Estimates of Ozone Return Dates from Chemistry-Climate Model Initiative Simulations

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Abstract. We analyse simulations performed for the Chemistry-Climate Model Initiative (CCMI) to estimate the return dates of the stratospheric ozone layer from depletion caused by anthropogenic stratospheric chlorine and bromine. We consider a total of 155 simulations from 20 models, including a range of sensitivity studies which examine the impact of climate change on

- 5 ozone recovery. For the control simulations (unconstrained by nudging towards analysed meteorology) there is a large spread (± 20 DU in the global average) in the predictions of the absolute ozone column. Therefore, the model results need to be adjusted for biases against historical data. Also, the interannual variability in the model results need to be smoothed in order to provide a reasonably narrow estimate of the range of ozone return dates. Consistent with
- 10 previous studies, but here for a Representative Concentration Pathway (RCP) of 6.0, these new CCMI simulations project that global total column ozone will return to 1980 values in 2049 (with a 1- σ uncertainty of 2043-2055). At Southern Hemisphere mid-latitudes column ozone is projected to return to 1980 values in 2045 (2039-2050), and at Northern Hemisphere mid-latitudes in 2032 (2020-2044). In the polar regions, the return dates are 2060 (2055-2066) in the Antarctic in
- 15 October and 2034 (2025-2043) in the Arctic in March. The earlier return dates in the NH reflect the larger sensitivity to dynamical changes. Our estimates of return dates are later than those presented in the 2014 Ozone Assessment by approximately 5-17 years, depending on the region, with the previous best estimates often falling outside of our uncertainty range. In the tropics only around half the models predict a return of ozone to 1980 values, at around 2040, while the other
- 20 half do not reach the 1980 value. All models show a negative trend in tropical total column ozone towards the end of the 21st century. The CCMI models generally agree in their simulation of the time evolution of stratospheric chlorine and bromine, which are the main drivers of ozone loss and recovery. However, there are a few outliers which show that the multi-model mean results for ozone recovery are not as tightly constrained as possible. Throughout the stratosphere the spread
- 25 of ozone return dates to 1980 values between models tends to correlate with the spread of the return of inorganic chlorine to 1980 values. In the upper stratosphere, greenhouse gas-induced cooling speeds up the return by about 10-20 years. In the lower stratosphere, and for the column, there is a more direct link in the timing of the return dates of ozone and chlorine, especially for the large Antarctic depletion. Comparisons of total column ozone between the models is affected
- 30 by different predictions of the evolution of tropospheric ozone within the same scenario, presumably due to differing treatment of tropospheric chemistry. Therefore, for many scenarios, clear conclusions can only be drawn for stratospheric ozone columns rather than the total column. As noted by previous studies, the timing of ozone recovery is affected by the evolution of N₂O and CH₄. However, quantifying the effect in the simulations analysed here is limited by the few
- 35 realisations available for these experiments compared to internal model variability. The large increase in N₂O given in RCP 6.0 extends the ozone return globally by ~15 years relative to N₂O fixed at 1960 abundances, mainly because it allows tropical column ozone to be depleted. The effect in extratropical latitudes is much smaller. The large increase in CH₄ given in the RCP 8.5 scenario compared to RCP 6.0 also lengthens ozone return by ~15 years, again mainly through its
- 40 impact in the tropics. Overall, our estimates of ozone return dates are uncertain due to both uncertainties in future scenarios, in particular of greenhouse gases, and uncertainties in models. The scenario uncertainty is small in the short term but increases with time, and becomes large by the end of the century. There are still some model-model differences related to well-known processes which affect ozone recovery. Efforts need to continue to ensure that models used for
- 45 assessment purposes accurately represent stratospheric chemistry and the prescribed scenarios of ozone-depleting substances, and only those models are used to calculate return dates. For future assessments of single forcing or combined effects of CO₂, CH₄, and N₂O on the stratospheric column ozone return dates, this work suggests that is more important to have multi-member (at least 3) ensembles for each scenario from every established participating model, rather than a large
- 50 number of individual models.

1) Introduction

It is well established that stratospheric ozone depletion over the past few decades has been mostly driven by the increase in stratospheric chlorine and bromine originating from the use of

- 5 halogenated ozone-depleting substances (ODSs). The production of these ODSs has now been largely controlled by the 1987 Montreal Protocol on Substances that Deplete the Ozone Layer, and its subsequent amendments and adjustments. For this reason, the protocol has been recognised as one of the most successful international environmental treaties ever enacted. Following its implementation, the atmospheric burdens of many ODSs have peaked and are now declining at a
- 10 rate dependent on their respective atmospheric lifetimes, which are typically many decades. This has led to the overall levels of tropospheric chlorine and bromine peaking in 1993 and 1998, respectively (WMO, 2014). These levels are now slowly decreasing and are expected to return to their 1980 values, an arbitrary reference date before the discovery of the Antarctic ozone hole, around the middle of this century. Accordingly, stratospheric ozone is expected to recover from
- 15 the effects of halogen-induced decreases on a similar timescale, which is already detectable in the Antarctic and upper stratosphere (e.g. see *Solomon et al.*, 2016; *Chipperfield et al.*, 2017; *Strahan and Douglass*, 2018 and references therein). However, other atmospheric changes, notably climate change through increasing levels of greenhouse gases (GHGs) are also expected to modify the rate of return of ozone and its subsequent evolution (e.g., *Shepherd and Jonsson*, 2008; *Eyring et al.*,
- 20 2010a). The effect of this climate impact is likely to be different in various latitudinal and altitudinal regions of the stratosphere and, for the polar regions, depends on the dynamical evolution of the polar vortices.

The prediction of ozone recovery and return therefore requires the use of three-dimensional (3D) coupled chemistry-climate models (CCMs) which contain details of the important stratospheric chemical processes, as well as a realistic, interactive representation of stratospheric temperature and dynamics (*Morgenstern et al.*, 2010, 2017 and references therein). In these models, the simulated composition of the atmosphere is fully interactive, wherein the radiatively active gases, e.g., carbon dioxide (CO₂), water vapour (H₂O), nitrous oxide (N₂O), methane (CH₄),

- 30 chlorofluorocarbons (CFCs), and ozone (O₃) affect model heating and cooling rates and therefore dynamics. The representation of stratospheric chemistry includes species and chemical reactions contained in the odd oxygen (O_x), nitrogen (NO_x), hydrogen (HO_x), chlorine (ClO_x), and bromine (BrO_x) chemical families as recommended, for example, by the NASA Chemical Kinetics and Photochemical Data for Use in Atmospheric Studies evaluation reports (e.g., *Sander et al.*, 2011).
- 35 These chemical recommendations also include heterogeneous processes on sulfate aerosols and polar stratospheric clouds (PSCs). In addition, many of the CCMs include a detailed representation of tropospheric chemistry.
- A process-oriented evaluation of CCMs was performed as part of the Stratospheric Processes And 40 their Role in Climate (SPARC) Chemistry Climate Model Validation Activity (CCMVal). Quantitative performance metrics (*Eyring et al.*, 2008; *Waugh and Eyring* 2008) were used in an extensive effort to evaluate the ability of CCMs to represent key processes related to dynamics, transport, chemistry, and climate. These CCMs have been used in support of several international assessments of ozone depletion and recovery (e.g., *Austin et al.*, 2003, 2010; WMO/UNEP 2003,
- 45 2007, 2011, 2014; *Eyring, et al.*, 2007, 2010a, b, 2013b). The models used in this current paper have benefitted from the testing and development that came out of these studies and many are direct updates of the models tested in CCMVal.

Radiative, dynamical and chemical processes affect ozone recovery and return date. The radiative effects of changing GHG concentrations and ozone-induced temperature changes can affect the wave driving of the atmosphere and subsequently can accelerate the Brewer-Dobson (*Brewer*, 1949) (BD) circulation (*Rind et al.*, 1990, 2001; *Butchart and Scaife*, 2001; *Sigmond et al.*, 2004,

- 5 *Eichelberger and Hartmann*, 2005; *Butchart et al.*, 2006, 2010; *Olsen et al.*, 2007; *Garcia and Randel*, 2008; *McLandress et al.*, 2010, *Polvani et al.*, 2017, 2018). These studies have shown that the acceleration of the BD circulation is a robust result across the majority of CCMs. This acceleration can affect ozone recovery by causing faster removal of ODSs and hence earlier ozone recovery, although decreases in tropical stratospheric ozone may result in column ozone remaining
- 10 permanently below the 1980 value at these latitudes. *Butchart and Scaife* (2001) estimated that the recovery of the global ozone layer could be brought forward by 8-10 years. This shortening of the ozone recovery was also found by *Morgenstern et al.* (2018) for the models represented in this study, although it is important to note that the use of surface mixing ratios in studies largely removes the feedback between circulation changes and ODS return dates. In addition, this
- 15 strengthening of the BD circulation leads to a decrease in the mean age of stratospheric air by about 0.05 years per decade (*Butchart et al.*, 2010).

Increases in GHGs lead to cooling of the upper stratosphere (e.g., *Fels et al.*, 1980; *Rind et al.*, 1990) which slows down temperature-dependent odd-oxygen loss processes and increases upper stratospheric ozone (e.g., *Haigh and Pyle*, 1982; *Brasseur and Hitchman*, 1988; *Pitari et al.*, 1992;

Rosenfeld et al., 2002; Waugh et al., 2009; Oman et al., 2010; Bekki et al., 2013; Marsh et al., 2016). Evolution of N₂O and CH₄ can also impact ozone recovery by both chemistry and climate processes (Randeniya et al., 1997; Chipperfield and Feng, 2003; Ravishankara et al., 2009; McLandress et al., 2010; Revell et al., 2012; Morgenstern et al., 2014; Revell et al., 2015; Kirner et al., 2015).

A number of studies have used detailed coupled CCMs to predict the future evolution of stratospheric column ozone. For example, using the GFDL model, *Austin and Wilson* (2006) showed that the October monthly mean Antarctic ozone return to 1980 abundances mainly depends on halogen loading and will not occur until ~2065. They also showed that Arctic ozone returns about 25-35 years earlier than Antarctic ozone does, while *Li et al.* (2009) showed that in the tropics ozone may never return to 1980 values due to changes in transport. However, due to various biases and weaknesses amongst various CCMs, our best estimates of ozone return dates come from multi-model assessments such as CCMI.

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The aim of this paper is to provide updated estimates of the return dates of stratospheric ozone in different latitude regions. The year at which column ozone is expected to return to a particular, historic level is a key metric for quantifying the recovery of the ozone layer (e.g., Table 2-5, WMO 2014). Note that we estimate return dates based on mean atmospheric behaviour, independent of

- 40 the extremes caused by dynamically driven variability, for example. Our estimates of return dates are based on analysis of the new, extensive range of CCMI simulations which have been produced using updated and improved models compared to previous studies such as CCMVal-2. All models compute the impact on ozone of future stratospheric cooling, the intensification of the Brewer-Dobson circulation, as well as numerous other dynamical and chemical factors. Three types of
- 45 uncertainties are considered: internal variability, model uncertainty, and scenario uncertainty. Internal variability refers to short-term variations in computed ozone, which are not dependent on long-term change driven by ODSs and GHGs. Model uncertainty represents the fact that various CCMs can provide different return dates, for the same prescribed time series of ODSs and GHGs, due to alternate representations of processes such as the impact of rising GHGs on stratospheric
- 50 circulation. Finally, scenario uncertainty means that the return date found by a CCM will be sensitive to the particular, prescribed time series of ODSs and GHGs used as model input.

The layout of the paper is as follows. Section 2 briefly summarises the models, scenarios, and simulations used in this study. Section 3 describes how the results of the CCM simulations are processed in order to smooth over interannual variability and adjust some model ozone values for

5 biases compared to observations. **Section 4** presents our main results on the return and recovery of stratospheric ozone and how it varies between different simulations and between models. Finally, **Section 5** presents our conclusions.

2) Models, Scenarios, and Simulations

- 10 The model simulations used in this work are taken from the Chemistry–Climate Model Initiative (CCMI), which is a joint activity from the International Global Atmospheric Chemistry (IGAC) project and SPARC. One of the main goals of CCMI is to provide support for the quadrennial Scientific Assessment of Ozone Depletion, produced by the World Meteorological Organisation (WMO) Global Ozone Research and Monitoring Project and United Nations Environment
- 15 Programme (UNEP). This is done through the provision of scenarios and input forcings that the models can use for standard experiments. This paper provides analysis to support the results that will be presented in the forthcoming 2018 assessment.
- Descriptions of the CCMI experiments used in this work are given in **Table 1**. These scenarios are discussed in detail in *Eyring et al.* (2013a) and *Morgenstern et al.* (2017). **Table 2** summarises the simulations performed by each model considered here. Briefly, there are three reference simulations designed to understand both past and future ozone evolution. The first (labelled REF-C1) is a hindcast simulation of the recent past [1960-2010] that is closely tied to the following observed time-dependent forcings: 1) greenhouse gases (GHGs) and ozone-depleting substances
- 25 (ODSs); 2) 11-year solar variability; 3) sulfate aerosol surface area density (including background and volcanically active periods); 4) sea surface temperatures (SSTs) and sea ice concentrations (SICs); 5) additional organic bromine from very short-lived substances (VSLS); and 6) tropospheric ozone and aerosol precursor emissions. The meteorology in REF-C1 is free-running. The second reference simulation (labelled REF-C1SD) has the same observed forcings as REF-
- 30 C1, with the additional constraint that model temperature and dynamics are nudged to analysed meteorology, i.e., specified dynamics (SD). The third reference scenario (labelled REF-C2) includes both a hindcast and forecast period [1960-2100]. It should be noted that for REF-C2, several of the models used in this study have an interactive ocean/sea ice modules (*Morgenstern et al.*, 2018). For this scenario, the hindcast forcings are similar to REF-C1 with the main exception
- 35 that the SSTs/SICs are based on model results (*Morgenstern et al.*, 2018). The forecast component of the REF-C2 scenario uses GHGs (i.e., CO₂, CH₄, and N₂O) that follow the Intergovernmental Panel on Climate Change (IPCC) Coupled Model Intercomparison Project Phase 5 (CMIP5) Representative Concentration Pathways 6.0 (RCP 6.0) scenario (*Meinshausen et al.*, 2011; *Masui et al.*, 2011).
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Table 2 shows the number of realisations that were available for each simulation from every model. For REF-C1 a total of 19 models performed 38 realisations, including 8 models which performed multi-member ensembles. For simulations REF-C1SD and REF-C2 the numbers are 13 models (13 realisations) and 19 models (33 realisations), respectively. In addition to these

45 reference simulations, eight additional sensitivity scenarios based on the REF-C2 simulation were defined by CCMI. SEN-C2-RCP26 (5 models, 5 realisations), SEN-C2-RCP45 (7, 9) and SEN-C2-RCP85 (8, 10) follow RCP 2.6, 4.5 and 8.5, respectively. These scenarios diverge from the REF-C2 definition in year 2000. SEN-C2-fODS (8, 12) and SEN-C2-fGHG (8, 13) are identical to REF-C2, except that concentrations of ODSs and GHGs, respectively, are fixed at 1960 levels. Finally, there are three scenarios that examine the sensitivity of ozone return to N_2O and CH_4 . Scenario SEN-C2-fN2O [1960-2100] is the same as REF-C2 but with surface N_2O mixing ratios fixed to 1960 values. This results in the N_2O surface abundance being ~40% higher in 2100 for the REF-C2 scenario versus the SEN-C2-fN2O scenario. Scenario SEN-C2-fCH4 [1960-2100] is

- 5 the same as REF-C2 but with surface CH₄ abundance fixed to 1960 values; the late 21st century REF-C2 surface abundance of CH₄ is 30-50% higher than in SEN-C2-fCH4. Scenario SEN-C2-CH4RCP85 [2000-2100] replaces the REF-C2 RCP 6.0 CH₄ surface abundance with that from RCP8.5. Since the RCP8.5 surface CH₄ is considerably larger than in REF-C2, there is 110-125% more CH₄ in SEN-C2-CH4RCP85 relative to REF-C2 in the late 21st century. There are 8 (8), 8
- 10 (8), 6 (6) CCMs (realisations) included for SEN-C2-fN2O, SEN-C2-fCH4 and SEN-C2-CH4RCP85, respectively. For scenarios SEN-C2-RCP26 and SEN-C2-CH4RCP85, in particular, the numbers of realisations available are limited.

