Sensitivity of stratospheric inorganic chlorine to differences in transport

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Abstract. Correctly modeling stratospheric inorganic chlorine (Cl₂) is crucial for modeling the past and future evolution of stratospheric ozone. However, comparisons of the chemistry climate models used in the latest international assessment of stratospheric ozone depletion have shown large differences in the modeled Cl₂, with these differences explaining many of the differences in the simulated evolution of ozone over the next century. Here in, we examine the role of transport in determining the simulated Cl₂ using three simulations from the same off-line chemical transport model that have the same lower tropospheric boundary conditions and the same chemical solver, but differing resolution and/or meteorological fields. These simulations show that transport plays a key role in determining the Cl₂ distribution, and that Cl₂ depends on both the time scales and pathways of transport. The time air spends in the stratosphere (e.g., the mean age) is an important transport factor determining stratospheric Cl₂, but the relationship between mean age and Cl₂ is not simple. Lower stratospheric Cl₂ depends on the fraction of air that has been in the upper stratosphere, and transport differences between models having the same mean age can result in differences in the fraction of organic chlorine converted into Cl₂. Differences in transport pathways result in differences in vertical profiles of CFCs, and comparisons of observed and modeled CFC profiles provide a stringent test of transport pathways in models.

Therefore, correctly modeling stratospheric Cl₂ and Br₂ is crucial for modeling past and future stratospheric ozone levels. However, comparisons of the coupled chemistry climate models (CCMs) used in the latest international assessment of stratospheric ozone (Chapter 6 of WMO, 2007) have shown large differences in the modeled Cl₂, both in terms of magnitude and the date that Cl₂ returns to pre-1980 values (Eyring et al., 2006, 2007). Furthermore, these differences appear to explain many of the differences in simulated ozone between the models, e.g., models with a later return of Cl₂ to pre-1980 values generally have later return of ozone to pre-1980 values. Given its importance for modeling ozone and the large differences between models, it is important to understand the causes of the differences in the simulations of Cl₂.

In the above CCM simulations the same time series of organic chlorine species (e.g., chlorofluorocarbons, CFCs) are specified in the lower troposphere and the same or similar chemical reactions are included in the models. This indicates that the differences in Cl₂ are not due to differences in source gases or stratospheric chemistry but are most likely due to differences in transport.

Here we explore this issue further by examining Cl₂ from three simulations from the same off-line chemical transport model (CTM). The simulations use the same lower tropospheric concentration boundary conditions and the same chemical solver, but differing resolution and/or meteorological fields. As the boundary conditions and chemistry are exactly the same any differences in Cl₂ are due to differences in transport.

The model and simulations are described in the next section. In Sect. 3 we compare the Cl₂ and mean age Γ from the CTM simulations, and Sect. 4 we compare the simulated Cl₂ to estimates using the formulation used by Newman et al. (2006) to calculate EESC. These comparisons show that transport plays a key role in determining the distribution of Cl₂, and that Cl₂ depends on both the time scales and pathways of transport.

1 Introduction

The observed changes in ozone over the last two decades, as well as the expected future increases in the next two decades, are primarily the result of changes in the concentration of stratospheric inorganic chlorine (Cl₂) and bromine (Br₂).

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2 Model description

The simulations examined are performed using the NASA Global Modeling Initiative (GMI) CTM (Douglass et al. (2004) and references therein). The CTM includes a full description of stratospheric chemistry, and can be driven by meteorological data from either a free-running general circulation model (GCM) or from a data assimilation system (DAS). We consider both cases here: Simulations are analyzed that used either the Goddard Earth Observing System (GEOS) DAS (Schubert et al., 1993) or the GEOS-4 GCM (Bloom et al., 2005).

We consider three different simulations with one pair differing in meteorological fields driving the CTM, and one pair differing in horizontal resolution and model top. Specifically, we consider simulations:

1. GCM winds, 2° latitude \( \times \) 2.5° longitude, top at 0.015 hPa with 33 levels (“GCM-HIGH”).
2. GCM winds, 4° latitude \( \times \) 5° longitude, top at 0.4 hPa with 28 levels (“GCM-LOW”).
3. DAS winds, 4° latitude \( \times \) 5° longitude, top at 0.4 hPa with 28 levels (“DAS-LOW”).

