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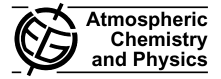
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Chemical ozone loss in the Arctic winter 1991–1992

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Abstract. Chemical ozone loss in winter 1991-1992 is recalculated based on observations of the HALOE satellite instrument, Version 19, ER-2 aircraft measurements and balloon data. HALOE satellite observations are shown to be reliable in the lower stratosphere below 400 K, at altitudes where the measurements are most likely disturbed by the enhanced sulfate aerosol loading, as a result of the Mt. Pinatubo eruption in June 1991. Significant chemical ozone loss (13-17 DU) is observed below 380 K from Kiruna balloon observations and HALOE satellite data between December 1991 and March 1992. For the two winters after the Mt. Pinatubo eruption, HALOE satellite observations show a stronger extent of chemical ozone loss towards lower altitudes compared to other Arctic winters between 1991 and 2003. In spite of already occurring deactivation of chlorine in March 1992, MIPAS-B and LPMA balloon observations indicate that chlorine was still activated at lower altitudes, consistent with observed chemical ozone loss occurring between February and March and April. Large chemical ozone loss of more than 70 DU in the Arctic winter 1991-1992 as calculated in earlier studies is corroborated here.

1 Introduction

The Arctic winter 1991-1992 was a climatological moderately warm winter. For this winter, chemical processes in the polar vortex were strongly influenced by the enhanced burden of sulfate aerosols after the eruption of Mt. Pinatubo in June 1991, in particular at altitudes below $16 \text{ km} (\approx 425 \text{ K})$ (Browell et al., 1993). Strong chlorine activation and significant chemical ozone loss was observed (e.g., Waters et al., 1993; Toohey et al., 1993; Proffitt et al., 1993; Salawitch et al., 1993; Brandtjen et al., 1994). The amount of chemical ozone loss was quantified in several previous studies (e.g., Proffitt et al., 1993; Rex et al., 1998; Müller et al., 2001). These studies were based on different data sets: ER-2 aircraft measurements from the Airborne Arctic Stratospheric Expedition II (AASE-II) (Anderson et al., 1991; Toohey et al., 1993), data from the European Arctic Stratospheric Ozone Experiment (EASOE) (Pyle et al., 1994), namely observations from balloon-borne whole air samplers and ozone sondes. Further, ozone loss was derived using satellite observations from the Halogen Occultation Experiment (HALOE) (Russell et al., 1993) and from the microwave limb sounder (MLS) measurements (Tilmes et al., 2004; Manney et al., 2003), both aboard the UARS satellite.

These studies consistently found ozone loss of $\approx 25\%$ in mid-winter (up to the end of January) and large ozone loss rates during January (Salawitch et al., 1993; von der Gathen et al., 1995). Further, model studies for January indicated largest chemical ozone loss in the outer part of the vortex due to a longer solar exposure (Lefèvre et al., 1994). Müller et al. (2001) and Tilmes et al. (2004) derived



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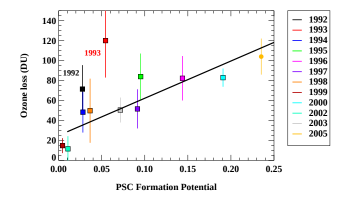


Fig. 1. Relation between the column ozone loss (DU) in March and the PSC formation potential (PFP) for the years 1992 to 2005 between 380–550 K using the tracer-tracer method (colored solid squares), taken from Tilmes et al. (2006) for the Arctic, only. Ozone loss values for March 1992 are taken from this work. The linear relation for the Arctic (calculated excluding winters 1992 and 1993 that are strongly impacted by the eruption of Mt. Pinatubo) is shown as a black line. Moreover, ozone loss values calculated from the O₃ measurements onboard M55 Geophysica (solid circle) (von Hobe et al., 2006) are shown. Here, 1992 refers to the Arctic winter 1991– 1992.

chemical ozone loss from HALOE satellite observations of 72 to 90 DU in February, March and April, using tracer-tracer correlations. Largest loss in column ozone was found below 450 K. However, the burden of aerosol particles in this year was strongly enhanced due to the volcanic eruption of Mt. Pinatubo in June, for altitudes up to 25 km (\approx 700 K) in mid-latitudes, and especially below 16 km (\approx 425 K) in polar regions (Browell et al., 1993). Retrieved HALOE O3 mixing ratios were strongly overestimated in situations of heavy aerosol loading before a correction had been applied (Hervig et al., 1995). After a correction of the data, the uncertainty of O3 mixing ratios was estimated to be 25% under normal conditions (Hervig et al., 1995). Brühl et al. (1996) performed a detailed evaluation of HALOE (Version 17) ozone profiles and reported differences to validation data of no more than 20% in the case of heavy aerosol loading in the lower stratosphere.

It is well established that enhanced stratospheric sulfate aerosol leads to greater halogen induced chemical ozone destruction (e.g., Cox et al., 1994; Tabazadeh et al., 2002; Rex et al., 2004; Tilmes et al., 2004; WMO, 2007). Model simulations by Tabazadeh et al. (2002) predicted a large increase of chemical ozone loss after a strong volcanic eruption, especially at lower altitudes (below 17 km). For the Arctic winters 1991–1992 and 1992–1993, after the eruption of Mt. Pinatubo, Tilmes et al. (2006) reported strong chemical ozone loss based on an analysis of HALOE satellite observations. These values are outliers from the compact empirical relationship between chemical ozone loss and the PSC (polar stratospheric cloud) formation potential $(PFP)^1$ for winters between 1991 and 2005, as shown in Fig. 1.

Winter 1991–1992 was characterized by a relatively small PFP value corresponding to a climatologically moderately warm winter. Based on this relationship, the influence of enhanced sulfate aerosols on chemical ozone loss is about 40 DU for altitudes between 380-550 K potential temperature and about 20 DU between 400-500 K potential temperature in winter 1991-1992 (Tilmes et al., 2006). However, up to now the question remains whether the enhanced chemical ozone loss derived in Tilmes et al. (2006) is the result of chemical processes, or whether observations are distorted by the enhanced sulfate aerosols and feign unreal chemical ozone loss. Rex et al. (2004) reported the impact of enhanced sulfate aerosol on chemical ozone loss to be smaller based on ozone sonde data than in Tilmes et al. (2006) (about 10 DU between 400–550 K) for this winter. For winter 1992–1993, Rex et al. (2006) do not find a significant impact of enhanced sulfate aerosol on ozone loss.