3) Methodology

- 15 We present an analysis based on seven latitude bands: Southern Hemisphere (SH) polar (90°S-60°S), SH mid-latitudes (60°S-35°S), tropics (20°S-20°N), Northern Hemisphere (NH) midlatitudes (35°N-60°N), NH polar (60°N-90°N), near-global (60°S-60°N) and global (90°S-90°N). For the SH and NH polar regions we only consider the months of October and March, respectively, which typically correspond to the end of the winter/spring ozone loss periods. For the other latitude
- 20 bands, we use annual means from each model simulation. We use 60°S-60°N for the near-global means, and 90°S-90°N for the global means, to allow for various latitudinal definitions of "global" ozone analyses in WMO ozone assessments (e.g. WMO 2011; WMO, 2014). In their analysis of CCMVal-2 simulations, *Eyring et al.* (2010b) used time-series additive model (TSAM) (*Scinocca et al.*, 2009) but here we show absolute ozone time series to identify models to be excluded as
- 25 outliers. As shown in **Table 2**, there are variations in the number of simulations for each reference/sensitivity experiment. Therefore, we decided to calculate the model averages (means and median) for each scenario using the following procedure.
- First, we calculate the zonal mean October, March or annual mean time series for each realisation. If more than one realisation is available for a particular model then we calculate an ensemble mean of monthly/annual mean time series. If only one realisation is available then we apply a 3-point (i.e. 3-year) boxcar smoothing to remove the short-term variations and somewhat mimic the effect of averaging an ensemble.
- These single time series for individual models are used to calculate the multi-model mean (MMM) and corresponding standard deviations for each year. We also calculate the '1-σ multi-model mean' (MMM1S) where we exclude models lying outside 1σ of the MMM for each year. Finally, all of the single time series are used to calculate the median model (MedM) along with the 10th and 90th percentiles.
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- To calculate smoothed and adjusted time series with respect to a reference year (e.g. 1960, 1980), we apply a 10-point boxcar smoothing to the individual model time series as well as the MMM, MMM1S and MedM time series.
- In order to make a robust estimate of ozone return dates we need to account for the different biases between the model simulations and observations. We calculate the mean biases between observational data and the REF-C2 MMM, MMM1S and MedM timeseries for the 1980-1984 time period. We choose the REF-C2 simulation as that is the reference that is used to estimate ozone return dates. An adjusted timeseries for each individual model is

then calculated by subtracting the respective observational bias. This procedure also results in the multi-model mean agreeing with the observations in the 1980-1984 period.

- Finally, using the MMM1S timeseries from REF-C2 as a reference line, the MMM1S timeseries from REF-C1, REF-C1SD, SEN-C2-fODS, SEN-C2-fGHG, SEN-C2-fCH4 and SEN-C2-fN2O are adjusted for the year 1980. The MMM1S timeseries from simulations starting in year 2000 (SEN-C2-RCP45, SEN-C2-RCP85) are adjusted for the year 2000 using the REF-C2 reference.
- 10 We compare the CCM simulations with selected observations to provide a basic evaluation of their performance. In particular, to test the height-resolved evolution of the modelled ozone fields we use BSVertOzone v1.0 (Bodeker Scientific Vertical Ozone, hereafter referred to as "BSVertOzone"), which is an updated and further developed version of the BDBP (Binary Database of Profiles) v1.1.0.6 that is described in detail in *Bodeker et al.* (2013). BSVertOzone
- 15 consists of monthly mean zonal mean ozone values on either pressure or altitude levels (from Earth's surface to about 70 km), where ozone is provided in mixing ratio or number density. For the data presented here the following improvements over the latest version of the BDBP were made: (1) ozone measurements from different data sources that are used as input for the monthly mean zonal mean calculation were updated (different satellite measurements and ozone
- 20 soundings); (2) additional data sources were added (Microwave Limb Sounder, MLS, ozone profiles and recent years of ozone soundings); (3) drifts and biases between measurements from different data sources are adjusted (using a chemical transport model as transfer standard); (4) uncertainties are propagated from individual measurements through all preparation and calculation procedure steps to the final product; and (5) the calculation of the monthly mean zonal mean values
- 25 was updated to correctly take into account the variable measurement frequencies of the different available data sources. The methodology of filling data gaps to construct a globally filled database is an updated version of the method described in *Bodeker et al.* (2013) where a pre-filling processing step was added. A more detailed description of BSVertOzone can be found in *Hassler et al.* (2018).
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BSVertOzone spans 70 pressure levels that are approximately 1 km apart (878.4 hPa to 0.046 hPa). For the calculation of the partial columns, ozone was interpolated to the exact boundaries of the partial columns from the two closest BSVertOzone pressure levels, and then ozone was integrated between the determined levels. The boundaries for the partial columns were defined as

- 35 follows: tropospheric column (surface-tropopause), lower stratospheric column (tropopause 10 hPa), upper stratospheric column (10 hPa and above; for BSVertOzone this means up to 0.046hPa). The tropopause pressure was defined as 100 hPa in the tropics (20°S-20°N), 150 hPa in the mid-latitudes (20°-60°N/S), and 200 hPa in the polar regions (60°-90°N/S). CCM partial columns were integrated between the same partial column boundaries, but directly from the CCM
- 40 pressure levels. No additional interpolation of CCM ozone profiles or BSVertOzone profiles was performed.

4) Results

4.1 Adjustment of model results

45 Figure 1 shows October mean total column ozone (TCO) from the REF-C2 simulations for the Antarctic to illustrate how the model simulations compare before and after adjustment to fit observations. (Other regions are shown in the Supplementary Material (SM) Figures S1-S5). Figure 1a shows the mean TCO time series from the individual models (with a 3-point boxcar smoothing if only 1 realisation is available) along with three estimates of the model mean. These

are the overall multi-model mean (MMM), the mean of the models which lie within 1σ of the MMM (MMM1S), and the median model (MedM). While all models show the characteristic behaviour of depletion followed by recovery, there is a large spread of around 150 DU between the model values at any time. This complicates the determination of the ozone return dates relative

- 5 to a baseline. Therefore, **Figure 1b** shows the same model runs after bias-correcting the individual model values to the observations over the period 1980-1984 and applying a 10-point boxcar smoothing. This correction to the models also forces the model means to fit the observations during this time period. With the bias correction, the individual models clearly give a more coherent picture and therefore, throughout this paper we generally show just the adjusted model
- 10 results (unless indicated otherwise). However, it is important to note that the values of the multimodel mean, which are used for our best estimate of ozone return dates, are similar between Figures 1a and 1b. Evidently, given enough model simulations, this approach of adjusting the model time series does not significantly alter the best estimate but does provide a smaller, more meaningful range of uncertainty. In other words, positive and negative model biases appear to be
- 15 equally represented in the CCMI ensemble, but removing them will allow comparison of model returns to the same common 1980 baseline. Concerning the different methods to estimate the model average, for the Antarctic October case shown here the results are very similar. The difference is in the estimated range of variability which is shown by the shading. The 10^{th} and 90^{th} percentiles of the median give the largest spread of around 100 DU. The 1σ variation of the MMM
- 20 is around ±40 DU. As expected the 1σ variation of the MMM1S, which has removed the outlying models, is smaller and around ±25 DU in this case. The MMM1S has an advantage over the median approach for the scenarios where there are not many realisations, and hence the $10^{th}/90^{th}$ percentiles cannot be derived robustly. Therefore, we use the MMM1S and the 1σ deviation (of the forecasts provided by the adjusted models) to determine our best estimate of return dates and

25 associated uncertainty, respectively.

4.2 Column ozone return dates

Figure 2 compares the MMM1S TCO from REF-C2 with the standard CCMI historical simulations REF-C1 and REF-C1SD, in which the models are nudged towards analysed
meteorology. Results are shown for both the direct and adjusted comparisons for the Antarctic and Arctic. The top panels show that, as expected, the REF-C1SD simulations reproduce better the observed evolution of TCO, as well as capturing much of the observed interannual variability. In particular, in the Antarctic the REF-C1SD mean reproduces the observed increase and then decrease in ozone after the year 2000. This was also shown in *Hardiman et al.* (2017) who

- 35 investigated the contributions of dynamics, transport and chemistry to these differences and is consistent with chemical transport model analyses in *Chipperfield et al.* (2015, 2017). For the Arctic it is noticeable that the REF-C1SD simulations give significantly lower TCO than the REF-C1 in the 1990s. This is related to the series of cold Arctic winters during this period when low TCO was caused through large chemical loss (e.g. 1994/95 and 1995/96) along with dynamical
- 40 contributions (e.g. 1996/97). The mean of the free-running REF-C1 simulations do not capture these low-column-ozone events. In contrast, **Figure 2** shows that the REF-C2 mean gives a more climatological picture of gradually varying ozone. The REF-C1 and REF-C2 means for the hindcast period are consistent. Therefore, within the limitations of the comparison of the heavily smoothed lines, the choice of ocean / sea ice representation does not seem to affect the
- 45 climatological TCO evolution in the regions shown, although as noted by *Zhang et al.* (2016) sea ice loss may affect zonally asymmetric TCO trends.

Figure 3 shows TCO results of the REF-C2 MMM1S for five latitude bands and the near-global (60°S-60°N) values. Also shown are the means from the sensitivity simulations SEN-C2-fGHG

and SEN-C2-fODS, which isolate the impacts of GHG and ODS changes. In all cases the shading gives the MMM1S 1σ variability. The TCO return dates from REF-C2, and the estimated range of that based on the 1σ variability, are summarised in **Table 3** and plotted in **Figure 4** (left panel). Globally the models predict a return to 1980 TCO values in 2049, with a 1σ spread from 2043-

- 5 2055. Earlier return dates are predicted in the NH mid-latitudes (2032), NH polar region (2034), and SH mid-latitudes (2045). In contrast, the return date from the large depletion in the SH polar region is much later at 2060. The 1σ variability gives the smallest range of return dates in the Antarctic (11 years) and SH mid-latitudes (11 years). The corresponding ranges in the NH, where dynamical variability is larger, are 24 years in mid-latitudes and 18 years near the pole. In the
- 10 tropics, the MMM1S shows a return to 1980 values towards 2058, followed by a turnaround and further decline. Note that individual models differ in whether they predict a return to 1980 values or not, before predicting the decrease at the end of this century (see SM **Figure S4**). The MMM1S near-global (60°S-60°N) column ozone also shows a decline after about 2080, which is mostly due to a decline in the tropics, with a small contribution from NH mid-latitudes. Dynamical decreases
- 15 in tropical TCO due to increased upwelling would be expected to cause increases in TCO at midhigh latitudes. For these variations in near global TCO after 2080 small changes of 2-3 DU in the tropospheric column, especially in the NH, are important factors (see Figure 9 below). The right panel of Figure 4 and Table 4 show the return dates of stratospheric column ozone (SCO, see below).

20

Our predictions of TCO return dates can be compared with previous estimates, particularly those used in past WMO Ozone Assessments. WMO (2011) used results from the CCMVal-2 experiments to derive ozone return dates and these are shown in **Table 3**. As no major update to CCMVal-2 had occurred, these same CCMVal-2 values were also used as the best estimates in the

- 25 subsequent WMO (2014) assessment. However, that assessment also showed results from a subset of four CCMVal-2 models, selected for their good representation of stratospheric circulation (Figures 2-21 and 3-15 from that assessment), and analysis of five CMIP5 models (Figure 2-23; *Eyring et al.*, 2013). We have included an analysis of these CMIP5 simulations in **Table 3**. Note that the CCMVal-2 runs used the A1b GHG scenario while the additional runs shown in WMO
- 30 (2014) used RCP 6.0 or 4.5. This study represents the first comprehensive update of TCO return dates since CCMVal-2. Compared to those values our return dates (albeit for a different GHG scenario) are later by values ranging from 4 years in the Arctic, 10 years in the Antarctic to 10-11 years at mid-latitudes. Similar differences are seen in the tropics, where not all models show a return. Interestingly, the subset of four CCMVal-2 models that performed new experiments for
- 35 WMO (2014), based on RCP 6.0 or 4.5, also showed later return dates than CCMVal-2, more in accord with our simulations. Compared to the CMIP5 models, which also use the RCP 6.0 scenario, our return dates are also later. The difference is small in northern mid-latitudes but it is as large as 14 years in the Antarctic (2046 compared to 2060).
- 40 There are three major differences in the assumptions used for the CCMVal-2 runs compared to the CCMI simulations, all of which likely contribute to our present estimate of up to a 17 year later return of total column ozone to the 1980 level. First, future atmospheric CO₂ rose more rapidly in the A1b scenarios used for CCMVal-2 compared to the RCP 6.0 scenario used by CCMI (**Figure S13**). Within A1b, CO₂ reaches 454 ppm in year 2032, which was the year given for
- 45 recovery of total column ozone in WMO (2011). This same mixing ratio for CO₂ is reached 9 years later within RCP 6.0. As a result, climate-driven changes in circulation within CCMI, which accelerate the recovery of the global ozone layer, will be similarly delayed. Next, the specification of CH₄ within A1b resulted in a more rapid rise compared to RCP 6.0 (**Figure S13**). For A1b in year 2032, atmospheric CH₄ is projected to be 44% larger than the 1980 value. For RCP 6.0, CH₄
- 50 is projected to rise by only 17% relative to 1980. As detailed in Section 4.5, the slower rise in CH₄ assumed for CCMI reduces the chemically-induced increase of ozone in both the stratosphere and

troposphere, contributing to a longer time for total column ozone to return to 1980 levels. The differences in the assumptions for future N_2O between A1b and RCP 6.0 are small, and therefore not responsible for our later return dates. Finally, the return of effective equivalent stratospheric chlorine to 1980 levels is delayed by about 5 years in the baseline WMO (2011) scenario (used in

- 5 this study) compared to the scenario used to drive CCMVal-2, reflecting new knowledge of lifetimes and emissions of various ODSs between the times these two scenarios were formulated. Since the return dates used in our study are based on analysis of results from 20 models, whereas the WMO (2011) return dates were based on results of 17 models, we have also examined the impact of restricting our return date estimates to a subset of the common models used by WMO
- 10 (2011). Return dates for global and near-global ozone found using only 9 common models differ by only 1 to 3 years compared to values given in **Table 3** for the complete set of models. Therefore, the primary causes of the later dates for global and near-global ozone to return to 1980 levels reported in **Table 3**, compared to WMO (2011) return dates, are assumptions regarding future atmospheric levels of CO₂, CH₄, and ODSs inherent to CCMVAL-2 and CCMI. As our estimates
- 15 are based on a large number of dedicated stratospheric simulations, with many models which have benefitted from further testing and development since CCMVal-2, we would argue that our values provide the best current estimates.
- The sensitivity simulations in **Figure 3** also illustrate the effect of climate change and ODS changes on total column ozone recovery, confirming the results of previous studies (e.g. *Morgenstern et al.*, 2018 and references therein). The smallest impact of climate change is predicted in the Antarctic where the simulation with fixed GHGs is very similar to REF-C2. In the mid-latitudes, increasing GHGs brings forward the return dates by about 10 (NH) to 20 (SH) years. In the Arctic, there is a larger impact. However, the variation of the MMM1S line for SEN-C2-
- 25 fGHG stays close to the 1980 reference line without crossing it, showing the challenge in extracting a return date (or range of dates) from the model runs. In the tropics, the competing effects of changes in ODSs, which increase ozone, and changes in GHGs, which decrease ozone, are clearly illustrated. In this region, the small rise in ozone due to ODS recovery is masked by decreases due to GHG increases. The net result is a decrease in tropical TCO in REF-C2.
- 30

