

1 Stratospheric ozone depletion inside the volcanic plume shortly after the 2 2022 Hunga Tonga eruption

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21 Abstract

22 **Abstract**
23 ~~Near-term in-~~plume ozone depletion was observed for about ten days by Microwave Limb
24 Sounder (Aura/MLS) right after the January 2022 Hunga Tonga-Hunga Ha'apai (HTHH)
25 eruption. This work analyzes the dynamic and chemical causes of this ozone depletion. The
26 results show that the large water injection (~150 Tg) from the HTHH eruption, with ~0.0013 Tg
27 injection of ClO (or ~0.0009 Tg of HCl), causes ozone loss due to strongly enhanced HOx and
28 ClOx cycles and their interactions. Aside from the gas phase chemistry, the heterogeneous
29 reaction rate for HOCl+HCl→Cl₂+H₂O increases to 10⁴ cm⁻³sec⁻¹ and is a major cause of
30 chlorine activation, making this event unique compared with the springtime polar ozone
31 depletion where HCl+ClONO₂ is more important. The large water injection causes relative
32 humidity over ice to increase to 70% - 100%, decreases the H₂SO₄/H₂O binary solution weight
33 percent to 35% compared with the 70% ambient value, and decreases the plume temperature by
34 2-6 K. These changes lead to high heterogeneous reaction rates. Plume lofting of ozone-poor air
35 is evident during the first two days after the eruption, but ozone concentrations quickly recover
36 because its chemical lifetime is short at 20 hPa. With such a large seawater injection, we expect
37 that ~5 Tg Cl was lifted into the stratosphere by the HTHH eruption in the form of NaCl, but
38 only ~0.02% of that remained as active chlorine in the stratosphere. ~~Lightning~~ NOx changes are
39 probably not the reason for the HTHH initial in-plume O₃ loss.

40 Key points:

- 41 • HOCl is identified as playing a large role in the in-plume chlorine balance and
42 heterogeneous processes, making this event unique compared with the ozone hole where
43 HCl+ClONO₂ is more important.
- 44 • The HTHH eruption enhanced the HOx/ClOx cycles and their interactions, which caused
45 in-plume O₃ depletion.
- 46

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- The injection of Cl, H₂O, and lightning NO_x modified the ambient chemistry.

1. Introduction

Stratospheric ozone concentrations change after volcanic eruptions for a variety of reasons. Enhanced polar ozone depletion occurs after large or medium volcanic eruptions [Hofmann and Oltmans, 1993; Portmann *et al.*, 1996; Solomon *et al.*, 2016] since heterogeneous reactions on volcanically enhanced sulfate aerosols result in amplified anthropogenic ClO_x and BrO_x induced ozone loss. Tie and Brasseur [1995] demonstrated that mid- and high latitude O₃ changes after a volcanic eruption largely depend on chlorine loading. For the pre-industrial era and in the absence of anthropogenic halogens in the stratosphere, O₃ would slightly increase in the middle atmosphere after a large volcanic eruption resulting from the suppression of NO_x-catalyzed destruction by heterogenous creation of HNO₃ on volcanic aerosols. After the 1991 Pinatubo eruption, the radiative heating caused by volcanic aerosols perturbed the local temperature and circulation, which lifted the ozone layer and caused equatorial ozone depletion [Kinnison *et al.*, 1994]. Wang *et al.* [2022] reported that, in the case of the Hunga-Tonga eruption, mid-latitude ozone reduction was primarily caused by anomalous upwelling. Enhanced water can also change O₃. In the lower most stratosphere, H₂O injection through deep convection or tropopause cirrus clouds could change the catalytic chlorine/bromine free-radical chemistry and shift the total available inorganic chlorine towards the catalytically active free-radical form, ClO [Solomon *et al.*, 1997; Anderson *et al.*, 2012].

Evan *et al.* [2023] report observations of decreased O₃ and HCl, and increased ClO in the first week following the HTHH eruption at 20 hPa, which is related to the injected H₂O exceeding the normal range of the stratospheric variability. Here we use the Whole Atmosphere Community Climate Model version 6 (WACCM6) model [Zhu *et al.*, 2022] to analyze the dynamic and chemical contributors to this initial in-plume ozone depletion, and to understand the climate model performance. A lofting plume can bring ozone-poor tropospheric air into the stratosphere and cause in-plume low ozone values compared with the surrounding stratospheric air [Yu *et al.*, 2019]. For a submarine volcanic eruption, the in-plume air composition is not only impacted by tropospheric air, but also by the seawater, and volcanic gases (including H₂O, CO₂, SO₂, HCl, HF, H₂S, S₂, H₂, CO, and SiF₄), and volcanic minerals. For the HTHH initial plume, besides high H₂O and high SO₂, Microwave Limb Sounder (MLS) observations indicate the in-plume air carried high CO (Figure A1), relatively low ozone, and high ClO, compared with the surrounding air. We constrain the initial plume chemical compounds based on observational data from MLS; then analyze how stratospheric chemistry changes the plume composition. We will answer the following scientific questions:

1. What are the initial conditions in the volcanic plume?
2. What are the main causes of in-plume ozone depletion?
3. How do volcanic injections impact heterogeneous reactions that cause chlorine activation in the plume?

2. Observational data description and model setup

The MLS instrument onboard the EOS Aura satellite was launched into a near-polar sun-synchronous orbit in 2004. This work uses MLS version 4 for O₃, ClO, temperature, and CO data during the first ten days after the eruption as recommended by Millán *et al.* [2022]. The vertical resolution of these MLS products is typically around 3-5 km in the stratosphere. All data used here were screened using the methodology indicated in Livesey *et al.* [2022]. We use the MLS

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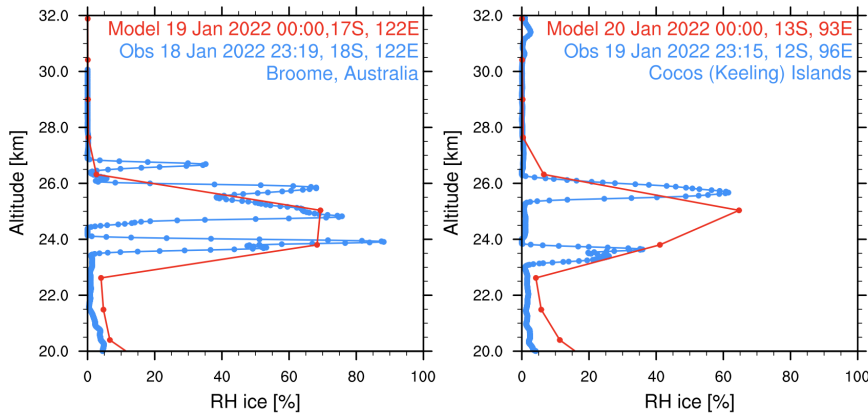
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99 H₂O data to identify the plume location and define it as regions with water vapor larger than 10
100 ppmv.

101 *Vömel et al.* [2022] provide water vapor radiosonde measurements during the first three
102 global circumnavigations of the plume. Here we calculate the relative humidity relative to ice
103 (RH_i) and compare the observed values with the simulated values.