As described above, the presence of the sulfate aerosol in the stratosphere caused by the eruption of Mt. Pinatubo has severely affected the remote sensing measurements of the radiometer channels of the HALOE experiment (Hervig et al., 1995). Here, we will address the question whether the large chemical ozone loss values during spring 1992 based on HALOE observations (Tilmes et al., 2006) are an artifact caused by uncertain observations, or if this much ozone loss can result from chemical processes, which are not included in the PFP value used for the linear relationship in Fig. 1.

The goal of this paper is to derive the most reliable value of chemical ozone loss for the winter 1991–1992. For the first time for this winter, we combine all relevant available insitu observations, which are balloon-borne tracer measurements (Schmidt et al., 1994; Müller et al., 2001) and high-altitude aircraft ER-2 measurements (Proffitt et al., 1993) to validate the HALOE satellite observations. We will show that HALOE observations in winter 1991–1992 are reliable, especially at lower altitudes. This finding should be also conferrable to measurements in winter 1992–1993, a winter that also shows a strongly enhanced aerosol burden.

Tracer-tracer correlations are used to derive chemical ozone loss as described in Sect. 2. Using this technique transport processes within the polar vortex are accounted for (e.g., Tilmes et al., 2004). Compared to other methods, this has the advantage that a possible mis-calculation of transport processes from models for a winter shortly after a volcanic eruption (Robock, 2001) do not affect the calculation of chemical ozone loss. We consider various observations, as there are aircraft, balloon and satellite observations to describe the reliability of the data sets, as described in Sect. 3. The meteorology of the Arctic winter 1991–1992 is described in Sect. 4.

¹PFP presents the fraction of the vortex, over an ozone loss season in Arctic or Antarctica, exposed to PSC temperatures (Tilmes et al., 2006).

Depending on the location of the observations, either inside or outside the polar vortex, air masses indicate different characteristics. The location of measurements with respect to the polar vortex will be discussed in detail in Sect. 5. Based on this information, chemical ozone loss is derived as described in Sect. 6. Further, a comparison of ozone loss profiles for different Arctic winters will be discussed.

Polar chemical ozone loss is a result of chlorine activation in connection with sunlight in the polar vortex. Starting in late December 1991, strong chlorine activation and enhanced ClO were observed until late February in the Arctic lower stratosphere (Waters et al., 1993; Toohey et al., 1993). Enhanced OClO is reported for as late as 11 March (Brandtjen et al., 1994). In Sect. 7, we use balloon measurements in mid-March 1992 (von Clarmann et al., 1993; Oelhaf et al., 1994; Wetzel et al., 1995) to further scrutinize the vertical structure of chlorine activation in March 1992 in the polar vortex.

2 Method

In this study, the tracer-tracer correlation method is used to derive chemical ozone loss in the Arctic polar vortex. The relationship between ozone and a long-lived tracer will not change unless active chlorine (formed by heterogeneous chemistry) causes substantial ozone loss, because of sufficiently long life-times of the tracers in the lower stratosphere in winter. Further, if the polar vortex is isolated and mixing across the vortex edge can be neglected, changes from the relation between ozone and a long-lived tracer in the early winter (the early winter reference function) can be assumed to be a result of chlorine activation and subsequent chemical ozone loss (e.g. Proffitt et al., 1993; Tilmes et al., 2004).

The validity of the tracer-tracer correlations (TRAC) technique as a method to deduce chemical ozone loss has been debated in the literature (e.g., Michelsen et al., 1998; Plumb et al., 2000; Salawitch et al., 2002; Tilmes et al., 2004; Müller et al., 2005, and references therein). Plumb et al. (2000) state that "estimates of ozone depletion inferred from O3:tracer relations are likely to be overestimates" due to the effect of mixing across the vortex edge. As summarized in the following, this concern is not valid under the conditions considered here. In the conceptual model used by Plumb et al. (2000) diffusivities for transport across the vortex edge are employed that are very likely too high by more than an order of magnitude (Müller et al., 2005). Moreover, the conceptual model is formulated in terms of artificial, chemically inert tracers χ_1 and χ_2 . Salawitch et al. (2002) have demonstrated that the model results for χ_1 and χ_2 should not be applied to the interpretation of the O_3/N_2O relation in the Arctic vortex. In the model used by Plumb et al. (2000), the development of the χ_1 - χ_2 relation is driven primarily by supply of air at the top of the vortex with near zero mixing ratios of both species. However, ozone mixing ratios at the top of the vortex are typically greater than those in the vortex after chemical ozone loss occurred. Even though air low in ozone exists in the mesosphere, photochemical model calculations indicate that O_3 is quickly regenerated to mixing ratios of 3–4 ppm by normal gas phase photochemistry as these air parcels descend to lower than \approx 40 km (Salawitch et al., 2002). Indeed ozone mixing ratios ranging between 3.6–5.6 ppm have been measured in the mesosphere in early 2003 (Müller et al., 2007).

To apply the technique carefully (Tilmes et al., 2004; Müller et al., 2005), the location of profiles has to be discussed with respect to the polar vortex edge to understand if they are influenced by air masses from outside the vortex. Profiles which are located within or outside the boundary region of the polar vortex show different characteristics. A detailed description of the technique and a discussion of the uncertainties due to mixing processes is given in Tilmes et al. (2004) and Müller et al. (2005). The polar vortex edge and the poleward edge of the vortex boundary region are calculated as defined by Nash et al. (1996), based on the PV gradient as a function of equivalent latitude. A more precise selection of vortex profiles is performed for aircraft observations with a large horizontal resolution, described in Sect. 5. The area poleward of the vortex boundary region is defined as the vortex core; the area of the vortex outside the vortex core is defined as the vortex boundary region (Tilmes et al., 2004).

