From the middle stratosphere to the surface, using nitrous oxide to constrain the stratosphere-troposphere exchange of ozone

Daniel J. Ruiz¹ and Michael J. Prather¹

¹Department of Earth System Science, University of California, Irvine, CA 92697-3100, USA

Correspondence to: Daniel J. Ruiz (djruiz@uci.edu)

Abstract

Stratosphere-troposphere exchange (STE) is an important source of tropospheric ozone, affecting all of atmospheric chemistry, climate, and air quality. Observations and the theoryThe study of tracer correlations provide only coarse (±20%) global-mean constraints. For impacts needs STE fluxes to be resolved by latitude and month, and for this we rely on global chemistry-transport models-(CTMs), and unfortunately, these, whose results diverge greatly. Overall, we lack guidance from model-measurement metrics that inform us about processes and patterns related to the STE flux of ozone, (O_3) . In this work, we use modeled tracers (N₂O, CFCl₃) whose distributions and budgets can be constrained by satellite and surface observations, allowing us to follow stratospheric signals across the tropopause. The satellite derived photochemical loss of N_2O on annual and quasi-biennial cycles can be matched by the models. The STE flux of N_2O depleted air in our CTM chemistry transport model drives surface variability that closely matches observed fluctuations on both annual and guasi-biennial cycles, confirming the modeled flux. The observed tracer correlations between N₂O and O₃ in the lowermost stratosphere provide a seasonal, hemispheric scaling of the N₂O STE flux to that of O₃. For N₂O and CFCl₃, we model greater southern hemispheric STE fluxes, a result supported by some metrics, but counter to prevailing theory of wave-driven stratospheric circulation. The STE flux of O₃, however, is predominantly northern hemispheric, but observational constraints showevidence shows that this is only caused by the Antarctic ozone hole. Here we show that reducing southern hemispheric O₃ STE by 14%. Our best estimate of the current STE O₃ flux based on a range of constraints is 400 Tg(O₃)/yr with a one-sigma uncertainty of $\pm 15\%$ and with a NH:SH ratio ranging from 50:50 to 60:40. We identify a range of observational metrics founded on observations that can better constrain the modeled STE O3 flux which will help guidein future model assessments.

1. Introduction & Background

The influx of stratospheric ozone (O₃) into the troposphere affects its distribution, variability,
lifetime, and thus its role in driving climate change and surface air pollution (Zeng et al., 2010;
Hess et al., 2015; Williams et al., 2019). The net stratosphere-to-troposphere exchange (STE)
flux of O₃ has a regular seasonal cycle in each hemisphere that is an important part of the
tropospheric O₃ budget (Stohl et al., 2003). Such fluxes are not directly observable, and we rely
on observational estimates using trace-gas ratios, in particular the O₃:N₂O ratio in the lower
stratosphere (Murphy and Fahey, 1994; McLinden et al., 2000), or dynamical calculations using
measured/modeled winds and O₃ abundances (Gettelman et al., 1997; Olsen et al., 2004; Yang et

47 al., 2016). The uncertainty in these estimates does not effectively constrain the wide range found 48 in the models being used to project future ozone (Young et al., 2013, 2018; Griffiths et al., 49 2021). Here we present the case for using the observed variations in nitrous oxide (N_2O) from 50 the middle stratosphere to the surface in order to constrain the STE flux of O₃. A similar case 51 has been made for the radionuclide ⁷Be (Liu et al., 2016), but N₂O has a wealth of model-52 observation metrics on hemispheric, seasonal, and interannual scales that constrains its STE flux very well (Prather et al., 2015; Ruiz et al., 2021)._ 53 54 55 Ozone-rich stratospheric air has been photochemically aged and is depleted in trace gases such as 56 N₂O and chlorofluorocarbons (CFCs). For these trace gases, the overall circulation from 57 tropospheric sources to stratospheric destruction and back is part of the lifecycle that maintains 58 their global abundance (Holton, 1990). For N_2O and CFCs, this cycle of (i) loss in the middle to 59 upper stratosphere, (ii) transport to the lowermost stratosphere (Holton et al., 1995), and then (iii) 60 influx into the troposphere produces surface variations not related to surface emissions 61 (Hamilton and Fan, 2000; Nevison et al., 2004; Hirsch et al., 2006; Montzka et al., 2018; Ray et 62 al., 2020; Ruiz et al., 2021). In this work we relate our modeled STE fluxes to variations at the 63 surface and throughout the stratosphere, linking the fluxes of N2O to O3 through stratospheric 64 measurements. Our goal is to develop a set of model metrics founded on observations that are 65 related to the STE O₃ flux and can be used with an ensemble of models to determine a better, 66 constrained estimate for the flux, including seasonal, interannual, and hemispheric patterns. This 67 approach is similar to efforts involving the ozone depletion recovery time (Strahan et al., 68 2011) and projections of future warming (Liang et al., 2020; Tokarska et al., 2020). 69 In a previous work (Ruiz et al., 2021, hence R2021) we showed that historical simulations with 70 three chemistry transport models (CTMs) were able to match the interannual surface variations 71 72 observed in N₂O. These were clearly driven by the stratospheric quasi-biennial oscillation 73 (QBO) which appears to be the major interannual signal in stratospheric circulation and STE 74 (Kinnersley and Tung, 1999; Baldwin et al., 2001; Olsen et al., 2019). In this work, we calculate the monthly latitudinal STE fluxes of O₃, N₂O, and CFCl₃ (F11), establish a coherent picture 75 76 relating fluxes to observed abundances, and summarize the methods in Section 2. In section 3, 77 we examine the annual and interannual cycles as well as geographic patterns of modeled STE 78 flux. In section 4, we relate the surface variability of N₂O to its STE flux. We find some 79 evidence to support our model result that the STE flux of depleted-N2O air is greater in the 80 southern hemisphere than in the northern, thus altering the asymmetry in surface emissions in the 81 source inversions (Nevison et al., 2007; Thompson et al., 2014). In section 5, we examine the 82 lowermost stratosphere to understand the large north-south asymmetry found in O_3 STE versus 83 N₂O or F11 STE, and find a clear signal of the Antarctic ozone hole in STE. In section 6, we 84 examine the consistency of the model calculations of STE flux and derive a best estimate for the O3 flux from this and previous studies. We summarize a sequence of model metrics, primarily 85 86 using O_3 and N_2O , that can narrow the range in the tropospheric O_3 budget terms for the multi-87 model intercomparison projects used in tropospheric chemistry and climate assessments. 88 89 2. Methods 90

91 <u>The modeled STE fluxes here are calculated with the UCI (University of California Irvine) CTM</u>
 92 driven by 3-hour forecast fields from the European Centre for Medium-range Weather Forecasts

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93 (ECMWF) Integrated Forecast System (IFS Cycle 38r1 T159L60) for years 1990-2017, as are 94 the calculations in R2021. The CTM uses the IFS native 160x320 Gauss grid (~1.1°) with 60 95 layers, about 35 of which are in the troposphere. The stratospheric chemistry uses the linearized 96 model Linoz v3 and includes O3, N2O, NOy, CH4, and F11 as transported trace gases (Hsu and 97 Prather, 2010; Prather et al., 2015; Ruiz et al., 2021). There is no tropospheric chemistry, but 98 rather a boundary-layer e-fold to a specified abundance, or a surface boundary reset to an 99 abundance. Equivalent effective stratospheric chlorine levels are high enough to drive an 100 Antarctic ozone hole, which is observed throughout this period. Thus, the ozone-hole chemistry 101 in Linoz v3 is activated for all years, and the amount of O3 depleted depends on the Antarctic 102 meteorology of that year (Hsu and Prather, 2010). 103 104 The STE flux is calculated using the e90 definition of tropospheric grid cells (Prather et al., 105 2011) and the change in tropospheric tracer mass from before to after each tracer transport step as developed at UCI (Hsu et al., 2005; Hsu and Prather, 2009; Hsu and Prather, 2014). This 106 107 method is precise and geographically accurate for O3 and is self-consistent with a CTM's tracer-108 transport calculation (Tang et al., 2013; Hsu and Prather, 2014). Extensive comparisons with 109 other methods of calculating STE are shown in Hsu and Prather (2014). Annual-mean STE 110 fluxes are calculated from the full 28-year (336 month) time series as 12-month running means, 111and the annual cycle of monthly fluxes is the average of the 28 values for each month.