104 We use the 70-layer WACCM model as described in *Zhu et al.* [2022], injecting SO₂
105 (0.42 Tg) and H₂O (150 Tg). The model has a horizontal resolution of 0.9° latitude × 1.25°
106 longitude. The injection plume in the model includes about 40 grid points. The model's vertical
107 resolution is about 1 km in the stratosphere. The model atmosphere is nudged to GEOS5
108 MERRA meteorological analysis [*Rienecker et al.*, 2008] until January 14, one day before the
109 eruption day. After January 15, we run the model freely with a fully interactive atmosphere and
110 ocean for ten days.

111 We constrain the simulated volcanic aerosol, H₂O, and chlorine by comparing to
112 observations during the first ten days after the eruption. *Zhu et al.* [2022] show that the simulated
113 aerosol backscatter coefficient agrees with the CALIPSO observations on January 17. The
114 simulated H₂O agrees with MLS [*Millán et al.*, 2022; *Zhu et al.*, 2022] from February 1 to April
115 1, 2022. Here, we compare the simulated H₂O with the radiosonde observations of humidity
116 [*Vömel et al.*, 2022] during the first week. **Figure 1** shows the RH_i on January 18 and January 19
117 observed by the radiosonde and from nearby simulated model output. Both the observations and
118 simulations show relative humidity between 70% to 100%. The radiosonde observations have a
119 much higher vertical resolution than the model. Therefore, they show multiple layers of water
120 enhancement, while the model only shows one.



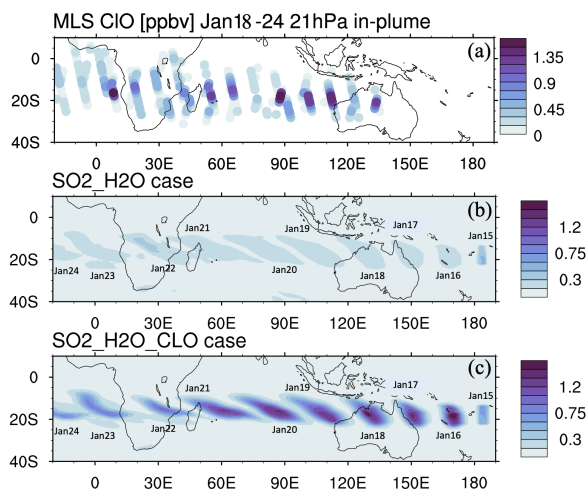
121 **Figure 1.** Relative humidity with respect to ice saturation vapor pressure from radiosondes (blue)
122 [*Vömel et al.*, 2022] and simulation (red). The profiles are picked at nearby locations. Note the
123 observations are about 45 minutes earlier in time than the simulations, which places them on a
124 different day.
125

126
127 We constrain the chlorine injection using MLS ClO observations at 20 hPa. **Figure 2a**
128 shows ClO from the MLS observations and the model simulations at 20 hPa from January 18 to
129 January 24. MLS values are selected from locations where water vapor is larger than 10 ppmv,
130 indicating these values are inside the volcanic plume. **Figures 2b** and **2c** show the simulated

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133 daytime ClO for one plume location for each day. The dates are marked next to each plume.
 134 MLS observations show elevated ClO, about 5 to 10 times higher than the ambient values
 135 (**Figure 2a**). If we only inject SO₂ and H₂O (The H₂O_SO₂ case defined in Table 1), we get a
 136 ClO amount about twice as large as the background (**Figure 2b**), which is much lower than
 137 observed. The change of ClO indicates that H₂O alters the Cl₂ partitioning. To match the
 138 observed values, we need to inject 0.0013 Tg of ClO (**Figure 2c**). This is equivalent to injecting
 139 ~0.0009 Tg of HCl (**Figure A2**). In our simulations, injecting ClO and HCl does not lead to
 140 different HOCl (**Figure A3**), ClO, and O₃ levels after January 15, indicating the balancing of
 141 ClO and HCl inside the HTHH plume happens very quickly. Unfortunately, the HOCl retrieval
 142 from MLS is not suitable for scientific use at this pressure level, so we cannot validate it. We
 143 choose the ClO injection case in our following analysis. Note that the MLS ClO vertical
 144 resolution is ~2 km near 20 hPa, which is coarser than the model vertical resolution (~1 km at 20
 145 hPa).
 146



147
 148 **Figure 2.** **a)** MLS in-plume ClO observations from January 18 - 24. “In-plume” is defined as the
 149 area with water vapor mixing ratios larger than 10 ppmv. MLS in-plume ClO data is not
 150 recommended for scientific use until January 18, 2022. **b)** and **c)** Simulated 10-day evolution of
 151 in-plume ClO in the SO₂_H₂O and SO₂_H₂O_CLO case. The modeled ClO concentrations are
 152 only taken during daytime each day (either 6 UTC or 12 UTC).
 153

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154 To investigate the O₃ decrease and its related chemical evolution during the first 10 days,
 155 we conduct several simulations as described in **Table 1**.
 156

157 **Table 1.** Model cases description.

Name	Description
Nonvolc	No injection of volcanic H ₂ O and SO ₂ .
H2O_SO2	H ₂ O and SO ₂ injection profile follows Zhu et al. [2022].

H2O_SO2_CIO	Besides H ₂ O and SO ₂ , injection of 0.00013 Tg of ClO. ClO injection profile is proportional to H ₂ O injection.
H2O_SO2_CIO_nohet	Same setting as H2O_SO2_CIO, but turn off the heterogeneous chemical reactions for HCl+HOCl, ClONO ₂ +H ₂ O, and ClONO ₂ +HCl
SO2_CIO	SO ₂ injection profile follows Zhu et al. (2022). No water injected. Injection of 0.00013 Tg of ClO using the same profile as H2O_SO2_CIO.
lowO3	Reduce the O ₃ to 75% of its original value at 20 hPa.
H2O_SO2_lowO3	H ₂ O and SO ₂ injection, plus reducing O ₃ to 75%.
H2O_SO2_CIO_lowO3	H ₂ O, SO ₂ and ClO injection, plus reducing O ₃ to 75%.
H2O_SO2_NO	Injection of 0.003 Tg of NO in addition to H ₂ O and SO ₂ .

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160

161 3. Results

162 *Evan et al.* [2023] show the HTHH in-plume ozone depletion at 20 hPa lasts at least ten
163 days after the HTHH eruption, which they attribute to the heterogeneous chlorine activation on
164 humidified volcanic aerosols. Here we analyze the contributions to this initial in-plume O₃
165 depletion considering three processes: 1) increasing H₂O injection may enhance the HOx
166 catalytic cycle and HOx/ClOx interactions; 2) increasing ClO during the injection phase may
167 deplete ozone due to both heterogeneous reactions and gas phase reactions; 3) the rising plume
168 from the troposphere may carry ozone-poor tropospheric air into the stratosphere.

