Drivers of hemispheric differences in return dates of mid-latitude stratospheric ozone to historical levels

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Abstract. Chemistry-climate models (CCMs) project an earlier return of northern mid-latitude total column ozone to 1980 values compared to the southern mid-latitudes. The chemical and dynamical drivers of this hemispheric difference are investigated in this study. The hemispheric asymmetry in return dates is a robust result across different CCMs and is qualitatively independent of the method used to estimate return dates. However, the differences in dates of return to 1980 levels between the southern and northern mid-latitudes can vary between 0 and 30 yr across the range of CCM projections analyzed. Positive linear trends in ozone lead to an earlier return of ozone than expected from the return of Cl₂ to 1980 levels. This forward shift is stronger in the Northern than in the Southern Hemisphere because (i) trends have a larger effect on return dates if the sensitivity of ozone to Cl₂ is lower and (ii) the trends in the Northern Hemisphere are stronger than in the Southern Hemisphere. An attribution analysis performed with two CCMs shows that chemically-induced changes in ozone are the major driver of the earlier return of ozone to 1980 levels in northern mid-latitudes; therefore transport changes are of minor importance. This conclusion is supported by the fact that the spread in the simulated hemispheric difference in return dates across an ensemble of twelve models is only weakly related to the spread in the simulated hemispheric asymmetry of trends in the strength of the Brewer–Dobson circulation. The causes for chemically-induced asymmetric ozone trends relevant for the total column ozone return date differences are found to be (i) stronger increases in ozone production due to enhanced NOₓ concentrations in the Northern Hemisphere lowermost stratosphere and troposphere, (ii) stronger decreases in the destruction rates of ozone by the NOₓ cycle in the Northern Hemisphere lower stratosphere linked to effects of dynamics and temperature on NOₓ concentrations, and (iii) an increasing efficiency of heterogeneous ozone destruction by Cl₂ in the Southern Hemisphere mid-latitudes as a result of decreasing lower stratospheric temperatures.

1 Introduction

In support of the WMO/UNEP (World Meteorological Organization/United Nations Environmental Programme) ozone assessment 2010 (WMO, 2011), the SPARC (Stratospheric Processes And their Role in Climate) CCMVal-2 activity coordinated simulations of the future evolution of atmospheric ozone using about a dozen chemistry-climate models (CCMs). These simulations have been extensively analyzed and evaluated (SPARC-CCMVal, 2010). As a result of the success of the Montreal Protocol and its amendments and adjustments, tropospheric concentrations of ozone depleting substances (ODSs) are decreasing (Montzka et al., 1999) and are expected to continue to decrease over the next century. In mid-latitudes, stratospheric Cl₂ is projected by CCMs to return to 1980 levels by the middle of the 21st century. Total column ozone (TOZ), on the other hand, is projected by CCMs to return to 1980 values earlier: by the early 2020s over northern mid-latitudes and by the mid-2030s over southern mid-latitudes (Austin et al., 2010; Eyring et al., 2010). Unlike Cl₂ where the return to 1980 values is hemispherically symmetric (Austin et al., 2010), the return of TOZ
to 1980 values is hemispherically asymmetric. In Austin et al. (2010) and Oman et al. (2010) it was shown that this hemispheric asymmetry results from changes in lower stratospheric ozone, specifically ozone below 20 hPa.

In addition to being influenced by ozone depleting substances (ODSs), ozone concentrations are also affected by changes in greenhouse gas (GHG) concentrations. While ODS concentrations increased strongly in the second half of the 20th century, and are expected to decline again in the 21st century, GHG concentrations are projected to keep rising monotonically. Eyring et al. (2010) used a multi-model set of sensitivity simulations, in which either ODS or GHG concentrations were held constant, to show that the hemispheric difference in mid-latitude ozone evolution is due to a stronger response of Northern Hemisphere (NH) ozone to increasing GHG concentrations compared to its Southern Hemisphere (SH) counterpart.

A number of processes determine the response of mid-latitude ozone to increasing GHG concentrations. In the upper stratosphere, increasing GHG concentrations cause temperatures to decrease which slows the \( O + O_3 \rightarrow 2O_2 \) reaction such that ozone increases (Portmann and Solomon, 2007). Increases in the concentrations of \( N_2O \) and \( CH_4 \) elevate the concentrations of the ozone precursors \( NO_x \) \((=NO+NO_2)\) and \( HO_x \) \((=H+OH+HO_2)\). However, the response of ozone to changes in \( N_2O \) and \( CH_4 \) depends on a number of coupled processes. For example, the ratio of \( NO_x \) to \( N_2O \) is temperature-dependent, such that decreasing temperatures lower the yield of \( NO_x \) from \( N_2O \) (Plummer et al., 2010; Revell et al., 2012b). The speed and path of air parcels through the Brewer–Dobson Circulation (BDC) also affects \( NO_x \) concentrations by determining the time available for photolysis of \( N_2O \) (Cook and Roscoe, 2012). Since CCMs consistently project a strengthening of the BDC (Butchart et al., 2010), this will also reduce the yield of \( NO_x \) from \( N_2O \) (Cook and Roscoe, 2012).

In previous studies, the hemispheric asymmetry in the response of ozone to increasing GHG concentrations was attributed to hemispheric asymmetries in changes in transport; the strength of the BDC is projected to increase more in the NH compared to the SH (WMO, 2011; Eyring et al., 2010; Austin et al., 2010). However, little evidence has been presented to support this hypothesis. In this study, the role of changes in transport on the hemispheric asymmetry of return dates of mid-latitude ozone to 1980 levels is reassessed. Two complementary approaches are followed: first, a multi-model ensemble of CCMVal-2 models is used to investigate the robustness of projections of the temporal evolution of ozone, and then two CCMs (described in Sect. 2) that incorporate the necessary diagnostics are used to conduct a detailed attribution of the drivers of mid-latitude ozone changes through the 21st century. These diagnostics, along with other methods used in the analysis, are described in Sect. 3. In Sect. 4, the multi-model analysis is presented, testing the consistency of the hemispheric asymmetry in return dates across models, evaluating the height dependence of the hemispheric differences and testing their connections to changes in the BDC. Based on two of the CCMs, an attribution is performed, allowing an explicit separation of the effects of chemistry and transport (Sect. 5). The chemically-induced ozone changes and their drivers are analyzed in more detail in Sect. 6. Uncertainties and the relevance of the processes found are discussed in Sect. 7. Conclusions are presented in Sect. 8.

## 2 Chemistry-climate models

Within CCMVal-2 (SPARC-CCMVal, 2010) a set of simulations was performed with a number of CCMs aiming at investigating the stratospheric ozone layer. Our analysis requires seamless simulations from the past to the future, and the necessary output of these simulations was available from 12 CCMs. The individual models are not identified as this is not required for the analyses that aim to explore the spread in the simulations rather than the performance of individual models. A description of models and the simulation setups can be found in Morgenstern et al. (2010). For 10 of the models, REF-B2 simulations are analyzed. The REF-B2 simulations are designated to consistently simulate past and future climate, extending from 1960 to 2100, and include anthropogenic forcing by GHG and ODS emissions. The GHG emissions follow the Special Report on Emissions Scenarios (SRES) A1B scenario. The SCB-B2d simulation, used for the remaining two models, differs from REF-B2 in that it includes natural variability, i.e., the solar cycle and the QBO (quasi-biennial-oscillation) (see Eyring et al., 2008). However, for the purposes of this study, the two simulations are equivalent.

Two CCMs that are equipped with the necessary diagnostics are used more extensively for the attribution analysis and are described in greater detail below.

### 2.1 NIWA-SOCOL

The NIWA-SOCOL (National Institute of Water and Atmospheric Research – SOLar Climate Ozone Links) CCM is a modified version of the CCM SOCOL_v2.0 (Schranner et al., 2008) in which stratospheric ozone is simulated, including a budget analysis of ozone production, loss from different catalytic cycles and transport. SOCOL comprises the MAECHAM4 global climate model and a modified version of the chemistry transport model MEZON (Egorova et al., 2003). MAECHAM4 is configured with a T30 spectral horizontal resolution with 39 vertical levels between Earth’s surface and 0.01 hPa (\( \sim 80 \) km). A hybrid transport scheme is employed to advect the chemical constituents. The chemical solver algorithm uses a Newton–Raphson iterative method, taking into account 41 chemical species, 140 gas-phase reactions, 46 photolysis reactions and 16 heterogeneous reactions.
The output used in this study originates from a NIWA-SOCOL REF-B2 simulation. This simulation is an updated version of the one performed for the SPARC CCMVal-2 activity in that the surface sea temperatures (SSTs) simulated by ECHAM5-MPI-OM are used. Otherwise the boundary conditions are as described in Morgenstern et al. (2010) and are consistent with the CCMVal-2 specifications. In the NIWA-SOCOL simulation, a minor error was made in the prescription of the NOx boundary condition, i.e., aircraft NOx emissions were inadvertently doubled. However, since NIWA-SOCOL is used to study ozone changes mostly above 100 hPa, this error is not expected to impact the results.