4.3 Ozone profile variations

We now consider ozone recovery of partial columns in the lower (tropopause-10hPa; **Figure 5**) and upper (10 hPa and above; **Figure 6**) stratosphere separately. The figures include partial column observations derived from the composite BSVertOzone data set (*Bodeker et al.*, 2013;

- 35 *Hassler et al.*, 2018). This composite dataset has the advantage, compared to observations from any single instrument, of being fully height resolved and available continuously over 1979-2016. BSVertOzone consists of several different data sets. We use here the Tier 1.4 data set that is based on observations, but has been created by applying a least squares regression model to attribute variability to various known forcing factors (natural and anthropogenic) for ozone. The variability
- 40 in this data set is reduced compared to pure observations, since it describes only the variability for which basis functions were included in the regression model. This data set is therefore optimised for the use in comparisons with CCM simulations that do not exhibit the same unforced variability as reality. **Figures S10** and **S11** in the Supplementary Material show results from the specific altitudes of 50 hPa and 5 hPa, respectively, compared to GOZCARDS observations (*Froidevaux*)
- 45 *et al.*, 2015). In the lower stratosphere, where ozone has a long photochemical lifetime, the adjusted results from the models show some variations, especially in the polar regions. Overall, in the extra-tropical regions the models follow the observed behaviour in the BSVertOzone dataset although the MMM1S appears to overestimate depletion in the Antarctic and underestimate it in the Arctic. Interestingly, in the tropics the BSVertOzone dataset indicates ongoing decreases after

the year 2000 while the models show a levelling off and turnaround. This observed decrease in tropical lower stratospheric ozone has been noted by *Ball et al.* (2018) and is not captured by the models shown here. There are also significant differences between the models in the tropics where the lower stratosphere column does not return to 1980 values in all REF-C2 simulations. This is

- 5 also the case for the near-global lower stratospheric column, reflecting the large influence of the tropics. In the upper stratosphere (**Figure 6**), ozone behaves more similarly in all regions and between all models (i.e. depletion followed by recovery to values larger than 1980) as dynamical variations are less important. Therefore, there is generally less spread in the model forecasts of ozone recovery than over the whole stratosphere, although the ULAQ-CCM, CHASER and
- 10 CESM1-CAM4 models are outliers in certain regions. Note that the models consistently underestimate the BSVertOzone values and appear to show larger depletion. At this altitude, the feedback of temperature changes on ozone becomes important (*Haigh and Pyle*, 1982) and there is a larger increase in ozone than determined by ODS changes alone due to stratospheric cooling. As the CCMs are based on similar photochemical data, they should be expected to exhibit similar
- 15 climate sensitivities, although they may predict different magnitudes of climate change. The reason for the disagreement between models and BSVertOZone in the upper stratosphere requires further investigation, but these upper stratospheric differences will not have a large impact on column ozone return years.
- 20 Since the main driver of past ozone depletion and ozone recovery is the evolution of stratospheric halogen loading, we pay particular attention to how well the CCMs model the time-dependent abundance of organic and inorganic chlorine and bromine. These are based on prescribed scenarios should be fields for which the CCMs agree well. **Figure 7** shows the evolution of the modelled inorganic chlorine ($Cly = HCl + ClONO_2 + HOCl + ClO + 2Cl_2O_2 + Cl + BrCl + OClO + ...$) and
- 25 total chlorine (Cly + organic) in the upper and lower stratosphere in the Antarctic from the REF-C2 simulations. This region was chosen as an example to illustrate model-model differences. Long-term variations in the modelled chlorine loading are determined by the specified surface mixing ratios of the ODS. Through this, the different models are constrained to show the same approximate timescale for chlorine to return to its values in e.g. 1980. If the stratospheric ODSs
- 30 were simulated using emissions, rather than specified surface mixing ratios, then the timing of the model return dates would vary depending on the speed of the model circulation (*Douglass et al.*, 2008). Despite this constraint, the models do show a variation in the Cly loading at any time, especially in the lower stratosphere where the fractional conversion of organic chlorine to Cly is smaller. Nevertheless, some models show lower Cly than would be expected based on differences
- 35 in circulation (EMAC-L47MA, EMAC-L90MA, CCSRNIES). Moreover, in the upper stratosphere, where most chlorine would be in the form of Cly, there are still large variations between the models. The reasons for this require further investigation beyond the scope of this study, but, if models are conserving chlorine in their chemistry and transport schemes, the evolution of Cly at 5 hPa should closely track that of the specified tropospheric chlorine with a
- 40 lag of around 3-6 years. Figure 7 also includes Microwave Limb Sounder (MLS) observations of the October vortex-mean volume mixing ratio of HCl + ClO. This provides a lower limit of the amount of Cly present, though it will be a good approximation in these locations. The year-to-year variation at 50 hPa (around 250 pptv) is due to variability in the polar vortex and chlorine activation, which is not an issue at 5 hPa. The comparison with the MLS data clearly shows that
- 45 the three models with low Cly are unrealistic.

Bromine loading is another driver of ozone depletion and recovery. **Figure S12** compares total inorganic bromine modeled at 5 and 50 hPa in the Antarctic from the models. The highest values of bromine are reported by the SOCOL3 and EMAC-L90MA models. For SOCOL3, the model

50 was run using surface mixing ratios of 1.63 and 1.21 ppt for CHBr₃ and CH₂Br₂, respectively, which results in a bromine content for these very short lived substances (VSLS) of 7.31 ppt. This

is larger than the 5 ppt estimate for total bromine of VSLS suggested for use in CCM models by *Eyring et al.* (2013a). Within EMAC-L90MA, bromine sources are explicitly represented by considering oceanic emission of CHBr₃ and CH₂Br₂, three other VSLS species, as well as sea salt (*Warwick et al.*, 2006). The tendency of Bry within EMAC-L90MA to be larger than found by

- 5 other models, and to exceed an empirical estimate based on field observations in the Tropical Western Pacific, has been noted in a recent model intercomparison study focused on bromine (*Wales et al.*, 2018). The EMAC-L90MA over-estimate is likely caused by the oceanic emission terms being too large. The outlier is CHASER-MIROC, which has a bromine loading that peaks near 9 ppt in the upper stratosphere of the Antarctic, which is more than a factor of two less than
- 10 surface mixing ratios of bromocarbons that reach the stratosphere (e.g., *Wales et al.*, 2018) as well as the mean value of peak Bry from the other models. The low value of Bry within CHASER-MIROC is due to the consideration of only CH₃Br, without scaling to represent either halons and VSLS bromocarbons, for the simulations conducted for CCMI. Within this model, the BrO+ClO cycle will have an unrealistically low effect on ozone.
- 15

The decline in stratospheric halogen loading is a main driver in the increase of stratospheric ozone. However, ozone is also affected by other stratospheric factors, notably driven by changing climate, such as rising temperature in the upper stratosphere. Moreover, differences in this estimated climate effect is a source of variations between the projections provided by the models analysed

- 20 here. This climate effect is illustrated by the difference in the return dates to 1980 levels of stratospheric chlorine versus ozone. **Figure 8** (top and middle rows) shows how these return dates compare for local changes in the polar upper stratosphere and lower stratosphere, as well as column ozone return versus lower stratospheric Cly. For the upper stratosphere in both polar regions, the ozone return dates are much earlier than the Cly return dates and span a slightly larger
- 25 range. In the lower stratosphere and for the column in the Antarctic there is a much closer correspondence between the Cly return date (at 50 hPa) and the ozone return date. For the Arctic, the ozone return dates are earlier than for Cly (see also Figure 3). Figure 8 also shows the large spread in return dates between individual models. However, these return dates do generally correlate with each other so that an earlier Cly return date corresponds to an earlier ozone return 30 date. Some exceptions to this occur e.g. for the CMAM and EMAC-L90 models.

Figure 9 shows the evolution of the tropospheric partial column ozone (PCO) from simulations based on scenario REF-C2. In the Antarctic (October), the MMM1S shows very little change from year 1980 through the forecast period. This is most likely due to limited tropospheric ozone precursor abundance in the polar SH region (e.g., low NO_x). In the other five regions, the MMM1S evolution typically shows 2-5 DU enhancement going from year 1980 to the peak tropospheric PCO in the forecast period. The models then generally show a decrease in the final few decades of the century. It is also interesting to note the large spread across participating CCMs in tropospheric PCO in the 21st century. For example, the SH and NH mid-latitude panels show, for

- 40 year 2060, the range across the models is ~8-10 DU. The range is even larger near the end-of-the 21^{st} century where the NH mid-latitudes and Arctic (March) ranges are ~15 DU and ~20 DU, respectively. Several models have less tropospheric PCO in year 2100 relative to year 1980. This is suggestive that the tropospheric chemistry sensitivity to mainly CH₄ is very different in the CCMs. There is more discussion of this topic in **Section 4.5**.
- 45

4.4 Sensitivity of ozone return to climate change

Figure 10 shows the sensitivity of the tropospheric PCO evolution to assumed GHG RCP scenarios. The CH₄ temporal trend is the largest in SEN-C2-RCP85, followed by REF-C2, and SEN-C2-RCP45 scenarios. As discussed later in **Section 4.5**, an increase in tropospheric CH₄ will

enhance the NO_x-HO_x-smog net ozone production (Haagen-Smit et al., 1950). Therefore, the larger future trend in CH₄ as represented in the SEN-C2-RCP85 scenario, relative to the REF-C2 scenario (Figure S12), increases the year 2100 global annual average MMM1S tropospheric PCO by ~5 DU. Previous studies (e.g. Shindell et al., 2009; Eyring et al., 2013b; Morgenstern et al.,

- 2018) have also found that the tropospheric impact of CH₄ increases is a major contributor to the 5 TCO changes. Because of this difference in tropospheric PCO between scenarios we will only consider the evolution of SCO for derivation of return dates in the RCPs and SEN-C2 simulations in the discussion below.
- Figure 11 shows the sensitivity of the SCO return dates to the assumed GHG RCP scenarios. The 10 relative change is smaller in the Antarctic, where recovery is largely determined by Cly loading, but larger in all other regions. However, the absolute changes between, for example, the Antarctic (October) and Arctic (March) are similar. Simulation SEN-C2-RCP26 (not shown), which assumes only small climate change but for which we only have 5 realisations, does not return to
- 1980 values at all except in the Arctic. Globally and at mid-low latitudes, simulation SEN-C2-15 RCP45 shows a behaviour between the RCP 2.6 and RCP 6.0 simulations. Compared to REF-C2, the simulation with the largest impact of climate change, SEN-C2-RCP85, shows a similar behaviour globally, but with regional differences, i.e. a positive effect on ozone at all latitudes except the tropics, where ozone decreases the most under RCP 8.5. The impact on return dates in
- different regions is summarized in Figure 12. This figure shows the return date for each model 20 (coloured dots) along with the MMM1S (red triangle) for each scenario, all for six regions. The grey triangle is the MMM1S for the REF-C2 scenario (see Figure 4, right panel). The uncertainty in the MMM1S is represented by the solid vertical line. The SCO magnitude and range for all scenarios is listed in Table 4. For all regions shown in Figure 12, the derived MMM1S return
- date for SEN-C2-RCP45 is within the uncertainties range of the MMM1S return date for REF-C2. 25 This is also true for SEN-C2-RCP85, although in the Antarctic (Oct.) region where the MMM1S return date for SEN-C2-RCP85 MMM1S is shortened by 12 years the uncertainly range only just overlaps with the MMM1S from REF-C2.

4.5 Sensitivity of ozone return to methane and nitrous oxide 30

We now focus on the sensitivity simulations (Table 1) which examine the individual roles of CH4 (SEN-C2-fCH4 and SEN-C2-CH4RCP85) and N₂O (SEN-C2-fN2O), and their combined impact with CO₂ (SEN-fGHG), on ozone recovery. We will only consider stratospheric ozone columns (SCO), thereby eliminating any impact from changes in tropospheric ozone discussed above. We

- first give a general overview based on prior studies of the expected ozone recovery impacts of 35 changing CH₄, N₂O and the combined GHGs. We then discuss the individual ozone recovery impacts of N₂O, CH₄, and GHGs based on this work. The derived impacts are compared to the REF-C2 recovery dates. We will not attempt here to diagnose the reason why models vary in the derived impacts; these details are beyond the scope of this study and will be addressed in future
- 40 work.

Many studies have investigated the impacts of CH₄, N₂O, and CO₂ evolution on ozone abundance and recovery (e.g., Haigh & Pyle, 1979; Le Texier et al., 1988; Rosenfield et al., 2002; Randeniya et al., 2002; Royer et al., 2002; Chipperfield and Feng, 2003; Portmann and Solomon, 2007;

Shepherd et al., 2008; Ravishankara et al., 2009; Oman et al., 2010; Fleming et al., 2011; Revell 45 et al. 2012, 2015, 2016; Kirner et al., 2015; Butler et al., 2016; Keeble et al., 2017). In summary, increases in CH₄ and N₂O will generate higher amounts of hydrogen oxides (HO_X) and nitrogen oxides (NO_X), respectively. It is well known that increased NO_X will enhance catalytic stratospheric ozone loss (Crutzen, 1970). Therefore, one would expect the ozone return date to be extended for temporal increases in N₂O, or shortened for decreases in N₂O. Nonetheless, the effect of future increases of N₂O varies with altitude and also depends on the temporal evolution of other GHGs (*Wang et al.*, 2014; *Revell et al.*, 2015). For changes in CH₄ the situation is more complicated. In a similar manner to NO_X, increased HO_X will decrease upper stratospheric ozone.

- 5 However, CH₄ can also affect the partitioning of reactive chlorine through the reaction of CH₄ + Cl \rightarrow HCl + CH₃, with more CH₄ leading to an increase in stratospheric ozone via a decrease in the abundance of reactive chlorine. Overall, temporal increases in CH₄ lead to increases in stratospheric column ozone (*Revell et al.*, 2012). In the troposphere, increases in CH₄ will enhance chemical production through NO_X-HO_X-smog process (*Haagen-Smit et al.*, 1950). This
- 10 tropospheric ozone net production and subsequent tropospheric partial column change is shown for the REF-C2 simulations in **Figure 9**. It should be noted that UMSLIMCAT shows a small tropospheric trend since the ozone is prescribed in the troposphere. UMUKCA-UCAM has only simplified tropospheric chemistry, whereas NIWA-UKCA has a representation of C_2 - C_3 -isoprene oxidation. In addition, the tropospheric trend is affected by the coupling to the stratosphere via
- 15 changes in stratotosphere-to-troposphere exchange and photolytic feedbacks. Although CO₂ is chemically inert below about 60 km, increases in its abundance (along with CH₄ and N₂O) will cool the stratosphere (*Haigh & Pyle*, 1979). This cooling will slow down the catalytic ozone destruction cycles and increase ozone, therefore temporal increases in CO₂, CH₄, and N₂O will shorten the ozone return date due to this process, which is most important in the upper stratosphere
- 20 and mesosphere. The warming of the troposphere and the cooling of the stratosphere can also affect the Brewer-Dobson circulation and therefore impact ozone through transport (e.g., *Polvani et al.*, 2017, 2018 and references therein). The cooling process operates throughout the stratosphere, but is most important for dynamical processes in the lower to middle stratosphere. Outside of the tropics, a speed-up of the Brewer-Dobson circulation would shorten the ozone

25 recovery date, while a slow-down of the BD circulation would extend it.