1. Introduction & Background

The influx of stratospheric ozone (O2) into the troposphere affects its distribution, variability, 116 117 lifetime, and thus its role in driving climate change and surface air pollution. The net 118 stratosphere to troposphere exchange (STE) flux of O₃-has a regular seasonal cycle in each 119 hemisphere that is an important part of the tropospheric O₃ budget (Stohl et al., 2003). Such 120 fluxes are not directly observable, and we rely on observational estimates using trace gas ratios 121 in the stratosphere (McLinden et al., 2000; Murphy & Fahey, 1994) or dynamical calculations 122 using measured/modeled winds and O3 abundances (Gettleman et al., 1997; M. A. Olsen et al., 123 2004; Yang et al., 2016). The uncertainty in these estimates does not effectively constrain the 124 wide range of the models being used to project future ozone (Young et al., 2013, 2018). Here we 125 present the case for using the observed variations in nitrous oxide (N2O) from the middle 126 stratosphere through to the surface in order to constrain the STE flux of O₃. A similar case has 127 been made for the radionuclide ⁷Be (Liu et al., 2016), but N₂O has a wealth of model observation 128 metrics on hemispheric, seasonal, and interannual scales that constrains its STE flux very well.-129 130 Ozone-rich stratospheric air has been photochemically aged and is depleted in trace gases such as 131 N2O and chlorofluorocarbons (CFCs).-For these trace gases, the overall circulation from 132 tropospheric sources to stratospheric destruction and back is part of the lifecycle that maintains 133 their global abundance (Holton, 1990). For N2O and CFCs, this cycle of (i) loss in the middle to 134 upper stratosphere, (ii) transport to the lowermost stratosphere (Holton et al., 1995), and then (iii) 135 influx into the troposphere produces surface variations not related to surface emissions 136 (Hamilton & Fan, 2000; Hirsch et al., 2006; Montzka et al., 2018; C. D. Nevison et al., 2004; 137 Ray et al., 2020; Ruiz et al., 2021). In this work we relate our modeled STE fluxes to variations

138 at the surface and throughout the stratosphere, linking the fluxes of N₂O to O₂ through

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139 stratospherie measurements. Our goal is to develop a set of model metrics founded on 140 observations that are related to the STE O2-flux and can be used with an ensemble of models to 141 determine a better, constrained estimate for the flux, including seasonal, interannual, and 142 hemispheric patterns. This approach is similar to efforts involving the ozone depletion recovery time (Strahan et al., 2011) and climate projections (Liang et al., 2020; Tokarska et al., 2020). 143 144 145 In a previous work (Ruiz et al., 2021), we showed that historical simulations with 146 three chemistry transport models (CTMs) were able to match the interannual surface variations 147 observed in N₂O. These were clearly driven by the stratospheric quasi biennial oscillation 148 (QBO) which appears to be the major interannual signal in stratospheric circulation and 149 STE (Baldwin et al., 2001; Kinnersley & Tung, 1999; M. A. Olsen et al., 2019). In this work, we 150 calculate the monthly latitudinal STE fluxes of O₃, N₂O, and CFCl₂-(F11) and establish a 151 coherent picture relating fluxes to observed abundances. In section 2, we examine the annual 152 and interannual cycles as well as geographic patterns of modeled STE flux. In section 3, we 153 relate the surface variability of N2O to its STE flux. We find some evidence to support our 154 model result that the STE flux of depleted N2O air is greater in the southern hemisphere than in 155 the northern, thus altering the asymmetry in surface emissions in the source inversions (Nevison 156 et al., 2007; Thompson et al., 2014). In section 4, we examine the lowermost stratosphere to 157 understand the large north south asymmetry found in O₂-STE versus N₂O or F11 STE, and find a 158 clear signal of the Antarctic ozone hole in STE. In section 5, we summarize the sequence of 159 model metrics, primarily using O₃ and N₂O, that that will usefully narrow the range in the 160 tropospheric O₂-budget terms like STE, for the multi-model intercomparison projects used in 161 tropospheric chemistry and climate assessments. 162 163

2. Annual and interannual cycles of modeled STE flux

165 The modeled STE fluxes here are calculated with the UCI-CTM driven by 3-hour forecast fields from the ECMWF Integrated Forecast System (IFS; Cy38r1 T159L60), as are the calculations in 166 167 R2021. The CTM uses the IFS native 160x320 Gauss grid (~1.1°) with 60 layers, about 35 in 168 the troposphere. The stratospheric chemistry uses the linearized model Linoz v3 and includes 169 O₃, N₂O, NO₄, CH₄, and F11 as transported trace gases (Hsu & Prather, 2010; Prather et al., 170 2015; Ruiz et al., 2021).- There is no tropospheric chemistry, but rather a boundary layer e-fold 171 to a specified abundance, or a surface boundary reset to an abundance. The STE flux is calculated using the e90 definition of tropospheric grid cells (Prather et al., 2011) and the 172 173 in tropospheric tracer mass from before to after each tracer transport step-(Hsu et al., 2005; Hsu 174 and Prather, 2009). This method is extremely robust for O₃ and self consistent with a CTM's 175 tracer transport (Hsu & Prather, 2014; Tang et al., 2013). 176

178 2.1. Model STE and tracer methods

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180 R2021 modeled the surface signal of stratospheric loss with the decaying tracers, N2OX and 181 F11X (e.g., (Hamilton and Fan, 2000; Hirsch et al., 2006). These X-tracers have the identical stratospheric chemical loss frequencies as N2O and CFCl3, respectively, but no surface sources 182 183 and are therefore affected only by the stratospheric sink and atmospheric transport. The multi-

184 decade (F11X) to century (N2OX) decays are easily rescaled on a month by month basis (using Formatted: Indent: Left: 0.25"

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186 320 ppb. We treat F11X like N2OX with the same initial conditions and molecular weight (i.e., 187 Budgets for N2OX are reported, as in N₂O studies (Tian et al., 2020), as Tg = TgN with 2 N's per 188 molecule).of N as N2O. These rescaled N2OX and F11X tracers we designate are designated 189 simply as N2O (not N2O) and F11. Our F11-derived STE fluxes are thus unrealistically large 190 compared to current CFCl₃ fluxes, but they can be easily compared with our N2OXN2O results. 191 192 Unfortunately, calculating When trying to calculate the STE flux of N2OX and F11X using N2O-193 depleted air across the tropopause, we found that the Hsu method was numerically noisy because 194 their gradients the gradient across the trop pause are minimal, unlike that of O_3 . We thus, was negligible. Thus, for this work we created the complementary tracers cN2OX and cF11X. For: 195 196 for each kg of the X-tracer (i.e., N2OX) destroyed by photochemistry, 1 kg of its complementary 197 tracer (cN2OX) is created. Air parcels that are depleted in N2OX (F11X) are therefore rich in 198 cN2OX (cF11X). After crossing the tropopause, cN2OX and cF11X are removed through rapid 199 uptake in the boundary layer, thus creating sharp gradients at the tropopause- in parallel with that 200 of O₃. As a check, we compared the boundary layer sinksinks of the c-tracers with their e90-201 derived STE fluxes and find that their sums are identical. The c-tracers and their STE fluxes 202 are rescaled as their corresponding-are the X-tracers to give them a stationary time series 203 corresponding to a tropospheric abundance of 320 ppb, we for their parallel X tracers. We designate these scaled tracers simply as cN2O and cF11. The inclusion of these new c-tracers 204 205 provides the missing link in R2021 by directly connecting the stratospheric loss signals to STE 206 flux and subsequent surface variability. 207

a 12-month smoothing filter) to give stationary results and a tropospheric mean abundance of

2.2 Mean 3. Modeled STE fluxes

The STE fluxes calculated at every time step for each latitude longitude grid column are
 integrated in time and longitude to give latitude by month resolved fluxes for years 1990 2017.
 3.1 Global and hemispheric means

Equivalent effective stratospherie chlorine levels are high enough to drive an Antaretic ozone hole, which is observed throughout this period. Thus, the ozone hole chemistry in Linoz v3 is activated for all years, and the amount of O₂-depleted depends on the Antaretic meteorology of that year. Annual mean STE fluxes are calculated from the full 28 year (336 month) time series, and monthly mean fluxes are calculated from the 28 values for each month.

