Variability in Antarctic ozone loss in the last decade (2004–2013): high-resolution simulations compared to Aura MLS observations

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Abstract. A detailed analysis of the polar ozone loss processes during 10 recent Antarctic winters is presented with high-resolution MIMOSA–CHIM (Modèle Isentrope du transport Méso-échelle de l’Ozone Stratosphérique par Advection avec CHIMie) model simulations and high-frequency polar vortex observations from the Aura microwave limb sounder (MLS) instrument. The high-frequency measurements and simulations help to characterize the winters and assist the interpretation of interannual variability better than either data or simulations alone. Our model results for the Antarctic winters of 2004–2013 show that chemical ozone loss starts in the edge region of the vortex at equivalent latitudes (EqLs) of 65–67°S in mid-June–July. The loss progresses with time at higher EqLs and intensifies during August–September over the range 400–600 K. The loss peaks in late September–early October, when all EqLs (65–83°S) show a similar loss and the maximum loss (> 2 ppmv – parts per million by volume) is found over a broad vertical range of 475–550 K. In the lower stratosphere, most winters show similar ozone loss and production rates. In general, at 500 K, the loss rates are about 2–3 ppbv sh⁻¹ (parts per billion by volume per sunlit hour) in July and 4–5 ppbv sh⁻¹ in August–mid-September, while they drop rapidly to 0 by mid-October. In the middle stratosphere, the loss rates are about 3–5 ppbv sh⁻¹ in July–August and October at 675 K. On average, the MIMOSA–CHIM simulations show that the very cold winters of 2005 and 2006 exhibit a maximum loss of ~3.5 ppmv around 550 K or about 149–173 DU over 350–850 K, and the warmer winters of 2004, 2010, and 2012 show a loss of ~2.6 ppmv around 475–500 K or 131–154 DU over 350–850 K. The winters of 2007, 2008, and 2011 were moderately cold, and thus both ozone loss and peak loss altitudes are between these two ranges (3 ppmv around 500 K or 150 ± 10 DU). The modeled ozone loss values are in reasonably good agreement with those estimated from Aura MLS measurements, but the model underestimates the observed ClO, largely due to the slower vertical descent in the model during spring.

1 Introduction

Since the early 1980s significant losses in stratospheric ozone have been observed in the Antarctic spring (WMO, 2014). The low temperatures (below 195 K) in the polar winter induce the formation of polar stratospheric clouds in the 15–21 km altitude region, on which chlorine and bromine are activated; when the sun returns over the region, the chlorine- and bromine-induced catalytic ozone loss takes place. Since the 1970s, anthropogenic activities have gradually increased the concentrations of ozone-depleting substances (ODSs) in the atmosphere, including chlorine- and bromine-containing compounds, until they peaked around 2000 in the polar regions. The springtime ozone loss, therefore, has correspondingly increased since the late 1980s and was saturated by the early 1990s (WMO, 2014). In the Southern Hemisphere, the land and ocean contrast is smaller than in the Northern Hemisphere, and, hence, the generation and propagation of large-
amplitude planetary waves are not very effective. Therefore, the polar vortex that forms in fall and winter in the stratosphere is relatively undisturbed and stable, and typically lasts for more than 7 (May–November) months. These dynamical conditions further aggravate the ozone loss in the southern polar vortex region. Because of the suppressed wave and dynamical disturbances, and the concomitant stability of the polar vortex, the interannual variation in Antarctic ozone loss since the 1990s has been small.