169 MLS observed in-plume low ozone concentration at 20 hPa (Figure 3a), especially
170 during these three days: ozone concentrations of 4.8 ppmv on January 17, 4.6 ppmv on January
171 20, and 5.1 ppmv on January 24. These are ozone anomalies of about 1.7 ppmv, 1.9 ppmv, and
172 1.4 ppmv, respectively. The anomalies are calculated using the background average values in this
173 area (6.5 ppmv) subtracting the low ozone values. Note that any interpretation of these O₃
174 anomalies needs to consider the coarse MLS vertical resolution (~3 km). Because the plume is
175 spatially small during the initial days, MLS tracks do not capture the maximum plume
176 perturbation every day. The simulation with the water injection (Figure 3b) accelerates the HOx
177 catalytic cycle and shows evident O₃ reduction, but less than observed. Once we inject ClO on
178 top of the massive water injection (Figure 3c), O₃ loss is significantly enhanced and is close to
179 the observations after January 18. The difference between Figure 3d and Figure 3c is caused by
180 heterogeneous reactions, which usually only happen in the stratospheric polar springtime where
181 they cause the Antarctic ozone hole and Arctic ozone depletion. Heterogeneous reactions
182 become important, despite the high non-polar temperatures because of the massive quantity of
183 water injected. The heterogeneous reaction rate is strongly related to the relative humidity [Shi et
184 al., 2001]. Usually, during the polar night, the relative humidity is higher (RH_i 60%-100%) than
185 in the non-polar stratosphere because of the low temperature (<195 K). Here, the water injection
186 increases the relative humidity (Figure 4c). Enhanced water causes the weight percent of H₂SO₄
187 of the sulfuric acid aerosol to decrease from 70% to 35% (Figure 4b). The massive water
188 injection also causes the in-plume temperature to drop about 2 to 6 K (Figure 4f) [Solomon et
189 al., 2016]. All these factors (temperature decrease, relative humidity increase, and particle H₂SO₄
190 dilution) can increase the three heterogeneous reaction probabilities (HCl+HOCl, ClONO₂+H₂O,
191 and ClONO₂+HCl). As shown in Figure 5, when the water vapor amount is near the

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Deleted: Figure 3b shows the simulated O₃ in the H2O_SO2 case using one model time step each day that occurs near local noon. Figure 3b

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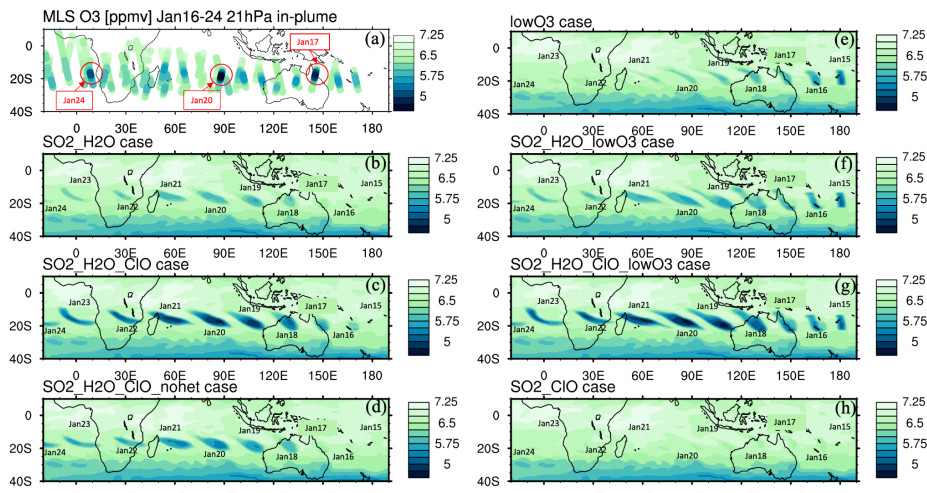
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206 climatological value of 6 ppmv, the heterogeneous reaction probability reaches 10^{-2} to 10^{-1} when
 207 the temperature is ~ 190 K. Meanwhile, the reaction probability is similar for temperatures of 215
 208 K when the water vapor is ~ 600 ppmv in the simulations, as was the case for the HTHH plume
 209 during the week following the eruption. COSMIC-2 radio occultation observed even higher
 210 water vapor during the first week: the maximum values over January 20-22 are ~ 1000 -2000
 211 ppmv [Randel et al., 2023]. Also, because the in-plume and the out-of-plume chemical
 212 concentrations are different, we apply both conditions (solid and dashed lines) to show how the
 213 different HCl, HOCl, and ClONO₂ conditions alter the HCl+HOCl and ClONO₂+HCl reactions
 214 probabilities by one order of magnitude. Volcanic sulfur injection also increases the sulfate
 215 surface area density (**Figure 4a**) that provides extra surfaces for heterogeneous reactions.