3 Observations

In this study, we use satellite observations taken from the HALOE instrument (Russell et al., 1993), Version 19. HALOE measurements made during January, February, March and April 1992 were within the polar vortex core (vortex boundary region for January). HALOE CH₄ and HF are suitable long-lived tracers to apply to the tracer-tracer correlation method as described in Tilmes et al. (2004). The satellite instrument uses gas filter channels to measure these long-lived tracers, different than ozone, which is observed using radiometer channels. The gas filter channels are only weakly affected by aerosol particles and, therefore, a correction of these species for aerosol interference was not necessary (Hervig et al., 1995). Based on a comprehensive validation study, Brühl et al. (1996) reported that ozone mixing ratios have an uncertainty of no more than 20% in the peak aerosol layer in the lower stratosphere (caused by the Mt. Pinatubo eruption) after a correction of the data (Hervig et al., 1995). Although Brühl et al. (1996) used an earlier HALOE version (Version 17) than the one used here (Version 19), the improvements made in the newer data version should not deteriorate the quality of the aerosol correction. Major modifications were made between Version 17 and 18

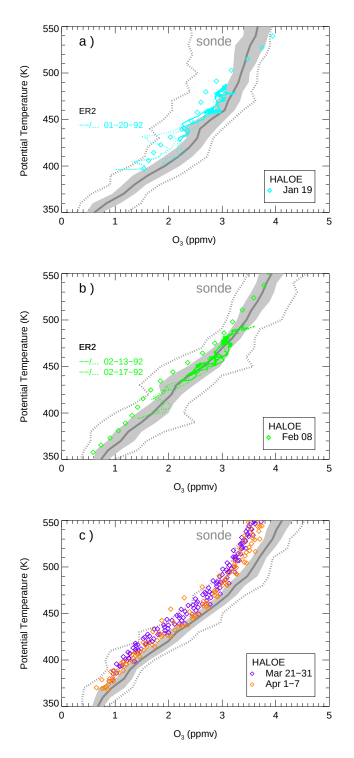


Fig. 2. Ozone mixing ratios are shown with respect to potential temperature taken by the HALOE satellite (diamonds) and ER-2 aircraft observations inside the vortex core (solid lines) and inside the entire vortex (dotted lines) in comparison to ozone sonde data inside the vortex core, for January (top), February (middle) and March 1992 (bottom). The averaged ozone sonde profile is shown as a grey solid line with a standard deviation shaded in grey and maximum and minimum mixing ratios as dotted lines.

and the Version 19 data. Version 19 data have a better accuracy for the retrieved species at altitudes below 70 hPa.

To quantify the uncertainty of HALOE ozone data in the Arctic lower stratosphere in spring 1992, we compare HALOE ozone mixing ratios with a large number of ozone sonde observations taken within the polar vortex core and with ER-2 observations (described below). Fig. 2 shows the average ozone sonde profile and the standard deviation derived for January (panel a), February (panel b) and March (panel c) 1992. Extreme values are shown. Using an established criterion by Nash et al. (1996) to identify vortex profiles, ozone sonde data taken within the polar vortex show a rather homogeneous distribution with a standard deviation between 0.1 and 0.2 ppm for January and March and up to 0.34 ppm in February (grey shaded area in Fig. 2)

For both January and February, only one HALOE profile is available in the vortex. In January, the profile is located close to the polar vortex edge. Ozone mixing ratios are slightly lower compared to the average ozone sonde profile for February for most of the considered altitude range. Especially between 400 and 450 K ozone mixing ratios are up to 0.4 ppm lower (around 420 K) than the average ozone sonde profile. Nevertheless, HALOE mixing ratios are still in the range of minimum observed ozone sonde data. Between 350 and 400 K, a region where HALOE ozone profiles were most strongly influenced by high aerosol loading, profiles in February agree well with ozone sonde data and deviate by less than 0.2 ppm from the mean ozone sonde profile. Additionally, ER2 observations agree well with ozone sonde data.

For March and the beginning of April, HALOE observations are in agreement with ozone sonde data within the range of observed ozone sonde profiles. As in February, HALOE ozone mixing ratios are slightly lower than the averaged ozone sonde profile, by about 0.2 and 0.3 ppm between 380 and 500 K, with less deviation towards lower latitudes. In summary, HALOE ozone measurements are within the range of the ozone sonde measurements in the polar vortex and show a deviation of less than 0.2 ppm below 400 K. This analysis corroborats our conclusions that HALOE ozone observations are reliable in the Arctic polar vortex and ozone loss can be derived from this data set. The influence of a possible uncertainty of 0.25 ppm on derived chemical ozone loss is discussed below.

Besides ozone sonde observations, balloons were launched in Kiruna, Sweden, during the Arctic winter between December 1991 and March 1992 employing cryogenic (Schmidt et al., 1987) and whole air grab sampling techniques (Bauer et al., 1994) that measured long-lived tracers like N₂O and CH₄. Ozone observations were concurrently taken by standard electrochemical-concentration-cell sondes (Pyle et al., 1994). In addition, we use ER-2 aircraft observations of N₂O and CH₄ taken by the ALIAS instrument (Webster et al., 1994) and the ATLAS instrument (Loewenstein et al., 1993) and O₃ from the dual beam photometer (Proffitt and McLaughlin, 1983).

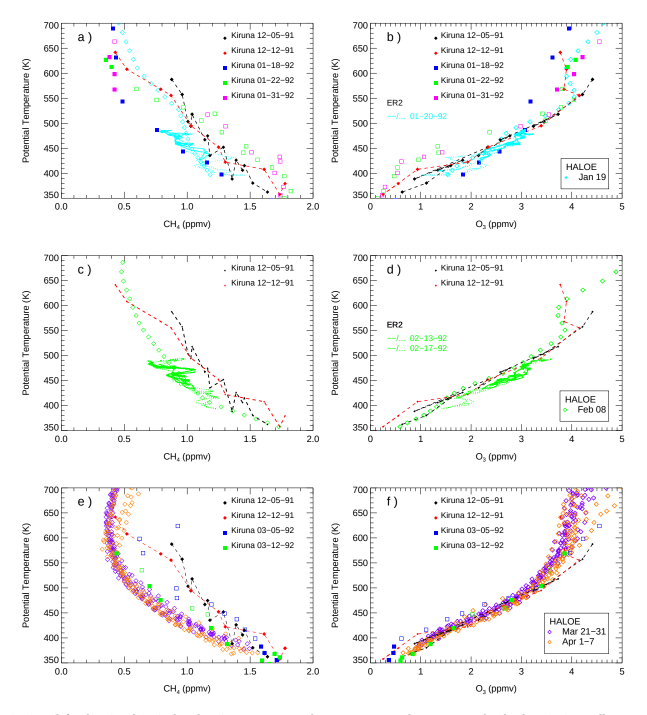


Fig. 3. CH_4 (left column) and O_3 (right column) mixing ratios with respect to potential temperature taken by the HALOE satellite, Kiruna balloons in the vortex core (filled symbols) and outside the vortex core (open symbols), and ER-2 aircraft observations within the vortex core (solid lines) and within the entire vortex (dotted lines), for different periods: January (top), February (middle) and March 1992 (bottom). Kiruna balloon observations during December (red and black symbols and dashed lines) are shown in each panel as early winter reference profiles.