Large variability in Antarctic ozone loss has been seen in the last few years (2004–2013) relative to other winters since 1992 (e.g., Tilmes et al., 2006; Huck et al., 2007; Yang et al., 2006; Santee et al., 2008a). For instance, the winters of 2004, 2010, and 2012 were relatively warm, with minor warmings and, hence, limited ozone loss (Santee et al., 2005; de Laat and van Weele, 2011; WMO, 2014). The first fortnight of August 2005 was unusually cold and showed a high rate of ozone loss and an unprecedented ozone hole (WMO, 2014). The winter of 2006 was one of the coldest and, hence, the Antarctic vortex experienced the largest ozone hole to date (Santee et al., 2011; WMO, 2014). The winters of 2007, 2009, and 2013 were characterized by average temperatures and, hence, ozone holes of a moderate size (Tully et al., 2008; Kuttippurath et al., 2013). However, the winters of 2008 and 2011 were again very cold and characterized by large ozone holes (Tully et al., 2011; WMO, 2014). Here, we provide a detailed view of these 10 winters in relation to polar processing and the chemistry of ozone loss. In this study, we discuss (i) the interannual variability in ozone loss and chlorine activation and (ii) horizontal, vertical, and seasonal variability in ozone loss in the Antarctic stratosphere during these (2004–2013) winters. We use high-resolution simulations for analyzing the polar processing and interannual changes in ozone loss in detail. Note that the simulations are highly resolved in the lower stratosphere (about 0.5 km between 425 and 550 K) too, to closely study the ozone loss features in those peak ozone loss altitude layers. Additionally, the past 10 winters offer a good opportunity to test the chemical and dynamical processes in numerical models. Furthermore, observations from the Aura microwave limb sounder (MLS) (Froidevaux et al., 2008; Santee et al., 2008b), one of the best satellite instruments currently available for sampling polar vortices, are compared to the model results. Therefore, for the first time ozone loss and chlorine activation can be studied with high-resolution measurements that have very good spatial and temporal coverage inside the Antarctic vortex. Previous satellite measurements were relatively limited to a small temporal and spatial area as far as high-latitude observations are concerned (e.g., Tilmes et al., 2006; Hoppel et al., 2005). While the Upper Atmosphere Research Satellite MLS (Waters et al., 1999) had a similar latitudinal coverage, the frequency of its polar measurements was lower than that of Aura MLS (e.g., Livesey et al., 2013; Froidevaux et al., 2008). Therefore, the study with high-resolution simulations (both horizontally and vertically) together with the high-resolution measurements offers some new insights into the polar processing and ozone loss features of the Antarctic stratosphere.

In this article, the MIMOSA–CHIM (Modèle Isentropique du transport Méso-échelle de l’Ozone Stratosphérique par Advection avec CHEMie) chemical transport model (CTM) (e.g., Kuttippurath et al., 2010; Tripathi et al., 2007) is used to simulate the chemical constituents for the period 2004–2013. The simulated results are then compared to the Aura MLS measurements. We first look at the ozone loss evolution within different equivalent latitudes (EqLs) averaged over 10 winters in Sect. 3.1. The assessment of the interannual variability in the ozone loss, chlorine activation, and ozone loss rates during the 10 (2004–2013) winters is presented in Sect. 3.2. Finally, Sect. 4 concludes with our main findings.

2 Simulations and measurements

We use the high-resolution MIMOSA–CHIM CTM for the simulations of chemical constituents (e.g., Kuttippurath et al., 2010; Tripathi et al., 2007). The model extends horizontally from 10° N to 90° S with 1° × 1° resolution, and there are 25 isentropic vertical levels between 350 and 950 K with a resolution of about 0.5–2 km, depending on altitude. The model is forced by European Centre for Medium-Range Weather Forecasts (ECMWF) operational analyses. The vertical levels of this data set have been changed from 60 to 91 levels from February 2006 to May 2013 and then to 137 levels from June 2013 onward. The chemical fields used to initialize MIMOSA–CHIM each year are obtained from a long-term simulation of the REPROBUS ( Reactive Processes Ruling the Ozone Budget in the Stratosphere) CTM (Lefèvre et al., 1994) driven by the same ECMWF analyses. The REPROBUS model is equivalent to a lower-resolution version of the MIMOSA–CHIM model, with its resolution being 2° × 2°. The chemical tendencies are calculated every 15 min and the output is written for every 6 h. The passive tracer is initialized only once at the beginning of the simulation, and thus the ozone loss is the cumulative loss in ppmv since the beginning of the simulation. The MIMOSA–CHIM CTM uses the Middle Atmosphere Radiation Scheme (MIDRAD) (Shine, 1987). Climatological H2O and CO2, but interactive O3 fields are used for the calculation of heating rates. The kinetics data are taken from Sander et al. (2011) but the Cl2O2 photolysis cross sections from Burkholder et al. (1990), with a log-linear extrapolation up to 450 nm (Stimpfle et al., 2004). These cross sections are in good agreement with the recent Cl2O2 measurements by Papanastasiou et al. (2009), which form the basis for the JPL 2011 recommendations (i.e., Sander et al., 2011). For each Antarctic winter the model was run from 1 May to 31 October.

The model has detailed polar stratospheric cloud (PSC) formation and growth and sedimentation schemes. The saturation vapor pressure provided by Hanson and Mauersberger (1988) and Murray (1967) is used to assume the existence
of nitric acid trihydrate (NAT) and ice particles, respectively. Liquid supercooled sulfuric acid aerosols, NAT, and ice particles are considered to be in equilibrium with the gas phase (Lefèvre et al., 1998). Equilibrium composition and volume of binary (H$_2$SO$_4$–H$_2$O) and ternary (HNO$_3$–H$_2$SO$_4$–H$_2$O) droplets are calculated using an analytic expression given by Carslaw et al. (1995). For NAT and ice particles, the number density is set to $5 \times 10^{-3}$ cm$^{-3}$, and the particle diameter is calculated within the scheme from the available volume of HNO$_3$ and H$_2$SO$_4$. A denitrification scheme is also incorporated to diagnose the sedimentation of HNO$_3$-containing particles. In this scheme, the NAT particles are assumed to be in equilibrium with gas phase HNO$_3$. All three types of NAT, ice, and liquid-aerosol particles are considered, and the sedimentation speed of the particles is calculated according to Pruppacher and Klett (2010).

