Detectability of the impacts of ozone-depleting substances and greenhouse gases upon stratospheric ozone accounting for nonlinearities in historical forcings

Justin Bandoro¹, Susan Solomon¹, Benjamin D. Santer², Douglas E. Kinnison³, and Michael J. Mills³

¹Department of Earth, Atmospheric, and Planetary Sciences, Massachusetts Institute of Technology, Cambridge, MA 02139, USA
²Program for Climate Model Diagnosis and Intercomparison (PCMDI), Lawrence Livermore National Laboratory, Livermore, CA 94550, USA
³Atmospheric Chemistry Observations and Modeling Laboratory, National Center for Atmospheric Research, Boulder, CO 80307, USA

Correspondence: Justin Bandoro (jbandoro@mit.edu)

Received: 26 June 2017 – Discussion started: 14 July 2017
Revised: 6 November 2017 – Accepted: 24 November 2017 – Published: 5 January 2018

Abstract. We perform a formal attribution study of upper- and lower-stratospheric ozone changes using observations together with simulations from the Whole Atmosphere Community Climate Model. Historical model simulations were used to estimate the zonal-mean response patterns ("fingerprints") to combined forcing by ozone-depleting substances (ODSs) and well-mixed greenhouse gases (GHGs), as well as to the individual forcing by each factor. Trends in the similarity between the searched-for fingerprints and homogenized observations of stratospheric ozone were compared to trends in pattern similarity between the fingerprints and the internally and naturally generated variability inferred from long control runs. This yields estimated signal-to-noise (S/N) ratios for each of the three fingerprints (ODS, GHG, and ODS + GHG). In both the upper stratosphere (defined in this paper as 1 to 10 hPa) and lower stratosphere (40 to 100 hPa), the spatial fingerprints of the ODS + GHG and ODS-only patterns were consistently detectable not only during the era of maximum ozone depletion but also throughout the observational record (1984–2016). We also develop a fingerprint attribution method to account for forcings whose time evolutions are markedly nonlinear over the observational record. When the nonlinearity of the time evolution of the ODS and ODS + GHG signals is accounted for, we find that the S/N ratios obtained with the stratospheric ODS and ODS + GHG fingerprints are enhanced relative to standard linear trend analysis. Use of the nonlinear signal detection method also reduces the detection time – the estimate of the date at which ODS and GHG impacts on ozone can be formally identified. Furthermore, by explicitly considering nonlinear signal evolution, the complete observational record can be used in the S/N analysis, without applying piecewise linear regression and introducing arbitrary break points. The GHG-driven fingerprint of ozone changes was not statistically identifiable in either the upper- or lower-stratospheric SWOOSH data, irrespective of the signal detection method used. In the WACCM simulations of future climate change, the GHG signal is statistically identifiable between 2020 and 2030. Our findings demonstrate the importance of continued stratospheric ozone monitoring to improve estimates of the contributions of ODS and GHG forcing to global changes in stratospheric ozone.

1 Introduction

Climate change detection and attribution (D&A) studies seek to identify and formally quantify an anthropogenic component of change in observed climate data. Formal identification of an anthropogenic climate change “fingerprint” has been successfully achieved with observations of atmosphere and ocean temperatures, sea level, ocean acidity, various components of the water cycle and the cryosphere, and certain climate extremes (Bindoff et al., 2013). To date, how-
ever, few formal D&A methods have been applied in studies involving stratospheric ozone (see Gillett et al., 2011, for one exception to this). There is evidence that stratospheric ozone is transitioning from an era of widespread and readily detectable depletion (linked to changes in anthropogenic chlorofluorocarbons) to an era characterized by early signs of recovery or healing (Solomon et al., 2016). Our motivation for this work is to determine whether formal D&A methods can provide a more confident and quantitative attribution of ozone depletion and recovery signals.

Global changes in the physical climate system are driven by both internal variability and external influences (Hegerl et al., 2007; Karl et al., 2006). Internal variability is generated through complex interactions of the coupled atmosphere–ocean system. External influences include human-caused changes in well-mixed greenhouse gases (GHGs), ozone-depleting substances (ODSs), and other radiative forcing agents, as well as natural fluctuations in solar irradiance and volcanic aerosols. Past D&A studies have found that each of these external influences has a unique fingerprint in the detailed zonal-mean latitude–altitude pattern of temperature change (Hansen et al., 2005; Karoly et al., 1994; Santer et al., 1996a; Tett et al., 1996; Thorne et al., 2002; Vinnikov et al., 1996). The use of such profiles of atmospheric temperature change has proved particularly useful in separating human, solar, and volcanic influences on climate, and in discriminating between externally forced signals and internal variability. In this study, we use the latitude/altitude patterns of both upper- and lower-stratospheric ozone change in response to individual and combined anthropogenic forcings to understand the relative detectability of ODS and GHG signals in observations.

Stratospheric ozone depletion has been a significant international concern since it was first recognized as a consequence of anthropogenic emissions of ODSs (Molina and Rowland, 1974). Following implementation of the Montreal Protocol, there was a decline in emissions of ODSs and a consequent decrease in halogen-containing compounds in the stratosphere. In the 14th World Meteorological Organization (WMO) Scientific Assessment of Ozone Depletion (WMO, 2014), a major topic of interest was the statistical significance of ozone trends over the last decade, and the extent to which “ozone recovery” is taking place. Assessing the statistical significance of observed ozone trends does not yield definitive information on the causes of trends. In addition to determining whether observed negative or positive trends in ozone are unusually large or small relative to model-based estimates of unforced trends in ozone, it is also important to understand and quantify the contributions of different climate forcings to the observed ozone changes and to assess whether the agreement between an externally forced fingerprint and observations could have been obtained by natural causes alone.

There is a clear scientific consensus that man-made chlorofluorocarbons were the dominant driver of global stratospheric ozone decline from the 1970s to 2000s (e.g., Solomon, 1999, and references therein). It is also well established that increasing GHGs contribute to a cooling of the middle to upper stratosphere (Boville, 1986; Fels, 1980). Colder temperatures result in a slowing down of gas-phase catalytic reactions that destroy ozone in the upper stratosphere, which in the absence of any other changes, leads to ozone increases. Thus, both decreasing ODS and increasing GHG concentrations can act to increase upper-stratospheric ozone. Understanding the extent to which GHG increases may confound the attribution of ozone changes due to ODSs is essential to the identification of ozone recovery at these altitudes (see WMO, 2014).

Another potential confounding factor is change in the Brewer–Dobson circulation (BDC), the meridional overturning circulation in the middle atmosphere that transports trace gases from the tropics to the poles (Brewer, 1949; Dobson, 1956). Previous studies have found that a strengthening of the BDC results in lower ozone concentrations in the tropical lower stratosphere and an increase in concentrations in the extratropics, and modeling studies have suggested that increases in GHG concentrations lead to a strengthening of the BDC (Eyring et al., 2010; Fleming et al., 2011; Garcia et al., 2008; Gillett et al., 2011; Oman et al., 2010). In contrast, a recent study by Polvani et al. (2017) found that trends in ODSs, and not in GHG levels, have been the primary driver of trends in tropical upwelling. This complicates the problem of attributing ozone changes that are primarily BDC-related since both GHG and ODS forcing may be implicated in driving changes in the BDC.

Another forcing that can affect the concentration of ozone in the stratosphere is the solar cycle. The total solar irradiance changes by about 0.1% during the 11-year solar cycle; however, UV radiation can change by about 4–8% (Lean, 2000). Stratospheric ozone changes during the solar cycle have been well documented (e.g., Merkel et al., 2011), with the largest percentage change in ozone between solar maximum and minimum occurring between 30 and 45 km. Stratospheric ozone is also influenced by internal variability, primarily via the quasi-biennial oscillation (QBO). The effect of the QBO on upper-stratospheric ozone has been shown to be small compared to its larger signal in interannual ozone variability in the lower and middle stratosphere (Hasebe, 1994; Zawodny and McCormick, 1991). These natural and internal changes must also be considered in attributing ozone changes to specific causes.

Most D&A studies have been focused on the attribution of changes in the climate system to anthropogenic GHG forcing, which has been increasing linearly since the early 20th century. In contrast, ODS forcing has had a unique nonlinear time evolution because of the implementation of the Montreal Protocol. For analyses spanning both the depletion and ozone recovery periods, purely linear trends are inadequate for capturing the more complex nonlinear evolution of ozone change in certain parts of the stratosphere (as we will...
show later). Many ozone trend studies address this nonlinear behavior by performing piecewise linear regressions with a break point around 1997 (Bourassa et al., 2014; Chehade et al., 2014; Jones et al., 2009; Kyrölä et al., 2013; Laine et al., 2014). The slopes of the piecewise trends are not constrained by physical and chemical considerations and are typically arbitrarily chosen to enhance the slopes for each of the eras. Another approach is to quantify changes over the entire record using regression to a nonlinear proxy, such as the halogen loading in the stratosphere, described by the equivalent effective stratospheric chlorine (EESC; see, e.g., Newchurch, 2003). Many studies have employed EESC-based regression to quantify trends in ozone (e.g. Langematz et al., 2016; Stolarski et al., 2006a; Wohltmann et al., 2007). This approach utilizes the entire observational record, with no arbitrary start or end points. While EESC-based regression offers a number of advantages for the detection of nonlinear ODS-driven ozone changes, it also has certain disadvantages (Kuttippurath et al., 2015). In Sect. 4, we use EESC-based regression to address the problem of nonlinear ODS evolution over the full observational record. Our D&A approach also accounts for the much smaller temporal changes in the GHG forcing time series.

To the best of the authors’ knowledge, only one previous study has considered the formal attribution of anthropogenic emissions to observed stratospheric ozone changes: Gillett et al. (2011). Their investigation examined zonal-mean solar backscatter ultraviolet (SBUV) ozone measurements (Mclinden et al., 2009) over the 27-year period from 1979 to 2005 in the middle and upper stratosphere (50–1 hPa). Model results were taken from simulations performed as part of the Chemistry–Climate Model Validation Activity (CCM-Val; Eyring et al., 2010). Gillett et al. analyzed multi-model simulations performed with GHGs only, combined anthropogenic factors only, and combined anthropogenic and natural external forcings. The ozone response to changes in ODSs and the response to changes in natural external forcings were estimated through subtraction of simulations. Gillett et al. (2011) used a standard space–time optimal regression methodology for signal detection and attribution (see, e.g., Allen and Tett, 1999). In this approach, the observations are modeled as a linear sum of simulated responses (fingerprints) to individual forcings, with each response scaled by an estimated regression coefficient (expressing the strength of the space–time response pattern in observations). A regression coefficient significantly greater than zero indicates a detectable response to the forcing, and a coefficient close to unity signifies that simulated and observed responses are similar in magnitude (attribution). The underlying premise here is that the observations can be well represented by a linear combination of the input model signal response fields and an additive noise term due to internal climate variability. It is also assumed that the response patterns to different individual forcings are statistically distinct (i.e., are separable) and that the sum of the individual responses is equivalent to the response obtained when all forcings are varied simultaneously.

Gillett et al. (2011) were unable to separate the individual ODS and GHG responses in the SBUV ozone data, but they found a clear combined anthropogenic signal that was consistent with observations. In their study, the authors hypothesized that the difficulty in separating the individual ODS and GHG responses was due to multiple factors: the limited ensemble size, and the degeneracy between the patterns of stratospheric ozone response to ODS and GHG forcing. Since the optimal regression methodology combines the spatial and temporal responses into a single space–time vector, it was not possible to determine whether the degeneracy between the ODS and GHG response patterns was primarily due to spatial similarity or to similarity in temporal evolution.

We use a different approach here to understand and quantify the relative contributions (and detectability) of the ozone responses to ODS and GHG forcing. Rather than combining spatial pattern and time evolution information in a single vector, we use pattern correlations to assess the time evolution of the spatial similarity between time-invariant fingerprints and time sequences of (1) observed ozone patterns and (2) model-based estimates of the natural variability in ozone. Such methods rely on some form of spatial covariance statistic (e.g., Santer et al., 1993, 1995) and may involve rotation of the fingerprint in a low-noise direction in order to optimize signal-to-noise ratios (Hasselmann, 1993; Hegerl et al., 1996). The conventional strategy in this method is to search for a long-term, positive trend in the pattern correspondence statistic, which would indicate an increasing expression of the searched-for signal in the observations. The underlying assumption here is that the spatial pattern of the fingerprint does not change markedly as a function of time, which is a reasonable assumption for historical GHG forcing, but is probably unreasonable for ODS forcing (see Solomon et al., 2016).

Our study differs from that of Gillett et al. (2011) in a number of ways. In addition to using a different D&A method and a longer ozone data set, we are also addressing different statistical questions. First, we seek to determine whether the observed changes in stratospheric ozone are unusual relative to estimates of both internal climate variability and total natural variability (Santer et al., 2013b). Second, we consider the time evolution of signal-to-noise ($S/N$) ratios, both for the individual fingerprints of ODS-forced and GHG-forced ozone changes, as well as for the ozone fingerprint arising from combined ODS + GHG forcing. This allows us to determine the detection time – the time at which $S/N$ exceeds and remains consistently above some stipulated significance level. Third, we calculate $S/N$ ratios separately for ozone changes in the upper and lower stratosphere rather than for a single region only (the 1 to 50 hPa region considered by Gillett et al., 2011). The reasons for this decision are explained below. Fourth, because we do not combine spatial
and temporal information, the spatial aspects of $S/N$ behavior are easier to evaluate (e.g., in terms of the pattern similarity between the searched-for fingerprints and the observations).