220 221 The 28-year mean of global O_3 STE is 390±16 Tg/yr (positive flux means stratosphere to 222 troposphere, the \pm values are the standard deviation of the 28 annual means), that and do not 223 represent uncertainty). This value is well within the uncertainty in the observation-based 224 estimates (Murphy and Fahey, 1994; Olsen et al., 2001), and far from the extreme ranges of the 225 34 models in the latest Tropospheric Ozone Assessment Report (Young et al., 2018), 150 to 940 226 Tg/yr. The global STE flux of cN2O is 11.5±0.7 Tg/yr, and that of cF11 is 23.5±1.5 Tg/yr. 227 These fluxes for cN2O and cF11 match the total long-term troposphere-to-stratosphere flux of 228 N2O and F11 as derived from their stratospheric losses. The cF11 budget is about twice as large 229 as cN2O, because F11 is photolyzed rapidly in the lower-middle stratosphere (~24 km) instead 230 of the upper stratosphere like N2O- (~32 km). The seasonal mean pattern of STE fluxes are

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shown in Figure 1. The large majority of STE flux enters the troposphere at 25° - 45° latitude in 231

232 each hemisphere, but there is a broadening of the northern flux to 65°N in Jun-Jul. The

233 importance of this region about the sub-tropical jet for STE is supported by satellite data where 234 stratospheric folding events (high O3 in the upper troposphere) are found at the bends of the jet

- 235 (Tang and Prather 2010, Atmos. Chem. Phys., 10, 9681-9688, 2010).
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Given the small STE fluxes in the core tropics, the northern hemisphere (NH) and southern 237

238 hemisphere (SH) fluxes are distinct. The annual mean of NH O3 STE is 208±11 Tg/yr (±

239 standard deviation over the 28 years) and is slightly larger than the SH mean of 182±11 Tg/yr.

240 This NH:SH ratio of 53:47 is typically found in other studies (Yang et al., 2016; Gettelman et al., 241

1997; Hsu and Prather, 2009; Yang et al., 2016), although some have higher ratios like 58:42 (Hegglin and Shepherd, 2009; Meul et al., 2018). In contrast, for cN2O and cF11, the NH flux 242

(5.1±0.4 Tg/yr and 10.6±0.8 Tg/yr, respectively) is smaller than the SH flux (6.4±0.5 Tg/yr and 243

12.9±1.0 Tg/yr, respectively), giving a NH:SH ratio of about 45:55. The established view on 244

245 STE is that the flux is wave-driven and under downward-sideways control, and thus the NH flux

is much greater than the SH flux (see Table 1 of Holton et al., 1995; also Appenzeller et al., 246

247 1996). Our unexpected results require further analysis including evidence for hemispheric

248 asymmetry in observations which is shown in section 4 along with other model metrics.

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Figure 1. The seasonal (latitude by month) cycle of STE flux (Tg/yr) for (a) O₃, (b) cN2O, and (c) cF11. Each month is averaged for years 1990-2017 (e.g., the 28 Januarys are averaged). The colorbar units are % of global, annual mean STE in each bin (1 month by ~1.1° latitude).

<u>3.</u>2.3 Seasonal cycle-of STE

The seasonal cycles of STE fluxes summed over global, NH, and SH are shown in Figure 2. The scales are given as the annual rate (as if the monthly rate were maintained for the year), and each species has a different axis. The right y-axes are kept at a N2O:F11 ratio of 1:2. Despite large differences in the stratospheric chemistry across all three species, the seasonal cycle of STE is highly correlated (>(Pearson's correlation coefficient cc > 0.98, except for SH O₃), indicating that all three enter the troposphere from a seasonally near-uniform mixture of O₃:N2O:F11 in the lowermost stratosphere.

Global STE peaks in June and reaches a minimum in November, but that merely reflects the 270 271 272 273 274 275 dominance of the NH seasonal cycle and hides the distinct patterns in each hemisphere. The two hemispheres have dramatically different seasonal amplitudes and somewhat opposite phases. NH peak STE for all 3 species occurs in the late boreal spring (May-JunJune), while that in the SH occurs at the start of austral spring (Sep-OctSeptember-October). In the NH O3 STE peaks a month before the c-tracers, and in the SH the whole annual cycle of O_3 is shifted a month earlier. The NH STE seasonal amplitude is very large for all species (~ 4:1 peakratio max-to-peakmin) 276 with exchange almost ceasing in the fall. In contrast, the SH STE is more uniform year-round 277 with seasonal amplitudes of a 1.5:1 ratio for cN2O and cF11, and 2.2:1 for O_3 . Other models 278 with similar NH and SH O₃ fluxes show different seasonal amplitudes and phasing (see Fig. 6 of 279 Tang et al., 2021), which will affect tropospheric O_3 abundances. It is important to develop 280 observational metrics that test the seasonality of the lowermost stratosphere related to STE 281 fluxes, and to establish monthly STE fluxes as a standard model diagnostic. 282

283 An interesting result here is the very tight correlation of the monthly cN2O and cF11 STE while 284 the O_3 STE is sometimes shifted. Loss of N₂O and F11 occurs at very different altitudes in the 285 tropical stratosphere (~32 km and ~24 km, respectively), but both have similar seasonality in

286 loss, driven mostly by the intensity of sunlight along the Earth's orbit (N₂O loss peaks in Feb and

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reaches a minimum in JulJuly, see Fig. 4 from Prather et al.,.. (2015). Photochemical losses of
N₂O and F11 drop quickly for air descending from the altitudes of peak loss in the tropics and
hence the relative cN2O and cF11 STE fluxes are locked in. O₃, however, continues to evolve
photochemically evolve from 24 km to 16 km (upper boundary of the lowermost stratosphere),
through net photochemical lossproduction in the tropics and loss at mid- and high-latitudes that
depends on sunlight and is thus seasonal. There may be observational evidence for the patterns
modeled here in the correlation of these three tracers in the lower (16-20 km) and lowermost (12-

294 16 km) extratropical stratosphere (see section 4).







Figure 2. The annual cycle of monthly STE (Tg/yr) of O_3 (black lines), cN2O (orange lines), and cF11 (blue lines). **(a)** Global STE fluxes, and **(b)** hemispheric STE fluxes (NH, solid lines; SH, dashed lines). Each month is averaged for years 1990-2017 (e.g., the 28 Januarys are averaged). Note the different y-axes for each tracer in each panel.

298 2.4<u>3.3</u> Interannual variability of STE

299 300 Interannual variability (IAV) of N₂O loss and its lifetime is associated primarily with the QBO 301 (most recently, R2021). When the QBO is in its easterly (westerly) phase the entire overturning 302 circulation is enhanced (suppressed) (Baldwin et al., 2001). This results in more (less) air rich in 303 N₂O and F11 being transported from the troposphere to the lower or middle stratosphere, thereby 304 increasing (decreasing) the N₂O and F11 sinks (Prather et al., 2015; Strahan et al., 2015). From 305 the tropical stratosphere, the overturning circulation transports air depleted in N₂O and F11 into 306 the lowermost extratropical stratosphere, where it enters the troposphere. R2021 showed that the 307 observed surface variability of N2O from this circulation can be modeled and has a clear QBO 308 signal, but one that is not strongly correlated with When the QBO is in its easterly (westerly) 309 phase the entire overturning circulation is enhanced (suppressed) (Baldwin et al., 2001). This results in more (less) air rich in N₂O and F11 being transported from the troposphere to the lower 310 311 or middle stratosphere, thereby increasing (decreasing) the N2O and F11 sinks (Prather et al., 312 2015; Strahan et al., 2015). From the tropical stratosphere, the overturning circulation transports 313 air depleted in N₂O and F11 into the lowermost extratropical stratosphere, where it enters the 314 troposphere. R2021 showed that the observed surface variability of N₂O from this circulation 315 can be modeled and has a clear OBO signal, but one that is not strongly correlated with the OBO 316 signal in stratospheric loss.

