4% (Fig. 2b). The increased stratospheric aerosol heats the stratosphere and amplifies tropical ascent by roughly 20%, leading to an additional uplift during a volcanic period of roughly 1.2 km, which compares reasonably with the uplift of the Pinatubo aerosol cloud by 1.8 km (DeFoor et al., 1992; Kinne et al., 1992).

- Applying a linearized transport-chemistry column model and using E39/C values for uplift, ozone production and destruction rates of ozone, the simulated vertical ozone profile can be reproduced within 5 to 25% accuracy. Introducing an additional uplift of 1.2 km over 8 months produces a vertical displacement of the ozone profile and a reduction of the ozone column of 5% (red stars in Fig. 2b), agreeing well with the findings of Kippe et al. (1002). This means that the ozone decline simulated during values in
- ¹⁰ of Kinne et al. (1992). This means that the ozone decline simulated during volcanic eruptions is dominated by dynamic effects.

Within IPCC (1999) the effects of a future (year 2015) supersonic aircraft fleet (500 aircraft) on stratospheric O_3 , H_2O and other species was conducted with various 2-D and 3-D CTMs (chemistry transport models). The maximum H_2O perturbations caused

- ¹⁵ by supersonic aircraft emissions occur in the northern hemisphere lower stratosphere (\approx 20 km) and range from 0.4 ppmv to 0.7 ppmv. The percentage changes in total column ozone averaged over the northern hemisphere are in the -0.3...-0.6% range. This scenario includes only H₂O emissions, NO_x emissions have been excluded (El(NO_x)=0). Performing further scenarios with different El(NO_x) all models
- predicted that the most significant supersonic impact on ozone is caused by the water vapor emissions and that the addition of NO_x emissions leads to a less ozone depletion (IPCC, 1999). The water vapor perturbation in H2O₋1 in the northern hemisphere lower stratosphere (≈50 hPa) range from 0.5 ppmv to 0.8 ppmv, depending on season (Fig. 5). The total ozone depletion averaged over the northern hemisphere is ≈-0.5% in H2O₋1. In this respect the results of H2O₋1 are within the range of IPCC (1999).

Figure 14 shows the total column ozone decrease for different geographical regions for simulation H2O_1 and H2O_5. In agreement with the above mentioned results the ozone response to the stratospheric water vapor increase is almost linear. Taking into account the simulated water vapor increase of +0.029 ppmv/yr between 1980 and

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2000 (Fig. 1), this increase corresponds to a stratospheric water vapor perturbation of approximately +0.6 ppmv. According to the linear behavior of the ozone response, at least on hemispheric scales, a perturbation of +0.6 ppmv leads to a global total column ozone reduction of -0.5% (indicated by the vertical dash-dotted line in Fig. 14).

- ⁵ The simulated ozone decline between 1980 and 2000 amounts –3.4%, considering the increasing chlorine loading and climate change. Thus the stratospheric water vapor increase plays a significant role in ozone decline. Dvortsov and Solomon (2001) analyzed the response of stratospheric temperatures and ozone to past (+1%/yr) and future increases in stratospheric humidity with a 2-D radiative-chemical-dynamical
- model. Their results show an additional depletion of mid-latitude column ozone of -0.3%/decade caused by the stratospheric water vapor increase. So our results are in good agreement with the findings of Dvortsov and Solomon (2001). A prolonged increase in stratospheric water vapor could have a significant effect on stratospheric ozone depletion: the chemical effect of the water vapor increase and the cooling effect
 on stratospheric temperatures together lead to a further ozone depletion, which could delay the ozone recovery caused by the expected future reduction of the stratospheric
 - chlorine loading (Shindell et al., 1998).

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Table 1. Overview of analyzed model experiments.

EXP	feedback mode	H ₂ O perturbation	simulation period
CNTL	without feedback	0 ppmv, reference simulation	11 years
VOLC	without feedback	2 ppmv, July and August, short-term increase	5 annual cycles July-June (last 5 years of CNTL)
H2O_1	without feedback	1 ppmv, long-term increase	11 years
H2O_5	without feedback	5 ppmv, long-term increase	11 years

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Table 2. Mixing ratios of greenhouse gases adopted for the year 2000 as recommended by IPCC (2001).