There are several disadvantages of our selected approach. Unlike Gillett et al. (2011), who performed a multi-model analysis, we rely on a single model only and are unable to evaluate the sensitivity of our results to possible errors in the model-based estimates of internal variability and the response to external forcing (North et al., 1998). Additionally, since we do not explicitly incorporate time evolution information in the searched-for fingerprints, pronounced differences in the time evolution of the ozone responses to ODS and GHG forcing cannot be used in the separation of these responses (at least not in the “linear trend” representation of these signals.

As noted above, we perform separate D&A analyses for ozone changes in the upper and lower stratosphere. Our premise is that different dominant processes control the changes in ozone concentrations in these two regions. In the lower stratosphere, particularly in the tropics and extratropics, ozone concentrations are affected by direct transport, both through interannual variability and via changes in the BDC (Shindell and Grewe, 2002). A further influence on lower-stratospheric ozone concentrations occurs by means of heterogeneous chemistry on the surfaces of polar stratospheric clouds and volcanic aerosols; the heterogeneous reactions are extremely sensitive to temperature changes (Solomon, 1999). The ozone chemical lifetime is on the order of weeks in the lower stratosphere in the Antarctic and on the order of months at midlatitudes.

In the upper stratosphere, the processes controlling ozone abundance are quite different; the photochemical lifetime of ozone is on the order of hours, so that direct transport of ozone is not important. The local concentrations of upper-stratospheric ozone are determined by gas-phase chemistry, which is well understood and well replicated in current chemistry–climate models (Austin et al., 2002). Transport of other GHGs (such as methane) can indirectly affect ozone concentrations in the upper stratosphere (Solomon and Garcia, 1984). The anticipated response from GHG forcing radiatively cools the upper stratosphere uniformly with latitude (Aquilla et al., 2016), increasing ozone concentrations, and potentially impacting atmospheric circulation through the BDC. A strengthening of the BDC would result in lower-stratospheric ozone decreases in the tropics and increases in the extratropics.

Based on this understanding from atmospheric physics and chemistry, we expect that different forcings should yield distinctly different latitudinal and vertical patterns of change. Below, we examine in detail the use of these patterns for attribution of the causes of ozone changes in the upper and lower stratosphere.

2 Model simulations and observed ozone data sets

The model used for all simulations analyzed in this study is version 1 of the Community Earth System Model, with version 4 of the Whole Atmosphere Community Climate Model as the atmospheric component (CESM1(WACCMM); see Marsh et al., 2013). WACCMM is a coupled chemistry–climate model that extends from the Earth’s surface to the lower thermosphere. WACCMM’s representation of heterogeneous chemistry has been shown to be in broad agreement with observations of polar ozone and related chemical species (Solomon et al., 2015). A wide range of experiments were performed with WACCMM:

1. Simulations that follow the experimental design of the Chemistry–Climate Model Intercomparison (CCMI) REFC1 experiment (Morgenstern et al., 2017), with volcanic aerosols prescribed according to CCMI were carried out (Arfeuille et al., 2013). These simulations include coupled chemistry and dynamics (Marsh et al., 2013) with sea surface temperatures prescribed according to observations for the period January 1955–December 2014. Concentrations of ODSs and GHGs vary over time, as specified in the REFC1 scenario. A set of five ensemble members was available, each with identical forcings but starting from slightly differing initial conditions. We refer to these simulations subsequently as ALL1.

2. Free-running simulations that follow the same REFC1 scenario as in ALL1, with the same atmospheric coupled chemistry and dynamics, but including an interactive fully coupled ocean component were carried out. These simulations begin in January 1960 and extend through December 2099. One set of runs includes both historical time-varying GHG and ODS concentrations (referred to herein as ALL2), while another keeps GHG concentrations fixed at their 1960 conditions but allows the evolution of time-varying ODS concentrations (referred to as FIXED GHG1960). A third set complements the second with ODS concentrations fixed at 1960 conditions but with time-varying GHG concentrations (FIXED ODS1960). Each of the aforementioned sets of experiments contains three ensemble members that were performed with slightly different initial conditions. FIXED GHG1960 and FIXED ODS1960 follow the convention of the CCMI REFC2 experiments (Eyring et al., 2013). It should be noted that the time-varying concentrations of ODSs in FIXED GHG1960 are radiatively active.

All sets of simulations in (1) and (2) include identical historical external forcings from solar insolation changes and prescribed CCMI volcanic aerosol evolution. As in many other general circulation models (see Xue et al., 2012), the QBO is imposed by nudging the tropical stratospheric zonal winds
to observed winds. From the sets of fully coupled simulations, we were able to estimate responses to GHGs alone (referred to herein as GHGonly) and ODSs (referred to herein as ODSonly) by independently differencing the individual ensemble anomalies from the FIXED GHG1960 and ODS1960 simulations from the ALL2 ensemble mean. In addition, we were able to isolate the combined response to volcanic aerosols, the solar cycle, and the QBO by differencing the sum of ensemble mean anomalies of the FIXED GHG1960 and ODS1960 simulations from the ensemble anomalies of ALL2. We designate this as the natural historical signal (referred to herein as NAT-h). The underlying assumption for our method of estimating the GHGonly, ODSonly and NAT-h responses is that the stratospheric ozone responses to individual forcings add linearly. Eyring et al. (2010) found this assumption to hold in the stratosphere, but with some departures from linearity in the tropical total column ozone. In Fig. S1 in the Supplement we show that, in the absence of the NAT-h response, the global-mean sum of the GHG and ODS responses in the upper- and lower-stratospheric ozone are close to the all forcing simulation.

There is reason to suspect that the NAT-h signal may not thoroughly represent the ozone response to volcanic eruptions in certain regions of the lower stratosphere. In the low-halogen-loading stratospheric conditions of FIXED ODS1960, increases in mid-stratospheric ozone following a large volcanic eruption are expected as the loss of ozone at these altitudes is dominated by odd nitrogen, due to enhanced halogen loading conditions of FIXED GHG1960 and ALL2. We designate this as the natural historical signal (referred to herein as NAT-h). The underlying assumption for our method of estimating the GHGonly, ODSonly and NAT-h responses is that the stratospheric ozone responses to individual forcings add linearly. Eyring et al. (2010) found this assumption to hold in the stratosphere, but with some departures from linearity in the tropical total column ozone. In Fig. S1 in the Supplement we show that, in the absence of the NAT-h response, the global-mean sum of the GHG and ODS responses in the upper- and lower-stratospheric ozone are close to the all forcing simulation.

3. Free-running fully coupled simulations of the last millennium (years 850 to 1850), in which only the estimated external forcing from historical solar variability is specified. This simulation (hereafter referred to as NAT) provides insight into the range of fluctuations in stratospheric ozone arising from the combined effects of the solar cycle and internal variability (other than the QBO; there is no nudging to observed stratospheric winds in this simulation).

4. Free-running fully coupled preindustrial control run length of 200 years. Unlike (3), there are no temporal changes in solar forcing and all variability is intrinsic to the climate system. This is the same experiment used by Marsh et al. (2013) to provide an estimate of internal climate variability alone and is referred to herein as CTL.

We rely primarily on the Stratospheric Water and Ozone Homogenized (SWOOSH) database (Davis et al., 2016) for observational estimates of stratospheric ozone changes. SWOOSH includes vertically resolved ozone from a subset of the limb profiling satellite instruments operating from 1984 to present day. SWOOSH’s ozone product is a gridded monthly-mean zonal-mean time series of mixing ratios on pressure levels ranging from 1 to 316 hPa. A key aspect of this merged product is that the source records are homogenized to account for inter-satellite biases and to reduce the impact of non-climatic artifacts. In this study, we use SWOOSH version 2.6 with a latitudinal resolution of 10°.

An advantage of the SWOOSH data set is that the vertical range enables investigation of the full stratosphere from January 1984 to December 2016 (Tummon et al., 2015) found that SWOOSH ozone concentrations from 1984 to 2011 were within ±10% of other data sets throughout much of the stratosphere, with the largest differences in the lower stratosphere.

For the signal-to-noise analysis, it was necessary to transform the ozone data from both observational grids and the model grid of WACCM to a common grid. The common horizontal grid chosen was SWOOSH’s 10° latitude zonal-mean grid from 85°S to 85°N. Transformation to a relatively coarse-resolution grid reduces the spatial dimensionality of the input data sets, which is of benefit in the estimation of empirical orthogonal functions (EOFs) used later in the fingerprint analysis. Regridding of the vertical coordinates was also performed: the WACCM output vertical levels were transformed to match SWOOSH’s vertical levels (which contain 31 pressure layers, spanning a pressure range from 1 to 316 hPa). Model ozone concentrations were masked at latitude bands and layers at which no data were present in SWOOSH. As a sensitivity test, we changed the definition of the upper- and lower-stratospheric regions used for the D&A analysis, using ±1 SWOOSH levels on either side of the original vertical ranges. When calculating the fingerprints for the upper and lower stratosphere, we use anomalies weighted by the cosine of latitude but do not employ vertical pressure weighting. The latter processing choice is reasonable since ozone concentrations do not vary by over an order
of magnitude within each region and the anomalies are expressed as percent changes.

3 Global mean ozone changes

Figure 1a and b show, separately for the upper and lower stratosphere, the time series of globally averaged annual-mean ozone anomalies for the SWOOSH observations and the five WACCM ensembles (ALL1, ALL2, GHGonly, ODSonly, and NAT-h). Anomalies are expressed in terms of percent changes relative to the first 10 years (1960–1969) of the ALL1 simulation. In the upper stratosphere, there is a steady decline in stratospheric ozone from the 1960s to late 1990s in both the ALL1 and ALL2 historical simulations, which is in agreement with the SWOOSH data. After the late 1990s, the decline slows down, and there is some indication of a recovery of stratospheric ozone. The ODSonly anomalies broadly track the temporal changes in ALL1 and ALL2 but reach larger depletion before beginning to increase, supporting the large body of literature indicating that the large decline was predominantly due to ODSs. The difference between the ALL1–ALL2 simulations and ODSonly can be explained by the steady increase in upper-stratospheric ozone in GHGonly, consistent with upper-stratospheric cooling. This supports the conclusion that upper-stratospheric ozone depletion would have been greater during the depletion era in the absence of increases in well-mixed GHGs (WMO, 2014). The NAT-h anomalies show the presence of the 11-year solar cycle in the upper stratosphere but no overall change in ozone from 1960 to 2016.

In the lower stratosphere (Fig. 1b), ozone changes are characterized by larger interannual variability. There is a dominant 2- to 3-year cycle that is due to the historically imposed QBO in all ensemble members of ALL1, ALL2, and NAT-h. Although the ODSonly and GHGonly simulations do not contain the natural and QBO responses present in the other simulations, we see a large inter-ensemble spread in the lower stratosphere when compared to the upper stratosphere. This is related to dynamical differences between ensemble members as a result of the direct transport of ozone in the lower stratosphere. The ODSonly evolution is similar to that in the upper stratosphere and broadly tracks the ALL1 and ALL2 runs. For global GHGonly ozone anomalies in the lower stratosphere, there is no significant trend. As noted by Li et al., 2009, in the lower stratosphere, enhanced ozone advection due to a strong BDC results in decreases in the tropical ozone and increases in the extratropical ozone, and in a global mean sense, the lower-stratospheric changes cancel out.

It is noteworthy that the historical simulations differ from SWOOSH after 2005, when the global-mean lower-stratospheric ozone in SWOOSH does not flatten or begin to turn around, in contrast to the recovery in ALL1 and ALL2. To further examine this divergent behavior, we calculate lower-stratospheric ozone changes averaged over the tropics (30° S to 30° N; see Fig. 1c) and over a global domain that excludes the tropics (Fig. 1d). It is clear from this comparison that the post-2005 observation–model difference in global-mean lower-stratospheric ozone is largely due to ozone differences in the tropics. This may be partly due to the large observational uncertainty in ozone loss in this region. As noted in the WMO (2014) report, there is considerable disagreement between observed data sets in terms of ozone changes in the tropical lower stratosphere since 2000. The report states that changes since 2000 "computed from different data sets in the tropical lowermost stratosphere remain an open question". Because of this observational uncertainty in tropical ozone changes (and because of the discrepancy between post-2000 tropical ozone changes in SWOOSH and the ALL1 and ALL2 simulations), our subsequent D&A analysis for the lower stratosphere is performed over two domains: a full global domain and a domain that excludes 30° S to 30° N.

To summarize, upper-stratospheric ozone in SWOOSH and the ALL1 and ALL2 simulations shows a large decline of 5–8 % decade$^{-1}$ through the middle 1990s, followed by an increase of 2.5–5 % decade$^{-1}$ over the last 10 to 15 years. This decline and recovery is in agreement with findings of the WMO (2014) report. In the lower stratosphere, larger interannual variability complicates the identification of long-term ozone changes, and there are post-2005 differences between the historical WACCM model simulations and SWOOSH data that are relevant to the interpretation of the D&A results. SBUV_CDR ozone anomalies in the upper stratosphere are similar to those of SWOOSH (see Fig. S2).

