Impacts of Stratospheric Sulfate Geoengineering on Tropospheric Ozone

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Abstract

A range of solar radiation management (SRM) techniques has been proposed to counter anthropogenic climate change. Here, we examine the potential effects of stratospheric sulfate aerosols and solar insolation reduction on tropospheric ozone and ozone at Earth’s surface. Ozone is a key air pollutant, which can produce respiratory diseases and crop damage. Using a version of the Community Earth System Model from the National Center for Atmospheric Research that includes comprehensive tropospheric and stratospheric chemistry, we model both stratospheric sulfur injection and solar irradiance reduction schemes, with the aim of achieving equal levels of surface cooling relative to the Representative Concentration Pathway 6.0 scenario. This allows us to compare the impacts of sulfate aerosols and solar dimming on atmospheric ozone concentrations. Despite nearly identical global mean surface temperatures for the two SRM approaches, solar insolation reduction increases global average surface ozone concentrations while sulfate injection decreases it. A fundamental difference between the two geoengineering schemes is the importance of heterogeneous reactions in the photochemical ozone balance with larger stratospheric sulfate abundance, resulting in increased ozone depletion in mid- and high latitudes. This reduces the net transport of stratospheric ozone into the troposphere and thus is a key driver of the overall decrease in surface ozone. At the same time, the change in stratospheric ozone alters the tropospheric photochemical environment due to enhanced ultraviolet radiation. A shared factor among both SRM scenarios is decreased chemical ozone loss due to reduced tropospheric humidity. Under insolation reduction, this is the dominant factor giving rise to the global surface ozone increase. Regionally, both surface ozone increases and decreases are found for both scenarios, that is, SRM would affect regions of the world differently in terms of air pollution. In conclusion, surface ozone and tropospheric chemistry would likely be affected by SRM, but the overall effect is strongly dependent on the SRM scheme. Due to the health and economic impacts of surface ozone, all these impacts should be taken into account in evaluations of possible consequences of SRM.
1 Introduction

1.1 Atmospheric Ozone

It is well known that sulfate aerosols in the stratosphere enhance heterogeneous chemical reactions that lead to enhanced ozone depletion after larger volcanic eruptions (Solomon, 1999). With present day anthropogenic halogen loading, the aerosols provide additional surface area for heterogeneous reactions that activate halogens and hence increase catalytic ozone destruction, especially in high latitudes (Tie and Brasseur, 1995). This has been modeled and observed following the large 1982 El Chichón and 1991 Pinatubo volcanic eruptions (Tie and Brasseur, 1995; Portman et al., 1996).

However, volcanic eruptions do not only affect stratospheric ozone, but also impact tropospheric composition, often due to stratosphere-troposphere coupled effects. The 1991 Pinatubo eruption, for example, has been linked to changes in stratosphere-troposphere exchange (STE) of ozone (Aquila et al., 2012; Aquila et al., 2013; Pitar et al., 2016). In addition, the stratospheric ozone decrease led to an invigorated photochemical environment in the troposphere due to enhanced downward chemically-active ultraviolet (UV) radiation (Tang et al., 2013).

This study focuses on tropospheric ozone, in particular surface ozone concentration changes. Surface ozone is of central importance to Earth’s environment and as an air pollutant it adversely impacts human health (e.g., Kampa and Castanas, 2008) and the ecosystem (e.g., Mauzeral and Wang, 2001; Ashmore, 2005; Ainsworth et al., 2012). There have been numerous studies of the observed surface ozone trend (e.g., Cooper et al., 2014), identifying ozone sources and sinks (e.g., Wild, 2007), predicting future changes (e.g., Young et al., 2013), and understanding the impacts of such changes (e.g., Silva et al., 2013). Global surface ozone concentrations are estimated to have doubled since the preindustrial period (Vingarzan, 2004),
mainly due to increased emissions of ozone precursors associated with industrialization (e.g., Forster et al., 2007). Differences in future tropospheric ozone concentrations will be strongly dependent on the emission pathway followed (Stevenson et al., 2006), which will determine both in-situ tropospheric chemical production of ozone and transport from the ozone-rich stratosphere (Collins et al., 2003; Wild et al., 2012; Neu et al., 2014).

1.2 Differences between sulfate and solar geoengineering

The progression of global warming, slow mitigation efforts, and our relatively limited adaptive capacity force consideration of SRM geoengineering as one possible strategy to avoid many of the most undesirable consequences of global warming (Crutzen, 2006; Wigley, 2006; Tilmes, 2016a). The above discussed factors controlling tropospheric ozone concentrations could be affected by SRM schemes (Nowack et al., 2016). Here we compare a proposed geoengineering scheme, stratospheric sulfur injection, to solar irradiance reduction. Both schemes would cool Earth’s surface by reducing sunlight reaching the surface, either by aerosols reflecting sunlight or by artificially reducing the solar constant in a climate model, but sulfate geoengineering would strongly heat the stratosphere and provide aerosol surfaces for chemical reactions. Previous studies have shown that injected sulfur chemically forms sulfate aerosols within a couple of weeks. The aerosol layer absorbs near infrared solar radiation as well as outgoing longwave radiation and results in stratospheric warming (e.g., Tilmes et al., 2009; Ammann et al., 2010; Jones et al., 2011). Additionally changes in ozone and advection impact the warming in the stratosphere (Richter et al., 2017, submitted). Under solar reduction, the stratosphere would be cooler due to reduced shortwave heating (Govindasamy and Caldeira, 2000), although simultaneous stratospheric ozone changes (if considered) may buffer this effect (Nowack et al., 2016).
One of the most important differences between the two scenarios is that if a permanently enhanced stratospheric aerosol layer is artificially created in an attempt to reduce anthropogenic global warming, the resulting strong ozone depletion, in particular in mid- and high latitudes, would have serious impacts on the biosphere, similar to the effects observed after large volcanic eruptions described above (Crutzen, 2006; Rasch et al., 2008a; Rasch et al., 2008b; Tilmes et al., 2008, 2009, 2012). This effect would have to be expected as long as there is anthropogenic halogen in the stratosphere. In the remote future, the decreasing burden of anthropogenic halogen will eventually result in the recovery of the ozone layer. Under such conditions additional stratospheric aerosols could actually have the opposite effect by deactivating ozone depleting nitrogen oxides, thus leading to an increase in ozone in the stratosphere (Tie and Brasseur, 1995; Pitari et al, 2014). Overall, such changes to the stratosphere would also have important implications for tropospheric composition. Decreasing stratospheric ozone leads to more UV propagating through, with increasing ozone having the opposite effect, which would thus alter the photochemical environment of the troposphere in different ways (Tilmes et al., 2012; Nowack et al., 2016).