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 β 18 We generate the IAV of STE fluxes for O₃, cN2O, and cF11 in Figure <u>33abc</u> with panels for

319 global, NH, and SH. Values are 12-month running means, and so the first modeled point at

1990.5 is the sum of STE for Jan through Dec of 1990. In Figures 3bc, we <u>also</u> show the seasonal amplitude of STE with double-headed arrows on the left (O₃) and right (cN2O a

seasonal amplitude of STE with double-headed arrows on the left (O₃) and right (cN2O and
 cF11). In a surprising result, the large NH-SH differences in seasonal amplitude are not reflected

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in the IAV where NH and SH amplitudes are similar for all three tracers. The QBO modulationof the lowermost stratosphere and STE appears to be unrelated to the seasonal cycle in STE.

325 326 Global STE for all three tracers is shows QBO-like cycling throughout the 1990-2017 time 327 series: cN2O and cF11 are well correlated (-(cc - 0.9), but <u>either species with O₃ is much less</u> 328 so $(\leftarrow (cc < 0.7))$. The hemispheric breakdown provides key information regarding O₃. In the NH 329 the STE IAV is similar across all three tracers with high correlation coefficients (cc = 0.82 for 330 O₃-cN2O, 0.83 for O₃-cF11, and 0.94 for cN2O-cF11). Conversely in the SH, O₃ STE diverges from the c-tracer fluxes, showing opposite-sign peaks in 2003 and 2016. The corresponding SH 331 332 correlations are (cc = 0.38, 0.65, 0.85). The loss of correlation between cN2O and cF11 is 333 unusual: cN2O STE drifts downward relative to cF11 STE, particularly after 2007; nevertheless, 334 the fine structure after 2007 is well matched in both tracers.

In the SH, the massive loss of O₃ within the Antarctic vortex, when mixed with the extra-polar lowermost stratosphere will systematically shift the O₃ STE to lower values, with lesser impact on the cN2O and cF11 STE. The IAV of the Antarctic winter vortex in terms of the amount of O₃-that is deplete (World Meteorlogical Organization (WMO), 2018, figure 4-4)less impact on the cN2O and cF11 STE. The IAV of the Antarctic winter vortex, in terms of the amount of the cN2O and cF11 STE. The IAV of the Antarctic winter vortex, in terms of the amount of the cN2O and cF11 STE. The IAV of the Antarctic winter vortex, in terms of the amount of O₃ that is depleted (see Fig. 4-4 of WMO, 2018), appears to drive the decorrelation of the SH STE fluxes and is analyzed in section 4.

344 In the NH, the high variability of the Arctic winter stratosphere can modulate the total O₂-STE 345 flux (e.g., (Hsu & Prather, 2009) but appears to maintain the same relative ratio with the cN2O 346 and cF11 fluxes. The model results here indicate that there is no differential IAV chemical signal 347 in these NH, and that the lowermost stratosphere is still combining the same chemical mixtures 348 of air masses from year to year. We know there is a large IAV in the Arctic winter activation of 349 halogen driven O₃ depletion (Manney et al., 2020), but the magnitude is still much smaller than 350 in the Antarctic, and it may not reach into the lowermost stratosphere (<380K potential 351 temperature). This model accurately simulates Antarctic O_3 -loss (section 4), but we have not 352 evaluated it for Arctic loss, and the Arctic conditions operate closer to the thresholds initiating 353 loss where Linoz v3 chemistry may be inadequate. The same meteorology and transport model 354 with full stratospheric chemistry is able to simulate Arctic O₃ loss (Oslo's CTM2: Isaksen et al., 355 2012), and thus it will be possible to re-evaluate the NH IAV with such models or with 356 lowermost stratosphere tracer measurements. 357

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360 In the NH, the high variability of the Arctic winter stratosphere can modulate the total O₃ STE flux (e.g., Hsu and Prather, 2009) but appears to maintain the same relative ratio with the cN2O 361 362 and cF11 fluxes. Model results here indicate that in the NH, the IAV of O₃, cN2O, and cF11 STE 363 fluxes are synchronized, and thus the air masses entering the lowermost stratosphere have the 364 same chemical mixtures from year to year. We know that cold-temperature activation of 365 halogen-driven O₃ depletion in the Arctic winter at altitudes above 400 K (potential temperature) 366 can produce large IAV in column ozone (Manney et al., 2011); but the magnitude is still much 367 smaller than in the Antarctic; and it may not reach into the lowermost stratosphere (<380K 368 potential temperature). This model accurately simulates Antarctic O₃ loss (section 4), but we







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381 <u>2.5. From 3.4 The link from</u> stratospheric loss to STE <u>flux</u> 382

383 What is unusual about the very tight correlation of cN2O and cF11 STE fluxes is that the

photochemical loss of N2O and F11 occurs at very different altitudes in the tropical stratosphere,
 which are not in phase with respect to the QBO as shown in R2021 (their Fig. 2). The separate

386 phasing of cN2O and cF11 production is lost, presumably by diffusive tracer transport, by the

time they reach the extratropical lowermost stratosphere. The overall synchronization of the

STE fluxes implies that the absolute STE flux is driven primarily by variations in venting of the

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³⁹⁰ 1995; Appenzeller et al., 1996) rather than by variations in the chemistry of the middle
 ³⁹¹ stratosphere.

392

393 This disconnect between the chemical signals generated by the prominent QBO signature of 394 wind reversals-and, upwelling in the tropical stratosphere, and the STE fluxes is also clear in the 395 magnitude of the loss versus STE. For N₂O, the IAV of cN2O production has a range of ± 0.5 396 Tg/yr, whether from Microwave Limb Sounder (MLS) observations or the model; whereas the 397 IAV of cN2O STE flux is ± 1.1 Tg/yr. The same is true in relative terms for cF11. Thus, the 398 modulation of the lowermost stratosphere by the QBO, which is clearly a part of the overall 399 changes in stratospheric circulation related to the QBO (Tung & Yang, 1994; Kinnersley and 400 Tung, 1999), Thus, the modulation of the lowermost stratosphere by the QBO is clearly a part of 401 the overall changes in stratospheric circulation related to the QBO (Tung and Yang, 1994a;

402 <u>Kinnersley and Tung, 1999) and</u> is the dominant source of IAV for these three greenhouse gases.

403 404

405 **2.6<u>3.5</u>**. The QBO signal of STE 406

407 To examine the QBO cycle in STE flux, we build a composite pattern (see R2021, Fig. 3 of N₂O 408 surface variations), by synchronizing the STE IAV in Figure 2 with the QBO cycle. The sync 409 point (offset = 0 months) is taken from one of the standard definitions of the OBO phase change, 410 i.e., the shift in sign of the 40 hPa tropical zonal wind from easterly to westerly (Newman, 411 2020). The 1990-2017 model period has 12 QBO cycles, but we restrict our analysis here to 412 vears 2001-2016 to overlap with the observed surface N₂O data. This period includes seven 413 OBO phase transitions (01/2002, 03/2004, 04/2006, 04/2008, 08/2010, 04/2013, 07/2015), but 414 the QBO centered on 08/2010 is highly anomalous (Coy et al., 2017; P. Newman et al., 2016; 415 Osprey et al., 2016), and we remove it from our comparison. The resulting QBO composites for 416 NH and SH in Figure 4 span 28 months. 417

418 In the NH, the QBO modulation of all three tracers is similar: STE flux begins to increase at an 419 offset of -8 months and continues to increase slowly for a year, peaking at offset = +4420 To examine the QBO cycle in STE flux, we build a composite pattern (see R2021, Fig. 3 of N_2O 421 surface variations), by synchronizing the STE IAV in Figure 2 with the QBO cycle. The sync 422 point (offset = 0 months) is taken from one of the standard definitions of the QBO phase change, i.e., the shift in sign of the 40-hPa tropical zonal wind from easterly to westerly (Newman, 423 2020). The 1990-2017 model period has 12 QBO cycles, but we restrict our analysis here to 424 425 years 2001-2016 to overlap with the observed surface N2O data. This period includes seven 426 QBO phase transitions (01/2002, 03/2004, 04/2006, 04/2008, 08/2010, 04/2013, 07/2015), but 427 the observed surface N₂O is highly anomalous during the QBO centered on 08/2010 (R2021), so 428 we remove it from our comparison for consistency with R2021 (see their Fig S4d). The resulting 429 QBO composites for NH and SH in Figure 4 span 28 months. 430

