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Influence of galactic cosmic rays on atmospheric composition and temperature

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Abstract

This study investigates the influence of the galactic cosmic rays (GCRs) on the atmospheric composition, temperature and dynamics by means of the 3-D Chemistry Climate Model (CCM) SOCOL v2.0. Ionization rates were parameterized according
to CRAC:CRII (Cosmic Ray induced Cascade: Application for Cosmic Ray Induced Ionization), a detailed state-of-the-art model describing the effects of GCRs in the entire altitude range of the CCM from 0–80 km. We find statistically significant effects of GCRs on tropospheric and stratospheric NO_x, HO_x, ozone, temperature and zonal wind, whereas NO_x, HO_x and ozone are annually averaged and the temperature and the zonal wind are monthly averaged. In the Southern Hemisphere, the model suggests the GCR-induced NO_x increase to exceed 10% in the tropopause region (peaking with 20% at the pole), whereas HO_x is showing a decrease of about 3% caused by enhanced conversion into HNO₃. As a consequence, ozone is increasing by up to 3% in the relatively unpolluted southern troposphere, where its production is sensitive to ad-

- ¹⁵ ditional NO_x from GCRs. Conversely, in the northern polar lower stratosphere, GCRs are found to decrease O₃ by up to 3%, caused by the additional heterogeneous chlorine activation via $CIONO_2$ +HCl following GCR-induced production of $CIONO_2$. There is an apparent GCR-induced acceleration of the zonal wind of up to 5 m/s in the Northern Hemisphere below 40 km in February, and a deceleration at higher altitudes with
- ²⁰ peak values of 3 m/s around 70 km altitude. The model also indentifies GCR-induced changes in the surface air, with warming in the eastern part of Europe and in Russia (up to 2.25 K for March values) and cooling in Siberia and Greenland (by almost 2 K). We show that these surface temperature changes develop even when the GCR-induced ionization is taken into account only above 18 km, suggesting that the stratospheri-
- cally driven strengthening of the polar night jet extends all the way down to the Earth's surface.



1 Introduction

Galactic cosmic rays (GCRs) are energetic particles (mostly protons and α -particles) which originate from outside of the solar system. While their flux outside the solar system can be regarded as roughly isotropic and time independent, at least on the time scales studied here (Usoskin et al., 2004), the intensity of GCRs near the Earth varies as a result of the modulation inside the heliosphere, i.e. the spatial region of about 200 Earth–Sun distances controlled by the solar wind and the solar magnetic field. Variations of the cosmic ray flux depend also on particle energy: the flux of less energetic (<1 GeV) particles varies by an order of magnitude modulated by the solar cycle, while energetic GCRs (above 100 GeV) are hardly modulated (Bazilevskava et al., 2008).

When galactic cosmic rays enter the Earth's atmosphere they collide with the ambient atmospheric gas molecules, thereby ionizing them. In this process they may produce secondary particles, which can be sufficiently energetic to contribute themselves to further ionization of the neutral gases. This leads to the development of an ionization

- ¹⁵ cascade (or shower). The intensity and penetration depth of the cascade depends on the energy of the primary cosmic particles. Cascades of particles with several hundred MeV of kinetic energy may reach the ground. However, due to their charge cosmic ray particles are additionally deflected by the geomagnetic field. Almost all particles can penetrate into the polar region, where the magnetic field lines are perpendicular to
- the ground, whereas only the rare most highly energetic particles with energies above 15 GeV are able to penetrate the lower atmosphere near the equator.

Early models of the cosmic ray induced ionization (CRII) were (semi)empirical (e.g., O'Brien, 1970; Heaps, 1978) or simplified analytical (Vitt and Jackman, 1996; O'Brien, 2005). Nicolet (1975), however, has used data from balloon soundings and ionization chambers to deduce the production rates of nitric oxide in the auroral region. State-ofthe-art models (Usoskin et al., 2004; Desorgher et al., 2005; Usoskin and Kovaltsov, 2006) are based on Monte-Carlo simulations of the atmospheric cascade and can provide 3-D time dependent computations of the CRII.