In the following sections, we describe the experimental set-up of the two geoengineering schemes and discuss some general climate impacts, followed by a detailed discussion of tropospheric and surface ozone changes. We also show that sulfate and solar geoengineering would impact the stratosphere differently, which implies further key differences in their potential influences on tropospheric composition. In this study, we examine the impacts of stratospheric sulfate geoengineering on tropospheric ozone for the first time.

2 Model and Experiment Design

We simulated both types of SRM schemes using the full tropospheric and stratospheric
chemistry version of the Community Earth System Model – Community Atmospheric Model 4 (CESM CAM4-chem) with horizontal resolution of 0.9° x 1.25° lat-lon and 26 levels from the surface to about 40 km (3.5 mb). The model has been shown to give a good representation of present-day atmospheric composition in the troposphere (Tilmes et al., 2016b) and stratosphere at 2° resolution (Fernandez et al., 2017). Similar to the 2° model version, the 1° horizontal resolution version of the model also produces reasonable stratosphere and troposphere ozone chemistry (Figs. S1-S2). CAM4-chem is fully coupled to the Community Land Model version 4.0 with prescribed satellite phenology (CLM4SP), the Parallel Ocean Program version 2 (POP2) ocean model, and the Los Alamos sea ice model (CICE version 4). The tropospheric chemical mechanism in CAM4-chem is based on the Model for Ozone and Related chemical Tracers (MOZART) version 4 (Emmons et al., 2010). The stratospheric chemical mechanism is described in Kinnison et al. (2007), Lamarque et al. (2012) and Tilmes et al. (2015), and the complete chemical reactions included (photolysis, gas-phase chemistry and heterogeneous chemistry) are listed in Tilmes et al. (2016b), Table A2. Reaction rates are updated following Jet Propulsion Laboratory 2010 recommendations (Sander et al., 2011). The model uses a nudged quasi-biennial oscillation (QBO), which means the QBO will not be modified by the radiative interaction of the aerosols. Interaction between aerosol burden and photolysis rates is not included in the model. Changes in photolysis rates in the troposphere depend on the stratospheric ozone column change (Kinnison et al., 2007). Increased ozone depletion as the result of geoengineering would therefore leads to an increase in UV in mid- and high latitudes. Since our model does not include the aerosol scattering effect on UV, expected UV reductions from the increased sulfate aerosol layer is not taken into account, which might result in an overestimation of the tropospheric photolysis. Volatile organic compound (VOC) emissions are
simulated by the Model of Emission of Gases and Aerosols from Nature (MEGAN v2.1) (Guenther et al., 2012). The dynamical ocean model does not include any biogeochemical feedbacks and only the atmospheric and land models are coupled to the atmospheric chemistry component. The model configuration used here, but at 2° resolution, is participating in the current phase of the Chemistry-Climate Model Initiative (Tilmes et al., 2016b, Morgenstern et al., 2017).

We compare three ensemble members each of the two geoengineering scenarios to a three-ensemble reference run with Representative Concentration Pathway 6.0 (RCP6.0; Meinshausen et al., 2011) anthropogenic forcing from 2020 to 2089. Both geoengineering scenarios include RCP6.0 forcings. Our sulfate aerosol implementation is the G4 Specified Stratospheric Aerosol (G4SSA) experiment (Tilmes et al., 2015), whereas solar reduction geoengineering is the solar analog (hereafter G4SSA-S) by imposing a solar irradiance reduction with the same negative radiative forcing at the top of the atmosphere (TOA) as in G4SSA. G4SSA uses a prescribed stratospheric aerosol surface area distribution to mimic the effects of continuous emission into the tropical stratosphere at 60 mb of 8 Tg SO₂ yr⁻¹ from 2020 to 2069. More details of this prescribed stratospheric aerosol distribution are given in Tilmes et al. (2015b) and Xia et al. (2016). The G4SSA scenario then continues from 2070 to 2089 without imposed aerosols to study the termination effect of geoengineering. During the sulfate injection period, the net solar flux at the TOA has been decreased by 2.5 W/m² compared to RCP6.0 (Fig. 1a). This number was obtained by a double radiation call in the model in calculating the direct forcing of the prescribed aerosol layer. To attain the same TOA solar flux reduction in G4SSA-S, we reduced the total solar insolation by 14.7 W/m² during 2020-2069 assuming a global average planetary albedo of 0.32 (14.7 W/m² = 2.5 W/m² × 4.10⁻⁰.32) (Fig. 1b). From 2070 on, we accordingly
reset the total solar insolation back to the reference level to simulate the abrupt termination of geoengineering.

