Strong Evidence of Heterogeneous Processing on Stratospheric Sulfate Aerosol in the Extrapolar Southern Hemisphere Following the 2022 Hunga Tonga-Hunga Ha'apai Eruption

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Abstract

The January 2022 eruption of Hunga Tonga-Hunga Ha'apai (HT-HH) caused the largest enhancement in stratospheric aerosol loading in decades and produced an unprecedented enhancement in stratospheric water vapor, which led to strong stratospheric cooling that in turn induced changes in the large-scale circulation. Here we use satellite measurements of gas-phase constituents together with aerosol extinction to investigate the extent to which the thick aerosol, excess moisture, and strong cooling enabled heterogeneous chemical processing. In the southern tropics, unambiguous signatures of substantial chlorine and nitrogen repartitioning appear over a broad vertical domain almost immediately after the eruption, with depletion of N2O5, NOx, and HCl accompanied by enhancement of HNO3, ClO, and ClONO2. After initially rising steeply, HNO3 and ClO plateau, maintaining fairly constant abundances for several months. These patterns are consistent with the saturation of N2O5 hydrolysis, suggesting that this reaction is the primary mechanism for the observed composition changes. The southern midlatitudes and subtropics show similar but weaker enhancements in ClO and ClONO2. In those regions, however, effects of anomalous transport dominate the evolution of HNO3 and HCl, obscuring the signs of heterogeneous processing. Perturbations in chlorine species are considerably weaker than those measured in the southern midlatitude stratosphere following the Australian New Year's fires in 2020. The moderate HT-HH-induced enhancements in reactive chlorine seen throughout the southern middle and low-latitude stratosphere following the Australian New Year's fires in 2020. The moderate HT-HH-induced enhancements in reactive chlorine seen throughout the southern middle and low-latitude stratosphere, far smaller than those in typical winter polar vortices, do not lead to appreciable chemical ozone loss; rather, extrapolar lower-stratospheric ozone remains primarily controlled by dynamical processes.

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Key Points:

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12	•	Satellite data show widespread stratospheric chlorine and nitrogen repartitioning
13		in 2022 following the Hunga Tonga-Hunga Ha'apai eruption
14	•	Observed composition changes are consistent with heterogeneous processing on
15		volcanic sulfate aerosol, in particular the hydrolysis of N_2O_5
16	•	Moderate enhancements in reactive chlorine in the southern mid- and low-latitude
17		stratosphere did not cause appreciable chemical ozone loss

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18 Abstract

The January 2022 eruption of Hunga Tonga-Hunga Ha'apai (HT-HH) caused the largest 19 enhancement in stratospheric aerosol loading in decades and produced an unprecedented 20 enhancement in stratospheric water vapor, leading to strong stratospheric cooling that 21 in turn induced changes in the large-scale circulation. Here we use satellite measurements 22 of gas-phase constituents together with aerosol extinction to investigate the extent to 23 which the thick aerosol, excess moisture, and strong cooling enabled heterogeneous chem-24 ical processing. In the southern tropics, unambiguous signatures of substantial chlorine 25 and nitrogen repartitioning appear over a broad vertical domain almost immediately af-26 ter the eruption, with depletion of N_2O_5 , NO_x , and HCl accompanied by enhancement 27 of HNO₃, ClO, and ClONO₂. After initially rising steeply, HNO₃ and ClO plateau, main-28 taining fairly constant abundances for several months. These patterns are consistent with 29 the saturation of N_2O_5 hydrolysis, suggesting that this reaction is the primary mecha-30 nism for the observed composition changes. The southern midlatitudes and subtropics 31 show similar but weaker enhancements in ClO and ClONO₂. In those regions, however, 32 effects of anomalous transport dominate the evolution of HNO_3 and HCl, obscuring the 33 signs of heterogeneous processing. Perturbations in chlorine species are considerably weaker 34 than those measured in the southern midlatitude stratosphere in 2020 following the Aus-35 tralian New Year's fires. The moderate HT-HH-induced enhancements in reactive chlo-36 rine seen throughout the southern middle and low-latitude stratosphere, far smaller than 37 those in typical winter polar vortices, do not lead to appreciable chemical ozone loss; rather, 38 extrapolar lower-stratospheric ozone remains primarily controlled by dynamical processes. 30

⁴⁰ Plain Language Summary

The January 2022 eruption of the Hunga Tonga-Hunga Ha'apai (HT-HH) volcano 41 caused the largest increase in stratospheric aerosol in decades and produced an unprece-42 dented increase in stratospheric water vapor, which led to strong stratospheric cooling 43 that in turn altered stratospheric winds. Here we use satellite measurements to inves-44 tigate the extent to which the thick aerosol, excess moisture, and strong cooling enabled 45 heterogeneous chemical processing. In the southern tropics, unambiguous signatures of 46 substantial changes in several chlorine and nitrogen species appear over a broad verti-47 cal range almost immediately after the eruption, with decreases in some species mirrored 48 by increases in others. The southern midlatitudes and subtropics show similar but weaker 49 evidence of chemical processing. In those regions, however, effects of anomalous wind pat-50 terns obscure the signs of heterogeneous processing in some species. Perturbations in chlo-51 rine are considerably weaker than those measured in the southern midlatitude strato-52 sphere following the Australian New Year's fires in 2020. The moderate HT-HH-induced 53 enhancements in reactive chlorine seen throughout the southern middle and low-latitude 54 stratosphere, far smaller than those in typical winter polar vortices, do not lead to ap-55 preciable chemical ozone loss; rather, outside the polar regions, lower-stratospheric ozone 56 remains primarily controlled by dynamical processes. 57

58 1 Introduction

The Hunga Tonga-Hunga Ha'apai (HT-HH) volcano in the Kingdom of Tonga be-59 gan an eruptive sequence in December 2021 that culminated in a paroxysmal explosion 60 on 15 January 2022 (e.g., Carn et al., 2022). It was the most explosive submarine erup-61 tion witnessed during the satellite era to date. Satellite and ground-based measurements 62 indicate that a substantial amount of material was lofted into the upper stratosphere; 63 although the bulk was deposited at altitudes between 20 and 30 km, the highest over-64 shooting tops of the eruption column penetrated into the lower mesosphere, reaching an 65 unprecedented \sim 55 km altitude (Carr et al., 2022; Khaykin et al., 2022; Millán et al., 2022; 66 Taha et al., 2022). The climactic blast was marked by the (previously undocumented 67

for a volcanic eruption) occurrence of a double umbrella cloud (that is, a cloud that spreads 68 laterally around the level of neutral buoyancy and rapidly transports the ejected volume 69 away from the volcano in all directions), with distinct signatures at about 30 km and near 70 the tropopause at $\sim 17 \,\mathrm{km}$ (Gupta et al., 2022). Also unprecedented was the mass of wa-71 ter vapor injected by HT-HH into the stratosphere, which has been estimated using Mi-72 crowave Limb Sounder measurements to have increased the total stratospheric water va-73 por burden by $\sim 10\%$ (Millán et al., 2022); other studies have calculated similar values 74 for the magnitude of the water vapor injection (Khaykin et al., 2022; Vömel et al., 2022; 75 Xu et al., 2022). Analysis of its isotopic composition shows that the injected water de-76 rived mainly from vaporized seawater, although some tropospheric moisture may also 77 have been entrained by the eruptive updraft (Khaykin et al., 2022). Peak enhancements 78 in water vapor were observed in the 25–30 km region (Khaykin et al., 2022; Millán et al., 79 2022; Sellitto et al., 2022; Vömel et al., 2022; Xu et al., 2022), consistent with being sourced 80 from the upper umbrella cloud (Gupta et al., 2022). The eruption column was also rich 81 in ice (Gupta et al., 2022; Khaykin et al., 2022); subsequent sedimentation and subli-82 mation of ice led to relatively humid conditions in the lower stratosphere (Khaykin et 83 al., 2022; Millán et al., 2022). 84

Although the explosive energy of HT-HH was similar to or even larger than that 85 of the 1991 eruption of Mt. Pinatubo (Poli & Shapiro, 2022; Wright et al., 2022; Yuen 86 et al., 2022), its SO₂ discharge into the stratosphere was $\sim 1-2$ orders of magnitude smaller 87 (Carn et al., 2022; Millán et al., 2022; Sellitto et al., 2022). SO₂ emissions from subma-88 rine volcanoes are typically lower than those from non-hydromagmatic eruptions of com-89 parable magnitude, presumably because of significant scrubbing of SO₂ in their water-90 rich eruption clouds (Carn et al., 2022). In addition, the SO₂ from HT-HH that did en-91 ter the stratosphere had a considerably shortened lifetime, undergoing unusually rapid 92 conversion to secondary sulfate aerosol (Legras et al., 2022; Sellitto et al., 2022). The 93 exceptionally fast formation of sulfate aerosol is also attributable to the abundant co-94 emitted water vapor, which provides a source of OH that in turn catalyzes the oxida-95 tion of SO_2 into H_2SO_4 , which then nucleates into sulfate aerosol (Zhu et al., 2022). With 96 its modest SO₂ emissions, HT-HH caused global-mean anomalies in stratospheric aerosol 97 optical depth smaller by about a factor of 6 than those resulting from Pinatubo (Khaykin 98 et al., 2022; Sellitto et al., 2022). Nevertheless, HT-HH led to the largest perturbation 99 in the stratospheric aerosol loading in the last 30 years (Khaykin et al., 2022; Sellitto et 100 al., 2022; Taha et al., 2022). 101

For the first few months after the eruption, the main volcanic cloud at $20-30 \,\mathrm{km}$ 102 remained largely confined within the tropical pipe as it encircled the globe; thereafter 103 it gradually dispersed meridionally, with more efficient transport in the Southern Hemi-104 sphere, such that by June 2022 it had spread from about 60°S to 30°N, with small frag-105 ments reaching higher northern latitudes (Coy et al., 2022; Legras et al., 2022; Millán 106 et al., 2022; Schoeberl et al., 2022, 2023; Taha et al., 2022). The strong transport bar-107 rier at the boundary of the Antarctic vortex prevented the plume from penetrating to 108 high southern latitudes until that vortex broke down in late 2022 (Manney et al., 2023). 109 Initially the enhanced aerosol and water vapor were spatially coincident, but they grad-110 ually diverged. Entrained within the ascending branch of the Brewer-Dobson circulation. 111 the moisture anomaly was carried upward in the months following its injection, whereas 112 the sulfate aerosols were subject to gravitational settling; hence the aerosol- and water-113 rich layers became increasingly separated in the vertical after mid-February (Legras et 114 al., 2022; Schoeberl et al., 2022). 115

Both the aerosol and the water vapor perturbations induced radiative effects, but infrared cooling from enhanced water vapor concentrations was shown to dominate (Legras et al., 2022; Sellitto et al., 2022; Jenkins et al., 2023), leading to the development of a strong and persistent low-temperature anomaly in the mid-stratosphere shortly after the eruption (Coy et al., 2022; Schoeberl et al., 2022; Vömel et al., 2022). The anomalous temperature structure in turn induced changes in the large-scale circulation; in particular, zonal winds in the mid-stratosphere strengthened, as did meridional flow, while descent in the middle and high latitudes slowed (Coy et al., 2022).

Enhanced sulfate aerosol loading following volcanic injection of SO_2 into the strato-124 sphere is known to perturb the partitioning of key chemical families controlling strato-125 spheric ozone (e.g., Solomon, 1999). The exceptional stratospheric hydration and ensu-126 ing temperature anomalies caused by HT-HH may have increased such chemical process-127 ing. Transport also plays a major role in shaping trace gas distributions, however, and 128 129 the alterations to the stratospheric circulation arising from the eruption may have either compounded or attenuated the chemical effects, complicating interpretation of ob-130 served changes in composition. Here we investigate the impact of the HT-HH eruption 131 on stratospheric composition using satellite measurements. 132

¹³³ 2 Data and Methods

We examine measurements of H_2O , HNO_3 , ClO, HCl, N_2O , and O_3 from the Aura 134 Microwave Limb Sounder (MLS) (e.g., Waters et al., 2006). The enormous enhancement 135 in stratospheric water vapor from HT-HH, with the largest concentrations more than an 136 order of magnitude greater than any previously observed by MLS, degraded the accu-137 racy of some of the products retrieved using the most recent MLS data processing al-138 gorithms, version 5 (v5). It also caused a large proportion of the measurements in the 139 plume to fail the recommended MLS quality screening for the first few weeks after the 140 eruption, with retrieval performance not returning to normal until 8 February 2022. For 141 these reasons, Millán et al. (2022) used v4 MLS data to study the early stages of HT-HH 142 plume evolution; moreover, they did not impose quality filtering on the data. Since v5 143 generally represents an improvement over v4, particularly for H_2O and N_2O , and since 144 our main focus here is on the perturbations in composition arising in the weeks to months 145 following the eruption, we use v5 MLS data for all species, with standard data screen-146 ing protocols applied (Livesey et al., 2020). 147

MLS v5 ClO suffers from non-negligible biases at the lowest retrieval levels, 68-148 147 hPa (Livesey et al., 2020). As no substantial enhancement in nightime ClO is ex-149 pected in the extrapolar regions of interest here, the ClO biases are mitigated by tak-150 ing ascending ($\sim 2-3$ PM at these latitudes) minus descending ($\sim 2-3$ AM) differences, 151 similar to the strategy adopted in a prior study of midlatitude chlorine partitioning by 152 Santee et al. (2022). MLS N₂O is subject to a negative drift that, while partially alle-153 viated in v5, remains about -3%/decade in the lower stratosphere (Livesey et al., 2021). 154 Accordingly, the MLS N_2O fields presented below have been "detrended" by removing 155 a linear fit to the daily values across the Aura mission at each gridpoint. 156

We use meteorological fields (temperature, potential temperature, potential vor-157 ticity (PV)) from the Modern Era Retrospective Analysis for Research and Applications 158 Version 2 (MERRA-2, Gelaro et al., 2017). Satellite data are cast into quasi-Lagrangian 159 PV-based equivalent latitude (EqL) (Butchart & Remsberg, 1986) and potential tem-160 perature coordinates to facilitate tracking stratospheric air motions and allow parcels 161 with similar dynamical histories to be grouped together. Each day on average ~ 330 MLS 162 profiles fall within each of the 16° EqL bands defined here: 38°S-54°S EqL, 22°S-38°S 163 EqL, and 6°S–22°S EqL, representing southern midlatitudes, subtropics, and tropics, re-164 spectively. 165

In addition to MLS data, we use version 4.1/4.2 measurements of N₂O₅, NO, NO₂, and ClONO₂ from the Atmospheric Chemistry Experiment-Fourier Transform Spectrometer (ACE-FTS; Boone et al., 2020), screened using the recommended quality flags (Sheese et al., 2015; Sheese & Walker, 2020). In conjunction with the trace gas measurements, we use the latest version (V2.1) aerosol extinction coefficient profiles obtained by the Suomi-

NPP Ozone Mapping and Profiler Suite Limb Profiler (OMPS-LP) at both 869 nm, as 171 recommended for tracking stratospheric injections from events such as volcanic eruptions 172 or wildfires (Taha et al., 2021), and 745 nm. According to Taha et al. (2022), the stan-173 dard V2.1 OMPS-LP algorithms provide accurate aerosol retrievals as long as the vol-174 canic cloud is below 36 km; our analysis here is confined to the region below $\sim 33 \text{ km} (900 \text{ K})$. 175 We also compared the NASA OMPS-LP V2.1 aerosol products with the tomographic re-176 trievals at 745 nm produced by the University of Saskatchewan (USask v1.2; Bourassa 177 et al., 2023). Although the NASA and USask aerosol products show substantial differ-178 ences at low latitudes below 500 K, those discrepancies have no bearing on our conclu-179 sions, and for simplicity only the NASA aerosol data at 869 nm are shown here. As in 180 the study by Santee et al. (2022), for interpolation to isentropic surfaces, the aerosol data 181 are first converted to cross section per mole of air (in units of $m^2 mol^{-1}$), which, unlike 182 the more commonly used extinction coefficient (in units of $\rm km^{-1}$), is conserved under 183 changes of atmospheric pressure in the absence of aerosol formation or loss. The con-184 version factor relating cross section to extinction is the local molar volume, RT/p (in units 185 of $m^3 mol^{-1}$), where R is the molar gas constant, T is temperature, and p is pressure. 186 We use the remotely sensed optical properties without conversion to aerosol surface area 187 or volume density, as would be necessary to use these data as inputs in chemical mod-188 els (e.g., Hervig & Deshler, 1998; Kovilakam & Deshler, 2015). 189

3 Aerosol-Mediated Chemical Repartitioning 190

Within a few weeks of an explosive volcanic eruption, oxidative processing of the 191 SO_2 injected into the stratosphere leads to the production of liquid sulfate particles, en-192 hancing the aerosol surface area density (SAD) available for heterogeneous chemical pro-193 cessing (e.g., Bekki & Pyle, 1994). Heterogeneous reactions of particular importance in 194 the stratosphere include (e.g., Solomon, 1999): 195

$$N_2O_5 + H_2O \to 2 HNO_3 \tag{R1}$$

$$ClONO_2 + H_2O \rightarrow HOCl + HNO_3$$
 (R2)

$$CIONO_2 + HCl \rightarrow Cl_2 + HNO_3$$
(R3)