Between the surface and 25–30 km CRII is the main source of the atmospheric ionization (Bazilevskaya et al., 2008) with the maximum ionization rate caused by the Bragg peak in the stopping power around 15 km (Pfotzer maximum), which is clearly visible in Fig. 1 based on the Usoskin et al. (2010) parameterization (red lines). The ionization rates calculated by means of the parameterization of Heaps (1978), represented by blue lines in Fig. 1, are still used often in modeling work (e.g., Schmidt et al., 2006). However, this parameterization does not reflect the Pfotzer maximum, because it does not cover the range from 0–18 km.

The CRII leads to the production of odd nitrogen. For example, fast secondary electrons (e^{*}) can dissociate the nitrogen molecule, $N_2 + e^* \rightarrow 2N(^2D) + e$, and almost all of the N atoms in the excited ²D state react with O₂, producing nitric oxide, $N(^2D)+O_2 \rightarrow NO+O$. Vitt and Jackman (1996) estimated CRII to produce 3.0 to 3.7×10^{33} molecules of odd nitrogen per year in the global stratosphere, which amounts to about 10% of the NO_x production following N₂O oxidation. They also mention that the northern polar/subpolar stratosphere (>50° N) is believed to be supplied with NO_x in equal amounts by GCRs (7.1 to 9.6×10^{32} molecules/yr) and by N₂O oxidation (9.4 to 10.7×10^{32} molecules/yr). In the deep polar winter stratosphere, when air masses experience sunlit periods only infrequently and photolysis of HNO₃ becomes negligible, CRII become the only source of NO_x, revealing the importance of GCRs in high latitudes.

Below the mesopause, where water cluster ions can form, CRII contributes to the formation of HO_x radicals. For example, molecular oxygen ions (O_2^+) produced by GCRs can via attachment of molecular oxygen form O_4^+ , which reacts with water: $O_4^++H_2O\rightarrow O_2^+ H_2O+O_2$. This hydrated ion quickly hydrates further to produce OH: $O_2^+ H_2O+H_2O\rightarrow H_3O^+ OH+O_2\rightarrow H_3O^++OH+O_2$ (Aikin, 1994). GCR-driven HO_x production competes with the most important source for HO_x in the atmosphere, the photolytically driven oxidation of water vapor (H₂O) by excited oxygen atoms, O(¹D), which are themselves produced from ozone photolysis. However, during polar night, HO_x is mainly produced by the GCRs given that no UV radiation is available for O(¹D)



production.

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The influence of GCRs on atmospheric chemistry has been studied by Krivolutsky et al. (2002) with a 1-D photochemical model. They found that ozone at 50° geomagnetic latitude might indeed be sensitive to the additional NO_x source. Their 1-D model

- ⁵ predicted maximum GCR-induced increases in NO_x of 4.5% around 10 km, enhancing tropospheric ozone by 0.6%, whereas above about 18 km ozone decreases with a maximum reduction of 0.5% close to 20 km. Above 35 km altitude they found no influence caused by the GCRs. The evaluation of the impact of GCRs on atmospheric temperature and dynamics, which adds to the chemical changes, cannot be performed with
- ¹⁰ 1-D model and requires the use of a 3-D chemistry-climate model (CCM) that is capable of describing the coupling between physicochemical processes and large-scale dynamics.

Here we study the effect of CRII using the recently developed CRAC:CRII (Cosmic Ray induced Cascade: Application for Cosmic Ray Induced Ionization) model, and then use the results of this event-based local model to force the global CCM SOCOL, focusing on the impact of CRII-induced NO_x and HO_x on chemistry, temperature and dynamics from the ground to 0.01 hPa barometric pressure (altitude of ~80 km).

We have also addressed the difference between the state-of-the-art parameterization of the ionization rate by Usoskin et al. (2010) and the more traditional parameterization given by Heaps (1978), which was based on fitting results from sealed ionization

tion given by Heaps (1978), which was based on fitting results from sealed ionization chambers flown continuously (yearly) on balloons extending to heights of 35 km. The parameterization by Heaps (1978) was and is widely used in various models (Verronen et al., 2002; Schmidt et al., 2006; Winkler et al., 2009)

Many studies of atmospheric chemistry and dynamics omit the influence of GCRs altogether, as was done for example in the first Chemistry-Climate Model Validation Activity (CCMVal) for coupled CCMs (Eyring et al., 2006) and in the most recent CCMVal report (see the homepage of SPARC: http://www.atmosp.physics.utoronto.ca/SPARC/ ccmval_final/index.php). Here, we use the CCM SOCOL, which is one of the CCMs that participated in the CCMVal activity, to investigate the consequences of this omission.