We use the Aura MLS measurements version (v) 3.3 (Livesey et al., 2013) for a comparison with simulations. The ozone measurements have a vertical resolution of 2.5–3 km over 215–0.02 hPa, and the vertical resolution of ClO measurements is about 3–3.5 km over 100–1 hPa. The time resolution of the measurements is about 3500 profiles per day. The estimated uncertainty of typical ozone and ClO retrievals is about 5–10% and 10–20%, respectively (Livesey et al., 2013; Santee et al., 2008a; Froidevaux et al., 2008). For a comparison with the measurements, the model results (6 h output) are interpolated to the measurement locations.

### 3 Results and discussion

The passive tracer method is applied to compute the ozone loss from the simulations and measurements. This requires the simulation of passive ozone (or a passive model tracer), i.e., ozone simulations without interactive chemistry. The ozone loss is computed as the difference between the modeled passive odd-oxygen tracer (defined as the sum of O$_3$, O(1D) and O$_3$(P)) and measured or simulated ozone. To derive the ozone loss inside the vortex, we use a vortex edge criterion of 65° EqL. The following equation is used to derive the vortex-averaged ozone loss rates and production rates from the model results.

$$ \frac{\delta O_3(\theta, j)}{\text{ppbv/sh}} = \frac{\sum_{\lambda_{eq}=65}^{\lambda_{eq}=90} \delta O_3(\theta, j, \lambda_{eq}) \times \text{sh}(\theta, j, \lambda_{eq})}{\sum_{\lambda_{eq}=65}^{\lambda_{eq}=90} \text{sh}(\theta, j, \lambda_{eq})} , \quad (1) $$

where $\delta O_3(\theta, j)$ is the ozone loss or production averaged within EqL ($\lambda_{eq} \geq 65\degree$ for each model vertical level ($\theta$) and day ($j$). $\delta O_3(\theta, j, \lambda_{eq})$ is the instantaneous ozone loss or production calculated by the model for each grid point defined by latitude ($\phi$) and longitude ($\psi$) for each $\theta$ and $j$. $\text{sh}(\theta, j, \lambda_{eq})$ is the sunlit hour, which is calculated with respect to solar zenith angle $< 95\degree$ and which varies between 0 and 1 for complete darkness to full illumination. The $\lambda_{eq}$ is computed for each model grid point ($\theta, \phi, \psi$) and for each day using potential-vorticity (PV) data.
3.1 Ozone loss: the 2004–2013 average

To elucidate various chemical ozone loss features, we analyze the vertical distribution of the average loss in 2004–2013 for different EqL bins estimated from the model and MLS data from May to October and shown in Fig. 1. The EqL-based analyses extend from 65 to 83° S in 2° increments as calculated for the MLS measurements. A detailed discussion on the calculation of EqLs and determination of the polar vortex edge can be found in Nash et al. (1996) and references therein. The simulations show that the chemical loss starts at lower EqLs of 65–67° S at the edge of the Antarctic vortex in May, above 600 K, as also shown by Lee et al. (2000), Godin et al. (2000), and Roscoe et al. (2012). It moves down to the lower altitudes by July, and the loss is largest at 65–67° S EqL. The loss continues to increase in August, with the EqLs of 65 and 83° S showing the largest and smallest loss, respectively, in accordance with the increase in incidence of sunlight over the region. A clear difference in the amount of ozone loss estimated at different EqLs is well simulated. At the edge of the vortex, a maximum modeled ozone loss of 2.1 ppmv (parts per million by volume) is found around 500–550 K, while in the 70–80° S EqL range the peak loss is found above 600 K.