3 Results and Discussion

3.1 Climatology in G4SSA and G4SSA-S

As a consequence of the same net all-sky TOA solar flux reduction in G4SSA and G4SSA-S (Fig. 1a), the two scenarios show approximately the same global mean surface temperature reduction of 0.8 K compared with RCP6.0 (Fig. 2a) (all values below are the average of years 2030-2069, the last 40 years of geoengineering). After the termination of geoengineering on 1 January 2070, the global mean surface temperature rapidly increases. Fig. 3 shows the surface temperature differences between G4SSA, G4SSA-S, and RCP6.0 in years 2030-2069. Consistent with the global average temperature change, the two geoengineering scenarios have similar temperature reduction patterns (Fig. 3a and 3b), and the differences between them are not significant in most regions (Fig. 3c). The similar warming in the North Atlantic under G4SSA and G4SSA-S relative to RCP6.0 (Fig. 3a and 3b) is due to the regional cooling under RCP6.0 as a result of slowing down of the Gulf Stream (Hartmann et al., 2013). The temperature difference between G4SSA and G4SSA-S (Fig. 3c) is larger in the Northern Hemisphere winter (Fig. S3). The warming over northern Europe and Asia in G4SSA relative to G4SSA-S is the characteristic “winter warming” from volcanic stratospheric aerosols (Robock, 2000). However, the zonal mean stratospheric temperatures in G4SSA and G4SSA-S differ substantially (Fig. 4). Sulfate aerosols in the stratosphere result in strong warming by 3 K in the tropics (Fig. 4a), while in G4SSA-S there is slight cooling (Fig. 4b), consistent with previous studies (Tilmes et al., 2009; Ammann et al., 2010; Jones et al., 2011). The slight warming in the lower stratosphere under G4SSA-S (Fig. 4b) might be a result of ozone changes and dynamical...
heating (discussion in Section 3.3.2). In both cases, the troposphere shows strong temperature reduction with similar patterns and ranges.

Global averaged precipitation and evaporation have similar reductions of 0.07 mm/day in the two scenarios (Fig. 2b and Fig. S4), with no statistically significant difference between them. Most of the evaporation terms show a larger reduction in G4SSA than in G4SSA-S, except for plant transpiration, which has the opposite pattern (Fig. S4). As shown by Xia et al. (2016), enhanced diffuse radiation in G4SSA increases photosynthesis, which produces stronger transpiration. Therefore, transpiration in G4SSA reduces less than in G4SSA-S.

The similar evaporation reduction in G4SSA and G4SSA-S can also be explained by the surface energy budget (Fig. 5b). Although we keep the net shortwave radiation at the TOA the same in the two schemes (Fig. 1a and Fig. 5a), surface net solar radiation reduces more in G4SSA than in G4SSA-S (Fig. 2c and Fig. 5b) due to the absorption by sulfate aerosols in the near-infrared. This stronger surface solar forcing in G4SSA-S is mainly balanced by larger net longwave radiation to the atmosphere (Fig. 5). As a result, latent heat changes in the two scenarios are similar.

Here, precipitation and evaporation changes are very similar under sulfate and solar geoengineering. This is different from previous studies by Niemeier et al. (2013) and Ferraro et al. (2014) who found that the effect on the hydrological cycle is larger for sulfate geoengineering. These differences are related to the experimental design. Niemeier et al. (2013) bias corrected all geoengineering scenarios to keep the net total flux at the TOA the same as that in 2020, while we keep the same net solar flux at the TOA in G4SSA and G4SSA-S (Fig. 1a). However, we found the net total fluxes at the top of the model in G4SSA and G4SSA-S are similar as well (Fig. 5a and Fig. S5). Therefore, differences in the TOA boundary conditions might not be the main
reason for the different hydrological cycle responses. In their studies, with the same magnitude
of surface cooling, the sulfate injection scenario led to a greater reduction of globally averaged
evaporation and precipitation as compared with the solar case. Ferraro et al. (2014) attributed
the enhanced hydrological cycle response to sulfate geoengineering to extra downwelling
longwave radiation because of stratospheric heating from the injected aerosols. Sulfate
geoengineering thus led to a relative stabilization of the troposphere (by heating the upper
troposphere more than the mid-lower troposphere) compared with the solar reduction case
(which we do not find, Fig. 4c). A more stratified troposphere, in turn, results in a stronger
reduction of latent heat fluxes and precipitation (similar to theoretical considerations by Bala et
al. (2008)). We find two possible reasons for the different response in our experiments. (1) The
column ozone change could play an important role. In Niemeier et al. (2013) and Ferraro et al.
(2014), the same prescribed ozone was used in all scenarios, while we used a fully coupled
atmosphere-chemistry model. As shown in section 3.2, total column ozone in G4SSA reduces
by about 5 DU (mainly in the lower stratosphere) compared with RCP6.0 and G4SSA-S (Fig. 6).
Less ozone in G4SSA will change its radiative forcing, surface radiative fluxes and atmospheric
lapse rate (Chiodo and Polvani, 2015; MacIntosh et al., 2016; Nowack et al., 2015, 2017) and
thus contribute to the differences between the two studies. (2) Enhanced transpiration in G4SSA
due to enhanced diffuse radiation reduces the evaporation difference in the two SRM schemes as
discussed above.