216 Comparing **Figure 3b** and **3c** with MLS observations, we can see that the chemical
 217 reactions do not explain the O₃ loss during the first three days of the eruption (January 15 -
 218 January 17, low O₃ near 160°E in MLS observation). This discrepancy suggests that the plume
 219 contains some ozone-poor tropospheric air after the injection into the stratosphere. We ran three
 220 cases with initial low ozone. For the low O₃ case (**Figure 3e**), we inject only ozone-poor air
 221 without volcanic H₂O and SO₂. It shows low O₃ as observed during the first couple of days, but
 222 ozone recovers quickly because the O₃ chemical lifetime is short at 20 hPa inside the plume
 223 (**Figure A4**). The H₂O_SO₂_lowO₃ case (**Figure 3f**) shows ozone loss similar to the
 224 observation in the first six or seven days. By adding the ClO and initial ozone-poor air (**Figure**
 225 **3g**), we obtain persistent low O₃ values that agree with the observational lowest values better
 226 than the other cases (**Figure 6a**). Compared with **Figure 3b**, **Figure 3d** has slightly more ozone
 227 depletion, indicating that the extra chlorine injection impacts O₃ even without heterogeneous
 228 chemistry. However, without including the high amounts of injected water, the additional ClO
 229 alone cannot deplete ozone much (**Figure 3h**).
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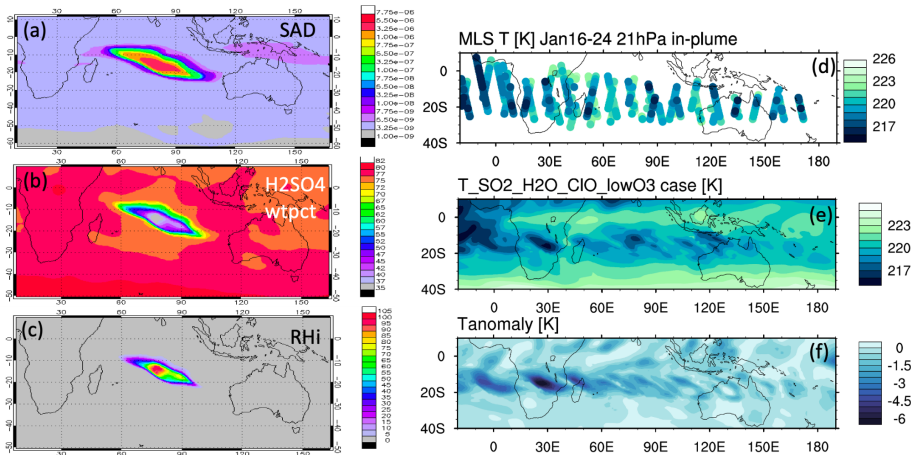
231 **Figure 3.** a) MLS in-plume O₃ observation from January 16 - 24. “In-plume” is defined as in
 232 **Figure 2**. Note that MLS ozone retrievals were unaffected by the plume leading to the addition of
 233 two extra days of data for this figure. The locations and days with low O₃ values used in **Figure**
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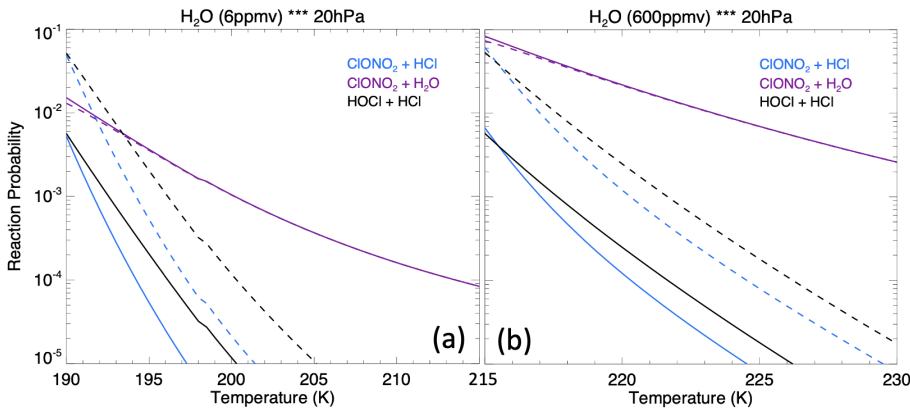
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237 **6** are marked with circles. **b-h)** Simulated 10-day evolution of in-plume O₃ in seven model cases
 238 with various injections of SO₂, H₂O, ClO, and low initial O₃. **Figure 3d** uses the same injection
 239 as **Figure 3c** but with heterogeneous reactions (i.e., HCl+HOCl, ClONO₂+H₂O, and
 240 ClONO₂+HCl) turned off. The simulated O₃ in the H₂O_SO₂ case uses one model time step
 241 each day that occurs near local noon.
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243 **Figure 4.** **a)** Simulated surface area density, **b)** simulated H₂SO₄/H₂O weight percent and **c)**
 244 relative humidity on January 20 at 20 hPa. **d)** Temperature evolution during the first ten days at
 245 20 hPa from MLS, **e)** simulated temperature evolution in the SO₂_H₂O_ClO_lowO₃ case; **f)**
 246 temperature difference between the SO₂_H₂O_ClO_lowO₃ case and the Nonvolc case.
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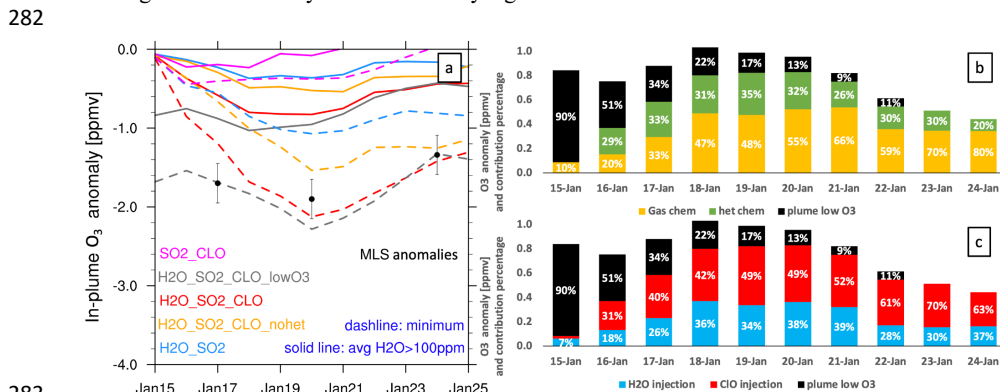
249 **Figure 5.** The heterogeneous reaction probability for three reactions on sulfate surfaces
 250 (ClONO₂+HCl, ClONO₂+H₂O and HOCl+HCl) as a function of water vapor assuming 0.4 μm
 251 particle size at 20 hPa. Panel **a)** assumes 6 ppmv of ambient water vapor and panel **b)** assumes
 252

253 600 ppmv of ambient water vapor. The solid lines use the out-of-plume chemical concentration
 254 on January 20: 1.0 ppbv of HCl, 0.03 ppbv of HOCl, and 0.5 ppbv of ClONO₂; the dashed lines
 255 use the in-plume chemical concentration: 0.1 ppbv of HCl, 1.0 ppbv of HOCl, and 0.05 ppbv of
 256 ClONO₂. These values are based on the simulation output.

257
 258 **Figure 6** shows the O₃ anomaly evolution from several model cases (a) and percentage
 259 contributions to the total ozone loss (b, c). The model case with all injections (initial low O₃,
 260 high H₂O, and high ClO) agrees well with MLS observations on the three days with the lowest
 261 O₃ values (**Figure 6a**). In **Figure 6b** and **6c**, the black bars represent the contribution from the
 262 low O₃ injection, which is significant during the first couple of days but diminishes quickly.
 263 From these percentage values, we conclude that the low O₃ carried in the plume lofting cannot be
 264 the reason for the low O₃ values after 3 days. Chemistry is the main reason that this O₃ depletion
 265 lasts so long.

266 There are two ways to look at the chemical contributors to ozone loss based on our model
 267 runs. The first is to separate the contributors due to various injections (**Figure 6c**): H₂O injection
 268 accounts for about 30-40% of the ozone loss most of the time (blue) and ClO injection accounts
 269 for 50% of the ozone loss most of the time (red). However, we cannot simply attribute the largest
 270 contribution to the ClO injection, because if we only inject ClO, it does not produce much ozone
 271 depletion (**Figure 6a**, magenta). It is the ClOx/HOx interactions that accelerate O₃ depletion.

272 A second way to look at the causes for ozone loss is to separate the contributions from
 273 the gas-phase chemistry and the heterogeneous chemistry (**Figure 6b**). The model run with the
 274 H₂O and ClO injections, but without the heterogeneous chemistry shows that the gas-phase
 275 chemistry (yellow bars) account for more than 47% of the ozone loss from January 18 - 24.
 276 Heterogeneous chemistry (green bars) destroys about 30% of the ozone. Hence, both
 277 heterogeneous chemistry and gas-phase chemistry are important for O₃ depletion. Once we turn
 278 off the heterogeneous chemistry, the partitioning between active chlorine and chlorine in the
 279 reservoirs is changed. The order in which the processes are accounted for can affect the resulting
 280 breakdown. Thus, we cannot simply say that gas phase chemistry contributions are larger than
 281 heterogeneous chemistry. Both are clearly significant.