The models and experimental setup are described in Sect. 2, the results containing the GCR effects on several chemical species and the comparison between the parameterizations by Usoskin et al. (2010) and Heaps (1978) are presented in Sect. 3. In Sect. 4 we give a short summary of the results.

5 2 Description of the model and experimental setup

Chemistry-climate modeling with SOCOL. The CCM SOCOL represents a combination of the global circulation model MA-ECHAM4 and the chemistry-transport model MEZON. MA-ECHAM4 (Manzini et al., 1997) is a spectral model with T30 horizontal truncation resulting in a grid spacing of about 3.75°; in the vertical direction the model has 39 levels in a hybrid sigma-pressure coordinate system spanning the model atmo-

sphere from the surface to 0.01 hPa.

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The chemical-transport part MEZON (Rozanov et al., 1999; Egorova et al., 2003) has the same vertical and horizontal resolution and treats 41 chemical species of the oxygen, hydrogen, nitrogen, carbon, chlorine and bromine groups, which are coupled by 140 gas-phase reactions, 46 photolysis reactions and 16 heterogeneous reactions in/on aqueous sulfuric acid aerosols, water ice and nitric acid trihydrate (NAT). The original version of the CCM SOCOL was described by Egorova et al. (2005).

An extensive evaluation of the CCM SOCOL (Egorova et al., 2005; Eyring et al., 2006, 2007) revealed model deficiencies in the chemical-transport part and led to the development of the CCM SOCOL v2.0 (which is applied in this study). The new features of the SOCOLv2.0 are: (i) all species are transported separately; (ii) the mass fixer correction after each semi-Lagrangian transport step is calculated for the chlorine, bromine and nitrogen families instead for individual family members, but then applied to each individual species; (iii) the mass fixer is applied to ozone only over the lati-

tude band 40° S–40° N to avoid artificial mass loss in the polar areas; (iv) the water vapor removal by the highest ice clouds (between 100 hPa and the tropical cold point tropopause) is explicitly taken into account to prevent an overestimation of stratospheric



water content; (v) the list of ozone-depleting substances is extended to 15 for the chemical treatment, while for the transport they are still clustered into three tracer groups; (vi) the heterogeneous chemistry module was updated to include HNO₃ uptake by aqueous sulfuric acid aerosols, a parameterization of the liquid-phase reactive uptake ⁵ coefficients and the NAT particle number densities are limited by an upper boundary of 5×10^{-4} cm⁻³ to take account of the fact that observed NAT clouds are often strongly supersaturated. A comprehensive description and evaluation of the CCM SOCOL v2.0 is presented by Schraner et al. (2008).

Cosmic ray induced ionization modeling. Here we study the effect of CRII using the recently developed CRAC:CRII model (see Usoskin et al., 2004; Usoskin and Kovaltsov, 2006), which has been extended from the stratosphere (Usoskin and Kovaltsov, 2006) to the upper atmosphere (Usoskin et al., 2010). The model is based on a Monte-Carlo simulation of the atmospheric cascade and reproduces the observed data within 10% accuracy in the troposphere and lower stratosphere (Bazilevskaya

- et al., 2008; Usoskin et al., 2009). In the mesosphere the agreement between observed and simulated ionizations rates are less easily assessed, because the ionization by other sources (solar radiation, precipitating soft particles of magnetospheric origin, etc.) becomes at least as important as by GCRs. The results of the CRAC:CRII model are parameterized to give ion pair production rate as a function of the altitude
- (quantified via the barometric pressure), geomagnetic latitude (quantified via geomagnetic cutoff rigidity) and solar activity (quantified via the modulation potential Φ), see Usoskin et al. (2005). In Fig. 1 we show the ionization rates for several geomagnetic latitudes as computed by the CRAC:CRII model (red line), compared to the ionization rates computed by the parameterization of Heaps (1978) (blue line). Solid lines show
 the ionization rates during solar minimum, the dashed lines during solar maximum.