A diagnostic implemented within the model tracks the change in the ozone concentration within each model grid cell, at each time step, resulting from net chemical production and net transport into the cell. The change in net chemical production is calculated explicitly, and can be further attributed to different catalytic cycles. Further attribution of odd oxygen ($O + O_3$) production, via the Chapman cycles, and destruction by the HOx, NOx and Clx cycles, across 15 different catalytic cycles, was performed within the model chemistry scheme using the rate limiting steps of the corresponding reaction cycles. This diagnostic, along with the detailed list of reaction cycles, is described in Revell et al. (2012b). In this paper, destruction by bromine-containing species is included in the Clx destruction term. The change in ozone due to transport is defined as the residual of the net chemical production subtracted from the total ozone change. Both terms are recorded and accumulated into monthly means for each grid cell. Since transport is determined as a residual, any numerical computation residual and renormalization of the ozone concentrations are folded into the transport term.

### 2.2 E39CA

The CCM ECHAM4.L39(DLR)/CHEM/ATTILA (E39CA) is an updated version of ECHAM4.L39(DLR)/CHEM (E39C) (Hein et al., 2001; Dameris et al., 2005) with the former semi-Lagrangian advection scheme replaced by the fully Lagrangian advection scheme ATTILA (Reitmeier and Sausen, 2002; Stenke et al., 2009). E39C is based on the spectral general circulation model ECHAM4.L39(DLR) (Land et al., 2002) and the chemistry-module CHEM (Steil et al., 1998). The model is run with a spectral horizontal resolution of T30, corresponding to $\sim 3.75^\circ \times 3.75^\circ$ on a transformed latitude–longitude grid. The model consists of 39 layers in the vertical, extending from the surface to the uppermost layer centered at 10 hPa. The chemistry module CHEM is based on a generalized family concept and includes homogeneous and stratospheric heterogeneous ozone chemistry and the most relevant chemical processes for describing tropospheric background chemistry. Ozone depletion cycles involving bromine were originally not implemented in the CHEM module, but subsequently a parameterization for the bromine chemistry was included. The parameterization is based on the photolysis of ClxO2 and described in detail in Stenke et al. (2009). Boundary conditions for the two families Clx (HCl + ClONO2 + ClO3) and NOy (= NOX+HNO3) are prescribed at the uppermost model level (10 hPa) to account for chemical processes above the model top. For a more detailed description of E39CA, see Stenke et al. (2009). The SCN-B2d simulation performed with E39CA, as submitted to the CCMVal-2 archives, is used here. The simulation extends from 1960 to 2049, and the boundary conditions including anthropogenic and natural forcings are described in Garny et al. (2009).

As in NIWA-SOCOL, ozone production and destruction rates from a prescribed set of chemical reaction cycles are calculated from the rate-limiting reaction steps, and are saved as part of the model output. The total chemical tendencies of ozone are then calculated off-line from the sum of the tendencies of the chemical cycles. The transport tendencies are calculated by subtracting the ozone chemical tendencies from the total ozone tendencies. Thus, transport tendencies are calculated as residuals in both models: in NIWA-SOCOL online during the simulation and in E39CA off-line after the simulation was performed. Therefore, transport tendencies include in both models any spurious contributions to ozone tendencies resulting from numerical artifacts within the model. Excessive diffusion due to the numerical representation of advection is thus also part of the transport tendency, as it should be since it is part of ozone transport in the models.

Chemical reaction cycles are grouped, as in NIWA-SOCOL, into production by the Chapman cycle, and destruction by the HOx, NOx and Clx cycles. As mentioned above, any bromine-related ozone destruction is included in the Clx loss term. Additionally, in E39CA the rate-limiting reaction rates of two of the main tropospheric ozone production cycles are saved as output. These rate-limiting reactions are (1) $HO_2 + NO \rightarrow OH + NO_2$ (in the following referenced as “PHN”) and (2) $CH_3O_2 + NO \rightarrow CH_3O + NO_2$ (referenced as “PCN”). Dry deposition at the surface is also considered for the budget calculation.

### 3 Methods

In this study, return dates of ozone to historical values are estimated using a linear regression approach, as described in the following in Sect. 3.1. Apart from providing a smooth time series that is needed for the return date calculation, the regression model is used for the statistical separation between that part of each time series that is congruent with the evolution of Clx and that part that is congruent with a linear trend. The linear trend is responsible for driving ozone return dates away from the return dates of Clx. We argue that the trends are responsible for the hemispheric asymmetric return dates (see Sect. 4.1). The physical processes responsible for the trends are further analyzed in Sect. 5. A method is used
that attributes ozone changes to changes in chemistry and transport (described in Sect. 3.2). Thus, the ozone trends can be separated into chemically-induced and transport-induced ozone trends.

3.1 Estimation of return dates

To estimate dates of return of ozone to 1980 levels, the time series need to be smoothed to remove the effects of unforced (internal) variability that would otherwise make the return date evaluation susceptible to climate noise. For the analysis in support of the 2010 Ozone Assessment (WMO, 2011), a time-series adaptive model analysis (TSAM) method (Scinocca et al., 2010) was used. This method separates the time series into a smooth signal and superimposed noise, based on a statistical non-parametric fit. An alternative approach is to iteratively smooth the time series by applying a 1 : 2 : 1 filter many times. A comparison of the TSAM method with the iterative 1 : 2 : 1 smoothing can be found in the supplement to Chapter 9 of the SPARC CCMVal-2 report (SPARC-CCMVal, 2010). This comparison shows that the multi-model mean estimate of mid-latitude ozone return dates is independent of the method used. For individual models, on the other hand, the choice of the method can lead to large differences in the return dates.

A conceptually different approach to separating the signal from the noise is to apply a linear least squares regression model with given geophysical predictors, i.e., using a priori information on the known drivers of the long-term changes in ozone. In practice, these predictors are the burden of Cl\(_y\) (known to be the cause of large depletion of ozone) and a linear trend, which is assumed to approximately represent the response to changes in GHG concentrations and other emissions that increase monotonically (e.g., NO\(_x\) emissions at the surface). Thus, the regression model is of the form

\[
O_3(t) = a_0 + a_1 \times Cl_y(t) + a_2 \times t + R(t),
\]

where \(a_0\), \(a_1\) and \(a_2\) are the offset, the \(Cl_y\) and the trend regression coefficients, respectively, and \(R\) is the residual. More details can be found in Appendix A. While this regression approach might be criticized for assuming ozone to be driven only by \(Cl_y\) or some linear process, it has the advantage of permitting a separation between these two terms. In particular, if the goal is to understand the deviation of ozone return dates from \(Cl_y\) return dates, the regression performed in this way separates the secular evolution of ozone into a component that is congruent with the \(Cl_y\) evolution (i.e., returns to 1980 values when \(Cl_y\) does) and a component that drives deviations away from the \(Cl_y\) return dates. This physical interpretation of the regression model attribution between its two basis functions requires a sufficiently long time series so that the \(Cl_y\) and linear trend basis functions are approximately orthogonal. This is achieved by considering the 1960 to 2049 period. The fitted time series from this regression model that attributes ozone changes to \(Cl_y\) and the linear trend term is then used to estimate return dates to 1980 values. As shown in Appendix A, this method of calculating return dates gives similar results to other methods; thus the overall results are independent of the method used for the return date estimation.

Dates of return of TOZ to 1980 values in the mid-latitudes (45–60°) of both hemispheres for twelve different CCMs are shown in Fig. 1. Compared to earlier estimates (Austin et al., 2010; Eyring et al., 2010), the multi-model mean return dates derived here are slightly earlier in both hemispheres. This might result from a different choice of the geographical region (60 to 45° N as opposed to 35° N), for reasons explained later, or from a different subset of CCMs used here. However, the 10 to 15 yr earlier return of northern mid-latitude TOZ to 1980 values compared to southern mid-latitude TOZ is in close agreement with estimates by Austin et al. (2010) and Eyring et al. (2010).

3.2 Ozone attribution method

Ozone changes are attributed to sources and sinks (chemical production and loss and transport of ozone) as described in Garny et al. (2011). A short summary of the method is given in the following. The required model output are the chemical ozone tendencies (the total ozone production and loss and transport of ozone) as described in Garny et al. (2011), changes in ozone between two time periods were attributed to changes in chemical production, destruction and transport. It was assumed that the annual mean ozone tendencies, averaged over the period of interest, were close to zero. Here, this method is extended so that it can be applied to time series of ozone, i.e., the method attributes year-to-year changes to changes in the sink and source terms.

