A closer look at Arctic ozone loss and polar stratospheric clouds

N. R. P. Harris¹, R. Lehmann², M. Rex², and P. von der Gathen²

¹European Ozone Research Coordinating Unit, Department of Chemistry, University of Cambridge, Lensfield Rd, Cambridge, CB2 1HE, UK
²Alfred Wegener Institute, Potsdam, Germany

Received: 24 December 2009 – Accepted: 8 February 2010 – Published: 10 March 2010
Correspondence to: N. R. P. Harris (neil.harris@ozone-sec.ch.cam.ac.uk)
Published by Copernicus Publications on behalf of the European Geosciences Union.
Abstract

The empirical relationship found between column-integrated Arctic ozone loss and the volume of polar stratospheric clouds inferred from meteorological analyses is updated and examined in more detail. The relationship is found to hold at different altitudes as well as in the column. Analysis of the photochemistry leading to the ozone loss shows that the early winter activation is limited by the photolysis of nitric acid. This step produces nitrogen dioxide which is converted to chlorine nitrate which in turn reacts with hydrogen chloride on any polar stratospheric clouds to form active chlorine. The rate-limiting step is the photolysis of nitric acid: this occurs at the same rate every year and so the interannual variation in the ozone loss is caused by the extent and persistence of the polar stratospheric clouds. In early spring the ozone loss rate increases as the solar insolation increases the photolysis of the chlorine monoxide dimer. However the length of the ozone loss period is determined by the photolysis of nitric acid which also occurs in the near ultraviolet. As a result of these compensating effects, the amount of the ozone loss is principally limited by the extent of original activation rather than its timing. In addition a number of factors, including the vertical changes in pressure and total inorganic chlorine as well as denitrification and renitrification, offset each other. As a result the extent of original activation is the most important factor influencing ozone loss. These results indicate that relatively simple parameterisations of Arctic ozone loss could be developed for use in coupled chemistry climate models.

1 Introduction

Polar ozone loss has been the subject of intense scientific and public interest since the discovery of the Antarctic ozone hole (Farman et al., 1985) and its relatively quick attribution to observations of chlorine compounds (de Zafra et al., 1987; Solomon et al., 1987; Anderson et al., 1989). Stratospheric ozone loss takes place in the polar vortex which forms over each pole in their respective winters. Marked differences in
stratospheric dynamics in the two hemispheres naturally lead to large interannual variations in vortex stability and in ozone in the dynamically active Arctic winter stratosphere and to small interannual variations in the dynamically less active Antarctic. These differences lead to higher natural average amounts of total ozone over the Arctic (∼450 Dobson Units – DU) than over the Antarctic (∼300 DU – Dobson, 1968; Newman and Rex, 2007). These differences in dynamics also lead to a much greater variability of polar ozone loss over the Arctic (where, for example, there were losses of <10% in 1998/99 (Schulz et al., 2001) and >65% in 1999/2000 (Rex et al., 2002) at around 18 km) than over the Antarctic where nearly complete ozone loss has taken place in nearly all winters since the early 1990s at altitudes between about 15 and 20 km. Even in the anomalous year of 2002 when the Antarctic vortex was disturbed and broke up in mid-September the minimum daily ozone value south of 40° S was ∼140 DU, 30% below the late July values (Bodeker et al., 2005). In percentage terms this loss is comparable with the largest losses observed in the Arctic.

This large variability in the Arctic dynamics coupled with a number of microphysical and photochemical thresholds has made it hard to reproduce ozone loss in past winters for which the meteorology is known and analyses are available, although 3-D chemical transport models (CTMs – e.g. Chipperfield et al., 2005) and data assimilation approaches (Jackson and Orsolini, 2008) are now doing a reasonable job. However the additional complication of calculating the stratospheric dynamics (coupled with the sensitivity of the ozone loss processes to the dynamics and transport) means that it is even harder to predict Arctic ozone losses for the coming decades using coupled chemistry climate models (CCMs) (Austin et al., 2003; Eyring et al., 2007). The sensitivity of ozone loss to changes in climate is thus hard to assess.