The two “LOW” simulations are described in Considine et al. (2004), while the GCM-HIGH simulation is described in Douglass et al. (2006).

In each case a single year of meteorological fields is used in the CTM: For the DAS simulation the fields from 1 July 1999 to 30 June 2000 are used, while for GCM simulations the fields are from a year with a cold Arctic vortex (see Considine et al., 2004). The GCM simulation used climatological sea-surface temperatures, volcanic aerosols and solar variations were not included, and there was no QBO. All three simulations use the same chemical solver (see Douglass et al., 2004) and the concentration of halogens in the lower troposphere are specified as in table 4B-2 of WMO (2003). The source gas boundary conditions vary between years. In the GCM-HIGH simulation the boundary conditions start in 1974 and end in 2025. The “LOW” simulations were spun up for 5 years at 1995 source gas boundary conditions and then integrated through the year 2030. Because 1995 boundary conditions were used rather than 1990 to 1994 conditions in the spin up of the “LOW” simulations the stratospheric chlorine is over-estimated in the early part of these simulations. However, the difference is much smaller than the inter-model differences discussed below. For all simulation, fields have been archived every 5 years.

In addition to the above, simulations of an age spectrum tracer were performed for each CTM. In these simulations the mixing ratio of a conserved tracer in the lowest two model levels was set to 1 for the first month and then set to 0 for the rest of the 20 year runs. This tracer allows calculation of the mean age of air in the stratosphere as well as the full age spectrum (Waugh and Hall, 2002).
3 Simulations of Cl\textsubscript{y} and mean age

We first consider the Cl\textsubscript{y} distribution in the three simulations. As shown in Fig. 1 (a–c; top panels) the general features of the Cl\textsubscript{y} distributions are the same, e.g., there are values larger than 3 ppb in the upper stratosphere and lower values in the tropics than in mid- and high latitudes. However, there are quantitative differences, especially in the wintertime (southern) polar regions. For example, the South Pole Cl\textsubscript{y} at 20 km varies from less than 1.5 ppb to around 3 ppb. As the surface concentrations of organic chlorine species and the chemistry are the same in all three simulations, this confirms the CCM results (Eyring et al., 2006) that transport plays a key role in determining Cl\textsubscript{y}.

As Cl\textsubscript{y} is formed by conversion of organic chlorine into inorganic chlorine species within the stratosphere, the time spent in the stratosphere is expected to play an important role in determining Cl\textsubscript{y}. We therefore next consider the distributions of the mean age \( \Gamma \). As described above, \( \Gamma \) is calculated from simulations of the age spectra, and represents the annual-mean \( \Gamma \), see Waugh and Hall (2002). As shown in Fig. 1 (d–f; bottom panels) the mean ages differ among all three simulations. The largest differences are between DAS-LOW and the two GCM simulations, with much younger ages in DAS-LOW. The younger ages in the DAS simulation are related to the excessive mixing in simulations using assimilated winds (e.g., Schoeberl et al., 2003). There are also differences in age between GCM-LOW and GCM-HIGH, but these are small in the lower stratosphere (e.g., the South Pole mean age at 20 km is \( \approx 4.7 \) yrs in both GCM simulations).

Comparison of the Cl\textsubscript{y} and \( \Gamma \) distributions shows that Cl\textsubscript{y} is, in general, larger for older \( \Gamma \). However, this comparison also suggests that differences in \( \Gamma \) do not explain all the differences in Cl\textsubscript{y}. This is clearly seen in vertical profiles within the spring Antarctic vortex. At 20 km the mean ages from the two GCM based simulations are very similar but there is a large difference in Cl\textsubscript{y}, see Fig. 2a, b. Conversely, above this altitude there are much larger differences in mean age than in Cl\textsubscript{y} between these two simulations. Hence, there is not a simple relationship between differences in mean age and those in Cl\textsubscript{y}.