3.1 Variability in WACCM natural and control simulations

The CTL and NAT simulations provide estimates of the amplitude and timescales of natural ozone variability in the WACCM model (see Fig. 2). In the CTL simulation (upper panels), fluctuations in upper- and lower-stratospheric ozone concentrations arise from internal variability alone. In the last millennium NAT simulation, ozone fluctuations result from both internal variability and solar forcing (lower panels). Previous work has shown that the solar cycle, El Niño–Southern Oscillation (ENSO), and the QBO are the main contributory factors to natural variability in stratospheric ozone, affecting ozone concentrations through chemistry and transport mechanisms (Kirgis et al., 2013; Nair et al., 2013). Below we show the different effects of these contributory factors in the upper and lower stratosphere.

In the upper stratosphere (Fig. 2a), the ozone variation caused by the 11-year solar cycle is clearly evident in the NAT simulation (see expanded time axis). The magnitude of the internally generated ozone variability in the CTL is smaller than ozone variability arising from the 11-year solar cycle. The NAT-derived estimate of the change in upper-
stratospheric ozone from solar maximum to solar minimum is approximately 2–4%. This is in agreement with results from previous observational and model-based studies (Lee and Smith, 2003; Newchurch et al., 2003; Randel and Wu, 2007). NAT also shows longer-timescale variability in upper-stratospheric ozone concentrations in response to solar insolation changes over multiple decades to centuries. The NAT results highlight the large contribution of solar forcing to ozone variability in the upper stratosphere and the smaller role played by internally generated variability. The NAT-h simulation (the yellow time series in Fig. 1a) indicates that there is limited influence of the QBO of 2–3 years on ozone variability in the upper stratosphere. This is consistent with a previous estimate that the QBO effect on upper-stratospheric ozone is less than half that of solar variability over the 11-year solar cycle (Egorova et al., 2004).

In the lower stratosphere (Fig. 2b), the interannual variability in ozone in the CTL is as large as the ozone variability in NAT. In this region of the atmosphere, internal variability in dynamics and transport is the dominant driver of natural fluctuations in ozone – not solar irradiance changes. In fact, the influence of the solar cycle on ozone is difficult to discern...
in the lower stratosphere (see expanded axis). This is consistent with results from previous studies, which have shown that the solar cycle effect on ozone maximizes around 35 km and weakens in the lower stratosphere (Egorova et al., 2004; Lee and Smith, 2003; Newchurch et al., 2003). The QBO-driven anomalies in NAT-h (Fig. 1b) are of much larger magnitude in the lower stratosphere; QBO-driven ozone anomalies range from 4 to 8%, in accord with earlier results (Hasebe, 1994; Randel et al., 1996).

Newchurch (2003) found that the large interannual variability in global-mean lower-stratospheric ozone cannot be explained by the solar cycle and the QBO only: a significant portion of the variance is related to monthly meteorological variability. Changes in interannual tropical upwelling and lower-stratospheric ozone concentrations have been previously linked to ENSO (Bronnimann et al., 2013; Oman et al., 2013). ENSO-induced variations in lower-stratospheric ozone were found to exceed 5%. ENSO-related variations in tropical upwelling are partly responsible for modulating the strength of planetary waves, leading to temperature and water vapor variations in the tropical lower stratosphere that can have impacts on the chemistry and transport of ozone (Randel et al., 2009). ENSO signals in ozone are strongest just above the tropical tropopause; at higher levels, they are weaker than the QBO-related ozone signal (Randel and Thompson, 2011).

Figure S3 shows the normalized power spectra of global upper- and lower-stratospheric ozone anomalies for NAT, NAT-h, and CTL. In the upper stratosphere, the 11-year solar cycle is the strongest signal present in the NAT and NAT-h simulations. In the lower stratosphere, the 2- to 3-year QBO signal is strongest in NAT-h (which has an imposed QBO).

By comparing the stratospheric ozone results in Figs. 1 and 2, we can obtain a rough estimate of the relative sizes of anthropogenically driven and purely natural changes. This simple comparison suggests that the upper stratosphere is the region where anthropogenic signal detection would be most feasible.

### 3.2 Observed versus model variability

Model estimates of internal climate variability are a key component of detection and attribution studies (Allen and Tett, 1999; Santer et al., 2013a). One common strategy is to estimate and remove externally forced climate signals from the observations and then compare the residual variability with control run internal variability (Hegerl et al., 1996). There are a number of uncertainties in such signal removal strategies (Santer et al., 1996b). Here we use an approach similar to Santer et al. (2011) and directly compare estimates of total variability (arising from both internal processes and natural external factors) in the observations, CTL, NAT, NAT-h, and ALL1 runs. If the model systematically underestimates the amplitude of stratospheric ozone variability on multi-decadal timescales, the $S/N$ ratios are likely to be inflated. Whether such a systematic error exists is difficult to determine because the global ozone observational records are relatively short. Observations cannot, therefore, provide an unambiguous constraint on model-based estimates of low-frequency ozone changes, but they can provide useful insights into the direction and size of model variability errors.

We investigate these issues with the modeled and observed monthly-mean time series of global-mean upper- and lower-stratospheric ozone anomalies. After detrending the modeled and observed time series, we isolate long-term variability by applying a band-pass filter with half-power cutoffs at 5 and 20 years to the residuals. We also used a high-pass filter with a half-power point at 3 years to exclude all variability on timescales longer than 5 years. All filtering operations were performed after detrending the time series with a low-pass filter with a half-power point of 30 years. The key point here is that the modeled and observed ozone data are filtered in exactly the same way. We use $S_{IV}$ (interannual variability) and $S_{DV}$ (decadal variability) to denote the temporal SDs of the high-pass- and band-pass-filtered ozone data, respectively.

The $S_{IV}$ and $S_{DV}$ results in Fig. 3 were calculated for SWOOSH (396 months), ALL1 (for the 372-month period from 1984 to 2014), NAT (12 000 months), CTL (2400 months), and NAT-h (2112 months). NAT (NAT-h) provides information on solar and internal variability in the absence (presence) of variability from the QBO. The CTL variability is solely generated by processes internal to the climate system and has no contribution from natural external forcing. Figure 3 shows that these simulations have systematic differences in the amplitude of variability. These differences are manifest in both the sub-3-year timescale and the 5- to 20-year timescales. In the lower stratosphere, the QBO is a dominant component of the variability on timescales < 3 years: only the ALL1 and NAT-h simulations (both of which have QBO-induced ozone changes) are close to the observed value of $S_{IV}$. NAT-h also includes volcanic variability, which is an important component for $S_{IV}$ in the lower stratosphere. CTL and NAT (which both lack QBO and volcanic-driven ozone changes) underestimate the observed interannual variability. In the upper stratosphere, the observed and modeled $S_{IV}$ values all fall within 0.60–0.85%.

The decadal variability is of key interest in D&A studies since it constitutes the background noise against which analysts attempt to identify gradually evolving anthropogenic signals. In the lower stratosphere, the CTL value of $S_{DV}$ is (as expected) lower than in the other three types of simulation. There is no evidence that the simulations most directly comparable to the observations (ALL1 and NAT-h) systematically underestimate the observed values of $S_{DV}$ obtained from the SWOOSH data. Such an underestimate would be concerning: it would spuriously inflate the $S/N$ ratios obtained in the D&A analysis (see below). In fact, both ALL1 and NAT-h yield values of $S_{DV}$ that are slightly larger than in observations (see Santer et al., 2013a).
In the upper stratosphere, however, all four types of model simulation have values of $S_{DV}$ that are smaller than the observed result. As in the lower stratosphere, CTL (which does not include solar forcing) has the lowest decadal variability. ALL1, which has the most realistic time-varying external forcing, is closest to SWOOSH, but still 11% less than the observed $S_{DV}$ value. One possible explanation for this difference in upper-stratospheric $S_{DV}$ between SWOOSH and ALL1 can be seen in Fig. 1a. In the mid- to late 1980s SWOOSH is prominently higher than ALL1, which would lead to a higher $S_{DV}$ value. There is also considerable observational uncertainty in $S_{DV}$ as different ozone data sets have significantly different representations of the magnitude of the solar cycle (Maycock et al., 2016). Chemistry–climate models are also known to have solar cycle variations in ozone that are towards the lower bounds of observational estimates (see chap. 8.5 of SPARC CCMVAL, 2010).

4 Latitude/altitude patterns of ozone change

Figure 1 shows that in the case of all simulations except GHGonly, a simple least-squares linear fit is not an adequate representation of ozone changes over the entire observational record (1984–2016). The nonlinear behavior of ozone in Fig. 1a occurs because ODS emissions were curtailed through implementation of the Montreal Protocol. As mentioned in the introduction, we address this nonlinearity by representing decadal changes in ozone using an EESC proxy as in Randel and Wu (2007) and Newchurch (2003) (see also WMO, 2002; Fioletov and Shepherd, 2005; Stolarski et al., 2006). This proxy isolates ozone changes associated with changes in the amount of ozone-depleting chlorine and bromine in the stratosphere. The EESC curve was taken from the NASA Goddard Automailer, which uses the Newman and Daniel (2007) EESC values updated according to WMO (2010). The EESC calculations assume that the mean age of stratospheric air is 5.5 years and age-of-air spectrum width $= 2.75$ years. Sensitivity to the choice of these two parameters (e.g., varying the MAA from 3 to 6 years) affected the magnitude but not the significance of the regression coefficients. The EESC fit removes some of the nonlinearity that is manifest in the ozone changes.

Figure 4a and b show the respective least-squares linear trends in ozone and in EESC over the entire period of observational record (1984–2016). Results are for the ensemble-mean model simulations and the SWOOSH ozone data. The purpose of the comparison is to determine the expected trend patterns from individual and combined forcings. We also compare our fingerprints, derived in the next section, to the patterns of long-term change.

We used the method outlined by Trenberth (1984) to determine the statistical significance of the trends. Here and subsequently, the stipulated significance level is 5%. When trends are directly calculated from the ozone data, GHGonly is the only simulation yielding an appreciable region of statistically significant results (Fig. 4a). This is the region where GHGonly shows a 1–2% decade$^{-1}$ increase in ozone in the upper stratosphere (1–10 hPa, Fig. 1a) and is the ex-
expected region of latitudinally coherent cooling. In GHG-only, there is a nonsignificant decrease in ozone in the tropical lower stratosphere and a nonsignificant increase in the extratropics, as expected from an enhanced BDC. WMO (2011, 2014) reported negative ozone trends in the tropical lower stratosphere between 1985 and 2005, and CCMVal simulations indicate a long-term increase in tropical upwelling and an increased BDC strength. We note, however, that the WMO (2014) analysis of shorter satellite data sets between 2002 and 2012 does not show significant tropical lower-stratospheric ozone trends, which may reflect the larger noise associated with substantial decadal variability in the lower stratosphere (see Fig. 3).

Very different results are obtained using the EESC regression (Fig. 4b). As expected, ODS forcing makes the largest contribution to the ozone-change pattern in ALL1 and ALL2. Large regions with significant effects of ODS changes on ozone are evident in ALL1, ALL2, and ODSonly. Each of these simulations shows the familiar pattern of midlatitude lobes in the upper stratosphere. Another common aspect of ALL1, ALL2, and ODSonly is a lower-stratospheric response in the Southern Hemisphere (SH), with an Antarctic ozone hole that is strong enough to persist in annual anomalies. The SWOOSH ozone data also display the significant two-lobe structure in the upper stratosphere and the lower-stratospheric response in the SH polar region. The latter feature, however, extends further equatorward in the ALL1, ALL2, and ODSonly simulations than in the observations. In the historical NAT-h simulation, there are no statistically significant trends in either the ozone or EESC data over the 1984 to 2016 period.

The key conclusion of this section is that the statistical significance of linear trends in stratospheric ozone behavior depends critically on whether trends account for nonlinearity in the ozone forcing. Previous studies with EESC-type regressions have shown a familiar pattern of minima in the upper stratosphere (~35–45 km) and the polar lower stratosphere (15–25 km), similar to that obtained here in the EESC-based ALL1, ALL2, and ODSonly simulations (Randel and Wu, 2007; Wang et al., 2002). The upper-stratospheric changes have a symmetric latitudinal structure, which is the fingerprint of gas-phase chlorine-induced ozone loss. For both ozone and EESC, there are no significant trends in NAT-h over 1984–2016. We conclude from our results that in the WACCM model, ozone changes over 1984 to 2016 are primarily forced by human-caused changes in ODS and cannot be explained by natural factors alone. Patterns similar to those found for the SWOOSH data (but with slightly larger magnitudes) were obtained using the SBUV_CDR observations (compare Figs. 4 and S4).