3.2 Surface and tropospheric ozone response

The ozone response is remarkably different in G4SSA and G4SSA-S. Global mean
surface ozone increases under G4SSA-S and decreases under G4SSA, relative to RCP6.0 (Fig.
6a). The total ozone column is dominated by stratospheric column ozone, and shows strong
reduction under G4SSA compared to RCP6.0, while the increase under G4SSA-S is small (Figs. 6b and 6d). The underlying upward trends of total column ozone as well as stratospheric ozone in all three scenarios are in line with the gradually declining stratospheric halogen content over time (Figs. 6b and 6d). The halogen loading in the three scenarios is the same, and more information can be found in Morgenstern et al. (2017). As there is less halogen in the stratosphere toward the end of the geoengineering, stratospheric ozone is recovering (Fig. 6d) and there is less reduction of the total ozone column in G4SSA relative to RCP6.0 (Fig. 6b). The agreement across the simulations concerning interannual and decadal variations is because of the imposed QBO and 11-year solar cycles in all the runs. The decreasing tropospheric ozone column and surface ozone after year 2060 in all scenarios results from decreases in global ozone precursor emissions following the RCP6.0 scenario (Young et al., 2013).

The surface ozone concentration distributions in the three scenarios are similar, with the highest concentration over the continents in the Northern Hemisphere (NH) (Fig. S6), while the concentration differences as well as the percentage difference between scenarios are spatially variable (Fig. 7 and Fig. S7). This highlights that the key driver behind the absolute surface ozone abundances is the underlying ozone precursor emissions following the RCP6.0 scenario. SRM is only a modulating factor, but depending on the SRM scheme even the sign of its impact can differ; global mean surface ozone concentrations in G4SSA are lower, relative to RCP6.0, whereas there are moderate surface ozone increases over the tropics (Fig. 7a). The strongest surface ozone reductions compared with RCP6.0 occur in NH mid-latitudes across all seasons (Figs. S8a-d) and Southern Hemisphere (SH) mid-to-high latitudes in MAM and JJA (Figs. S8b, c). In G4SSA-S, surface ozone also increases in the tropics relative to RCP6.0 (Fig. 7b), and this regional change is greater than in G4SSA (Fig. 7c). Surface ozone decreases, however, are
found at NH mid-latitudes over the continents during all seasons (Fig. S8e-h). Comparing the
two types of geoengineering experiments directly, surface ozone concentrations are generally
lower in G4SSA than in G4SSA-S (Fig. 7c), with peak differences in terms of absolute changes
(ppb) at SH mid-to-high latitudes in MAM and JJA (Fig. S8i, j) and at NH mid-to-high latitudes
in DJF (Fig. S8l).

3.3 Mechanisms of surface ozone change

Surface ozone concentrations are determined by chemical production and loss controlled
by emissions of ozone precursors and the composition of the atmosphere, loss due to surface
deposition of ozone, and transport of ozone from other regions of the atmosphere (Monks et al.,
2015). Since all scenarios considered here are based on the same RCP6.0 emission scenario of
ozone precursors, such as nitrogen oxide (NOx) and methane (CH4), the differences in surface
ozone must necessarily be driven by changes in climate in response to the geoengineering
interventions, which include changes in temperature, humidity, atmospheric dynamics, and the
photochemical environment. To understand the differences mechanistically, it is helpful to
consider the impact of geoengineering on the tropospheric ozone budget.

The upper part of Table 1 shows the sources (production and net transport from the
stratosphere (stratosphere-troposphere-exchange, STE)) and sinks (loss rates and dry deposition)
of tropospheric ozone. Both G4SSA and G4SSA-S show positive net chemical change of
tropospheric ozone (chemical production minus loss) and negative change in STE of ozone
relative to RCP6.0. However, the magnitude of these changes is significantly different.
Compared with RCP6.0, tropospheric ozone net chemical change increases by ~125 Tg yr\(^{-1}\) and
~40 Tg yr\(^{-1}\) in G4SSA and G4SSA-S, respectively, whereas STE of ozone decreases by ~140 Tg
yr\(^{-1}\) (~25%) and ~30 Tg yr\(^{-1}\) (~5%) in G4SSA and G4SSA-S, respectively. The positive net
chemical changes are the result of reductions in both chemical ozone production and loss under G4SSA and G4SSA-S relative to RCP6.0, with larger reductions in ozone loss reactions (Table 1). Specifically, G4SSA-S shows a ~90 Tg yr\(^{-1}\) larger decrease in ozone chemical production, whereas ozone loss budgets are reduced by similar magnitudes for the two SRM schemes (262.5 Tg yr\(^{-1}\) and 269.5 Tg yr\(^{-1}\)). Combining the chemical and transport changes, the tropospheric ozone budget decreases under G4SSA and increases under G4SSA-S relative to RCP6.0, which is consistent with the overall surface ozone changes.

The reasons for these specific changes are discussed in detail in the following two sections. Then, the impacts of the factors are combined to explain regional surface ozone differences, as shown in Fig. 7.

3.3.1 Chemical ozone production and loss in the troposphere

Changes in tropospheric water vapor concentrations and the tropospheric photolysis environment under G4SSA and G4SSA-S are key to understand the differences in tropospheric ozone production and loss. This result is consistent with results of a previous study for the case of solar geoengineering under a more idealized forcing scenario (Nowack et al., 2016).