The ozone loss continues through to September, when all EqLs show a large loss. The largest modeled loss is still found in the lower EqLs of 65–69° S, reaching about 3 ppmv at 500 K. The higher EqLs (77–83° S) show the smallest ozone loss and it peaks in the middle stratosphere (600 K), while other EqLs show their peak loss below 575 K. Above 600 K, all EqLs show a similar ozone loss of about 1.4 ppmv. The maximum loss is about 3 ppmv over 65–70° S at 500 K, 2.5 ppmv over 70–75° S at 550 K, 1.7 ppmv over 75–80° S at 575 K, and 1.3 ppmv over 80–83° S at 600 K, and thus, the altitude of maximum loss increases with EqL up to September. As expected, the maximum ozone loss is found in October, and all EqLs show more or less the same loss (about 3 ppmv) at the peak loss altitude of around 500 K. In addition, all EqLs show a more or less similar amount of ozone loss with altitude below 500 K in October. Above 500 K, the ozone loss shows slight differences, with the largest loss occurring at the highest EqLs, unlike in months earlier than September. Note that the ozone loss presented here is the monthly mean ozone loss, not the instantaneous loss (see later analyses in Sect. 3.2.3 and 3.2.5), which explains the large ozone loss estimated for late September and October after the chlorine activation period (mid-June to mid-September). In addition, in October, the mixing processes affecting the air mass within the vortex when the ozone loss cycles are stopped also homogenize the ozone distributions.

The analyses with model results below 500 K are in good agreement with those of the MLS measurements. Nevertheless, the loss estimated from the observations is more compact with altitude in October, and the model–measurement differences are larger in September than in other months.

3.2 Interannual variations

We have discussed the general features of the measured and modeled ozone loss evolution in the Antarctic stratosphere. Now we look into the details of the interannual variations in Antarctic ozone loss. It is well-known that low temperatures initiate PSC formation, enabling chlorine activation, which then triggers ozone loss. Therefore, we first analyze the minimum temperature and meteorological situation of each winter in 2004–2013. The link between the prevailing meteorology and chlorine activation of the winters is discussed in the following section, as the amount and extent of activated chlorine are a good indicator of the yearly changes in ozone loss. The year-to-year changes in ozone loss, i.e., the vertical profile of ozone loss, column ozone loss, and the instantaneous ozone loss are discussed in detail with respect to the aforementioned factors in the subsequent sections.

3.2.1 Meteorological conditions

Figure 2 shows the minimum polar cap (50–90° S) temperature extracted from the ECMWF operational analyses for each winter since 2004 at 500 K (∼19 km) potential temperature. The $T_{\text{Nat}}$ and $T_{\text{ICE}}$ thresholds are also marked.

The comparison of ozone loss above 550 K shows that the model overestimates the ozone loss there. The ozone loss of about 0.05–0.08 ppmv that is derived from the measurements in May at 360–370 K is within the estimated error bars of the measurements and therefore insignificant. The differences between measurements and simulations will be discussed further in Sect. 3.2.

Figure 2. Evolution of polar cap (50–90° S) minimum temperature derived from the ECMWF operational analyses for each winter from 2004 to 2013 at 500 K (∼19 km) potential temperature. The $T_{\text{Nat}}$ and $T_{\text{ICE}}$ thresholds are also marked.
In 2006, the temperatures are higher (around 184 K) than in other winters in late June and July, but the September and October temperatures (183–199 K) are very low. In 2007, 2008 and 2009, the evolution of temperature is very similar, but there are minor differences in the details, such as a sudden increase of about 181–183 K in July in 2007. In 2010, minor warmings are apparent in early August (182–186 K) and mid-September (185–190 K). The lowest mid-August to October temperatures at this level are found in 2011 as in the case of 2006, but the highest temperatures in October are found in 2012 with values of 194–212 K. In addition, the winter of 2013 shows the lowest July–August temperatures, with about 177–179 K. In brief, the winters of 2004, 2010 and 2012 can be termed warm winters (de Laat and van Weele, 2011) and 2005, 2006, 2011, and 2013 cold or very cold winters (WMO, 2014). Since the temperature evolution is between these extremes, the other winters can be called moderately cold or moderately warm. Note that the winters are categorized with respect to the temperature evolution at this particular potential temperature, and hence the classification can be slightly different at other altitude levels. Further details about the temperature distribution and meteorology of the winters of 2010–2013 can be found in WMO (2014).
Figure 4. The vortex-averaged (defined by $\geq 65^\circ$ EqL) vertical and temporal evolution of ClO, HCl, HNO$_3$, N$_2$O, O$_3$ and ozone loss from the MIMOSA–CHIM model and MLS measurements. The model results are interpolated with respect to the time and location of the MLS observations inside the vortex and then averaged for the corresponding time period. The data are the average of 10 Antarctic winters (2004–2013) and are smoothed for 7 days. ClO data are not available for early May 2006 and late July and early August 2007. The ClO measurements are selected for 10:00–16:00 for local solar time and solar zenith angles below $89^\circ$. The $y$ axis represents potential temperature in K and is on a logarithmic scale. The white dotted lines represent 500 K ($\sim 19$ km) and 675 K ($\sim 26$ km).