All of the above-mentioned chemical, radiative, and dynamical impacts are represented within the REF-C2 simulations (using RCP 6.0 GHGs for the future period). Here we examine the sensitivity scenarios for N_2O and CH_4 individually, along with the combined GHGs scenario impacts relative

- 30 to the REF-C2 scenario. It should be noted that for the two CH₄ and one N₂O sensitivity scenarios, there is only one realisation available for each model, whereas for many models the REF-C2 scenario has multiple ensemble members (**Table 2**). The SEN-C2 temporal abundances compared to REF-C2 are shown for N₂O, CH₄, and CO₂ in **Figure S13.** For N₂O (SEN-C2-fN2O), the abundance is approximately 290 ppbv and 405 ppbv for 1960 and 2100, respectively, an increase
- 35 of 115 ppbv (~40%). For CH₄ (SEN-C2-fCH4) the abundance is 1.24 ppmv in 1960 and a maximum of 1.96 ppmv in the 2070s, an increase of 57%. The SEN-C2-RCP85 scenario increases CH₄ over that given by the REF-C2 scenario by 2.1 ppmv in 2100, an increase of 128%. The CO₂ change in REF-C2 from 1960 to 2100 is 352 ppmv, approximately a 110% increase.
- 40 **Figure 13** shows the evolution of October mean SCO in the Antarctic region for the REF-C2, SEN-C2-fN2O (fixed N₂O at 1960 conditions), SEN-C2-fCH4 (fixed CH₄ at 1960 conditions), and SEN-C2-CH4RCP85 (RCP8.5 CH₄ abundance) scenarios. The SCO observations are again based on the BSVertOzone data set (*Bodeker et al.*, 2013). The panels show results from eight different models. Solid lines show 10-year smoothed SCO for a given simulation. Shading
- 45 indicates the 1- σ standard derivation from an ensemble of realisations (or 3-box smoothed line if only one realisation is available). In addition, the SEN-C2-fODS (fixed ozone-depleting substances in 1960) is shown. This scenario shows the behaviour of SCO without the evolution of halogens (i.e., no ozone depletion and recovery due to halogens). This scenario does include the previously discussed impacts of N₂O, CH₄, and CO₂ on SCO. However, since there is no ozone
- 50 depletion period from ODSs in this simulation, it does not make sense to calculate an ozone recovery date. The red lines in all panels show the evolution of SCO for the REF-C2 scenario,

which can be directly compared to the BSVertOzone dataset. For each model, the REF-C2 simulations have been bias corrected to the BSVertOzone dataset for the 1980-1984 period. The SEN-C2 simulations are then adjusted to the bias corrected REF-C2 ensemble mean for the 1960 period. Comparison of the SCO for the REF-C2 simulations show that models generally compare

- 5 well to BSVertOzone for the depletion period, although CMAM and UMSLIMCAT appear to be biased low. In summary, when one examines the relative impact on the ozone return date across the eight models from the four SEN-C2 scenarios, there is no consistent pattern. Therefore, the result suggests that the Antarctic region is not sensitive to the perturbations presented in this work.
- 10 This is not the case when one examines the annual average SH mid-latitude region (**Figure 14**). Generally, across the models the ozone return date varies as follows. The SEN-C2-fN2O (light blue line) simulations return before the REF-C2 simulations. This is consistent with our understanding that less NO_X produced from a fixed 1960 N₂O abundance will allow the SCO to recover earlier than the increasing NO_X in a REF-C2 scenario. However, it should be noted that
- 15 there can be a smaller impact of N₂O on the return date due to cancellation of the upper and lower stratospheric response of N₂O on ozone (*Morgenstern et al.*, 2018). The SCO from the SEN-C2-CH4RCP85 simulations (dark blue line), for the four models which performed this run, also tends to have a recovery date that is earlier than the REF-C2 simulations. Again, with more CH₄ specified in the 21st century, ozone will recover faster due to the sequestering of reactive chlorine
- 20 into HCl and the stratospheric cooling effect of slowing down ozone loss rates. The impact of increased HO_x production from increased CH₄, causing more ozone depletion to extend the recovery, does not dominate over these two processes. In contrast, the SEN-C2-fCH4 simulation (purple line), has less CH₄ in the 21st century than the REF-C2 simulation and therefore has later return date. Finally, the SEN-C2-fGHG simulation generally has the latest ozone return date. The
- 25 corresponding time evolution results for the SEN-C2 scenarios for the Arctic (March), annual average NH mid-latitude region, and annual average near-global (60°S-60°N) are shown in Figures S14, S15, and S16, respectively.
- The SCO return dates for the simulations based on the four sensitivity scenarios are also summarised and compared to the REF-C2 scenario in **Figure 12** (see above) and **Table 4**. The individual model details regarding the SCO return date (similar in format to **Figure 8**) between the SEN-C2 simulations and the REF-C2 simulations are shown in **Figures S17-S20**. We first discuss the change in SCO return dates between the SEN-C2-fGHG and REF-C2 simulations. The Antarctic (October) region difference between the two scenarios is small, within 2 years. The
- 35 uncertainty range for both scenarios are approximately ± 12 years. The SH mid-latitudes region shows that the MMM1S SCO recovery date is extended in the SEN-C2-fGHG case by ~16 years relative to the REF-C2 case. This extended SCO recovery period is even larger in the NH midlatitudes and Arctic (March) by ~25 years. This is consistent with the GHG cooling impact on ozone loss rates and a lack of strengthening of the BD circulation. In this comparison, having 1960
- 40 abundances of GHGs compared to the REF-C2 evolution means less cooling of the stratosphere and therefore an extension of the SCO recovery date. In addition, the CH₄ abundance is less in SEN-C2-fGHG which also decreases sequestering of reactive chlorine into HCl and acts to extend the SCO recovery date. Both of these factors override the direct impact of less production of NO_x and HO_x from N₂O and CH₄ which would shorten the SCO recovery. Interestingly the near-global
- 45 (annual) average SCO return date is not that different between the two scenarios. This is most likely due to the fact that when the BD circulation strengthens, tropical ozone is reduced and extratropical ozone is increased. Therefore, the net impact on the stratospheric column ozone return date cancels out for this process in the global average. **Figure 12** and **Table 4** shows the comparison of the SCO return dates for the SEN-C2-fN2O and the REF-C2 simulations. In this
- 50 comparison, one would expect that SEN-C2-fN2O with 1960 abundances of N₂O would bring forward the SCO recovery date. This is certainly true for the near-global (annual) average

comparison, where the MMM1S SEN-C2-fN2O SCO recovery date is shortened by ~20 years relative to the REF-C2 case. This is mostly due to a shortening of the return date in the tropics; at mid-high latitudes there is little change. As noted above, the future rise in N₂O can lead to significant increases in lower stratospheric ozone, particularly for regions where the loss rate of

- 5 ozone due to halogens exceeds that due to NO_x prior to the perturbation of N_2O . The effect of N_2O on ozone varies as a function of latitude and altitude (*Wang et al.*, 2014), complicating the sensitivity to the ozone return date to variations in N_2O (*Morgenstern et al.*, 2018). Figure 12 and Table 4 also shows the comparison of the SCO return dates for the SEN-C2-fCH4 simulations. Here all regions except the Antarctic (and the tropics, for which the return date is undefined) show
- 10 an extension of the SCO return date. This is consistent with the discussion above for the fixed GHG scenario. The near-global (annual) average SCO return date is extended by ~5 years. For the SEN-C2-CH4RCP85 scenario the MMM1S near-global (annual) average SCO return date is reduced relative to the REF-C2 scenario by ~15 years. As expected, this is an opposite effect to that derived from the SEN-C2-fCH4 scenario. In the other regions for SEN-C2-CH4RCP85
- 15 MMM1S there is less separation between the REF-C2 reference and the CH₄ perturbation scenario. Unfortunately for this work, there was only one realisation from each modelling group for the SEN-C2-fN2O, SEN-C2-fCH4, and SEN-C2-CH4RCP85 scenarios. Therefore, the signal from the perturbation on the SCO return date may be affected by the internal variability of each CCM. For future assessments of single forcing or combined effects of CO₂, CH₄, and N₂O on SCO
- 20 the return date, a recommendation based on this work is to have at least three ensemble members per scenario type.

5) Summary and Conclusions

We have analysed simulations performed for the Chemistry-Climate Modelling Initiative (CCMI) to estimate the return dates of the stratospheric ozone layer from depletion caused by anthropogenic stratospheric chlorine and bromine. CCMI represents an extensive multi-model exercise to study the future evolution of the ozone layer under changing climate conditions. Here we consider a total of 155 simulations from 20 models, including a range of sensitivity studies. For the control simulations (unconstrained by nudging towards analysed meteorology) there is a

- 30 large spread in the predictions of the absolute ozone column. Therefore, the model results need to be adjusted for biases against historical data. Also, the interannual variability in the model results need to be smoothed in order to provide a reasonable and useful estimate of the range of ozone return dates.
- 35 The total column ozone return dates calculated here differ from those presented in the previous WMO/UNEP assessment (WMO, 2014). The differences could be explained by the choice of GHG scenario for the baseline estimate (A1b in WMO (2014) versus RCP 6.0 here), and some model updates including realistic tropospheric chemistry. In addition, the time series for surface ODSs used here results in a return of stratospheric halogen loading to the 1980 value that is about
- 40 5 years later than assumed for the CCM runs examined in WMO (2014). The CCMI models project that global total column ozone will return to 1980 values in 2049 (with a 1- σ uncertainty of 2043-2055). At mid-latitudes, Southern Hemisphere ozone is projected to return to 1980 values in 2045 (2039-2050), and Northern Hemisphere ozone in 2032 (2020-2044). In the polar regions, the return dates are 2060 (2055-2066) in the Antarctic in October and 2034 (2025-2043) in the Arctic
- 45 in March. The earlier return dates in the NH reflect larger impact of dynamics on ozone in this hemisphere. In the tropics only around half the models predict a return to 1980 values, at around 2040, while the other half do not show a return to 1980 values (giving the mean of 2058). All models show a negative trend in tropical total column ozone towards the end of the 21st century.

An important result from the simulations presented here is the strong regional differences in the future evolution of total column ozone due to the effects of climate change. These climate effects are least evident in the Antarctic spring where future ozone depends largely on halogen loading. In contrast, in the NH the models predict a super-recovery while in the tropics the models predict

5 a possible further decrease in column ozone, possibly without any return to 1980 values.

There is different behaviour in the partial column ozone between the lower and upper stratosphere. In the lower stratosphere, where ozone is long-lived and affected by dynamics, there are differences in the timescale for recovery between the polar regions and mid-latitudes. Moreover,

- 10 in the tropics, increased upwelling prevents the return of partial column ozone (PCO) in many models. In contrast in the upper stratosphere the predicted behaviour is similar in all regions. Ozone returns to values larger than in 1980 by 2040 and carries on increasing due to the effect of stratospheric cooling. For the upper stratosphere, the CCM predictions do not vary a lot and are in good agreement with past observations, indicating that relevant processes are represented
- 15 adequately in the models. For the lower stratosphere, some obvious differences are seen between the CCM results, indicating possible inadequate descriptions of dynamical (transport) and chemical (heterogeneous) processes due to temperature biases (in the polar regions and tropics) in the CCMs.
- 20 The CCMI models generally agree in their simulation of the time evolution of stratospheric inorganic chlorine (Cly), which is the main driver of ozone loss and recovery, although there is some inter-model variability. The situation is similar for the simulation of inorganic bromine but with a larger relative spread between the models due to varying treatments of short-lived species and a model without a representation of halons. However, there do appear to be issues with the
- 25 chemistry and/or transport in a few of the model simulations. Throughout the stratosphere the spread of ozone return dates to 1980 values between models tends to correlate with the spread of the return of Cly to 1980 values. In the upper stratosphere, greenhouse gas-induced cooling speeds up the return by about 10-20 years. In the lower stratosphere, and for the column, there is a more direct link in the timing of the return dates, especially for the large Antarctic depletion.
- 30 Comparisons of total column ozone between the models is affected by different predictions of the evolution of tropospheric ozone within the same scenario, presumably due to differing treatment of tropospheric chemistry. Therefore, for some scenarios clear conclusions can only be drawn for stratospheric ozone columns.
- 35 As noted by previous studies, the timing of ozone recovery is affected by the evolution of N₂O and CH₄. However, the effect in the simulations analysed here is small and at the limit of detectability from the small number of realisations available for these experiments compared to internal model variability. The sensitivity scenarios examined in this study did show that a future decline of N₂O below an RCP6.0 projection (i.e., back to 1960 abundances) could reduce the global SCO return date by up to 15 years. In the opposite sense, an increase of CH₄ above the RCP6.0 projection (i.e., using RCP8.5 abundances) could again reduce the global SCO return date by up to 15 years. Both the N₂O and CH₄ global column ozone sensitivities are mainly realised through chemical impacts in the tropics.
- 45 Overall, our estimates of ozone return dates are uncertain due to both uncertainties in future scenarios, in particular of GHGs, and uncertainties in models. The scenario uncertainty is small in the short term but increases with time, and is large by the end of the century. For the models, while it is possible that they all may be missing important but unknown processes, there are still some model-model differences related to known first-order processes which affect ozone recovery.
- 50 Work needs to continue to ensure that models used for assessment purposes accurately represent stratospheric chemistry and the prescribed scenarios of ozone-depleting substances, and only those

models are used to calculate return dates. Nevertheless, the agreement between the results presented here and previously published work gives some confidence that we can model the future evolution of the ozone layer. For future assessments of single forcing or combined effects of CO_2 , CH_4 , and N_2O on the stratospheric column ozone return dates, this work suggests that is more

5 important to have multi-member (at least 3) ensembles for each scenario from each established participating model, rather than a large number of individual models.

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References

Austin, J., and N. Butchart, Coupled chemistry-climate model simulation for the period 1980 to 2020: Ozone depletion and the start of ozone recovery, *Q. J. R. Meteorol. Soc.*, 129, 3225–

3249, 2003.

5

40

50

- Austin, J., & Wilson, R. J.: Ensemble simulations of the decline and recovery of stratospheric ozone. J. Geophys. Res., 111(D16), D16314, doi:10.1029/205JD006907, 2006.
- Austin, J., J. Scinocca, D. Plummer, L. Oman, D. Waugh, H. Akiyoshi, S. Bekki, P. Braesicke, N. Butchart, M. P. Chipperfield, D. Cugnet, M. Dameris, S. Dhomse, V. Eyring, S. Frith, R.
- 10 Garcia, H. Garny, A. Gettelman, S.C. Hardiman, D. Kinnison, J. F. Lamarque, E. Mancini, M. Marchand, M. Michou, O. Morgenstern, T. Nakamura, S. Pawson, G. Pitari, J. Pyle, E. Rozanov, T. G. Shepherd, K. Shibata, H. Teyssedre, R.J. Wilson, and Y. Yamashita , The decline and recovery of total column ozone using mult-model time series analysis, *J. Geophys. Res.*, doi:10.1029/2010JD013857, 2010.
- 15 Bednarz, E. M., Maycock, A. C., Abraham, N. L., Braesicke, P., Dessens, O., and Pyle, J. A.: Future Arctic ozone recovery: the importance of chemistry and dynamics, *Atmos. Chem. Phys.*, 16, 12159-12176, https://doi.org/10.5194/acp-16-12159-2016, 2016.
 - Ball, W. T., Alsing, J., Mortlock, D. J., Staehelin, J., Haigh, J. D., Peter, T., Tummon, F., Stübi, R., Stenke, A., Anderson, J., Bourassa, A., Davis, S. M., Degenstein, D., Frith, S., Froidevaux,
- 20 L., Roth, C., Sofieva, V., Wang, R., Wild, J., Yu, P., Ziemke, J. R., and Rozanov, E. V.: Continuous decline in lower stratospheric ozone offsets ozone layer recovery, *Atmos. Chem. Phys.*, 18, 1379-1394, doi:10.5194/acp-2017-862, 2018.
 - Bekki, S., A. Rap, V. Poulain, S. Dhomse, M. Marchand, F. Lefevre, P. M. Forester, S. Szopa, M.P. Chipperfield, Climate impact of stratospheric ozone recovery. *Geophy. Res. Lett.*, 40(11),
- 25 2796–2800, doi:10.1002/grl.50358, 2013.
 - Bodeker, G.E., Hassler, B., Young, P.J., and Portmann, R.W.: A vertically resolved, global, gapfree ozone database for assessing or constraining global climate model simulations, *Earth Syst. Sci. Data*, 5, 31–43, doi:10.5194/essd-5-31-2013, 2013.