To explain this, we briefly re-iterate that tropospheric ozone (O\(_3\)) production is driven by photolysis of nitrogen dioxide (NO\(_2\)) and the subsequent formation of ozone via a three-body-reaction with resulting ground state atomic oxygen O\((^3P)\) (Monks, 2005),

\[
\text{NO}_2 + h\nu (\lambda < 420 \text{ nm}) \rightarrow \text{NO} + \text{O}(^3\text{P}) \tag{1}
\]

\[
\text{O}(^3\text{P}) + \text{O}_2 + M \rightarrow \text{O}_3 + M \tag{2}
\]

where M is an inert collision partner (mostly molecular nitrogen). NO\(_2\) formation in turn is crucially dependent on the oxidation of NO by reaction with peroxides present in the troposphere, for example,
where $R$ represents general organic residues such as CH$_3$ (row 6 in Table 1). RO$_2$ in turn is produced by oxidation reactions between VOCs and the hydroxyl radical OH. Tropospheric OH is formed primarily by ozone photolysis and subsequent reaction of excited atomic oxygen O($^1$D) with water vapor,

$$O_3 + \text{hv (} \lambda < 320 \text{ nm)} \rightarrow O(^1D) + O_2$$  \hspace{1cm} (5)$$

$$O(^1D) + H_2O \rightarrow 2 \text{ OH}$$ \hspace{1cm} (6)

Reaction (6) competes with several other reactions due to the high reactivity of O($^1$D). However, most importantly, the majority of O($^1$D) is quenched by collision with inert molecules such as molecular nitrogen to ground state atomic oxygen O($^3$P), which subsequently undergoes reformation to O$_3$ via three-body-reaction (2). Therefore, tropospheric ozone production and loss is strongly linked to concentrations of water vapor and the photochemical environment (availability of UV) in the troposphere.

In the case of clean (low NO$_x$) environments, lower water vapor concentrations (mainly in the tropical region 30°N – 30°S) (Fig. S9) lead to less ozone loss via reactions (5) and (6) and additional reactions with the formed HOx species (r-O($^1$D)-H$_2$O, r-OH-O$_3$, and r-HO$_2$-O$_3$ in Table 1). This happens at the expense of more quenching of O($^1$D) and subsequent recycling of ozone, thus resulting in ozone increases. In contrast, in polluted (high NO$_x$) environments, less OH formation under lower atmospheric water vapor concentrations leads to reduced formation of RO$_2$ and HO$_2$. Therefore, less NO$_2$ is produced via reactions (3) and (4), resulting in less catalytic ozone production via reactions (1) and (2) (r-NO-HO$_2$ and e.g. r-CH$_3$O$_2$-NO in Table 1).
Consequently, ozone production is reduced in NOx-polluted environments under lower atmospheric water vapor concentrations.

With these fundamentals in mind, it is possible to understand the sign of the tropospheric ozone loss and production rate changes summarized in Table 1. Under both G4SSA and G4SSA-S, the key objective is to achieve surface temperature decreases. Tropospheric water vapor concentrations (or specific humidity) are strongly coupled to surface temperatures, because relative humidity does not change much with climate change (Soden and Held, 2006; Dessler and Sherwood, 2009), so that the surface cooling simultaneously reduces tropospheric specific humidity by 5-20% depending on region and altitude. As a result, less water vapor in both G4SSA and G4SSA-S reduces ozone chemical loss by ~150 Tg yr\(^{-1}\) through reactions (5) and (6). The resulting decrease in HO\(_x\) leads to further reductions in ozone loss, i.e., via reaction with OH (~20 Tg yr\(^{-1}\)) and HO\(_2\) (~60-70 Tg yr\(^{-1}\)). Overall, these water vapor/HO\(_x\)-related reactions explain ~90% of the overall reduction in ozone loss under SRM compared to a future RCP6.0 simulation.

The reduction in atmospheric humidity also affects ozone production, but to a smaller degree. Here, ozone production via reaction between NO and HO\(_2\) is the key factor in driving these changes, with reductions of ~55 and 120 Tg yr\(^{-1}\) for G4SSA and G4SSA-S, respectively. The signal of reduced OH production propagates through all other NO\(_x\)-catalyzed ozone production pathways involving RO\(_2\) via reactions (4) and subsequently (1) and (2). NO oxidation via the CH\(_3\)O\(_2\)-NO pathway decreases by ~27 and 49 Tg yr\(^{-1}\) in G4SSA and G4SSA-S. Changes in natural NO\(_x\) emissions by lightning play a minor role in comparison. In both SRM schemes, the reduction of lightning induced NO\(_x\) is not significant in most regions, and there is no significant difference between the two SRM schemes (Fig. S10).
The changes in chemical ozone production rates tend to be smaller in the sulfate G4SSA experiment than in the case of a solar constant reduction in G4SSA-S. There are three possible factors that contribute to this:

1. The entire reaction cycle depends on the availability of sunlight to photolyse O$_3$ and NO$_2$. Since SRM schemes modulate the intensity of sunlight (here by 1%) reaching the troposphere in order to mitigate tropospheric warming, this will necessarily also play a role in all changes to ozone production and loss reactions in our SRM simulations. More importantly, however, the sulfate injection geoengineering alters stratospheric ozone concentrations, which ultimately impacts the photochemical environment of the troposphere by changing radiative fluxes into the troposphere (DeMore et al., 1997; Nowack et al., 2016). For example, a reduced stratospheric column will help to stimulate the tropospheric photochemistry by allowing more radiation relevant reactions (1) and (5) to propagate into the troposphere.