To explore the differences in Cl\textsubscript{y} in more detail we compare the time series of lower stratospheric Cl\textsubscript{y} at two locations where \( \Gamma \) is similar in the two GCM based simulations (Fig. 3). In mid-latitudes the two GCM simulations have very similar Cl\textsubscript{y} (Fig. 3a), but at the polar location there is a large difference (Fig. 3b). At both locations \( \Gamma \) and Cl\textsubscript{y} from the DAS simulation are smaller than from the GCM simulations. Note, as discussed above, the two “LOW” simulations overestimate Cl\textsubscript{y} in 1995 because of the initial conditions used, and this results in higher Cl\textsubscript{y} in 1995 than in 2000 in these simulations.

The differences in Cl\textsubscript{y} could be due to differences in the simulated total amount of chlorine Cl\textsubscript{tot}(the sum of organic and inorganic chlorine) in the stratosphere. However, Cl\textsubscript{tot} is very similar among all three simulations (dashed curves in Fig. 3). There is a slight time shift between the DAS and two GCM simulations, which is consistent with the approximately 2 year difference in the mean age. However, the magnitude of Cl\textsubscript{tot} is very similar and differences in Cl\textsubscript{tot} do not explain the differences in Cl\textsubscript{y}. As well as the time shift in Cl\textsubscript{tot} between the DAS and GCM simulations there are slight differences in the peak values, which are due to differences in the width of the age spectra. However these differences are insignificant compared to the differences in Cl\textsubscript{y}.

The above shows that the time spent in the stratosphere is important but this is not the sole transport factor determining stratospheric Cl\textsubscript{y}(and differences between the simulations). Hence, the paths taken to get to a particular location must also be important for determining Cl\textsubscript{y}.

4 Fractional release rates

To understand the differences in simulated Cl\textsubscript{y} we estimate the Cl\textsubscript{y} using the formulation of Newman et al. (2006, 2007). In this formulation Cl\textsubscript{y} is estimated by summing the contribution from each CFC, i.e.,

\[
\text{Cl}_{y}^{*}(t) = \sum_{i} n_{i} F_{i} \tilde{\rho}_{i}(t)
\]
where \( n_i \) is the number of chlorine atoms in source gas \( i \), \( \hat{\rho}_i \) is the mixing ratio in the stratosphere that would occur if the source gas \( i \) was perfectly conserved, and \( F_i \) is the fraction of species dissociated while it has been in the stratosphere. The mixing ratio \( \hat{\rho}_i \) depends on the tropospheric concentrations \( \rho_{i,trop} \) and the troposphere to stratosphere transport time scales:

\[
\hat{\rho}_i(t) = \int_{-\infty}^{t} \rho_{i,trop}(t') G(t - t') dt'.
\]

where \( G(t) \) is the age spectrum. The fractional release is given by

\[
F_i = (\hat{\rho}_i - \rho_i) / \hat{\rho}_i,
\]

where \( \rho_i \) is the actual concentration of the source gas at given stratospheric location.

We focus here on lower stratospheric \( Cl_y \) and, as in Newman et al. (2006), assume that the fractional releases are solely a function of the mean age, i.e. \( F_i = F_i(\Gamma) \). Although we assume a dependence only on the mean age it is important to know that the release rates also depend on pathways, and the functional relationships are not necessarily the same for all simulations. \( F_i(\Gamma) \) are determined separately for the three simulations using Eqs. (2) and (3).