The basic mechanisms leading to ozone loss are the same over the two poles and are generally well understood (e.g. Newman and Rex, 2007). As the polar vortex is established, the temperatures drop below a critical point, and polar stratospheric clouds (PSCs) can form. HCl and ClONO$_2$ react on the surface of PSCs, with these unreactive chlorine species being converted to active forms (ClO$_x$=Cl+ClO+Cl$_2$O$_2$) which can
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Rapidly destroy ozone. In the absence of further exposure to PSCs, the ClO (chlorine monoxide) formed continues to destroy ozone while it is gradually converted back to unreactive forms. The observed evolution of the main chemical species in the Arctic can be seen in Fig. 1 for the 2004/05 winter. The timing of the growth in ClO and ClO\(_x\) coincides with the onset of PSCs. Continued PSC occurrence leads to continued formation of ClO and ClO\(_x\). Reversion to ClONO\(_2\) and HCl occurs after the vortex has warmed and the PSCs have evaporated. Occasionally in Arctic winters, particularly large ozone losses can occur when air masses become depleted in HNO\(_3\) as PSCs fall to lower altitudes during prolonged cold periods, a process that delays the deactivation of the active chlorine species. The persistence of PSCs depends very strongly on the dynamical situation in that winter and so is the main reason that ozone loss can vary so much from year to year in the Arctic.

Given the mechanistic complexity and large variability, it came as a surprise to find that a compact, linear relation exists between ozone loss and the calculated volume of PSCs (\(V_{PSC}\)) when each is integrated over the period of vortex existence (Fig. 2 – see also Rex et al., 2004, 2006; Tilmes et al., 2004). The relation holds over a wide range of ozone losses and PSC volumes implying that the effects of many influences on ozone loss (e.g. denitrification, solar exposure, initial chemical fields, descent rates, in-mixing, vortex inhomogeneities, and vertical extent) must be offsetting to some extent. The existence of this relation (calculated from temperature fields and vortex average descent rates derived from meteorological analyses and ozonesonde measurements) has been confirmed using HALOE satellite measurements and an independent approach to calculating the ozone loss (Tilmes et al., 2004). Both Rex et al. (2004) and Tilmes et al. (2004) found larger than expected chemical ozone losses in the 1991/92 and 1992/93 winters which they attributed to the increase in stratospheric aerosol following the eruption of Mt Pinatubo.

Otherwise there has been little mechanistic investigation, and the explanation for the existence of the relation is not detailed, being simply that large ozone losses occur in cold winters when PSCs are widespread and long-lasting, while small ozone
losses occur in warm winters when PSCs are less common. This explanation does not really address either the compactness or the near-linearity of the relation. The lack of explanation is one of the main reasons why it is not being used to evaluate the coupled climate models (CCMs) in the CCM Validation Activity for SPARC (CCMVal) (see Eyring et al., 2005, updated at http://www.pa.op.dlr.de/CCMVal/CCMVal_EvaluationTable.html), despite its empirical robustness and its implication that there is sensitivity of 15 DU column ozone loss for each 1°C cooling (Rex et al., 2006).

The relation has however been used to evaluate (a) the improvements made to the SLIMCAT 3-D Chemical Transport Model (CTM) which used to show a compact, linear relation of the wrong slope (Rex et al., 2004), and now can reproduce the slope as well (Chipperfield et al., 2005; Fig. 4.13 in Newman and Rex, 2007), and (b) the ozone loss calculated in the Whole Atmosphere Community Climate Model (WACCM) (Tilmes et al., 2007).

The aim of the current study is to explain the compactness and linearity of this relationship and to identify its sensitivity to the range of possible influences discussed above. Our earlier work (Harris et al., 2009) found that the rate-determining step for chlorine activation in early winter is the photolysis of HNO₃, while the subsequent ozone loss depended on the competition between the photolysis of Cl₂O₂ (leading to ozone loss) and HNO₃ (leading to deactivation). Since both processes go faster as the solar zenith angle decreases, the integrated ozone loss depends primarily on the extent of the initial chlorine activation and not on the speed of the ozone loss. Here we investigate the empirical Vₚₛₖ/ozone loss relationship in more detail and extend the photochemical analysis to investigate the effect of multiple activations, denitrification and the vertical distribution of available chlorine (Clᵥ).

In the next section the methodology and data sources are described. In Sect. 3, the altitude variation of the relation is investigated and the sensitivity to the assumptions about PSC composition are discussed. The critical photochemical steps involved in the activation and deactivation steps are then described and illustrated in Sect. 4 by photochemical calculations on a single surface. Three dimensional aspects are
discussed in Sect. 5. Finally the relevance and context of these results is discussed and summarised in Sect. 6.