This parameterization of the ionization rates cannot be directly used in CCM SOCOL, which has no explicit treatment of ion chemistry and requires the conversion of the ionization rates into NO_x and HO_x production rates. Following Porter et al. (1976), we assumed that 1.25 NO_x molecules are produced per ion pair, and 45% of this NO_x



production is assumed to yield ground state atomic nitrogen N(⁴S), whereas 55% yields the electronically excited state atomic nitrogen N(²D). While the ground state may lower the overall NO_x concentration via N(⁴S)+NO \rightarrow N₂+O, N(²D) converts instantaneously to NO (see Introduction).

The production of HO_x has been studied by Solomon and Crutzen (1981) with a 1-D time-dependent model of neutral and ion chemistry. They parameterized the number of odd hydrogen particles produced per ion pair as a function of altitude and ionization for daytime, polar summer conditions of temperature, air density and solar zenith angle. We implement their parameterization in the CCM SOCOL to take into account the GCR
 induced production of HO_x from the ground up to the height of 0.01 hPa barometric pressure (altitude of ~80 km).

For this study, we have carried out three 27-yr long runs of CCM SOCOL v2.0 from 1976 to 2002. The control run has been performed without the influence of the galactic cosmic rays, while two experiment runs include GCRs using the ionization rates given

¹⁵ by Usoskin et al. (2010) and Heaps (1978). The first two years of the runs have been omitted from the analysis to eliminate possible spin up problems of the model. In a final section we compare the results with runs using the often applied CRII parameterization of Heaps (1978).

3 Results

Figures 2–5 show the annual mean response of the zonal mean NO_x , HO_x , HNO_3 and ozone to the GCRs calculated as a relative deviation of the experiment run from the reference run. The figures are limited to the range from 1000 hPa to 1 hPa even though the model reaches up to 0.01 hPa, because there is little influence of the GCRs above 1 hPa.



Chemical species

 NO_x . The galactic cosmic rays produce substantial amounts of NO_x during all seasons (not shown). In the annual mean the simulated NO_x increase affects most of the troposphere, exceeding 20% or 10 pptv in the region extending from the south pole to 60°

- ⁵ N around 8–12 km altitude (see Fig. 2, significant changes at 95% level are marked by hatching). There is also a significant impact on the tropical and subtropical middle stratosphere. The reason for smaller effects on the upper stratosphere lies in the vertical distribution of the ionization rates shown in Fig. 1: the ionization rate is the highest between 15 and 20 km, rendering the production of odd nitrogen by GCRs highest. The difference in circuit the neutron to parthere hereighter to parthere.
- ¹⁰ difference in significance between the southern to northern hemispheric troposphere is explained by the fact that more NO_x is produced anthropogenically in the Northern Hemisphere (NH) than in the Southern Hemisphere (SH), making the GCR-induced signal most relevant in the remote regions in the SH.

The annual mean NO_x production by GCRs in the southern hemispheric polar region (up to 5 pptv NO_x in January south of 70° S) is comparable to or even more important than the natural production through lightning (up to 2 pptv NO_x in January south of 70° S, Penner et al., 1998).

 HO_x and HNO_3 . Figure 3 represents the response of annual mean zonal mean HO_x to GCRs. The HO_x increase due to GCR-induced production is largely compensated or overcompensated by NO_x production followed by HO_x deactivation. Thus, there is no statistically significant HO_x increase except in the upper tropical stratosphere, rather a broad area of significant GCR-induced HO_x reduction in the tropical/mid-latitude UTLS. The inner hatched areas, representing 95% statistical significance, show a decrease of about 3% or 0.1 pptv over the southern hemispheric mid-latitudes at an alti-

tude of ~20 km. The outer hatched areas, representing 80% significance, show a decrease of up to 3%. This broad area of HO_x decrease coincides with a region of the highest NO_x enhancements and can be explained by the more intensive removal of OH via OH+NO₂+M→HNO₃+M, resulting in a significant HNO₃ increase of about 8%



(Fig. 4).