5 Fingerprint estimation

In most applications, the climate change fingerprint is a geographical pattern (Hegerl et al., 1996; Santer et al., 2003), a vertical profile through the atmosphere or ocean (Barnett et al., 2005; Santer et al., 1996b; Tett et al., 1996) or a vector with information on the combined spatial and temporal properties of the signal (Gillet, 2002; Stott et al., 2000; Tett et al., 2002). Here, the fingerprint is a time-invariant latitude–altitude pattern; ozone changes are zonally averaged along latitude bands. The fingerprint provides an estimate of the multi-decadal response to external forcing by combined and individual human forcings. The implicit assumption in this approach is that the spatial pattern of response does not change markedly over time (Santer et al., 2013). We examine the adequacy of this assumption for the specific problem of interest here – the identification of a human-caused
fingerprints in the SWOOSH observations. The assumption of timescale invariance of the fingerprint is tested by defining the fingerprint over different time intervals. We use a standard method (Hasselmann, 1979; Santer et al., 1995) to determine whether model-predicted patterns of externally forced stratospheric ozone changes can be identified in SWOOSH. Fingerprinting is performed separately for the upper and lower stratosphere.

Let $\chi(i, x, p, t)$ represent the annual-mean percent ozone anomalies at latitude band $x$, pressure $p$, and year $t$ for the $i$th ensemble realization for each of the ALL1, ODSonly, and GHGonly simulations:

$$i = 1, \ldots, N_i \quad \text{(number of ensemble members, ranging from 3 to 5)}$$

$$x = 1, \ldots, N_x \quad \text{(total number of latitude bands, 17)}$$

$$p = 1, \ldots, N_p \quad \text{(total number of pressure layers, 10 and 6 for the upper and lower stratosphere, respectively)}$$

$$t = 1, \ldots, N_t \quad \text{(time in years)}$$

Note that we are not using ALL2 for fingerprint estimation because the ODSonly and GHGonly responses were derived from ALL2. As in previous work (Santer et al., 2003, 2013a), we define the fingerprint $F(x, p)$ by first averaging ozone changes over individual ensemble members and then calculating the leading EOF of the covariance matrix of $\chi(x, p, t)$. Many fingerprint studies seek to rotate $F(x, p)$ in a direction that maximizes the signal strength relative to the control run noise (Gillett et al., 2011; Tett et al., 2002). Optimization of $F(x, p)$ generally leads to enhanced detectability of the signal. In this study, we were able to achieve high signal-to-noise levels (see below) without any optimization of $F(x, p)$, and only non-optimized results are discussed. The searched-
for fingerprints were computed using ozone anomalies from 1960 to 2016 for ODSonly and GHGonly and from 1955 to 2014 for ALL1. Sensitivity of our results to the choice of time period for fingerprint calculation was tested by using both a longer period of time (1950 to 2050 for ODSonly and GHGonly) and a shorter time period (1984 to 2016). The fingerprint patterns were found to be relatively insensitive to the choice of time period for estimating $F(x, p)$.

The fingerprint patterns for the upper and lower stratosphere are shown in Figs. 5 and 6 (respectively) for ODSonly (panel a), GHGonly (panel b), and ALL1 (panel c). Below each fingerprint is the associated principal component (PC) time series showing the temporal changes in the strength and sign of the pattern in the model simulation. For both the upper and lower stratosphere, the EOF patterns are similar to the latitude–altitude trend patterns presented previously, with GHGonly EOF1 closely matching the GHGonly linear trends in ozone (Fig. 4a), and the leading ODSonly and ALL1 EOFs closely corresponding to trends computed from the EESC proxy.

In the upper stratosphere, the leading ODSonly and GHGonly EOFs tend to have a maximum amplitude at high latitudes in both hemispheres (ODSonly) and in the tropics (GHGonly). These differences are expected based on the spatial structure and time evolution of long-term changes in ozone in ODSonly and GHGonly (see Fig. 4a and b). On the global scale, however, we note that the EOF1 patterns in both ODSonly and GHGonly have the same sign at virtually all grid points in the upper stratosphere. The standardized PC time series indicate that the ODSonly and ALL1 results are qualitatively similar to the EESC curve, while PC 1 for GHGonly is more linear. We rely on these PC time series in our subsequent $S/N$ analysis.

In the lower stratosphere, the familiar Antarctic ozone hole is visible in the ODSonly and ALL1 fingerprints (Fig. 6a and c). The GHGonly fingerprint (Fig. 6b) is qualitatively similar to the long-term linear trend in ozone in the GHGonly simulation (Fig. 4a), with ozone changes of opposite sign between 100 and 50 hPa in the tropical lower stratosphere.

A common feature of both the upper- and lower-stratospheric results in Figs. 5 and 6 is the similarity between the ALL1 and ODSonly fingerprints. This similarity indicates that in the model simulations, changes in ODS are the primary driver of the changes in stratospheric ozone over the past 50 years. A noticeable difference between the upper- and lower-stratospheric results in Figs. 5 and 6 is that results for the lower stratosphere are noisier. This increased noise is manifest in two ways. First, relative to the upper stratosphere, EOF1 of ODSonly, GHGonly, and ALL1 consistently explains less of the overall variance in lower-stratospheric
ozone changes. Second, the amplitude of the variability in the ODSonly, GHGonly, and ALL1 PC time series is systematically larger in the lower stratosphere. As noted above, these differences between the upper and lower stratosphere reflect the influence of different processes. In the upper stratosphere, local concentrations of ozone are primarily driven by gas-phase chemistry. In contrast, ozone concentrations in the lower stratosphere receive a substantial influence from dynamical transport, which introduces larger interannual variability.

Before presenting the results of the S/N analysis, we first examine the major modes of naturally forced and internal variability estimated from the NAT-h, NAT, and CTL simulations. This is carried out separately for upper- and lower-stratospheric ozone (see Figs. 5d–f and 6d–f, respectively). For NAT and CTL we use the full simulations for calculating EOFs and PCs, while for NAT-h we examine the ensemble mean of the individual members prior to EOF and PC estimation, and we rely on the years 1960 to 2016. Below each NAT and CTL EOF pattern, we show a 50-year segment of the associated PC for a 50-year slice, thus facilitating comparisons with the PC time series of ODSonly, GHGonly, and ALL1.

In the upper stratosphere, the 11-year solar cycle is present in NAT (EOF1) and NAT-h (EOF3). EOF1 from the CTL is also manifest in NAT (EOF2) and appears to be associated with interannual changes in tropical upwelling. Tropical upwelling influences temperature and therefore also affects ozone via temperature-dependent reaction-rate chemistry. In the lower stratosphere, solar-induced ozone changes are not clearly visible in the three leading modes of variability computed from NAT-h, NAT, and CTL. The dominant mode is similar in these simulations and appears to be associated with changes in upwelling and direct transport of ozone through the lower branch of the BDC. EOF2 of NAT-h displays the 2- to 3-year QBO influence on lower-stratospheric ozone (recall that NAT-h contains an imposed QBO to match observations).

In both the upper and lower stratosphere, the patterns of the dominant modes of variability in NAT-h, NAT, and CTL are noticeably different from the searched-for fingerprint patterns. As noted above, the fingerprints estimated from the ODSonly, GHGonly, and ALL1 simulations show upper-stratospheric ozone changes that have the same sign at all grid points. In contrast, the leading three modes of variability estimated from the NAT-h, NAT, and CTL runs do not have the same spatial coherence of ozone change and are generally characterized by ozone changes of opposite sign on smaller spatial scales. The sole exception is the coherent, same-signed EOF pattern associated the 11-year solar cycle (EOF3 in NAT-h and EOF1 in NAT; see Fig. 5d and e). To quantify these similarities and differences, we calculated pattern correlations between the searched-for fingerprints and the leading noise modes (see Table 1). We rely on these correlations later for interpreting results from the S/N analysis (see below).

### 6 Signal-to-noise ratio estimates

We now seek to identify the model-predicted fingerprints of anthropogenically forced stratospheric ozone changes in observations. We project the time-varying annual-mean latitude–height ozone anomalies from the SWOOSH data, denoted here by \( O(x, p, t) \), onto \( F(x, p) \) the time-invariant fingerprint from the ODSonly, GHGonly, or ALL1 simulation (see Figs. 5a–c and 6a–c). This projection step yields the signal time series \( c[F, O](t) \):

\[
c[F, O](t) = \sum_{x=1}^{N_x} \sum_{p=1}^{N_p} F(x, p) O(x, p, t). \tag{1}
\]

This projection is equivalent to a spatially uncentered covariance between the patterns \( F(x, p) \) and \( O(x, p, t) \) at time \( t \). The signal time series \( c[F, O](t) \) provides information on both the amplitude and sign of the fingerprint in observational data. We can then analyze how \( c[F, O](t) \) changes with time – i.e., whether the searched-for fingerprint pattern becomes increasingly similar to observed latitude–altitude patterns of ozone change.

Several approaches can be used to assess the significance of the changes in \( c[F, O](t) \): direct comparison of actual \( c[F, O](t) \) values with a null distribution (e.g., Wigley et al., 1998) or comparison of the trends in \( c[F, O](t) \) with a null distribution of trends (e.g., Santer et al., 2003). We use the latter approach here. To assess trend significance, we require a case in which \( O(x, p, t) \) is replaced by a record in which we know a priori that any spatial correspondence with the fingerprint occurs by chance alone. Here, we use the noise data set \( N(x, p, t) \), which is constructed by concatenating together the NAT, NAT-h, and CTL simulations. The associated noise time series \( c[F, N](t) \) is the spatially uncentered covariance of \( F(x, p) \) and \( N(x, p, t) \):

\[
c[F, N](t) = \sum_{x=1}^{N_x} \sum_{p=1}^{N_p} F(x, p) N(x, p, t). \tag{2}
\]

Estimates of signal to noise (S/N) are conventionally evaluated (e.g., see Santer et al., 2003) by fitting linear least-squares trends of increasing length \( L \) to \( c[F, O](t) \) and then comparing these trends with the SD of the distribution of \( L \)-length trends found in the noise time series \( c[F, N](t) \). Signal detection is obtained when the trend in \( c[F, O](t) \) exceeds and remains above a stipulated significance level. The test is one-tailed and assumes a Gaussian distribution of trends in \( c[F, N](t) \). Here we use the 1% significance level, which corresponds to a S/N ratio close to 2.33. S/N ratios that consistently exceed this level are highly unlikely to be due to the combined effects of natural internal variability and natural external forcing.
Table 1. Correlation coefficients between the leading signal and noise patterns for the upper stratosphere (bold) and lower stratosphere. Signal patterns are the leading EOFs for ALL1, GHGonly, and ODSonly. The noise patterns are the leading EOFs for NAT-h, NAT, and CTL (see Figs. 5 and 6).

|        | ALL1 | GHGonly | ODSonly | NAT-h | NAT   | CTL   |
|--------|------|---------|---------|-------|-------|-------|
| ALL1   |      |         | 0.99    | -0.18 | -0.42 | -0.32 |
| GHGonly| 0.19 | -0.72   |         | 0.18  | -0.10 | 0.19  |
| ODSonly| 0.72 | -0.18   | -0.13   | -0.35 | -0.38 |       |
| NAT-h  | 0.19 | -0.35   | -0.13   |       |       | 0.19  |
| NAT    | -0.25| 0.32    | 0.05    | -0.93 | -      | -0.16 |
| CTL    | 0.09 | 0.24    | -0.03   | 0.90  | 0.97  |       |

Since the sign of the fingerprint EOFs in Figs. 5 and 6 is arbitrary (and since the EOF patterns are very similar to the patterns of ozone trends in Fig. 4), we stipulate that the sign of the leading EOF in ODSonly, GHGonly, or ALL1 should match the sign of the corresponding simulation’s ozone trend pattern. With this stipulation, negative $S/N$ ratios indicate that the signal of the overall change in ozone in the anthropogenically forced simulation is inconsistent with the observed ozone change.

We estimate $S/N$ ratios by fitting least-squares trends to $L$-length segments of $c[F, O](t)$ and then comparing these signal trends with $s(L)$, the SD of the sampling distribution of $m$ maximally overlapping $L$-length trends in $c[F, N](t)$ (i.e., for overlap by all but 1 year). As noted earlier, $c[F, N](t)$ is the concatenation of NAT, NAT-h, and CTL, and due to the discontinuities between the simulations, we discard trends that span two different noise simulations. As $L$ increases, $m$ decreases. We use maximally overlapping trends to guard against excluding the largest changes in ozone from our analysis. We use a minimum trend length of $L = 10$ years; thus, the first $S/N$ ratio (and the earliest possible detection time) is for 10-year trends starting in 1984 and ending in 1993.

One innovative feature of this work is that we derive estimates of $S/N$ behavior using two types of approach. The regression is of the standard form:

$$y = X\beta + \epsilon,$$

where $y$ can be either $c[F, O](t)$ or $c[F, N](t)$, $X$ contains the regressors, $\beta$ contains the reported trend or regression coefficients, and $\epsilon$ is an error term.

The first method we employ for $S/N$ estimation is of the conventional form used in such studies (see, e.g., Santer et al., 2003), where $X$ is the time coordinate in years:

$$X = \begin{pmatrix} 1 & t_0 \\ \vdots & \vdots \\ 1 & t_0 + L \end{pmatrix}, \beta = \begin{pmatrix} \beta_0 \\ \beta_1 \end{pmatrix}. \quad (4)$$

$t_0$ is the starting year (1984), $\beta_0$ is the intercept, and $\beta_1(L)$ is the reported trend coefficient of interest, with units of percent change in ozone per year.