The ozone budget equation for the annual mean change in ozone for two periods \(p_1\) and \(p_2\) is the following (see Eq. 4 of Garny et al., 2011):

\[
\begin{align*}
\frac{\partial O_3}{\partial t}^{p_1} & = [P]^{p_1} - [DO_3]^{p_1} + [T]^{p_1} \\
\frac{\partial O_3}{\partial t}^{p_2} & = [P]^{p_2} - [DO_3]^{p_2} + [T]^{p_2}. 
\end{align*}
\]

The integration denoted by \([\cdot]\) is the integral over the time periods \(p_1\) and \(p_2\), respectively. \(O_3\) is the ozone mixing ratio within some model grid cell, \(P\) is the ozone production within the grid cell, and \(D\) is the fraction of the ozone destroyed in that time step within the grid cell. \(T\) is the net amount of ozone transported into or out of the grid cell. In contrast to Garny et al. (2011), the term \(\frac{\partial O_3}{\partial t}\) is not assumed to be zero, allowing periods to be a single year.
For $[DO_3] \approx [D][O_3]$, the equations above can be transformed to describe the relative change in ozone as

$$\frac{[O_3]^p_1 - [O_3]^p_2}{[O_3]^p_1} = \frac{[D]^p_1 - [D]^p_2}{[D]^p_2}$$

relative ozone change  \hspace{2cm} \text{Destruction-induced O}_3 \text{ change}


Production-induced O$_3$ change  \hspace{2cm} \text{Transport-induced O}_3 \text{ change}

$$\frac{\Delta [P^p_1] - \Delta [T]^p_1}{[P]^p_1 + [T]^p_1 - \Delta [T]^p_1} + \frac{\delta D \times \delta P}{\text{Nonlinear, chemistry}}$$

Imbalance term  \hspace{2cm} \text{Nonlinear, mixed}

$$\frac{\delta D \times (\delta T + \delta \Delta)}{}.$$  \hspace{2cm} \text{(2)}$$

The first three terms on the right-hand side of the equation are the contributions to ozone changes due to changes in destruction, production and transport, respectively. $\Delta$ is the ozone tendency $\frac{\partial O_3}{\partial t}$ averaged over the respective periods, and the fourth term is therefore the contribution to ozone changes due to a difference in the total gain or loss of ozone over one period compared to the other. Hereafter this is referred to as “imbalance” term. The last two terms in the equation describe the nonlinear contributions, where the $\delta$ denotes the change in ozone due to changes in the respective source or sink, i.e., the first to fourth terms. The nonlinear terms consist of the changes in the destruction rates applied to all other terms (production, transport and the imbalance term). For example, if there is more ozone production from some chemical cycle during a time step, but it is accompanied by an enhanced destruction rate from a different cycle during the same time step, the elevated ozone is immediately subjected to the enhanced destruction, reducing the effect of the additional production. Similarly, stronger transport of ozone into a grid cell might also be mitigated by local changes in the ozone destruction rate, as described by the nonlinear term. The nonlinear terms are one order of magnitude smaller than the other change terms, so as long as changes are small ($< 0.1$), this term is one order of magnitude smaller, and thus not important.

In the analysis presented below, transport and chemically-induced ozone changes are separated. The changes due to transport are diagnosed from the third term in Eq. (2). The changes due to chemistry are the sum of the production, destruction and the first nonlinear term (i.e., the first, second and fifth terms in Eq. 2). Hereafter, when referring to the nonlinear term, only the mixed transport–chemistry term (6th term in Eq. 2) is meant.

In Fig. 2, an example of the application of the method is demonstrated for lower stratospheric, partial column, zonal mean ozone at 45° S from 1960 to 2049. Ozone changes in each year ($= p_2$) are calculated relative to the mean over 1960–1969 ($= p_1$), and are reasonably well reproduced by the sum of the terms on the right-hand side of Eq. (2). The
individual components of these terms are shown, indicating the extent to which changes in the respective quantity contribute to the relative change in ozone in a particular year. For example, the red line indicates how much ozone would have changed as a result of changes in transport alone. The largest contributions to ozone changes are from changes in chemistry and transport. The imbalance-term (Δ) has a minor contribution to interannual variability and none to the long-term decrease in ozone over this period at this location. The nonlinear term has a negligible contribution in this case.

4 Multi-model analysis

4.1 Height dependence of hemispheric difference in return dates

The dates of return of mid-latitude TOZ to 1980 levels are found to be hemispherically asymmetric in the multi-model mean shown in WMO (2011). Here, we use twelve of the CCMs that were used in the WMO/UNEP ozone assessment to investigate inter-model differences in the asymmetry of the return dates and which altitude regions are primarily responsible for the asymmetry.

Mid-latitude ozone is defined as the area weighted average over the latitude band 45 to 60° of either hemisphere. This is narrower compared to the WMO (2011) definition of mid-latitude ozone (35 to 60°). As shown in Sect. 5.2, strengthening of the BDC tends to induce negative ozone trends over low latitudes and positive ozone trends at middle to high latitudes. Because the zero trend in ozone is usually around 45° N/S, this is used as the equatorward boundary of the mid-latitudes to minimize any reduction in the diagnosed effects of transport on ozone trends.

The hemispheric differences in return dates of mid-latitude TOZ for the twelve models are shown in the leftmost panel of Fig. 3, calculated from the regression model fit as described in Sect. 3.1. The multi-model mean ozone returns to 1980 levels about 13 yr earlier in the NH compared to the SH, in agreement with WMO (2011), and this results is robust against using different methods for the return date calculation (see Appendix A). However, there is a large spread in the hemispheric differences among the individual models, ranging from 0 to close to 30 yr.

To evaluate the relative contributions of different altitude regions to the hemispheric differences in return dates, partial ozone columns were calculated for (i) the surface to 10 hPa, (ii) the tropopause to 10 hPa and (iii) the region from 100 to 10 hPa. Thus, the altitude domain of regarded ozone columns is stepwise reduced, excluding first the upper stratosphere, further the troposphere and in a third step also the lowermost stratosphere (i.e., region between the tropopause and 100 hPa). The upper boundary of the lower stratosphere at 10 hPa was chosen to separate the region where both dynamics and chemistry control ozone from the region where ozone is controlled purely by chemistry (as will be shown for ozone trends in Fig. 10). As shown in Fig. 3, excluding the region above 10 hPa slightly amplifies the hemispheric asymmetry in return dates. As ozone above 10 hPa is primarily chemically controlled and ozone changes tend to be correlated with temperature changes which are generally hemispherically symmetric, including the region above 10 hPa obscures some of the hemispheric asymmetry. Additionally excluding the tropospheric ozone column also has a small effect, in this case slightly reducing the hemispheric asymmetry in return dates. As shown by panel f in Fig. 3, the hemispheric asymmetry induced by tropospheric ozone shows a large spread between models that even show differences in sign. Given that tropospheric chemistry is treated quite differently, and in many cases rather simplistically in the CCMs, the large spread is not surprising. The contribution of the lowermost stratosphere (LMSTR; hereafter defined as the layer between the tropopause and 100 hPa), on the other hand, accounts for about half of the hemispheric difference in return dates (the exact contribution depending on the method used to determine the return date). In all but two models, excluding the LMSTR causes the hemispheric differences in return dates to decrease. One might suspect that the time evolution of tropospheric and lowermost stratospheric partial ozone columns
Fig. 3. Hemispheric difference ((45° to 60° S) − (45° to 60° N)) in return dates of ozone to 1980 values for twelve different models (back crosses with uncertainty bars) and the multi-model mean (triangle) for (a) total column ozone, and partial column ozone (b) from the surface to 10 hPa, (c) from the tropopause to 10 hPa and (d) 100 hPa to 10 hPa. Furthermore the impact on hemispheric differences in return dates from the upper stratosphere (e = a−b), from the tropopause (f = b−c) and from the lowermost stratosphere (g = c−d) is shown. Multi-model means are also shown for the two additional methods described in Appendix A: method 2 (red) and 3 (blue). The green markers indicate the hemispheric difference in return dates that would result if the linear trend in the NH was the same than in the SH.

are affected by trends in the height of the tropopause. However, when calculating the partial ozone columns with a fixed tropopause at its mean 1960s value, the results on return date differences are unaffected in the multi-model mean. Thus, trends in the tropopause height cannot be made responsible for introducing hemispheric asymmetries in ozone return dates, probably because tropopause trends themselves do not have a strong hemispheric asymmetry. Ozone evolution in the lower stratosphere (LSTR; 100 to 10 hPa) accounts next to the LMSTR for the other half of the hemispheric difference in the return dates. A large spread between the models is found in the LSTR, and in about half the models the hemispheric difference is only marginally significant when considering the uncertainty resulting from the different methods used to calculate 1980 return dates. These results suggest that it is the LMSTR and LSTR that make the largest contribution to the hemispheric asymmetry in TOZ return dates, in agreement with Oman et al. (2010) and Austin et al. (2010). However, large model-to-model differences and uncertainties in return date estimates exist.