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Ozone. Significant increase of NO_x in the southern hemispheric troposphere leads to the statistically significant ozone enhancement. As mentioned above ozone photochemistry in the southern hemispheric troposphere is in large parts NO_x -limited,

- ⁵ so that the CRII relaxes this limitation leading to up to 3% or 1 ppbv ozone increase (see Fig. 5). Conversely, in the northern polar lowermost stratosphere a significant ozone decrease of more than 3% is caused by the additional production of ClONO₂ via ClO+NO₂+M→ClONO₂+M, which in a second step reacts in heterogeneous reactions on polar stratospheric cloud particles or cold sulfate aerosols to enhance chlorine ac-
- tivation, CIONO₂+HCl^{het}Cl₂+HNO₃ (with subsequent photolysis Cl₂+hv→Cl+Cl). As a secondary effect, higher HNO₃ concentrations in the polar winter stratosphere lead to enhanced polar stratospheric cloud occurrences, and hence to faster heterogeneous chemical processing. The ozone decrease by the activated chlorine is negligible in the southern hemispheric polar region because the background concentration of chlorine
 is too high.

The latitudinal average of our results for O_3 and NO_x resemble the results of simple 1-D model calculations published by Krivolutsky et al. (2002). For ozone they modeled a maximum increase in the troposphere at a height of approximately 10 km and a maximum decrease at about 20 km. For NO and NO_2 , their peak is visible at 10 km. In their work the influence of the GCRs vanishes above 35 km. The hemispheric asymmetries discussed above could, of course, not be retrieved in their 1-D calculation. Also, because Krivolutsky et al. (2002) did not discuss HO_x in their paper, it is not possible to make a quantitative comparison with our results.

Temperatures and winds

The effects of GCRs on monthly mean ozone, temperatures and winds are noticeable year-round. However, significance is highest for winter/spring, hence results are displayed for this season in Fig. 6. The upper panel in Fig. 6 shows the monthly mean



zonal mean changes for ozone during February. The significant area and the percentage of decrease are similar to the annual mean results shown in Fig. 5. A decrease of up to 5% or more than 60 ppbv is visible in the NH polar region between 20 and almost 30 km. The influence of the GCRs on the SH is strongest in the troposphere, but remains statistically insignificant on the 95% level. As discussed above, the reason for the ozone depletion in the NH polar region is the additional GCR-induced chlorine activation.

Temperature profile. The center panel of Fig. 6 shows zonal mean response of the temperature in February. There is a cooling in the NH lowermost stratosphere (below 20 km altitude), resulting from the radiative cooling caused by the ozone loss. This cooling is facing a warming at low altitudes at about 40° N. These two effects lead to an increase in the latitudinal temperature gradient in the lowermost stratosphere. In addition, there is a significant warming between 40 and 50 km in the NH polar region due to an intensification of the polar vortex which leads in turn to the increase of air descent and adiabatic warming of the upper stratosphere.

Zonal wind profile. The influence of GCRs on the monthly mean zonal wind for February shows a significant increase of up to 5 m/s in the NH polar region, peaking in the lower stratosphere and extending all the way to the ground (see lower panel in Fig. 6). The acceleration is caused by the cooling of the polar lower stratosphere due to the GCR-induced polar ozone depletion, opposed to the warming of the northern mid-

the GCR-induced polar ozone depletion, opposed to the warming of the northern midlatitude lowermost stratosphere. These changes increase the meridional temperature gradient, leading to acceleration of the zonal wind in agreement with the thermal wind balance. Intensification of the polar vortex leads in turn to the increase of air descent and adiabatic warming of the upper stratosphere, in turn causing deceleration of the zonal wind (Limpasuvan et al., 2005).

Comparison of the Heaps and CRII parameterizations. As mentioned above, a major difference between the Heaps parameterization and Usoskin's model-based approach is that the ionization rate calculated with the Heaps parameterization is applicable only at altitudes above 18 km, whereas the ionization rates derived by Usoskin extend to



the ground. As described above, a proper description of the ionization rate in the upper troposphere and lower stratosphere is required for a correct simulation in of atmospheric composition, in particular of free tropospheric ozone.