Brasseur, G., and M. H. Hitchman, Stratospheric response to trace gas perturbations: Changes in ozone and temperature distributions, *Science*, 240, 367–634, 1988.

Brewer, A. W., Evidence for a world circulation provided by the measurements of helium and water vapour distribution in the stratosphere, *Q. J. Roy. Meteorol. Soc.*, 75(326), 351–363, 1949.

Butchart, N., and Scaife, A. A., Removal of chlorofluorocarbons by increased mass exchange

- between the stratosphere and troposphere in a changing climate. *Nature*, *410*(6830), 799–802, doi:10.1038/35071047, 2001.
 - Butchart, N., A.A. Scaife, M. Bourqui, J. de Grandpré, S.H.E. Hare, J. Kettleborough, U. Langematz, E. Manzini, F. Sassi, K. Shibata, D. Shindell, and M. Sigmond: Simulations of anthropogenic change in the strength of the Brewer-Dobson circulation. *Clim. Dyn.*, 27, 727-741, doi:10.1007/s00382-006-0162-4, 2006.
 - Butchart, N., I. Cionni, V. Eyring, T. G. Shepherd, D. W. Waugh, H. Akiyoshi, J. Austin, C. Bruhl, M. P. Chipperfield, E. Cordero, M. Dameris, R. Deckert, S. Dhomse, S. M. Frith, R. R. Garcia, A. Gettelman, M. A. Giorgetta, D. E. Kinnison, F. Li, E. Mancini, C. McLandress, S. Pawson, G. Pitari, D. A. Plummer, E. Rozanov, F. Sassi, J. F. Scinocca, K. Shibata, B. Steil, and W.
- 45 Tian, Chemistry-climate model simulations of 21st century stratospheric climate and circulation changes, *J. Climate*, 23, 5349-5374, doi: 10.1175/2010JCLI3404.1, 2010.
 - Butler, A. H., Daniel, J. S., Portmann, R.W., Ravishankara, A. R., Young, P. J., Fahey, D. W., and Rosenlof, K. H.: Diverse policy implications for future ozone and surface UV in a changing climate, *Environ. Res. Lett.*, 11, 064017, https://doi.org/10.1088/1748-9326/11/6/064017, 2016.

Chipperfield, M.P., S. Bekki, S. Dhomse, N.R.P. Harris, B. Hassler, R. Hossaini, W. Steinbrecht,

R. Thieblemont and M. Weber, Detecting recovery of the stratospheric ozone layer, *Nature*, 549, 211-218, doi:10.1038/nature23681, 2017.

Chipperfield, M.P., S.S. Dhomse, W. Feng, R.L. McKenzie, G. Velders and J.A. Pyle, Quantifying the ozone and UV benefits already achieved by the Montreal Protocol, *Nature Communications*, 6, 7233, doi:10.1038/ncomms8233, 2015.

- 6, 7233, doi:10.1038/ncomms8233, 2015.
 Chipperfield, M. P., & Feng, W., Comment on: Stratospheric Ozone Depletion at northern midlatitudes in the 21st century: The importance of future concentrations of greenhouse gases nitrous oxide and methane. *Geophys. Res. Lett.*, 30(7), doi:10.1029/2002GRL16353, 2003.
- Crutzen, P.: The influence of nitrogen oxides on atmosphere ozone content, Q. J. R. Meteorol.
 Soc., 96, 320, 1970.
 - Douglass, A., R. S. Stolarski, M. R. Schoeberl, C. H. Jackman, M. L. Gupta, P. A. Newman, J. E. Nielsen, and E. L. Fleming, relationship of loss, mean age of air and the distribution of CFCs to stratospheric circulation and implications for atmospheric lifetimes, *J. Geophys. Res.*, 113, D14309, doi:10.1029/2007JD009575, 2008.
- 15 Eichelberger, S. J., and D. L. Hartmann, Changes in the strength of the Brewer–Dobson circulation in a simple AGCM. *Geophys. Res. Lett.*, 32, L15807, doi:10.1029/2005GL022924, 2005.

Eyring et al., D.W. Waugh, G.E. Bodeker, E. Cordero, H. Akiyoshi, J. Austin, S.R. Beagley, B. Boville, P. Braesicke, C. Bruehl, N. Butchart, M.P. Chipperfield, M. Dameris, R. Deckert, M.

- 20 Deushi, S.M. Frith, R.R. Garcia, A. Gettelman, M. Giorgetta, D.E. Kinnison, E. Mancini, E. Manzini, D.R. Marsh, S. Matthes, T. Nagashima, P.A. Newman, J.E. Nielsen, S. Pawson, G. Pitari, D.A. Plummer, E. Rozanov, M. Schraner, J. F. Scinocca, K. Semeniuk, T.G. Shepherd, K. Shibata, B. Steil, R. Stolarski, W. Tian and M. Yoshiki, Multi-model projections of ozone recovery in the 21st century, *J. Geophys. Res.*, 112, D16303, doi:10.1029/2006JD008332,
- 25 2007.
 - Eyring, V., M. P. Chipperfield, M. A. Giorgetta, D. E. Kinnison, K. Matthes, P. A. Newman, S. Pawson, T. G. Shepherd, and D. W. Waugh, Overview of the new CCMval reference and sensitivity simulations in support of upcoming ozone and climate assessments and the planned SPARC CCMVal report, *SPARC Newsletter*, 30, 20–26, 2008.
- 30 Eyring, V., I. Cionni, J. F. Lamarque, H. Akiyoshi, G. E. Bodeker, A. J. Charlton-Perez, S. M. Frith, A. Gettelman, D. E. Kinnison, T. Nakamura, L. D Oman, S. Pawson, and Y. Yamashita, Sensitivity of 21st century stratospheric ozone to greenhouse gas, *Geophys. Res. Lett.*, doi:10.1029/2010GL044443, 2010a.
 - Eyring, V., I. Cionni, G. E. Bodeker, A. J. Charlton-Perez, D. E. Kinnison, J. F. Scinocca, D. W. Waugh, H. Akiyoshi, S. Bekki, M. P. Chipperfield, M. Dameris, S. Dhomse, S. M. Frith, H.
- Waugh, H. Akiyoshi, S. Bekki, M. P. Chipperfield, M. Dameris, S. Dhomse, S. M. Frith, H. Garny, A. Gettelman, A. Kubin1, U. Langematz1, E. Mancini, M. Marchand, T. Nakamura, L. D. Oman, S. Pawson, G. Pitari, D. A. Plummer, E. Rozanov, T. G. Shepherd, K. Shibata, W. Tian, P. Braesicke1, S. C. Hardiman, J. F. Lamarque, O. Morgenstern, J. A. Pyle and Y. Yamashita, Multi-model assessment of ozone return dates and ozone recovery in CCMVal-2 models, *Atmos. Chem. Phys.*, 10, 9451-9472, doi:10.5194/acp-10-9451-2010, 2010b.
- Eyring, V. J.-F. Lamarque, P. Hess, F. Arfeuille, K. Bowman, M. Chipperfield, B. Duncan, A. Fiore, A. Gettelman, M. Giorgetta, C. Granier, M. Hegglin, D. Kinnison, M. Kunze, U. Langematz, B. Luo, R. Martin, K. Matthes, P. A. Newman, T. Peter, A. Robock, T. Ryerson, A. Saiz-Lopez, R. Salawitch, M. Schultz, T. G. Shepherd, D. Shindell, J. Staehelin, S.
- 45 Tegtmeier, L. Thomason, S. Tilmes, J-P Vernier, D. W. Waugh, P. J. Young, Overview of IGAC/SPARC Chemistry-Climate Model Initiative (CCMI) Community Simulations in Support of Upcoming Ozone and Climate Assessments, SPARC Newsletter, no. 40, pg48-66, January 2013, 2013a.
- Eyring, V., J. M. Arblaster, I. Cionni, J. Sedlacek, J. Perlwitz, P. J. Young, S., Bekki, D.
 Bergmann, P. Cameron-Smith, W. J. Collins, G. Faluvegi, K.-D. Gottschaldt, L. W. Horowitz1,
 D. E. Kinnison, J.-F. Lamarque, D. R. Marsh, D. Saint-Martin, D. T. Shindell, K. Sudo1, S.

Szopa, and S. Watanabe, Long-term changes in tropospheric and stratospheric ozone and associated climate impacts in CMIP5 simulations, *J. Geophys. Res.*, 118, 5029-5060, doi:10.1002/jgrd.50316, 2013b.

- Fels, S. B., J. D. Mahlman, M. D. Schwarzkopf, and R. W. Sinclair, Stratospheric Sensitivity to
 Perturbations in Ozone and Carbon Dioxide: Radiative and Dynamical Response, *J. Atmos. Sci.*, 37, 2265-2297, 1980.
 - Fleming, E. L., Jackman, C. H., Stolarski, R. S., and Douglass, A. R.: A model study of the impact of source gas changes on the stratosphere for 1850–2100, *Atmos. Chem. Phys.*, 11, 8515–8541, https://doi.org/10.5194/acp-11-8515-2011, 2011.
- 10 Frith, S. M., Stolarski, R. S., Kramarova, N. A., and McPeters, R. D.: Estimating uncertainties in the SBUV Version 8.6 merged profile ozone data set, *Atmos. Chem. Phys.*, 17, 14695-14707, https://doi.org/10.5194/acp-17-14695-2017, 2017.

Froidevaux, L., J. Anderson, H.-J. Wang, R. A. Fuller, M. J. Schwartz, M. L. Santee, N. J. Livesey, H. C. Pumphrey, P. F. Bernath, J. M. Russell III, and M. P. McCormick, *Atmos. Chem. Phys.*, 15, 10471–10507, doi:10.5194/acp-15-10471-2015.

- 15, 10471–10507, doi:10.5194/acp-15-10471-2015.
 Garcia, R. R., and W. J. Randel, Acceleration of the Brewer-Dobson circulation due to increases in greenhouse gases, *J. Atmos. Sci.*, 65(8), 2731–2739, 2008.
 - Haagen-Smit, A. J.: The air pollution problem in Los Angeles, *Eng. Sci.*, 14, 7-13, Calif. Inst. Of Technol., Pasadena, 1950.
- 20 Haigh, J. D., & Pyle, J. A.: A two-dimensional calculation including atmospheric carbon dioxide and stratospheric ozone. *Nature*, 279, 222–224. doi:10.1038/279222a0, 1979.

Haigh, J. D., and J. A. Pyle, Ozone perturbation experiments in a two-dimensional circulation model, *Q. J. Roy. Meteorol. Soc.*, 108, 551-574, 1982.

- Hardiman, S. C., Butchart, N., O'Connor, F. M., and Rumbold, S. T.: The Met Office HadGEM3-
- 25 ES chemistry–climate model: evaluation of stratospheric dynamics and its impact on ozone, *Geosci. Model Dev.*, 10, 1209-1232, https://doi.org/10.5194/gmd-10-1209-2017, 2017.

Hassler, B., Kremser, S., Bodeker, G.E., Lewis, J., Nesbit, K., Chipperfield, M., and Dameris, M.: An updated version of a gap-free monthly mean zonal mean ozone database, *Earth Syst. Sci. Data Discuss.*, doi:10.5194/essd-2018-56, (in review), 2018.

30 Hegglin, M. I. and J.-F. Lamarque: The IGAC/SPARC Chemistry-Climate Model Initiative Phase-1 (CCMI-1) model data output, NCAS British Atmospheric Data Centre, date of citation. <u>http://catalogue.ceda.ac.uk/uuid/9cc6b94df0f4469d8066d69b5df879d5</u>, 2015.

Keeble, J., E. M. Bednarz, A. Banerjee, N. L. Abraham, N. R. P. Harris, A. C. Maycock, J. A. Pyle, Diagnosing the radiative and chemical contributions to future changes in trooical column

35 ozone with the UM-UKCA chemistry-climate model, *Atmos. Chem. Phys.*, 17, 13801-13818, doi.org/10.5194/acp-17-13801-2017, 2017.

40

50

Kirner, O., Ruhnke R., and Sinnhuber, B.-M.: Chemistry–Climate Interactions of Stratospheric and Mesospheric Ozone in EMAC Long-Term Simulations with Different Boundary Conditions for CO₂, CH₄, N₂O, and ODS, *Atmosphere-Ocean*, 53, 140-152, doi:10.1080/07055900.2014.980718, 2015.

Lamarque, J. F., et al., Historical (1850–2000) gridded anthropogenic and biomass burning emissions of reactive gases and aerosols: methodology and application. *Atmos. Chem. Phys.*, 10, 7017-7039, 2010.

Le Texier, H., Solomon, S., & Garcia, R. R.: The role of molecular hydrogen and methane

- 45 oxidation in the water vapour budget of the stratosphere. *Q. J. Roy. Meteorol. Soc.*, 114, 281–295. doi:10.1002/qj.49711448002, 1988.
 - Li, F., Stolarski, R. S., & Newman, P. A.: Stratospheric ozone in the post-CFC era. *Atmos. Chem. Phys.*, *9*(6), 2207–2213, doi:10.5194/acp-9-2207-2009, 2009.

Livesey, N., et al. Version 4.2x Level 2 data quality and description document, Tech. Rep. JPL D-33509 Rev. C, Jet Propulsion Lab, 2017.

22

Marsh, D. R., Lamarque, J.-F., Conley, A. J., and Polvani, L. M., Stratospheric ozone chemistry feedbacks are not critical for the determination of climate sensitivity in CESM1(WACCM). *Geophys. Res. Lett.*, 43(8), 3928–3934, doi:10.10002/2016GL068344, 2016.