2 Methodology

This study uses the values for ozone loss and \( V_{\text{PSC}} \) calculated in Rex et al. (2006) updated with values for 2005/06, 2006/07 and 2007/08 using the same methodology. Briefly, vortex averaged profiles of ozone loss have been determined as the differences between the end of March and early January, with an adjustment for the vortex average descent being made using diabatic heating rates from the SLIMCAT CTM. These have been converted into concentration versus altitude profiles, using the vortex averaged temperature and pressure profiles from late March. The total column loss was calculated as the vertical integral of the loss profiles between 14 and 24 km. The lower limit of this range (~380 K) is close to the bottom of the well-isolated part of the polar vortex. For most winters ozone loss at this level is small. Also, the effect of any chemical loss in the vertical region below 14 km on the total ozone column in the Arctic would be limited because of rapid exchange with mid latitude air. Formation of PSCs above the vertical range considered here is unlikely and consequently significant chemical loss of ozone is not expected above 24 km. The estimated uncertainty in the integrated ozone loss is ~10–15 DU (Rex et al., 2002), mainly due to uncertainties in the calculated cooling rates and the potential impact of mixing across the vortex edge.

PSCs are assumed to be nitric acid trihydrate (NAT), and \( V_{\text{PSC}} \) is calculated from the laboratory observations of Hanson and Mauersberger (1988), a water vapour mixing ratio of \( \text{H}_2\text{O}=5 \text{ ppm} \), and an observed profile of \( \text{HNO}_3 \) (Rex et al., 2002 for more details of the methodology). In this paper temperatures from ECMWF ERA-Interim re-analyses are used in order to have a consistent vertical resolution and data quality across the whole period. The validity of this approach is discussed further in Sect. 3.

In order to investigate the underlying processes, the Alfred Wegener Institute (AWI) photochemical box model was used. This model contains 48 chemical species and
174 reactions. The formation of solid and liquid PSC particles is simulated according to Murray (1967), Hanson and Mauersberger (1988) and Carslaw et al. (1995). For the photolysis of Cl₂O₂, the absorption cross sections from Burkholder et al. (1990) are used, which are close to the values found in the most recent laboratory studies (Chen et al., 2009; Papanastasiou, et al., 2009; von Hobe et al., 2009). All remaining information on rate constants is taken from the NASA-JPL 2006 Evaluation (Sander et al., 2006). The overhead ozone needed for the calculation of photolysis frequencies is the average of all ozonesonde measurements made between January and March from 1992–2007 at Ny-Alesund (79° N). The numerical integration is performed using the kinetic preprocessor KPP (Damian et al., 2000).

Idealised trajectories are used which correspond approximately to the range of isentropic surfaces on which the observed ozone loss is derived. The trajectories are designed to mimic air toward the edge of an Arctic vortex which is displaced off the North pole by 10°: it is centred at 80° N and has a sinusoidal 6 day cycle of 20°. The main set of runs is performed at 50 hPa (~475 K). Runs investigating the vertical dimension were performed at 550 K (~30 hPa), 500 K (~40 hPa) and 450 K (~55 hPa). The analysis presented here is split into three main parts: the activation period (Sect. 4.1); the ozone loss period (Sect. 4.2); and aspects of the vertical dimension process (Sect. 5).

3 Altitude variation of relation and PSC composition

The original plots of integrated column ozone loss against integrated PSC volume (Rex et al., 2004, 2006) were calculated using ECMWF operational analyses. The PSC volumes shown in Fig. 2 are calculated using ERA-Interim re-analyses. These give greater consistency between years. The general features of the plot are unchanged. The most notable is that the largest value for V_{PSC} is now calculated to be in 1995/96 rather than 2004/05. However the changes are largely within the estimated uncertainties.
Figure 3 shows the ozone loss on individual isentropic surfaces (400, 450, 500 and 550 K) plotted against the area of PSCs \( A_{\text{PSC}} \), both integrated over the course of the winter. (As for the column ozone loss and \( V_{\text{PSC}} \), the isentropic surfaces “descend” over the course of the winter according to the diabatic heating rates calculated in the SLIMCAT model. The isentropic surface referred to is the one at the end of the winter.)

The ozone losses in Fig. 3 are shown in molec cm\(^{-3}\) (not ppm), so that their relative contributions to the column can be more easily seen. The plots for these levels are generally similar to the column plot of ozone loss vs \( V_{\text{PSC}} \), being reasonably linear and compact. The slopes (in molec cm\(^{-3}\) km\(^{-2}\) or \( 10^{15}\) molec km\(^{-1}\)) are greatest at 400 K and 450 K, with smaller values at 500 K and 550 K, showing the importance of the lower levels to the integrated column loss. The 550 K plot has the largest relative scatter. This is partly due to the smaller ozone losses found there and partly due to the poorer data quality (a significant fraction of ozonesondes do not reach that high especially early in winter as the descending layer starts at 600 or 700 K).