The importance of the accuracy of the GCR parameterizations for ozone is illustrated in Fig. 7. The left panel represents the annual mean effect of GCRs on the zonal average ozone at 70°–90° N given in percent averaged from 1978 to 2002. It reveals that the Heaps parameterization (Heaps, blue line; Usoskin, red line) clearly underestimates the ozone decrease due to the additional NO_x that is produced in the UTLS region, because it neglects the ionization below 18 km altitude. Conversely, the right panel of Fig. 7 shows the importance of the Usoskin scheme for correctly describing the GCR-induced ozone production in the southern hemispheric troposphere.

The upper panel in Fig. 8 shows the monthly mean zonal mean effect of the GCRs on ozone at 70° – 90° N for November and December given in percent whereas the lower panel depicts the changes for February and March. The larger and significant decrease

- ¹⁵ in November at about 30 km with Heaps parameterization is caused through the fact that the ionization rate is larger in the middle stratosphere (see Fig. 1) and that the PCS chemistry is not important yet. This changes in February and March: the lower panel in Fig. 8 shows that the parameterization with Usoskin below altitudes of 20 km shows a larger impact on ozone than the Heaps parameterization which stops at 18 km.
- ²⁰ Finally, we investigate the influence of the two GCR parameterizations on the surface air temperature (SAT) and its connection with the Arctic Oscillation. The upper panels of Fig. 9 show the March monthly mean and the annually averaged changes in SAT for ionization rates calculated by Usoskin et al. (2010), and the lower panels show the January monthly mean and the annually averaged changes but using the parameterization
- by Heaps (1978). The general patterns and intensities for both parameterization are in good agreement: both show a warming over the eastern part of Europe and Russia and a cooling in the high Arctic.

Additionally, we see that the effects of the galactic cosmic rays result in an alternating warming/cooling pattern resembling the typical response of the SAT caused by an



intensification of the polar vortex known as positive phase of Arctic Oscillation (Thompson and Wallace, 1998), termed AO^+ . A resulting interesting question is whether this response is primarily due to GCR-induced stratospheric changes or due to the penetration of the GCRs into the troposphere.

⁵ The presence of the AO⁺-like warming-cooling pattern also for the Heaps parameterization, which ignores GCR-effects below 18 km, corroborates the interpretation of Thompson and Wallace (1998), namely "that under certain conditions, dynamical processes at stratospheric levels can affect the strength of the polar vortex all the way down to the earth's surface...".

10 4 Summary

Based on the 3-D CCM SOCOL v2.0 and on CRAC:CRII (the "Cosmic Ray induced Cascade: Application for Cosmic Ray Induced Ionization") model, we present in this paper a modeling study of the influence of the galactic cosmic rays on atmospheric composition, winds and temperature from 0.01 hPa or approximately 80 km down to the ground.

Our calculations indicate that GCR-induced ionization leads to the following modifications in atmospheric composition, winds (U), atmospheric temperatures (T) and surface air temperatures (SAT). Only results with 95% level of statistical significance are given:

- 20 Southern hemispheric troposphere, pristine conditions:
 - NO_x : increases by more than 20% in the polar region,
 - HO_x : decreases of ~3% in the mid-latitude upper troposphere,
 - HNO₃: increases by more than 10% between the South Pole and subtropics,
 - $O_3:$ increases by up to ~3% throughout the troposphere to 20 km between the South Pole to 20° N,



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- SAT: small patches of (significant) warming up to 0.5 K in Antarctica.

Northern hemispheric troposphere, anthropogenically preconditioned:

- HNO_3 : marginally significant increases in the mid-latitude upper troposphere,
- O3: marginally significant decreases in the polar upper troposphere,
- U: enhancements of the polar night jet by up to 5 m/s at the tropopause with perturbations reaching all the way to the ground,
 - SAT: warming by up to 2.25 K in the eastern part of Europe and Russia and decreases by almost 2 K over Greenland.