CO ₂ (ppmv)	CH ₄ (ppmv)	N ₂ O (ppbv)	Cl _y (ppbv)
375	1.76	316	2.7

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Table 3. NO_x emissions adopted for the year 2000 (*taking into account IPCC (2001) rates of increase).

NO _x source	Global source (Tg(N)/yr)	Reference
Air traffic	0.7	Schmitt and Brunner (1997)*
Lightning	~5	Grewe et al. (2001)
Industry, Traffic	33.0	Benkovitz et al. (1996)*
Soils	5.6	Yienger and Levy (1995)*
Biomass burning	7.1	Hao et al. (1990)*

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Table 4. Ozone destroying cycles/reactions at 50 hPa, different latitudes and seasons. CNTL: Contribution of each reaction to the total ozone destruction [%]. VOLC, H2O_1 and H2O_5: Changes compared to CNTL [%]. The term O_3 -Loss includes all ozone destroying reactions considered in E39/C.

Annual Mean, Tropics					Annual Mean, Mid-Latitudes				
O ₃ -Loss	HO_{x}	NO_x	CIO _x	$H_2O + O(^1D)$	O_3 -Loss	HO_x	NO_x	CIO _x	$H_2O + O(^1D)$
_	77.1	14.0	1.4	2.9	-	60.8	19.6	8.4	1.4
+5.1	+6.4	-1.6	+7.8	+17.0	+2.4	+4.1	-2.2	+2.7	+12.7
+25.8	+29.0	-7.7	+114.9	+86.0	+11.0	+19.0	-7.0	+3.4	+64.0
+5.0	+6.0	-0.2	+5.9	+17.0	+0.6	+0.9	-0.07	+0.03	+2.6
+1.8	+2.3	-1.2	+4.0	+5.6	+0.9	+1.5	-0.8	+1.3	+4.4
Arctic Spring, Polar Regions				ons	Antarctic Spring, Polar Regions				
O ₃ -Loss	HOx	NO_x	CIO _x	H ₂ O + O(¹ D)	O_3 -Loss	HO_x	NO_x	CIO _x	$H_2O + O(^1D)$
-	26.8	47.7	7.9	0.3	-	6.7	4.0	88.7	0.07
-0.9	+5.6	-6.7	+12.4	+10.5	+38.9	-3.4	-47.4	+46.4	+4.3
-1.5	+23.3	-17.5	+16.5	+55.7	+126.6	-22.8	-94.9	+149.1	+35.9
_	O ₃ -Loss - +5.1 +25.8 +5.0 +1.8 O ₃ -Loss - -0.9 -1.5	$\begin{array}{c c} & & & \\ & & \\ \text{O}_3\text{-Loss} & \text{HO}_x \\ \hline & & & \\ +5.1 & +6.4 \\ +25.8 & +29.0 \\ \hline & +5.0 & +6.0 \\ +1.8 & +2.3 \\ \hline & & \\ \text{Arctic S} \\ \hline & & \\ \text{O}_3\text{-Loss} & \text{HO}_x \\ \hline & & \\ \hline & & \\ \text{O}_3\text{-Loss} & \text{HO}_x \\ \hline & & \\ \hline & & \\ - & & \\ 26.8 \\ -0.9 & +5.6 \\ -1.5 & +23.3 \\ \end{array}$	$\begin{array}{c c c c c c c c c c c c c c c c c c c $	$\begin{array}{c c c c c c c c c c c c c c c c c c c $	$\begin{array}{c c c c c c c c } \hline & \text{Annual Mean, Tropics} \\ \hline O_3\text{-Loss} & HO_x & NO_x & CIO_x & H_2O+O(^1D) \\ \hline - & 77.1 & 14.0 & 1.4 & 2.9 \\ +5.1 & +6.4 & -1.6 & +7.8 & +17.0 \\ +25.8 & +29.0 & -7.7 & +114.9 & +86.0 \\ \hline +5.0 & +6.0 & -0.2 & +5.9 & +17.0 \\ +1.8 & +2.3 & -1.2 & +4.0 & +5.6 \\ \hline & \text{Arctic Spring, Polar Regions} \\ \hline O_3\text{-Loss} & HO_x & NO_x & CIO_x & H_2O+O(^1D) \\ \hline - & 26.8 & 47.7 & 7.9 & 0.3 \\ -0.9 & +5.6 & -6.7 & +12.4 & +10.5 \\ -1.5 & +23.3 & -17.5 & +16.5 & +55.7 \\ \hline \end{array}$	$\begin{array}{c c c c c c c c c c c c c c c c c c c $	$\begin{array}{c c c c c c c c c c c c c c c c c c c $	$\begin{array}{c c c c c c c c c c c c c c c c c c c $	$\begin{array}{c c c c c c c c c c c c c c c c c c c $