The values for \( V_{\text{PSC}} \) and \( A_{\text{PSC}} \) in Figs. 2 and 3 were calculated on the assumption that PSCs are composed of nitric acid trihydrate. The true nature of PSCs, especially near the PSC formation threshold, has been studied extensively with considerable evidence that liquid sulphate aerosol and supercooled ternary solutions (STS) play an important role in chlorine activation in the Arctic (e.g. Drdla et al., 2002). STS is present at different temperatures to NAT so that the NAT threshold temperatures used here might be misleading. The composition of large-scale fields of PSCs has been recently investigated using the MIPAS infrared sounder and the CALIPSO/CALIOP lidar measurements (Höpfner et al., 2009; Pitts et al., 2007, 2009). They find that the PSC fields can be described by four main types (ice, NAT-STS mixtures, STS-ICE mixtures, and just STS). The overall evolution of the observed PSCs is similar to that deduced from meteorological analyses, but it is hard to make quantitative comparisons. Pitts et al. (2009) use observed HNO\(_3\) and H\(_2\)O fields to calculate possible PSC existence while we assume a temporally unvarying concentration profile. Their PSC occurrence in the Antarctic is 15% higher than their earlier value which was 50% lower than that
inferred from meteorological temperatures and the NAT thermodynamics. A closer examination of these issues in the Arctic is necessary to justify a change in our definition of PSC occurrence which relies solely on meteorological reanalyses and assumptions about the mixing ratios of H$_2$O and HNO$_3$.

The effect of this assumption on the ozone loss/$V_{\text{PSC}}$ relation was investigated by Tilmes et al. (2008) who found significant differences only in winters with high sulphate aerosol loading, i.e. 1991/92 and 1992/93 following the Mt Pinatubo eruption. For comparison, we have used the temperature dependent cold aerosol formation to calculate $A_{\text{PSC}}$ (as in Fig. 3, but not shown). No real difference is found except at 400 K, where a slightly more compact relation is found. This is presumably related to the higher amount of background aerosol at these altitudes. This is consistent with the CALIOP analysis in the Antarctic which shows the biggest discrepancy in a 1–2 km band at altitudes below 15 km at the beginning of winter (Fig. 14c in Pitts et al., 2009). A similar feature can be seen in the comparison of calculated PSC occurrence using $T_{\text{NAT}}$ with ground-based lidar observations in 1999/2000 Arctic winter (Fig. 2 in Rex et al., 2002).

Overall, we found no reason to change from using the NAT definition in our calculation of $A_{\text{PSC}}$ and $V_{\text{PSC}}$.

4 Idealised case: a single layer

We now examine the relationship of the ozone loss to the photochemical steps involved in the activation and deactivation of $\text{ClO}_x$ ($=\text{ClO}+2\times\text{Cl}_2\text{O}_2+2\times\text{Cl}_2$) at 50 hPa using the AWI box model. In Sect. 4.1 we discuss the effects of the frequency of occurrence and temperature of the PSCs; while in Sect. 4.2 we investigate the competition between the ozone loss and $\text{ClO}_x$ deactivation chemistry.
4.1 Activation

Figure 4 shows the mixing ratios of the chlorine reservoirs, ClONO$_2$ and HCl and the activated forms ClO$_x$ from 1 December to 14 February assuming that PSCs are continuously present. The assumed temperature is 194 K which is representative of the vortex as a whole. Two phases can be seen: (i) a rapid initial rise in ClO$_x$ associated with equal decreases in both HCl and ClONO$_2$; and (ii) a slower continued increase in ClO$_x$ accompanied by a continued decrease in HCl. By the end of the period, nearly all the available chlorine is in the form of ClO$_x$. The two steps are now discussed in turn.

ClO$_x$ rises initially as a result of the fast heterogeneous reaction between ClONO$_2$ and HCl, with the extent of the initial activation being determined by the amount of ClONO$_2$ initially present (Müller et al., 1994; Santee et al., 2008). The heterogeneous reaction of ClONO$_2$ + HCl proceeds quicker at lower temperatures, while the photolysis of HNO$_3$ has only a weak temperature dependence. The main effect of lower temperatures is thus to accelerate the initial activation (not shown).