TT 0

$$HOCI + HCI \to Cl_2 + H_2O \tag{R4}$$

 (\mathbf{D}, \mathbf{i})

$$BrONO_2 + H_2O \rightarrow HOBr + HNO_3$$
 (R5)

Nitrogen, chlorine, and hydrogen photochemistry are strongly coupled, thus changes in 196 the partitioning of one chemical family affect species in other families. Substantial per-197 turbations in chemical partitioning can persist, as it can take a year or more for volcanic 198 sulfate aerosols to be removed from the stratosphere through sedimentation. To set the 199 stage for the interpretation of satellite measurements in Section 4, we first summarize 200 the expected impacts of R1–R5 on several key nitrogen and chlorine species affecting strato-201 spheric ozone, as well as ozone itself, under volcanically enhanced aerosol conditions. The 202 chemical processes discussed here are described in greater detail by, for example, Bekki 203 and Pyle (1994), Kinnison et al. (1994), Tie et al. (1994), Coffey (1996), Solomon (1999), 204 Anderson et al. (2012), Berthet et al. (2017), and Robrecht et al. (2019). For brevity, 205 these references are not cited for all relevant points in the following text. 206

3.1 HNO₃, N_2O_5 , and NO_x

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 HNO_3 is the major reservoir for odd nitrogen, NO_y , in the lower stratosphere. In 208 the immediate aftermath of the HT-HH eruption, a few individual MLS profiles showed 209 strong stratospheric enrichments in several trace gases, including HNO₃ (Millán et al., 210 2022, their Figure S1). Although it has been measured in volcanic plumes near the ground 211 (e.g., Mather et al., 2004; Oppenheimer et al., 2010; Voigt et al., 2014), in the lower strato-212

sphere volcanogenic HNO_3 has been conclusively detected only once, when a peak mix-213 ing ratio of ~ 3 ppbv above background was measured in an encounter with the fresh vol-214 canic cloud from the February 2000 eruption of Hekla (Hunton et al., 2005; Rose et al., 215 2006). Major explosive eruptions can enhance stratospheric HNO₃ through injection, ei-216 ther of HNO_3 itself or of nitrogen-bearing gases (e.g., ammonia) from which it can be 217 generated. Volcanic lightning can also induce HNO_3 formation, and, in addition to its 218 other unprecedented characteristics, HT-HH produced prodigious amounts of lightning 219 (Yuen et al., 2022). Thus, a direct volcanic origin for the anomalous post-eruption MLS 220 HNO_3 profiles is possible. However, they are more likely to be retrieval artifacts aris-221 ing from contamination of the MLS HNO_3 spectral signature by SO_2 . In any case, these 222 few highly localized and transient HNO₃ enhancements do not significantly affect the 223 broad EqL-band averages analyzed in this study, and they are not considered further. 224

Hydrolysis of N_2O_5 (R1) is fairly insensitive to temperature (Figure S1 in Support-225 ing Information) and takes place rapidly under virtually all stratospheric conditions. Its 226 rate is intensified dramatically when a major volcanic eruption enhances aerosol SAD. 227 Since (i) N_2O_5 is the main nighttime reservoir for reactive nitrogen, NO_x (defined here 228 as $NO+NO_2$, (ii) the HNO₃ produced through R1 returns to the gas phase, and (iii) the 229 rate at which HNO_3 undergoes conversion back to NO_x is unchanged, an increase in SAD 230 is generally expected to be accompanied by observed decreases in the abundances of both 231 N_2O_5 and NO_x and a corresponding increase in HNO₃. Similarly, R5 (hydrolysis of BrONO₂) 232 can be active at all latitudes and seasons during periods of high aerosol loading. 233

 $ClONO_2$ hydrolysis (R2) and reaction with HCl (R3) on sulfate aerosol are also po-234 tential sinks of NO_x and sources of gas-phase HNO₃. Unlike for R1 and R5, the reac-235 tion probabilities of R2 and R3 are strongly dependent on particle water content and hence 236 temperature (Figure S1 in Supporting Information), and typically they only become com-237 petitive with R1 in affecting nitrogen (and chlorine) partitioning under cold $(T < \sim 200 \text{ K})$ 238 conditions, such as in the winter polar regions, where they take place on the surfaces of 239 polar stratospheric cloud (PSC) particles. These two reactions, however, can come into 240 play to convert NO_x to HNO_3 in the midlatitude lower stratosphere when SAD and wa-241 ter vapor mixing ratios (and hence the aerosol water fraction) are sufficiently elevated 242 (Keim et al., 1996). Whereas condensed HNO₃ remains sequestered in PSCs in the win-243 ter polar regions, at higher temperatures a large proportion of the HNO_3 formed by these 244 reactions is released to the gas phase (Robrecht et al., 2019). 245

Consistent with heterogeneous processing, substantial reductions in NO_x and/or 246 N₂O₅ have been seen following several large eruptions (e.g., Fahey et al., 1993; Mills et 247 al., 1993; Koike et al., 1994; Rinsland et al., 1994; Coffey, 1996; Adams et al., 2017; Berthet 248 et al., 2017; Zambri et al., 2019). Significant decreases in NO_x observed in the months 249 after severe wildfires in Australia in the austral summer of 2019/2020 have also pointed 250 to the occurrence of heterogeneous processing on smoke particle surfaces (Solomon et 251 al., 2022; Strahan et al., 2022). Observational evidence for associated increases in HNO₃ 252 has been more ambiguous, however. Strongly enhanced HNO_3 (as high as 40-50% above 253 unperturbed background values in some cases) was measured in aged stratospheric vol-254 canic clouds sampled weeks or months after major eruptions (Arnold et al., 1990; Koike 255 et al., 1994; Rinsland et al., 1994; Jurkat et al., 2010). In addition, declining trends in 256 HNO_3 abundances as the sulfate aerosol loading from the eruption of Mt. Pinatubo slowly 257 decayed in subsequent years were attributed to the diminishing impact of R1 (David et 258 al., 1994; Kumer et al., 1996; Slusser et al., 1998; Rinsland et al., 2003; Santee et al., 2004). 259 In other cases, by contrast, no clear volcanic enhancement in HNO₃ was seen (Coffey, 260 1996; Adams et al., 2017; Berthet et al., 2017). Nor were HNO₃ abundances elevated fol-261 lowing the 2019/2020 Australian fires (Santee et al., 2022; Strahan et al., 2022). 262

Zambri et al. (2019) used coupled chemistry-climate model simulations in conjunc tion with MLS and other satellite measurements to investigate the impact on stratospheric
 composition of multiple moderate-magnitude eruptions in the post-Pinatubo era. Neg-

ative anomalies (with respect to unperturbed background values) in response to erup-266 tive events were seen in modeled N_2O_5 and measured and modeled NO_x , particularly 267 at 50–70 hPa, where anomalies exceeded 20–25%. The model showed a clear relationship 268 between SAD and HNO_3 changes through much of the stratosphere, but the perturbations in HNO_3 were not entirely congruent with those in NO_x . In particular, positive anoma-270 lies in HNO_3 extended over a larger altitude range, exceeding 10% during periods of in-271 creased sulfate aerosol SAD at levels from 70 to 5 hPa, with the exception of 50 hPa, where 272 fractional changes were much smaller than those in NO_x and confined mainly to the trop-273 ics. MLS data indicated similar, albeit weaker, aerosol-induced HNO_3 increases, again 274 except near 50 hPa. Zambri et al. (2019) offered no explanation for the behavior seen in 275 both measured and modeled HNO_3 fields at 50 hPa. Other recent modeling studies have 276 also found post-eruption reductions in NO_x and N_2O_5 of as much as 25-50% but con-277 comitant increases in HNO_3 of only 5–15% (Adams et al., 2017; Berthet et al., 2017). 278

It may not be surprising that a strong correlation between SAD/NO_x and HNO_3 is not consistently observed; HNO_3 concentrations in the lower stratosphere are much larger than those of NO_x , and thus comparable changes in its abundance have only modest relative effects. Small changes are difficult to detect against the backdrop of natural dynamical variability, which is of order 10% for HNO_3 in the lower stratosphere, making HNO_3 a less sensitive indicator of the occurrence of R1 than NO_x (e.g., Adams et al., 2017; Solomon et al., 2022; Strahan et al., 2022).

3.2 ClO

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ClO is the main form of ozone-destroying reactive chlorine. In the week following 287 the HT-HH eruption, MLS recorded strongly enhanced ClO in a handful of profiles (Millán 288 et al., 2022). The only halogenated compounds known to have been volcanically injected 289 into the stratosphere are HCl (discussed in Section 3.3) and OClO (Theys et al., 2014), 290 and the measured high ClO values were thought to largely reflect MLS retrieval artifacts 291 arising from SO_2 spectral interference, rather than direct stratospheric injection of ClO. 292 Given its rapid transformation into sulfate aerosol, the SO₂ from HT-HH no longer com-293 promised the reliability of MLS retrievals after late January, when abundances reverted 294 to background levels (Millán et al., 2022). As we are concerned here with the signatures 295 of spatially extensive chemical processing that manifest weeks to months after the HT-HH 296 injection, the initial enhancements in a few ClO profiles are not considered further. 297

Post-eruption decreases in NO_x impede $CIONO_2$ formation, shifting chlorine par-298 titioning toward ClO. These gas-phase processes affect ClO and ClONO₂ abundances 299 while leaving HCl unchanged. In addition, photolysis of the gas-phase HNO₃ and HOBr 300 resulting from the hydrolysis of N₂O₅, ClONO₂, and BrONO₂ (R1, R2, and R5) is a source 301 of reactive hydrogen, HO_x (OH+HO₂), and reduced NO_x concentrations also bring about 302 increases in HO_x by inhibiting the rate of NO₂+OH+M \rightarrow HNO₃+M. In the case of HT-HH, 303 moreover, simulations show that the massive injection of water vapor rapidly increased 304 stratospheric OH abundances (Zhu et al., 2022). Elevated OH abundances in turn ac-305 celerate conversion of HCl to reactive chlorine via $HCl+OH\rightarrow Cl+H_2O$. Thus, small en-306 hancements in stratospheric ClO may occur following major volcanic eruptions from R1 307 and R5 alone, under conditions unfavorable for R2–R4 (which may also occur on vol-308 canic sulfate aerosol where ambient temperatures and water vapor abundances allow). 309 Although in general the consequences of increased OH on stratospheric chlorine parti-310 tioning are expected to be relatively minor, in the extremely water-rich environment af-311 ter HT-HH, the gas-phase oxidation of HCl may be intensified. 312

Enhancements in ClO of a few tens to 100 ppt observed outside of the polar regions in the months following the eruption of Mt. Pinatubo were attributed to heterogeneous processing on sulfate aerosol, in particular R1 (Avallone et al., 1993; Dessler et al., 1993; Fahey et al., 1993; Toohey et al., 1993; Wilson et al., 1993; Keim et al., 1996). In their study of the effects of moderate eruptions since Pinatubo, Zambri et al. (2019) found responses of as much as 20–50 pptv in simulated ensemble-mean monthly mean zonal-mean
ClO from 70–5 hPa, but they showed an overall weaker correlation with SAD than that
of HNO₃. In many cases, changes similar to those simulated were not seen in Aura MLS
v4.2 bias-corrected ClO data. Those authors asserted that the precision of the MLS ClO
measurements was too poor to permit detection of small anomalies, basing that statement on the MLS single-profile precision without considering the benefits of averaging.

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3.3 HCl and ClONO₂

HCl and ClONO₂ are the main reservoirs of chlorine in the stratosphere. As discussed in Section 3.2, in the weeks following an explosive eruption, HCl can decrease through destruction by enhanced OH (since the rate of HCl loss through that mechanism is faster than its formation via Cl+CH₄ \rightarrow HCl+CH₃), and OH abundances may be especially elevated after the HT-HH water vapor injection. Heterogeneous chlorine chemistry (R2– R4) may also contribute if conditions allow. To our knowledge, however, HCl depletion induced by heterogeneous reactions on volcanic sulfate aerosol has never been observed.

MLS has captured direct volcanic injection of HCl into the stratosphere by a num-332 ber of moderate eruptions (Prata et al., 2007; Theys et al., 2014; Carn et al., 2016), al-333 though peak volcanic HCl concentrations are likely underestimated since the vertical ex-334 tent of the plumes is typically much smaller than the vertical resolution of MLS mea-335 surements. Only eight HCl profiles in mid-January 2022 exceeded the threshold used by 336 Millán et al. (2022) to identify enhancements (7σ , corresponding to ~5 ppbv of HCl through 337 much of the domain), so no spikes can be expected to stand out in the HCl zonal means 338 presented below. Scavenging by hydrometeors (liquid water drops or ice particles) or ash 339 in eruption columns prevents much of the HCl emitted by major eruptions from reach-340 ing the stratosphere (Tabazadeh & Turco, 1993; Textor et al., 2003). Klobas et al. (2017) 341 suggested that the very weak stratospheric HCl injection from Mt. Pinatubo (Mankin 342 et al., 1992; Wallace & Livingston, 1992) is attributable to the extremely wet conditions 343 caused by the coincidental passage of a tropical cyclone during the eruption, which may 344 have allowed most of the degassed HCl to be scrubbed from its plume. Noting the con-345 trast in HCl signals between the Sarychev and Kasatochi events despite their compara-346 ble magnitudes, Carn et al. (2016) postulated that the weaker HCl injection from the 347 latter arose through more effective scavenging by abundant water in its plume since it 348 erupted through a pre-existing crater lake. Similarly, the relatively modest HCl injec-349 tion from HT-HH may have been a consequence of removal in the exceptionally moist 350 and ice-rich environment in its plume (Carn et al., 2022). Nevertheless, it is conceivable 351 that volcanogenic HCl could compensate to some degree any depletion caused by het-352 erogeneous processing in the early post-eruption period. 353

If R2 or R3 are active to any significant extent, then ClONO₂ should decrease. On the other hand, some model simulations have shown mild increases in ClONO₂ (e.g., 20 pptv, \sim 16%) under enhanced volcanic SAD conditions, since ClO (the limiting reactant in ClO+NO₂+M→ClONO₂+M) increases (Kinnison et al., 1994; Berthet et al., 2017), while other modeling studies have found no discernible response in ClONO₂ (Zambri et al., 2019).

3.4 Ozone

360

The volcanically induced changes in stratospheric chlorine and nitrogen partitioning described above exacerbate chemical ozone destruction by the HO_x and ClO_x catalytic cycles (of primary importance in the lower stratosphere) but impede that by the NO_x cycle (the dominant ozone loss mechanism in the middle stratosphere). In addition to those direct chemical perturbations, increases in aerosol SAD affect radiative balance and hence the large-scale stratospheric circulation, with potentially considerable conse-

quences for the distribution of ozone (and other trace gases). Enhanced aerosol loading 367 can also alter photolysis rates (e.g., Tie et al., 1994), further modifying ozone abundances. 368 The net impact of these (in some cases competing) effects depends on the amount of sul-369 fate aerosol produced from the emitted SO_2 and the latitude, altitude, and timing of the 370 injection. Substantial reductions in lower-stratospheric and total column ozone have been 371 observed and modeled after several previous volcanic eruptions (e.g., Dhomse et al., 2015; 372 Hofmann & Solomon, 1989; Kilian et al., 2020; Millard et al., 2006; Naik et al., 2017; Ran-373 del et al., 1995; Solomon et al., 2016; Stone et al., 2017; Wilka et al., 2018, see also the 374 citations listed at the beginning of this Section, and references therein). On the other 375 hand, significant enhancement in ozone was observed in the southern midlatitudes above 376 ~ 26 km following Pinatubo; model simulations confirmed the important role of the NO_x 377 loss cycle—and its suppression through R1 on volcanic sulfate aerosol—in determining 378 the ozone budget at higher altitudes (Mickley et al., 1997). 379

380 4 Results

The unprecedented stratospheric impacts of HT-HH are clear in Figure 1, which 381 provides an overview of the evolution of aerosol cross section from OMPS-LP (panel (a)) 382 and water vapor from MLS (panel (b)) over their respective missions at 620 K potential 383 temperature, a representative lower-stratospheric level that corresponds to $\sim 20-30$ hPa 384 or $\sim 24-26$ km, depending on the EqL band and month. The previously reported initial 385 low-latitude confinement, early northward extension, and eventual more efficient south-386 ward dispersion of the aerosol and water vapor plumes (Coy et al., 2022; Legras et al., 387 2022; Millán et al., 2022; Schoeberl et al., 2022; Taha et al., 2022) are evident. Large anoma-388 lies in other chemical constituents also arise in the aftermath of the eruption. A strong 389 enhancement in HNO_3 quickly develops at the lowest latitudes and then gradually spreads 390 poleward in both hemispheres over the next several months (Figure 1c). HNO_3 anoma-391 lies of comparable magnitude are not infrequent in the 18^+ -year MLS record, however, 392 and around mid-year the anomaly abruptly shifts sign, becoming strongly negative. In 393 contrast, at this altitude the enhancement in ClO in the months after the eruption is un-394 paralleled (Figure 1d), while HCl is largely unaffected initially but exhibits a severe low 395 anomaly later in the year (Figure 1e). In the following subsections, we explore the ex-396 tent to which these perturbations can be attributed to heterogeneous chemical process-397 ing. To elucidate the role of dynamics in controlling trace gas distributions, we also ex-398 amine N_2O , a long-lived transport tracer whose vertical and meridional gradients in much 399 of the lower stratosphere are opposite to those of HNO_3 , HCl, and O_3 and which is there-400 fore generally anticorrelated with those species in that region (Figure 1f). Finally, Fig-401 ure 1g shows the unremarkable evolution of ozone in 2022. 402

We focus this analysis on the Southern Hemisphere (SH) midlatitudes, 38°S–54°S 403 EqL, and the SH tropics, 6°S–22°S EqL. The former allows comparison of the volcano-404 induced perturbations in chlorine and nitrogen partitioning with those engendered by 405 the Australian New Year's (ANY) fires in late 2019 / early 2020 (Santee et al., 2022; Solomon 406 et al., 2022, 2023; Strahan et al., 2022) and is also characterized by somewhat better ACE-FTS 407 data coverage; the latter experiences the largest HT-HH impacts on stratospheric com-408 position but suffers from poor ACE-FTS sampling. We also briefly examine the inter-409 mediate region of the SH subtropics, 22°S–38°S EqL. Equivalent latitudes poleward of 410 54°S are ignored to minimize the confounding influence of seasonal chemical processing 411 inside the Antarctic winter polar vortex. 412

In Figure 2 for the SH midlatitudes and similar figures for the other two EqL bands, we place the 2022 observations into climatological context and examine the evolution of chemical species in relation to sulfate aerosol loading, humidity, temperature, and transport. In the SH midlatitudes, we show 500 K (corresponding to ~47–57 hPa or ~20–21 km) to enable ANY comparisons and because the 2022 perturbations in ClO in that EqL band are largest there. In the SH subtropics and tropics, as in Figure 1, we again show 620 K, where the ClO enhancements at those latitudes peak. The two preceding years, 2020 and
2021, are also highlighted. Figure 3 and similar figures provide information on the vertical extent of the perturbations. Anomalies are calculated as described in the relevant
figure captions. Normalized anomalies are shown in the Supporting Information to facilitate comparison of changes across different quantities and EqL bands.