Southern hemispheric stratosphere:

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- NO_x : increases by up to 4% in the tropical middle stratosphere,
 - HO_x : decreases by up to 3% caused by $OH+NO_2$ producing HNO_3 in the low latitude lower stratosphere,
 - HNO₃: largely mirroring the HO_x changes, with increases by 4% in the low latitude lower stratosphere,
- ¹⁵ Northern hemispheric stratosphere:
 - NO_x: increases by up to 4% in the tropical middle stratosphere,
 - HO_x and HNO_3 : similar to Southern Hemisphere,
 - O₃: strong loss in the polar lower stratosphere with annual mean mixing ratios decreasing by 3% due to additional chlorine activation (specifically in February decreases up to 5%, corresponding to a loss of >60 ppbv),
 - T: cooling by up to -1.5 K in the lower polar stratosphere, opposed to a slight warming (<+0.5 K) in the tropical lower stratosphere and a moderate warming (<+1.5 K) in the upper polar stratosphere,

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 U: enhancements of the polar night jet by up to 5 m/s resulting from the enhanced meridional temperature gradient in the lower stratosphere, and a decrease by 3 m/s in the mesosphere.

We conclude that for NO_x-limited regions it is important to have a parameterization
for the GCRs that extents to the surface, otherwise important consequences for tropospheric ozone (Fig. 5) and for the oxidation capacity of the troposphere (Fig. 3) will be neglected. Conversely, Galactic cosmic rays appear to affect winds and temperatures in the middle and lower atmosphere in a manner that is governed by the ionization processes in the middle atmosphere alone, i.e. a detailed description of the ionization processes in the troposphere appears to be less important. The comparison between the often applied parameterization of ionization rates derived by Heaps (1978) and the

- state-of-the-art modeling work by Usoskin et al. (2010), which agree largely above but differ below 18 km, reveals that changes in the surface air temperature are to first order independent of the choice of parameterization. This suggests that the changes in
- tropospheric meteorology depend on changes in the stratosphere, i.e. that the acceleration of the polar night jet reaches all the way down to the Earth's surface. This constitutes an example of stratosphere-troposphere coupling. Conversely, tropospheric NO_x and ozone depend strongly on a correct description of the GCRs down to the lowest parts of the troposphere. The simulations with the 3-D chemistry-climate model SO COL show that the influence of the GCRs should not be neglected in investigations of
- the tropospheric and stratospheric chemistry and dynamics.

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Fig. 1. Red lines: ionization rates (number of ion pairs produced per air mass and time unit) for several geomagnetic latitudes as computed by the CRAC:CRII model (Usoskin et al., 2010). Blue lines: ionization rate computed by Heaps (1978). Solid lines: ionization rate during solar maximum. Dashed lines: ionization rate during solar minimum. Note different scales on abscissas in dependence on geomagnetic latitude.



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Interactive Discussion



Fig. 2. Annual mean effect of GCRs on zonal mean NO_x , $([NO_x]_{GCR} - [NO_x]_{control})/[NO_x]_{control}$, in percent $([NO_x]=[NO]+[NO_2])$. Results are averaged from 1978–2002 (after allowing for a 2-yr model spin-up) with appropriate accounting for solar minimum and maximum periods. Solid contours indicate positive, dotted contours negative changes. Hatched areas (enclosed by solid contours) indicate changes with at least 95% statistical significance.































Fig. 7. GCR-induced effects on ozone, $([O_3]_{GCR}-[O_3]_{control})/[O_3]_{control}$, given in percent. Left panels shows the annual mean averaged for 70°–90° N and for 50° S (right). Red line: parameterization by Usoskin et al. (2010). Blue line: parameterization by Heaps (1976). Results are averaged from 1978–2002 (all seasons, after allowing for a 2-yr model spin-up).



Fig. 8. GCR-induced effects on ozone, $([O_3]_{GCR}-[O_3]_{control})/[O_3]_{control}$, given in percent, for 70°– 90° N. Upper panel: November and December; lower panel: February and March. Red lines: parameterization by Usoskin et al. (2010). Blue lines: parameterization by Heaps (1976). Thick solid lines: altitudes where the changes in ozone are significant at 95% level for the respective parameterization. Results are averaged from 1978–2002.





Fig. 9. Effect of GCRs on SAT, $[SAT]_{GCR}-[SAT]_{control}$, given in Kelvin for January monthly mean (left) and annual mean (right). Upper panels: using ionization rate modeled by Usoskin et al. (2010). Lower panels: using parameterization by Heaps (1978). Results are averaged from 1978–2002 (after allowing for a 2-yr model spin-up). Reddish colors: positive changes. Bluish colors: negative changes. Hatched areas (enclosed by thick solid contours) indicate changes with at least 95% statistical significance.