If the initial mixing ratio of HCl is greater than that of ClONO$_2$, further activation depends on the formation of more ClONO$_2$ (e.g. Müller et al., 1994) which depends in turn on the photolysis of HNO$_3$ producing NO$_2$ which reacts with ClO to form ClONO$_2$. In the presence of PSCs, the ClONO$_2$ reacts rapidly with HCl to produce more ClO$_x$. In the Arctic winter the slow step in the overall activation mechanism is the photolysis of HNO$_3$, with a timescale of days to weeks. The slow, continued rise in ClO$_x$ in Fig. 4 thus depends principally on the photolysis frequency, J(HNO$_3$), and this factor is responsible for the acceleration in the activation after mid-winter. Complete activation would take several weeks in the Arctic in the continued presence of PSCs. The full mechanism is thus

\[
\text{ClONO}_2 + \text{HCl} \rightarrow \text{Cl}_2 + \text{HNO}_3 \text{ heterogeneous, so fast if PSCs are present} \quad (R1)
\]

\[
\text{HNO}_3 + h\nu \rightarrow \text{NO}_2 + \text{OH} \quad \text{slow} \quad (R2)
\]

\[
\text{NO}_2 + \text{ClO} + \text{M} \rightarrow \text{ClONO}_2 + \text{M} \quad \text{relatively fast in the presence of sunlight.} \quad (R3)
\]

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In the real Arctic vortex, the initial activation is patchy depending on the altitudes and regions where the PSCs first form as well as on their extent. Vortex average ClO$_x$ initially rises slower in reality than in the idealised case, as indicated in Fig. 1. To illustrate this, Fig. 5 shows the same model calculation as in Fig. 4, except that the PSCs occur episodically (1 day in 6). The slow rise is still apparent, but the effect of the fast initial activation is smeared out over time. The stabilisation of HCl between PSC exposures is a result of the lack of PSCs. Similarly each of the small rises in ClO$_x$ results from exposure to PSCs, and the decreases occur during exposure to sunlight (orange). However none of these individual fast processes really influence the overall shape which is still determined by J(HNO$_3$). Further, the initial rapid activation could be responsible for the non-zero intercepts in Figs. 2 and 3, as any small-scale or sub-grid process leading to PSC formation (e.g. mountain waves) could rapidly activate a small but significant amount of air while barely contributing to $V_{PSC}$. However this mechanism might not be the correct interpretation since a non-zero intercept is also seen in the SLIMCAT model calculations.

Overall, our analysis of the chlorine activation shows that activation in Reaction (R1) is fast and that the rate limiting factor for any continued activation is the photolysis of HNO$_3$ to form NO$_2$ in Reaction (R2) and then ClONO$_2$ in Reaction (R3). To first order the photolysis of HNO$_3$ does not vary much from year to year, and so interannual variations in activation depend on the presence of PSCs for activation to occur through Reaction (R1).

### 4.2 Deactivation and ozone loss

There have not been many studies explicitly looking at how the amount of ozone loss depends on the timing and extent of the initial activation. The main ozone loss period starts early in the year as the insolation increases. The instantaneous ozone loss rates have been observed to peak in late January with a gradual reduction thereafter (von der Gathen et al., 1995). The daily ozone loss rate maximises a little later as it is also determined by the length of the day which is increasing in this period. As discussed
in Harris et al. (2009), both the ozone loss and the chlorine deactivation are driven by sunlight and so the ozone loss and chlorine deactivation processes accelerate as the insolation increases. The rate limiting step for ozone loss in the Arctic (for a given amount of total inorganic chlorine (Cl\textsubscript{y})) is:

\[
\text{Cl}_2\text{O}_2 + h\nu \rightarrow \text{ClOO} + \text{Cl}
\]  
(R4)

while in the absence of PSCs, Reactions (R2) and (R3) become the main deactivation mechanism.

Both HNO\textsubscript{3} and Cl\textsubscript{2}O\textsubscript{2} are photolysed in the UV, and the atmospheric photolysis rates of both (R2) and (R4) in the Arctic polar lower stratosphere depend mainly on wavelengths longer than 300 nm as shorter wavelengths are absorbed by stratospheric ozone.

Figure 6 shows these photolysis rates (integrated from 250–400 nm) as a function of solar zenith angle. They both increase markedly as the solar zenith angle decreases as winter turns into spring. To first order both the chlorine deactivation rate and the ozone loss rate are proportional to the solar intensity (as well as to each other), with both processes having a timescale of several days at 70–75° N in January and February. As a result the integrated ozone loss is, to first order, independent of the solar intensity and the timing of the activation, but is dependent on the original level of activation.

To illustrate this, Fig. 7 shows the ozone mixing ratios for three cases calculated by the AWI model. All cases have an initial activation of 3 ppb of Cl\textsubscript{y} with the same hypothetical trajectory used before (50 hPa centred at 80° N with 20° deviation with a 6 day cycle). The only difference is in the start dates which are set three weeks apart: 11 January (red), 1 February (blue) and 22 February (green). Despite the large differences in the start dates, the ozone losses are within ±10%.