424

4.1 Southern Hemisphere Midlatitudes (38°S–54°S EqL)

We begin by looking at the SH midlatitudes, focusing on 500 K. Appreciable aerosol 425 from HT-HH starts arriving in mid-April in this region, where it swiftly surpasses that 426 resulting from ANY in 2020 and vastly exceeds the OMPS-LP mission envelope of be-427 havior (Figure 2a). These results are not sensitive to the choice of aerosol data set ex-428 amined (not shown; see Section 2). A corresponding positive anomaly in water vapor dis-429 plays a slight lag in its arrival relative to aerosol in this EqL band; although water va-430 por mixing ratios at 500 K are above the climatological mean through most of 2020 and 431 2021 and remain so into 2022 (as noted also by Manney et al., 2023), they do not turn 432 sharply upward at this level until mid-June (Figure 2b). Comparable delays in the up-433 turn in water vapor relative to that in aerosol occur at most levels (cf. Figures 3h and 434 3i; see also Figure S2 in Supporting Information). Khaykin et al. (2022) similarly found 435 slightly earlier appearance of aerosol than water vapor anomalies in the SH midlatitudes, 436 again based on OMPS-LP and MLS data. The increases in midlatitude aerosol and wa-437 ter vapor seen at 500 K likely result from a combination of poleward flow along that isen-438 trope and transport at higher levels (where the main volcanic cloud was initially deposited, 439 as noted in the Introduction) followed by downward motion in this EqL band, either through 440 gravitational settling in the case of aerosol or via diabatic descent for water vapor. We 441 hypothesize that the discrepancy in the onset of the enhancements in aerosol and wa-442 ter vapor in the SH midlatitudes arises from a faster aerosol sedimentation rate as com-443 pared to the rate of water vapor descent in the stratospheric circulation. 444

⁴⁴⁵ Climatological temperatures in this EqL band at 500 K decrease from January into ⁴⁴⁶ July; in 2022, temperatures are below average after mid-April and reach record lows in ⁴⁴⁷ mid-August (Figure 2c). A sizeable negative anomaly in N₂O through much of 2021 con-⁴⁴⁸ tinued into 2022, leading to low N₂O values at levels around 500 K for the first half of ⁴⁴⁹ the year (Figure 2d; see also Figures 3f and 3m). Another transport tracer measured by ⁴⁵⁰ MLS, CH₃Cl, exhibits similar, albeit weaker, departures from climatology in early 2022 ⁴⁵¹ (not shown), corroborating the picture seen in N₂O.

The evolution of the nitrogen species over the first few months of 2022 is more or 452 less typical; although mixing ratios of N_2O_5 (Figure 2f), NO_x (NO+NO₂, Figure 2g), 453 and HNO_3 (Figure 2h) at 500 K are initially below average, they all generally track the 454 normal seasonal behavior at first. By early June, however, N₂O₅ values drop well be-455 low any previously observed in this region by ACE-FTS, and NO_x is also slightly below 456 average. HNO_3 gradually increases relative to climatology until a very small positive anomaly 457 develops in early June (Figures 2h). Slightly larger enhancements are visible at higher 458 altitudes (540–620 K) from mid-April to mid-June (Figures 3c, 3j), but none stand out 459 against the considerable dynamically driven year-to-year variability. 460

ClO at 500 K closely follows climatological mean evolution for the first few months, 461 but it starts to climb steeply upward along with aerosol (but before water vapor) in early 462 May (Figure 21; the timing of the onset of the increases in ClO, aerosol, and water va-463 por can also be compared in Figure S2 in Supporting Information). Modest ClO enhancements of 10–20 pptv are seen on isentropic surfaces as low as 440 K and at least as high 465 as 660 K, with peak positive anomalies approaching 40 pptv in early August at 500 K (Fig-466 ures 3d, 3k). Non-negligible CIO anomalies (positive and negative) are also evident at 467 higher altitudes, but MLS frequently records comparable vertically extensive features (Fig-468 ure 3d); they likely arise through descent of changes in the mixing ratios at the secondary 469

peak in the ClO profile at around 1200 K in the upper stratosphere (not shown). Thus 470 we focus on the ClO anomalies at 660 K and below. Given that those anomalies are sig-471 nificant at the $>2\sigma$ level for several months over a range of altitudes (Figure 3k), there 472 is little doubt that they reflect real atmospheric features and not merely MLS measure-473 ment noise. Notable though they are, however, these enhancements are still substantially 474 weaker than the maximum ClO perturbation of ~ 80 pptv seen in the same EqL band 475 in 2020 in the wake of the ANY fires (Figures 2l, 3d) and attributed to heterogeneous 476 chlorine activation on smoke particles (Santee et al., 2022; Solomon et al., 2023; Stra-477 han et al., 2022). Indeed, the 2022 CIO enhancements are not especially distinctive ex-478 cept near the peak in the anomaly profile at 500 K, where they lie at or above the range 479 of variability observed over 2005–2019 from June to September and even exceed those 480 in 2020 during August (Figure 21). Consistent with the moderate increase in ClO, as dis-481 cussed in Section 3.3, $ClONO_2$ shows small enhancements during the intervals of ACE-FTS 482 sampling of this EqL band from June to August (Figure 2m). 483

Although HCl starts off quite a bit lower than normal, the suppressed values in early 484 January 2022 reflect the continuation of a pre-existing negative anomaly that began in 485 2020 and persisted through 2021 (Figures 2n, 3e). The enduring below-average HCl abun-486 dances at the turn of the year may reflect a combination of transport effects and resid-487 ual chlorine repartitioning after ANY (since aerosol, water vapor, and ClO also remained 488 slightly perturbed through most of 2021; Figures 2a, 2b, 2l), but in any case they pre-489 date the eruption and could not have been caused by it. HCl increases fairly steadily for 490 the first few weeks of 2022, reaching average values by mid-February. Thereafter, HCl 491 generally follows the typical pattern of behavior until starting to drop sharply at most 492 levels in mid-May (Figure 2n; also seen in Figures 3e, 3l). The downturn in HCl over May 493 to August 2022 roughly parallels the much deeper ANY-induced depletion in HCl that 494 occurred in earlier months in 2020 (Figure 2n). ACE-FTS indicates a very similar deficit 495 in HCl starting in July and persisting through the end of 2022 (not shown). At face value, 496 the significant concurrent but opposing changes in ClO and HCl point to the occurrence 497 of substantial chemical processing. The story is not so simple, however. 498

At about the same time as the changes in the chlorine species, rather than increas-499 ing as would be expected from R1-R3 (see Section 3.1), HNO₃ rapidly decreases (Fig-500 ure 2h), falling to very low values from July onward that at some levels redefine the bot-501 tom of the MLS mission envelope (not shown). This behavior is at odds with the changes 502 in N_2O_5 , whose values fall to near (or even below) zero, well outside the range previously 503 observed during the ACE-FTS sampling intervals throughout the second half of the year 504 (Figure 2f), and NO_x , which also sets new lower limits in October and November (Fig-505 ure 2g). The lifetimes of N_2O_5 and NO_x are sufficiently short that their anomalies are 506 almost completely driven by chemical perturbations, whereas HNO_3 is also subject to 507 dynamical control. The abrupt shift in HNO_3 , evident over a broad vertical range in the 508 lower and middle stratosphere (Figure 3c), is mirrored by corresponding rapid increases 509 in the transport tracer N_2O (Figures 2d, 3f). Scatter plots (Figure 4, left column) show 510 reasonably tight anticorrelation between HNO₃ and N₂O through much of the lower strato-511 sphere, including in 2022. At the levels where the mild positive HNO_3 anomaly is seen 512 from mid-April to mid-June (540–620 K), two distinct clusters—one in the middle of the 513 distribution from the early months of 2022, the other forming an extreme tail in the dis-514 tribution later in the year—are linked by the points undergoing a steep decline in HNO_3 515 and growth in N_2O in July. But at no time during the year do the 2022 values stray dra-516 matically from the historical HNO_3/N_2O relationship, indicating that the observed vari-517 ations in HNO₃ in the SH midlatitudes are governed to a large extent by transport pro-518 cesses. The levels that show anomalously low values in the latter part of 2022 fall near 519 or below below the peak in the HNO_3 profile, which is generally situated at around 600 K 520 in the SH (e.g., Santee et al., 2004), while at potential temperatures above 700 K, HNO₃ 521 anomalies are slightly positive at that time (Figures 3c, 3j). These signatures are con-522

sistent with the markedly weaker diabatic descent and stronger poleward flow in the SH midlatitudes in the aftermath of HT-HH reported by Coy et al. (2022).

To help diagnose chemical effects, we approximate odd nitrogen, NO_y , as $2 \times N_2O_5$ 525 + NO_x + HNO₃ + ClONO₂, with HNO₃ taken from MLS and the others from ACE-FTS; 526 together, these species account for about 97% of the stratospheric NO_y budget (e.g., Berthet 527 et al., 2017). While the ACE-FTS (averages of sunrise and sunset occultations) and MLS 528 measurements of these diurnally varying species are obtained at different local times, com-529 plicating precise quantification of NO_y , our purpose here is merely to qualitatively com-530 pare the behavior observed in 2022 with that in prior years. The ratios of N_2O_5 and NO_x 531 to NO_y (Figures 2i, 2j) confirm that stratospheric nitrogen partitioning is shifted away 532 from those species in the latter half of 2022. In contrast, HNO_3 makes up an anomalously 533 large fraction of NO_{u} from June onward, redefining the previously observed range in early 534 October and November (Figure 2k). These results suggest that heterogeneous HNO_3 pro-535 duction does take place in this EqL band in the months following HT-HH, but the sig-536 nature of such processing in Figures 2h and 3c is overwhelmed by large countervailing 537 transport effects. 538

The evolution of HNO_3 in the SH midlatitudes in 2022 differs from that observed 539 by the predecessor MLS instrument on the Upper Atmosphere Research satellite (UARS). 540 launched in September 1991, a few months after the June eruption of Mt. Pinatubo. UARS 541 $MLS HNO_3$ retrievals in the equatorial region suffered from contamination by strongly 542 volcanically enhanced SO_2 for the first ~100 days of the mission, but retrievals at higher 543 latitudes were unaffected by this artifact. UARS MLS first observed middle and high south-544 ern latitudes in November, by which time HNO₃ abundances were elevated throughout 545 a broad swath of the hemisphere, with values at 585 K well outside the range measured 546 by UARS MLS at similar times and locations during the rest of the mission (Santee et 547 al., 2004). HNO₃ mixing ratios did not return to normal at the highest EqLs until the 548 following June (1992), and the enhancement lingered for several months longer at mid-549 latitudes, in line with residual volcanic aerosol there. Following Pinatubo, heating from 550 absorption of longwave radiation by the volcanic aerosol altered the radiative balance 551 and dynamics of the stratosphere, intensifying both the upwelling in the tropics and the 552 downwelling in the SH extratropics (Aquila et al., 2012, 2013). This aerosol-induced per-553 turbation in the mean circulation caused a positive anomaly in ozone in the SH midlat-554 itudes (Aquila et al., 2013) and presumably had a similar effect on HNO₃, adding to any 555 chemical enhancement that may have occurred. In contrast, as noted earlier, the radia-556 tive perturbation from the exceptional water vapor injection by HT-HH counteracted 557 that from aerosol, resulting in net longwave cooling and weaker extratropical descent. 558 The contrasting circulation changes likely account to a large extent for the apparently 559 different HNO₃ responses to the two eruptions. 560

The strong influence of dynamics on HNO_3 in the second half of 2022 calls into ques-561 tion a primary chemical provenance for the steep mid-year dive in HCl. As with HNO_3 , 562 the rapid decrease in HCl coincides with an increase in N_2O (cf. Figures 2n and 2d). Scat-563 ter plots for HCl also show a bimodal structure in the lower stratosphere in 2022, with 564 two discrete clumps linked by the points from July (Figure 4, middle column). In the 565 case of HCl, however, notable deviations from the overall climatological HCl/N_2O re-566 lationship are seen at some levels. The conspicuous departure from anticorrelation in the 567 grey dots at 460–620 K is a signature of heterogeneous chlorine activation on smoke par-568 ticles from ANY (Santee et al., 2022). While behavior as clearly anomalous as that in 569 2020 is not evident in 2022, points from July / August onward at 460-660 K (the domain 570 over which ClO displays enhancements) do fall in positions that were previously unoc-571 cupied in the distribution to a much greater extent than they did for HNO_3 . To help dis-572 entangle dynamical and chemical effects, we define inorganic chlorine, Cl_y , as $ClO + ClONO_2$ 573 + HCl, with ClO and HCl from MLS and ClONO₂ from ACE-FTS. Stratospheric trans-574 port is known to play a key role in controlling the Cl_{y} distribution (Strahan et al., 2014). 575

The ratios of ClO and ClONO₂ to Cl_y (Figures 20, 2p) confirm that stratospheric chlorine partitioning is shifted toward those species, especially ClO, in the second half of 2022. The HCl/Cl_y ratios, on the other hand, reveal an anomalously small contribution to inorganic chlorine from HCl in early June, July, and August (Figure 2q). We surmise that, although the pronounced reduction in SH midlatitude HCl in the latter part of 2022 is chiefly governed by transport, HCl abundances are also being suppressed by chemical processing.