- Masui T, Matsumoto, K., Hijioka, Y., Kinoshita, T., Nozawa, T., Ishiwatari, S., Kato E., Shukla
 P. R., Yamagata, Y., Kainuma, M., *An emission pathway for stabilization at 6 Wm⁻² radiative forcing. Climatic Change*, 109, 59-76. doi:10.1007/s10584-011-0150-5, 2011.
- McLandress, C., Jonsson, A. I., Plummer, D. A., Reader, M. C., Scinocca, J. F., Shepherd, T. G.: Separating the dynamical effects of climate change and ozone depletion. Part I: Southern Hemisphere stratosphere. J. Climate, 23(18), 5002–5020. doi:10.1175/2010JCLI3586.1, 2010.
- 10 Meinshausen, M., Smith, S. J., Calvin, K., Daniel, J. S., Kainuma, M. L., Lamarque, J.-F., Matsumoto, K., Montzka, S. A., Raper, S. C., Riahi, K., Thomson, A., Velders, G. J., and van Vuuren, D. P.: The RCP greenhouse gas concentrations and their extensions from 1765 to 2300, *Clim. Change*, 109, 213–241, doi:10.1007/s10584-011-0156-z, 2011.
- Morgenstern, O., M.A. Giorgetta, K. Shibata, V. Eyring, D. Waugh, T. G. Shepherd, H. Akiyoshi,
 J. Austin, A. J. G. Baumgaertner, S. Bekki, P. Braesicke, C. Bruhl, M. P. Chipperfield, D. Cugnet, M. Dameris, S. Dhomse, S. M. Frith, H. Garny, A. Gettelman, S. C. Hardiman, M. I. Hegglin, P. Jockel, D. E. Kinnison, J. F. Lamarque, E. Mancini, E. Manzini, M. Marchand, M. Michou, T. Nakamura, J. E. Nielsen, D. Olivie, G. Pitari, D. A. Plummer, E. Rozanov, J. F. Scinocca, D. Smale, H. Teyssedre, M. Toohey, W. Tian, and Y. Yamashita Review of present-
- 20 generation stratospheric chemistry-climate models and associated external forcings, *J. Geophys. Res.*, 115, D00M02, doi:10.1029/2009JD013728, 2010.
 - Morgenstern, O., G. Zeng, N. L. Abraham, P. J. Telford, P. Braesicke, J. A. Pyle, S. C. Hardiman, F. M. O'Connor, and C. E. Johnson, Correction to "Impacts of climate change, ozone recovery, and increasing methane on surface ozone and the tropospheric oxidizing capacity", *J. Geophys. Res. Atmos.*, 119, 5028–5036, doi:10.1002/2014JD021515, 2014.
- *Res. Atmos.*, 119, 5028–5036, doi:10.1002/2014JD021515, 2014.
 Morgenstern, O., M.I. Hegglin, E. Rozanov, F.M. O'Connor, N.L. Abraham, H. Akiyoshi, A.T. Archibald, S. Bekki, N. Butchart, M.P. Chipperfield, M. Deushi, S.S. Dhomse, R.R. Garcia, S.C. Hardiman, L.W. Horowitz, P. Joeckel, B. Josse, D. Kinnison, M. Lin, E. Mancini, M.E. Manyin, M. Marchand, V. Marecal, M. Michou, L.D. Oman, G. Pitari, D.A. Plummer, L.E.
- 30 Revell, D. Saint-Martin, R. Schofield, A. Stenke, K. Stone, K. Sudo, T.Y. Tanaka, S. Tilmes, Y. Yamashita, K. Yoshida and G. Zeng: Review of the global models used within phase 1 of the Chemistry–Climate Model Initiative (CCMI), *Geosci. Model Dev.*, 10, 639-671, doi:10.5194/gmd-10-639-2017, 2017.
- Morgenstern, O., Akiyoshi, H., Yamashita, Y., Kinnison, D. E., Garcia, R. R., Plummer, D. A.,
 Scinocca, J., Zeng, G., Rozanov, E., Stenke, A., Revell, L. E., Pitari, G., Mancini, E., Di
 Genova, G., Dhomse, S. S., and Chipperfield, M. P.: Ozone sensitivity to varying greenhouse
 gases and ozone-depleting substances in CCMI-1 simulations, *Atmos. Chem. Phys.*, 18, 1091-1114, doi:10.5194/acp-18-1091-2018, 2018.
- Olsen, M. A., Schoeberl, M. R., and Nielsen, J. E.: Response of stratospheric circulation and
 stratosphere-troposphere exchange to changing sea surface temperatures, *J. Geophys. Res.*, 112, D16104, doi:10.1029/2006JD008012, 2007.

Oman, L., D. W. Waugh, S. R. Kawa, R. S. Stolarski, A. R. Douglass, and P. A. Newman, Mechanisms and feedbacks causing changes in upper stratospheric ozone in the 21st century, *J. Geophys. Res.*, 115, D05303, doi:10.1029/2009JD012397, 2010.

- 45 Oman, L. D., D. A. Plummer, D. W. Waugh, J. Austin, J. F. Scinocca, A. R. Douglass, R. J. Salawitch, T. Canty, H. Akiyoshi, S. Bekki, P. Braesicke, N. Butchart, M. P. Chipperfield, D. Cugnet, S. Dhomse, V. Eyring, S. Frith, S.C. Hardiman, D.E. Kinnison, J.-F. Lamarque, E. Mancini, M. Marchand, M. Michou, O. Morgenstern, T. Nakamura, J. E. Nielsen, D. Olivie, G. Pitari, J. Pyle, E. Rozanov, T. G. Shepherd, K. Shibata, R. S. Stolarski, H. Teyssedre, W. Tian,
- 50 Y. Yamashita, and J. R. Ziemke, Mult-model assessment of the factors driving stratospheric ozone evolution over the 21st century, *J. Geophys. Res.*, 115, D24306,

doi:10.1029/2010JD014362, 2010.

10

- Pitari, G., S. Palermi, and G. Visconti, Ozone response to a CO₂ doubling: Results from a stratospheric circulation model with heterogeneous chemistry, *J. Geophys. Res.*, 97, 5953–5962, 1992.
- 5 Polvani, L. M., L. Wang, V. Aquila, and D. W. Waugh, The impact of ozone-depleting substances on tropical upwelling, as revealed by the absence of lower-stratospheric cooling since the late 1990s, *Journal of Climate*, *30*(7), 2523–2534, 2017.

Polvani, L. M., M. Abalos, R. Garcia, D. Kinnison, and W. J. Randel, Significant weakening of Brewer-Dobson circulation trends over the 21st century as a consequence of the Montreal Protocol, *Geophys. Res. Lett.*, 45, 401-409, doi:10.1002/2017GL075345, 2018.

Portmann, R. W., and S. Solomon, Indirect radiative forcing of the ozone layer during the 21st century, *Geophys. Res. Lett.*, 34, L02813, doi:10.1029/2006GL028252, 2007.

Randeniya, L. K., Vohralik, P. F., Plumb, I. C., and Ryan, K. R.: Heterogeneous BrONO₂ hydrolysis: Effect on NO₂ columns and ozone at high latitudes in summer. *J. Geophys. Res.*, 102, 23543–23557, 1997.

- Randeniya, L. K., P. F. Vohralik, and I. C. Plumb, Stratospheric ozone depletion at northern midlatitudes in the 21st centrury: The importance of future concentrations of greenhouse gases nitrous oxide and methane, *Geophys. Res. Lett.*, 29, 1041, doi.org/10.1029/2001GL014295, 2002.
- 20 Ravishankara, A. R., Daniel, J. S., and Portmann, R. W.: Nitrous Oxide (N₂O): The dominant ozone-depleting substance emitted in the 21st century. *Science*, *326*(5949), 2009.
 - Revell, L. E., Bodeker, G. E., Huck, P. E., Williamson, B. E. and Rozanov, E.: The sensitivity of stratospheric ozone changes through the 21st century to N₂O and CH₄, *Atmos. Chem. Phys.*, 12, 11309-11317, 2012.
- 25 Revell, L. E., F. Tummon, R. J. Salawitch, A. Stenke, and T. Peter: The changing ozone depletion potential of N₂O in a future climate, *Geophys. Res. Lett.*, 42, 10,047–10,055, doi:10.1002/2015GL065702, 2015.

Revell, L. E., A. Stenke, E. Rozanov, W. Ball, S. Lossow, and T. Peter, The role of methane in projections of 21st century stratospheric water vapor, *Atmos. Chem. Phys.*, 16, 13067–13080, doi:10.5104/sep.16.12067.2016.2016

Rind, D., R. Suozzo, N. Balachandran, and M. Prather, Climate change and the mid- dle atmosphere. Part I: The doubled CO₂ climate, *J. Atmos. Sci.*, *47*(4), 475–494, 1990.

Rind, D., J. Lerner, and C. McLinden, Changes of tracer distributions in the doubled CO₂ climate. *J. Geophys. Res.*, 106, 28,061–28,080, 2001.

- 35 Rosenfeld, J. E., A. R. Douglass, and D. B. Considine, The impact of increasing carbon dioxide on ozone recovery, *J. Geophys. Res.*, 107, 4049, 10.1029/2001JD000824, 2002.
- Royer, J.-F., D. Cariolle, F. Chauvin, M. Deque, H. Douville, R.-M. Hu, S. Planton, A. Rascol, J.-L. Ricard, D.S.Y. Melia, F. Sevault, P. Simon, S. Somot, S. Tyteca, L. Terray, and S. Valcke, Simulation of climate changes during the 21st century including stratospheric ozone. *C. R.*

40 *Geoscience*, 334, 147-154, 2002.

Sander, S. P. et al., Chemical Kinetics and Photochemical Data for Use in Atmospheric Studies, Evaluation Number 17, JPL Publication 10-6, 2011.

Scinocca, J. F., Stephenson, D. B., Bailey, T. C., & Austin, J.: Estimates of past and future ozone trends from multimodel simulations using a flexible smoothing spline methodology, J. Geophys. Res., 115, doi:10.1029/2009JD0136222010, 2009.

- Geophys. Res., 115, doi:10.1029/2009JD0136222010, 2009.
 Shepherd, T. G. and Jonsson, A. I.: On the attribution of stratospheric ozone and temperature changes to changes in ozone- depleting substances and well-mixed greenhouse gases, *Atmos. Chem. Phys.*, 8, 1435–1444, https://doi.org/10.5194/acp-8-1435-2008, 2008.
- Shindell, D. T., Faluvegi, G., Koch, D. M., Schmidt, G. A., Unger, N., and Bauer, S. E.: Improved
 attribution of climate forcing to emissions, *Science*, 326, 716–718, doi:10.1126/science.1174760, 2009.

³⁰ doi:10.5194/acp-16-13067-2016, 2016.

Sigmond, M., Siegmund, P.C., Manzini, E., Kelder, H.: A simulation of the separate climate effects of middle atmospheric and tropospheric CO₂ doubling. *J Clim.*, 17(12), 2352–2367, 2004.

- Solomon, S., D. J. Ivy, D. E. Kinnison, M. J. Mills, R. R. Neely III, and A. Schmidt, Emergence of healing in the Antarctic ozone layer, *Science*, 353, 269-274, 2016.
- Strahan, S. E., and A. R. Douglass, Decline in Antarctic Ozone Depletion and Lower Stratospheric Chlorine Determined From Aura Microwave Limb Sounder Observations, *Geophys. Res. Lett.*, 45, 382-390, 10.1002/2017GL074830, 2018.
- Wales, P.A., et al., Stratospheric Injection of Brominated Very Short-Lived Substances: Aircraft
 Observations in the Western Pacific and Representation in Global Models, *J. Geophys. Res.*, doi:10.1029/2017JD027978, (in press), 2018.
 - Wang, W., Tian, W., Dhomse, S., Xie, F., Shu, J., and Austin, J.: Stratospheric ozone depletion from future nitrous oxide increases, *Atmos. Chem. Phys.*, 14, 12967-12982, doi:10.5194/acp-14-12967-2014, 2014.
- 15 Waters, J.W., et al., The Earth Observing System Microwave Limb Sounder (EOS MLS) on the Aura satellite, IEEE Trans. *Geosci. Remote Sens.*, 44, 1075–1092, doi:10.1109/TGRS.2006.873771, 2006.
 - Waugh, D. W. and Eyring, V.: Quantitative performance metrics for stratospheric-resolving chemistry-climate models, *Atmos. Chem. Phys.*, 8, 5699-5713, doi:10.5194/acp-8-5699-2008, 2008.
- 20

25

Waugh, D. W., Oman, L., Kawa, S. R., Stolarski, R. S., Pawson, S., Douglass, P. A. Newman, J. E. Nielsen, Impacts of climate change on stratospheric ozone recovery. *Geophys. Res. Lett.*, 36(3), doi:10.1029/2008GL036223, 2009.

- World Meteorological Organization, Scientific assessment of ozone depletion: 2006. Global Ozone Research and Monitoring Project Report No. 50, Geneva, Switzerland, 2007.
 - World Meteorological Organization, Scientific Assessment of Ozone Depletion: 2010. Global Ozone Research and Monitoring Project-Report No. 52, Geneva, Switzerland, 2011.
- 30 World Meteorological Organization, Scientific Assessment of Ozone Depletion: 2014. Global Ozone Research and Monitoring Project-Report No. 55, Geneva, Switzerland, 2014.
 - Zhang, J., W. Tian, M.P. Chipperfield, F. Xie and J. Huang, Persistent shift of the Arctic polar vortex towards the Eurasian continent in recent decades, *Nature Climate Change*, **6**, 1094-1099, doi:10.1038/nclimate3136, 2016.

35

World Meteorological Organization, Scientific assessment of ozone depletion: 2002. Global Ozone Research and Monitoring Project Report No. 47, Geneva, Switzerland, 2003.

Tables

Simulations	Period	GHGs	ODSs	SSTs / SICs	Bqckground & Volcanic Aerosols	Solar Variability	VSLS Bromine	QBO	Ozone and Aerosol precursors
REFERENCE SCENARIOS									
REF-C1	1960- 2010	OBS + CMIP5 updated to 2010	OBS (WMO- 2011)	OBS, HadlSST1	OBS Sulfate SAD & radius	Spectrally resolved irradiance, Proton ionization, Ap	YES ~5 pptv additional Bry	OBS or internally generated	OBS Based on Lamarque, (2010).
REF-C1SD	1980- 2010	REF-C1	REF-C1	Consistent with met fields.	REF-C1	REF-C1	REF-C1	Consistent with met fields.	REF-C1
REF-C2	1960- 2100	Obs + RCP6.0	REF-C1	Model SSTs	Obs and background for forecast period.	REF-C1, repeating SV in the future	REF-C1	Consistent with REF-C1 approach	REF-C1 until + RCP6.0 scn in the future
RCP SENSITIVITY									
SEN-C2- RCP26	2000- 2100	RCP2.6	REF-C2	Model SSTs (RCP2.6)	REF-C2	REF-C2	REF-C2	REF-C2	Based on RCP2.6 in the future
SEN-C2- RCP45	2000- 2100	RCP4.5	REF-C2	Model SSTs (RCP4.5)	REF-C2	REF-C2	REF-C2	REF-C2	Based on RCP4.5 in the future
SEN-C2- RCP85	2000- 2100	RCP8.5	REF-C2	Model SSTs (RCP8.5)	REF-C2	REF-C2	REF-C2	REF-C2	Based on RCP8.5 in the future
ODS and GHG SENSITIVITY									
SEN-C2-fODS	1960- 2100	REF-C2	Fixed ODSs at 1960	REF-C2	REF-C2	REF-C2	REF-C2	REF-C2	REF-C2
SEN-C2-fGHG	1960- 2100	Fixed GHGs at 1960	REF-C2	REF-C2	REF-C2	REF-C2	REF-C2	REF-C2	REF-C2
SINGLE FORCING									
SEN-C2-fCH4	1960- 2100	REF-C2, w/Fixed CH ₄ at 1960	REF-C2	REF-C2	REF-C2	REF-C2	REF-C2	REF-C2	REF-C2
SEN-C2-fN2O	1960- 2100	REF-C2, w/Fixed N ₂ O at 1960	REF-C2	REF-C2	REF-C2	REF-C2	REF-C2	REF-C2	REF-C2
SEN-C2- CH4RCP85	2000- 2100	REF-C2, w/CH4 from RCP8.5	REF-C2	REF-C2	REF-C2	REF-C2	REF-C2	REF-C2	REF-C2

Table 1. Definitions of CCMI scenarios. Adapted from *Eyring et al.*, (2013a).

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Table 2. CCMI simulations analysed in this study. The numbers indicate the number of realisations by each model for each simulation.