283 **Figure 6. a)** O₃ anomaly in different model cases. The solid lines are the average O₃ anomaly at
 284 20 hPa on each day near local noon where water vapor is larger than 100 ppmv. 100 ppmv here
 285 is suggested by *Evan et al.* [2023], who found that O₃ anomalies are not significant for a 10
 286

287 ppmv but significant for a 100 ppmv threshold. The dashed lines are the simulated maximum O₃
 288 anomaly on each day at 20 hPa. The black dots show the three days during which MLS measures
 289 the lowest O₃ values (explained in **Figure 3a, b**) The percentage contributions to ozone loss
 290 from gas phase chemistry (orange) (H₂O_SO₂_CLO_nohet), heterogeneous chemistry (green,
 291 H₂O_SO₂_CLO minus H₂O_SO₂_CLO_nohet), and low O₃ air carried into the stratosphere
 292 (black, H₂O_SO₂_CLO_lowO₃ minus H₂O_SO₂_CLO). **c**) The percentage contributions to
 293 ozone loss from H₂O injection (blue, H₂O_SO₂ minus Nonvolc), ClO injection (red,
 294 H₂O_SO₂_CLO minus H₂O_SO₂), and low O₃ air carried into the stratosphere (black,
 295 H₂O_SO₂_CLO_lowO₃ minus H₂O_SO₂_CLO).

296
 297 To better understand which reactions are critical in the HTHH plume, we investigate the
 298 simulated reaction rates related to HOx and chlorine compounds (**Figure 7**). These reactions
 299 reflect how the water and ClO injections strengthen the in-plume HOx/ClOx interactions,
 300 chlorine activation, and the relative importance of each heterogeneous reaction rate. The
 301 WACCM model uses the methods developed by *Shi et al.* [2001] for heterogeneous reaction rate
 302 calculations. **Figure 7a** shows the HOx cycle inside and outside the water plume on January 20,
 303 daytime, at 20 hPa. The HO₂+O₃ reaction rate increases **by a factor of four** (from 5x10⁴ to 2x10⁵
 304 cm⁻³sec⁻¹); OH+O increases **by a factor of ~four** (from 2x10⁴ to 7.5x10⁴ cm⁻³sec⁻¹); HO₂+O
 305 increases **by a factor of four** (from 2x10⁴ to 8x10⁴ cm⁻³sec⁻¹). In addition, the extra HOx plays a
 306 large role in chlorine activation. **Figure 7b** shows the chlorine compound reactions inside the
 307 HTHH initial plume. The HOCl photolysis rate increases **by a factor of ~30 inside the plume**
 308 (from 6x10³ cm⁻³sec⁻¹ outside the plume to 2x10⁵ cm⁻³sec⁻¹) **due to the high HOCl mixing ratio,**
 309 which is the dominant process causing the increase in chlorine activation to Cl. The HOCl
 310 concentration remains high due to the enhanced ClOx/HOx interaction (i.e.,
 311 ClO+HO₂→HOCl+O₂ reaction), as well as the increase of the heterogeneous reaction rate of
 312 ClONO₂+H₂O **by five orders of magnitude** (from 0.3 to 1x10⁴ cm⁻³sec⁻¹). The large amounts of
 313 HOCl also make the heterogeneous reaction of HOCl+HCl faster than the ClONO₂+HCl
 314 reaction, while the latter reaction is known as the major reaction contributing to the chlorine
 315 activation that contributes to the polar ozone depletion. **Figure A5** shows the uptake coefficient
 316 for the three heterogeneous reactions HCl+HOCl, ClONO₂+H₂O, and ClONO₂+HCl on January
 317 20. The reaction **probability** of ClONO₂+HCl is increased **by eight orders of magnitude** (from the
 318 background value of 10⁻¹⁰ to 10⁻²). This value is even higher than *Evan et al.* [2023] suggested,
 319 who estimate that enhanced water increases the uptake coefficient of ClONO₂+HCl to 10⁻⁴ cm⁻³
 320 sec⁻¹. The reaction probability of HCl+HOCl and ClONO₂+H₂O increases to 10⁻². Furthermore,
 321 inside the plume, the reactions that convert Cl back to HCl are slower than their activation rate.

322 **Besides the ozone loss reactions, ozone production reactions are also significantly altered**
 323 by the water plume (**Figure 7c**). HO₂+NO is usually not an important process for O₃ production
 324 in the stratosphere (more important in the troposphere). The reaction rate **doubles inside the**
 325 **plume** (from 1x10⁵ cm⁻³sec⁻¹ to 2x10⁵ cm⁻³sec⁻¹). **Note that we don't inject lightning NOx in this**
 326 **case, a possible scenario during the eruption phase, that can also further increase the O₃**
 327 **production (detailed in the discussion section).**

328 **Comparing the partitioning of Cly**
 329 (Cl+ClO+2Cl₂+2Cl₂O₂+OCIO+HOCl+ClONO₂+HCl+BrCl) **reveals the in-plume chlorine**
 330 **activation processes (Figure 8)**. Outside the plume, HCl and ClONO₂ are dominant, indicating
 331 that most of the Cl is in reservoirs. While inside the water plume, both the H₂O_SO₂ and
 332 H₂O_SO₂_CLO cases show strong depletion of the reservoirs HCl and ClONO₂, and most of the

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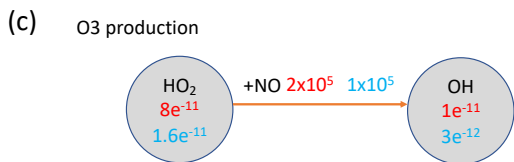
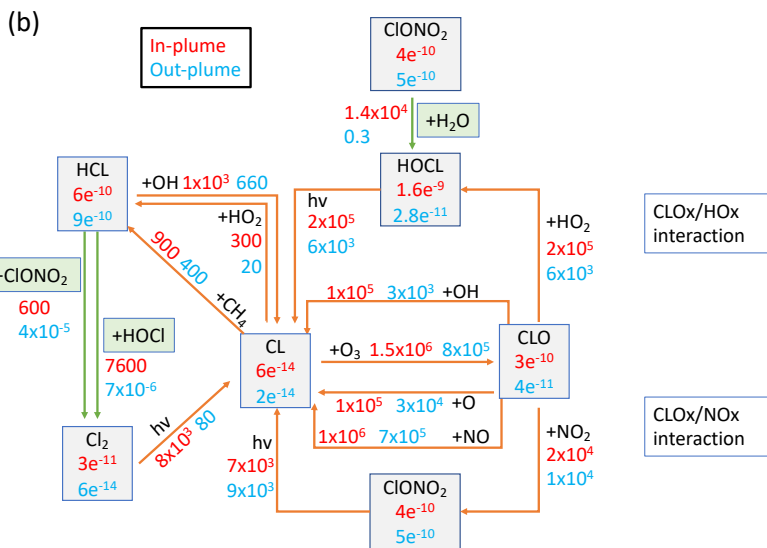
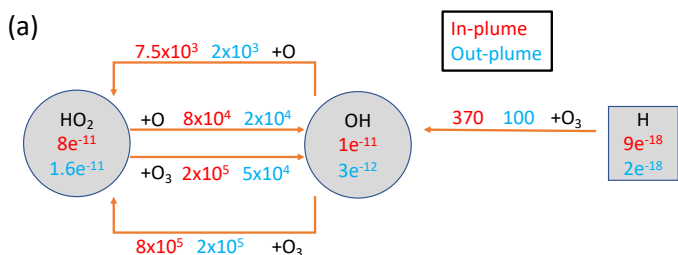
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355 Cly is either in the form of HOCl (a short-lived reservoir) or is activated in the form of ClO.
 356 Unlike the chlorine activation process in the polar winter, HOCl is the highest in the HTHH
 357 plume because heterogeneous chemistry is not fast enough to destroy HOCl to produce ClO. In
 358 the case without heterogeneous chemistry, HCl and ClONO₂ are dominant in the plume,
 359 indicating that heterogeneous chemistry is the main process of converting HCl to active chlorine.
 360 Comparing total Cly and ClO in all panels, ClO does not exceed a quarter of the Cly, indicating
 361 adding 0.00013Tg of ClO through injection is one way to produce the observed ClO. There is a
 362 possibility that ClO is converted from other Cly species through chemical reactions we are not
 363 aware of because this was a very unusual eruption.