To obtain a qualitative sense of the onset of anomalous post-HT-HH partitioning 583 via reactions R1–R5 at 500 K, where the observed perturbations in the chlorine species 584 in this EqL band are largest, Figure 5 compares their respective first-order rate constants 585 in 2022 to those in 2021. The rate constant for a heterogeneous reaction is proportional 586 to its reaction probability (γ value, Figure S1 in Supporting Information), SAD, and the 587 thermal velocity of the molecule. The calculations in Figure 5 are based on the water 588 vapor and temperature values shown in Figure 2, but they use constant abundances for 589 the other gas-phase species as well as a fixed SAD of $2\,\mu\text{m}^2\,\text{cm}^{-3}$, representing background 590 conditions. Thus, these results are intended to illuminate only the influence of water va-591 por and temperature on R1–R5; they take no account of the enhanced aerosol loading, 592 the true chlorine species mixing ratios, or the evolution of those quantities over the year. 593 While using an enhanced SAD value for 2022 would more accurately characterize post-594 HT-HH conditions, doing so would affect the rate constants for R1–R5 proportionally 595 and thus would reveal no additional information about the relative effectiveness of those 596 reactions beyond that provided in Figure 5. Calculation of actual reaction rates, and hence 597 full quantification of the individual contributions of R1–R5 to the observed repartition-598 ing, requires detailed chemical modeling beyond the scope of this analysis. Nevertheless, 599 Figure 5 provides useful insight. The rate constants for R2–R4 are slightly higher in 2022 600 than in the preceding year from late July onward, not dissimilar to the timing of the per-601 turbations in ClO and HCl (i.e., the HCl/Cl_y ratio). This might suggest a role for those 602 reactions. However, their rate constants are still several orders of magnitude smaller than 603 those for R1 and R5. Taking the reciprocal of the maximum 2022 rate constants (which 604 are in units of s^{-1}) in Figure 5 yields approximate chemical lifetimes for the gas-phase 605 reactants of 1 day for R1, 10 months for R2, 60 years for R3, 40 years for R4, and 5 hours 606 for R5. Our results are in line with the modeling work by Robrecht et al. (2019), who 607 demonstrated that R2 and R3 (and by implication also R4) do not become effective un-608 til temperatures fall below 205 K even for very high water vapor mixing ratios of 20 ppmy 609 and volcanic conditions of $10 \times$ background SAD. Although at its peak the EqL-mean 610 aerosol cross section in this region is enhanced by about a factor of 20, the maximum 611 water vapor values are just over 5 ppmv, and minimum temperatures are $\sim 210 \,\mathrm{K}$. We 612 conclude that, despite the heavy aerosol loading, excess moisture, and strong cooling in 613 the 2022 SH midlatitudes following HT-HH, ambient conditions remain unfavorable for 614 R2–R4, and only R1 and R5 are at play in producing the observed moderate chlorine 615 activation. This is in contrast to the situation following ANY in 2020, when the high sol-616 ubility of HCl in aged stratospheric smoke particles apparently triggered heterogeneous 617 chlorine activation, in particular via R3, under relatively warm conditions in the SH mid-618 latitudes (Solomon et al., 2023). 619

Finally, an obvious question is whether the mild enhancement in reactive chlorine 620 in 2022 had a discernible impact on ozone concentrations. As noted by Santee et al. (2022), 621 although the strong and sustained midlatitude chlorine activation that occurred in the 622 wake of ANY was unprecedented in the satellite record, it was still an order of magni-623 tude or more weaker than that in a typical winter polar vortex. They found no conclu-624 sive observational evidence of significant ANY-induced chemical ozone destruction; rather, 625 they attributed the anomalously low ozone observed in the SH midlatitudes in 2020 (and 626 until midway through 2021) largely to transport effects, as did Strahan et al. (2022). In 627 contrast, other studies argued that heterogeneous chemistry on ANY smoke particles did 628 give rise to detectable ozone loss (e.g., Rieger et al., 2021; Solomon et al., 2022, 2023; 629

Yu et al., 2021). Given that the degree of chlorine activation in 2022 is about half that following ANY and the large-scale dynamical perturbations are far greater, we expect the evidence of chemical ozone destruction from HT-HH to be even more equivocal.

Figure 2e shows that the behavior of ozone in the SH midlatitude lower stratosphere 633 in 2022 is not dissimilar to that in 2020, with ozone mixing ratios generally low but not 634 outside the range of previously observed variability. Below 600 K, the changes in ozone 635 track those in HNO_3 and HCl; above that level, ozone becomes positively correlated with 636 N_2O in this EqL band and increases in the second half of the year (Figures 1g, 3g, 3n). 637 Although the $2022 O_3/N_2O$ correlations show two distinct populations in the early and 638 late months of the year at lower potential temperatures, as they do also for HNO_3 and 639 HCl, for the most part the points do not lie outside the typical transport-controlled dis-640 tribution (Figure 4, right column). Thus MLS measurements indicate that the minor in-641 creases in ClO provoked by HT-HH do not result in appreciable chemical ozone loss in 642 the SH midlatitude lower stratosphere in 2022. 643

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4.2 Southern Hemisphere Subtropics (22°S–38°S EqL)

The evolution in the SH subtropics at 620 K is similar to that at midlatitudes and 645 is summarized only briefly. Aerosol increases rapidly almost immediately after the erup-646 tion, reaching maximum values between April (620-660 K) and August (460-500 K) that 647 are larger at a given level than those at more poleward EqLs (Figure 6a; Figures S3a, 648 S3h in Supporting Information). HNO_3 ticks upward in late January above 560 K (Fig-649 ure 6h; Figures S3c, S3j in Supporting Information). This positive anomaly, though still 650 small, starts earlier, achieves a larger magnitude (>0.6 ppbv at 580–660 K in April), and 651 extends to higher altitudes than seen in the $38^{\circ}S-54^{\circ}S$ EqL band. Maximum values pro-652 trude above the mission envelope (Figure 6h), albeit only marginally and for a short in-653 terval in some cases. HNO_3 values continue to slowly rise in subsequent months at the 654 highest altitudes (Figure S3j in Supporting Information). At lower levels, however, they 655 decline, with a particularly precipitous drop in May–June over 540–700 K, and then re-656 main anomalously low through the end of the year. The downturn starts sooner but is 657 less abrupt than that seen at midlatitudes, and for the most part HNO_3 values do not 658 dip below the MLS mission envelope. Although the sparser sampling of ACE-FTS in this 659 EqL band compromises identification of anomalies, slightly depressed N_2O_5 (Figures 6f, 660 6i) and NO_x (Figures 6g, 6j) in later months again suggest repartitioning within the odd 661 nitrogen family, and elevated HNO_3/NO_y ratios (Figure 6k) are consistent with some 662 chemical production of HNO₃. 663

HCl follows the climatological mean for the first few months of 2022 before falling 664 fairly steeply to values lower than any previously observed above $550 \,\mathrm{K}$ (Figure 6n); be-665 low that level, the record HCl deficits in 2020 arising from ANY still stand. The decrease 666 in HCl begins about a month earlier than in the $38^{\circ}S-54^{\circ}S$ EqL band. As before, scat-667 ter plots (Figure S4 in Supporting Information, left and middle columns) show two sep-668 arated clusters in 2022 in much of the lower stratosphere, connected by points with rapidly 669 decreasing HNO₃ and HCl but increasing N₂O, in this case during May–June. Here, how-670 ever, above 460 K the 2022 HCl values lie in previously unpopulated areas of the distri-671 bution to a much greater extent than at midlatitudes. In addition, the HCl/Cl_{y} ratios 672 are indicative of chlorine repartitioning from April onward (Figure 6q). ClO shows an 673 even more prominent signature of chemical processing, with significant positive anoma-674 lies appearing over 460–700 K as early as March and reaching maxima of as much as 30– 675 35 pptv in May–July, depending on the level, as the peak in the ClO anomaly profile grad-676 ually descends in concert with the downward progression of the sulfate aerosol (Figures S3d, 677 S3k in Supporting Information). At most levels, the 2022 ClO abundances sit near or 678 above the top of the mission envelope for a few months (e.g., Figure 61); at and above 679 620 K, they exceed the ANY enhancements at all times, while at lower levels they are 680 initially smaller than 2020 values but then stay elevated through the end of the year (Fig-681

⁶⁶² ure S3d in Supporting Information). The timing of the buildup, peak values, and decay
⁶⁶³ of the ClO anomaly coincides closely with that of the aerosol anomaly but generally pre⁶⁶⁴ cedes that of the water vapor anomaly (Figure S2 in Supporting Information). Although
⁶⁶⁵ the general picture is similar to that in the SH midlatitudes, the relationship between
⁶⁶⁶ active chlorine and aerosol cross section is clearer in this EqL band.

As with HNO₃ and HCl, ozone mixing ratios briefly dip below the climatological mean at 620 K in June–July, mirroring the bump up in N₂O; thereafter, ozone rises steadily until it skirts the top of the observed range from October onward (Figure 6e). None of the analyses, including the scatter plots in Figure S4 in Supporting Information (right column), show patterns indicative of significant chemical ozone loss in 2022.

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4.3 Southern Hemisphere Tropics (6°S–22°S EqL)

In the SH tropics, aerosol (Figures 7a, 8a, 8h) and water vapor (Figures 7b, 8b, 8i) are abruptly enhanced through much of the lower stratosphere immediately after the eruption. At the levels of interest here, the choice of aerosol dataset used makes no material difference to these results. A warming trend throughout the stratosphere in this EqL band prior to the event is reversed, with temperatures dropping by $\sim 2 \text{ K}$ during February (Figure 7c; see also Schoeberl et al., 2022).

Concomitant with these changes, HNO₃ increases rapidly above 500 K (Figures 8c, 699 8j), overshooting the previously observed range by early February, with values then near 700 or above the top of the envelope through the first half of the year (Figure 7h). Large anoma-701 lies of $\sim 0.7-0.8$ ppbv are seen from 580–800 K (Figures 8j). The HNO₃/N₂O scatter plots 702 (Figure 9, left column) show that the March–July points lie at the outer edge of or slightly 703 apart from the main distribution at most levels, suggesting that transport is not the sole 704 factor controlling HNO_3 abundances at this time; rather, the observed behavior is in-705 dicative of heterogeneous processing. This determination is substantiated by the very 706 low N_2O_5 (Figures 7f, 7i) and NO_x (Figures 7g, 7j) values, as well as the greatly elevated 707 HNO_3/NO_4 ratios (Figure 7k) throughout 2022. Although the picture is unfortunately 708 fragmentary because of the poor ACE-FTS sampling at low latitudes, the 2022 values 709 of all of these quantities are clearly highly anomalous. 710

After rising steadily for a few weeks, however, HNO₃ mixing ratios then level off 711 and remain approximately flat until June, when they start to increase again, largely fol-712 lowing the typical seasonal pattern (Figure 7h). We interpret the March–May plateau 713 in HNO₃ as evidence of the saturation of N_2O_5 hydrolysis. The amount of N_2O_5 avail-714 able to participate in R1 is limited by its formation (mainly from the recombination of 715 NO_2 and NO_3 at night, whose rate depends quadratically on NO_x). When aerosol SAD 716 exceeds a certain threshold, N_2O_5 is hydrolyzed as fast as it can be formed, and further 717 increases in SAD do not affect the rate of conversion of NO_x to HNO_3 by R1 (e.g., Fa-718 hey et al., 1993; Tie et al., 1994). Although ACE-FTS coverage in the tropics is too sparse 719 to permit diagnosis of saturation in the NO_x data, this mechanism may explain why HNO_3 720 production appears to stall by early March at many levels (Figure 8j), while aerosol (Fig-721 ure 8h) and water vapor (Figure 8i) are still increasing. It may also account for the con-722 tinuous gradual increase in HNO_3 in April at higher potential temperatures (660–850 K), 723 as well as the lack of strong enhancement at lower potential temperatures $(460, 500 \, \text{K})$ 724 despite the presence of substantial aerosol there, since saturation occurs for relatively 725 lower aerosol amounts at lower altitudes (e.g., Mills et al., 1993). Similarly, Santee et al. 726 (2004) reported considerably greater impact from Pinatubo on HNO₃ abundances at higher 727 (585 K) than at lower (420, 465 K) potential temperatures based on UARS MLS mea-728 surements and attributed the differences to R1 saturation. 729

The evolution of ClO is also consistent with substantial sustained heterogeneous processing in this EqL band in 2022. Significant enhancements appear by mid-January from 500 to 700 K, with maxima as large as 40–45 pptv at 580–660 K in March (Figure 8k),

far exceeding the previously observed range of behavior (Figures 71, 8d; see also Figure 1d). 733 The enhancement slowly decays after April, but mixing ratios remain elevated over much 734 of the domain through the end of the year. A secondary MLS ClO product retrieved from 735 radiances in a different spectral band than that used in generating the standard ClO data 736 indicates a similar (albeit noisier) persistent strong enhancement (not shown). $CIONO_2$ 737 is also high during the brief windows of ACE-FTS sampling through most of the year 738 (Figures 7m, 7p). As with HNO₃, after an initial steep rise, ClO values then remain more 739 or less steady for the next several months (Figures 7l, 8k). A similar nonlinear response 740 to increasing aerosol SAD seen in airborne in situ ClO measurements after the eruption 741 of Mt. Pinatubo was attributed to the saturation of N_2O_5 hydrolysis on sulfate aerosols 742 (Avallone et al., 1993; Fahey et al., 1993). The aircraft campaigns also found the largest 743 ClO enhancements at the lowest latitudes sampled (Avallone et al., 1993). Thus, our re-744 sults are qualitatively in accord with the earlier in situ findings. In addition, the MLS-745 based CIO enhancements are comparable to post-eruption values simulated in several 746 modeling studies (e.g., Kinnison et al., 1994; Tie et al., 1994; Zambri et al., 2019). On 747 the other hand, Zambri et al. (2019) also examined MLS ClO measurements after mul-748 tiple moderate-magnitude eruptions in the post-Pinatubo era (see Section 3.2) and found 749 no enhancements matching those modeled, possibly because the HT-HH impact on strato-750 spheric chlorine activation in the SH tropics greatly surpasses that of any prior event. 751

In the immediate aftermath of the HT-HH eruption, as HNO_3 and ClO shoot up, 752 HCl displays a modest (<0.08 ppbv) but fairly sharp decrease over a limited altitude range 753 (540–660 K; Figure 81). This dip in January, which is not seen at higher latitudes, places 754 the 2022 HCl values at or near the bottom of the envelope at those altitudes in Febru-755 ary and March (Figure 7n). Given that N_2O is generally low at this level at this time 756 in 2022 (Figures 7d, 8f), correspondingly high HCl mixing ratios would have been ex-757 pected. After its initial decrease, HCl follows climatological patterns (i.e., deviations from 758 the mean remain essentially flat) for a few months, echoing the plateau in ClO (Figures 7n, 759 81) until dynamically driven changes—negative anomalies below 660 K, positive anoma-760 lies above (Figures 8e, 8l)—develop in the latter part of 2022. Although the HCl/N_2O 761 correlations are not especially unusual for the first half of the year, from about July on-762 ward they stand apart from the typical relationships at and above 620 K (Figure 9, mid-763 dle column). This behavior likely signifies continuing mid-year chemical suppression of 764 HCl, in consonance with the evolution of ClO. The extremely low HCl/Cl_{y} ratios seen 765 during the intervals of ACE-FTS coverage (Figure 7q) confirm the depletion of HCl through 766 heterogeneous chemical processing. 767

In the 620–660 K layer, close to the altitude of the maximum HT-HH water vapor 768 injection (Figure 8i; Millán et al., 2022), the steep increases in HNO₃ and ClO and the 769 decrease in HCl in the first weeks after the eruption occur nearly simultaneously with 770 the pulses in both water vapor and aerosol (Figure S2 in Supporting Information). Be-771 low that layer, however, HNO_3 , CIO, and HCl change more or less in unison with the 772 increase in water vapor, while the aerosol curves slope upward more gently and peak weeks 773 to months later. This behavior in the SH tropics differs from that in the subtropics and 774 midlatitudes, where, as discussed in the previous section, ClO enhancement generally 775 tracks the evolution of aerosol more closely than that of water vapor at altitudes below 776 600 K. Climate model simulations have shown that the extraordinary magnitude of the 777 HT-HH water vapor anomaly halved the SO₂ lifetime, leading to more rapid coagula-778 tion of larger sulfate particles than is typical (Zhu et al., 2022). The slower buildup in 779 aerosol at lower potential temperatures in the tropics may reflect the longer time required 780 for sulfate formation at less water-rich levels and/or the timescale for sedimenting par-781 ticles from above to reach those levels. The observed latitudinal differences in the rela-782 tionships between aerosol, water vapor, and reactive chlorine may arise because, by the 783 time a substantial portion of the HT-HH plume disperses to the SH subtropics, the con-784 version to sulfate aerosol is well underway. Nevertheless, sufficient sulfate aerosol to en-785 able heterogeneous processing is clearly present soon after the eruption even at lower al-786

titudes in the tropics, as evidenced by the rapid rise in HNO₃ and ClO there (Figures 8j,
8k; Figure S2 in Supporting Information).