3.2.2 Activated chlorine

Figure 3 compares the simulated and measured ClO at the MLS sampling points inside the vortex, defined here by the area within 65–90$^\circ$ S EqL and with a solar zenith angle (SZA) of less than $89^\circ$, for the Antarctic winters of 2004–2013. In MIMOSA–CHIM, relatively high chlorine activation is found in 2005, when a maximum of 1.6–1.8 ppbv (parts per billion by volume) is simulated from June to late July and early August, consistent with the lower temperatures in that winter (WMO, 2014). The other winters show a more or less similar distribution of ClO and thus, chlorine activation, except in the warm winters of 2010 and 2012. The simulated ClO stands in contrast to the temperature structure and PSC observations in each winter (Pitts et al., 2009), which showed the largest areas of PSCs in the coldest winters of 2005 and 2006. It should be noted that chlorine activation does not depend only on temperature and the occurrence of PSCs but also on the available chlorine reservoirs, which vary from one year to the next (Strahan et al., 2014).

However, when we examine the MLS measurements, they mostly follow the temperature history of each winter, as the observations show the strongest chlorine activation in 2005 and 2006 and the weakest in 2010 and 2012. The highest ClO values ($>1.5$ ppbv) are found from July to the end of September over 450–600 K in the colder winters but around 550 K episodically in July–August in the warmer winters in the observations. Therefore, in contrast to the model results, the vortex-averaged MLS measurements display a clear interannual variation in the chlorine activation. These comparisons show that the model underestimates the observed ClO over the 450–600 K range.

In order to find out the reasons for the differences between simulated and measured ClO, we compared the simulated HCl, N$_2$O, and HNO$_3$ with the MLS observations. Figure 4 illustrates the 10-year average of the vortex mean (defined by $\geq 65^\circ$ EqL) ClO, HCl, HNO$_3$, N$_2$O, O$_3$ and ozone loss from the MIMOSA–CHIM simulations and MLS measurements. The N$_2$O comparisons show that the simulated values are higher than the measured ones, as illustrated by the 50
and 100 ppbv isopleths. This bias in the simulations implies that the vertical diabatic descent in the model is slower than that deduced from MLS observations in polar spring. Consequently, the Cl\textsubscript{v} and thus ClO in the model are relatively smaller. Therefore, the available chlorine or HCl for conversion to activated ClO is smaller in the model. The HCl comparison corroborates this feature of the simulations, as HCl is smaller in the simulations than in the measurements in early and midwinter and in late spring in the lower stratosphere. The deficiency in the vertical descent in the model has clearly affected the ozone simulations. For instance, the statistics of the ozone comparison reveal that the differences in May are negligible at all altitudes. In June–August, the model underestimates the measured ozone by about 0.2–0.5 ppmv below 500 K. In September–October, the model overestimates (0.2–0.4 ppmv) the measured ozone below 450 K, with a peak at around 400 K. However, this overestimation is well below the peak ozone loss altitudes. Above 500 K, the model consistently underestimates the measured ozone from about 0.2 ppmv in June to 0.5–1 ppmv in August–October. Hence the differences at these altitudes are also qualitatively consistent with the slower vertical descent of ozone in the model.

**Figure 5.** Vertical distribution of the vortex-averaged (defined by $\geq 65^\circ$ EqL) ozone loss estimated for the Antarctic winters in 2004–2013. The model results are interpolated with respect to the time and location of the MLS measurements inside the vortex and then averaged for the corresponding day. Left: the ozone loss derived from the difference between the passive ozone and the chemically integrated ozone by MIMOSA–CHIM. Right: the ozone loss derived from the difference between the MIMOSA–CHIM passive ozone and the ozone measured by MLS. Both model results and MLS data are smoothed for 7 days. The $y$ axis represents potential temperature in K. The white dotted lines represent 500 K ($\sim$19 km) and 675 K ($\sim$26 km).
A simulation with a different Cl$_2$O$_2$ recombination rate constant (Nickolaisen et al. (2006) instead of the JPL recommendation, as suggested by von Hobe et al. (2007)), was performed to test the sensitivity of the simulations and to diagnose whether the new simulations reduce the model—measurement differences. However, the ClO results did not improve significantly and hence the original simulations are presented. The HNO$_3$ comparisons also indicate that the denitrification in the model is overestimated, due to the equilibrium PSC scheme of the model, as it forms large NAT particles too readily, causing more sedimentation of PSCs and thus more denitrification.