While this analysis highlights the importance of the two photolysis Reactions (R2) and (R4) in determining ozone loss in the Arctic, other processes do play a limited role. For example, in Fig. 7 a smaller ozone loss is calculated when the 22 February start date is used. This is consistent with the greater sensitivity of the Cl\textsubscript{2}O\textsubscript{2} photolysis
rate to decreasing SZA at high SZA (∼90°) and the greater sensitivity of the HNO$_3$ photolysis rate at lower SZA (∼70–80°) shown in Fig. 6. Also, the conversion of HNO$_3$ to NO$_x$ via the alternative channel

$$\text{OH} + \text{HNO}_3 \rightarrow \text{NO}_3 + \text{H}_2\text{O} \quad \text{(R5)}$$

depends on $J(\text{HNO}_3)$ since HNO$_3$ photolysis is one of the main ways in which OH is formed.

In addition, reactions which play an important role in Antarctic ozone loss are less significant in most Arctic winters. The highly temperature-dependent, heterogeneous reactivation of ClONO$_2$ through

$$\text{ClONO}_2 + \text{H}_2\text{O} \rightarrow \text{HOCl} + \text{HNO}_3 \quad \text{(R6)}$$

is unimportant in most Arctic winters.

We have so far shown the direct relationship between $V_{\text{PSC}}$ and ClO$_x$. A further important point about the degree of activation is that the ozone loss is close to linear with ClO$_x$. The blue line in Fig. 8 shows the dependence of the accumulated ozone loss at 500 K (close to 50 hPa, the level of the calculations shown in Fig. 7) on the initial amount of ClO$_x$. The accumulated ozone losses for 1, 2 and 3 ppb ClO$_x$ at 500 K are 0.27, 0.59 and 1.00, respectively, so that there is a small, positive non-linearity in this relationship.

5 Extending to the 3-D view

The analysis discussed in Sect. 4 is concerned only with a single layer in the Arctic vortex. In this section we look at some of the greater complexities of the real atmosphere by investigating the effects of a number of the three-dimensional processes which affect ozone loss: (1) pressure (altitude); (2) the increasing Cl$_y$ with altitude; (3) the vertical redistribution of NO$_y$ in de/renitrification; (4) the interannual variation of transport.

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5.1 Pressure

For fixed ClO\textsubscript{x}, the potential for ozone loss depends slightly on pressure, with larger ozone losses calculated at lower potential temperatures as shown in Fig. 8. For an initial ClO\textsubscript{x} of 3 ppb, the accumulated ozone losses at 450, 500 and 550 K are 1.2, 1.0 and 0.8 ppm. The reasons for the pressure dependence are as follows.

(a) At higher altitudes the ClO and BrO concentrations are smaller for the same mixing ratios, as is the overall pressure, thus reducing the rates of the reactions

\[
\text{ClO} + \text{ClO} + \text{M} \rightarrow \text{Cl}_2\text{O}_2 + \text{M} \quad (R7)
\]

\[
\text{ClO} + \text{BrO} \rightarrow \text{Br} + \text{Cl} + \text{O}_2. \quad (R8)
\]

(b) At higher altitudes the chlorine deactivation rate is higher, because NO\textsubscript{2} production through both HNO\textsubscript{3} photolysis and Reaction (R5) is faster. An increase of the HNO\textsubscript{3} mixing ratio with altitude, not considered in the present model runs, would amplify this effect.

Two counteracting effects at higher altitudes are the faster rate of the reaction

\[
\text{ClO} + \text{O} \rightarrow \text{Cl} + \text{O}_2 \quad (R9)
\]

because the O concentration increases with altitude; and the faster photolysis of Cl\textsubscript{2}O\textsubscript{2} (Reaction R4). However these are small compared to points (a) and (b).

5.2 Available chlorine

The amount of available chlorine increases rapidly with altitude in the Arctic vortex (Schmidt et al., 1994). As a result, more chlorine is activated for a given PSC exposure at higher altitudes. The black crosses in Fig. 8 indicate the values of Cl\textsubscript{y} measured toward the end of the 1991/92 winter (Schmidt et al., 1994). Taking these as representative upper limits for the amount of ClO\textsubscript{x} initially available shows that as altitude
increases the additional Cl\textsubscript{y} offsets the decreasing efficiency of the ozone loss processes. The combined effect is to limit the importance of any interannual variations of PSC altitudes on the integrated ozone loss.