The second method relies on linear regression between $c[F, O](t)$ or $c[F, N](t)$ and a selected PC$_1$ time series. There are a total of six PC$_1$ time series, one for each layer (upper and lower stratosphere) and for each of the ODSonly, GHGonly, and ALL1 fingerprints. In this second method, the regressor $X$ contains the PC$_1$ time series for the selected trend years:

$$X = \begin{pmatrix} 1 & \text{PC}_1(t_0) \\ \vdots & \vdots \\ 1 & \text{PC}_1(t_0 + L) \end{pmatrix}, \beta = \begin{pmatrix} \beta_0 \\ \beta_{\text{PC}_1} \end{pmatrix}, \quad (5)$$

where $\beta_{\text{PC}_1}(L)$ is the reported trend coefficient and has units in percent change per unit SD of PC$_1$. For example, if both $c[F, O](t)$ and PC$_1$ show similar nonlinear behavior over a common analysis period (such as 1984 to 2016), the regression coefficient $\beta_{\text{PC}_1}(L)$ will be unusually large relative to values of the regression coefficient estimated with $c[F, N](t)$. In cases in which PC$_1$ exhibits change that is nearly linear with time, then the change in $\beta_1(L)$ (the linear trend representation of ozone change) and $\beta_{\text{PC}_1}(L)$ with increasing $L$ will be similar.

The advantage of the second method is that it accommodates forcings whose time evolution is markedly nonlinear over the observational record, whereas the first method implicitly assumes that the forcing evolution is quasi-linear over the period of interest. Like the first method, however, the second method still assumes that the spatial structure of the searched-for fingerprint is essentially unchanged with time (an assumption that is justifiable; see above). In summary, our second method of estimating $S/N$ behavior uses information on both the model-derived fingerprint and its time evolution to search for the fingerprint in SWOOSH data. Since the ODSonly and GHGonly fingerprints have different time evolution properties (particularly in the upper stratosphere; compare Fig. 5a and b), explicit consideration of this time evolution information can be useful in separating the ODS- and GHG-induced ozone change signals in observations. In the following, we refer to our first $S/N$ method as the linear or $\beta_1$ trends method, and we refer to the second method as either the PC$_1$ or nonlinear signal method.
The $S/N$ results are shown in Figs. 7 and 8 for the upper and lower stratosphere, respectively. The left panels show $S/N$ estimates from the $\beta_1$ trends, and the right panels show the $S/N$ estimates based on $\beta_{PC_1}$ trends. Results are for the three fingerprints: ALL1, GHGonly, and ODSonly. Since the $\beta_{PC_1}$ trends are regressions to the associated PC1 of the fingerprints, the ALL1 simulation (which ends in December 2014) yields $S/N$ ratios that end in 2014, while the ODSonly and GHGonly simulations span the full observational record, allowing $S/N$ ratios to be calculated through to 2016 (inclusive).

First we consider $S/N$ ratios for the upper stratosphere and for results based on the first method of estimating trends (see left panels of Fig. 7). The use of $\beta_1$ yields virtually identical signal trends, noise trends, and $S/N$ ratios for the ALL1 and ODSonly fingerprints. Recall that the signal trends are calculated from the time series of the projections of the SWOOSH data onto the fingerprints. The ALL1 and ODSonly signal trends are large and positive in the first 10–20 years. These large positive trends arise during the depletion era, when upper-stratospheric ozone decreased steadily until the late 1990s. The positive sign of the signal trends that end in the depletion era reflects the consistency in sign between the observed ozone loss and the loss of ozone captured in the ALL1 and ODSonly simulations (see Fig. 5a and c). As the trend length $L$ extends into the 21st century, the signal trends for ALL1 and ODSonly decline towards zero. This is a result of the stabilization of ozone loss and the emerging recovery of stratospheric ozone. Such nonlinear behavior is not well described by fitting a straight line through the entire $c(F, O)(t)$ time series.

The time evolution of the $\beta_1$ signal trends obtained with the GHGonly fingerprint mirrors the results for ODSonly and ALL1, but is of opposite sign (Fig. 7a). The negative sign of the GHG signal trend arises because of the sign mismatch between upper-stratospheric ozone changes in SWOOSH and the GHGonly simulation (see Fig. 1a). The SDs of the noise trends (Fig. 7b) are very similar for the three fingerprints, which is expected given the similarity between the ODSonly, GHGonly, and ALL1 fingerprint patterns (see Table 1). The decrease in the amplitude of the noise trends with increasing trend-fitting period is typical behavior for many different climate variables (see, e.g., Santer et al., 2013a, b).

Because of early 21st century ozone recovery, and the impact of recovery on $\beta_1$ signal trends, $S/N$ ratios obtained with the ODSonly and ALL1 fingerprints decline from the late 1990s through 2016 (2014 in the case of ALL1; see Fig. 7c). For both of these fingerprints, however, $S/N$ ratios remain above the stipulated 1% significance threshold, even for signal trends sampling the recovery era. This re-
result suggests that even with an emerging “healing” signal in the early 21st century, the overall loss in upper-stratospheric ozone over the entire 1984 to 2014 analysis period is still significantly larger than can be explained by WACCM-based estimates of internally generated and solar-forced changes in ozone. The close agreement between the ALL1 and ODSonly $S/N$ ratios indicates that human-caused changes in stratospheric ozone are the dominant contributor to forced ozone changes in ALL1. In the GHGonly case, the $S/N$ ratio becomes less negative as the trend length $L$ increases. This reflects the observed increase in upper-stratospheric ozone in the early 2000s, which projects positively onto the GHGonly fingerprint (Fig. 5b). In summary, use of a standard fingerprint identification method allows us to positively detect (in SWOOSH ozone data) the upper-stratospheric ozone fingerprints in response to ODS forcing alone and in response to combined ODS and GHG forcing. For all three fingerprints, however, the $\beta_1$ signal trends display pronounced nonlinear behavior because of the observed depletion followed by recovery. This nonlinear behavior is not accounted for in the first ($\beta_1$ based) signal detection method and decreases $S/N$ ratios as the analysis period lengthens.

To address this problem, we also calculated $S/N$ ratios using the above-described method 2, which relies on regressing the $c(F, O)(t)$ time series onto each of the PC time series for the individual ODSonly, GHGonly, and ALL1 fingerprints (see Fig. 5a–c). These “signal” regression coefficients, $\beta_{PC_1}$, are shown as a function of $L$, the analysis period length (Fig. 7d–f). It is instructive to compare the $S/N$ results in the left and right panels of Fig. 7, which highlights differences between the linear and nonlinear signal detection approaches (i.e., between methods 1 and 2, respectively). As $L$ increases and the analysis periods sample both ozone depletion and recovery, use of method 2 markedly increases $S/N$ ratios relative to a purely linear representation of signal trends. This enhancement of $S/N$ occurs because method 2 incorporates information about the nonlinear behavior of the upper-stratospheric ozone signal (behavior that is common to the real world and the WACCM ODSonly and ALL1 simulations). Further information on the enhancement of $S/N$ is provided in Table 2.

There are several other noteworthy features of the method 2 results in Fig. 7. First, the method 1 and method 2 $S/N$ ratios for the GHGonly fingerprint show qualitatively and quantitatively similar behavior. This similarity is due to the fact that $PC_1$ for the GHGonly fingerprint (Fig. 5b) is well described by a linear trend; thus, the regression coefficients between the linear $PC_1$ and the nonlinear $c(F, O)(t)$ time series change markedly as $L$ increases. Second, $S_L$, the SD of the sampling distribution of noise trends, shows some differences in the method 1 and method 2 cases (Fig. 7b and e). In method 1, $S_L$ decreases in amplitude with increasing $L$; this holds for all three fingerprints. In contrast, the method 2 results show that for the ODSonly and ALL1 fingerprints, $S_L$ has a local minimum for $L = 18$ years, and then increases slightly for longer analysis periods. This local minimum in $S_L$ is absent in the GHGonly results for method 2.

The differences between the method 1 and method 2 $S_L$ results are probably related to multiple factors. These include (1) the strong influence of solar forcing on upper-stratospheric ozone, (2) the periodicity in solar forcing in the NAT simulation, (3) the fact that the dominant mode of ozone variability in NAT is more similar to the ODSonly and ALL1 fingerprint patterns than to the GHGonly fingerprint (see pattern correlation results in upper triangle of Table 1), and (4) the quasi-linearity of the $PC_1$ time series for the GHGonly fingerprint and the nonlinearity of the $PC_1$ time series for the ODSonly and ALL1 fingerprints (see Fig. 5).\(^1\)

For the upper stratosphere, we also explored whether our $S/N$ results were robust to the choice of ozone data set. This involved replicating the SWOOSH-based $S/N$ analysis with the SBUV$_{CDR}$ upper-stratospheric ozone data set, which was available from 1979 to 2015. The SBUV $S/N$ analysis was performed with the same model fingerprints shown in Fig. 5a–c. Relative to the SWOOSH results, $S/N$ ratios were generally slightly larger when calculated with SBUV data. As in the SWOOSH case, the SBUV results show that use of method 2 yields a noticeable enhancement of $S/N$ ratios for the ODSonly and ALL1 fingerprints (see Fig. S5). In summary, the method 2 results for the upper stratosphere show (relative to method 1) higher $S/N$ ratios and more confident identification of the ODSonly and ALL1 fingerprints in observations. Even with method 2, however, the

\(^{1}\)Recall that for method 2, the noise regression coefficients are calculated between maximally overlapping $L$-year segments of $c(F, N)(t)$ (the projection of the combined NAT and CTL data onto the ODSonly, GHGonly, or ALL1 fingerprint) and a $L$-year segment of the $PC_1$ time series for the ODSonly, GHGonly, or ALL1 fingerprint. As the $L$-year analysis window is being advanced through $c(F, N)(t)$, there will be (for certain values of $L$) times of random agreement in the phasing of solar variability and the nonlinear behavior in the ODSonly and ALL1 $PC_1$ time series. Such agreement would be expected to yield more complex behavior in $S_L$ as a function of $L$. 

---

**Table 2.** Signal-to-noise ratios for the longest $L$-year analysis periods shown in Figs. 7 and 8. Results are for the linear ($\beta_1$) and nonlinear signal methods ($\beta_{PC_1}$). For ODSonly and GHGonly, the longest analysis period is 33 years (1984–2016). For ALL1, the longest analysis period is 31 years (1984–2014). The figures in brackets for the lower stratosphere indicate the signal-to-noise ratios when the tropics are excluded.

| $F(x, p)$ | Lower stratosphere | Upper stratosphere |
|-----------|--------------------|---------------------|
| $\beta_1$ | $\beta_{PC_1}$ | $\beta_1$ | $\beta_{PC_1}$ |
| ALL1 | 3.43 | 9.66 | 6.72 (4.68) | 6.93 (5.82) |
| ODSonly | 3.27 | 7.68 | 5.35 (4.43) | 3.21 (5.42) |
| GHGonly | −3.49 | −3.81 | 0.18 (−4.13) | 0.32 (−4.89) |
Lower stratosphere

| Linear signal method | Nonlinear signal method |
|----------------------|-------------------------|
| (a) Signal trends in \(c(F,O)(t)\) | (d) Signal trends in \(c(F,O)(t)\) |
| (b) SD of noise trends in \(c(F,N)(t)\) | (e) SD of noise trends in \(c(F,N)(t)\) |
| (c) Signal-to-noise ratio | (f) Signal-to-noise ratio |

**Figure 8.** Same as Fig. 7 but for the lower stratosphere. The solid line shows results for the global lower stratosphere, while the dashed lines exclude the tropics (30° S to 30° N).

GHGonly ozone signal cannot be detected by the final year of the SWOOSH record (2016).

Next, we examine the lower stratosphere. Because of the above-described differences in the post-2005 behavior of tropical lower-stratospheric ozone in ALL1 and SWOOSH (see Fig. 1c), we partition our lower-stratospheric S/N analysis into two cases: for a global domain and for a domain poleward of 30° S to 30° N (i.e., excluding the tropics). Figure 8a–f show the signal trends, noise trends, and S/N ratios for the lower stratosphere, estimated with methods 1 and 2. Consider the \(\beta_l\) signal trends first. As in the case of the upper stratosphere, the behavior of \(\beta_l\) as a function of the trend length \(L\) is similar for \(c\{F, O\}(t)\) time series obtained with the ODSonly and ALL1 fingerprints. These similarities are greater when the tropics are excluded from the analysis. Inclusion or exclusion of the tropics also impacts the \(\beta_l\) signal trends obtained with the GHGonly fingerprint: in the global analysis, trends ending after 2000 are slightly positive, while in the “tropics excluded” case, trends ending after 2000 are slightly negative. The S/N ratios for the \(\beta_l\) trends (Fig. 8c) show that in the lower stratosphere, both the ALL1 and ODSonly fingerprint patterns are detectable in the SWOOSH observations. This holds for the global and tropics excluded domains. The GHGonly fingerprint is not statistically identifiable in either domain by the end of the SWOOSH record.