In contrast to R1, reactions R2 and R5 are not subject to saturation in the lower 789 stratosphere. Thus, the fact that HNO_3 and CIO do not continue to increase in tandem 790 with aerosol implies that R1 is playing a dominant role in their production. Moreover, 791 a larger decrease in HCl, as well as a decrease (rather than an increase) in $CIONO_2$, would 792 be expected if direct heterogeneous chlorine activation were proceeding. The results of 793 our rate constant calculations (Figure 10) support this supposition. Although the ambient (temperature, water vapor) conditions in 2022 increase the rate constants for R2– 795 R4 considerably over their respective 2021 values starting in late January / early Febru-796 ary, they remain several orders of magnitude smaller than those for R1 and R5. There-797 fore, we conclude that, as exceptional as the aerosol and water vapor enhancements in 798 this EqL band are, they are not sufficient to promote R2–R4 to any significant extent 799 under the colder than normal but still relatively (compared to polar winter) warm con-800 ditions that prevail in the first half of 2022 (Figure 7c). 801

Figures 5 and 10 show that the rate constant for R5 is as much as 5-6 times faster 802 than that for R1. Photolysis of the HOBr and HNO₃ resulting from BrONO₂ hydrol-803 ysis is an important source of OH, rivalling that from nitrogen chemistry under condi-804 tions of volcanically enhanced aerosol; indeed, about half of the simulated changes in strato-805 spheric chlorine species after the Sarychev eruption were attributed to elevated OH abun-806 dances from R5 (Berthet et al., 2017). In addition to $ClONO_2$ and $BrONO_2$ hydroly-807 sis as a source of OH, the excess water vapor from HT-HH likely amplifies direct OH for-808 mation from H_2O oxidation; model simulations by Zhu et al. (2022) show increases in 809 OH in the SH tropics immediately following the eruption compared to a run with no wa-810 ter vapor injection. Unfortunately, MLS no longer measures OH itself, but measurements 811 of HO₂ (the other main member of the HO_x family) are available from MLS in the mid-812 dle stratosphere; those data indicate some enhancement in early 2022 (Figure S5 in Sup-813 porting Information). As noted above, the small decreasing tendency in HCl levels off 814 at approximately the same time as the increases in HNO_3 and ClO; this behavior may 815 point to saturation and hence the influence of enhanced OH arising from R1, rather than 816 from R5 or from the additional water vapor from HT-HH. Again, full quantification of 817 the effectiveness of the different processes contributing to OH production (and hence chlo-818 rine repartitioning) requires detailed chemical modeling beyond the scope of this study. 819

Finally, although heterogeneously driven chemical loss typically plays little or no 820 role in determining ozone abundances in the tropics, we investigate whether the unprece-821 dented degree of chlorine activation in the 500–700 K layer following HT-HH leads to per-822 ceptible reductions in ozone. At 620 K, ozone displays substantial variability through-823 out 2022, rising and falling fairly rapidly at various times and occasionally redefining the 824 top of the mission envelope (Figure 7e). Of particular note is the steep decline in March 825 and early April, which coincides with the maximum enhancement in ClO (Figure 71) and 826 thus could be a sign of chemical loss. However, ozone mixing ratios at this level barely 827 fall below the climatological mean before beginning to increase again at the end of May, 828 and surrounding levels also see only small decreases (Figure 8n). Ozone remains largely 829 anticorrelated with N_2O in the tropics at this altitude (cf. Figures 8f and 8g), and the 830 O_3/N_2O scatter plots show no hint of abnormally low O_3 values suggestive of depletion 831 at any point during the year (Figure 9, right column). Therefore, any signature of chlorine-832 catalyzed ozone loss that may be taking place in the lower stratosphere is masked by much 833 larger changes due to transport. On the other hand, points from the latter half of 2022 834 fall well outside the previously observed O_3/N_2O distribution at 660 K (and higher lev-835 els; not shown). These anomalously high ozone values may reflect diminished efficiency 836 of the NO_x catalytic loss cycle, which dominates ozone destruction in the middle strato-837 sphere (see Section 3.4). 838

5 Summary and Conclusions

The eruption of HT-HH in January 2022 caused a substantial perturbation in strato-840 spheric aerosol loading, which, although not unprecedented in the modern instrumen-841 tal era, was nevertheless the largest in the last several decades. Even more noteworthy 842 was the uniquely extreme enhancement in stratospheric water vapor. The submarine vent 843 erupted at a "Goldilocks" depth—sufficiently shallow that the pressure of the overlying 844 water did not dampen the power of the blast, sufficiently deep to allow extensive magma-845 seawater interaction (Witze, 2022). In addition, the plume was injected at an advanta-846 geous location, as lofting into lower altitudes or higher latitudes where air is colder would 847 have resulted in smaller water vapor anomalies (Schoeberl et al., 2022). The excess mois-848 ture led to strong stratospheric cooling that in turn induced anomalous circulation pat-849 terns. 850

In this study, we use constituent profile measurements from Aura MLS and ACE-FTS, 851 aerosol extinction from OMPS-LP, and meteorological parameters from MERRA-2 to 852 investigate the extent to which the dense aerosol, high humidity, and strong cooling brought 853 about by HT-HH prompted heterogeneous chemical reactions to occur in regions where 854 they are not normally active to a significant degree. We look for signatures of chlorine 855 and nitrogen repartitioning in the SH midlatitudes, subtropics, and tropics over a range 856 of altitudes in the lower and middle stratosphere. Calculated rate constants for the het-857 erogeneous reactions R1–R5 provide insights into their relative importance in altering 858 trace gas distributions. Ozone is also examined for evidence of chemical loss as a result 859 of the eruption. 860

In the SH midlatitudes ($38^{\circ}S-54^{\circ}S$ EqL), N₂O₅ and NO_x exhibit considerable de-861 creases at 500 K in the second half of 2022 that are consistent with heterogeneous pro-862 cessing on HT-HH aerosol. Although enhancements in HNO_3 would be expected in association with the reductions in N_2O_5 and NO_x , strong dynamical effects instead lead 864 to unusually low HNO_3 values that in some cases redefine the previously observed range, 865 and HNO₃ production only becomes apparent in the HNO_3/NO_y ratios. Signs of het-866 erogeneous chlorine activation are more compelling, with a positive anomaly in ClO reach-867 ing 35–40 pptv at its peak near 500 K from June through August, accompanied by an 868 increase in ClONO₂. Although significant at the $>2\sigma$ level, however, the ClO anomaly 869 is only marginally outside the envelope of previous behavior over a narrow layer for a 870 few months. Transport effects dominate the evolution of HCl in the latter part of 2022, 871 but chemically induced deficits in HCl that mirror the enhancements in ClO and ClONO₂ 872 are evident in the HCl/Cl_{y} ratios. Perturbations in all chlorine species are considerably 873 weaker than those measured at these levels in this EqL band in 2020 in the aftermath 874 of the ANY fires. Rate constant calculations suggest that, despite the unusually moist 875 and cool conditions in the SH midlatitude stratosphere following HT-HH, heterogeneous 876 chlorine activation reactions R2–R4 are not at work to any significant extent, and R1 877 and R5 (hydrolysis of N_2O_5 and BrONO₂, respectively) are primarily responsible for the 878 observed composition changes. Evidence of widespread and persistent HT-HH-induced 879 chlorine and nitrogen repartitioning is even clearer in the SH subtropics $(22^{\circ}S-38^{\circ}S EqL)$, 880 but the signatures of heterogeneous processing in HNO_3 and HCl are again obscured by 881 dynamical influences. 882

In the SH tropics (6°S–22°S EqL), unambiguous signatures of substantial hetero-883 geneous processing appear over a broad vertical domain almost immediately after the 884 eruption. Substantial repartitioning of both the nitrogen and the chlorine families oc-885 curs, as evidenced by depletion of N_2O_5 and NO_x together with enhancement of HNO_3 , 886 ClO, and $ClONO_2$. Unlike at higher latitudes, in this EqL band HCl also decreases rapidly 887 right after the eruption, and, in the second half of 2022, it displays extremely low HCl/Cl_{u} 888 ratios, as well as notable departures from the typical HCl/N_2O correlations. These pat-889 terns all point to significant chemical suppression of HCl. To our knowledge, volcanically 890 induced HCl depletion has not been reported previously. After initially rising steeply, 891

HNO₃ and ClO essentially plateau, maintaining fairly constant anomalies (of as much 892 as 0.7–0.8 ppbv and 40–45 pptv, respectively) for several months. This behavior is con-893 sistent with the saturation of N_2O_5 hydrolysis, suggesting that R1 is the primary mech-894 anism for the production of HNO_3 and, after associated gas-phase chemistry, the increases 895 in ClO and ClONO₂. Enhancements in OH brought about by the excess water vapor from 896 HT-HH may also play a role in the chlorine repartitioning. As at midlatitudes, rate con-897 stant calculations imply that reactions R2-R4 play no significant role in the observed 898 composition changes. Quantification of the relative contributions of the different pro-899 cesses perturbing stratospheric composition in the wake of HT-HH will require detailed 900 modeling efforts. 901

The moderate enhancements in reactive chlorine seen throughout the SH mid- and 902 low-latitude lower stratosphere following HT-HH do not lead to appreciable chemical loss 903 of ozone; rather, lower-stratospheric ozone in those regions remains primarily controlled 904 by transport processes. The lack of chemical ozone depletion in 2022 substantial enough 905 over an area extensive enough to be perceptible in the EqL means examined here is not 906 surprising. Although the 2022 ClO anomalies are unprecedented in the tropics and sub-907 tropics and are exceeded only by those induced by ANY in the midlatitudes, they are 908 still only about half as large as those arising from that event. Moreover, the ClO per-909 turbations caused by ANY were themselves more than an order of magnitude weaker than 910 the enhancements in typical winter polar vortices. In the middle stratosphere, by con-911 trast, reduced NO_x may have decreased the efficiency of that loss cycle, leading to mild 912 increases in ozone at and above 660 K. 913

The strong positive stratospheric aerosol anomaly from HT-HH has already sub-914 stantially abated, and, assuming no further major perturbations, stratospheric aerosol 915 loading is expected to return to pre-eruption levels by around mid-2023 (Khaykin et al., 916 2022). In contrast, stratospheric water vapor abundances remain historically high, with 917 the extreme enhancement likely to endure for several more years (Khavkin et al., 2022) 918 Millán et al., 2022). The HT-HH water vapor plume was effectively excluded from the 919 2022 Antarctic polar vortex and thus had little or no impact on that year's ozone hole; 920 similarly, it did not reach northern high latitudes in time to influence Arctic ozone loss 921 in boreal spring 2023 (Manney et al., 2023). However, the sustained HT-HH water va-922 por enhancement is expected to significantly affect Antarctic and Arctic vortex chem-923 istry and ozone loss during their upcoming 2023 and 2023/2024 winter/spring seasons. 924 Our ability to assess the impact of HT-HH on stratospheric chemistry and polar ozone 925 depletion over the next few years will depend strongly on the continuing availability of 926 satellite observations from Aura MLS and ACE-FTS. 927

6 Data Availability Statement

⁹²⁹ The data sets used here are publicly available, as follows:

930 •	Aura MLS Level 2 data (Lambert, Read, & Livesey, 2020; Manney et al., 2020;
931	Santee et al., 2020; Froidevaux et al., 2020; Lambert, Livesey, & Read, 2020; Schwartz
932	et al., 2020; Wang et al., 2020): https://disc.gsfc.nasa.gov/datasets?page=
933	1&keywords=AURA%20MLS
934 •	Aura MLS Derived Meteorological Products (DMPs; Manney & Millán, 2007–present):
935	https://mls.jpl.nasa.gov/eos-aura-mls/dmp (registration required)
936 •	ACE-FTS 4.1/4.2 data: http://www.ace.uwaterloo.ca (registration required
937	at https://databace.scisat.ca/l2signup.php)
938 •	ACE-FTS v4.1/4.2 error flags (Sheese & Walker, 2020): https://doi.org/10.5683/
939	SP2/BC4ATC
940 •	OMPS-LP L2 Aerosol Extinction Vertical Profile swath multi-wavelength daily
941	3slit Collection 2 V2.0 data (Taha, 2020): https://doi.org/10.5067/CX2B9NW6FI27

 MERRA-2 (Global Modeling and Assimilation Office (GMAO), 2015): https:// disc.sci.gsfc.nasa.gov/uui/datasets?keywords=%22MERRA-2%22

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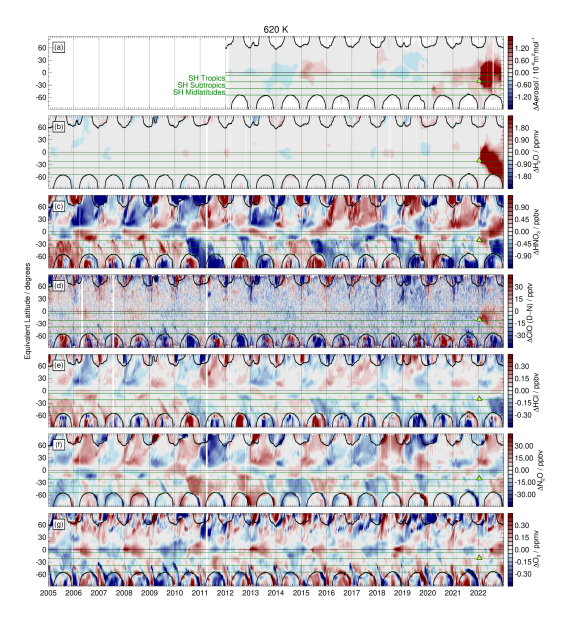


Figure 1. Equivalent latitude (EqL) / time series at 620 K of anomalies (calculated by subtracting from the daily data in each EqL bin the corresponding climatological value over 2005– 2021 for MLS and 2012–2021 for OMPS-LP for that day of the year) in (a) NASA OMPS-LP aerosol cross section at 869 nm and MLS (b) H_2O , (c) HNO_3 , (d) CIO (day minus night), (e) HCl, (f) N₂O, and (g) O₃. To ameliorate a negative drift in the MLS N₂O measurements (see Section 2), the N₂O anomalies (panel (f)) have been "detrended" by removing a linear fit to the daily values across the Aura mission at each EqL/potential temperature gridpoint. Blank spaces represent data gaps. Black overlays identify the approximate boundary of the winter polar vortices (as defined by Lawrence et al., 2018). The black horizontal line marks the equator; green horizontal lines mark the EqLs defining the regions examined in this study: the southern tropics (6°S–22°S), subtropics (22°S–38°S), and midlatitudes (38°S–54°S). The bright green triangle marks the time and latitude of the main HT-HH eruption.

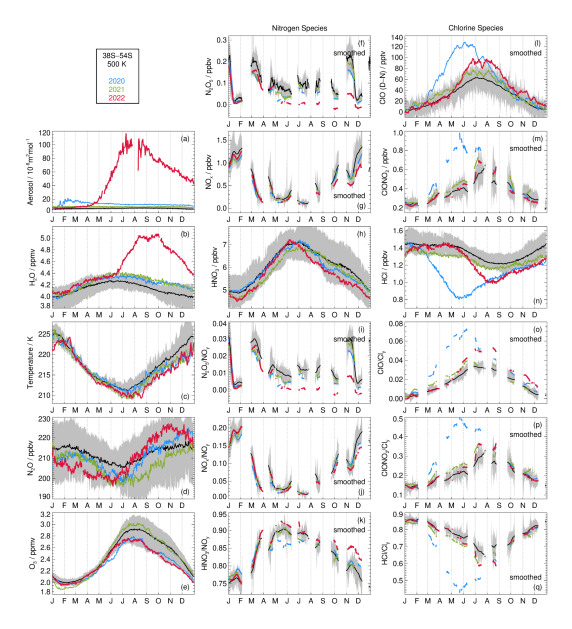


Figure 2. Averages over $38^{\circ}S-54^{\circ}S$ EqL (southern midlatitudes) at 500 K of (a) NASA OMPS-LP aerosol cross section at 869 nm, (b) MLS H₂O, (c) MERRA-2 temperature, (d) MLS N₂O, (e) MLS O₃, (f) ACE-FTS N₂O₅, (g) ACE-FTS NO_x (NO+NO₂), (h) MLS HNO₃, (i) N₂O₅/NO_y (where NO_y=N₂O₅+NO_x+HNO₃+ClONO₂), (j) NO_x/NO_y, (k) HNO₃/NO_y, (l) MLS ClO (day minus night), (m) ACE-FTS ClONO₂, (n) MLS HCl, (o) ClO/Cl_y (where Cl_y=ClO+ClONO₂+HCl), (p) ClONO₂/Cl_y, and (q) HCl/Cl_y. Values for 2022 are shown in red, for 2021 in green, and for 2020 in blue. Grey shading depicts the envelope of behavior and black lines the mean over 2005–2019 (thus excluding the highlighted years). ClO is one of the noisier MLS products; in addition, for this analysis we are taking day minus night differences, so fewer individual points contribute to these EqL-band averages than for the other species measured by MLS. Consequently, the ClO field displays a fairly large degree of day-to-day variability. To enhance clarity, the ClO results (highlighted years and mean, but not the envelope) have been smoothed using a boxcar of width 10 days. Although such smoothing attenuates extreme values, it facilitates detection of persistent anomalies. Similarly, all ACE-FTS fields, as well as all quantities derived using them, have also been smoothed with a 10-day boxcar average.

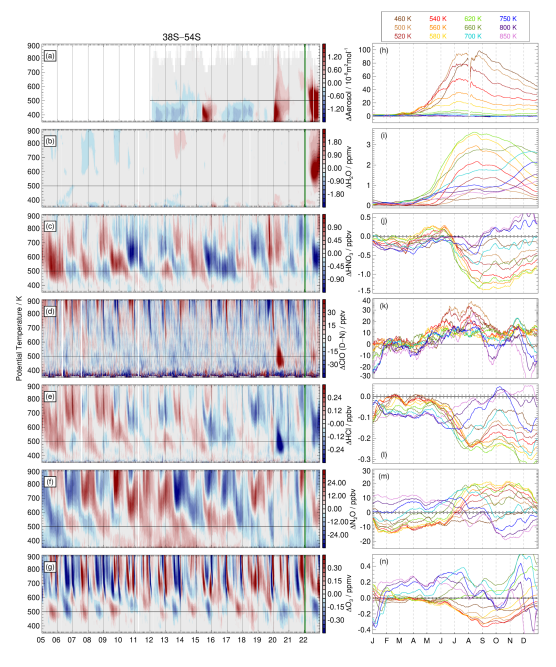


Figure 3. Potential temperature / time sections of anomalies (calculated as in Figure 1) in the 38°S–54°S EqL band of (a) NASA OMPS-LP aerosol cross section at 869 nm and MLS (b) H₂O, (c) HNO₃, (d) ClO (day minus night), (e) HCl, (f) N₂O, and (g) O₃. The vertical green line marks the date of the main HT-HH eruption. Panels (h)–(n) show the 2022 anomalies for each species at multiple potential temperatures (see legend). To enhance clarity, all curves in (h)–(n) except ClO have been smoothed using a boxcar of width 10 days; for ClO the smoothing window is 20 days. This smoothing removes some of the small-scale structure seen in the unsmoothed EqL-band averages shown in Figure 2. Solid black horizontal lines mark zero. Dashed black horizontal lines on the bottom five panels denote the 2σ contributions (± values) of MLS measurement noise to the smoothed EqL-band average anomalies, taken to indicate likely statistical significance for individual points in the anomaly timeseries.