The contributions of ozone at each level to TOZ (i.e., the partial column of ozone over the model layer centered at the respective level) and its response to Cl$_2$ and the linear trend is shown in Fig. 4. The models agree well in the climatological profile (upper left panel of Fig. 4). The Cl$_2$ regression coefficients derived for each of the twelve CCMs (middle panel) also agree relatively well, with a negative response of ozone to Cl$_2$ maximizing at 50 hPa. The response of mid-latitude ozone to Cl$_2$ is stronger in the SH than in the NH, which is well known (Solomon, 1999). The linear trend (upper right panel) shows significant inter-model differences in both amplitude and vertical structure. Most models show a positive trend in the troposphere; in the multi-model mean the trend is stronger in the NH than in the SH. The multi-model mean trends show a local maximum at ~150 hPa. Above this level, trends in both hemispheres decrease with altitude up to ~70 hPa. In the SH, about half of the models have a negative ozone trend between 70 and 50 hPa. Another local maximum, with positive trends in both hemispheres, is found at 30 hPa. Above 30 hPa, trends decrease towards zero and become hemispherically symmetric in the multi-model mean. The structure of the trends agrees well with the differences in cumulative ozone columns from the mid-20th to the end of the 21st century shown by Plummer et al. (2010). They used sensitivity simulations to attribute these trends almost fully to changes in GHG concentrations, thus validating our
method of separating the effects of ODSs and GHGs through linear regression.

How do the linear trends in ozone translate to ozone return dates? If there is no linear trend, ozone returns to 1980 values just when Cl\textsubscript{y} does. Further, if the return dates of Cl\textsubscript{y} are the same in both hemispheres, ozone in the NH and SH would also return to its 1980 values at the same time, independent of its sensitivity to Cl\textsubscript{y}. As shown in Fig. 1, the multi-model mean Cl\textsubscript{y} returns to 1980 levels at the same time in each hemisphere, and the hemispheric asymmetry in ozone return dates can in none of the models be fully explained by Cl\textsubscript{y} return date differences. If ozone is subject to a positive linear trend, the return date of ozone is shifted forward. However, as illustrated in Fig. 5, the amount by which the return date is shifted depends on the sensitivity of ozone to Cl\textsubscript{y}. Therefore, even if linear trends in ozone are the same in the NH and SH, the return dates to 1980 values can be different. Larger linear trends in the NH than in the SH, as found for most models (see Fig. 4), cause even larger differences in the ozone return dates between the NH and SH (illustrated in the middle panel in Fig. 5). The dependence of the return date on the combination of the Cl\textsubscript{y} and the trend regression coefficient can be expressed in the following way: the ozone anomaly relative to 1980 is described by \( \Delta O_3(t) = a_1 \times Cl_y(t) + a_2 \times t \), and thus ozone returns when \( t = -a_1/a_2 \times Cl_y(t) \). Thus, the ratio of the Cl\textsubscript{y} to the trend regression coefficient determines \( \Delta O_3(t) \).
the shift of the return dates (see lower left in Fig. 5). As illustrated in the lower right panel in Fig. 5, given a stronger Cl$_y$ sensitivity in the SH than in the NH (assuming 2 and 1 DU ppb$^{-1}$, respectively), the linear trend in the NH would need to be smaller than in the SH by the same ratio (i.e., here 1/2) to result in identical return dates. However, the CCMs simulate a stronger positive trend in NH ozone. Thus, overall both the stronger sensitivity of SH ozone to Cl$_y$ and the stronger positive trend in NH ozone lead to earlier ozone return dates in the NH. The relative role of these effects is quantified by calculating ozone return dates for the NH assuming that they would be subject to the same (weaker) trend as ozone in the SH. The resulting hemispheric differences in return dates are indicated in Fig. 3 as green markers. These show that the hemispheric differences in ozone return dates would be about half as strong if trends were the same in both hemispheres. Thus, both the different effect of positive trends on return dates in the two hemispheres as well as the hemispheric differences in the trends are causing the hemispheric differences in ozone return dates. Therefore, it is important to understand the origin of these linear trends in ozone in both hemispheres. In Sect. 5 the physical drivers of trends will be investigated in detail.

4.2 Relation of hemispheric differences in the BDC to ozone return dates

The annual mean downward mass flux in each hemisphere, calculated from the residual circulation stream-function, is used as a measure of the strength of the BDC. The sum of the two hemispheric values is the total downward mass flux, equaling, in absolute value, the total upward mass flux, or tropical up-welling, which is often used to quantify the strength of the BDC. The total downward mass flux in each hemisphere equals the meridional mass flux, which is responsible for transporting ozone from the tropics to mid-latitudes.
Trends in total downward mass flux in each hemisphere are shown in Fig. 6 (left) as a function of pressure. The trends are calculated using the same linear regression model as used to calculate the ozone trends. The mass flux increases in both hemispheres and in all models, consistent with the BDC strengthening, as shown by Butchart et al. (2010). The multimodel mean trend in the NH is stronger than in the SH between 90 and 30 hPa (see Fig. 6, right panel). At 100 hPa, however, the trend is slightly stronger in the SH than in the NH.

If the hemispheric differences in the strengthening of the BDC were the cause of the hemispheric differences in ozone return dates, it would be expected that in models with a stronger hemispheric difference in the strengthening of the BDC, the hemispheric return dates would show greater separation. The relation between hemispheric differences in the strengthening of the BDC and hemispheric differences in ozone return dates is shown for the set of twelve CCMs in Fig. 7. The LSTR return date differences are plotted against the 70 hPa hemispheric differences in the mass flux trends (i.e., the 70 hPa values shown in Fig. 6 (left); 70 hPa is chosen since the hemispheric difference maximizes there). No statistically significant relationship between hemispheric differences in changes of the BDC and hemispheric differences in return dates are found (the T test value is 0.6, where significance at the 80% level is obtained at T = 1.36 for this sample size). Even when excluding the outlier seen on the lower right of the upper panel of Fig. 7 (with stronger circulation changes in the SH than NH), the correlation is not significant. The LMSTR return date differences are compared with the hemispheric differences in the mass flux trends at 100 hPa in the lower panel of Fig. 7. In the LMSTR a significant positive correlation at the 80% significance level is found. While the statistical significance is still rather weak, it provides evidence for some influence of hemispheric asymmetry in BDC changes being responsible for the hemispheric asymmetry in ozone return dates. Half of the models have a negative residual mass flux trend difference, i.e., the SH mass flux increases more than its NH counterpart (see also right panel of Fig. 6). The return date differences for five of these models are, however, positive, i.e., return dates in the NH are earlier than in the SH. This might seem contradictory, however, according to the arguments presented around Fig. 5, a smaller trend in the NH can still have a larger effect on ozone return dates due to its smaller Cl sensitivity as compared to the SH. Since generally stronger ozone trends can lead to larger hemispheric differences due to their different effects in NH and SH (see Fig. 5, lower panels), one might suspect that the trends in the BDC itself, rather than its hemispheric differences, might correlate better with the
ozone return date differences. However, this is neither the case in the LSTR or the LMSTR. Overall, these results suggest that other processes than BDC changes play a significant role.

5 The role of transport and chemistry in driving ozone changes

The results presented in the previous section suggest that hemispheric differences in the strengthening of the BDC are unlikely to completely explain the hemispheric differences in ozone return dates, as has been previously thought (WMO, 2011). In this section, two CCMs, E39CA and NIWA-SOCOL, each of which incorporate a diagnostic to separate between chemical- and transport-induced changes in ozone (see Sect. 3.2), are used to investigate the processes leading to hemispheric asymmetric ozone trends.

5.1 Lowermost stratosphere and troposphere

The E39CA model is suitable for examining the ozone changes in the LMSTR since (i) the model behaves in a similar fashion to the multi-model mean in this region (see Figs. 4 and 6) and (ii) all relevant reaction cycles for this region are saved as output. Unfortunately, the tropospheric reaction cycles are not diagnosed in NIWA-SOCOL; hence only E39CA is used here.

The regression model including a Cl and linear trend term is applied to the time series of relative deviations of ozone from the 1960s (as shown exemplarily in Fig. 2 for the time series at 45° S). In Fig. 8, the trend coefficient of the regression model applied to mid-latitude ozone (averaged over 45 to 60°) is shown as a function of height. The resulting units of the trends are percentage change in ozone per year.