431 In the NH, the QBO modulation of all three tracers is similar: STE flux begins to increase at an

432 <u>offset of -8 months and continues to increase slowly for a year, peaking at an offset of +4</u>

 $\begin{array}{l} \mbox{433} \\ \mbox{months}; \mbox{ thereafter it decreases more rapidly in about 1/2 year (offset = +10). The rise-and-fall \\ \mbox{cycle takes about 18 months. In the SH, the pattern for cN2O and cF11 is more sinusoidal and is \\ \end{array}$

435 shifted later by ~3 months. The SH amplitude of the c-tracers is slightly larger relative to the

- 436 hemispheric mean flux than in the NH, and thus the SH QBO signal is larger than the NH by
- 437 about 40%. Thus, over the typical QBO cycle centered on the sync point, more depleted N₂O
- 438 and F11 is entering the SH than in the NH. For O₃, the SH modulation of STE is irregular and
- 439 reduced compared with the NH. Our hypothesis here, consistent with the annual cycle of STE
- 440 (Figure 1), is that the breakup of the Antarctic ozone hole has a major impact on STE,
- particularly that of O₃, and that its signal has large IAV that does not synchronize with the other
- 442 source of IAV, the QBO. Surprisingly, the large wintertime IAV in the NH Arctic, in the form
- 443 of sudden stratospheric warmings, does not seem to have a major role in STE fluxes as noted
- 444 above. This model may miss some of the Arctic O₃ depletion, but it accurately simulates the
- 445 warmings, which must have a small impact on STE because they do not disrupt the clear QBO
- 446 signal in the c-tracers.
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	Figure 4. QBO composites of the STE of O_3 (black lines; left y-axes), cN2OX (orange lines; orange right y-axes), and cF11X (blue lines; blue right y-axes) for the (a) NH (0°-90°N; solid lines) and (b) SH (0°-90°S; dashed lines). These composites are averages centered on the QBO phase transition at 40 hPa throughout the period of surface observations (years 2001-2016, excluding the 08/2010 observed anomaly, for a total of 6 QBOs). Note: the y-axes limits are different for each panel, but the interval scale is consistent for each tracer.	
448 449 450	34 . Surface variability of N ₂ O related to STE flux	
451 452 453 454 455 456 457	Surface variability of N ₂ O is driven by surface emissions, stratospheric loss, and a transport that mixes the first two signals. R2021 explored the variability originatin stratospheric chemistry using the decaying tracer N2OX-and. Here, we use surf-N the surface abundances of N2OX when corrected to steady state. R2021 showed t independent chemistry-transport models produced annual and QBO patterns in sur simply from stratospheric loss. In this paper we link surf-N2O to the STE cN2O f	ng only from 2O to denote hat three face N ₂ O

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459 460

The observed surface N2O (denoted obs-N2O and taken from the NOAA network (Dlugokencky

461 et al, 2019)) shows a slowly increasing abundance (-0.9 ppb/yr) with a clear signal of annual and 462 interannual variability at some latitudes (see R2021). We calculate annual and QBO-composite

463 obs N_2O after de-trending and restrict analysis in this section to model years 2001–2016 to be

464 consistent with the surface data. The latitude by month pattern of obs-N₂O includes the impact

465 of both stratospheric loss (~13.5 Tg/yr) and surface emissions (~17 TgN/yr), with the

466 preponderance of emissions being in the NH (Tian et al., 2020). Total emissions are not

467 expected to have large IAV but may have a seasonal cycle. The seasonal variation of surface

468 N2O can also be driven by seasonality in the interhemispheric mixing of the NH SH gradient (-1 469 ppb).

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linked above to the STE O₃ flux.

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472 3 The observed surface N₂O, denoted obs-N₂O and taken from the NOAA network (Dlugokencky 473 et al., 2019), shows a slowly increasing abundance (~0.9 ppb/yr) with a clear signal of annual 474 and interannual variability at some latitudes (see R2021). We calculate annual and QBO-475 composite obs-N₂O after de-trending and restrict analysis in this section to model years 2001-476 2016 to be consistent with the surface data. The latitude-by-month pattern of obs-N₂O includes 477 the impact of both stratospheric loss (\sim 13.5 Tg/yr) and surface emissions (\sim 17 TgN/yr), with the 478 preponderance of emissions being in the NH (Tian et al., 2020). Total emissions are not expected to have large IAV but may have a seasonal cycle. The seasonal variation of surface 479 480 N₂O can also be driven by seasonality in the interhemispheric mixing of the NH-SH gradient (~1 481 ppb). 482

484 <u>4</u>.1 Annual cycle

485 486 Figure 5 replots the hemispheric mean annual cycles of cN2O STE flux alongside the annual 487 cycles of surf-N2O and obs-N₂O. As noted above, the STE in each hemisphere is almost in opposite phase, as is the modeled surf-N2O (taken from Fig. 5 of R2021). The NH:SH 488 489 amplitude ratio is about 2.4:1 for both STE and surf-N2O. The lag from peak STE flux of cN2O 490 (negative N_2O) to minimum surf-N2O is about 3 months. Such a 90° phase shift is expected for 491 the seasonal variation of a long-lived tracer relative to a seasonal source or sink. The time lag 492 between the signal at the tropopause and at the surface, the tropospheric turnover time, should be 493 no more than a month. Surprisingly, the cN2O STE seasonal amplitude is much larger in the NH 494 $(\pm 3.4 \text{ Tg/yyr})$ than in the SH $(\pm 1.3 \text{ Tg/yyr})$, although the SH mean (6.5 Tg/yyr) is larger than the 495 NH (5.2 Tg/yr). Essentially, there is more variability of air depleted in N_2O entering the NH, but 496 air entering the SH has a larger overall deficit. Thus in our model, the stratosphere creates a NH-497 SH gradient of +0.3 ppb at the surface, which is a significant fraction of the observed N-S 498 difference of +1.3 ppb (R2021). This important result needs to be verified with other models or 499 analyses because it constrains the NH-SH location of sources.

500

501 In the NH, as noted in R2021, the two surface abundances, surf-N2O and obs-N₂O, have the 502 same amplitude and phase, implying that, if the model is correct, the emissions-driven surface

503 signal has no seasonality-, although we know that some important emissions are seasonal

504 (Butterbach-Bahl et al., 2013). In the SH, the surf-N2O signal is much smaller, in parallel with 505 the small seasonal amplitude in cN2O STE, but it is out of phase with the obs-N₂O. This result

the small seasonal amplitude in cN2O STE, but it is out of phase with the obs-N₂O. This result implies that the SH has some highly seasonal sources, or simply that the forcing of SH surf-N2O

507 by the seasonal cycle of cN2O is weak. Indeed, this is what we might expect from Figure 3: In

the NH the seasonal amplitude in N_2O overwhelms the IAV amplitude and is driving the obs-

509 N₂O; but in the SH, both amplitudes are comparable. Given the quasi-<u>regular</u> nature of the

510 QBO, it would interfere with the seasonal cycle and likely change its phase (as found for other

- 511 models in R2021).
- 512

513 In the NH, the annual cycle of O_3 and cN2O STE are clearly linked. If we accept that the obs-514 N₂O NH seasonal cycle is simply driven by the STE flux, then how will tropospheric O_3 respond

515 seasonally? A mole-fraction scaling of the STE fluxes gives an O₃:N₂O ratio of ~25, and thus

- 516 scaling the surf-N2O amplitude gives a large O₃ surface seasonality of ~18 ppb. However, the
- 517 residence time of a tropospheric O₃ perturbation is ~1 month, and thus the peak surface
- abundance will lag the peak STE flux by only about a month and not by 3 months as for N_2O .
- 519 O₃ will equilibrate with the flux on monthly timescales and not accumulate. Thus, our estimate
- 520 is that NH 30°-90° surface ozone might increase about 5 ppb, peaking in June, due to the STE
- flux. In the SH, seasonal patterns are weaker and not well defined, and thus no obvious STE O_3 signal is expected.
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Figure 5. The annual cycle of O_3 and cN2O STE (black and orange lines; left y-axes), and the surf-N2O and obs-N₂O (red and blue knotted lines; right y-axes) taken from R2021 (see their figure 5) for the **(a)** NH and **(b)** SH. cN2O, surf-N2O, and obs-N₂O has been rescaled to reflect that of a tropospheric abundance of 320 ppb. The hemispheric domains for STE is defined as 0°-90° while the surf-N2O and obs-N₂O is from 30°-90° N/S. Note: the left y-axes limits are different between the tracers, but the interval scale is the same.