During January and February, the ER-2 observed air masses well within the polar vortex core. In-situ balloon and ER-2 observations of ozone mixing ratios are not influenced by enhanced sulfate aerosols in the lower stratosphere and can be used as a comparison to HALOE satellite observations (as shown in Fig. 2).

We compare O_3 and CH_4 profiles for the different measurements as well as the relationship between O_3/CH_4 . N_2O balloon and aircraft observations were converted to CH_4 mixing ratios using a CH_4/N_2O tracer relation derived using

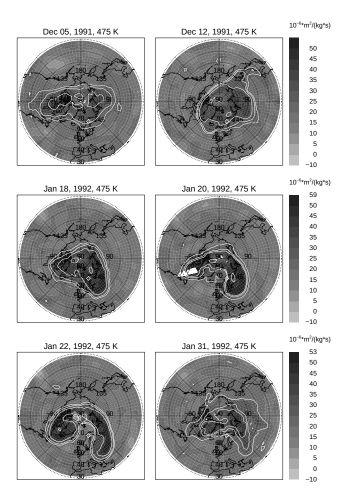


Fig. 4. Potential vorticity derived from MetO analysis on the 475 K potential temperature level valid for 5, 12 December 1991, and 18, 20, 22 and 31 January 1992, 12:00 UTC. White lines indicate the location of the polar vortex core, edge, and outer edge of the vortex as defined by Nash et al. (1996). The location of Kiruna is marked as a white diamond. For the same altitude the location of ER-2 aircraft measurements are presented as white triangles for 20 January, only (taken within 8 hr at the day shown).

whole air sampler measurements, reported in Tilmes et al. (2006). The uncertainty of CH₄ using the CH₄/N₂O tracer relation is $\sigma \approx 50$ ppb and will be included in the error estimate of chemical ozone loss derived below.

Additionally, two balloon-borne experiments were launched from Kiruna on 14 March 1992. MIPAS-B took measurements of several trace gases including CH_4 and $CIONO_2$ and LPMA measured HCl and CH_4 (von Clarmann et al., 1993; Camy-Peyret, 1994; Oelhaf et al., 1994; Wetzel et al., 1995). MIPAS-B is a cryogenic Fourier Transform Spectrometer recording limb emission spectra and LPMA performs solar occultation infrared measurements. For these instruments broadband high spectral resolution spectroscopy was used. The aerosol contributions in the spectra can well be accounted for in the retrieval (e.g., Echle et al., 1998). Earlier, HCl/tracer relations based on HALOE satellite observations were investigated by Tilmes et al. (2004) to estimate possible chlorine activation in the polar vortex.

4 Meteorology of the Arctic vortex 1991–1992

The polar vortex in winter 1991-1992 was cold between November and January and disturbed by several warming pulses (Naujokat et al., 1992). Temperatures were below 195 K (favorable conditions for chlorine activation by PSCs) only during January (Newman et al., 1993; Manney et al., 2003). Owing to the strongly enhanced sulfate aerosol densities in the lower stratosphere, the threshold temperature for chlorine activation increased significantly (Katja Drdla, pers. communication). Temperatures during the first part of February and the first half of March 1992 below 450 K are below this threshold and a potential of chlorine activation exists during this period. At the end of January, a major warming resulted in weaker, westerly zonal winds at 60° N for altitudes between 380 and 550 K (Naujokat et al., 1992). Transport of air into the vortex was reported by Grooß and Müller (2003). During February and March, the vortex was stable and broke down during April.

5 Location of the observations

The characteristic of the distribution of long-lived tracers in the polar region strongly depends on the location of observations. In the polar vortex, air masses descend most significantly at the beginning of the winter. Descent results in larger ozone and lower CH₄ mixing ratios within the polar vortex than outside the vortex below about 600 K, the region important for chemical ozone loss due to catalytic ozone destruction. Additionally, chemical ozone loss occurs during winter and spring within the Arctic vortex, if vortex temperatures reach values allowing chlorine activation, resulting in a decrease of ozone mixing ratios. In the vortex boundary region airmasses are influenced by air from outside the vortex as a result of isentropic mixing and show different distributions compared to airmasses within the polar vortex core. In order to discuss the reliability of satellite data, it is important to compare similar airmasses. We therefore define the location of various observations, based on whether they were observed within the vortex core, in the vortex boundary region or outside the vortex. In Fig. 3, we compare CH₄ and O₃ profiles of available observations in winter 1991–1992.

For HALOE satellite observations, we use the Nash et al. (1996) criterion to distinguish observations that are taken within the polar vortex core, the vortex boundary region and outside the vortex. In the following, only satellite profiles in the vortex and the vortex boundary region (denoted as "entire vortex") are considered. For ER-2 observations, only the part of the flight path within the entire polar vortex is considered. We use the potential vorticity PV, interpolated

on the flight path to distinguish between measurements that have been taken inside and outside the polar vortex, as described in the following. The modified potential vorticity (Lait, 1994; Müller and Günther, 2003) $\Pi = PV \cdot (\theta_0/\theta)^{-\varepsilon}$ is employed, where θ is the potential temperature, $\theta_0 = 475 \text{ K}$ is a reference potential temperature and the exponent ε may be chosen to adjust the scaling to the prevailing temperature profile (Müller and Günther, 2003). Here, we use $\varepsilon = 4.5$, based on the approximately isothermal temperature profiles measured by radiosonde and by the Microwave Temperature Profiler (MTP) aboard the ER-2 in January/February 1992. The potential vorticity at the edge of the vortex is calculated to be PV=29 PVU (potential vorticity units: 1 PVU=10⁻⁶ K $m^2/(kg s)$ and at the edge of vortex core to be PV=34 PVUat 475 K during January and February 1992, using MetO data based on the Nash et al. (1996) criterion. Measurements with Π larger 29 PVU are assigned to the entire vortex (dotted lines) and measurements with Π larger than 34 PVU are assigned to the vortex core (solid lines in Fig. 3).