Ozone increases linearly in the LMSTR and troposphere and the trend is stronger in the NH. The chemistry–transport diagnostic allows these trends to be attributed to trends in chemistry (blue lines) and trends in transport (red lines), as calculated using Eq. (2). The nonlinear term (6th term in Eq. 2) is also shown in Fig. 8 and it can be seen that this term becomes relevant only for large values of the chemistry and transport changes, e.g., around 250 hPa. The sum of the chemistry, transport and nonlinear term is shown as the control trace (grey line). If this control line deviates from the net ozone trend, the imbalance term becomes important, and the results can no longer be easily interpreted. This occurs when the variability in the time series is large.

As shown in Fig. 8, in both hemispheres, between about 100 and 150 hPa, the positive trends in ozone result from positive trends in both transport- and chemically-induced changes in ozone. Below 150 hPa, changes in chemistry drive a positive trend in ozone which maximizes at around 250 hPa. However, the chemically-induced positive trend is largely offset by transport-induced negative ozone trends. The compensation of chemical effects by transport can be understood as follows: in the mean over the 1960s, at 250 hPa, production (of 0.32 DU month⁻¹) is balanced by transport (of −0.20 DU month⁻¹) and by destruction (of −0.12 DU month⁻¹). Production increases to 0.53 DU month⁻¹ in the 2040s, and this increase in
production is mainly balanced by increased transport away from the source region (i.e., transport decreases to \(-0.37\text{ DU month}^{-1}\)), with the remainder balanced by destruction (decreasing to \(-0.16\text{ DU month}^{-1}\)). Thus, enhanced production leads to more ozone being available at any time step, and therefore more ozone can be transported away from this region leading to a negative effect on ozone values by transport. This effect becomes even more evident when employing an ozone origin diagnostic, as presented for E39CA in Garny et al. (2011) (see their Fig. 10).

The strong chemically-induced trends in ozone occur close to the tropopause, which moves upward from the 1960s to the 2040s. One might suspect that the trend in the tropopause height is in some way responsible for the ozone trends. We calculated the trends in ozone profiles in vertical coordinates relative to the tropopause height (not shown). The slight reduction in positive trends in ozone at and above the tropopause in both the NH and SH seen in Fig. 8 vanishes when calculating the trends relative to the tropopause, as can be expected from an upward shift of the tropopause. Otherwise, the results are unaffected, and we can conclude that the chemically-induced positive ozone trend around 250 hPa is not the result of a vertical shift of the tropospheric regime.

Below 500 hPa in the NH and around 300 hPa in the SH, the transport term becomes positive, reinforcing the positive ozone trend. This suggests that ozone transport into this region dominates transport out of this region, resulting from the stronger local production of ozone. It is most likely that it is ozone produced above these levels, and transported downward, possibly together with horizontal transport, that leads to the overall positive contribution to ozone changes. Close to the surface, particularly in the NH, the increase in ozone due to chemistry again dominates. Again, transport changes offset the chemically-induced positive trend.

The hemispheric differences are highlighted in the lower-most panel in Fig. 8, showing that ozone trends are stronger in the NH compared to the SH, in particular in the upper troposphere. The stronger increase in ozone in the NH can be attributed to stronger chemically-induced ozone trends. The contribution from changes in transport largely offsets the contribution from changes in chemistry, but does not itself lead to asymmetric ozone trends in the LSTR and troposphere. The induced asymmetry by chemistry is due to ozone production by NOx, which increases more strongly in the NH, as will be shown in Sect. 6.1.

### 5.2 Lower stratosphere

Causes of ozone trends in the LSTR are investigated in more detail using the NIWA-SOCOL model. The ozone trends in this model behave similarly to the multi-model mean in the LSTR (see Figs. 4 and 6). E39CA, on the other hand, does not simulate the earlier return of NH ozone in the LSTR to 1980 levels, and the ozone trend profile is very different to most other models.

First, the latitudinal structure of trends in the LSTR partial ozone column from NIWA-SOCOL is shown in Fig. 9. Ozone trends are negative in the tropics, and positive at middle to high latitudes. Extratropical ozone trends are slightly stronger in the NH compared to the tropics, and positive at middle to high latitudes. Extratropical ozone trends are slightly stronger in the NH compared to the SH, consistent with earlier ozone return dates in the NH. The attribution of ozone trends shows that transport-induced changes lead to a decrease in ozone in the tropics and increases in the extratropics. The transport-induced trends change sign at about 45° N/S, motivating the choice of that latitude as equatorward boundary for our definition of mid-latitudes. The transport-induced ozone trends are consistent with an enhanced strength of the BDC (see Fig. 6). However, the stronger northern mid-latitude trends compared to southern mid-latitudes in NIWA-SOCOL cannot be explained by differences in transport. Rather, transport changes induce stronger positive ozone trends in the southern mid-latitudes compared to the northern mid-latitudes. The transport-induced trends in the SH are, partially, compensated by strong negative trends induced by chemistry changes.

To elucidate this point further, profiles of mid-latitude (45–60° N/S) ozone trends, attributed to changes in chemistry and transport within NIWA-SOCOL, are shown in Fig. 10. Ozone trends at 100 hPa are positive in both hemispheres, decreasing with height up to 70 hPa, where the trend in the SH is slightly negative. Above 70 hPa trends increase with height and are positive throughout the LSTR.

Ozone trends induced by transport are positive at all levels in the LSTR in both hemispheres, but maximize at 100 hPa and are close to zero above 20 hPa. Ozone trends due to
changes in chemistry are negative to around 40 to 30 hPa, and dominate the positive ozone trend above 20 hPa in both hemispheres. The positive trend induced by transport is again consistent with a strengthened BDC, and the decrease of the effect with height is expected from the decreasing lifetime of ozone with height.

The hemispheric differences in ozone trends, as shown in the lower panel of Fig. 10, are positive throughout the LSTR, maximizing at 50 hPa. Changes in transport are the main cause of relatively stronger NH trends below 80 hPa, and are slightly negative below. The hemispheric differences in transport-induced trends appear contradictory to the hemispheric differences in trends in the residual mass fluxes (see Fig. 6). However, transport of ozone is determined not only by the strength of the mass fluxes, but also by the available background ozone, which is influenced by chemistry. The negative chemically-induced ozone trends below 30 to 40 hPa lead to less ozone being available for transport. Thus, the stronger negative chemically-induced trend in the SH than in the NH above 80 hPa might be the cause for the stronger positive transport-induced trend. Below 50 hPa, the sum of the attribution terms deviates from the actual ozone trend differences, making the attribution less reliable. As discussed in Sect. 7, the dominant role of chemically-induced changes is robust against the uncertainties introduced by the method.

Integrated over the lower stratospheric column, transport-induced trends are positive in both hemispheres, but stronger in the NH. Chemistry-induced trends, on the other hand, are positive in the NH but negative in the SH, and these large hemispheric differences lead to the overall stronger trend in the NH (see Fig. 9). These different trends are then reflected in the return dates, as summarized in Fig. 16. The causes for asymmetric ozone trends induced by chemistry in the lower stratosphere are investigated in Sect. 6.2.

6 Hemispheric differences in chemically-induced ozone trends

6.1 Lowermost stratosphere and troposphere: enhancement in production by NOx

In the LSTR and upper troposphere, strong positive trends in ozone are induced by changes in chemistry. Chemically-induced trends are stronger in the NH than in the SH. The contributions of individual reaction cycles to the chemically-induced increase in ozone are averaged between 300 and 250 hPa and are shown in Fig. 11. The primary contributor to ozone trends in this region, and the primary cause of hemispheric differences in ozone trends, is an increase in the production of ozone by NOx, in particular via the HO2 + NO cycle (the bars labeled “PHN” in Fig. 11), which is significantly stronger in the NH than in the SH. NOx concentrations increase in the troposphere due to stronger emissions, and the increase is stronger in the NH, affecting tropospheric ozone (Lelieveld and Dentener, 2000). Comparisons of NOx trends in the troposphere and LSTR between different CCMs showed clear discrepancies between individual models (not shown), most likely as a result of differences in the prescribed emissions (or concentrations) and differences in the way that tropospheric chemistry is simulated in the CCMs. In E39CA, NOx trends ∼ 200 hPa are stronger compared to other models. Therefore, the effect of enhanced production from the NOx cycles on ozone trends, as shown here, might be overestimated. Consistently, ozone trends below 150 hPa are stronger in E39CA compared to the multi-model mean (see Fig. 4). However, four of the models show tropospheric
Fig. 11. Contributions to the chemically-induced ozone trends (blue bars) in the NH (top) and SH (middle) mid-latitudes, and the hemispheric difference (bottom), integrated between 250 and 300 hPa in E39CA. Reaction cycles are the production via HO$_2$ + NO ("PHN") and CH$_3$O$_2$ + NO (PCN), and the HO$_x$ destruction cycle. The Chapman production cycles and the NO$_x$ and ClO$_x$ destruction cycles are omitted since their contributions are close to zero.

ozone trends close to zero (most likely models that neglect tropospheric chemistry), thus biasing the multi-model mean to lower values. In general, tropospheric chemistry is often treated in a simplified manner in CCMs, introducing uncertainty on the results shown here, as will be further discussed in Sect. 7.