2. Diffuse radiation under G4SSA promotes the photosynthesis rate and increases canopy transpiration (Fig. S4). Therefore, we expect that water vapor concentration over the continents with plants would be slightly higher in G4SSA relative to G4SSA-S (Fig. S11). Those regions with higher water vapor (East Asia, South Asia, North America, South Africa) are consistent with high NO$_x$ regions (Fig. S12). Therefore, the slightly smaller reduction of water vapor under G4SSA in the regions above increases ozone chemical production compared with G4SSA-S, and hence G4SSA shows less reduction of ozone chemical production than that in G4SSA-S.

3. Different biogenic VOC emissions under G4SSA and G4SSA-S, which, due to their central role in forming NO$_2$, are highly important for ozone production. In both scenarios, lower temperatures reduce the heat stress on the emitting plants and therefore reduces their VOC
emissions (Tingey et al., 1980; Sharkey and Yeh, 2001; Lathière et al., 2005; Bornman et al., 2015) (e.g., bio-emitted isoprene, Fig. S13). However, at the same time enhanced diffuse radiation under G4SSA increases biogenic VOC emissions compared with G4SSA-S (Wilton et al., 2011) (Fig. S13i, j, k and l). In Table 1, biogenic VOC-related ozone chemical production is generally very similar between G4SSA with G4SSA-S (e.g., r-ISOP O2-NO, r-MACRO2-NOa, r-MCO3-NO and r-TERPO2-NO), and contributes less than 2% to the overall difference between G4SSA and G4SSA-S.

3.3.2 Changes in stratosphere-troposphere exchange

Stratospheric chemical and dynamical changes can impact tropospheric ozone not only by changing the tropospheric photochemical environment, but also by changing the actual transport of ozone from the stratosphere into the troposphere (Hegglin and Shepherd, 2009; Neu et al., 2014). This can be either caused by changes in ozone concentrations in the stratosphere, or by changes in the rate of exchange of air masses between stratosphere and troposphere (i.e., the strength of the Brewer–Dobson (B-D) circulation and tropopause folds).

Fig. 8 shows seasonal latitude-height cross-sections of differences in ozone volume mixing ratios between G4SSA and RCP6.0 as well as G4SSA-S and RCP6.0 for altitudes above the 500 mb pressure level. Under G4SSA, heterogeneous reactions on the aerosol surfaces lead to increased halogen activation and with that an enhancement of ozone depletion in mid to high latitudes (60°-90° N/S) in the lower stratosphere (70-150 mb) (Tilmes et al., 2008, 2009, 2012; Heckendorn et al., 2009). On the other hand, heterogenous reactions reduce the NOx to NOy ratio, which results in an increase in ozone mixing ratios, mainly in the middle stratosphere (10-30 mb) (Tie and Brasseur, 1995) (Fig. 8a). In addition, changes in stratospheric temperature (warming in G4SSA and cooling in G4SSA-S) also change the photochemistry of ozone.
Altogether, this results in year-round lower stratospheric ozone loss worldwide that peaks during the return of sunlight at high SH latitudes (Fig. S14). In comparison, the solar reduction in G4SSA-S does not enhance stratospheric heterogeneous reactions. The much smaller change (increase) in ozone (Fig. 8b) is driven by the change of homogeneous chemistry due to slight temperature reduction (Fig. 4b). However, in Fig. 4b, there is a slight warming around 50 mb in the tropics, where ozone concentration also shows a stronger increase (Fig. 8b). As tropospheric cooling results in a slow-down of the B-D circulation (Fig. 9b) (Lin and Fu, 2013; Nowack et al., 2015; Shepherd and McLandress, 2011), there is an increase of ozone in the tropical upwelling region, which leads to increasing temperatures there as ozone is a strong shortwave and longwave absorber. The net result is small ozone increases in the tropical lower stratosphere and decreases in both extratropical lower stratospheres (Fig. 8b).

Age of air is used to indicate the strength of the B-D circulation (Fig. 9). Here, it is calculated relative to the zonal mean of 1°N at 158.1 mb (Garcia and Randel, 2008; Waugh, 2002). Older air indicates a slow-down of the B-D circulation. Compared with RCP6.0, both G4SSA and G4SSA-S show older air in the stratosphere indicating a slowdown of the circulation. The cooling effect in two SRM scenarios correlates with a weakening of tropical upwelling. However, in G4SSA, the heating of the tropical stratosphere results in enhanced lifting, which counteracts the weakening of the B-D circulation (Figs. 9a and 9c). Previous studies show controversial results on how the B-D circulation changes due to extra aerosols in the atmosphere. Aquila et al. (2012) modeled stronger tropic upwelling after the eruption of Mt. Pinatubo, and other studies also found enhanced simulated B-D circulation after this volcano eruption (Aquila et al., 2013; Pitari et al., 2016). The differences between previous studies and our result may be because some previous studies used fixed ozone, with different stratospheric heating rates. In
addition, in previous studies, the QBO was interactively simulated and the models had a higher model top. However, with extra black carbon in the stratosphere, the tropical upwelling weakens due to the simultaneous effect of tropospheric cooling (Shepherd and McLandress, 2011; Mills et al., 2014). We hope that future studies will address the potential model-dependency of this result.