We first compare \( Cl^*_y \) calculated using (1) with the directly simulated \( Cl^*_y \). In these calculations of \( Cl^*_y \) we use the age spectrum \( G(t) \) simulated within the CTMs (see Sect. 2), the \( \rho_{i,trop} \) used in the CTMs, and the above estimates of \( F_i \) calculated from model fields. As shown in Fig. 4 the \( Cl^*_y \) estimated from \( G(t) \) and \( F \) matches the time evolution of \( Cl_y \) and the differences between the simulations. There are some differences between \( Cl^*_y \) and \( Cl_y \) for GCM-LOW, but these differences are much smaller than model-model differences, and \( Cl^*_y \) is a good representation of \( Cl_y \) to within a few percent. Note that virtually the same \( Cl^*_y \) is obtained if the approximate age spectrum used by Newman et al. (2006) (an inverse Gaussian age spectrum with width equal to half the mean age) is used rather than the true model age spectrum. The dashed curves in Fig. 4 show \( Cl^*_y \) from these calculations, and a virtually indistinguishable from the solid curves.

The good agreement between \( Cl^*_y \) and the directly simulated \( Cl_y \) supports the use of the method of Newman et al. (2006) for calculating EESC. If \( \Gamma \) and \( F_i \) are known then the Newman et al. (2006) method can accurately predict EESC. Both \( \Gamma \) and \( F_i \) can change in the future, and this will add uncertainties to estimates of EESC (Newman et al., 2007).

Given this general agreement we can now explore the differences in the individual contributions to \( Cl^*_y \). As \( \rho_{i,trop} \) is the same in all simulations and the differences in \( Cl^*_y \) are

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**Fig. 3.** Time series of October-mean \( Cl_y \) (solid) and \( Cl_{tot} \) (dashed) at 56.6 hPa for (a) mid-latitude (46° S) and (b) polar (82° S) regions.

**Fig. 4.** Comparison of directly simulated \( Cl_y \) at 56.6 hPa and 82° S (symbols) and reconstruction using Newman et al. method (curves), for the three simulations. The solid curves show estimates using simulated age spectra, whereas the dashed curves, which generally cannot be distinguished from the solid curves, show estimates using inverse Gaussian age spectra.
small, differences in fractional release are the main cause for differences in Cl$_y$.

Figure 5 shows the variation of lower stratospheric concentration (a, b; top panels) and fractional release $F$ (c, d; bottom panels) with mean age $\Gamma$, for CFC-11 (left) and CFC-12 (right). Very similar results are found for other CFCs. Focusing first on the two GCM based simulations we see that the CFC concentrations and $F$ are very similar for $\Gamma<4$ years, but differences occur for older ages. For old ages (polar air) the CFC concentrations are lower for GCM-HIGH, which means that the fraction of CFC converted into Cl$_y$ is larger (i.e. $F$ is larger). Thus, differences in transport can result in differences in the fraction of organic chlorine converted into Cl$_y$, and hence differences in Cl$_y$, for air with the same mean age.

The fractional release rates for the DAS simulation differs from both the GCM simulations. The maximum values of both $\Gamma$ and $F$ are smaller in the DAS simulation (the smaller $F$ is consistent with the smaller lower stratospheric Cl$_y$ in the DAS simulation). For given $\Gamma$, $F$ is larger from the DAS simulation than the GCM simulations. In the lower stratosphere, a given mean age will be found at higher latitude in the DAS simulation compared with the GCM simulations, e.g., air at 20 km with $\Gamma=2.4$ yrs in the DAS simulation is at the South Pole, whereas $\Gamma=2.4$ yrs is in the subtropics in both GCM simulations (Fig. 1). A higher latitude location generally has a higher percentage of air that has been in CFC loss regions, and so higher latitude air with the same age will have higher $F$. If comparisons between $F$ are made at the same location then $F$ is smaller in the DAS simulation than the GCM simulations, and as a result Cl$_y$ is smaller.

The differences in the fractional release of CFCs in the Antarctic vortex can be clearly seen in the vertical profiles of the CFCs, see Fig. 2c, d. One consequence of this is that observations of CFCs can probably be used to differentiate model simulations of Cl$_y$. For example, the solid circles in Fig. 2c, d show the monthly average CFCs near 80$^\circ$ S from the Atmospheric Chemistry Experiment (ACE) (Bernath et al., 2005) observations in September 2005. (The model and ACE CFCs are from September rather than October as ACE samples south of 79$^\circ$ S in early September but not later.) All three simulations overestimate the observed CFC abundances, with the best agreement for the GCM-HIGH simulation. This suggests that GCM-HIGH has the most realistic simulation of Cl$_y$.