6.2 Lower stratospheric chemistry

Changes in chemistry were shown to be the primary contributor to hemispheric differences in LSTR ozone trends. The chemically-induced trend profiles from Fig. 10 are further divided into individual reaction cycles in Fig. 12.

Below 30 to 40 hPa, ozone trends induced by changes in chemistry are negative, caused primarily by changes in rates of ozone destruction by the HO$_x$ cycles. The destruction rates of the HO$_x$ cycle are affected by increasing CH$_4$ concentrations, with CH$_4$ being the primary source of stratospheric HO$_x$ (Wuebbles and Hayhoe, 2002). Stratospheric water vapor is projected to increase in a future climate (Gettelman et al., 2010), leading to enhanced ozone destruction (Stenke and Grewe, 2005), consistent with the negative ozone trends induced by HO$_x$ in Fig. 12.

Changes in the rates of ozone destruction by the NO$_x$ cycles cause an increase in ozone throughout the lower stratosphere, dominating the total chemically-induced ozone trends above around 40 hPa. Changes in rates of ozone destruction by the NO$_x$ cycles were analyzed in detail, based on the same NIWA-SOCOL simulation as used here by Revell et al. (2012b). They showed that the decrease in ozone destruction by NO$_x$, despite an increase in N$_2$O, is caused by a decrease in the ratio of NO$_x$ to N$_2$O and a reduction
in the available atomic oxygen, both in turn largely due to GHG-induced stratospheric cooling.

The largest hemispheric differences in chemically-induced ozone trends occur at 50 hPa, and are primarily due to ozone destruction by the (i) NO\textsubscript{x} cycles and (ii) Cl\textsubscript{y} cycles (see lower panel in Fig. 12). Ozone destruction by the HO\textsubscript{x} cycles leads to stronger negative ozone trends around 100 hPa in the NH. Overall, HO\textsubscript{x} chemistry counteracts the effects of the NO\textsubscript{x} and Cl\textsubscript{y} cycles in the lower stratosphere (see Fig. 16) and plays a minor role for hemispheric asymmetries in ozone trends. The HO\textsubscript{x}-induced trends are thus not further discussed here. The increase in ozone caused by reduced ozone destruction by NO\textsubscript{x} is stronger in the NH compared to the SH. The hemispheric differences in ozone trends due to changes in the Cl\textsubscript{y} destruction cycles maximize at 70 hPa. While the changes caused by the Cl\textsubscript{y} destruction cycles are slightly positive in the NH, they are negative in the SH at 70 hPa. The drivers of the trends in the destruction rates of the Cl\textsubscript{y} and NO\textsubscript{x} cycles are investigated below.

6.2.1 Changes in NO\textsubscript{x} destruction rates

As discussed above, Revell et al. (2012b) showed that the effectiveness of N\textsubscript{2}O in depleting ozone is determined by NO\textsubscript{x} concentrations and by the availability in atomic oxygen. The former is strongly influenced by cooling of the stratosphere (Rosenfield and Douglass, 1998) and by changes in the strength of the BDC (Cook and Roscoe, 2012).

In Fig. 13, the vertical profile of linear trends in mid-latitude NO\textsubscript{x} over the period 1960 to 2049 are shown, calculated in the same manner as the ozone trends, i.e., using a regression model taking an offset, Cl\textsubscript{y} and a linear trend into account. Below 30 hPa, NO\textsubscript{x} mixing ratios decrease slightly in the NH, whereas trends in the SH are close to zero. Above 30 hPa, trends are positive in both hemispheres and increase with height. Despite this increase in NO\textsubscript{x} in both hemispheres above 30 hPa, ozone loss by the NO\textsubscript{x} cycle decreases in this region. As explained by Revell et al. (2012b), this results from a decrease in the availability of atomic oxygen, which in turn is due to decreasing temperatures and to competition with other ozone destruction cycles, in particular the HO\textsubscript{x} cycles.

The difference in the NO\textsubscript{x} trends between the hemispheres, shown in the right panel, closely matches the difference in the NO\textsubscript{x}-destruction-induced ozone trends shown in Fig. 12; the stronger increase in NH ozone compared to the SH at 50 hPa is consistent with a slight decrease of NO\textsubscript{x} in the NH and close to zero trend in NO\textsubscript{x} in the SH. At 20 hPa, the sign of the hemispheric difference in NO\textsubscript{x} trends is reversed (but uncertainties are large and trends are not statistically different), which is reflected in a slightly negative hemispheric difference in NO\textsubscript{x}-destruction-induced ozone trends (as shown in Fig. 12). At 10 hPa the NO\textsubscript{x} trend in the SH is significantly stronger compared to the NH. This hemispheric difference in NO\textsubscript{x} trends increases further up to 5 hPa (not shown). The stronger positive ozone trend, due to reduced NO\textsubscript{x}-induced ozone destruction changes in the NH compared to the SH at 10 hPa (see Fig. 12), can again be explained by the asymmetric NO\textsubscript{x} trends. Overall, the hemispheric differences in the trends in ozone destruction rates by the NO\textsubscript{x} cycle are consistent with hemispheric differences in NO\textsubscript{x} trends.

6.2.2 Changes in Cl\textsubscript{y} destruction rates

In the LSTR, the efficiency of homogenous Cl\textsubscript{y} ozone destruction cycles is expected to decrease with decreasing temperatures (Rosenfield et al., 2002). As shown in Fig. 12, ozone trends due to changes in destruction rates by the Cl\textsubscript{y} cycles are slightly positive throughout the lower stratosphere in the NH, consistent with decreasing temperatures. In the SH, on the other hand, a reduction in ozone resulting from an increase in the Cl\textsubscript{y} destruction rates is found at 70 hPa.

The increase in ozone destruction rates cannot be due to changes in the abundance of Cl\textsubscript{y}, since any changes in ozone congruent with Cl\textsubscript{y} changes are included in the Cl\textsubscript{y} regression coefficient. Therefore, the trend in ozone destruction rates must be caused by changes in the efficiency of a given amount of Cl\textsubscript{y} to destroy ozone. The efficiency is determined by temperature, with lower temperatures slowing homogenous reaction cycles. However, at very low temperatures, as those occurring in the Antarctic vortex, ozone can be depleted efficiently by heterogeneous chemistry. Hence, any expansion in space or time of the region where the conditions for heterogeneous chemistry occur can cause an increase in the efficiency of the Cl\textsubscript{y} ozone destruction cycles.

Trends in the rates of ozone destruction rates by the Cl\textsubscript{y} cycles in southern mid-latitudes at 70 hPa are shown as a
function of month in Fig. 14. Destruction rates increase (i.e., larger negative values) in midwinter, and the trend maximizes in July. When disaggregating the destruction rates into individual reaction cycles, it can be seen that the largest contributor to the destruction rate increase in midwinter is the ClO-dimer reaction cycle. Therefore, it is mostly heterogeneous chemistry that is responsible for the increased efficiency of ozone destruction. In midwinter, the polar night prohibits heterogeneous ozone depletion south of 66.5° S for varying periods since sunlight is required for the photochemical reactions. Figure 15 shows the trend in temperatures over the southern middle to high latitudes at 70 hPa in July, where the increase in heterogeneous ozone depletion maximizes. Temperature trends are negative over the polar region and maximize ∼ 60° S and 150° W. The pattern is suggestive of an intensification and shift of the polar vortex center away from the pole. Through this process, polar vortex air with low temperatures reaches latitudes equatorward of 60° S, where sunlight is available in midwinter, allowing for heterogeneous ozone depletion. This process can explain the negative trend in ozone at southern mid-latitudes at 70 hPa in NIWA-SOCOL.

7 Discussion

The implications of and uncertainties in the results presented in the last sections will be discussed in the following. The results obtained with the attribution analysis (Sects. 5 and 6) are subject to two kinds of uncertainties: (1) uncertainties in the attribution method itself, and (2) uncertainties in the model used for the attribution analysis. The latter refers to the question of whether we can expect the processes identified in E39CA and NIWA-SOCOL to be of similar importance in other CCMs. To answer this question, the attribution analysis would have to be applied to other models as well, which requires knowledge of the chemical loss and production rates. However, the comparison of NIWA-SOCOL and E39CA to the ensemble analysis of CCMs presented in the first half of the paper and to earlier studies allows us some estimations.