Recall that in the upper stratosphere, the use of method 2 yielded S/N ratios for the ODSonly and ALL1 fingerprints that were markedly larger than for method 1. A large enhancement of S/N ratios is not evident in the lower stratosphere (Fig. 8c and f). A small S/N enhancement occurs for ODSonly and ALL1 fingerprints, but only when the tropics are excluded. The smaller differences between the method 1 and method 2 S/N results (relative to the upper stratosphere) are likely due to the larger variability in ozone in the lower stratosphere (see Fig. 3). As in the case of method 1, use of method 2 leads to consistent detection of the ODSonly and ALL1 fingerprints in observed lower-stratospheric ozone data, but does not yield positive identification of the GHGonly fingerprint.

In summary, we find that for both the upper and lower stratosphere, the observed decline and emerging recovery of stratospheric ozone is strongly influenced by secular changes in anthropogenic chlorofluorocarbons. These secular changes reflect scientific recognition of the serious consequences of ozone depletion and the eventual formulation and implementation of the Montreal Protocol. We were able to identify the model-predicted latitude–altitude patterns of ozone depletion in observations of upper- and lower-stratospheric ozone loss. We were not able to identify the model-predicted ozone changes arising from human-caused increases in well-mixed GHGs. Our results show that in the upper stratosphere, ozone depletion and recovery introduce pronounced nonlinearity in both modeled and observed ozone changes. Accounting for this common nonlinear behavior in our signal detection method (in our method 2) substantially amplifies S/N ratios. This amplification is largest in the upper stratosphere, where the nonlinear behavior is clearest and noise levels are lowest (see Table 2).

### 7 Discussion

In this study, we used the SWOOSH stratospheric ozone data set and simulations performed with the Whole Atmo-
sphere Community Climate Model to evaluate the relative detectability of ozone changes arising from forcing by ODSs, GHGs, and combined changes in ODSs and GHGs. Our focus was on the period covered by the SWOOSH data (1984 to 2016). Our detection study relied on zonal-mean latitude-height profiles (fingerprints) of ozone change. The lower and upper stratosphere were considered separately in this investigation. This separation was made because the forcing by ODSs and GHGs has different ozone-change signatures in the lower and upper stratosphere, and because the amplitude and patterns of ozone variability differ in these two atmospheric regions.

The credibility of our findings rests on the reliability of the WACCM-based estimates of natural climate variability. Before conducting our signal detection study, we first investigated the skill of the WACCM model in capturing the observed variability in global-mean lower- and upper-stratospheric ozone. This comparison was performed using high-pass- and band-pass-filtered data, which isolated variability on short (less than 3 years) and long (5–20 years) timescales, respectively. We found that in the lower stratosphere, the WACCM simulation incorporating variability associated with the QBO yielded short-term variability that was closest to the SWOOSH-based estimate. Reliable simulation of the longer-timescale variability in ozone required the inclusion of solar forcing, particularly in the upper stratosphere (where solar-forced ozone changes are largest). In the simulation of historical climate change with combined ODS and GHG forcing (which is the simulation most relevant for comparison with observations), the amplitude of the longer-timescale variability is larger than in SWOOSH in the lower stratosphere and slightly smaller than in SWOOSH in the upper stratosphere.

In global-mean terms, the upper- and lower-stratospheric ozone changes from the ODS + GHG simulation best match the observed 20th century ozone depletion and early 21st century ozone recovery (see Fig. 1). In the full pattern analysis of zonal-mean latitude–altitude changes in ozone, all three anthropogenic fingerprints (ODS, GHG, and ODS + GHG) have coherent, large-scale structure. The three fingerprints are spatially dissimilar to the smaller-scale (and opposite-signed) structure of the dominant modes of internal and solar variability. The ODS and GHG fingerprints have similar large-scale structure in the upper stratosphere, but are of opposite sign, and they have distinctly different time evolution over the observational record (see Fig. 5a–c).

We applied two different fingerprint identification methods. The first (our method 1) has been routinely used in a number of detection and attribution studies. It assumes that the time evolution of the fingerprint pattern is quasi-linear over the length of the observational record. In the case of stratospheric ozone, however, the time evolution of the ODS and ODS + GHG fingerprints (which we refer to as signal time series) is markedly nonlinear because of 20th century stratospheric ozone depletion and early 21st century ozone recovery. This means that method 1 – which relies on linear fits to both signal and noise time series – is sub-optimal for assessing the S/N properties of stratospheric ozone changes.

In contrast, our method 2 explicitly accounts for the nonlinear time evolution of the ODS and ODS + GHG fingerprints. In the upper stratosphere, where this nonlinear behavior is clearest, method 2 yields S/N ratios for ODS and ODS + GHG fingerprints that are markedly larger than those obtained with method 1. In both the upper and lower stratosphere, only the ODS + GHG and ODSonly fingerprint patterns were detectable before the current final year (2016) of the SWOOSH ozone data set. The GHG-driven fingerprint of ozone changes was not statistically identifiable in either the upper- or lower-stratospheric SWOOSH data, irrespective of whether we employed method 1 or method 2.

Our results illustrate the importance of explicitly considering the forced, nonlinear temporal changes in the signal of interest. Here, both the 20th century ozone depletion and the emerging early 21st century ozone recovery are driven by well-understood temporal changes in ODS forcing. The time structure of the ODS ozone signal is very different from the much more linear increase in ozone caused by increases in GHG levels (see Fig. 5a–c). Our findings suggest that these differences in time evolution are key to confident separation of the ODS and GHG ozone signals. While one previous detection and attribution study involving stratospheric ozone considered time information, it was not able to clearly separate ODS and GHG signals (Gillett et al., 2011). The key advantage here is that we are dealing with an observational record that extends to 2016 compared to 2005 in Gillett et al., and thus are better able to capture the emerging recovery of stratospheric ozone and the nonlinear behavior of the ODS-driven ozone signal. Figure 9 shows a schematic comparing method 1 and our method 2, which can be applied to attribution studies in which there is evidence that the climate forcing has a temporal component that changes nonlinearly over historical records. Although our study was confined to using spatial fingerprints in the latitude–altitude domain, the method is equally suitable for other domains, such as for longitude–latitude fingerprints. Not shown in Fig. 9 is the calculation of the SD of the sampling distribution for the noise regression coefficients, s(L). Noise calculations use the same approach applied for the signal: projection of the combined noise simulation data onto the fingerprint to retrieve c(F, N)(t), followed by the calculation of regression coefficients. The regressions rely on L-year segments of either c(F, N)(t) and time (in the case of method 1) or c(F, N)(t) and the leading PC time series of the model ALL1, ODSonly, or GHGonly simulation (in the case of method 2).

2As described above, noise time series are obtained by projecting model estimates of naturally caused changes in ozone (both internally generated and arising from solar forcing) onto the ODS, GHG, and ODS + GHG fingerprints.
Figure 9. Schematic of the linear and nonlinear signal methods used for fingerprint identification. $O(x,t)$ denotes the observed time-varying patterns of zonal-mean ozone anomalies; $x$ is an index over the spatial dimensions (latitude and pressure). $F(x)$ is the searched-for fingerprint – the leading empirical orthogonal function of the zonal-mean ozone response in the ODS-only, GHG-only, or ALL1 simulation. For each of the three fingerprints, there is an associated leading principal component time series ($PC_1$) that spans the time period of the observational record. The observations are projected onto each fingerprint, yielding the signal time series, $c\{F,O\}(t)$. In the linear signal method, it is assumed that the time evolution of the fingerprint pattern is quasi-linear over the length of the observational record, and $L$-length linear regression coefficients are calculated between $c\{F,O\}(t)$ and the time in years; these are the coefficients $\beta_{l1}(L)$. In the nonlinear method, the regression is between $L$-year segments of $c\{F,O\}(t)$ and the model $PC_1$ of the corresponding fingerprint. These are the regression coefficients $\beta_{PC_1}(L)$. As noted in Sect. 2, there is a discrepancy between simulated and observed SWOOSH tropical lower-stratospheric ozone post-2005. This is also a region where there is considerable divergence between different observational estimates of ozone changes (WMO, 2014). Both of these factors (model–data differences and observational uncertainty) motivated the additional analysis with exclusion of ozone changes in the tropical lower stratosphere from our $S/N$ analysis. In the tropics excluded case, there is high confidence in our detection of the model ODS signal in the lower stratosphere. For the two reasons outlined above, we have less confidence in the interpretation of our $S/N$ results for the global "tropics included" case, especially for method 2 for which the PC time series are used. Due to the noticeable divergence between simulated and observed post-2005 ozone changes in the tropical lower stratosphere, inclusion of the tropics reduces $S/N$ ratios for our method 2 (i.e., the temporal evolution of ozone differs over the last decade in observations and in the ODS-only signal, thus reducing the regression coefficient in method 2). The above-described results pertain to the identification of the model anthropogenic fingerprints in observations. A related question is the detection time of the GHG signal – the point in the future at which the currently undetectable GHG signal might become identifiable if we continued to monitor stratospheric ozone. We can obtain a purely model-based estimate of this detection time by using the ALL2 ensemble members that extend to the end of the 21st century and treating these ensemble members as surrogate observations. As noted above, $S/N$ ratios for the GHG fingerprint were approaching the 1 % significance level (our stipulated signal detection threshold) near the end of SWOOSH observational.
Figure 10. Same as in Figs. 7 and 8 but the ALL2 ensemble members (three total) are used as surrogate observations, yielding information regarding the estimated detection time for the ODSonly and GHGonly fingerprints in model data. The $\beta$ and $\beta_{PC_1}$ results in (a) and (b) are for the upper stratosphere (respectively); results in (c) and (d) are the corresponding method 1 and method 2 for the lower stratosphere.
fiable in method 2, with model world detection times similar to those estimated for method 1. The model results suggest, therefore, that the GHG fingerprint may not be identifiable in the SWOOSH data for at least another 5 to 10 years.

Our study relies on a single global climate model (WACC). It would be important to determine whether the $S/N$ results obtained here are consistent with those inferred from other climate models. Such a multi-model assessment will require use of models with a well-resolved stratosphere and with reasonable representation of observed ozone variability. This will be the focus of subsequent work.

**Data availability.** The CCM model simulations from WACC can be requested at https://www.earthsystemgrid.org/search.html?Project=CCMII.

The Supplement related to this article is available online at https://doi.org/10.5194/acp-18-143-2018-supplement.

**Competing interests.** The authors declare that they have no conflict of interest.

**Acknowledgements.** The authors would like to thank Sean Davis and Karen Rosenlof, from the National Oceanic and Atmospheric Administration, Chemical Sciences Division, for access to the SWOOSH ozone data set. The National Center for Atmospheric Research is funded by the National Science Foundation.

Edited by: Martin Dameris
Reviewed by: three anonymous referees

**References**

Allen, M. R. and Tett, S. F. B.: Checking for model consistency in optimal fingerprinting, Clim. Dynam., 15, 419–434, https://doi.org/10.1007/s003820050291, 1999.

Aquila, V., Swartz, W. H., Vaugh, D. W., Colarco, P. R., Pawson, S., Polvani, L. M., and Stolarski, R. S.: Isolating the roles of different forcing agents in global stratospheric temperature changes using model integrations with incrementally added single forcings, J. Geophys. Res.-Atmos., 121, 8067–8082, https://doi.org/10.1002/2015JD023841, 2016.

Arfeuille, F., Luo, B. P., Heckendorn, P., Weisenstein, D., Sheng, J. X., Rozanov, E., Schraner, M., Brönnimann, S., Thomason, L. W., and Peter, T.: Modeling the stratospheric warming following the Mt. Pinatubo eruption: uncertainties in aerosol extinctions, Atmos. Chem. Phys., 13, 11221–11234, https://doi.org/10.5194/acp-13-11221-2013, 2013.

Austin, J., Bourqui, M., Braesicke, P., Brühl, C., Butchart, N., Considine, D. B., Cunnold, D., Fahey, D. W., Fleming, E. L., De, P. M., Forster, F., Geller, M. A., Godin-Beekmann, S., Grewe, V., Haigh, J. D., Jackman, C. H., Johnston, P., Lange-matz, U., Law, K. S., Liley, J. B., Megretskaia, I. A., Miller, A. J., Nagashima, T., Pitari, G., Portmann, R. W., Ramaswamy, V., Rognesr, B., Rosenfield, J. E., Ross, M. N., Schmidt, S., Schwarzkopf, M. D., Shindell, D. T., Smith, C. A., Smythinyay, S., Stolarski, R. S., Velders, G. J. M., Wang, R., Weisenstein, D. K., Wu, F., Zander, R. J., Chipperfield, M. P., Randel, W. J., Bodeker, G. E., Dameris, M., Fioletov, V. E., Friedl, R. R., Harris, N. R. P., Logan, J. A., Mcpeters, R. D., Muthama, N. J., Peter, T., Shepherd, T. G., Shine, K. P., Solomon, S., Thomason, L. W., and Zawodny, J. M.: Global Ozone: Past and Future, in: Scientific Assessment of Ozone Depletion: 2002, Geneva, 2002.

Barnett, T. P., Pierce, D. W., AchutaRoan, K. M., Gleckler, P. J., Santer, B. D., Gregory, J. M., and Washington, W. M.: Peningration of human-induced warming into the world’s oceans, Science, 309, 284–287, 2005.