Some of the available Kiruna balloon profiles represent the airmass composition at various locations relative to the vortex boundary as described by Bauer et al. (1994) and Schmidt et al. (1994). Observations taken inside the vortex core in January and March are shown as solid symbols, data outside the vortex core are shown as open symbols, in the following.

In Fig. 3, CH₄ and O₃ mixing ratios are shown for different periods and different observations. In each panel, a to f, two Kiruna balloon data are shown as black and red dashed lines. The profile taken on 5 December 1991, black line, was located well inside the vortex for all altitudes. The profile taken on 12 December 1991, red line, was partly located inside the polar vortex, for altitudes above 475 K, and partly outside but close to the vortex edge for altitudes at and below 475 K (compare Fig. 4, top row). Kiruna observations within the entire vortex are used as a reference for early winter conditions observed during December.

Figure 3, panels a and b, show ER-2, Kiruna and HALOE observations during January 1992 compared to the two Kiruna balloon observations in December. The Kiruna balloon observations on 18 January are located well inside the vortex, see also Fig. 4, middle, left panel.

Smaller CH₄ mixing ratios compared to the reference profiles are a result of descent of the polar vortex between December and January. Profiles taken on 22 and 31 January 1992, over Kiruna, are located outside the polar vortex at altitudes below 500 K (see Fig. 4, bottom panels). Above 500 K, the measurements show vortex characteristics with smaller CH₄ mixing ratios compared to the reference profiles. The flight direction of the balloon, especially for the 31 of January, moved toward the vortex core. Additionally, ER-2 aircraft observations were taken at locations between outside the vortex and deep inside the polar vortex on 20 January (Fig. 4, middle right panel, white triangles). Observations taken in the boundary area of the vortex show larger CH₄ mixing ratios (dotted lines in Fig. 3, panel a). ER-2 CH₄ mixing ratios and their deviation from the reference in the vortex core are in good agreement with the balloon observations over Kiruna on 18 January 1992. One HALOE satellite profile, taken on 19 January within the vortex boundary region (panel a and b, cyan diamonds), does not show lower CH_4 mixing ratios and was influenced by air from outside the vortex.

The corresponding O_3 profiles observed in January 1992 indicated minor deviations from the reference profile (see Fig. 3, panel b). ER-2 ozone mixing ratios and the balloon profile on January 18, observed within the vortex core, are slightly larger compared to the reference function due to the descent of vortex air masses. Kiruna balloon data on 22 and 31 January and HALOE observations in January show smaller ozone mixing ratios below 500 K. These profiles were influenced by air masses with smaller ozone mixing ratios from outside the vortex, as also discussed by Müller et al. (2001). The smaller ozone mixing ratios on 18 January above 550 K are a result of the influence of out of vortex air.

In Fig. 3 panel c to f, we compare CH_4 and O_3 mixing ratios of HALOE satellite observations taken within the vortex with Kiruna balloon observations (panel e and f) during March and ER-2 observations during February (panel c and d). All HALOE observations show smaller CH_4 mixing ratios compared to the reference profile (panel c and e) and therefore indicate the descent of airmasses within the polar vortex between December and March/April. One HALOE profile was partly observed within the vortex core (below 500 K) on February 8 and several profiles were observed at the end of March and the beginning of April in the polar vortex, as shown in Fig. 5.

ER-2 observations are only available within the polar vortex during January and February. All ER-2 observations during March were obtained outside the polar vortex (not

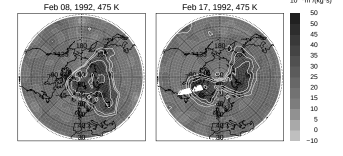


Fig. 5. Potential vorticity derived from MetO analysis on the 475 K potential temperature level valid for 8, 17 February 1992, 12:00 UTC. White lines indicate the location of the polar vortex core, edge, and outer edge of the vortex as defined by Nash et al. (1996). One available HALOE satellite profile inside the vortex core is represented as white squares and ER-2 aircraft measurements are presented as white triangle. The HALOE satellite profiles were repositioned to 12:00 UTC through trajectory calculations, black square.

10⁻⁶*m²/(kg*s)

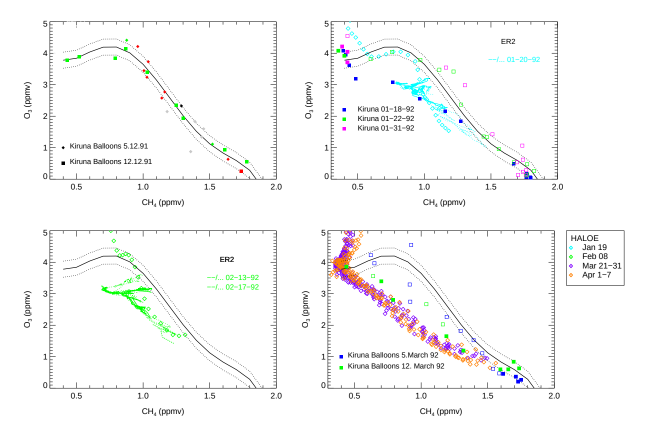


Fig. 6. O_3/CH_4 relations, the early winter reference function, shown as a black line in all four panels, was derived using Kiruna balloon data from 5 and 12 December 1991 (top left panel). Different colors indicate the position of the data, vortex core (green), entire vortex (red), vortex boundary region (black) and outside the vortex (grey). Top right panel: O_3/CH_4 relations during January 1992 using ER-2 aircraft measurements within the entire vortex (cyan dotted lines) and within the vortex core (cyan solid lines), Kiruna balloon data within the vortex core (filled colored squares) and outside the vortex core (open colored squares) and HALOE satellite observations (cyan diamonds) within the polar vortex. Bottom panel: O_3/CH_4 relations during February (right panel) and March / April (left) in 1992 using ER-2 available aircraft measurements within the vortex (green lines) and HALOE satellite observations (colored diamonds) within the polar vortex.

shown). As for January, observations during February were taken at locations in the outer vortex and the vortex core. Observations show smallest CH_4 mixing ratios and largest O_3 mixing ratios at particular theta levels if they were taken deep inside the vortex core. CH_4 profiles taken within the vortex core are in good agreement with HALOE observations. ER-2 ozone mixing ratios (Fig. 3 panel d) are slightly larger (0.2–0.3 ppm) in February compared to HALOE observations for the same potential temperature level. This can be a result of a sampling of air masses that experienced less ozone depletion in the observed part of the vortex compared to HALOE ozone observations in February. Slightly inhomogeneous ozone depletion in the vortex is also reflected by a larger standard deviation of ozone profiles in February, as discussed above (see Fig. 2, panel b).