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<u>34</u>.2. QBO cycle

529 The QBO composite of hemispheric mean cN2O STE flux from Figure 4 is compared with the 530 composite of surface abundances (surf-N2O and obs-N2O) in Figure 6. The peak in cN2O flux is 531 broad and flat, but centers on +2 months for the NH and +4 months for the SH. Unlike the 532 annual cycle, the QBO cycle in STE flux is almost in phase in both hemispheres, with the NH 533 preceding the SH. This phasing of the QBO cycle in surface N₂O was seen with the three 534 models in R2021. In both hemispheres, the modeled surf-N2O peaks before the rise in cN2O 535 and then decreases through most of the period with elevated cN2O flux as expected. The 536 amplitude of the QBO STE flux is smaller in the NH than SH by about half, and the amplitude of 537 surf-N2O is likewise smaller. The ratio of the amplitudes of surf-N2O to cN2O STE flux is 538 similar in both hemispheres (~ 0.4 ppb per Tg/yr), which is encouraging. This ratio is larger than 539 the corresponding one from the annual cycles (~ 0.1 ppb per Tg/yr) because the length of the 540 QBO cycle leads to longer accumulation of N2O-depleted air from the cN2O flux. 541

542 In the SH, where the QBO cycle in cN2O flux has a large amplitude, the modeled surf-N2O 543 matches obs-N₂O in amplitude and phase as reported in R2021. In the NH, the comparison of 544 surf-N2O with obs-N2O is not so good: obs-N2O has a much smaller amplitude and a different 545 phase. This OBO cycle pattern is similar, but reversed, to that of the annual cycle and can be 546 understood in the same way. The NH QBO cycle has relatively small amplitude and thus the 547 interference with the large-amplitude annual cycle adds noise, obscuring the QBO cycle. In the 548 SH it is the opposite, with its weak annual cycle, the SH QBO cycle is clear. The modeled cN2O 549 fluxes enable us to understand the large-scale variability of the observations.

550

551 Thus, for both annual and QBO fluctuations, when the variation in STE flux is dominated by 552 either cycle, the surface variations are clearly seen and modeled for that cycle. This further

- supports the findings in R2021 and other studies, that hemispheric surface N_2O variability is
 - 22



555 STE flux. Given the connection between O₃ and cN2O STE, this relational metric can be used to

556 constrain the O_3 STE for a model ensemble.





45. Lowermost stratosphere

561 4.1. The O₃:N₂O slopes and STE fluxes

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563 If we accept that matching the observed annual and QBO cycles in surface N₂O constrains the 564 modeled STE cN2O flux, then how can we use that to also constrain the modeled STE O₃ flux? 565 All evidence, theoretical, observational, and modeled, shows that the STE flux is simultaneous 566 for all species (e.g., Figure S1) and in proportion to their relative abundances in the lowermost 567 stratosphere (Plumb & Ko, 1992). We can test this]) and in proportion to their relative 568 abundances (i.e., tracer:tracer slopes) in the lowermost stratosphere, defined roughly as the 569 region 100-200 hPa in each hemisphere outside the tropics (Plumb and Ko, 1992).

571 <u>5.1. The O₃:N₂O slopes and STE fluxes</u> 572

573 We can test the Plumb and Ko hypothesis in our model framework by comparing the relative

574 STE fluxes for O₃, cN2O and cF11 with the modeled tracer-tracer slopes in the lowermost

575 stratosphere. These slopes can then be tested using SCISAT-1 ACE-FTS (Scientific Satellite-1

576 Atmospheric Chemistry Experiment-Fourier Transform Spectrometer) measurements of O₃ and

577 N_2O in the lowermost stratosphere to establish the ratio of the two STE fluxes. 578

579 Figure 7ab shows the N₂O-O₃ slope in each hemisphere taken from the ACE climatology dataset 580 and the UCI CTM. The current ACE dataset (version 3.5) has been curated from measurements 581 made by ACE-FTS from February 2004 to February 2013 (Koo et al., 2017). The SCISAT orbit 582 results in irregular season-latitude coverage, and thus we average the lowermost stratosphere 583 data over a wide range of latitudes centered on the peak STE flux (20° 60° in both hemispheres). 584 For both ACE data and the CTM we keep to the lowermost stratosphere (200-100 hPa) and 585 average over the 4-month peak of STE flux, Feb May in the NH and Sep-Dec in the SH (see 586 Figure +1 The current ACE dataset (version 3.5) has been curated from measurements made by ACE-FTS from February 2004 to February 2013 (Koo et al., 2017). The SCISAT orbit results in 587 588 irregular season-latitude coverage, and thus we average the lowermost stratosphere data over a 589 wide range of latitudes centered on the peak STE flux $(20^{\circ}-60^{\circ})$ in both hemispheres). For both 590 ACE data and the CTM we keep to the lowermost stratosphere (200-100 hPa) and average over 591 the 4-month peak of STE flux, Feb-May in the NH and Sep-Dec in the SH (see Figure 1). 592 Extending into the upper tropical troposphere at 20° helps define the tropospheric end-point of 593 the slope (low O₃, high N₂O). 594 595 Based on the long-term mean STE fluxes in the model, we would expect an O₃:N₂O slope of

about -24 (ppb/ppb) in the NH and -17 in the SH. The slopes fitted to our modeled grid-cell values of O₃ and N2O in the lowermost stratosphere are <u>remarkably</u> similar<u>but smaller</u>: <u>21:</u> <u>23,2</u> (NH) and <u>-1517,5</u> (SH). The ACE data are more scattered but show similar<u>smaller</u> slopes of -19.4 (NH) and -15.3 (SH). Thus, the NH-SH asymmetry in O₃ versus N₂O STE fluxes is clearly reflected in the tracer-tracer slopes, both modeled and observed. <u>Hegglin and Shepherd</u> (2007) had already identified these NH:SH differences when comparing their model to the ACE-FTS observations (their Fig. 13cd), but implications for STE fluxes were not brought forward.

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604	In the modeled SH (Figure 7b), one can see strings of points that are samples along neighboring
605	cells and reflect a linear mixing line between two different end points, one of which has
606	experienced extensive O ₃ depletion (i.e., the Antarctic O ₃ hole). We know that there is some
607	chemical loss of O3 in the NH lowermost polar stratosphere during very cold winters (Isaksen et
608	al., 2012; Manney et al., 2011), We know that there is some chemical loss of O_3 in the NH
609	lowermost polar stratosphere during very cold winters (Manney et al., 2011; Isaksen et al.,
610	<u>2012)</u> , but it is not extensive enough to systematically affect the $O_3:N_2O$ slope over the mid-
611	latitude lowermost stratosphere in either the ACE observations or the CTM simulations.
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628	5.2. IAV of the Antarctic ozone hole and the SH STE O ₃ flux
629	



4.2. IAV of the Antarctic ozone hole and the SH STE O3 flux

634 The Antarctic ozone hole appears to be the source of the NH-SH asymmetry in the STE fluxes of 635 O3 versus N2O. TheIt is known that the massive chemical depletion of O3 inside the Antarctic 636 vortex between about 13 and 23 km altitude creates an air mass with lower O3:N2O ratios than 637 usually found in the mid-latitude lowermost stratosphere. -When the vortex breaks up, nominally 638 in late November, much of this O3-depleted air mixes with the rest of can mix along isentropes 639 into the mid-latitude lowermost stratosphere-and reduces, changing the O3:N2O ratios and 640 reducing the SH STE O₃ flux.