### 5.3 Vertical redistribution of NO\textsubscript{y}

The vertical redistribution of NO\textsubscript{y} through denitrification at higher altitudes and renitrification at lower altitudes has been widely cited as having a major impact on the accumulated ozone loss as the removal of NO\textsubscript{y} limits the deactivation of ClO\textsubscript{x} through Reaction (R3). Significant denitrification in the Arctic only occurs in a few winters (1994/95, Sugita et al., 1998; 1995/96, Rex et al., 1997; 1999/2000, e.g. Popp et al., 2001 and 2004/05, Kleinböhl et al., 2005), which are the winters with the larger ozone losses in the top right of Fig. 2. Denitrification in the majority of winters is negligible, even non-existent. When large-scale denitrification does occur, the enhanced ozone loss in the denitrified layer tends to be offset by reduced ozone loss in a lower, renitrified layer (e.g. Rex et al., 1997). The magnitude of this effect is investigated here by assuming a denitrification at 550 K of 5 ppb (Fig. 9a) and a concurrent renitrification of 5 ppb at 450 K (Fig. 9b). For a given ClO\textsubscript{x}, there is a greater sensitivity of ozone loss to changes in NO\textsubscript{y} at 450 K. However at this level, the Cl\textsubscript{y} and ClO\textsubscript{x} are lower and so the overall effect of this hypothetical de/renitrification on the vertically integrated ozone loss is limited. Interestingly (but not conclusively) if one looks at particular years in Fig. 3, one can see indications of both denitrification and renitrification. For example in 1999/2000, a year of extensive ozone loss and well observed denitrification, Fig. 3b shows higher than “expected” ozone loss at 450 K and less than “expected” at 400 K, consistent with the altitudes of the observed denitrification/renitrification (Popp et al., 2001).

### 5.4 Interannual variations in transport

A further process which could affect the chemical recovery is any in-mixing of air with a different chemical composition from outside the vortex. Overall this does not seem to
be a significant influence on ozone loss in the Arctic vortex (Newman and Pyle, 2003) despite the identification of individual events (e.g. Pyle et al., 1994).

6 Discussion and summary

We have updated and extended the previous studies reporting the empirical relation between accumulated ozone loss and PSC volume in the Arctic vortex (Rex et al., 2004, 2006; Tilmes et al., 2004). Not only do the calculated results still hold with additional winters, but it is also shown to hold on individual potential temperature surfaces albeit with greater scatter. At 400 K, the relationship is found to be slightly more significant when cold aerosol activation is assumed in the place of NAT. This is consistent with the larger amounts of aerosol at 400 K than at higher altitudes. At higher potential temperatures, the correlation with PSC areas deduced from NAT existence is better. This finding implies a smaller sensitivity of ozone loss to stratospheric aerosol loading than that found by Tilmes et al. (2008) in their assessment of the deliberate maintenance of an enhanced stratospheric aerosol layer as a geo-engineering response to climate change.

The compactness and linearity of the empirical results are interpreted using a photochemical box model with standard photochemistry. The aim is to identify the important mechanisms, not to reproduce the losses from winter to winter. More complete reconstructions have been successfully reported elsewhere using a 3-D CTM and analysed meteorological fields (Chipperfield et al., 2005; Tilmes, 2007). We show that the principal timescale for the extensive activation of chlorine is not the fast reaction of ClONO$_2$ and HCl on NAT particles but is the re-supply of ClONO$_2$ with photolysis of HNO$_3$ as the rate determining step. As a result, extensive activation in the Arctic vortex takes place with a timescale of days to weeks, and the overall activation depends on the continued (though not necessarily the continuous) presence of PSCs. The use of $V_{\text{PSC}}$ is thus a sensible measure of the activation process.
The extent of ozone loss in any particular winter is found to depend most strongly on the degree of initial activation and not so much on its timing or vertical distribution. This somewhat surprising finding occurs as a result of a number of offsetting factors. For any given air mass, there is the almost cancelling competition between ozone loss and chlorine deactivation, both of which rely on the photolysis in the near UV and so accelerate as the Sun becomes higher in the sky in early spring. In the vertical, the effect of the decreasing number density with altitude is offset by the increasing levels of Cl$_y$, and in the few winters where denitrification occurs the increased ozone loss at the denitrified altitudes is offset by the decreased ozone loss at the lower altitudes where re-nitrification takes place. The baroclinicity of the vortex affects the degree of denitrification (Mann et al., 2003): this would affect our conclusions about denitrification if the cold region were sufficiently deep that the renitrification occurred below the region of ozone loss. However this is very rare in the Arctic and has probably not occurred even in the years of extensive denitrification.