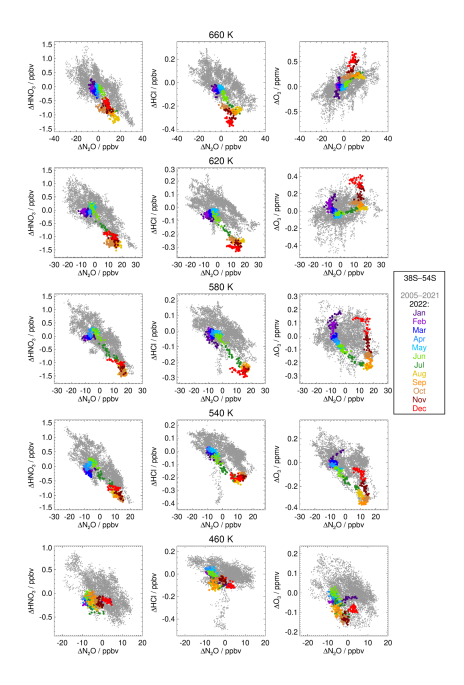


Figure 4. Scatter plots of daily HNO₃ (left), HCl (middle), and O₃ (right) vs. N₂O anomalies in the $38^{\circ}S-54^{\circ}S$ EqL band at various potential temperature levels. Grey dots depict data from 2005–2021; 2022 data are color-coded by month (see legend).

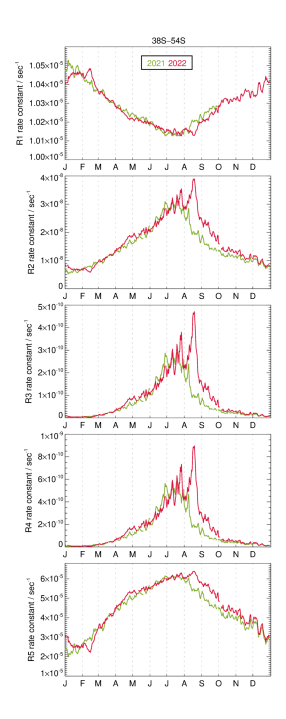


Figure 5. First-order rate constants for reactions R1–R5 in the $38^{\circ}S-54^{\circ}S$ EqL band at 500 K, based on the daily mean temperature and water vapor abundances observed in this region and the reactive uptake coefficients in Figure S1 in Supporting Information, assuming fixed abundances of 2 ppbv HCl, 0.1 ppbv ClONO₂, 0.1 ppbv HOCl, and 0.1 ppbv H₂SO₄ for a mean particle radius of 0.1μ m and an SAD of $2\,\mu$ m² cm⁻³. Values for 2022 are shown in red and for 2021 in green.

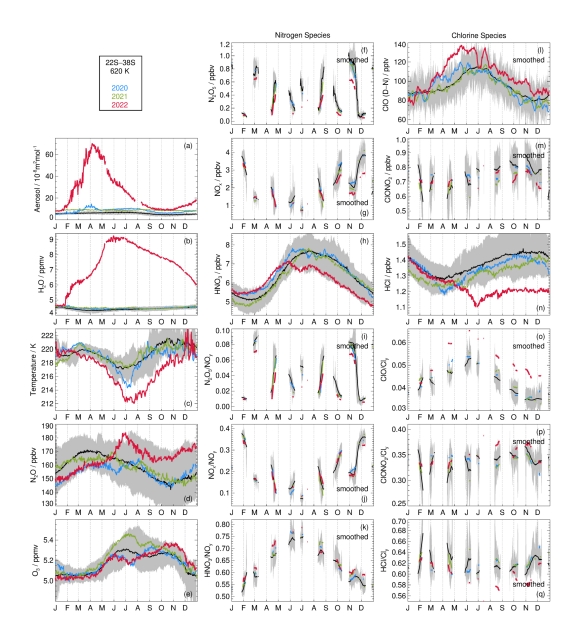


Figure 6. As Figure 2, but for 22°S–38°S EqL (southern subtropics) at 620 K.

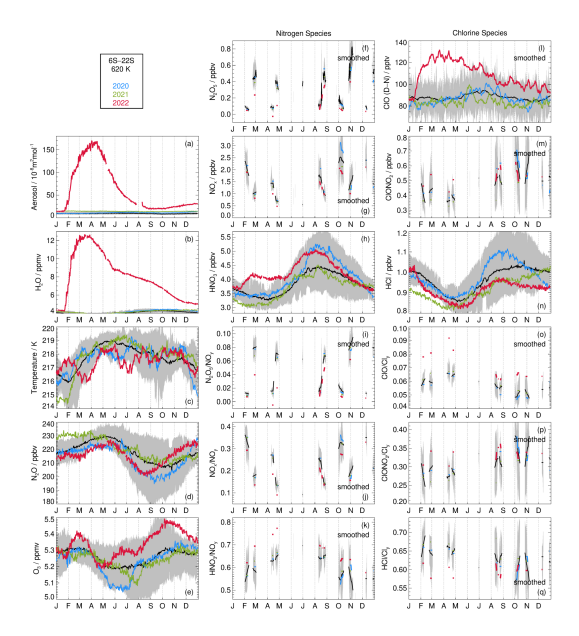


Figure 7. As Figure 2, but for 6°S–22°S EqL (southern tropics) at 620 K.

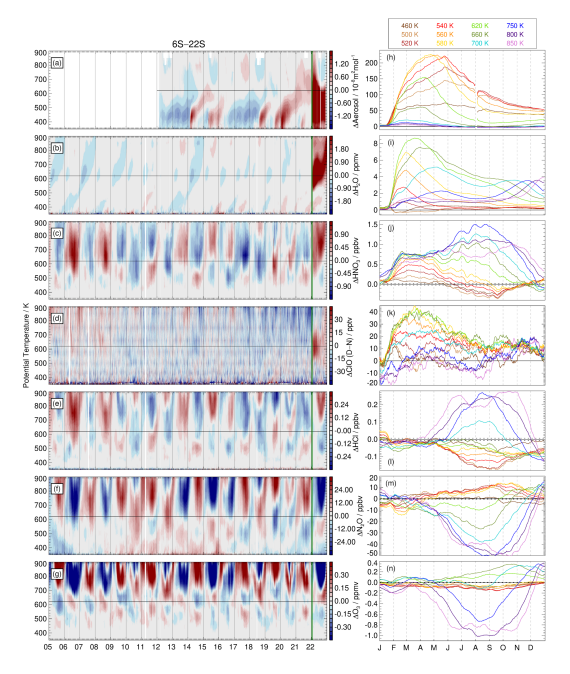


Figure 8. As Figure 3, but for 6°S–22°S EqL.

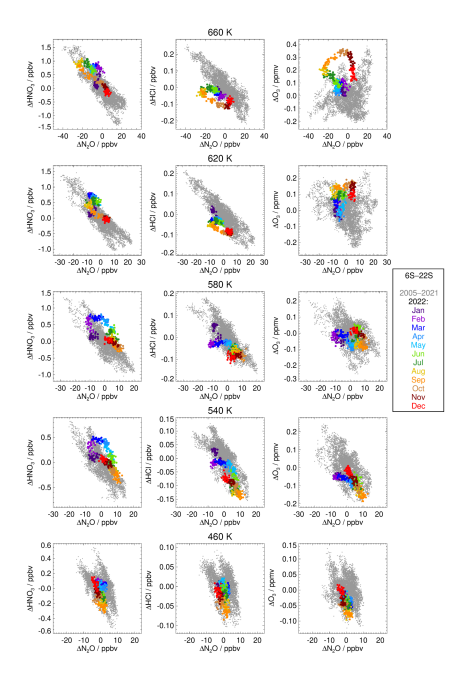


Figure 9. As Figure 4, but for $6^{\circ}S-22^{\circ}S$ EqL.

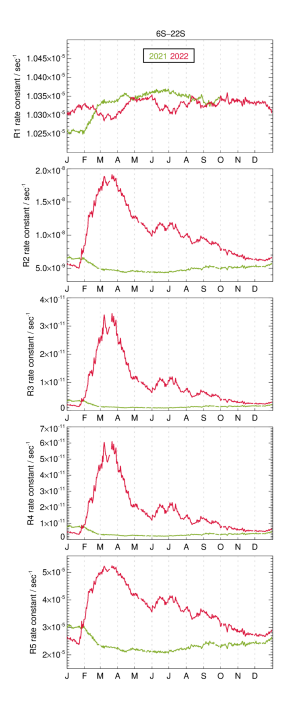


Figure 10. As Figure 5, but for $6^{\circ}S-22^{\circ}S$ EqL at 620 K.

Strong Evidence of Heterogeneous Processing on Stratospheric Sulfate Aerosol in the Extrapolar Southern Hemisphere Following the 2022 Hunga Tonga-Hunga Ha'apai Eruption

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Key Points:

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12	• Satellite data show widespread stratospheric chlorine and nitrogen repartitioning
13	in 2022 following the Hunga Tonga-Hunga Ha'apai eruption
14	• Observed composition changes are consistent with heterogeneous processing on
15	volcanic sulfate aerosol, in particular the hydrolysis of N_2O_5
16	• Moderate enhancements in reactive chlorine in the southern mid- and low-latitude
17	stratosphere did not cause appreciable chemical ozone loss

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18 Abstract

The January 2022 eruption of Hunga Tonga-Hunga Ha'apai (HT-HH) caused the largest 19 enhancement in stratospheric aerosol loading in decades and produced an unprecedented 20 enhancement in stratospheric water vapor, leading to strong stratospheric cooling that 21 in turn induced changes in the large-scale circulation. Here we use satellite measurements 22 of gas-phase constituents together with aerosol extinction to investigate the extent to 23 which the thick aerosol, excess moisture, and strong cooling enabled heterogeneous chem-24 ical processing. In the southern tropics, unambiguous signatures of substantial chlorine 25 and nitrogen repartitioning appear over a broad vertical domain almost immediately af-26 ter the eruption, with depletion of N_2O_5 , NO_x , and HCl accompanied by enhancement 27 of HNO_3 , ClO, and ClONO₂. After initially rising steeply, HNO_3 and ClO plateau, main-28 taining fairly constant abundances for several months. These patterns are consistent with 29 the saturation of N_2O_5 hydrolysis, suggesting that this reaction is the primary mecha-30 nism for the observed composition changes. The southern midlatitudes and subtropics 31 show similar but weaker enhancements in ClO and ClONO₂. In those regions, however, 32 effects of anomalous transport dominate the evolution of HNO_3 and HCl, obscuring the 33 signs of heterogeneous processing. Perturbations in chlorine species are considerably weaker 34 than those measured in the southern midlatitude stratosphere in 2020 following the Aus-35 tralian New Year's fires. The moderate HT-HH-induced enhancements in reactive chlo-36 rine seen throughout the southern middle and low-latitude stratosphere, far smaller than 37 those in typical winter polar vortices, do not lead to appreciable chemical ozone loss; rather, 38 extrapolar lower-stratospheric ozone remains primarily controlled by dynamical processes. 30

⁴⁰ Plain Language Summary

The January 2022 eruption of the Hunga Tonga-Hunga Ha'apai (HT-HH) volcano 41 caused the largest increase in stratospheric aerosol in decades and produced an unprece-42 dented increase in stratospheric water vapor, which led to strong stratospheric cooling 43 that in turn altered stratospheric winds. Here we use satellite measurements to inves-44 tigate the extent to which the thick aerosol, excess moisture, and strong cooling enabled 45 heterogeneous chemical processing. In the southern tropics, unambiguous signatures of 46 substantial changes in several chlorine and nitrogen species appear over a broad verti-47 cal range almost immediately after the eruption, with decreases in some species mirrored 48 by increases in others. The southern midlatitudes and subtropics show similar but weaker 49 evidence of chemical processing. In those regions, however, effects of anomalous wind pat-50 terns obscure the signs of heterogeneous processing in some species. Perturbations in chlo-51 rine are considerably weaker than those measured in the southern midlatitude strato-52 sphere following the Australian New Year's fires in 2020. The moderate HT-HH-induced 53 enhancements in reactive chlorine seen throughout the southern middle and low-latitude 54 stratosphere, far smaller than those in typical winter polar vortices, do not lead to ap-55 preciable chemical ozone loss; rather, outside the polar regions, lower-stratospheric ozone 56 remains primarily controlled by dynamical processes. 57

58 1 Introduction

The Hunga Tonga-Hunga Ha'apai (HT-HH) volcano in the Kingdom of Tonga be-59 gan an eruptive sequence in December 2021 that culminated in a paroxysmal explosion 60 on 15 January 2022 (e.g., Carn et al., 2022). It was the most explosive submarine erup-61 tion witnessed during the satellite era to date. Satellite and ground-based measurements 62 indicate that a substantial amount of material was lofted into the upper stratosphere; 63 although the bulk was deposited at altitudes between 20 and 30 km, the highest over-64 shooting tops of the eruption column penetrated into the lower mesosphere, reaching an 65 unprecedented \sim 55 km altitude (Carr et al., 2022; Khaykin et al., 2022; Millán et al., 2022; 66 Taha et al., 2022). The climactic blast was marked by the (previously undocumented 67

for a volcanic eruption) occurrence of a double umbrella cloud (that is, a cloud that spreads 68 laterally around the level of neutral buoyancy and rapidly transports the ejected volume 69 away from the volcano in all directions), with distinct signatures at about 30 km and near 70 the tropopause at $\sim 17 \,\mathrm{km}$ (Gupta et al., 2022). Also unprecedented was the mass of wa-71 ter vapor injected by HT-HH into the stratosphere, which has been estimated using Mi-72 crowave Limb Sounder measurements to have increased the total stratospheric water va-73 por burden by $\sim 10\%$ (Millán et al., 2022); other studies have calculated similar values 74 for the magnitude of the water vapor injection (Khaykin et al., 2022; Vömel et al., 2022; 75 Xu et al., 2022). Analysis of its isotopic composition shows that the injected water de-76 rived mainly from vaporized seawater, although some tropospheric moisture may also 77 have been entrained by the eruptive updraft (Khaykin et al., 2022). Peak enhancements 78 in water vapor were observed in the 25–30 km region (Khaykin et al., 2022; Millán et al., 79 2022; Sellitto et al., 2022; Vömel et al., 2022; Xu et al., 2022), consistent with being sourced 80 from the upper umbrella cloud (Gupta et al., 2022). The eruption column was also rich 81 in ice (Gupta et al., 2022; Khaykin et al., 2022); subsequent sedimentation and subli-82 mation of ice led to relatively humid conditions in the lower stratosphere (Khaykin et 83 al., 2022; Millán et al., 2022). 84

Although the explosive energy of HT-HH was similar to or even larger than that 85 of the 1991 eruption of Mt. Pinatubo (Poli & Shapiro, 2022; Wright et al., 2022; Yuen 86 et al., 2022), its SO₂ discharge into the stratosphere was $\sim 1-2$ orders of magnitude smaller 87 (Carn et al., 2022; Millán et al., 2022; Sellitto et al., 2022). SO₂ emissions from subma-88 rine volcanoes are typically lower than those from non-hydromagmatic eruptions of com-89 parable magnitude, presumably because of significant scrubbing of SO₂ in their water-90 rich eruption clouds (Carn et al., 2022). In addition, the SO₂ from HT-HH that did en-91 ter the stratosphere had a considerably shortened lifetime, undergoing unusually rapid 92 conversion to secondary sulfate aerosol (Legras et al., 2022; Sellitto et al., 2022). The 93 exceptionally fast formation of sulfate aerosol is also attributable to the abundant co-94 emitted water vapor, which provides a source of OH that in turn catalyzes the oxida-95 tion of SO_2 into H_2SO_4 , which then nucleates into sulfate aerosol (Zhu et al., 2022). With 96 its modest SO₂ emissions, HT-HH caused global-mean anomalies in stratospheric aerosol 97 optical depth smaller by about a factor of 6 than those resulting from Pinatubo (Khaykin 98 et al., 2022; Sellitto et al., 2022). Nevertheless, HT-HH led to the largest perturbation 99 in the stratospheric aerosol loading in the last 30 years (Khaykin et al., 2022; Sellitto et 100 al., 2022; Taha et al., 2022). 101

For the first few months after the eruption, the main volcanic cloud at 20-30 km 102 remained largely confined within the tropical pipe as it encircled the globe; thereafter 103 it gradually dispersed meridionally, with more efficient transport in the Southern Hemi-104 sphere, such that by June 2022 it had spread from about 60°S to 30°N, with small frag-105 ments reaching higher northern latitudes (Coy et al., 2022; Legras et al., 2022; Millán 106 et al., 2022; Schoeberl et al., 2022, 2023; Taha et al., 2022). The strong transport bar-107 rier at the boundary of the Antarctic vortex prevented the plume from penetrating to 108 high southern latitudes until that vortex broke down in late 2022 (Manney et al., 2023). 109 Initially the enhanced aerosol and water vapor were spatially coincident, but they grad-110 ually diverged. Entrained within the ascending branch of the Brewer-Dobson circulation. 111 the moisture anomaly was carried upward in the months following its injection, whereas 112 the sulfate aerosols were subject to gravitational settling; hence the aerosol- and water-113 rich layers became increasingly separated in the vertical after mid-February (Legras et 114 al., 2022; Schoeberl et al., 2022). 115

Both the aerosol and the water vapor perturbations induced radiative effects, but infrared cooling from enhanced water vapor concentrations was shown to dominate (Legras et al., 2022; Sellitto et al., 2022; Jenkins et al., 2023), leading to the development of a strong and persistent low-temperature anomaly in the mid-stratosphere shortly after the eruption (Coy et al., 2022; Schoeberl et al., 2022; Vömel et al., 2022). The anomalous temperature structure in turn induced changes in the large-scale circulation; in particular, zonal winds in the mid-stratosphere strengthened, as did meridional flow, while descent in the middle and high latitudes slowed (Coy et al., 2022).