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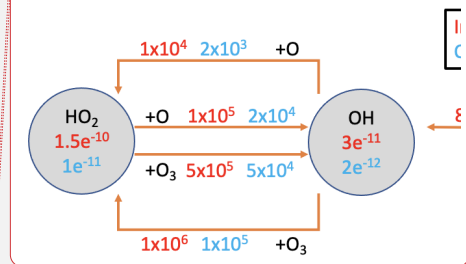


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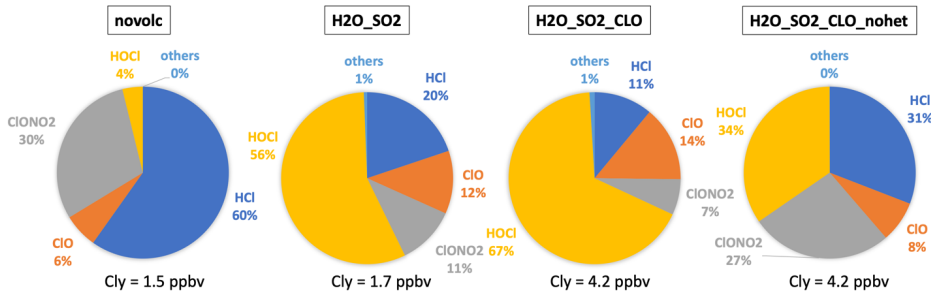
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370 **Figure 7.** Reactions inside and outside the plume in $\text{cm}^{-3}\text{sec}^{-1}$ and compound concentrations in
 371 mol/mol. Red numbers represent values inside the plume, blue numbers outside the plume. **a)**
 372 HOx balance and its interaction with Ox during daytime at 20 hPa on January 20, 2022. **b)**
 373 Chlorine compound reactions in the H2O_SO2_CIO case. **c)** HOx cycle impact on O₃
 374 production. Green arrows represent the heterogeneous reactions for chlorine activation. H₂O is ~
 375 600 ppm inside the plume and ~5.5 ppm outside the plume. Cly is ~4.2 ppbv inside the plume
 376 and 1.5 ppbv outside the plume.

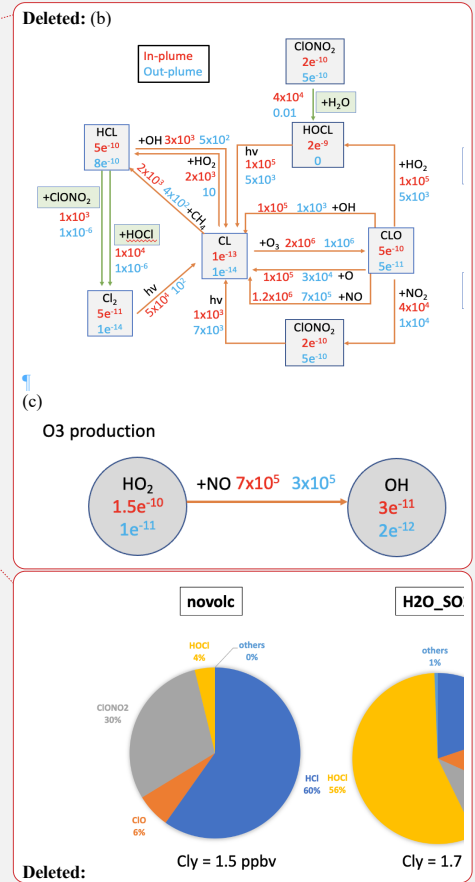


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 379 **Figure 8.** The percentage of each inorganic chlorine compound
 380 ($\text{Cly} = \text{Cl} + \text{ClO} + 2\text{Cl}_2 + 2\text{Cl}_2\text{O}_2 + \text{OCIO} + \text{HOCl} + \text{ClONO}_2 + \text{HCl} + \text{BrCl}$) inside and outside the plume.
 381 The slight difference between novolc Cly and H₂O_SO2 Cly is because H₂O injection changes
 382 the plume dynamics in the free-running simulations.

384 4. Discussion

385 The ozone loss inside the HTHH plume during the first ten days provides a unique
 386 opportunity to study stratospheric chemistry and to understand the performance of the WACCM
 387 state-of-the-art climate model, because the HTHH injected ClO and H₂O exceed the normal
 388 range of the stratospheric variability. These volcanic injections strongly altered the ClOx/HOx
 389 interactions and heterogeneous reaction rates, producing different chemical pathways for
 390 chlorine activation and ozone depletion from what occurs in the Antarctic ozone hole or Arctic
 391 ozone depletion in the polar stratospheric winter and spring. HOCl is identified as playing a large
 392 role in the in-plume chlorine balance and heterogeneous processes. The high HOCl
 393 concentrations are a result of the very high in-plume water vapor content, which makes this event
 394 different from chemistry in the Antarctic ozone hole, where ClONO₂ is more important.