Bindoff, N., Stott, P. A., AchutaRao, K. M., Allen, M. R., Gillett, N., Gutzler, K., Hansingo, K., Hegerl, G., Hu, Y., Jain, S., Mokhov, I. I., Overland, J., Perlwitz, J., Ssebb, R., and Zhang, X.: Detection and attribution of climate change, in: Climate Change 2013: The Physical Basis. Contribution of Working Group I to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change, Cambridge, UK, 2013.

Bouassa, A. E., Degenstein, D. A., Randel, W. J., Zawodny, J. M., Kyröla, E., McLinden, C. A., Sioris, C. E., and Roth, C. Z.: Trends in stratospheric ozone derived from merged SAGE II and Odin-OSIRIS satellite observations, Atmos. Chem. Phys., 14, 6983–6994, https://doi.org/10.5194/acp-14-6983-2014, 2014.

Boville, B.: Wave-mean flow interactions in a general circulation model of the troposphere and stratosphere, J. Atmos. Sci., 43, 1711–1725, 1986.

Brewer, A. W.: Evidence for a world circulation provided by the measurements of helium and water vapour distribution in the stratosphere, Q. Roy. Meter. Soc., 75, 351–363, 1949.

Brönnimann, S., Bhend, J., Franke, J., Flückiger, S., Fischer, A. M., Bleisch, R., Bodeker, G., Hassler, B., Rozanov, E., and Schraner, M.: A global historical ozone data set and prominent features of stratospheric variability prior to 1979, Atmos. Chem. Phys., 13, 9623–9639, https://doi.org/10.5194/acp-13-9623-2013, 2013.

Chehade, W., Weber, M., and Burrows, J. P.: Total ozone trends and variability during 1979–2012 from merged data sets of various satellites, Atmos. Chem. Phys., 14, 7059–7074, https://doi.org/10.5194/acp-14-7059-2014, 2014.

Davis, S. M., Rosenlof, K. H., Hassler, B., Hurst, D. F., Read, W. G., Vömel, H., Selkirk, H., Fujiwara, M., and Damadeo, R.: The Stratospheric Water and Ozone Satellite Homogenized (SWOOSH) database: a long-term database for climate studies, Earth Syst. Data, 8, 461–490, https://doi.org/10.5194/essd-8-461-2016, 2016.

Dobson, G. M. B.: Origin and distribution of the polyatomic molecules in the atmosphere, P. Roy. Soc. A-Math. Phy., 236, 187–193, https://doi.org/10.1098/rspa.1956.0127, 1956.

Egorova, T., Rozanov, E., Manzini, E., Haberreiter, M., Schmutz, W., Zubov, V., and Peter, T.: Chemical and dynamical response to the 11-year variability of the solar irradiance simulated with a chemistry-climate model, Geophys. Res. Lett., 31, L05119, https://doi.org/10.1029/2003GL019294, 2004.
Eyring, V., Cionni, I., Lamarque, J. F., Akiyoshi, H., Bodeker, G. E., Charlton-Perez, A. J., Frith, S. M., Gettelman, A., Kinnison, D. E., Nakamura, T., Oman, L. D., Pawson, S., and Yamashita, Y.: Sensitivity of 21st century stratospheric ozone to greenhouse gas scenarios, Geophys. Res. Lett., 37, L16807, https://doi.org/10.1029/2010GL044443, 2010.

Eyring, V., Lamarque, J. F., and Hess, P.: Overview of IGAC/SPARC Chemistry–Climate Model Initiative (CCMI) community simulations in support of upcoming ozone and climate assessments, in: SPARC Newsletter No. 40, 48–66., 2013.

Fels, S.: Stratospheric sensitivity to perturbations in ozone and carbon dioxide: radiative and dynamical response, J. Atmos. Sci., 37, 2265–2297, https://doi.org/10.1175/1520-0469(1980)037<2265:SSTPIO>2.0.CO;2, 1980.

Fioletov, V. E. and Shepherd, T. G.: Summertime total ozone variations over middle and polar latitudes, Geophys. Res. Lett., 32, L04807, https://doi.org/10.1029/2004GL022080, 2005.

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.

Garcia, R. R., Randel, W. J., Garcia, R. R., and Randel, W. J.: Acceleration of the Brewer–Dobson Circulation due to increases in greenhouse gases, J. Atmos. Sci., 65, 2731–2739, https://doi.org/10.1175/2008JAS2712.1, 2008.

Gillett, N. P.: Detecting anthropogenic influence with a multi-model ensemble, Geophys. Res. Lett., 29, 1970, https://doi.org/10.1029/2002GL015836, 2002.

Gillett, N. P., Akiyoshi, H., Bekki, S., Braesicke, P., Eyring, V., Garcia, R., Karpechko, A. Y., McLinden, C. A., Morgenstern, O., Plummer, D. A., Pyle, J. A., Rozanov, E., Scinocca, J., and Shibata, K.: Attribution of observed changes in stratospheric ozone and temperature, Atmos. Chem. Phys., 11, 599–609, https://doi.org/10.5194/acp-11-599-2011, 2011.

Hansen, J., Sato, M., Ruedy, R., Nazarenko, L., Lacis, A., Schmidt, G. A., Russell, G., Aleinov, I., Bauer, M., Bauer, S., Bell, N., Cairns, B., Canuto, V., Chandler, M., Cheng, Y., Del Genio, A., Faluvegi, G., Fleming, E., Friend, A., Hall, T., Jackman, C., Kelley, M., Kiang, N., Koch, D., Lean, J., Lerner, J., Lo, K., Menon, S., Miller, R., Minnis, P., Novakov, T., Oinas, V., Perlwitz, J. J., Perlwitz, J. J., Rind, D., Romanou, A., Shindell, D., Stone, P., Sun, S., Tausnev, N., Thresher, D., Wielicki, B., Wong, T., Yao, M., and Zhang, S.: Efficacy of climate forcings, J. Geophys. Res., 110, D18104, https://doi.org/10.1029/2005JD006577, 2005.

Hasebe, F.: Quasi-biennial oscillations of ozone and diabatic circulation in the equatorial stratosphere, J. Atmos. Sci., 51, 729–745, https://doi.org/10.1175/1520-0469(1994)051<0729:QBOOOA>2.0.CO;2, 1994.

Hasselmann, K.: On the signal-to-noise problem in atmospheric response studies, in: Meteorology of Tropical Oceans, 251–259, 1979.

Hasselmann, K.: Optimal fingerprints for the detection of time-dependent climate change, J. Climate, 6, 1957–1971, https://doi.org/10.1175/1520-0442(1993)006<1957:OFFTD>2.0.CO;2, 1993.

Hegerl, G., Von Storch, H., Hasselmann, K., Santer, B. D., Cubasch, U., and Jones, P. D.: Detecting greenhouse-gas-induced climate change with an optimal fingerprint method, J. Climate, 9, 2281–2306, 1996.

Hegerl, G. C., Zwiers, F., Braconnot, P., Gillet, N. P., Luo, Y., Marengo, J., Nicholls, N., Penner, J., and Stott, P. A.: Understanding and attributing climate change, in: Climate Change 2007: The Physical Basis. Contribution of Working Group I to the Fourth Assessment Reprot of the Intergovernmental Panel on Climate Change, edited by: Solomon S., et al., Cambridge University Press, Cambridge, UK, 2007.

Jones, A., Urban, J., Murtagh, D. P., Eriksson, P., Brohede, S., Haley, C., Degenstein, D., Bourassa, A., von Savigny, C., Sonkaew, T., Rozanov, A., Bovensmann, H., and Burrows, J.: Evolution of stratospheric ozone and water vapour time series studied with satellite measurements, Atmos. Chem. Phys., 9, 6055–6075, https://doi.org/10.5194/acp-9-6055-2009, 2009.

Karl, T. R., Hassol, S. J., Miller, C. D., and Murray, W.: Temperature Trends in the Lower Atmosphere: Steps for Understanding and Reconciling Differences. A report by the U.S. Climate Change Science Program and the Subcommittee on Global Change Research, Asheville, NC, 2006.

Karlovy, D. J., Cohen, J. A., Meehl, G. A., Mitchell, J. F. B., Oort, A. H., Stouffer, R. J., and Wetherald, R. T.: An example of fingerprint detection of greenhouse climate change, Clim. Dynam., 10, 97–105, https://doi.org/10.1007/BF00210339, 1994.

Krigs, G., Leblanc, T., McDermid, I. S., and Walsh, T. D.: Stratospheric ozone interannual variability (1995–2011) as observed by lidar and satellite at Mauna Loa Observatory, HI and Table Mountain Facility, CA, Atmos. Chem. Phys., 13, 5033–5047, https://doi.org/10.5194/acp-13-5033-2013, 2013.

Kuttippurath, J., Bodeker, G. E., Roscoe, H. K., and Nair, P. J.: A cautionary note on the use of EESC-based regression analysis for ozone trend studies, Geophys. Res. Lett., 42, 162–168, https://doi.org/10.1002/2014GL062142, 2015.

Kyrölä, E., Laine, M., Soiæva, T., Tamminen, J., Päävirinta, S.-M., Tukiainen, S., Zawodny, J., and Thomason, L.: Combined SAGE II–GOMOS ozone profile data set for 1984–2011 and trend analysis of the vertical distribution of ozone, Atmos. Chem. Phys., 13, 10645–10658, https://doi.org/10.5194/acp-13-10645-2013, 2013.

Laine, M., Latva-Pukkilä, N., and Kyrölä, E.: Analysing time-varying trends in stratospheric ozone time series using the state space approach, Atmos. Chem. Phys., 14, 9707–9725, https://doi.org/10.5194/acp-14-9707-2014, 2014.

Langematz, U., Schmidt, F., Kunze, M., Bodeker, G. E., and Brasseics, P.: Antarctic ozone depletion between 1960 and 1980 in observations and chemistry–climate model simulations, Atmos. Chem. Phys., 16, 15619–15627, https://doi.org/10.5194/acp-16-15619-2016, 2016.

Lean, J.: Evolution of the Sun’s spectral irradiance since the Maunder Minimum, Geophys. Res. Lett., 27, 2425–2428, https://doi.org/10.1029/2000GL000043, 2000.

Lee, H. and Smith, A. K.: Simulation of the combined effects of solar cycle, quasi-biennial oscillation, and volcanic forcing on stratospheric ozone changes in recent decades, J. Geophys. Res., 108, 4049, https://doi.org/10.1029/2001JD001503, 2003.

Li, F., Stolarski, R. S., and Newman, P. A.: Stratospheric ozone in the post-CFC era, Atmos. Chem. Phys., 9, 2207–2213, https://doi.org/10.5194/acp-9-2207-2009, 2009.
Newchurch, M. J., Yang, E., Cunnold, D. M., Reinsel, G. C., Zan- 
Nair, P. J., Godin-Beekmann, S., Thiéblemont, R., and 
Hood, L.: The representation of solar cycle signals in strato- 
spheric ozone – Part I: A comparison of recently updated 
satellite observations, Atmos. Chem. Phys., 16, 10021–10043, 
https://doi.org/10.5194/acp-16-10021-2016, 2016.

McLinden, C. A., Tegtmeier, S., and Fioletov, V.: Technical Note: 
A SAGE-corrected SBUV zonal-mean ozone data set, Atmos. 
Chem. Phys., 9, 7963–7972, https://doi.org/10.5194/acp-9-7963- 
2009, 2009.

Merkel, A. W., Harder, J. W., Marsh, D. R., Smith, A. K., 
Fontenla, J. M., and Woods, T. N.: The impact of solar spectral ir- 
radiance variability on middle atmospheric ozone, Geophys. Res. 
Lett., 38, L13802, https://doi.org/10.1029/2011GL047561, 2011.

Molina, M. J. and Rowland, F. S.: Stratospheric sink for chloroflu- 
oromethanes: chlorine atom-catalysed destruction of ozone, Na- 
ture, 249, 810–812, 1974.

Morgenstern, O., Hegglin, M. I., Rozanov, E., O’Connor, F. M., 
Abraham, N. L., Akiyoshi, H., Archibald, A. T., Bekki, S., Butchart, 
N., Chipperfield, M. P., Deushi, M., Dhomse, S., Garcia, R. R., Hardiman, S. C., Horowitz, L. W., Jöckel, P., Josse, B., 
Kinnison, D., Lin, M., Mancini, E., Manyin, M. E., Marchand, M., Marécal, V., Michou, M., Oman, L. D., Pitiäri, G., 
Plummer, D. A., Revell, L. E., Saint-Martin, D., Schofield, R., 
Stenke, A., Stone, K., Sudo, K., Tanaka, T. Y., Tilmes, S., Yamashita, Y., Yoshida, K., and Zeng, G.: Review of the global models used within phase 1 of the Chemistry–Climate Model Initiative (CCMI), Geosci. Model Dev., 10, 639–671, 
https://doi.org/10.5194/gmd-10-639-2017, 2017.

Nair, P. J., Godin-Beekmann, S., Kuttipurath, J., Ancellet, G., 
Goutail, F., Pazmiño, A., Fraúdevalx, L., Zawodny, J. M., Evans, R. D., Wang, H. J., Anderson, J., and Pastel, M.: Ozone trends derived from the total column and vertical profiles at a northern mid-latitude station, Atmos. Chem. Phys., 13, 10373–10384, https://doi.org/10.5194/acp-13-10373-2013, 2013.