Note that the relatively low model top could also influence the slower descent or vertical transport in the model. A detailed discussion of the differences in measured and modeled ClO and HCl, chlorine partitioning in the model, and the influence of the lower model top in the simulated results will be presented in a separate paper. Also, there can be small interannual differences in the diabatic descent, depending on the accuracy of the wind fields. These have to be kept in mind while interpreting the simulations. However, in a similar study, Santee et al. (2008a) compared the MLS measurements to SLIMCAT (Single Layer Isentropic Model for Chemistry And Transport) model (Chipperfield, 1999) results and found that their simulations slightly overestimate the ClO measurements for the Antarctic winters of 2004 and 2005. They attributed these differences to the equilibrium PSC scheme of the model. In contrast, our model ClO results underestimate the observations, although using a very similar PSC scheme. It suggests that even if the models use similar PSC schemes, the difference in model dynamics can induce significant changes in the simulated results. Nevertheless, note that the model has performed better in northern hemispheric simulations, where the ClO simulations overestimate the MLS measurements in the 2011 winter (Kuttippurath et al., 2012) but slightly underestimatethem in 2005–2010 (Kuttippurath et al., 2010).

### 3.2.3 Ozone loss: vertical and temporal features

Figure 5 shows the vortex-averaged ozone loss estimated from the model and MLS at the MLS sampling locations inside the vortex in 2004–2013. As discussed in Sect. 3.1, the ozone loss onset (i.e., ozone loss > 0.5 ppmv) in the model occurs in mid-June at altitudes above 550 K and gradually moves down to the lower stratosphere by mid-August. The loss intensifies by mid-August, peaks by late September–early October and slows down thereafter as the ozone recovers through dynamical processes.

In the simulations, as expected, the colder winters of 2005 and 2006 show the early onset of ozone loss, in mid-June. The estimated loss is less than 0.5 ppmv above 675 K until mid-August, increases to 1–1.5 ppmv by mid-September in the lower stratosphere, and peaks at 2.5–3.6 ppmv over 450–600 K by early October, consistent with the temporal and vertical extent of activated ClO in these winters. A maximum ozone loss of around 3.5 ppmv is derived around 550 K in 2005–2006 and about 3 ppmv around 500 K in 2007, 2008 and 2011. A smaller ozone loss is found in the warmer winters of 2004, 2010 and 2012, where the peak loss is about 2.6 ppmv, around 475 K. A similar range of ozone loss, but in a slightly broader vertical extent of 450–600 K, is simulated in 2009 and 2013. Therefore, the center of the peak ozone loss altitude (i.e., loss > 2 ppmv) also shows corresponding variations in agreement with the meteorology of the winters, as it is located around 550 K in very cold winters (e.g., 2005 and 2006), around 500 K in moderately cold winters (e.g., 2007), and around 475 K in warm winters (e.g., 2004 and 2012).

In general, the timing and vertical range of ozone loss in the simulations are similar to those of the observations. The modeled ozone loss onset is in mid-June, except for the colder winters (when it is in mid-May), as discussed previously, and the loss strengthens by August–September and reaches a maximum in early to mid-October, consistent with the maxima of the observations. The large ozone losses observed above 550 K in September–October in the colder winters (e.g., 2005, 2006, 2007, 2008, and 2011) are also reproduced by the model. In contrast to the observed loss, the simulated ozone loss in 2007 is slightly higher than that of 2005, consistent with the temperature during these years at 500 K. However, the model consistently overestimates the measured ozone loss in the middle stratosphere in all years by about 0.2–0.5 ppmv in spring, as the model underestimates the measured ozone by the same amount at these altitudes, primarily due to the slower descent in the model during the period as discussed before. Therefore, the interannual variations are more prominent in the measurements than in the model simulations. Note that since the same passive model
Figure 7. Vortex-averaged (defined by $\geq 65^\circ$ EqL) chemical ozone loss and production rates at 675 K ($\sim 26$ km) and 500 K ($\sim 19$ km) in ppbv per sunlit hour (ppbv sh$^{-1}$) for the Antarctic winters 2004–2013 estimated from the MIMOSA–CHIM model simulations.

tracer affected by the slower diabatic descent in spring is used to derive ozone loss from the measurements, the measured ozone loss is also overestimated. Both the simulations and measurements, however, provide consistent results for the peak ozone loss altitudes in each winter.

The ozone loss derived from the SCanning Imaging Absorption spectroMeter for Atmospheric CHartographY (SCIAMACHY) (Bovensmann et al., 1990) ozone profiles using the vortex descent method shows comparable values to that of the simulations and MLS data for the Antarctic winters of 2004–2008 (Sonkaew et al., 2013). There is also good agreement in peak ozone loss values (around 3–3.5 ppmv) and the differences in the altitudes of maximum loss for various winters. In addition, the large loss above 500 K found in the model and MLS data is also inferred from the SCIAMACHY measurements.