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642 We have additional information on the SH O₃ STE flux from the year-to-year variations in the 643 size of the ozone hole. The best measure of the scale of Antarctic ozone depletion is the October 644 mean ozone column (DU) averaged from the pole to 63°S equivalent latitude (see Figure 4.5 of 645 WMO (2018)). Fig. 4-5 of WMO, 2018). When we compare the CTM with the observations 646 (Figure 8), we find remarkable verisimilitude in the model: the *rmsroot-mean-squared* difference 647 is 9 DU out of a standard deviation of 29 DU and the correlation coefficient is 0.96. Thus, we 648 have confidence that we are simulating the correct IAV of the ozone hole. Next, we plot the 649 modeled O₃ STE flux (summed over the 12 months following the peak ozone hole, Nov-650 Oct)November-October) with the modeled October ozone column and find a fairly linear 651 relationship. If we estimate the STE O_3 flux before the O_3 hole, when the mean October O_3 652 column was about 307 DU, then our O3 flux isincreases to 209 Tg/yr (see Figure 78, red marker), 653 eliminating the hemispheric asymmetry in O₃ STE flux. 654 655 The annual deficit in SH STE O₃ flux brought on by the Antarctic ozone hole ranges from about 656 5 to 55 Tg/yr and with a central value of 30 Tg/yr or 14% of the total. Using the decadal trends 657 1965-2000 from Hegglin and Shepherd (2009), this deficit is 8%; and from Meul et al. (2018), 658 5%. Since both of these models calculate a much larger SH flux (~300 Tg/yr), we estimate their 659 absolute change in O3 flux to be 24 and 15 Tg/yr, respectively. Because the ozone hole 660 effectively removes a fixed, rather than proportional, amount of ozone that presumably is 661 mapped onto the STE flux the following year, we believe the absolute change is the best 662 measure. Thus the three models estimate the ozone hole causes a deficit in the SH O₃ STE flux 663 in the range of 15-30 Tg/yr. The UCI CTM's ability to match the observed IAV of the ozone 664 hole, and to match that linearly with the deficit in STE flux provides support for the upper end of 665 the range. Note that the difference in O₃:N₂O slopes between NH and SH in Figure 7 is about 5. 666 If we attribute that solely to the ozone hole and split the flux of N₂O-depleted air evenly between 667 hemispheres, then the ozone-hole-driven O_3 STE flux difference is about 55 Tg/yr, about twice 668 that derived from the variability in our model. This difference in estimated flux indicates that even without chlorine-driven ozone depletion, the O3:N2O slopes may be inherently different 669 simply because of the strong descent inside the wintertime Antarctic vortex. This can be readily 670 671 investigated with further model studies. 672

We looked for any relationship between ozone hole IAV and the STE fluxes of cN2O or cF11
and found mostly a scatter plot with no clear relationship. Given the analysis above, we expect
that much of the scatter is related to QBO cycles.

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timescales (Tang et al., 2021). These changes in O3 overhead can have a direct influence on O3

transport to the troposphere, but the link requires further analysis. Tang et al. (2021) showed the

685 UCI CTM is able to capture the observed annual cycle of stratospheric O₂-column (extracted 686 from total column using Ziemke method; Ziemke et al., 2019). QBO modulation of stratospheric 687 column O3 has not been fully investigated but its magnitude, like that of the annual cycle, is 688 comparable to the magnitude of O3-STE and is clearly somehowStratospheric column O3 (DU) 689 varies on annual and QBO timescales. These changes in O3 overhead can have a direct influence 690 on O₃ transport to the troposphere, but the link requires further analysis. Tang et al. (2021) 691 showed the UCI CTM is able to capture the observed annual cycle of stratospheric O₃ column as 692 extracted from total column using the Ziemke et al. (2019) method. QBO modulation of 693 stratospheric column O₃ has not been fully investigated since Tung and Yang (1994b). Yet, the 694 fluctuations in mass over the annual cycle are comparable to the corresponding variability in O₃ 695 <u>STE flux (1 DU = 10.9 Tg) and likely</u> connected (Figure 9).







Figure 9. Stratospheric O₃ column residuals taken from MLS **(a, c)** and UCI CTM **(b, d)** for their mean annual cycle **(a, b)** and mean QBO cycle **(c, d)** during years 2005-2017. Residuals are defined at each latitude with a mean of zero DU.

56. Conclusions

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701 This work examines how closely O₃ STE is linked to STE fluxes of other trace gases. By 702 including our complementary N₂O and F11 tracers, we can follow stratospheric loss of these 703 gases along with stratospheric O_3 across the tropopause. The magnitudes of the fluxes are 704 proportional to their abundances in the lower stratosphere as expected (Plumb & Ko, 1992), The 705 magnitudes of the fluxes are proportional to their abundances in the lower stratosphere as 706 expected (Plumb and Ko, 1992), and their variability is highly correlated with one another, 707 indicating that they are entering the troposphere simultaneously. Even during QBOs, which have 708 their own-the distinct QBO pattern of STE fluxes, we find that the link between is consistent 709 across O₃, N₂O and F11-STE, remains consistent. We further constrain the N₂O transport 710 pathway by linking STE of depleted-N₂O air with surface fluctuations of N₂O abundance. The 711 surface response in modeled N₂O matches well with the observed surface variability in the 712 **NHSH**, indicating that surface variability is driven largely by STE flux. 713 714 Consistency of STE O₃ flux. As summarized here, there are a number of model diagnostics and

715 observational constraints that provide a reality check on the consistency of the modeled O₃ STE 716 flux. In Table 1, we examine these for our model and also for the CMAM model (Hegglin and 717 Shepherd, 2007, 2009) because it is one of the few with enough published results. For UCI we 718 calculate NH:SH fluxes of O₃ (208:182 Tg-O₃/yr) and N₂O (5.1:6.4 Tg-N/yr). Thus the mole 719 fraction slopes in the lowermost stratosphere should be -23.8 (NH) and -16.6 (SH). Our model 720 O₃:N₂O slopes are -23.2 (NH) and -17.5 (SH). Given the seasonal variability and scatter in the 721 correlation plots (Figure 7), we count this as consistent. For CMAM, the modeled O₃:N₂O 722 slopes, -21 (NH) and -18 (SH) are similar to ours and to the ACE-FTS observations as analyzed 723 by Hegglin and Shepherd (2007), -22 (NH) and -15 (SH), or by us, -19 (NH) and -15 (SH). 724 CMAM does not report the STE N₂O fluxes, but their model seems to match observations and so 725 we assume that the MLS-derived N₂O fluxes are a close estimate (12.9 Tg-N/yr). Just using the 726 CMAM global numbers for O3 STE flux, we calculate the O3:N2O slope in the lowermost 727 stratosphere should average to -30. We conclude that their diagnosis of the STE O_3 flux, 655 728 Tg/yr, is inconsistent with the circulation that generated the $O_3:N_2O$ slopes and is 50% too large.