Newchurch, M. J., Yang, E., Cunnold, D. M., Reinsel, G. C., Zaw- 
odny, J. M., and Russell, J. M.: Evidence for slowdown in stratospheric ozone loss: first stage of ozone recovery, J. Geo-
phys. Res., 110, 4507, https://doi.org/10.1029/2005JD006347, 
2003.

Newman, P. A., Daniel, J. S., Waugh, D. W., and Nash, E. R.: A 
new formulation of equivalent effective stratospheric chlorine (EESC), Atmos. Chem. Phys., 7, 4537–4552, 
https://doi.org/10.5194/acp-7-4537-2007, 2007.

North, G. R., Stevens, M. J., North, G. R., and Stevens, M. J.: De- 
tection climate signals in the surface temperature record, J. Climate, 11, 563–577, https://doi.org/10.1175/1520-
0442(1998)011<0563:DCSITS>2.0.CO;2, 1998.

Oman, L. D., Plummer, D. A., Waugh, D. W., Austin, J., Scinocca, J. F., Douglass, A. R., Salawitch, R. J., Canty, T., Akiyoshi, H., Bekki, S., Braesicke, P., Butchart, N., Chipper- 
field, M. P., Cugnet, D., Dhomse, S., Eyring, V., Frith, S., 
Hardiman, S. C., Kinnison, D. E., Lamarque, J.-F., Mancini, E., Marchand, M., Michou, M., Morgenstern, O., Nakamura, T., Nielsen, J. E., Oliivié, D., Pitiäri, G., Pyle, J., Rozanov, E., Shepherd, T. G., Shibata, K., Stolarski, R. S., Teyssèdre, H., 
Tian, W., Yamashita, Y., and Ziemke, J. R.: Multimodel as-
essment of the factors driving stratospheric ozone evolution over the 21st century, J. Geophys. Res., 115, D24306, 
https://doi.org/10.1029/2010JD014362, 2010.

Oman, L. D., Douglass, A. R., Ziemke, J. R., Rodriguez, J. M., 
Waugh, D. W., and Nielsen, J. E.: The ozone response to ENSO in Aura satellite measurements and a chemistry-climate simulation, J. Geophys. Res.-Atmos., 118, 965–976, 
https://doi.org/10.1029/2012JD018546, 2013.

Polvani, L. M., Wang, L., Aquila, V., Waugh, D. W., Polvani, L. M., 
Wang, L., Aquila, V., and Waugh, D. W.: The impact of ozone-depleting substances on tropical upwelling, as revealed by the absence of lower-stratospheric cooling since the late 1990s, J. Climate, 30, 2523–2534, https://doi.org/10.1175/JCLI-D-16-
0532.1, 2017.

Randel, W. J. and Thompson, A. M.: Interannual variability and trends in tropical ozone derived from SAGE II satellite data and SHADOZ ozonesondes, J. Geophys. Res., 116, D07303, 
https://doi.org/10.1029/2010JD015195, 2011.

Randel, W. J. and Wu, F.: A stratospheric ozone profile data set for 1979–2005: variability, trends, and comparisons with column ozone data, J. Geophys. Res., 112, D06313, 
https://doi.org/10.1029/2006JD007339, 2007.

Randel, W. J., Wu, F., Randel, W. J., and Wu, F.: Isolation of the ozone QBO in SAGE II data by singular-value decomposition, J. Atmos. Sci., 53, 2546–2559, https://doi.org/10.1175/1520-
0469(1996)053<2546:ITQOQ>2.0.CO;2, 1996.

Randel, W. J., Shine, K. P., Austin, J., Barnett, J., Claud, C., 
Gillett, N. P., Keckhut, P., Langematz, U., Lin, R., Long, C., 
Mears, C., Miller, A., Nash, J., Seidel, D. J., Thompson, D. W. J., Wu, F., and Yoden, S.: An update of observed stratospheric temperature trends, J. Geophys. Res., 114, D20107, 
https://doi.org/10.1029/2008JD010421, 2009.

Santer, B., Wigley, T., and Jones, P.: Correlation methods in fingerprint detection studies, Clim. Dynam., 8, 265–276, 1993.

Santer, B., Wigley, T. M. L., Meehl, G. A., Wehner, M. F., Mears, C., Schabel, M., Wentz, F. J., Ammann, C., Arblaster, J., Bettge, T., Washington, W. M., Taylor, K. E., Boyle, J. S., Bruggemann, W., and Doutriaux, C.: Influence of satellite data uncertainties on the detection of externally forced climate change, Science, 300, 1280–1284, 2003.

Santer, B. B., Painter, J. F. J., Mears, C. a, Doutriaux, C., Caldwell, P., Arblaster, J. M., Cameron-Smith, P. J., Gillett, N. P., Gleckler, P. J., Lanzante, J., Perlwitz, J., Solomon, S., Stott, P. a, Taylor, K. E., Terray, L., Thorne, P. W., Wehner, M. F., Wentz, F. J., Wigley, T. M. L., Wilcox, L. J., and Zou, C.-Z.: Identifying human influences on atmospheric temperature, P. Natl. Acad. Sci. USA, 110, 26–33, https://doi.org/10.1073/pnas.1210514109, 2013a.

Santer, B. D., Taylor, K. E., Wigley, T. M. L., Penner, J. E., Jones, P. D., and Cubasch, U.: Towards the detection and attribution of an anthropogenic effect on climate, Clim. Dynam., 12, 77–100, https://doi.org/10.1007/ BF00237222, 1995.

Santer, B. D., Taylor, K. E., Wigley, T. M. L., Johns, T. C., 
Jones, P. D., Karoly, D. J., Mitchell, J. F. B., Oort, A. H., 
Penner, J. E., Ramaswamy, V., Schwarzkopf, M. D., Stouf- 
er, R. J., and Tett, S.: A search for human influences on the thermal structure of the atmosphere, Nature, 382, 39–46, 
https://doi.org/10.1038/382039a0, 1996a.
Santer, B. D., Wigley, T. M. L., Barnett, T. P., and Anyamba, E.: Detection of climate change and attribution of causes, edited by: Santer, B. D., Wigley, T. M. L., Barnett, T. P., and Anyamba, E., Cambridge University Press, New York, NY, 1996b.

Santer, B. D., Mears, C., Doutriaux, C., Caldwell, P., Gleckler, P. J., Wigley, T. M. L., Solomon, S., Gillett, N. P., Ivanova, D., Karl, T. R., Lanzante, J. R., Meehl, G. A., Stott, P. A., Taylor, K. E., Thorne, P. W., Wehner, M. F., and Wenzt, F. J.: Separating signal and noise in atmospheric temperature changes: the importance of timescale, J. Geophys. Res.-Atmos., 116, D22105, https://doi.org/10.1029/2011JD016263, 2011.

Santer, B. D., Painter, J. F., Bonfils, C., Mears, C. A., Solomon, S., Wigley, T. M. L., Gleckler, P. J., Schmidt, G. A., Doutriaux, C., Gillett, N. P., Taylor, K. E., Thorne, P. W., and Wentz, F. J.: Human and natural influences on the changing thermal structure of the atmosphere, P. Natl. Acad. Sci. USA, 110, 6–11, https://doi.org/10.1073/pnas.1305332110, 2013b.

Shindell, D. T. and Grewe, V.: Separating the influence of halogen and climate changes on ozone recovery in the upper stratosphere, J. Geophys. Res., 107, 3–10, 2002.

Solomon, S.: Stratospheric ozone depletion: a review of concepts and history, Rev. Geophys., 37, 275–316, 1999.

Solomon, S. and Garcia, R. R.: On the distributions of long-lived tracers and chlorine species in the middle atmosphere, J. Geophys. Res., 89, 11633, https://doi.org/10.1029/JD089iD07p11633, 1984.

Solomon, S., Kinnison, D., Bandoro, J., and Garcia, R.: Simulation of polar ozone depletion: an update, J. Geophys. Res.-Atmos., 120, 7958–7974, https://doi.org/10.1002/2015JD023365, 2015.

Solomon, S., Ivy, D. J., Kinnison, D., Mills, M. J., Neely, R. R., and Schmidt, A.: Emergence of healing in the Antarctic ozone layer, Science, 353, 269–274, https://doi.org/10.1126/science.aee0061, 2016.

Stolarski, R. S., Douglass, A. R., Steenrod, S., and Pawson, S.: Trends in stratospheric ozone: lessons learned from a 3D chemical transport model, J. Atmos. Sci., 63, 1028–1041, https://doi.org/10.1175/JAS3650.1, 2006.

Stott, P. A., Tett, S. F. B., Jones, G. S., Allen, M. R., Mitchell, J. F. B., and Jenkins, G. J.: External control of 20th century temperature by natural and anthropogenic forcings, Science, 290, 2133–2137, https://doi.org/10.1126/science.290.5499.2133, 2000.

Tett, S. F. B., Mitchell, J. F. B., Parker, D. E., and Allen, M. R.: Human influence on the atmospheric vertical temperature structure: detection and observations, Science, 274, 1170–1173, 1996.

Tett, S. F. B., Jones, G. S., Stott, P. A., Hill, D. C., Mitchell, J. F. B., Allen, M. R., Ingram, W. J., Johns, T. C., Johnson, C. E., Jones, A., Roberts, D. L., Sexton, D. M. H., and Woodward, M. J.: Estimation of natural and anthropogenic contributions to twentieth century temperature change, J. Geophys. Res., 107, 4306, https://doi.org/10.1029/2000JD000228, 2002.

Thorne, P. W., Jones, P. D., Osborn, T. J., Davies, T. D., Tett, S. F. B., Parker, D. E., Stott, P. A., Jones, G. S., and Allen, M. R.: Assessing the robustness of zonal mean climate change detection, Geophys. Res. Lett., 29, 26-1-26-4, https://doi.org/10.1029/2002GL015717, 2002.

Tie, X. and Brasseur, G.: The response of stratospheric ozone to volcanic eruptions: sensitivity to atmospheric chlorine loading, Geophys. Res. Lett., 22, 3035–3038, 1995.

Trenberth, K.: Some effects of finite sample size and persistence on meteorological statistics. Part I: Autocorrelations, Mon. Weather Rev., 112, 2459–2368, 1984.

Tunmon, F., Hassler, B., Harris, N. R. P., Staellein, J., Steinbrecht, W., Anderson, J., Bodeker, G. E., Bourassa, A., Davis, S. M., Degenstein, D., Frith, S. M., Froidevaux, L., Kyrölä, E., Laine, M., Long, C., Penckwitt, A. A., Sioris, C. E., Rosenlof, K. H., Roth, C., Wang, H.-J., and Wild, J.: Intercomparison of vertically resolved merged satellite ozone data sets: interannual variability and long-term trends, Atmos. Chem. Phys., 15, 3021–3043, https://doi.org/10.5194/acp-15-3021-2015, 2015.

Vinnikov, K. Y., Robock, A., Stouffer, R. J., and Manabe, S.: Vertical patterns of free and forced climate variations, Geophys. Res. Lett., 23, 1801–1804, https://doi.org/10.1029/96GL01736, 1996.

Wang, H. J., Cunnold, D. M., Thomason, L. W., Zawodny, J. M., and Bodeker, G. E.: Assessment of SAGE version 6.1 ozone data quality, J. Geophys. Res. Atmos., 107, ACH 8–1–ACH 8–18, https://doi.org/10.1029/2002JD002418, 2002.

Wigley, T. M. L., Jaumann, P. J., Santer, B. D., and Taylor, K. E.: Relative detectability of greenhouse-gas and aerosol climate change signals, Clim. Dynam., 14, 781–790, https://doi.org/10.1007/s003820050254, 1998.

WMO (World Meteorological Organization): Scientific Assessment of Ozone Depletion: 2002, Global Ozone Research and Monitoring Project – Report No. 47, 498 pp., Geneva, Switzerland, 2003.

WMO (World Meteorological Organization): Scientific Assessment of Ozone Depletion: 2010, Global Ozone Research and Monitoring Project-Report No. 52, 516 pp., Geneva, Switzerland, 2011.

WMO (World Meteorological Organization): Scientific Assessment of Ozone Depletion: 2014, Global Ozone Research and Monitoring Project – Report No. 55, 416 pp., Geneva, Switzerland, 2014.

Wohltmann, I., Lehmann, R., Rex, M., Brunner, D., and Mäder, J. A.: A process-oriented regression model for column ozone, J. Geophys. Res., 112, D12304, https://doi.org/10.1029/2006JD007573, 2007.

Xue, X.-H., Liu, H.-L., and Dou, X.-K.: Parameterization of the inertial gravity waves and generation of the quasi-biennial oscillation, J. Geophys. Res. Atmos., 117, D06103, https://doi.org/10.1029/2011JD016778, 2012.

Zawodny, J. M. and McCormick, M. P.: Stratospheric Aerosol and Gas Experiment II measurements of the quasi-biennial oscillations in ozone and nitrogen dioxide, J. Geophys. Res., 96, 9371, https://doi.org/10.1029/91JD00517, 1991.