The sum of both effects, stratospheric chemical changes and the impact of B-D circulation change on STE of ozone, is shown in Fig. 10. In G4SSA, ozone transported from the stratosphere to the troposphere is significantly decreased by ~25% relative to RCP6.0. In G4SSA-S the reduction is small. Since the air mass transported from the stratosphere to the troposphere is reduced in both scenarios, and is even more strongly reduced under G4SSA-S (Fig. 9), we find that the enhanced stratospheric ozone depletion in G4SSA is the dominant reason for the strong reduction of STE of ozone. This is also supported by a stratospheric ozone tracer from the model, $O_3^{\text{Strat}}$, which is set to ozone mixing ratios in the stratosphere and experiences only chemical loss in the troposphere without chemical production (Fig. S15). We thus conclude that the significant changes in STE of ozone in G4SSA are mainly driven by enhanced stratospheric ozone depletion catalysed through the aerosols (see also Table 1).

### 3.3 Balance of the different mechanisms and uncertainties

In summary, there are two main factors that determine the tropospheric ozone responses in our SRM and RCP6.0 simulations: (a) changes in tropospheric ozone chemical production/loss due to water vapor changes and impacts on the photochemical environment of the troposphere as a result of both changes in stratospheric ozone and (to a smaller degree) the actual dimming of sunlight depending on the geoengineering scheme, and (b) changes in stratosphere-troposphere exchange of ozone.
These factors can also be used to explain the big picture behind the surface ozone changes shown in Fig. 7. In G4SSA-S the reduced tropospheric humidity leads to stronger reductions of ozone loss than the decreases in ozone production, leading to global increases in surface ozone, but particularly in clean air oceanic environments in the tropics. This net increase in ozone chemical change is not cancelled out by the slight reduction of ozone transport from the stratosphere (Fig. 10). In G4SSA, reduction of ozone transport from the stratosphere is the controlling factor, which overwhelms the increase in net ozone production. The effect is particularly pronounced at mid-to-high latitudes (Fig. S14a), thus giving rise to surface ozone decreases there (Fig. 7). In contrast, the effect of reduced tropospheric humidity is relatively more important in the tropics than in other regions, which results in a local increase in surface ozone under G4SSA. Regionally HOx-NOx induced reductions in ozone production (Table 1) can become important to explain surface ozone decreases in NOx-polluted land areas in the NH for both scenarios (Figs. 7 and S6). Further minor contributions to the differences in surface ozone between G4SSA and G4SSA-S are a consequence of changes in water vapor due to regional canopy transpiration effects and biogenic VOC emissions (e.g., isoprene, Table 1 and Fig. S13).

This study may be biased by the following factors: (1) using prescribed stratospheric aerosols does not allow the simulation of the full interactions between chemistry, aerosol microphysics, and dynamics. A fully interactive model including those interactions would be important. (2) The vertical resolution is not sufficient to produce an interactive QBO in the model used, which may also affect transport processes. (3) The model does not include the scattering effect of aerosols on tropospheric photolysis rates, which might lead to an overestimate of the UV enhancement in the troposphere.
4. Conclusions

Tropospheric ozone changes are to be expected in a geoengineered climate with consequent impacts on air pollution and crop yields. However, for the scenarios considered here, solar and sulfate geoengineering could have entirely different impacts, even in terms of the sign of the response, a rare discrepancy for a surface signal between these two types of geoengineering. There have been many studies using solar irradiance reductions to illustrate SRM. However, it turns out that different SRM strategies would have different impacts on hydrology, atmospheric dynamics, the terrestrial carbon sink, and as investigated in this paper, tropospheric chemistry. These results also depend on the scenario of future ozone precursor and halogen emissions.

We have identified and explained the mechanisms by which stratospheric sulfate geoengineering would change surface ozone concentrations. We find that geoengineering might have the potential to significantly reduce some climate impacts, but it cannot fix the problem of air pollution. To reduce air pollution effectively, changes in surface emissions are key, with changes in climate (including geoengineering) being only a modulating factor (Monks et al., 2015; Stevenson et al., 2013; Young et al., 2013). More importantly, the surface ozone reduction between 2030 and 2070 in G4SSA is primarily the result of decreased STE of ozone following ozone depletion in the stratosphere. The rather mild pollution benefit under the RCP6.0 background would thus be bought at the expense of a delay of stratospheric ozone recovery, which would result in enhanced UV penetration to Earth’s surface and with that serious impacts on human health (e.g., skin cancer) and the ecosystem. In the future, potential increases of stratospheric ozone as a result of geoengineering may result in an increase of surface ozone,
causing more ozone pollution. However, further analysis on air pollutants other than ozone are needed.

As shown by Pitari et al. (2014), impacts on ozone from stratospheric geoengineering can be highly model dependent. Therefore, we consider the results here to be a GeoMIP testbed experiment, and encourage others to compare our results to those from other climate models to evaluate the robustness of the results presented here.

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Table 1. Tropospheric ozone production and loss rates (Tg yr\(^{-1}\)) over the period of years 2030-2069 (average of three ensemble members). For chemical ozone production and ozone loss the net impacts of only the most important reaction pathways are listed.