The Kiruna balloon profile observed on 12 March 1992 (Fig. 3, panel e, green squares) indicates a similar characteristic as the HALOE observations between March and April between 350 and 420 K, at 475 K and above 550 K. During its flight, the balloon moved towards the edge of the vortex

for the other altitudes as described by Bauer et al. (1994) and Schmidt et al. (1994). Ozone mixing ratios of the balloon profile in 12 March 1992 and satellite observations in March and April 1992 are in agreement, see panel f. The balloon profile taken over Kiruna on 5 March was located in the vortex boundary region (Fig. 3, panel e, blue squares) and does not show significant deviations from the reference profiles (black and red lines in Fig. 3) below 500 K. As described above, a deviation from the reference CH₄ profile indicates the measured air mass has descended diabatically. The agreement of the balloon profile on 5 March and the reference profiles, below 500 K, indicates that measured air masses were located outside the vortex where descent did not occur. Therefore, this profile is characterized by air from outside the vortex below 500 K and should not be compared to observations inside the polar vortex.

In summary, CH_4 and O_3 mixing ratios of the considered observations agree well if observed within the polar vortex core. Profiles observed outside the polar vortex core show larger CH_4 mixing ratios and smaller O_3 mixing ratios as a result of the influence of outer vortex air for altitudes below 600 K. It is shown that HALOE O_3 and CH_4 observations are reliable between 350 and 700 K. Observations are in good agreement with ER-2 observations below 550 K potential temperature and with Kiruna balloon observations taken within the polar vortex core in January and March.

6 Chemical ozone loss

6.1 Tracer-tracer correlations

To estimate chemical ozone loss, we derived an O_3/CH_4 early winter reference function based on balloon observations taken on 5 and 12 December 1991, that are located in the entire polar vortex (Fig. 6, top panel). On 12 December, balloon data were measured well inside the polar vortex core (green symbols). On 5 December most of the observations were located within the entire polar vortex (red symbols). Profiles around 400 K were located outside the vortex boundary region (grey squares) and are not included in the calculation of the early winter reference function (Fig. 6, top panel).

This reference function is constructed as a simple unweighted least square polynominal fit to the data points of the order of 5, which is slightly improved compared to Tilmes et al. (2004) and Müller et al. (2001).

$$O_{3} = -12.1793 \cdot (CH_{4})^{5} + 69.3824 \cdot (CH_{4})^{4}$$
(1)
-145.932 \cdot (CH_{4})^{3} + 136.509 \cdot (CH_{4})^{2}
-56.2627 \cdot CH_{4} + 12.1302

The standard deviation of the derived reference function is $\sigma = 0.255$ ppm. The value used to calculate the uncertainty of the loss in column ozone is 0.39 ppm. It was derived by combining (as root mean square) the uncertainty of the reference function $\sigma = 0.255$ ppm, the uncertainty of the CH₄ to N₂O conversion (of 50 ppb CH₄), which results in 0.14 ppm uncertainty of ozone loss, plus the uncertainty of HALOE ozone observations in comparison to ozone sonde data in spring of 0.25 ppm.

During January, ER-2 aircraft observations indicate significant deviation from the early winter reference function, (Fig. 6, top, right panel). Kiruna balloon observations taken within the vortex core on 18 January, as discussed above, agree well with ER-2 observations. Kiruna profiles that are influenced by outer vortex air scatter above the reference function. HALOE observations in January show less deviation from the reference function than ER-2 observations due to the location outside the polar vortex core.

During February, ER-2 observations indicate a larger deviation from the early winter reference function compared to January. Therefore, further chemical ozone loss has occurred. Later in March and April, ER-2 aircraft observations are not considered, because they were taken far outside of the polar vortex (not shown). HALOE satellite ob-

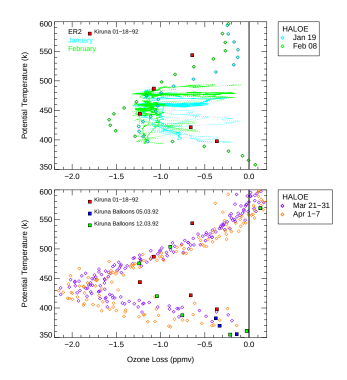


Fig. 7. Ozone loss profiles derived from O_3/CH_4 tracer relations between mid-December 1991 and January/February 1992 (top panel) and mid-December and March/April (bottom panel). HALOE and balloon data observed in the vortex core are shown, as well as ER2 aircraft data in the entire vortex (dotted lines) and in the vortex core (solid lines).

servations during February are in agreement with ER-2 data (Fig. 6 green diamonds and lines). Further, Kiruna balloon data on 12 March show similar deviations from the early winter reference function as HALOE observations obtained in March and April, within the polar vortex core between 350 and 420 K, at 475 K, and above 550 K (as discussed above).