Model name	REF-C1	REF-C1SD	REF-C2	SEN-C2-fGHG	SEN-C2-fODS	SEN-C2-RCP26	SEN-C2-RCP45	SEN-C2-RCP85	SEN-C2-fCH4	SEN-C2-CH4RCP85	SEN-C2-fN2O	Total Simulations
										P85		0
ACCESS CCM	1		2	2	2							7
CCSRNIES MIROC3.2	3	1	2	1	1	1	1	1	1		1	13
CESM1 CAM4-CHEM	3		3									6
CESM1 WACCM	5	1	3	3	3		1	3	1	1	1	22
CHASER (MIROC-ESM)		1	1	1	1	1		1	1		1	8
СМАМ	3	1	1	1	1	1	1	1	1	1	1	13
CNRM-CM5-3	4	1	2									7
EMAC-L47	1	1	1									3
EMAC-L90	1	1	1									3
GEOSCCM	1		1						1	1	1	5
GFDL-CM3/AM3	1	1	1				3	1				7
HadGEM3-ES	1	1	1									3
IPSL-LMDZ-REPROBUS	1	1	1									3
MRI	1	1	1									3
NIWA-UKCA	3		5	3	2				1		1	15
SOCOL3	3		1			1	1	1	1	1	1	10
TOMCAT (CTM)	1	1										2
ULAQ CCM	3		3	1	1	1	1	1	1	1	1	14
UMSLIMCAT	1		1	1	1		1	1		1		7
UMUKCA-UCAM	1	1	2									4
Total realisations	38	13	33	13	12	5	9	10	8	6	8	155
Total models	19	13	19	8	8	5	7	8	8	6	8	

Table 3. Total column ozone (TCO) return dates to 1980 baseline from REF-C2 simulations using different averaging methods. Values in brackets indicate recovery dates based on either $1-\sigma$ standard deviation or 10^{th} and 90^{th} percentile estimates. The number *2100* in italics indicates that the estimated ozone uncertainty range has not returned to the 1980 values within the time range of the model simulation. The MMM for RCP 6.0 derived from the CMIP5 models (*Erying et al.*, 2013b) is shown in column two.

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	WMO ¹	CMIP5 ²	CCMI REF-C2 (this work)				
	(2011, 2014)	Eyring et al. (2013)	MMM	Median	MMM1S		
SH pole	2050	2046	2062	2061	2060		
(October)	(2045-2060)	(2040-2055)	(2051-2082)	(2042-2069)	(2055-2066)		
SH Mid-	2035	2041	2046	2046	2045		
latitudes	(2030-2040)	(2033-2046)	(2038-2053)	(2038-2071)	(2039-2050)		
Tropics	2042 (2028-)	N/A	2100 (2034-2100)	2058 (2013-2100)	2058 (2038-2100)		
NH Mid-	2021	2032	2033	2032	2032		
latitudes	(2017-2026)	(2026-2039)	(2011-2047)	(2010-2048)	(2020-2044)		
NH pole	2030	2028	2039	2034	2034		
(March)	(2025-2035)	(2020-2033)	(2021-2050)	(2011-2058)	(2025-2043)		
Near Global		2043	2045	2046	2047		
(60°S-60°N)		(2035-2050)	(2034-2064)	(2025-2100)	(2042-2051)		
Global	2032 (2027-2038)		2046 (2035-2058)	2048 (2040-2073)	2049 (2043-2055)		

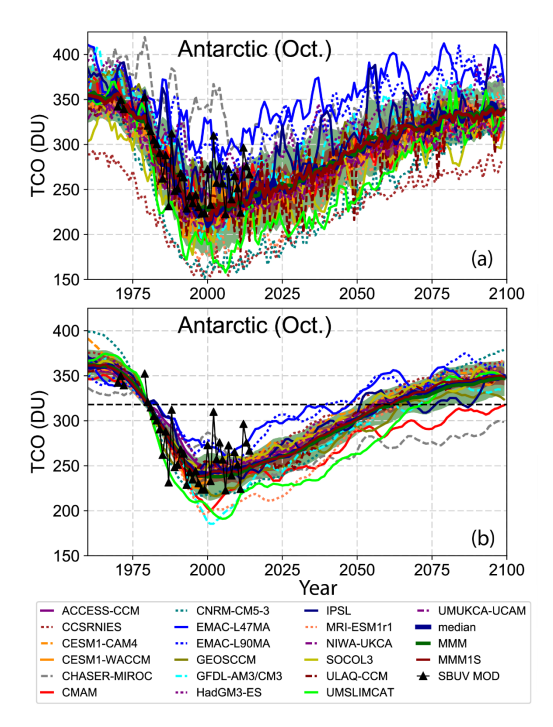
1. Based on CCMVal-2 model simulations (A1b GHG scenario) and reported in WMO (2011) and Table 2-5 and Figure 3-16 of WMO (2014).

10 2. Based on CMIP5 models used in Figure 2-23 of WMO (2014) with the point-wise 95% confidence interval. This approach to estimating uncertainties was also used in *Eyring et al.* (2013b), *Eyring et al.* (2010b) and Chapter 3 of WMO (2011).

Table 4. Stratospheric column ozone (SCO) return dates to 1980 baseline from various simulations. Values in brackets indicate the range of recovery dates based on 1-σ standard deviation. The number *2100* in italics indicates that the estimated ozone uncertainty range has not
returned to the 1980 values within the time range of the model simulation. Simulations starting in year 2000 (SEN-C2-CH4RCP85, SEN-C2-RCP45 and SEN-C2-RCP85) use 1980 baseline from REF-C2 simulations.

	REF-C2	SEN-C2- fGHG	SEN-C2- fN2O	SEN-C2- fCH4	SEN-C2- CH4RCP85	SEN-C2- RCP45	SEN-C2- RCP85
SH pole	2061	2059	2061	2062	2056	2052	2049
(October)	(2054- 2067)	(2052- 2064)	(2055- 2066)	(2058- 2068)	(2051- 2062)	(2046- 2060)	(2045- 2055)
SH Mid-	2045	2061	2043	2054	2042	2043	2042
latitudes	(2039- 2049)	(2055- 2066)	(2037- 2047)	(2050- 2060)	(2036- 2048)	(2036- 2049)	(2036- 2048)
	N/A	2049	2054	N/A	N/A	N/A	N/A
Tropics		(2043- 2068)	(2039- 2100)				
NH Mid-	2035	2060	2026	2043	2034	2035	2032
latitudes	(2028- 2045)	(2055- 2079)	(2022- 2033)	(2039- 2047)	(2027- 2043)	(2029- 2042)	(2027- 2044)
NH pole	2031	2058	2030	2037	2030	2028	2027
(March)	(2025- 2036)	(2040- 2064)	(2011- 2037)	(2027- 2041)	(2022- 2036)	(2022- 2044)	(2023- 2038)
Near	2058	2060	2040	2063	2044	2066	2056
Global	(2046- 2077)	(2056- 2075)	(2035- 2047)	(2056- 2088)	(2038- 2050)	(2044- 2100)	(2045- 2080)
Global	2049	2058	2042	2057	2043	2055	2047
	(2045- 2059)	(2052- 2064)	(2035- 2048)	(2051- 2074)	(2037- 2049)	(2044- 2078)	(2042- 2056)

Figures



- 5 Figure 1. (a) Total column ozone time series (DU) for Antarctic in October from 19 individual CCMs for the REF-C2 simulations along with observations from the Solar Backscatter Ultraviolet (SBUV) merged ozone dataset (MOD) (*Frith et al.*, 2017). The MMM, median (MedM) and MMM1S are shown with thick green, blue and red lines, respectively. The light blue shaded region indicates the 10th and 90th percentile range. Light green and red regions show
- 10 1-σ variability w.r.t. MMM and MMM1S lines, respectively. (b) Same as panel (a) but adjusted total ozone time series w.r.t. mean 1980-1984 observations and after application of 10-point boxcar smoothing. The dashed black line indicates 1980 reference value.

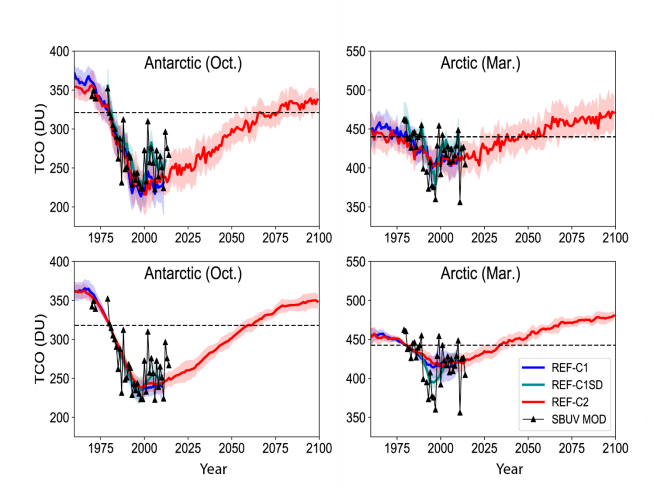


Figure 2. MMM1S total column ozone time series (DU) from REF-C1 (blue), REF-C1SD (dark
cyan) and REF-C2 (red) simulations for the (left) SH polar (October) and (right) NH polar (March) regions. The dashed black lines show the 1980 reference value for each latitude band. The shaded regions show 1-σ variability w.r.t. the MMM1S lines of the same colour. The top row shows the unadjusted modelled values and the bottom row shows the time series adjusted w.r.t. mean 1980-1984 observations and after application of a 10-point boxcar smoothing. Also shown are the merged SBUV observations.

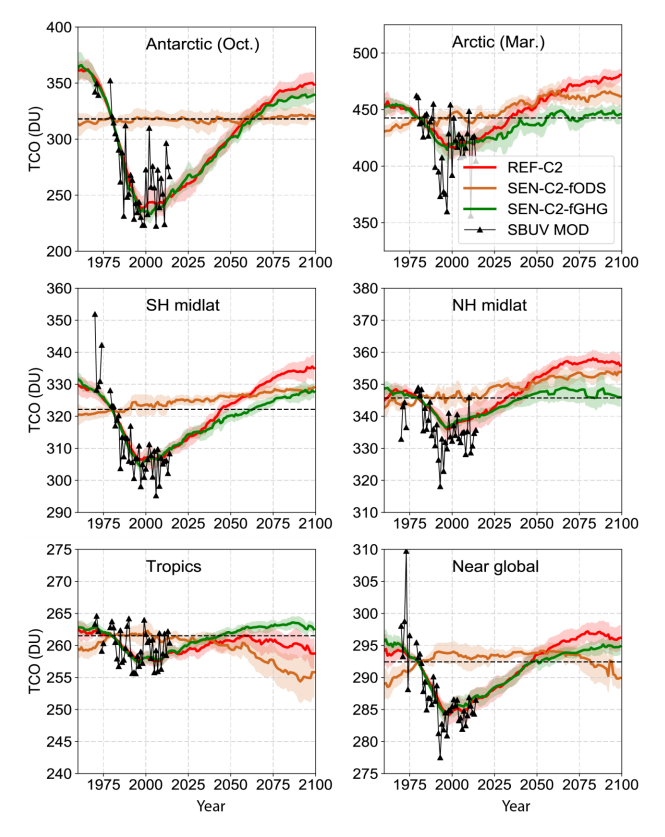


Figure 3. MMM1S total column ozone time series (DU) from REF-C2 (red), SEN-C2-fGHG
5 (dark green), and SEN-C2-fODS (brown) simulations for five latitudinal bands and the near-global (60°S-60°N) mean (see main text). The dashed black lines show the 1980 reference value for each latitude band. Also shown are the merged SBUV observations.

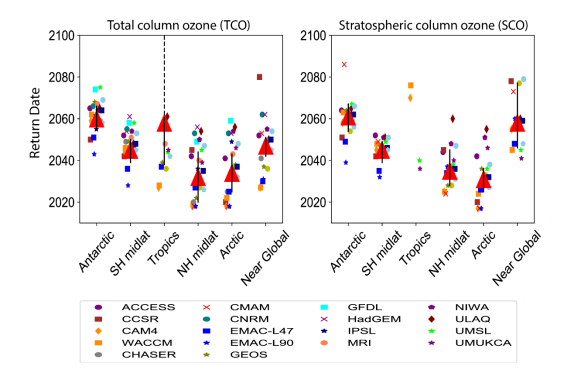


Figure 4. Estimated MMM1S return dates (red triangles) from the REF-C2 simulations for (left)
total column ozone (TCO) and (right) stratospheric column ozone (SCO) for different latitude bands. The estimated 1-σ uncertainties are shown with vertical black lines. Estimates for individual models are shown with coloured dots. Some individual models do not predict a return of column ozone in the tropics and so this uncertainty is indicated by a dashed line.

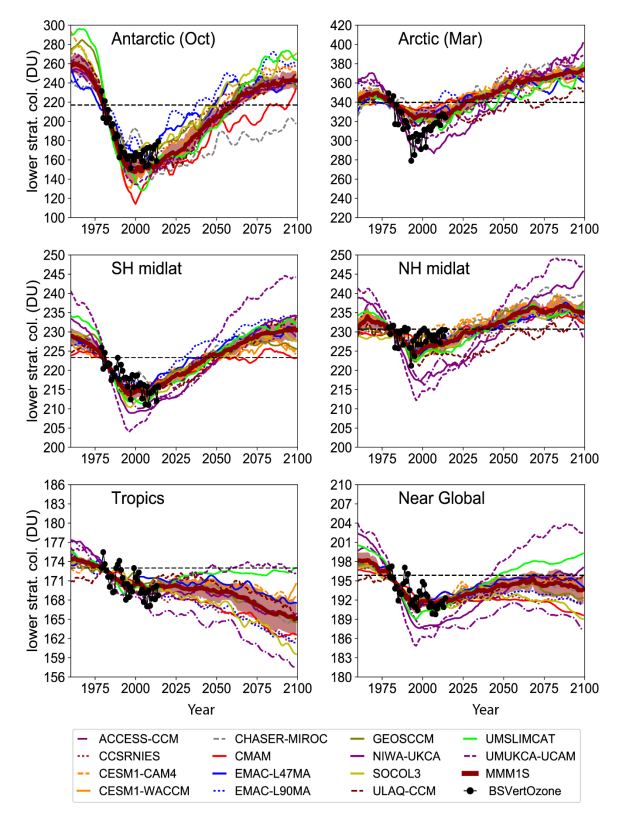


Figure 5. Evolution of partial column ozone (DU) for the lower stratosphere (tropopause - 10
5 hPa) from the REF-C2 simulations from 14 individual models, along with the MMM1S. Also shown are estimates of the partial column from the Bodeker Scientific Vertical Ozone (BSVertOzone) database, which is based on a compilation of satellite, balloon and ground-based measurements (*Bodeker et al.*, 2013).

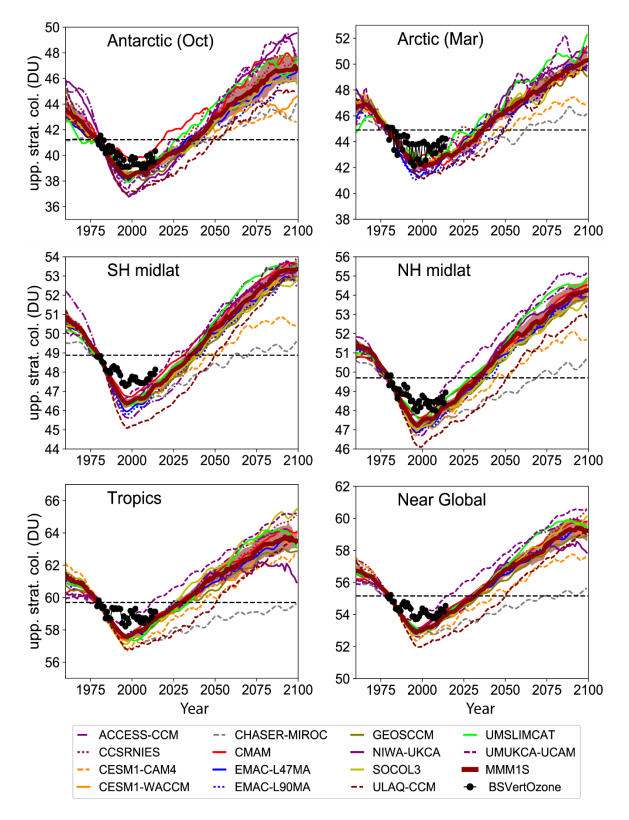


Figure 6. As **Figure 5** but for the upper stratosphere (≥ 10 hPa).