These findings raise a number of interesting possibilities for the development of simple parameterisations of stratospheric Arctic (or even polar) ozone loss in coupled chemistry climate models (CCMs), a step beyond the use of simple models to investigate future sensitivities (Knudsen et al., 2004; Tilmes et al., 2008). For example, the relationship shown in Fig. 2, which is derived in a period of high and nearly constant Cl$_y$, could be scaled by the levels of Cl$_y$ calculated in the CCMs. As Cl$_y$ decreases so will ozone loss – but the effect of any changes in the occurrence of PSCs resulting from climate change would be allowed for. A slightly more sophisticated, but still very simple approach would be to develop a photochemical model based around the main factors described above, namely photolysis of nitric acid and Cl$_2$O$_2$, an activation involving ClONO$_2$ on PSCs and initial levels of Cl$_y$. Both of these schemes would depend on good representations of PSCs, which implies a greater need for improvements to the dynamical schemes used in CCM which determine temperature and vortex characteristics, rather than to the photochemical schemes. These comments only apply to polar ozone loss, because of the unique nature of the stratosphere over the poles.
Additionally this work underlines the value of more analysis of the existing models to see where simplifying assumptions are justified and where parameterisations might be made.

Acknowledgements. This work was supported by the EU Integrated Project SCOUT-O3 (505390-GOCE-CT-2004) and by the BMBF DYCHO project (FKZ07ATC08). NRPH thanks NERC for an Advanced Research Fellowship. We thank the many ozonesonde personnel who have launched the ozonesondes used in this analysis. Michelle Santee and Peter Bernath provided useful and helpful advice. Meteorological data were provided by ECMWF and FU-Berlin. The ACE mission is supported primarily by the Canadian Space Agency. EOS Aura MLS is supported by NASA.

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Fig. 1. Evolution of chlorine species inside the Arctic vortex at 460 K in the 2004/05 winter. The top panel shows the area of PSCs as a percentage of the vortex area (light blue) and the vortex average sunlit time per day in percent (pink). The middle panel (based on Santee et al., 2008) is the vortex average HCl from Aura MLS, the vortex average ClONO$_2$ from ACE FTS, and an estimate of the vortex average ClO$_x$ found by subtracting the sum of the HCl and ClONO$_2$ from 2.8 ppb, a representative value of Cl$_y$ for 460 K in that winter. ClONO$_2$ and ClO$_x$ have been smoothed with a 10 day running mean in order to compensate for the sampling biases from ACE FTS, which has the typical sampling issues of any solar occultation instrument. The bottom panel shows the ozone loss rates found from the Match campaign for that year (Rex et al., 2006).
Fig. 2. Integrated ozone loss as a function of $V_{PSC}$ for 1992/93 to 2008/09. No values are shown for the winters 2000/01, 2001/02, 2003/04, 2005/06 and 2008/09 due to major warmings and/or lack of ozonesonde data. Ozone losses are derived from the Arctic ozonesonde network using the vortex average approach. The error bars on $V_{PSC}$ are based on the sensitivity to a temperature perturbation of $\pm 1$ K in the ECMWF data. $V_{PSC}$ is derived from ECMWF ERA-Interim re-analyses using the temperature of formation of nitric acid trihydrate. (Updated from Rex et al., 2006).
Fig. 3. Integrated ozone loss as a function of $A_{PSC}$ for 1992/93 to 2008/09. Other details given in caption for Fig. 2.
Fig. 4. The evolution of ClO\textsubscript{x} (=ClO+2Cl\textsubscript{2}O\textsubscript{2}+2Cl\textsubscript{2}), HCl and ClONO\textsubscript{2} for an idealised trajectory at 194 K and 50 hPa with continued activation from 1 December to 14 February. The latitude of the trajectory oscillates sinusoidally around 80° N with a 20° amplitude. The orange shading indicates the sunlight exposure experienced by the air parcel.
Fig. 5. The evolution of ClO$_x$, HCl and ClONO$_2$ for an idealised trajectory at 50 hPa with a one day period of activation at 194 K every six days from 1 December to 14 February.
Fig. 6. The photolysis rates of HNO$_3$ and Cl$_2$O$_2$ at 50 hPa in the Arctic vortex as a function of solar zenith angle.
Fig. 7. Ozone loss calculated for an air parcel with an initial chlorine activation of 3 ppb ClO\textsubscript{x} on 11 January (red), 1 February (blue) and 22 February (green).
Fig. 8. Integrated ozone loss as a function of initial activation on the 450 K (green), 500 K (blue) and 550 K (red) potential temperature surfaces. The crosses represent the amount of available inorganic chlorine in March 1992 inferred by Schmidt et al. (1994).
Fig. 9. The integrated ozone loss as a function of initial activation on the (a) 550 K and (b) 450 K potential temperature surfaces. The green lines show the effect of HNO$_3$=5 ppb, the red lines for HNO$_3$=10 ppb and the blue lines for HNO$_3$=15 ppb. The arrows indicate the effect of a dentrification of 5 ppb at 550 K in (a) and a renitrification of 5 ppb at 450 K in (b).