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Fig. 1. Time series of individual water vapor soundings with the CMDL frostpoint hygrometer between 24 and 26 km at Boulder, CO (black) (Oltmans et al., 2000), and of monthly mean water vapor mixing ratios from ECHAM4.L39(DLR)/CHEM at 40° N, 20 hPa (red). The linear trend with the 95% confidence interval is 0.044±0.012 ppmv/yr for Boulder and 0.029±0.007 ppmv/yr for the model simulation. The dashed line indicates enhanced water vapor mixing ratios over Boulder in 1992 after the Pinatubo eruption.



Fig. 2. (a) Modeled total ozone deviations, deseasonalized with respect to the period 1979– 1987 and expressed as percentages of the ground-based time average for the period 1964-1980, area-weighted mean 90° N– 90° S (see Fioletov et al. (2002) for details of analysis). Red line shows a 13-month running mean. (b) Modeled total ozone deviations within the tropics (10° N– 10° S) averaged over the last three major volcanic eruptions, expressed as percentages of the average for the period 1964–1980. Month 0 indicates the minimum ozone value after the volcanic eruptions. The eruptions occurred between 8 and 23 months earlier. The red stars indicate the estimated reduction of the ozone column caused by the additional uplift during a volcanic period applying a linearized transport-chemistry column model.

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Fig. 3. Zonally and monthly averaged distribution of OH $(10^5 \text{ molecules/cm}^3)$ for January (a), April (b), July (c) and October (d) as modeled with E39/C. The figure shows a 10-year-climatology, 2000 climate.

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Fig. 4. Five annual cycles of the zonal mean water vapor perturbation (ppmv) at 89 hPa, 30° N/S (upper panel) and 60° N/S (lower panel) of the simulation VOLC. Solid line: northern hemisphere; dashed line: southern hemisphere.

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Fig. 5. Zonally averaged volume mixing ratio of the water vapor perturbation (ppmv). This figure displays the mean over 5 analyzed years, simulation H2O_5. The gray-shaded area marks the tropical "source region" of the water vapor perturbation.

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Fig. 6. Relative changes (%) of zonal mean OH concentration at 50 hPa (upper panel) and 80 hPa (lower panel) depending on latitude and season, VOLC-CNTL/CNTL. Heavy (light) shaded areas indicate the 99% (95%) significance level of the displayed changes (*t*-test).

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Fig. 7. Monthly and zonally averaged changes of OH (%) for August, VOLC-CNTL/CNTL. Shaded areas again indicate the significance level.

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Fig. 8. Percentage increase/decrease of different ozone destroying cycles/reactions in VOLC compared to CNTL at 50 hPa. Upper panel: tropics, lower panel: sum of northern and southern mid-latitudes.

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Fig. 9. Relative changes (0.01%) of zonal mean total column ozone depending on latitude and season, VOLC-CNTL/CNTL. Heavy (light) shaded areas indicate the 99% (95%) significance level of the displayed changes (*t*-test).

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Fig. 12. Relative changes (%) of climatological zonal mean total column ozone depending on latitude and season, simulation H2O_5-CNTL/CNTL. Heavy (light) shaded areas indicate the 99% (95%) significance level of the displayed changes (*t*-test).

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Fig. 13. Relative changes (%) of the surface dose at 309 nm (lower panel) in H2O_5 compared to CNTL. Black areas indicate missing data during polar night.

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Fig. 14. Total column ozone reduction (%) for different geographical regions depending on the stratospheric water vapor perturbation.