Enhanced sulfate aerosol loading following volcanic injection of SO_2 into the strato-124 sphere is known to perturb the partitioning of key chemical families controlling strato-125 spheric ozone (e.g., Solomon, 1999). The exceptional stratospheric hydration and ensu-126 ing temperature anomalies caused by HT-HH may have increased such chemical process-127 ing. Transport also plays a major role in shaping trace gas distributions, however, and 128 129 the alterations to the stratospheric circulation arising from the eruption may have either compounded or attenuated the chemical effects, complicating interpretation of ob-130 served changes in composition. Here we investigate the impact of the HT-HH eruption 131 on stratospheric composition using satellite measurements. 132

¹³³ 2 Data and Methods

We examine measurements of H_2O , HNO_3 , ClO, HCl, N_2O , and O_3 from the Aura 134 Microwave Limb Sounder (MLS) (e.g., Waters et al., 2006). The enormous enhancement 135 in stratospheric water vapor from HT-HH, with the largest concentrations more than an 136 order of magnitude greater than any previously observed by MLS, degraded the accu-137 racy of some of the products retrieved using the most recent MLS data processing al-138 gorithms, version 5 (v5). It also caused a large proportion of the measurements in the 139 plume to fail the recommended MLS quality screening for the first few weeks after the 140 eruption, with retrieval performance not returning to normal until 8 February 2022. For 141 these reasons, Millán et al. (2022) used v4 MLS data to study the early stages of HT-HH 142 plume evolution; moreover, they did not impose quality filtering on the data. Since v5 143 generally represents an improvement over v4, particularly for H_2O and N_2O , and since 144 our main focus here is on the perturbations in composition arising in the weeks to months 145 following the eruption, we use v5 MLS data for all species, with standard data screen-146 ing protocols applied (Livesey et al., 2020). 147

MLS v5 ClO suffers from non-negligible biases at the lowest retrieval levels, 68-148 147 hPa (Livesey et al., 2020). As no substantial enhancement in nightime ClO is ex-149 pected in the extrapolar regions of interest here, the ClO biases are mitigated by tak-150 ing ascending ($\sim 2-3$ PM at these latitudes) minus descending ($\sim 2-3$ AM) differences, 151 similar to the strategy adopted in a prior study of midlatitude chlorine partitioning by 152 Santee et al. (2022). MLS N₂O is subject to a negative drift that, while partially alle-153 viated in v5, remains about -3%/decade in the lower stratosphere (Livesey et al., 2021). 154 Accordingly, the MLS N_2O fields presented below have been "detrended" by removing 155 a linear fit to the daily values across the Aura mission at each gridpoint. 156

We use meteorological fields (temperature, potential temperature, potential vor-157 ticity (PV)) from the Modern Era Retrospective Analysis for Research and Applications 158 Version 2 (MERRA-2, Gelaro et al., 2017). Satellite data are cast into quasi-Lagrangian 159 PV-based equivalent latitude (EqL) (Butchart & Remsberg, 1986) and potential tem-160 perature coordinates to facilitate tracking stratospheric air motions and allow parcels 161 with similar dynamical histories to be grouped together. Each day on average ~ 330 MLS 162 profiles fall within each of the 16° EqL bands defined here: 38°S-54°S EqL, 22°S-38°S 163 EqL, and 6°S–22°S EqL, representing southern midlatitudes, subtropics, and tropics, re-164 spectively. 165

In addition to MLS data, we use version 4.1/4.2 measurements of N₂O₅, NO, NO₂, and ClONO₂ from the Atmospheric Chemistry Experiment-Fourier Transform Spectrometer (ACE-FTS; Boone et al., 2020), screened using the recommended quality flags (Sheese et al., 2015; Sheese & Walker, 2020). In conjunction with the trace gas measurements, we use the latest version (V2.1) aerosol extinction coefficient profiles obtained by the Suomi-

NPP Ozone Mapping and Profiler Suite Limb Profiler (OMPS-LP) at both 869 nm, as 171 recommended for tracking stratospheric injections from events such as volcanic eruptions 172 or wildfires (Taha et al., 2021), and 745 nm. According to Taha et al. (2022), the stan-173 dard V2.1 OMPS-LP algorithms provide accurate aerosol retrievals as long as the vol-174 canic cloud is below 36 km; our analysis here is confined to the region below $\sim 33 \text{ km} (900 \text{ K})$. 175 We also compared the NASA OMPS-LP V2.1 aerosol products with the tomographic re-176 trievals at 745 nm produced by the University of Saskatchewan (USask v1.2; Bourassa 177 et al., 2023). Although the NASA and USask aerosol products show substantial differ-178 ences at low latitudes below 500 K, those discrepancies have no bearing on our conclu-179 sions, and for simplicity only the NASA aerosol data at 869 nm are shown here. As in 180 the study by Santee et al. (2022), for interpolation to isentropic surfaces, the aerosol data 181 are first converted to cross section per mole of air (in units of $m^2 mol^{-1}$), which, unlike 182 the more commonly used extinction coefficient (in units of $\rm km^{-1}$), is conserved under 183 changes of atmospheric pressure in the absence of aerosol formation or loss. The con-184 version factor relating cross section to extinction is the local molar volume, RT/p (in units 185 of $m^3 mol^{-1}$), where R is the molar gas constant, T is temperature, and p is pressure. 186 We use the remotely sensed optical properties without conversion to aerosol surface area 187 or volume density, as would be necessary to use these data as inputs in chemical mod-188 els (e.g., Hervig & Deshler, 1998; Kovilakam & Deshler, 2015). 189

3 Aerosol-Mediated Chemical Repartitioning 190

Within a few weeks of an explosive volcanic eruption, oxidative processing of the 191 SO_2 injected into the stratosphere leads to the production of liquid sulfate particles, en-192 hancing the aerosol surface area density (SAD) available for heterogeneous chemical pro-193 cessing (e.g., Bekki & Pyle, 1994). Heterogeneous reactions of particular importance in 194 the stratosphere include (e.g., Solomon, 1999): 195

$$N_2O_5 + H_2O \to 2 HNO_3 \tag{R1}$$

$$ClONO_2 + H_2O \rightarrow HOCl + HNO_3$$
 (R2)

$$CIONO_2 + HCl \rightarrow Cl_2 + HNO_3$$
(R3)

TT 0

$$HOCI + HCI \to Cl_2 + H_2O \tag{R4}$$

 (\mathbf{D}, \mathbf{i})

$$BrONO_2 + H_2O \rightarrow HOBr + HNO_3$$
 (R5)

Nitrogen, chlorine, and hydrogen photochemistry are strongly coupled, thus changes in 196 the partitioning of one chemical family affect species in other families. Substantial per-197 turbations in chemical partitioning can persist, as it can take a year or more for volcanic 198 sulfate aerosols to be removed from the stratosphere through sedimentation. To set the 199 stage for the interpretation of satellite measurements in Section 4, we first summarize 200 the expected impacts of R1–R5 on several key nitrogen and chlorine species affecting strato-201 spheric ozone, as well as ozone itself, under volcanically enhanced aerosol conditions. The 202 chemical processes discussed here are described in greater detail by, for example, Bekki 203 and Pyle (1994), Kinnison et al. (1994), Tie et al. (1994), Coffey (1996), Solomon (1999), 204 Anderson et al. (2012), Berthet et al. (2017), and Robrecht et al. (2019). For brevity, 205 these references are not cited for all relevant points in the following text. 206

3.1 HNO₃, N_2O_5 , and NO_x

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 HNO_3 is the major reservoir for odd nitrogen, NO_y , in the lower stratosphere. In 208 the immediate aftermath of the HT-HH eruption, a few individual MLS profiles showed 209 strong stratospheric enrichments in several trace gases, including HNO₃ (Millán et al., 210 2022, their Figure S1). Although it has been measured in volcanic plumes near the ground 211 (e.g., Mather et al., 2004; Oppenheimer et al., 2010; Voigt et al., 2014), in the lower strato-212

sphere volcanogenic HNO_3 has been conclusively detected only once, when a peak mix-213 ing ratio of ~ 3 ppbv above background was measured in an encounter with the fresh vol-214 canic cloud from the February 2000 eruption of Hekla (Hunton et al., 2005; Rose et al., 215 2006). Major explosive eruptions can enhance stratospheric HNO₃ through injection, ei-216 ther of HNO_3 itself or of nitrogen-bearing gases (e.g., ammonia) from which it can be 217 generated. Volcanic lightning can also induce HNO_3 formation, and, in addition to its 218 other unprecedented characteristics, HT-HH produced prodigious amounts of lightning 219 (Yuen et al., 2022). Thus, a direct volcanic origin for the anomalous post-eruption MLS 220 HNO_3 profiles is possible. However, they are more likely to be retrieval artifacts aris-221 ing from contamination of the MLS HNO_3 spectral signature by SO_2 . In any case, these 222 few highly localized and transient HNO₃ enhancements do not significantly affect the 223 broad EqL-band averages analyzed in this study, and they are not considered further. 224

Hydrolysis of N_2O_5 (R1) is fairly insensitive to temperature (Figure S1 in Support-225 ing Information) and takes place rapidly under virtually all stratospheric conditions. Its 226 rate is intensified dramatically when a major volcanic eruption enhances aerosol SAD. 227 Since (i) N_2O_5 is the main nighttime reservoir for reactive nitrogen, NO_x (defined here 228 as $NO+NO_2$, (ii) the HNO₃ produced through R1 returns to the gas phase, and (iii) the 229 rate at which HNO_3 undergoes conversion back to NO_x is unchanged, an increase in SAD 230 is generally expected to be accompanied by observed decreases in the abundances of both 231 N_2O_5 and NO_x and a corresponding increase in HNO₃. Similarly, R5 (hydrolysis of BrONO₂) 232 can be active at all latitudes and seasons during periods of high aerosol loading. 233

 $ClONO_2$ hydrolysis (R2) and reaction with HCl (R3) on sulfate aerosol are also po-234 tential sinks of NO_x and sources of gas-phase HNO₃. Unlike for R1 and R5, the reac-235 tion probabilities of R2 and R3 are strongly dependent on particle water content and hence 236 temperature (Figure S1 in Supporting Information), and typically they only become com-237 petitive with R1 in affecting nitrogen (and chlorine) partitioning under cold $(T < \sim 200 \text{ K})$ 238 conditions, such as in the winter polar regions, where they take place on the surfaces of 239 polar stratospheric cloud (PSC) particles. These two reactions, however, can come into 240 play to convert NO_x to HNO_3 in the midlatitude lower stratosphere when SAD and wa-241 ter vapor mixing ratios (and hence the aerosol water fraction) are sufficiently elevated 242 (Keim et al., 1996). Whereas condensed HNO_3 remains sequestered in PSCs in the win-243 ter polar regions, at higher temperatures a large proportion of the HNO_3 formed by these 244 reactions is released to the gas phase (Robrecht et al., 2019). 245

Consistent with heterogeneous processing, substantial reductions in NO_x and/or 246 N₂O₅ have been seen following several large eruptions (e.g., Fahey et al., 1993; Mills et 247 al., 1993; Koike et al., 1994; Rinsland et al., 1994; Coffey, 1996; Adams et al., 2017; Berthet 248 et al., 2017; Zambri et al., 2019). Significant decreases in NO_x observed in the months 249 after severe wildfires in Australia in the austral summer of 2019/2020 have also pointed 250 to the occurrence of heterogeneous processing on smoke particle surfaces (Solomon et 251 al., 2022; Strahan et al., 2022). Observational evidence for associated increases in HNO₃ 252 has been more ambiguous, however. Strongly enhanced HNO_3 (as high as 40-50% above 253 unperturbed background values in some cases) was measured in aged stratospheric vol-254 canic clouds sampled weeks or months after major eruptions (Arnold et al., 1990; Koike 255 et al., 1994; Rinsland et al., 1994; Jurkat et al., 2010). In addition, declining trends in 256 HNO_3 abundances as the sulfate aerosol loading from the eruption of Mt. Pinatubo slowly 257 decayed in subsequent years were attributed to the diminishing impact of R1 (David et 258 al., 1994; Kumer et al., 1996; Slusser et al., 1998; Rinsland et al., 2003; Santee et al., 2004). 259 In other cases, by contrast, no clear volcanic enhancement in HNO₃ was seen (Coffey, 260 1996; Adams et al., 2017; Berthet et al., 2017). Nor were HNO₃ abundances elevated fol-261 lowing the 2019/2020 Australian fires (Santee et al., 2022; Strahan et al., 2022). 262

Zambri et al. (2019) used coupled chemistry-climate model simulations in conjunc tion with MLS and other satellite measurements to investigate the impact on stratospheric
 composition of multiple moderate-magnitude eruptions in the post-Pinatubo era. Neg-

ative anomalies (with respect to unperturbed background values) in response to erup-266 tive events were seen in modeled N_2O_5 and measured and modeled NO_x , particularly 267 at 50–70 hPa, where anomalies exceeded 20–25%. The model showed a clear relationship 268 between SAD and HNO_3 changes through much of the stratosphere, but the perturbations in HNO_3 were not entirely congruent with those in NO_x . In particular, positive anoma-270 lies in HNO_3 extended over a larger altitude range, exceeding 10% during periods of in-271 creased sulfate aerosol SAD at levels from 70 to 5 hPa, with the exception of 50 hPa, where 272 fractional changes were much smaller than those in NO_x and confined mainly to the trop-273 ics. MLS data indicated similar, albeit weaker, aerosol-induced HNO_3 increases, again 274 except near 50 hPa. Zambri et al. (2019) offered no explanation for the behavior seen in 275 both measured and modeled HNO_3 fields at 50 hPa. Other recent modeling studies have 276 also found post-eruption reductions in NO_x and N_2O_5 of as much as 25-50% but con-277 comitant increases in HNO_3 of only 5–15% (Adams et al., 2017; Berthet et al., 2017). 278

It may not be surprising that a strong correlation between SAD/NO_x and HNO_3 is not consistently observed; HNO_3 concentrations in the lower stratosphere are much larger than those of NO_x , and thus comparable changes in its abundance have only modest relative effects. Small changes are difficult to detect against the backdrop of natural dynamical variability, which is of order 10% for HNO_3 in the lower stratosphere, making HNO_3 a less sensitive indicator of the occurrence of R1 than NO_x (e.g., Adams et al., 2017; Solomon et al., 2022; Strahan et al., 2022).

3.2 ClO

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ClO is the main form of ozone-destroying reactive chlorine. In the week following 287 the HT-HH eruption, MLS recorded strongly enhanced ClO in a handful of profiles (Millán 288 et al., 2022). The only halogenated compounds known to have been volcanically injected 289 into the stratosphere are HCl (discussed in Section 3.3) and OClO (Theys et al., 2014), 290 and the measured high ClO values were thought to largely reflect MLS retrieval artifacts 291 arising from SO_2 spectral interference, rather than direct stratospheric injection of ClO. 292 Given its rapid transformation into sulfate aerosol, the SO₂ from HT-HH no longer com-293 promised the reliability of MLS retrievals after late January, when abundances reverted 294 to background levels (Millán et al., 2022). As we are concerned here with the signatures 295 of spatially extensive chemical processing that manifest weeks to months after the HT-HH 296 injection, the initial enhancements in a few ClO profiles are not considered further. 297

Post-eruption decreases in NO_x impede $CIONO_2$ formation, shifting chlorine par-298 titioning toward ClO. These gas-phase processes affect ClO and ClONO₂ abundances 299 while leaving HCl unchanged. In addition, photolysis of the gas-phase HNO₃ and HOBr 300 resulting from the hydrolysis of N₂O₅, ClONO₂, and BrONO₂ (R1, R2, and R5) is a source 301 of reactive hydrogen, HO_x (OH+HO₂), and reduced NO_x concentrations also bring about 302 increases in HO_x by inhibiting the rate of NO₂+OH+M \rightarrow HNO₃+M. In the case of HT-HH, 303 moreover, simulations show that the massive injection of water vapor rapidly increased 304 stratospheric OH abundances (Zhu et al., 2022). Elevated OH abundances in turn ac-305 celerate conversion of HCl to reactive chlorine via $HCl+OH\rightarrow Cl+H_2O$. Thus, small en-306 hancements in stratospheric ClO may occur following major volcanic eruptions from R1 307 and R5 alone, under conditions unfavorable for R2–R4 (which may also occur on vol-308 canic sulfate aerosol where ambient temperatures and water vapor abundances allow). 309 Although in general the consequences of increased OH on stratospheric chlorine parti-310 tioning are expected to be relatively minor, in the extremely water-rich environment af-311 ter HT-HH, the gas-phase oxidation of HCl may be intensified. 312

Enhancements in ClO of a few tens to 100 ppt observed outside of the polar regions in the months following the eruption of Mt. Pinatubo were attributed to heterogeneous processing on sulfate aerosol, in particular R1 (Avallone et al., 1993; Dessler et al., 1993; Fahey et al., 1993; Toohey et al., 1993; Wilson et al., 1993; Keim et al., 1996). In their study of the effects of moderate eruptions since Pinatubo, Zambri et al. (2019) found responses of as much as 20–50 pptv in simulated ensemble-mean monthly mean zonal-mean
ClO from 70–5 hPa, but they showed an overall weaker correlation with SAD than that
of HNO₃. In many cases, changes similar to those simulated were not seen in Aura MLS
v4.2 bias-corrected ClO data. Those authors asserted that the precision of the MLS ClO
measurements was too poor to permit detection of small anomalies, basing that statement on the MLS single-profile precision without considering the benefits of averaging.