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729 We do not view this as a critical assessment of CMAM since it involves us combining 730 diagnostics from two separate publications, but it is an example of how we might expect future 731 studies of the STE O₃ flux to self-evaluate. 732 733 Uncertainty Quantification in STE O₃ flux. Deriving a best estimate and uncertainty from this 734 work involves expert judgment. Changes in meteorological data used by the UCI CTM (IFS 735 Cycles 29r1, 36r1, and 38r1, all at 60-layer 1.1° resolution, see Table 1) give a standard 736 deviation in STE of 13% (only 3 values). If we use observations to derive a value as in Murphy 737 and Fahey (1994), we must expand our dimensions to the uncertainty in the NH:SH split of N₂O 738 flux to calculate each hemisphere's O3 flux. The factors are: (1) total STE N2O flux is 12.9 Tg-739 N/yr from the MLS data and we assign a ±10% one-sigma uncertainty; (2) the NH:SH split of 740 the N2O flux is 44:56 in our current model, was not diagnosed for previous ones, and so we 741 assume a value of 50:50 that ranges from 40:60 to 60:40; (3) analysis of the ACE-FTS 742 observations (ours and Hegglin and Shepherd, 2007) gives O3:N2O slopes of about -21 (NH) and 743 -15 (SH) to which we assign a one-sigma uncertainty of ± 3 . Propagating these as root-mean 744 square errors, we find a $\pm 15\%$ uncertainty in the global value, 400 ± 60 Tg/yr. Uncertainty in the 745 hemispheric values is more difficult to assess, and from a range of model results shown in Table 746 1, we can only estimate that the NH:SH ratio is between 60:40 and 50:50, a range that bounds 747 our and CMAM results plus 2%. Note that this estimate is for current conditions with a regularly occurring Antarctic ozone hole. We believe the low 50:50 ratio is plausible because we have 748 749 shown that our large SH STE N₂O flux is consistent with the surface OBO variability in N₂O. 750 For pre-1980, and for when the ozone hole recovers later this century, we anticipate that the SH 751 O₃:N₂O slope will revert to -18 to -21, and the total STE O₃ flux to 430-460 Tg/yr. This 752 753 simplistic estimate is based on a fixed atmospheric circulation. 754 A major surprise from our model is that the STE flux of O₃ is predominantly NH biased 755 currently, only because of the Antarctic ozone hole. Prior to 1980, and after 2060, it would/will 756 be symmetric between the hemispheres. Our model calculates slightly greater STE fluxes for 757 trace gases like N_2O or F11 in the SH, which is counter to prevailing theory that the wave-driven 758 fluxes force relatively greater STE in the NH. This difference cannot be directly tested with 759 observations of trace gases, but a range of N₂O hemispheric observations are well modeled and 760 support this premise. More extensive work with multi-model ensembles that include both 761 chemical and dynamical diagnostics in the stratosphere would be needed to overturn the 762 established theory. Our work reemphasizes the importance of trace-gas correlations in the 763 lowermost stratosphere as a key observational metric for climate models that may be able to 764 constrain total STE fluxes. The tracer slopes may go beyond just relative STE fluxes because we 765 have other measurements from the upper stratosphere to the surface that constrain, for example, 766 the absolute flux of N₂O better than we first did using just the modeled lifetime (Murphy & Fahey, 1994; McLinden et al., 2000). 767 768

769 In Table $\frac{12}{2}$, we propose gather a set of observation-based model metrics that relate to STE fluxes 770 and will help the community build more robust models to better derive the STE flux of O₃. 771

Table 1. Metrics from Measurements or Constrained Values for CCMs related to Stratosphere-+ **Formatted Table** Troposphere ExchangeTable 1. Summary of key results for the STE flux of O3 and N2O presented here (bold), SH Global NH notes

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STE O flux (TaO /)	208	192	200	IES Cy29r1 yrs 1000 2017 (this paper Dyin at al. 2021)
<u>STE O₃ flux (TgO₃/yr)</u>	<u>208</u>	<u>182</u>	<u>390</u>	IFS Cy38r1, yrs 1990-2017 (this paper; Ruiz et al., 2021)
	<u>239</u>	<u>198</u>	<u>437</u>	IFS Cy36r1, yrs 2000-2007 (Hsu & Prather, 2014)
	<u>301</u>	<u>233</u>	<u>534</u>	IFS Cy29r1, yrs 2000-2006 (Hsu & Prather, 2014)
	<u>383</u>	272	<u>655</u>	CMAM, yrs 1995-2005 (Hegglin & Shepherd, 2009)
<u>STE N₂O flux (TgN/yr)</u>	5.1	<u>6.4</u>	11.5	yrs 1990-2017, scaled to 320 ppb
			<u>12.9</u>	using MLS lifetime of 119 yr and 320 ppb
LMS O3:N2O slope*	-23.2	-17.5		UCI model
	<u>-19.4</u>	-15.3		ACE-FTS observations
	-23.0	-17.5		CMAM model, Fig 13 of (Hegglin & Shepherd, 2007)
	-22.0	-15.0		ACE-FTS observations, ibid
			-20.0	(Murphy & Fahey, 1994)
			-22.0	(McLinden et al., 2000)
STE flux O3:N2O				
(mole/mole)	<u>-23.8</u>	-16.6		UCI model, calculated from entries above
				CMAM (Hegglin & Shepherd, 2009), using MLS N2O
			<u>-29.6</u>	lifetime
Best Estimate	<u>60%</u>	<u>40% to</u>	$400 \pm$	current Antarctic ozone hole conditions, see text
<u>STE O₃ flux (Tg/yr)</u>	<u>to</u>	<u>50%</u>	<u>60</u>	
	<u>50%</u>			

* LMS = lowermost stratosphere. For UCI model, months are selected for highest STE (FMAM in NH, SOND in SH, Fig. 1). For CMAM, annual averages from their Fig. 13cd. Where no reference is given, the source is this paper.

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Name	Metric	Measured values	Model requirements	Example figure	·K
N ₂ O loss	Annual and QBO	Monthly N ₂ O loss	Stratospheric chemistry	Fig. 4	
	cycles of global	calculated from MLS	for N ₂ O as tracer; a QBO	(P2015Prather et	No.
	mean stratospheric	profiles (2005-	cycle; monthly mean	al., 2015); Fig. 2	X
	N ₂ O loss	present)	diagnostics	(R2021Ruiz et al.,	\mathbb{X}
				2021);	
				Fig. 3 (R2022 this	
				paper)	
STE slopes	Matching O ₃ :N ₂ O	ACE FTS profiles	Stratospheric O ₃ and N ₂ O	Fig. 7 (R2022 this	
	slopes in lowermost	(2004-2013)	calculation, possibly also	paper)	1
	stratosphere		CFCs; monthly snapshots	· · ·	\sum
Strat O ₃	Annual and QBO	Monthly zonal mean	Stratospheric O ₃	Fig. 9 (R2022 this	
column	composite cycles of	stratospheric O ₃	chemistry; a QBO cycle;	paper)	
	stratospheric O ₃	column from Z2019	monthly mean		11
	column	analysisZiemke et	diagnostics; separate		N
		<u>al., 2019</u> (2005-	strat & trop O ₃ columns		
		present)			1

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N₂O loss at	Annual and QBO	NOAA surface N ₂ O	Stratospheric N ₂ O	Fig. 3 (R2021 Ruiz	1	Form
surface	composite cycles of	observations	chemistry; N2OX as a	et al., 2021); Fig. 5		Form
	surface N ₂ O solely from stratospheric		tracer; monthly mean diagnostics	(R2022 this paper)	\bigvee	Form
	loss				X	Form
		Constrained				Form
		(modeled) values				Forma
STE flux of O ₃	-	Monthly, latitude or	Run O3strat as a tracer;	Fig. 1 & 2		
	A	hemispheric	diagnose monthly flux	(R2022 this paper)		Forma
		resolved, net O ₃ flux	into troposphere, at			Forma
			tropopause or through		M)	Forma
			trop-loss of O3strat			Form
STE flux of	-	Monthly, latitude or	Run cN2O (cF11) as a	Fig. 1 & 2	A	
N ₂ O depleted	•	hemispheric	tracer; diagnose monthly	(R2022this paper);	V /	Forma
air (also CFC-		resolved, STE flux of	flux into troposphere			Forma
11)		N ₂ O (CFC-11)				Forma
SH O₃ hole	-	Change in SH O ₃ STE	IAV of ozone hole; daily	Fig 7 (R2022 this		Forma
and flux	_	flux with size of	total O₃ column (lat,	paper)	M I	
		ozone hole; observed	long); monthly SH O ₃ STE			Forma
		IAV of O₃ hole	flux			Forma
Notes: Constra	ained values are key, n	nodel-only, derived quan	tities that can be diagnosed	from CCMs or CTMs.	MIN	Forma
Reference sł	northand: P2015 =	Prather et al., 2015; I	R2021 = Ruiz et al., 2021	L ; R2022 = this		Forma
paper; Z2019	9 = Ziemke et al., 2 0) <u>19</u>				Forma
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Author Contributions:

DJR and MJP designed and carried out the study and prepared the manuscript for publication.

Competing interests:

The authors declare that they have no conflict of interest. 798

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