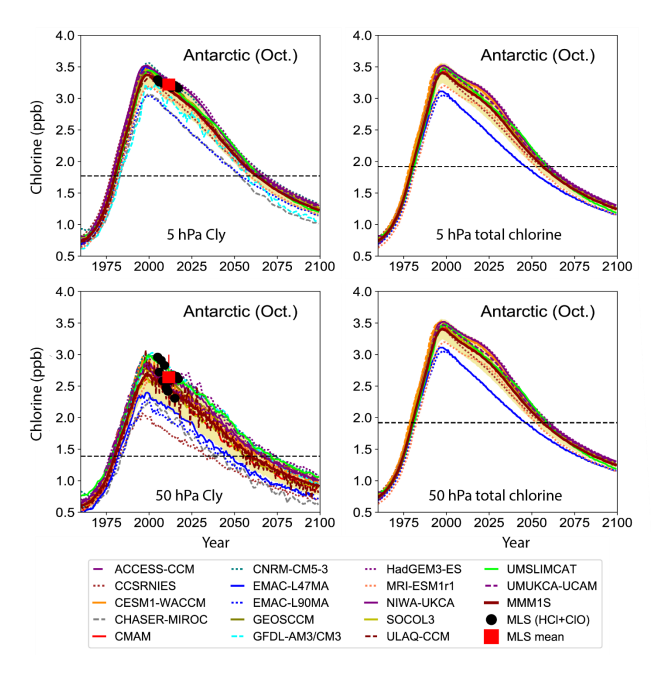


Figure 7. Evolution of (left) inorganic chlorine (Cly, ppb) and (right) total (organic + inorganic)
5 chlorine from the REF-C2 simulations for 17 individual models and the MMM1S over the Antarctic in October at (top) 5 hPa and (bottom) 50 hPa. The dashed black lines show the 1980 reference value. Also shown in the left panels are observed October mean values of the sum of HCl and ClO from version 4 of the Microwave Limb Sounder (MLS) data (*Waters et al.*, 2006; *Livesey et al.*, 2017) from 2005 to 2017 (black dots) and the mean value over that period (red

10 square). Note that not all models are plotted in the right-hand panels.

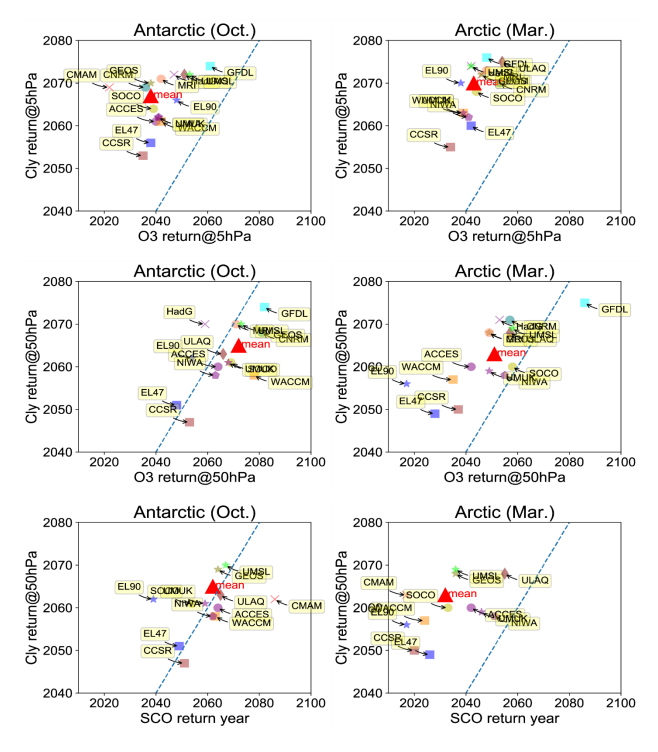


Figure 8. Correlation plots of ozone return dates against Cly return dates for (left) the Antarctic
and (right) the Arctic from REF-C2 simulations for individual models and the MMM1S at (top)
hPa, (middle) 50 hPa and (bottom) stratospheric column (SCO). The red triangle is the multimodel mean. The dashed blue line is the 1:1 line between Cly and ozone return dates. The model symbols are the same as those used in Figure 4.

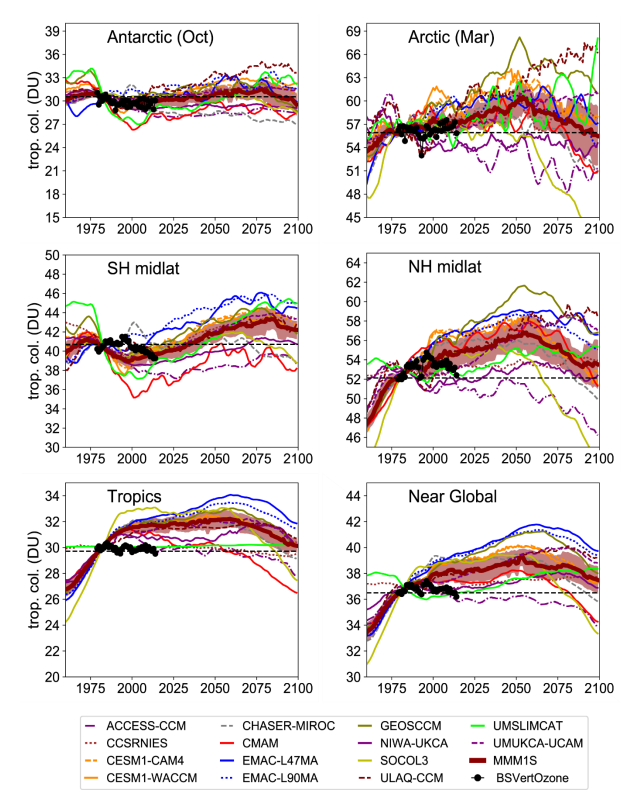


Figure 9. Evolution of tropospheric partial column ozone (DU) (surface - tropopause) from 14
individual models and the MMM1S for the REF-C2 simulations. Also shown is the tropospheric partial column ozone derived from BSVertOzone data.

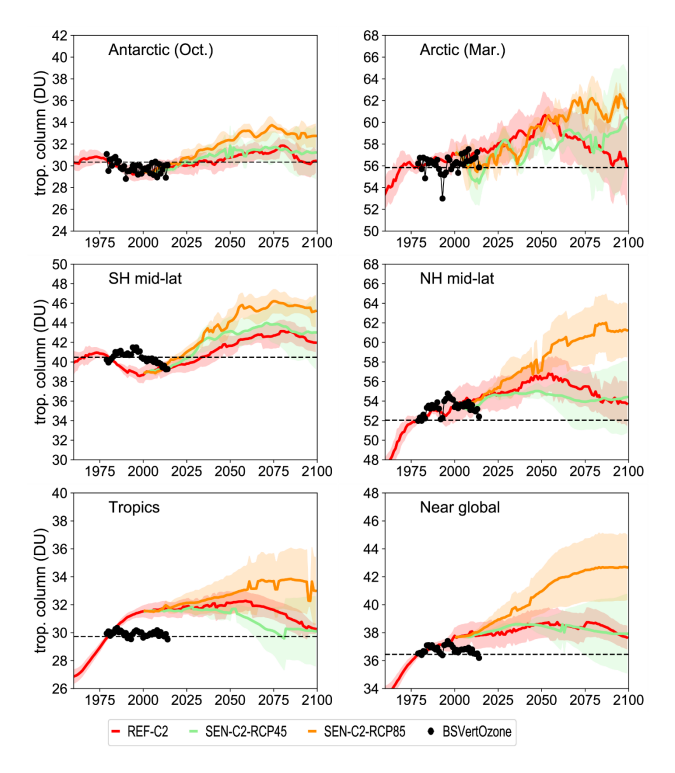


Figure 10. Evolution of tropospheric partial column ozone (DU) (surface-tropopause) MMM1S
for REF-C2 and the RCP scenarios SEN-C2-RCP45 and SEN-C2-RCP85. Also shown is the tropospheric partial column ozone derived from BSVertOzone data.

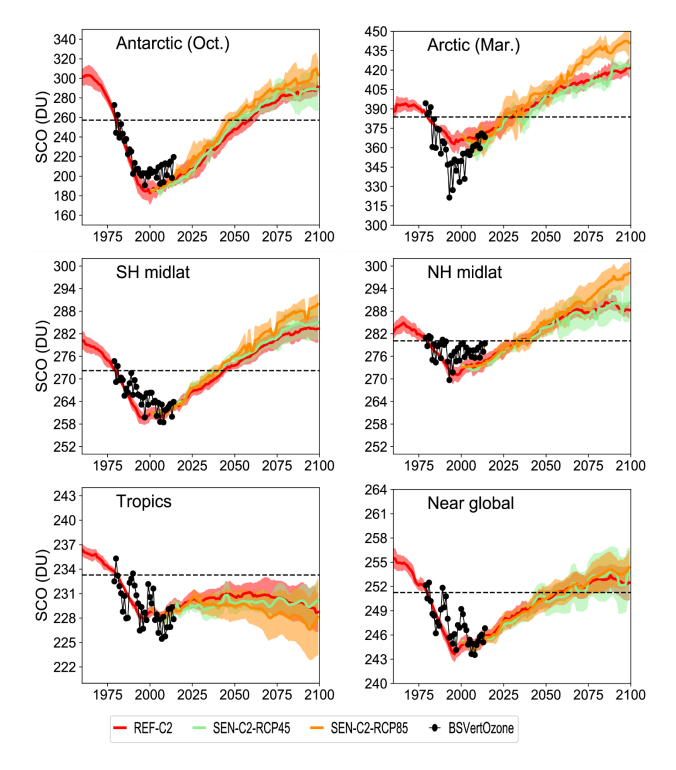


Figure 11. As Figure 10 but for stratospheric column ozone (SCO).

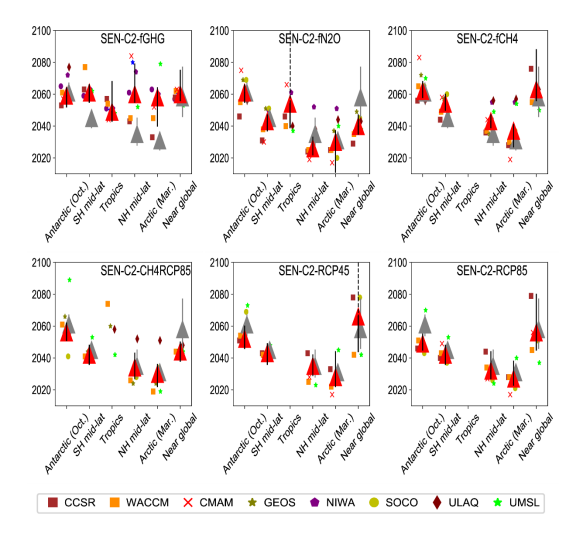


Figure 12. Estimated MMM1S return dates (red triangles) from stratospheric column ozone time
series for (top) SEN-C2-fGHG, SEN-C2-fN2O and SEN-C2-fCH4 and (bottom) SEN-C2-CH4RCP85, SEN-C2-RCP45 and SEN-C2-RCP85. Estimated uncertainties are shown with
vertical black lines. Grey triangles indicate SCO return dates from REF-C2 (Figure 4).
Estimates for individual models are shown with coloured dots. Points with return dates (and uncertainties) that are greater than year 2100 are not shown.

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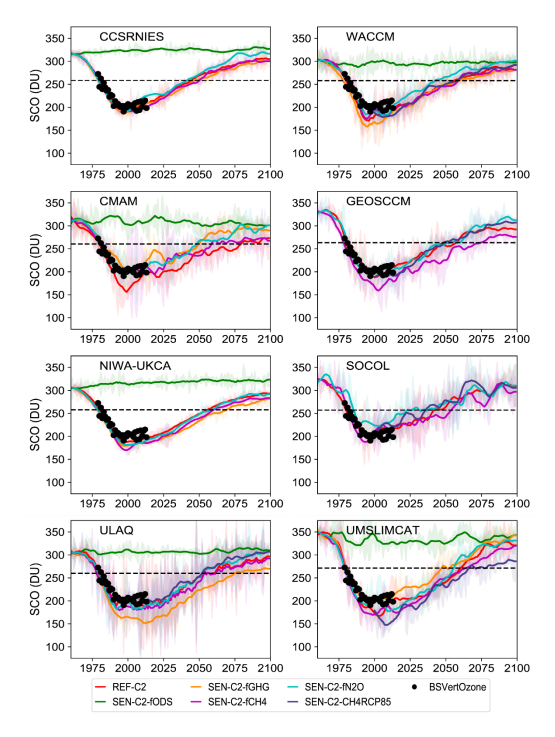


Figure 13. Evolution of Antarctic October mean stratospheric column ozone (DU) from 8

- 5 selected models for the REF-C2, SEN-C2-fCH4, SEN-C2-fN2O, SEN-C2-fODS, SEN-C2-fGHG and SEN-C2-CH4RCP85 simulations. Each panel gives the name of the model shown. The solid lines are the 10-year smoothed SCO for a given simulation. The shading on the lines shows either the standard deviation from an ensemble of realisations from that model, or the deviation from a 3-box smoothed line if only 1 realisation is available. Note that not all models
- 10 have performed all simulations. Also shown in each panel is the SCO derived from BSVertOzone data (filled black circles).

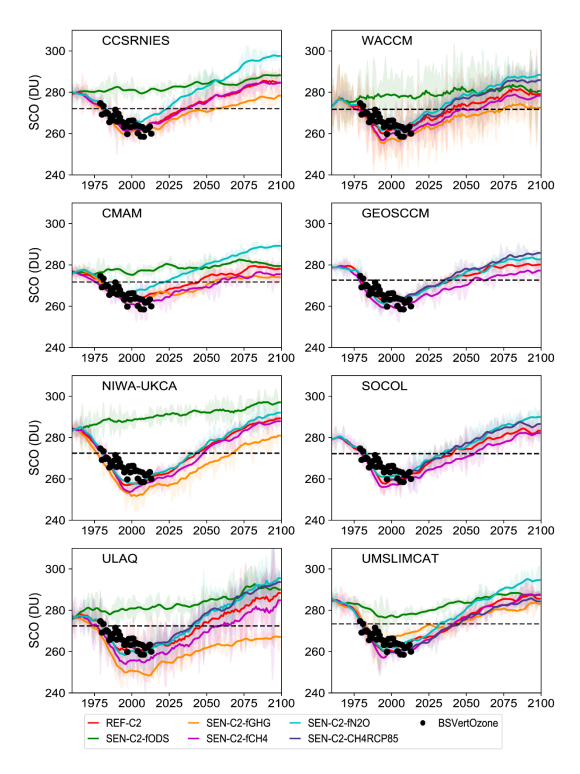


Figure 14. As Figure 13 but for SH mid-latitude annual mean.