3.2.4 Partial column ozone loss

In order to gain further insights into the interannual variability in ozone loss, we now analyze the column ozone loss derived from simulations and measurements. Since there are also other published results available for comparisons, analyzing partial column loss will assist with the interpretation of interannual changes. We have calculated the column ozone loss from the simulations and observations at the MLS footprints inside the vortex for each winter for the complete altitude range of the model and for the altitude region over which peak loss occurs (400–600 K); results are given in Fig. 6. The ozone loss as computed are the average over the maximum loss period in the Antarctic; from late September to early October (25 September–5 October). The simulated results show the lowest ozone loss in 2004 and the highest in 2005 at 350–850 K, depending on the meteorology and chlorine activation during the winters. However, note that the ozone hole in 2006 was record-breaking, but both the model simulations and measurements show only slightly lower total column ozone loss compared to 2005, due to the comparatively larger chlorine activation in early winter in 2005, as also illustrated in Fig. 3 (see the difference in chlorine activation in 2006 and 2005). Large ozone loss of about 167 DU is also simulated in the winter of 2007 due to relatively strong chlorine activation in that winter (Fig. 3). The other winters show a column ozone loss of about $155 \pm 10$ DU over the same altitude range. On the other hand, the ozone loss derived from MLS observations shows the highest column ozone loss of about 162–166 DU in the very cold winters of 2005 and 2006 and the lowest of around 127–134 DU in the moderately cold winter of 2013 and the warm winters of 2010 and 2012 at 350–850 K; these are in agreement with the meteorology of the winters. Despite the differences in the values, the ozone column loss computed for 400–600 K also shows similar patterns as those discussed for the 350–850 K range. Note that there was strong and prolonged chlorine activation in the coldest winter of 2006 (i.e., ClO was enhanced for slightly longer in this winter than in most other winters). Also, Santee et al. (2011) showed that there was unusually strong chlorine activation in the lowermost stratosphere (around 375 K) that contributed to the record-setting ozone hole and ozone loss in that winter. Therefore, the data exhibit a clear interannual variation of ozone column loss as discussed for the ozone loss profile comparisons in Sect. 3.2.3.
The average partial column ozone depletion above the 550 K level computed from the model and data for the 10 winters is about 50±5 DU. This column ozone loss has to be considered when deriving partial column loss from profiles. This is slightly different from the Arctic, where significant ozone loss occurs mostly in the lower stratosphere over 350–550 K in colder winters and where the depletion above 550 K is limited to ∼19±7 DU (Kuttippurath et al., 2010). The larger Antarctic ozone column loss contribution from higher altitudes (above 550 K) is consistent with the loss estimated above these altitudes, as shown by the ozone profiles in this study for 2004–2013 and in Lemmen et al. (2006) and Hoppel et al. (2005) for a range of Antarctic winters prior to 2004. It is also evident from the maximum ozone loss altitudes, as most Antarctic winters have their peak loss altitudes around 525 K as opposed to 475 K in the Arctic (e.g., Kuttippurath et al., 2012; Tripathi et al., 2007; Grooß et al., 2005a; Rex et al., 2004).

The partial column loss estimated from the Halogen Occultation Experiment ozone measurements (∼172 DU) over 350–600 K (Tilmes et al., 2006) is larger than our results for 2004. Our loss estimates over 350–850 K for 2004–2010 are in reasonable agreement with those derived from ground-based and other satellite total ozone observations in the Antarctic (Kuttippurath et al., 2013). The ozone loss computed from a bias-corrected satellite data set using a parameterized tracer by Huck et al. (2007) for the winter of 2004 also shows a similar estimate. The slight differences amongst various ozone loss values can be due to the differences in the altitude of ozone loss estimates, vortex definition, vortex sampling, and the method used to quantify the loss by the respective studies.

### 3.2.5 Ozone loss and production rates

The interannual variability in ozone loss is further analyzed with the ozone loss and production rates in the model simulations. Figure 7 shows the instantaneous loss and production rates at 675 and 500 K. In general, at 500 K, the loss rates are about 2–3 ppbv sh⁻¹ (ppbv/sunlit hour) in mid-June during the onset of ozone loss, about 3–4 ppbv sh⁻¹ in July as the loss advances to the vortex core (i.e., inside the vortex at higher EqLs) and about 4–5 ppbv sh⁻¹ from August to mid-September during the peak loss period. The loss rates decrease from late September onward and reach 0 by mid-October and stay at near-zero values thereafter. Since the loss rate during the mid-September to October period depends on photochemical ozone production and loss, interannual variability is small in that period, and most winters show loss rates of about 2–5 ppbv sh⁻¹. However, significant year-to-year variations are noted from mid-June to mid-August, as the loss rate depends on the chlorine activation and, hence, on the meteorology of the winters.