395 This study also raises an interesting question of where the Cl comes from in the volcanic
 396 injection. Seawater contains 3.5% sea salt, which implies that about 5 Tg of NaCl could have
 397 been injected assuming that the injected 150 Tg of H₂O came from sea water. However, we only
 398 need to inject 0.00013 Tg of ClO to match the MLS ClO observations during the first few days
 399 after the eruption. We also conducted a test injecting an equivalent amount of HCl (0.0009 Tg),
 400 which resulted in a similar HOCl, ClO, and O₃ pattern (**Figure A2 and A3**). If we inject more
 401 HCl or ClO, ClO would exceed the observed concentration, causing depletion of OH, and
 402 slowing down the SO₂ oxidation. Evidently, if the water came from seawater, most NaCl was not
 403 converted to HCl but stayed in the stratosphere as particles. Vernier *et al.* [2023] sampled NaCl
 404 particles eight months after the eruption near Brazil. Based on their sampled NaCl concentration,
 405 we estimate 0.5 to 1 Tg of NaCl may have been injected and stayed in the atmosphere. There are



412 several possibilities why this event did not inject 5 Tg of NaCl in the stratosphere: Remote
413 sensing particle size estimations [Khaykin *et al.*, 2022] and in-situ measurements [Asher *et al.*,
414 2023] indicates that the particles were submicron sized. However, sea salt particles injected into
415 the lower troposphere by wind are mainly particles larger than 10 μm . Hence, if the volcanic
416 injection had similar sized NaCl particles, most of them may have quickly fallen out of the
417 stratosphere. In addition, the majority of NaCl might have been washed out during the first
418 couple of hours of plume injection by acting as nuclei for ice particles. It is also possible that the
419 reactions that might release Cl from NaCl may not efficiently lead to reactive Cl. For example,
420 HNO_3 can react on sea salt heterogeneously very quickly in the troposphere to release HCl (De
421 Haan and Finlayson-Pitts, 1997; Guimbaud *et al.*, 2002; Murphy *et al.*, 2019). This reaction may
422 be accelerated by HTHH high humidity even if the temperature is low in the stratosphere. HCl
423 could be removed by condensing in supercooled water, which would reduce HCl vapor
424 concentrations by up to four orders of magnitude, preventing substantial stratospheric chlorine
425 injection [Tabazadeh and Turco, 1993]. Finally, it may be that the water injected came from
426 magmatic water, or seawater that percolated into the volcano and was released as steam. Such
427 water would not be rich in NaCl. In that case Cl observed by Vernier *et al.* [2023] may have been
428 bound up in minerals of the volcanic ash. Other halogen species such as bromine and iodine are
429 often observed after volcanic eruptions (large amounts of BrO were observed after HTHH in the
430 troposphere [Li *et al.*, 2023]). However, they can lead to much stronger ozone depletion if they
431 persist in the stratosphere. Since the elevated Cl in the model can well explain the O_3 depletion,
432 the impact of bromine and iodine on stratospheric O_3 is minimal for this eruption.

433 In addition, NOx can be produced by lightning inside or around the volcanic plume.
434 Observations show there was a record number of lightning events in this volcanic plume. Almost
435 400,000 flashes were observed by the GLD360 network over the 6 hours of the most active
436 eruption period (and ~590,000 total flashes) [Global Volcanism Program, 2022]. Considering
437 that tropospheric global models use a lightning source of 5 Tg(N)/yr and an average flash the
438 OTD/LIS satellite sensors produced an average global flash rate of 44 ± 5 flashes per second, an
439 injection of N of ~0.001- 0.003 Tg (0.002 - 0.006 Tg of NO) would be expected for the HTHH
440 eruption. We conducted a model run with H_2O , SO_2 , and an injection of 0.003 Tg of NO (the
441 H₂O SO₂ NO case), showing that this additional NO has little impact on the O_3 loss and ClO
442 levels during the first ten days (**Figure A6**). Compared to the H₂O SO₂ case, the simulated O_3
443 loss in the H₂O SO₂ NO case increased by $\sim 5 \times 10^5$ molecules/cm³/sec, but at the same time,
444 the O_3 production rate increased by $\sim 5 \times 10^5$ molecules/cm³/sec. The NO+HO₂ reaction rate in
445 the H₂O SO₂ NO case increases 5 times compared with the H₂O SO₂ case. Therefore, lightning
446 NOx probably does not contribute to the HTHH initial in-plume O_3 loss. Because of the high
447 water, NO would convert to HNO_3 in the first couple of days. Unfortunately, we lack
448 observations of HNO_3 , NO, or NO_2 right after the eruption. MLS observations in February
449 (**Figure A7**) and the model simulations with H_2O injection or $\text{H}_2\text{O}+\text{NO}$ injections show elevated
450 HNO_3 compared with the background.

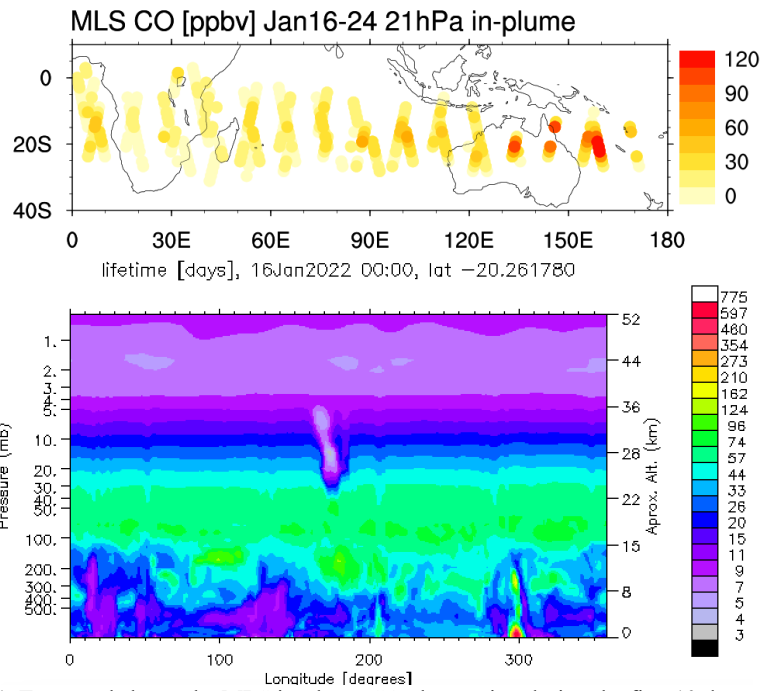
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453 **Appendix A**

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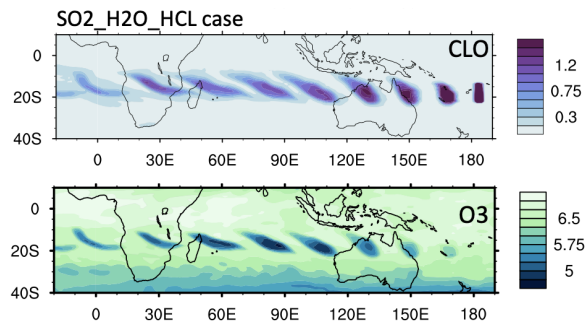
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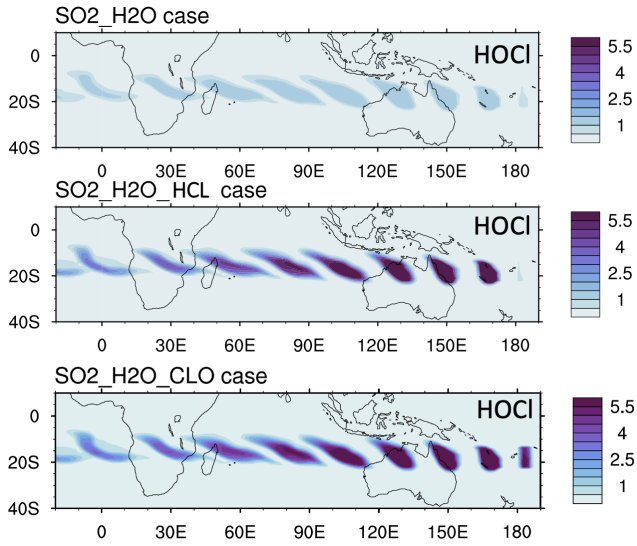
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Figure A1. Top panel shows the MLS in-plume CO observation during the first 10 days after the eruption. The bottom panel shows the CO lifetime on Jan 16 at 20°S is shortened from a month to a few days because of the volcanic water plume. The observed CO mixing ratios of around 120 ppmv seem incompatible with typical CO levels over oceanic regions, indicating the production of CO within the magma chamber or in the hot plume itself.