Unfortunately, the corresponding profiles of Cl$_y$ cannot be determined directly from ACE as Cl$_2$O (which is a significant reservoir of Cl$_y$ inside Antarctic lower stratospheric vortex in September) is not measured. However, measurements of HCl and ClO in October 2005 by Aura MLS indicate that at 20 km Cl$_y$ is $\approx$3.3 ppb (Fig. 4-8 of WMO, 2007). Consistent with the above CFC comparisons, the modeled Cl$_y$ are too low, with GCM-HIGH the most realistic.
5 Transport pathways

Previous analysis of models has shown that $\Gamma$ is very important for correctly estimating Cl$_y$. However, the above comparisons have shown that although the mean age in the polar lower stratosphere may be the same in two simulations, this does not mean that Cl$_y$ will be the same. It is important to consider the pathways that the air has taken, which can be different even if $\Gamma$ is the same. Even if two parcels have the same $\Gamma$ one parcel could have lower CFC concentrations, higher fractional releases, and larger Cl$_y$ if this parcel spent more time in regions with larger CFC photolysis. This is illustrated schematically in Fig. 6 which shows two parcels with same transit time from the tropopause but different pathways. As a result the two parcels have different photochemical exposure (Schoeberl et al., 2000) and Cl$_y$.

The larger CFC concentrations, and hence smaller Cl$_y$, in the GCM-LOW simulation could be explained by a smaller fraction of the polar air having been in the middle-upper stratosphere where CFCs are photolysed. This is supported by the analysis of Strahan and Polansky (2006). They showed that with lower horizontal resolution there are leakier transport barriers (in subtropics and edges of polar vortices), which results in more rapid transport of young air to polar regions. The leakier barriers also allow more recirculation of air between the tropics and extratropics, allowing air to become old without spending time in the upper stratosphere. The greater influence of young air in the GCM-LOW simulation can also be seen in the age spectra, see Fig. 7 (see also Fig. 14 of Strahan and Polansky, 2006). In the GCM-LOW simulation there is a much higher percentage of young air (<2 years) than in the GCM-HIGH simulation. Also, air arrives continuously in GCM-LOW whereas the GCM-HIGH simulation shows an episodic annual injection of air into the polar region in the transport (Strahan and Polansky, 2006).

6 Conclusions

The CTM simulations presented here show that transport plays a key role in determining the distribution of Cl$_y$, and that large differences in Cl$_y$ can occur in models using the same surface concentrations of source gases and same chemistry. The time air spends in the stratosphere (e.g., the mean age, $\Gamma$) is an important factor but this is not the sole transport factor determining stratospheric Cl$_y$. Stratospheric Cl$_y$ also depends on where the air has been. Differences in transport pathways can result in differences in the fraction of air that has been in the upper stratosphere, where organic chlorine is converted into Cl$_y$, even for air with the same mean age. Thus there can be differences in Cl$_y$ for air with the same $\Gamma$.

This analysis shows that to correctly model Cl$_y$ it is necessary to correctly simulate both the time scales and pathways for stratospheric transport. The mean age provides a stringent test of the transport time scales, and can be inferred from observations. A complementary test of transport pathways is also required. Hall (2000) introduced the “maximum path height” distribution which quantifies transport pathways and complements the age spectrum. However, this quantity cannot be observed. We therefore have to rely on simulations of chemical species to assess the model transport. As shown above transport-induced differences in Cl$_y$ can be clearly seen in profiles of CFCs, and observations of CFCs (e.g., Fig. 2c, d) are available to assess the reality of the transport pathways in models. Hence, comparison of simulated mean age and CFCs with observations provide complementary tests that assess the transport time scales and pathways in models. The complementary nature of tracers with different lifetimes for evaluating transport can also be seen in Schoeberl et al. (2005), where multiple tracers are used to constrain the age spectrum.

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