<table>
<thead>
<tr>
<th>Pathway</th>
<th>RCP6.0</th>
<th>G4SSA</th>
<th>G4SSA-S</th>
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<tr>
<td>O(_3) Net Chemical Change</td>
<td>346.1</td>
<td>472.7</td>
<td>384.8</td>
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<td>O(_3) Tropospheric Dry Deposition</td>
<td>901.5</td>
<td>891.5</td>
<td>909.4</td>
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<td>O(_3) STE*</td>
<td>555.4</td>
<td>418.8</td>
<td>524.6</td>
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<td>O(_3) Production</td>
<td>4895.8</td>
<td>4764.0</td>
<td>4671.8</td>
</tr>
<tr>
<td>r-NO-HO(_2)</td>
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<td>3031.0</td>
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<tr>
<td>r-CH(_2)O(_2)-NO</td>
<td>1132.3</td>
<td>1105.2</td>
<td>1083.1</td>
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<tr>
<td>r-PO(_2)-NO</td>
<td>21.8</td>
<td>20.1</td>
<td>19.9</td>
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<tr>
<td>r-CH(_3)CO(_3)-NO</td>
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<tr>
<td>r-C(_2)H(_5)O(_2)-NO</td>
<td>6.6</td>
<td>6.7</td>
<td>6.7</td>
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<tr>
<td>0.92* r-ISOPO(_2)-NO</td>
<td>149.8</td>
<td>135.3</td>
<td>134.0</td>
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<td>r-MACRO(_2)-NO(_a)</td>
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<td>4.1</td>
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<td>r-jonitr</td>
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<td>1.2</td>
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<td>O(_3) Loss</td>
<td>4421.1</td>
<td>4158.6</td>
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<tr>
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<td>r-OH-O(_3)</td>
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<td>r-HO(_2)-O(_3)</td>
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<td>11.5</td>
<td>11.5</td>
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<td>71.4</td>
<td>58.0</td>
<td>57.6</td>
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<td>0.8*r-MVK-O(_3)</td>
<td>18.6</td>
<td>15.5</td>
<td>15.7</td>
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<tr>
<td>0.8*r-MACR-O(_3)</td>
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<td>2.9</td>
<td>2.9</td>
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<td>r-C(_10)H(_16)-O(_3)</td>
<td>37.0</td>
<td>31.5</td>
<td>31.6</td>
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</table>

*O\(_3\) STE is ozone transported through the Stratosphere-Troposphere Exchange. We calculated this value using equation –

\[
O_{3, STE} + O_{3, net tropospheric chemical change} + O_{3, dry tropospheric deposition} = 0
\]

Tropospheric ozone is defined as ozone concentration less than 150 ppb.
Figure 1. (a) Global averaged annual net solar flux on the top of the atmosphere (W/m²) and (b) downwelling solar flux on the top of the atmosphere (W/m²). Dashed lines are ensemble members, and solid lines are the average of three ensemble members. Geoengineering starts at January 1st 2020 and ends at January 1st 2070. The 11-year periodicity is imposed as a prediction of the sunspot cycle. In (b) the G4SSA curve exactly covers the RCP6.0 curve.
Figure 2. (a) Global averaged annual surface air temperature (K), (b) global averaged annual precipitation (mm/day), and (c) downwelling surface solar radiation (W/m²). Dashed lines are ensemble members, and solid lines are the average of the three ensemble members. Geoengineering starts at 1 January 2020 and ends at 1 January 2070.
Figure 3. Global maps of surface temperature differences (K) between (a) G4SSA and RCP6.0, (b) G4SSA-S and RCP6.0, and (c) G4SSA and G4SSA-S over the period 2030-2069. Hatched regions are areas with $p > 0.05$ (where changes are not statistically significant based on a paired $t$-test).
Figure 4. Zonal mean temperature differences (K) in the geoengineering experiments (a) G4SSA minus RCP6.0, (b) G4SSA-S minus RCP6.0, and (c) G4SSA minus G4SSA-S. These are averaged for three ensemble members for years 2030-2069. Hatched regions are insignificant, with $p > 0.05$. The yellow dashed lines in (a) and (c) are the injected sulfate aerosol (surface area density =10 µm$^2$ cm$^{-3}$).
**Figure 5.** Energy flux on the top of the atmosphere (TOA) / the top of the model top (TOM) (a) and at the surface (surf) (b), shown as G4SSA minus RCP6.0 and G4SSA-S minus RCP6.0 for 2030-2069. For all fluxes, downwelling is positive. ΔSW is the net shortwave flux, ΔLW is the net longwave flux, R is the sum of ΔSW and ΔLW, SH is sensible heat, LH is latent heat, and Δsurf is the sum of ΔSWsurf, ΔLWsurf, SH, and LH; as is all sky and cs is clear sky.
Figure 6. (a) Global averaged annual surface ozone concentrations (ppb), (b) total column ozone (DU), (c) tropospheric column ozone (DU), and (d) stratospheric column ozone (DU). Ozone concentration of 150 ppb is used as the boundary of tropospheric ozone and stratospheric ozone. Dashed lines are ensemble members, and solid lines are the average of the three ensemble members. Geoengineering starts at 1 January 2020 and ends at 1 January 2070.
Figure 7. Global maps of surface ozone concentration differences (ppb) between (a) G4SSA and RCP6.0, (b) G4SSA-S and RCP6.0, and (c) G4SSA and G4SSA-S for 2030-2069. Hatched regions are insignificant, with $p > 0.05$. 
Figure 8. Zonal mean ozone concentration differences (ppb) in the geoengineering experiments, averaged for three ensemble members for 2030-2069. Hatched regions are insignificant, with $p > 0.05$. The yellow dashed lines in (a) and (c) are the upper and lower limits of the prescribed sulfate aerosol (surface area density = 10 $\mu$m$^2$ cm$^{-3}$).
Figure 9. Zonal mean age of air differences (years) between (a) G4SSA and RCP6.0, (b) G4SSA-S and RCP6.0, and (c) G4SSA and G4SSA-S. They are averaged for three ensemble members for 2030-2069. Hatched regions are insignificant, with $p > 0.05$. 
Figure 10. Global annual averaged ozone transported from the stratosphere to the troposphere (STE of ozone) in Tg yr$^{-1}$. Geoengineering starts at 1 January 2020 and ends at 1 January 2070.