7.1 What is the role of transport for hemispheric asymmetric ozone trends?

Both the analysis of the relation of hemispheric differences in mid-latitude ozone return dates to asymmetric trends in the strength of the BDC and the attribution analysis to chemical and transport changes showed evidence that changes in transport of ozone plays a smaller role in explaining hemispherically asymmetric return dates than previously thought. In the lower stratosphere, no significant correlation of ozone return date differences to asymmetric trends in the BDC strength is found across a set of twelve models. The attribution of ozone trends to transport and chemistry changes applied to NIWA-SOCOL showed that transport changes induce stronger positive ozone trends in the NH than SH at levels around 100 hPa. However, above 80 hPa, transport-induced ozone trends are stronger in the SH than in the NH.
Fig. 16. Summary of the results: hemispheric differences in mid-latitude return dates to 1980 values are about 13 yr in the multi-model mean (MMM) for TOZ. Upper stratospheric ozone contributed negatively to this difference, while return dates of lower stratospheric ozone (100 to 10 hPa) alone have a hemispheric difference of about 10 yr, and ozone below 100 hPa contributes another 5 yr. In the LSTR, in NIWA-SOCOL changes in chemistry only would lead to an earlier return in the NH of close to 20 yr, and changes in transport of ozone offset the chemical effects. Attribution of chemically-induced trends in the LSTR show that the earlier return of NH ozone is due to changes in the NOx and Cly cycles. In the lowermost stratosphere and troposphere, hemispheric differences modeled in E39CA, somewhat above-average, are attributed to changes in chemistry, and transport offsets the chemically-induced changes. Enhanced production of ozone in the NH by the NOx cycles is found to account for the stronger positive trend there.

Figure 16 shows how the lower stratospheric ozone trends translate to return dates. The positive transport-induced trends shift return dates forward in both hemispheres, but more so in the SH. The positive chemically-induced trend in the NH shifts the return date even further forward, while the chemically-induced trend is negative in the SH, leading to a later return date here. Thus, overall, chemically-induced trends are the major cause of the hemispheric differences in ozone return dates.

However, to obtain meaningful results with the attribution method, the changes in annual mean ozone tendencies needs to be small (i.e., the “imbalance” term negligible), so that the chemically- and transport-induced changes together with the nonlinear contribution equal the total change in ozone. As seen in Fig. 10, this is not the case below 50 hPa in the Southern Hemisphere, resulting in a discrepancy also in the hemispheric difference. These are the levels where transport-induced trends are of major importance; thus could we underestimate transport effects due to problems with the method in exactly the regions where transport seems to matter most? Let us assume that the difference between the estimated and actual ozone changes (i.e., the grey and black solid line in Fig. 10) would be entirely due to transport changes. Thus, we increase the hemispheric differences in transport-induced trends so that the estimated and actual ozone trends match. Then, the overall effect of transport changes on hemispheric differences in ozone trends in the lower stratospheric column would be slightly positive instead of negative. However, chemically-induced asymmetry in return dates in the lower stratospheric column would still dominate over the transport-induced asymmetry by a factor of 10. Thus, the conclusion that chemistry is the dominant factor in driving hemispheric asymmetries in lower stratospheric ozone trends is insensitive to the uncertainties in the attribution method.

Concerning uncertainties stemming from the choice of the model, NIWA-SOCOL simulates increases in the strength of the mass fluxes that are above average (Fig. 6). The hemispheric differences in the 70 hPa mass fluxes are slightly larger compared to most models (Fig. 7). Thus, if transport were to play an important role in driving asymmetric ozone trends, it should be apparent in the NIWA-SOCOL model. In general, current CCMs are run with a rather coarse resolution due to restrictions in computational costs. Transport might be dependent on the model resolution and the chosen advection scheme, and in particular the role of smaller-scaled transport processes in the tropopause region might not be
captured properly. How strongly transport of ozone is scale-dependent, and whether this has any implications for the simulated trends, is an open question future studies will have to address.

In the lowermost stratosphere, we found a weak correlation between the hemispheric differences in the BDC changes at 100 hPa and the differences in return dates for the ensemble of twelve CCMs. A closer investigation with E39CA showed that the positive trends in this region are in large part caused by chemically-induced changes, which are stronger in the NH and thus introduce hemispheric differences. The attribution method works reasonably well in this region for E39CA, as seen from the approximate agreement of the ozone changes and control line (black dashed) in Fig. 8. The model E39CA simulates stronger than average strengthening of the downward mass flux at 100 hPa (Fig. 6), but hemispheric differences here are small (Fig. 7). A group of four models simulate stronger hemispheric differences in the mass flux trends and larger asymmetries in the return dates. Thus, for this group of models, transport changes might play a larger role. Another group of five models simulates two to five years earlier return dates of LMSTR ozone in the NH, but stronger mass flux increases in the SH than to the NH. For this group of models, the correlation between return date difference and asymmetric circulation changes apparently breaks down. Thus, it appears likely that at least in this group of models, chemically-induced changes play the leading role in introducing the hemispheric differences.

Overall the results suggest that the hemispheric differences in ozone trends, and thus return dates in the lowermost stratosphere are caused by a combination of (i) enhanced ozone production by increasing NO\textsubscript{x} concentrations, which is stronger in the NH, and (ii) hemispheric different transport changes due to the hemispheric asymmetric BDC trends. The relative importance of the mechanisms probably varies from model to model.

7.2 What is the relevance of the identified chemical processes?

7.2.1 Lowermost stratosphere and troposphere: NO\textsubscript{x}-related production

In the lowermost stratosphere and upper troposphere, analysis of the E39CA model reveals an important contribution to positive ozone trends from ozone production by the NO\textsubscript{x} production cycles. Due to stronger increases in NO\textsubscript{x} concentrations in the NH, the production increase is larger there compared to the SH, and thus leads to asymmetric ozone trends. Even though the absolute amount of NO\textsubscript{x} seems to be simulated well in E39CA (as indicated by a brief comparison to a tropospheric chemistry model and to observations, not shown), the trends are larger compared to other CCMs. Thus, the effect of NO\textsubscript{x}-related ozone production might be less relevant in other CCMs. In addition, despite E39CA having been used to examine variability in tropospheric ozone, the model incorporates only background tropospheric chemistry, which limits the capabilities to accurately simulate trends (Grewe, 2007). The increase in the tropospheric ozone burden from 2000 to 2030 of around 10\% in E39CA compares reasonably well with the estimates of an ensemble of models focused on tropospheric chemistry of 6 to 15\%, depending on the scenario, as given by Stevenson et al. (2006). However, more recent results indicate that tropospheric ozone is more likely to decrease until 2030 under a moderate emission scenario (Young et al., 2013). Observations also indicate that the positive trends, in particular in NH tropospheric ozone from 1970 to about 1990, flattened or even reversed over the last 10–15 yr (Oltmans et al., 2013). Thus, the positive ozone trends in the troposphere simulated by most CCMs used in this study over the period 1960 to 2049 might be unrealistic. Tropospheric ozone trends are found to be strongly coupled to trends in NO\textsubscript{x} also in models with more sophisticated tropospheric chemistry schemes (Young et al., 2013). It remains unclear how much tropospheric ozone trends in CCMs are affected by the prescribed NO\textsubscript{x} emissions (or concentrations), or by simplified tropospheric chemistry schemes. Projections with CCMs with improved tropospheric chemistry schemes will need to be analyzed to conclude with more certainty on the role tropospheric ozone trends play for return dates in total column ozone.

7.2.2 Lower stratosphere: NO\textsubscript{x} destruction cycles

The hemispheric differences in the decrease of ozone destruction by the NO\textsubscript{x} cycles are found to be controlled by hemispheric differences in the trends in NO\textsubscript{x} concentrations. The detailed analysis of the causes for the hemispheric differences in NO\textsubscript{x} trends are beyond the scope of this study. From previous studies, it is known that trends in stratospheric NO\textsubscript{x} concentrations are influenced by the speed of the BDC (Plummer et al., 2010; Cook and Roscoe, 2012). Thus, one possible explanation of the hemispheric differences in NO\textsubscript{x} trends could be a feedback of changes in the circulation on the efficiency of the production of NO\textsubscript{x}. As shown by Cook and Roscoe (2012) in a simplified model approach, an increase in the BDC causes a decrease in the formation of NO\textsubscript{x} (since the time available for photolysis decreases, and this effect dominates over the enhanced influx of the tropospheric source gas N\textsubscript{2}O). Climatologically, the BDC is stronger in the NH compared to the SH, and in agreement with the work of Cook and Roscoe (2012), climatological NO\textsubscript{x} concentrations in NIWA-SOCOL are higher in the SH than in the NH (not shown). A stronger increase in the strength of the BDC in the middle stratosphere of the NH could thus explain weaker positive NO\textsubscript{x} trends in the NH compared to the SH.