6.2 Local chemical ozone loss

Accumulated ozone loss profiles between December 1991 and observations in January to April 1992 are shown in Fig. 7. During January, local chemical ozone loss up to 1.3 ppm is reached between 400 K and 480 K based on ER-2 data and Kiruna balloon data (red squares). In the vortex boundary region, one HALOE profile shows less ozone loss (up to 0.8 ppm) in January. The observed chemical ozone loss during January is in agreement with the large ozone loss values observed in mid-winter (e.g. von der Gathen et al., 1995). Further, HCl/CH₄ profiles observed by HALOE in the vortex boundary region indicate significant chlorine activation (Tilmes et al., 2004) and strongly enhanced ClO mixing ratios were observed (Waters et al., 1993; Toohey et al., 1993). In January and February a significant increase of local ozone loss is obvious only for one HALOE profile (19 January and 8 February, respectively) measured within the polar

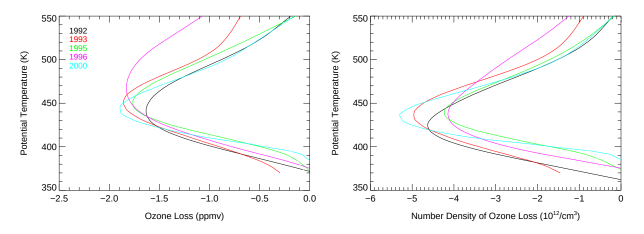


Fig. 8. Averaged ozone loss profiles for March/April for winter 1991–1992, 1992–1993 and cold Arctic winters 1994–1995, 1995–1996 and 1999–2000.

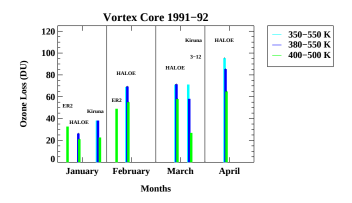


Fig. 9. Accumulated chemical loss in column ozone for different altitude intervals and different months in winter 1991–1992, derived using ER-2 aircraft measurements, HALOE satellite observations, and Kiruna balloon measurements within the vortex core, for different partial columns (see legend). Measurements are placed on the chart based on the month during which they were obtained; the placement within the month is not meant to indicate the precise timing of the observations.

vortex core. Later in March and April, HALOE observations show local ozone loss values between 1.6 and 2.1 ppm between 420 and 460 K. Additional ozone loss is possible during March and April due to possible chlorine activation during March, as described in Sect. 7, and the increase in solar illumination on the polar vortex in spring. Local ozone loss derived using the Kiruna balloon profile on 12 March 1992, (green squares) is in agreement with HALOE observations below 400 K and above 470 K, at altitudes where the balloon was located within the polar vortex core, as discussed above. Especially at altitudes below 380 K, Kiruna balloon observations show large local ozone loss. All observations shown in Fig. 7 are located inside the polar vortex edge.

The impact of enhanced sulfate aerosols in the lower stratosphere after the Mt. Pinatubo eruption is obvious in comparing averaged ozone loss profiles for March/April of different winters between 1991 and 2003 (see Fig. 8). The winter 1991–1992 and 1992–1993 show enhanced ozone loss at altitudes below 400 K, black and red line, respectively. Further, the maximum ozone loss in winter 1991–1992 is located about 10 K below the maximum of the winter 1992–1993 and 1999–2000.

6.3 Chemical loss in column ozone

Chemical ozone loss in winter 1991–1992 is summarized in Fig. 9. Derived ozone loss values have an uncertainty of $\pm 32-38$ DU for altitudes between 350 and 550 K, $\pm 24-$ 27 DU for altitudes between 380 and 550 K, and $\pm 15-16$ DU for altitudes between 400 and 500 K potential temperature. This uncertainty is estimated from the combination of the uncertainty of the early winter reference function (see Fig. 6, black dotted lines), the uncertainty of the N₂O to CH₄ conversion of balloon observations in the early winter, and an uncertainty of HALOE ozone observations in comparison to ozone sonde data in spring, as described above.

The standard deviation of the column ozone loss distribution (using the large amount of HALOE profiles in the vortex core) is 24 DU in March and 13 DU in April, between 380 and 550 K, and 19 DU in March and 7 DU in April, between 400 and 500 K potential temperature.

Between 400–500 K, chemical ozone loss increases from 33 DU in January (observed by ER-2 within the entire vortex) towards 65 DU in April (observed by HALOE), as shown in Fig. 9, green columns. Between 380 and 550 K, ozone loss reached 71 DU in March and 85 DU in April (blue columns). Balloon data in March show smaller ozone loss values in 380 and 550 K, because of missing data between 420 and 470 K in the vortex core. Between 350–380 K, as can be seen in the difference between the cyan and blue columns in Fig. 9, balloon data and HALOE observations indicate

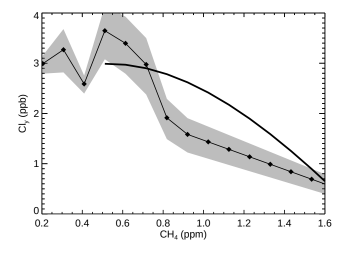


Fig. 10. The sum of ClONO₂ (measured by MIPAS-B on 14/15 March 1992) and HCl (measured by LPMA on 14 March 1992) against methane mixing ratios, solid diamonds. The grey scale indicates the combined uncertainty range of ClONO₂ and HCl. Also shown is the Cl_y^*/CH_4 relation from Grooß et al. (2002) (solid black line) where Cl_y^* has been reduced by 12%, to account for the lower chlorine loading in 1992 than in 2000 (WMO, 2007).

significant chemical ozone loss (13–17 DU) between December and March.

7 Chlorine activation deduced from MIPAS-B and LPMA measurements

Chemical ozone loss during February and March in the lower polar stratosphere is only possible if chlorine is still activated. Information about the two major chlorine reservoir species, ClONO₂ and HCl, is available from the balloon-borne measurements MIPAS-B and LPMA on 14 March 1992. By that time, chlorine deactivation in the Arctic vortex had proceeded and the majority of the active chlorine, ClO_x ($ClO+2 \times Cl_2O_2$), had been converted to $ClONO_2$ (e.g., Toohey et al., 1993; Müller et al., 1994).