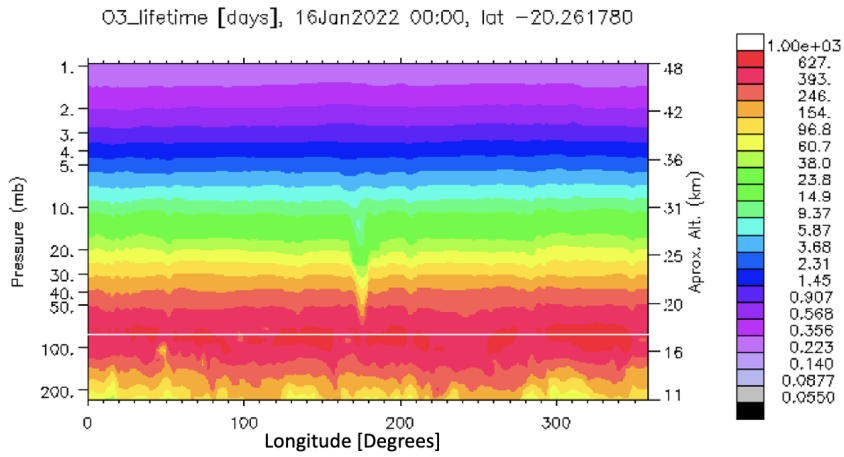


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Figure A2. The O₃ and ClO evolution from the model case with an HCl injection of 0.000092 Tg (equivalent to 0.00013 Tg of ClO injection).



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468 **Figure A3.** The HOCl evolution from the three model cases.



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470 **Figure A4.** O₃ chemical lifetime is about 1 to 2 months at 20 hPa and is reduced to 10 days at the
471 HTHH location.
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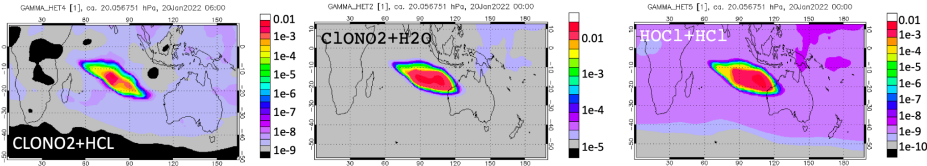


Figure A5. Heterogeneous reaction probabilities for the three heterogeneous reactions on January 20 at 20 hPa.

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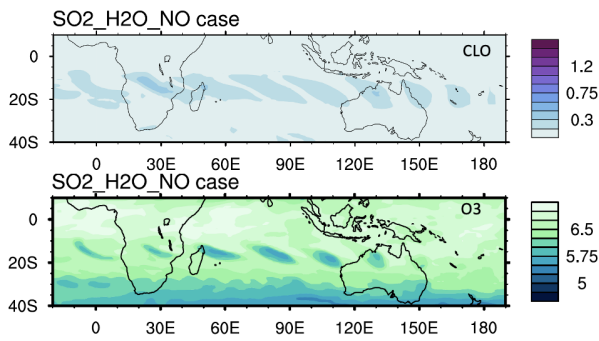
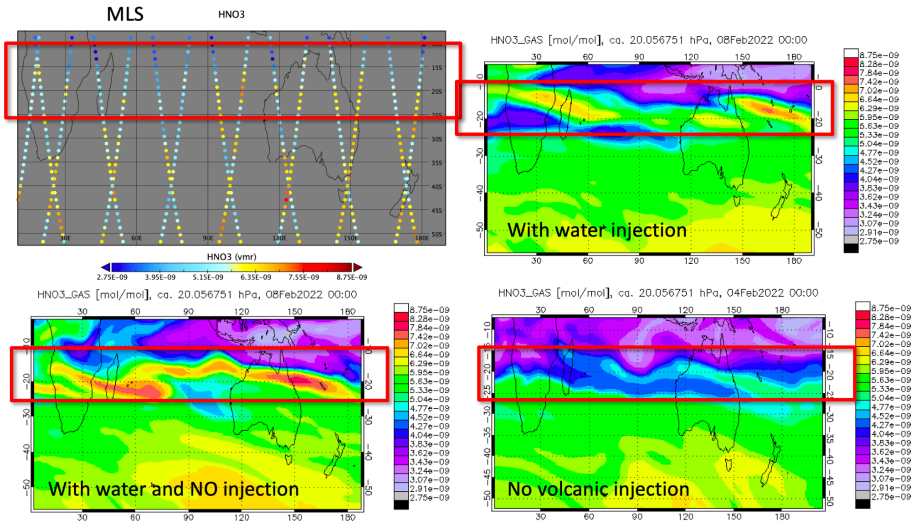


Figure A6. O₃ and ClO evolution from the model case with NO injection of 0.003 Tg, which is identical to the SO₂_H₂O case. The ClO and O₃ enhancement are due to the H₂O injection.

481



482 **Figure A7.** HNO₃ observed by MLS on February 8, 2022 compared to the model simulation
483 with water and NO injection, as well as the no volcanic injection case. MLS shows similar
484 elevated HNO₃ as the simulation case with H₂O injection or with H₂O/NO injection.

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488 **Code availability:** The CESM2 model is available on the CESM trunk to any registered user at
489 www.cesm.ucar.edu.

490 **Data availability:** The main simulation data generated during this study are available at
491 (<https://osf.io/f69ns/>) with a permanent DOI 10.17605/OSF.IO/F69NS. Aura MLS v4 data is
492 available at <https://disc.gsfc.nasa.gov/datasets?page=1&keywords=AURA%20MLS>. Water
493 vapor radiosonde data is available at <https://doi.org/10.5065/p328-z959> (26).

494 **Author contribution:** YZ, RWP, DK, and KHR designed the experiments and YZ performed
495 the simulations. YZ prepared the manuscript with contributions from all co-authors. DK
496 examined the sensitivity of the stratospheric H₂O abundance on the reaction probability (Figure
497 5). LM, HV and SE provided observational data and analysis. RWP, DK, OBT, JZ, ST, CGB,
498 XW, WJR and KHR participated in the modeling data analysis.

499 **Competing interests:** At least one of the (co-)authors is a member of the editorial board
500 of Atmospheric Chemistry and Physics.

501

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