Compared to other models, trends in NO\textsubscript{x} concentrations at 10 hPa in NIWA-SOCOL are of similar magnitude. In addition, most models show a stronger NO\textsubscript{x} trend in the SH.
than in the NH (not shown), indicating that the hemispheric asymmetric changes in NO\textsubscript{x} concentrations can be a relevant factor in determining the hemispheric different ozone trends and thus return dates.

As opposed to most other models, E39CA simulates a slightly earlier return of lower stratospheric ozone in the SH than in the NH (see Fig. 7), and ozone trends are negative above 30 hPa in contrast to the remaining group of CCMs analyzed here (see Fig. 4). The attribution analysis in the lower stratosphere for E39CA (not shown) reveals that the chemically-induced ozone trend, and in particular the NO\textsubscript{x} destruction cycles, are responsible for the decrease in ozone above 30 hPa in E39CA. Positive trends in NO\textsubscript{x} in the lower stratosphere are about 4 times stronger than in NIWA-SOCOL, explaining the strong ozone decrease induced by the NO\textsubscript{x} destruction cycles. Furthermore, the NO\textsubscript{x} trends are identical in the two hemispheres in E39CA, and consequently also the hemispheric differences in chemically-induced ozone trends are much smaller compared to NIWA-SOCOL. Since in E39CA, NO\textsubscript{x} (\(=\) NO\textsubscript{y} + HNO\textsubscript{3}) concentrations are prescribed at the upper level at 10 hPa, and concentrations are assumed to increase at the same rate as N\textsubscript{2}O, any influence of changes in temperature and dynamics on NO\textsubscript{x} production is not accounted for. Thus, the impact of dynamical and temperature changes on NO\textsubscript{x} concentrations can be concluded to be crucial for both the response of ozone to N\textsubscript{2}O changes, in agreement with earlier studies (Plummer et al., 2010; Revell et al., 2012b), and the hemispheric asymmetry of ozone trends, as the results of this study suggest.

### 7.2.3 Lower stratosphere: Cl\textsubscript{y} destruction cycles

Changes in the efficiency of Cl\textsubscript{y} to deplete ozone is found to be one of the causes for hemispheric asymmetries in ozone trends. Enhanced efficiencies are found in the SH, where an expansion or shift of the polar vortex provides the conditions for heterogeneous ozone depletion in midwinter north of 60° S. This effect was noted earlier for southern polar ozone, for example by Plummer et al. (2010), who found a nonlinearity of the response of ozone to ODSs in the Antarctic lower stratosphere under changing GHG concentrations. Similarly, Revell et al. (2012a) reported an increase of Antarctic lower stratospheric ozone in a scenario with lower GHG concentration increases as compared to the standard REF-B2 simulation. The results presented here suggest that the effect of GHG-induced cooling on homogenous ozone depletion is of importance in particular for the evolution of ozone in the mid-latitudes.

The trend analysis of the ensemble of CCMs reveals negative trends in SH mid-latitude ozone around 70 to 50 hPa in about half of the models, indicating that the effect of an increasing efficiency of destruction by Cl\textsubscript{y} chemistry is of importance not only in NIWA-SOCOL. The strength of this effect will depend strongly on the pattern of dynamical changes, in particular in the location and timing of the Antarctic vortex, simulated in each individual model. Thus, the variety of the simulated changes in the polar vortex likely accounts at least in part for the spread in the ozone trends and thus ozone return dates simulated by the different models.

### 8 Conclusions

The detailed investigation of mid-latitude return dates of TOZ to 1980 values from an ensemble of twelve CCMVal models, with particular focus on the hemispheric differences, confirmed recent studies (e.g. WMO, 2011) in that the earlier return date in the northern mid-latitudes compared to the southern mid-latitudes is a robust result. However, we found that there are large differences between individual models, with hemispheric return date differences ranging from about 0 to 30 yr for TOZ. The method used to estimate the return dates can cause deviations of the results for the hemispheric difference in return dates of up to 5 to 10 yr for individual models. Although results for individual models slightly depend on the applied method, the multi-model mean is barely affected by the method used.

Return dates in both hemispheres occur earlier than would be expected from the return of Cl\textsubscript{y} to 1980 levels. The forward shift is caused by positive linear trends in ozone. This forward shift is larger in the NH than in the SH because (i) the effect of a certain linear trend on the return date depends on the sensitivity of ozone to Cl\textsubscript{y}, and (ii) the trend in ozone is stronger in the NH than in the SH.

As summarized in Fig. 16, the hemispheric differences in total column return dates are due to differences in the evolution of ozone in the lower stratosphere and lowermost stratosphere/troposphere. Based on the models NIWA-SOCOL and E39CA, processes driving hemispheric differences in return dates are identified. The key result of this study is that changes in chemical production and loss play the major role in driving hemispheric differences in ozone return dates. The statement in WMO (2011) that the “more pronounced strengthening of the poleward transport of ozone” is the cause of the earlier return of ozone in the NH could not be confirmed in the present study. The chemically-induced changes identified in the lower stratosphere are, however, likely indirectly influenced by asymmetric BDC changes, which influence transport of ozone precursors – the rate at which these are produced by photolysis and temperatures that affect reaction rates. In particular, the main chemical processes identified here are the relatively less efficient destruction of ozone by NO\textsubscript{x} chemistry in the NH, caused by likely dynamical influences on NO\textsubscript{x} concentrations, and the relatively more efficient destruction of ozone by Cl\textsubscript{y} in the SH due to an expansion of the polar vortex into mid-latitudes in midwinter.

Furthermore, ozone trends in the lowermost stratosphere and troposphere are found to contribute significantly to the hemispheric asymmetry in TOZ return dates. Enhanced
production of ozone due to increasing NOx emissions are identified to introduce the hemispheric asymmetries in ozone trends. Since tropospheric chemistry is treated in a simplified manner in the CCMs used here, studies with improved representations of tropospheric chemistry will be necessary to validate this result. Overall it can be concluded that the uncertainties introduced by the treatment of tropospheric chemistry and ozone precursor emissions are not negligible for the correct simulation of the temporal development of TOZ.

Appendix A

Methods of return date estimation

To test the sensitivity of ozone return dates to the method used for their estimation, three different methods are intercompared:

1. Simple regression model
A linear least squares regression of the form
\[
O_3(t) = a_0 + a_1 \times Cl_y(t) + a_2 \times t + R(t)
\]
is applied to the annual mean ozone time series from \( t = 1960 \) to 2049. Cl\(_y\) mixing ratios at 50 hPa and at the same latitude as the ozone time series are used as the basis function. \( a_0, a_1 \) and \( a_2 \) are the offset, the Cl\(_y\) and the trend regression coefficients, respectively, and \( R \) is the residual. The uncertainties on the regression coefficients are calculated taking auto-correlation into account.

2. Regression model with quadratic terms
Again a linear regression model is used, but including Cl\(_y\) and trend terms that are quadratic in Cl\(_y\) and \( t \):
\[
O_3(t) = a_0 + a_1 \times Cl_y(t) + a_2 \times Cl_y(t)^2 + a_3 \times t + a_4 \times t^2 + R(t).
\]
These additional terms allow for more degrees of freedom in the fitted time series, but a physical interpretation of the attributed variance is no longer straightforward.

3. Smoothing with a 1 : 2 : 1 filter
A 1 : 2 : 1 filter is applied iteratively to the ozone time series 20 times. The number of iterations of applying the filter is chosen as to obtain a smooth time series from which variability in the interannual to decadal time scale is removed but the long-term signal is maintained.

For methods 1 and 2, uncertainties on the return dates are calculated based on the uncertainty on the regression fit. The regression model uncertainties are estimated using a bootstrap procedure: first, the regression model is run on the original time series, obtaining the smooth regression fit and the residuals (original data minus fit). Then, a set of 10 000 perturbed time series are constructed by adding noise to the regression fit. The added noise is generated by randomly selecting, with replacement, from the residuals. The regression model is fitted to each of the 10 000 time series, and for each the return date is estimated. The error bars shown in Fig. 1 are then one standard deviation over these 10 000 return date estimates.

Differences in estimated multi-model mean return dates between the three methods are significantly smaller than the return date differences between models (Fig. 1). The simple regression (method 1) yields a multi-model mean return date in the NH ∼ 3 yr earlier than the other two methods, but otherwise the agreement is good. For each individual model, however, the differences in return dates can be 5 to 10 yr, depending on the method used. Our results and those of SPARC-CCMVal (2010) (see their Fig. 9, p. 49) suggest that using different methods for deriving return dates will not bias the multi-model results, but only return date estimates for individual models. This suggests that the differences in the estimates from various methods are caused by the large amount of variability in the individual model time series, adding ambiguity to the retrieval of the signal.

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References