Here, we estimate the extent of chlorine activation still present in mid-March 1992. The abundance of inorganic chlorine Cl_y can be estimated by the observed sum of HCl and ClONO₂ from the balloon-borne measurements. Grooß et al. (2002) reported a Cl_y^*/CH_4 relation for deactivated conditions in 2000. Cl_y^* has been reduced by 12% to account for the lower chlorine loading in 1992 than in 2000 (WMO, 2007). The increase in methane (2%) from 1992 to 2000 (Simpson et al., 2002) has only a minor effect but is also taken into account. The maximum Cl_y^* of about 3 ppb at the lowest CH_4 mixing ratios is in very good agreement with Cl_y deduced independently from whole air sampler measurements for winter 1991/1992 (Schmidt et al., 1994). The comparison of the Cl_v^*/CH_4 relationship with the observed re-

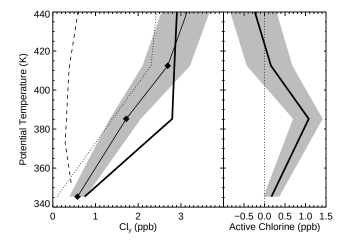


Fig. 11. Similar as in Fig. 10, but here the sum of $CIONO_2$ and HCl mixing ratios (solid diamonds) is shown against potential temperature. The grey scale indicates the combined uncertainty range of $CIONO_2$ and HCl. Solid black line shows the adjusted Cl_y^{\star} reference from Grooß et al. (2002), using the MIPAS-B methane measurements. Also shown are $CIONO_2$ mixing ratios (from MIPAS-B, dotted line) and HCl (from LPMA, dashed line).

lation between (HCl + ClONO₂)/CH₄ is shown in Fig. 10. For methane mixing ratios greater than ≈ 0.8 ppm, the sum of ClONO₂ and HCl is much lower than the estimated Cl_y. This indicates that chlorine is not yet completely deactivated in mid-March 1992.

In Fig. 11, left panel, the sum of the measured ClONO₂ and HCl mixing ratios is compared against the estimated inorganic chlorine Cl^{*}_y on a potential temperature scale. Between about 345 and 400 K, Cl^{*}_y exceeds the sum of ClONO₂ and HCl, showing that at those altitudes chlorine was still activated. The vertical profiles of HCl and ClONO₂ on 14 March 1992 (dashed and dotted lines, respectively in Fig. 11) confirms that the deactivation that has occurred up to this date is dominated by the formation of ClONO₂ (e.g., Toohey et al., 1993; Müller et al., 1994). HALOE HCl/CH₄ relations support this finding (Tilmes et al., 2004). Figure 11, right panel, shows the vertical profile of active chlorine on 14 March found as the difference between the estimated Cl^{*}_y and the observed sum of ClONO₂ and HCl on 14 March 1992. Maximum active chlorine of ≈ 1 ppb occurred around 380 K.

Box model calculations (e.g., Canty et al., 2005) indicate that, for 1 ppb of ClO_x present at 65° N, 380 K on 20 March, calculated chemical loss rates are about 40 ppb per day, which equals a loss of 4% per day when 1 ppm of ozone is present [Tim Canty, private communication, 2007]. The presence of activated ClO_x in middle to late March in combination with the long period of sunlight leads to rapid ozone loss. This likely explains why ozone losses observed by HALOE in early April exceed those in March, particularly below 400 K (region of elevated ClO_x).

8 Discussion

The comparison of different observations in the winter 1991– 1992 results in a consistent picture of chemical ozone loss derived using tracer-tracer correlations. HALOE ozone observation were shown to agree well with observed ozone sonde observations in the Arctic polar vortex between January and March. The deviation between a monthly averaged ozone sonde profile and HALOE ozone is on average 0.3 ppm at altitudes above 400 K potential temperature. Below 400 K, an altitude which was very likely influenced by high aerosol loading after the eruption of Mt. Pinatubo, these data do not deviate by more than 0.2 ppm.

The detailed discussion about the location of profiles in Sect. 5 was performed to distinguish between profiles observed within the vortex core and the vortex boundary region. Profiles located within the polar vortex core show largest chemical ozone loss values, as observed by ER-2 during January and February and by HALOE between February and April. Kiruna balloon observations confirm larger local ozone loss values for observations taken within the polar vortex core.

During March, Kiruna balloon measurements and HALOE observations indicate chemical ozone loss of 13-17 DU below 380 K. By mid-March 1992, chlorine deactivation had substantially proceeded, dominated by the formation of ClONO₂. Nevertheless, balloon-borne measurements (Fig. 11) demonstrate that chlorine was still activated in the altitude range between 345 and 400 K potential temperature, with a maximum active chlorine of ≈ 1 ppb occurring at \approx 380 K. These findings are in agreement with very low HCl mixing ratios observed by HALOE in the polar vortex between January and February below 450 K (Tilmes et al., 2004). Assuming active chlorine of ≈ 1 ppb, box model simulations show rapid ozone destruction of about 4% per day in March, which explains the additionally derived ozone loss between March and April 1992 based on HALOE observation.

The comparison of averaged ozone loss profiles of Arctic winters between 1991–1992 and 2002–2003, derived using HALOE observations, shows significantly larger ozone loss values at altitudes below 400 K for the years shortly after the Mt. Pinatubo eruption. Between 400 and 500 K, chemical ozone loss values of 65 ± 16 DU are in agreement with values derived by Rex et al. (2004). Therefore, the influence of enhanced sulfate aerosols in altitudes between 400–550 K described by Rex et al. (2004) is in agreement with the results found here.

The comparison between ER-2, balloon and HALOE observations show a good agreement around and below 400 K. This is the altitude where the uncertainty of HALOE was presumed to be largest due to aerosol interference. Further, column ozone loss between 400–500 K is in agreement for ER-2 and HALOE observations. Therefore, we conclude the influence of large aerosol loadings on HALOE measurements is small and ozone loss values calculated between February and April are reliable with and uncertainty of 15–16 DU of the total loss. Chemical ozone loss in winter 1991–1992 is significantly larger than in comparable moderately warm winters, due presumably to enhanced chlorine activation caused by the highly enhanced volcanic aerosol loading.

9 Conclusions

In the moderately warm winter 1991-1992 significant chemical ozone loss was found as a result of the enhanced burden of sulfate aerosols in the lower stratosphere after the Mt. Pinatubo eruption in June 1991. The comparison of different observations corroborates results derived earlier from HALOE satellite observations, especially for altitudes below 400 K. Between 400 and 500 K, 50–55 ± 16 DU of chemical ozone loss was estimated between December and February based on ER-2 and HALOE observations and up to 65±16 DU between December and April based on HALOE observations. Between 380 and 550 K potential temperature, 71 ± 26 DU of chemical ozone loss in March and 86 ± 27 DU during April is deduced from HALOE observations. For altitudes below 380 K, ozone loss of 13 DU in March was derived based on Kiruna balloon observations and 17 DU in April based on HALOE observations, in agreement with activated chlorine of ≈ 1 ppb occurring at ≈ 380 K in mid-March 1992.

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