The very cold winter of 2006 exhibits an extended period of loss rates of about 4 ppbv sh⁻¹ until early October, while the winter of 2009 shows the shortest span of high loss rates, only until mid-August. The colder winter of 2008 also exhibits high loss rates in most months, May–August in particular. In some winters (e.g., 2004 and 2008) the loss rates in August are also higher than those in September. The lidar measurements of Godin et al. (2001) in the Antarctic winters of 1992–1998, and model studies of Tripathi et al. (2007) and Frielers et al. (2007) in the Antarctic winter of 2003, also show comparable loss rates at 475 K. Our analyses are consistent with the loss rates found in the very cold Arctic winters (1994/1995, 1999/2000, 2004/2005, and 2010/2011) during the peak loss rate period in January–February, for which loss rates of about 5–8 ppbv sh⁻¹ around 450–500 K are estimated (Kuttippurath et al., 2012, 2010; Frielers et al., 2007). No significant ozone production is found at this altitude level.

At 675 K, large interannual variability is found in the ozone loss rates from June to August, which are about 2–5 ppbv sh⁻¹, depending on the day of year and winter. The loss rates are typically about 2 ppbv sh⁻¹ in September and then increase rapidly to 4–6 ppbv sh⁻¹ thereafter. For instance: in 2005, the largest ozone loss rates of 3–5 ppbv sh⁻¹ are simulated in early winter, whereas about 3 ppbv sh⁻¹ is calculated in 2008. The lowest loss rates (about 1–2 ppbv sh⁻¹) among the 10 winters during the ozone hole period (June–September) are found in the warm winters of 2010 and 2012. Note that a similar range of loss rates of 2–7 ppbv sh⁻¹ is also calculated for the colder Arctic winters in late March and mid–to late April in 2010/2011, February–March in 2008/2009 and March in 2004/2005, depending on the day of year (Kuttippurath et al., 2010, 2012).

The production rates at 675 K show large variations from one year to the next, from 0 in mid-August to 7 ppbv sh⁻¹ in late October. These substantial production rates in the September–October period offset the large loss rates during the same period. The high production rates at the end of winter are expected as small disturbances (toward the final warming) shift the polar vortices to sunlit parts of the midlatitudes. The analyses of the ozone production and loss rates at 675 K imply that the ozone loss in the middle stratosphere also depends on the position of the polar vortex in sunlight and the dynamics of the winter.

### 4 Conclusions

The interannual variability in the Antarctic winter meteorology was relatively large in the last decade (2004–2013), which included an extremely cold winter (2006), three moderately cold winters (2005, 2008, and 2011), and three warm winters (2004, 2010, and 2012). As analyzed from the average of the 10 winter simulations, ozone loss in the Antarctic starts at the edge of the vortex at low EqLs (65–67° EqL) by mid-June, consistent with the findings of Lee et al. (2000). Ozone loss progresses with time and advances to higher EqLs (69–83° EqL), with the largest loss at lower EqLs (65–
69° EqL) in June–August in agreement with the exposure of the vortex to sunlight. The maximum ozone loss is attained in the mid-September to mid-October period. The peak ozone loss (> 2 ppmv) is found over a broad altitude range of 475–550 K. The maximum modeled ozone loss is about 3.5 ppbv around 550 K in 2005 and 2006, the coldest winters with the largest loss. In contrast, the maximum loss in the warmer winters of 2004, 2010 and 2012 was restricted to 2.6 ppmv. The modeled column loss shows the largest value of 173 DU in 2005 and the lowest of 110 DU in 2004 over 350–850 K, consistent with the meteorology of the winters. The comparison between simulated and observed trace gas evolution during the winters suggests that the diabatic descent during spring is slower in the model. Therefore, the amount of chlorine available to be activated in spring is lower and hence, the simulated ClO is smaller than the measurements in the lower stratosphere below 550 K, leading to less simulated chemical ozone loss. However, the slower descent also leads to less ozone-rich air being brought down from above. Simulated ozone values reflect a balance between these two effects.

In the lower stratosphere at 300 K, the ozone loss rates have a comparable distribution in all winters, with about 2–3 ppbv sh\textsuperscript{−1} in July and 4–5 ppbv sh\textsuperscript{−1} from August to late September. However, as expected, the very cold winters are characterized by slightly larger and extended periods of high loss rates. In the middle stratosphere at 675 K, a loss rate of about 2–5 ppbv sh\textsuperscript{−1} in July–September, and a production rate of about 4–9 ppbv sh\textsuperscript{−1} in September–October, are simulated. Therefore, these higher production rates largely outweigh the loss rates during the same period.

Our study finds large interannual variability in Antarctic ozone loss in the recent decade with a number of winters showing shallow ozone holes but also with the year with the maximum ozone hole in the last decades. These smaller ozone holes or ozone losses are mainly related to the year-to-year changes in dynamical processes rather than the variations in anthropogenic ODSs, as the change in ODS levels during the study period was very small.

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