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3.3 HCl and ClONO₂

HCl and ClONO₂ are the main reservoirs of chlorine in the stratosphere. As discussed in Section 3.2, in the weeks following an explosive eruption, HCl can decrease through destruction by enhanced OH (since the rate of HCl loss through that mechanism is faster than its formation via Cl+CH₄ \rightarrow HCl+CH₃), and OH abundances may be especially elevated after the HT-HH water vapor injection. Heterogeneous chlorine chemistry (R2– R4) may also contribute if conditions allow. To our knowledge, however, HCl depletion induced by heterogeneous reactions on volcanic sulfate aerosol has never been observed.

MLS has captured direct volcanic injection of HCl into the stratosphere by a num-332 ber of moderate eruptions (Prata et al., 2007; Theys et al., 2014; Carn et al., 2016), al-333 though peak volcanic HCl concentrations are likely underestimated since the vertical ex-334 tent of the plumes is typically much smaller than the vertical resolution of MLS mea-335 surements. Only eight HCl profiles in mid-January 2022 exceeded the threshold used by 336 Millán et al. (2022) to identify enhancements (7σ , corresponding to ~5 ppbv of HCl through 337 much of the domain), so no spikes can be expected to stand out in the HCl zonal means 338 presented below. Scavenging by hydrometeors (liquid water drops or ice particles) or ash 339 in eruption columns prevents much of the HCl emitted by major eruptions from reach-340 ing the stratosphere (Tabazadeh & Turco, 1993; Textor et al., 2003). Klobas et al. (2017) 341 suggested that the very weak stratospheric HCl injection from Mt. Pinatubo (Mankin 342 et al., 1992; Wallace & Livingston, 1992) is attributable to the extremely wet conditions 343 caused by the coincidental passage of a tropical cyclone during the eruption, which may 344 have allowed most of the degassed HCl to be scrubbed from its plume. Noting the con-345 trast in HCl signals between the Sarychev and Kasatochi events despite their compara-346 ble magnitudes, Carn et al. (2016) postulated that the weaker HCl injection from the 347 latter arose through more effective scavenging by abundant water in its plume since it 348 erupted through a pre-existing crater lake. Similarly, the relatively modest HCl injec-349 tion from HT-HH may have been a consequence of removal in the exceptionally moist 350 and ice-rich environment in its plume (Carn et al., 2022). Nevertheless, it is conceivable 351 that volcanogenic HCl could compensate to some degree any depletion caused by het-352 erogeneous processing in the early post-eruption period. 353

If R2 or R3 are active to any significant extent, then ClONO₂ should decrease. On the other hand, some model simulations have shown mild increases in ClONO₂ (e.g., 20 pptv, \sim 16%) under enhanced volcanic SAD conditions, since ClO (the limiting reactant in ClO+NO₂+M→ClONO₂+M) increases (Kinnison et al., 1994; Berthet et al., 2017), while other modeling studies have found no discernible response in ClONO₂ (Zambri et al., 2019).

3.4 Ozone

360

The volcanically induced changes in stratospheric chlorine and nitrogen partitioning described above exacerbate chemical ozone destruction by the HO_x and ClO_x catalytic cycles (of primary importance in the lower stratosphere) but impede that by the NO_x cycle (the dominant ozone loss mechanism in the middle stratosphere). In addition to those direct chemical perturbations, increases in aerosol SAD affect radiative balance and hence the large-scale stratospheric circulation, with potentially considerable conse-

quences for the distribution of ozone (and other trace gases). Enhanced aerosol loading 367 can also alter photolysis rates (e.g., Tie et al., 1994), further modifying ozone abundances. 368 The net impact of these (in some cases competing) effects depends on the amount of sul-369 fate aerosol produced from the emitted SO_2 and the latitude, altitude, and timing of the 370 injection. Substantial reductions in lower-stratospheric and total column ozone have been 371 observed and modeled after several previous volcanic eruptions (e.g., Dhomse et al., 2015; 372 Hofmann & Solomon, 1989; Kilian et al., 2020; Millard et al., 2006; Naik et al., 2017; Ran-373 del et al., 1995; Solomon et al., 2016; Stone et al., 2017; Wilka et al., 2018, see also the 374 citations listed at the beginning of this Section, and references therein). On the other 375 hand, significant enhancement in ozone was observed in the southern midlatitudes above 376 ~ 26 km following Pinatubo; model simulations confirmed the important role of the NO_x 377 loss cycle—and its suppression through R1 on volcanic sulfate aerosol—in determining 378 the ozone budget at higher altitudes (Mickley et al., 1997). 379

380 4 Results

The unprecedented stratospheric impacts of HT-HH are clear in Figure 1, which 381 provides an overview of the evolution of aerosol cross section from OMPS-LP (panel (a)) 382 and water vapor from MLS (panel (b)) over their respective missions at 620 K potential 383 temperature, a representative lower-stratospheric level that corresponds to $\sim 20-30$ hPa 384 or $\sim 24-26$ km, depending on the EqL band and month. The previously reported initial 385 low-latitude confinement, early northward extension, and eventual more efficient south-386 ward dispersion of the aerosol and water vapor plumes (Coy et al., 2022; Legras et al., 387 2022; Millán et al., 2022; Schoeberl et al., 2022; Taha et al., 2022) are evident. Large anoma-388 lies in other chemical constituents also arise in the aftermath of the eruption. A strong 389 enhancement in HNO_3 quickly develops at the lowest latitudes and then gradually spreads 390 poleward in both hemispheres over the next several months (Figure 1c). HNO_3 anoma-391 lies of comparable magnitude are not infrequent in the 18^+ -year MLS record, however, 392 and around mid-year the anomaly abruptly shifts sign, becoming strongly negative. In 393 contrast, at this altitude the enhancement in ClO in the months after the eruption is un-394 paralleled (Figure 1d), while HCl is largely unaffected initially but exhibits a severe low 395 anomaly later in the year (Figure 1e). In the following subsections, we explore the ex-396 tent to which these perturbations can be attributed to heterogeneous chemical process-397 ing. To elucidate the role of dynamics in controlling trace gas distributions, we also ex-398 amine N_2O , a long-lived transport tracer whose vertical and meridional gradients in much 399 of the lower stratosphere are opposite to those of HNO_3 , HCl, and O_3 and which is there-400 fore generally anticorrelated with those species in that region (Figure 1f). Finally, Fig-401 ure 1g shows the unremarkable evolution of ozone in 2022. 402

We focus this analysis on the Southern Hemisphere (SH) midlatitudes, 38°S–54°S 403 EqL, and the SH tropics, 6°S–22°S EqL. The former allows comparison of the volcano-404 induced perturbations in chlorine and nitrogen partitioning with those engendered by 405 the Australian New Year's (ANY) fires in late 2019 / early 2020 (Santee et al., 2022; Solomon 406 et al., 2022, 2023; Strahan et al., 2022) and is also characterized by somewhat better ACE-FTS 407 data coverage; the latter experiences the largest HT-HH impacts on stratospheric com-408 position but suffers from poor ACE-FTS sampling. We also briefly examine the inter-409 mediate region of the SH subtropics, 22°S–38°S EqL. Equivalent latitudes poleward of 410 54°S are ignored to minimize the confounding influence of seasonal chemical processing 411 inside the Antarctic winter polar vortex. 412

In Figure 2 for the SH midlatitudes and similar figures for the other two EqL bands, we place the 2022 observations into climatological context and examine the evolution of chemical species in relation to sulfate aerosol loading, humidity, temperature, and transport. In the SH midlatitudes, we show 500 K (corresponding to ~47–57 hPa or ~20–21 km) to enable ANY comparisons and because the 2022 perturbations in ClO in that EqL band are largest there. In the SH subtropics and tropics, as in Figure 1, we again show 620 K, where the ClO enhancements at those latitudes peak. The two preceding years, 2020 and
2021, are also highlighted. Figure 3 and similar figures provide information on the vertical extent of the perturbations. Anomalies are calculated as described in the relevant
figure captions. Normalized anomalies are shown in the Supporting Information to facilitate comparison of changes across different quantities and EqL bands.

424

4.1 Southern Hemisphere Midlatitudes (38°S–54°S EqL)

We begin by looking at the SH midlatitudes, focusing on 500 K. Appreciable aerosol 425 from HT-HH starts arriving in mid-April in this region, where it swiftly surpasses that 426 resulting from ANY in 2020 and vastly exceeds the OMPS-LP mission envelope of be-427 havior (Figure 2a). These results are not sensitive to the choice of aerosol data set ex-428 amined (not shown; see Section 2). A corresponding positive anomaly in water vapor dis-429 plays a slight lag in its arrival relative to aerosol in this EqL band; although water va-430 por mixing ratios at 500 K are above the climatological mean through most of 2020 and 431 2021 and remain so into 2022 (as noted also by Manney et al., 2023), they do not turn 432 sharply upward at this level until mid-June (Figure 2b). Comparable delays in the up-433 turn in water vapor relative to that in aerosol occur at most levels (cf. Figures 3h and 434 3i; see also Figure S2 in Supporting Information). Khaykin et al. (2022) similarly found 435 slightly earlier appearance of aerosol than water vapor anomalies in the SH midlatitudes, 436 again based on OMPS-LP and MLS data. The increases in midlatitude aerosol and wa-437 ter vapor seen at 500 K likely result from a combination of poleward flow along that isen-438 trope and transport at higher levels (where the main volcanic cloud was initially deposited, 439 as noted in the Introduction) followed by downward motion in this EqL band, either through 440 gravitational settling in the case of aerosol or via diabatic descent for water vapor. We 441 hypothesize that the discrepancy in the onset of the enhancements in aerosol and wa-442 ter vapor in the SH midlatitudes arises from a faster aerosol sedimentation rate as com-443 pared to the rate of water vapor descent in the stratospheric circulation. 444

⁴⁴⁵ Climatological temperatures in this EqL band at 500 K decrease from January into ⁴⁴⁶ July; in 2022, temperatures are below average after mid-April and reach record lows in ⁴⁴⁷ mid-August (Figure 2c). A sizeable negative anomaly in N₂O through much of 2021 con-⁴⁴⁸ tinued into 2022, leading to low N₂O values at levels around 500 K for the first half of ⁴⁴⁹ the year (Figure 2d; see also Figures 3f and 3m). Another transport tracer measured by ⁴⁵⁰ MLS, CH₃Cl, exhibits similar, albeit weaker, departures from climatology in early 2022 ⁴⁵¹ (not shown), corroborating the picture seen in N₂O.

The evolution of the nitrogen species over the first few months of 2022 is more or 452 less typical; although mixing ratios of N_2O_5 (Figure 2f), NO_x (NO+NO₂, Figure 2g), 453 and HNO_3 (Figure 2h) at 500 K are initially below average, they all generally track the 454 normal seasonal behavior at first. By early June, however, N₂O₅ values drop well be-455 low any previously observed in this region by ACE-FTS, and NO_x is also slightly below 456 average. HNO_3 gradually increases relative to climatology until a very small positive anomaly 457 develops in early June (Figures 2h). Slightly larger enhancements are visible at higher 458 altitudes (540–620 K) from mid-April to mid-June (Figures 3c, 3j), but none stand out 459 against the considerable dynamically driven year-to-year variability. 460

ClO at 500 K closely follows climatological mean evolution for the first few months, 461 but it starts to climb steeply upward along with aerosol (but before water vapor) in early 462 May (Figure 21; the timing of the onset of the increases in ClO, aerosol, and water va-463 por can also be compared in Figure S2 in Supporting Information). Modest ClO enhancements of 10–20 pptv are seen on isentropic surfaces as low as 440 K and at least as high 465 as 660 K, with peak positive anomalies approaching 40 pptv in early August at 500 K (Fig-466 ures 3d, 3k). Non-negligible CIO anomalies (positive and negative) are also evident at 467 higher altitudes, but MLS frequently records comparable vertically extensive features (Fig-468 ure 3d); they likely arise through descent of changes in the mixing ratios at the secondary 469

peak in the ClO profile at around 1200 K in the upper stratosphere (not shown). Thus 470 we focus on the ClO anomalies at 660 K and below. Given that those anomalies are sig-471 nificant at the $>2\sigma$ level for several months over a range of altitudes (Figure 3k), there 472 is little doubt that they reflect real atmospheric features and not merely MLS measure-473 ment noise. Notable though they are, however, these enhancements are still substantially 474 weaker than the maximum ClO perturbation of ~ 80 pptv seen in the same EqL band 475 in 2020 in the wake of the ANY fires (Figures 2l, 3d) and attributed to heterogeneous 476 chlorine activation on smoke particles (Santee et al., 2022; Solomon et al., 2023; Stra-477 han et al., 2022). Indeed, the 2022 CIO enhancements are not especially distinctive ex-478 cept near the peak in the anomaly profile at 500 K, where they lie at or above the range 479 of variability observed over 2005–2019 from June to September and even exceed those 480 in 2020 during August (Figure 21). Consistent with the moderate increase in ClO, as dis-481 cussed in Section 3.3, $ClONO_2$ shows small enhancements during the intervals of ACE-FTS 482 sampling of this EqL band from June to August (Figure 2m). 483

Although HCl starts off quite a bit lower than normal, the suppressed values in early 484 January 2022 reflect the continuation of a pre-existing negative anomaly that began in 485 2020 and persisted through 2021 (Figures 2n, 3e). The enduring below-average HCl abun-486 dances at the turn of the year may reflect a combination of transport effects and resid-487 ual chlorine repartitioning after ANY (since aerosol, water vapor, and ClO also remained 488 slightly perturbed through most of 2021; Figures 2a, 2b, 2l), but in any case they pre-489 date the eruption and could not have been caused by it. HCl increases fairly steadily for 490 the first few weeks of 2022, reaching average values by mid-February. Thereafter, HCl 491 generally follows the typical pattern of behavior until starting to drop sharply at most 492 levels in mid-May (Figure 2n; also seen in Figures 3e, 3l). The downturn in HCl over May 493 to August 2022 roughly parallels the much deeper ANY-induced depletion in HCl that 494 occurred in earlier months in 2020 (Figure 2n). ACE-FTS indicates a very similar deficit 495 in HCl starting in July and persisting through the end of 2022 (not shown). At face value, 496 the significant concurrent but opposing changes in ClO and HCl point to the occurrence 497 of substantial chemical processing. The story is not so simple, however. 498

At about the same time as the changes in the chlorine species, rather than increas-499 ing as would be expected from R1-R3 (see Section 3.1), HNO₃ rapidly decreases (Fig-500 ure 2h), falling to very low values from July onward that at some levels redefine the bot-501 tom of the MLS mission envelope (not shown). This behavior is at odds with the changes 502 in N_2O_5 , whose values fall to near (or even below) zero, well outside the range previously 503 observed during the ACE-FTS sampling intervals throughout the second half of the year 504 (Figure 2f), and NO_x , which also sets new lower limits in October and November (Fig-505 ure 2g). The lifetimes of N_2O_5 and NO_x are sufficiently short that their anomalies are 506 almost completely driven by chemical perturbations, whereas HNO_3 is also subject to 507 dynamical control. The abrupt shift in HNO_3 , evident over a broad vertical range in the 508 lower and middle stratosphere (Figure 3c), is mirrored by corresponding rapid increases 509 in the transport tracer N_2O (Figures 2d, 3f). Scatter plots (Figure 4, left column) show 510 reasonably tight anticorrelation between HNO₃ and N₂O through much of the lower strato-511 sphere, including in 2022. At the levels where the mild positive HNO_3 anomaly is seen 512 from mid-April to mid-June (540–620 K), two distinct clusters—one in the middle of the 513 distribution from the early months of 2022, the other forming an extreme tail in the dis-514 tribution later in the year—are linked by the points undergoing a steep decline in HNO_3 515 and growth in N_2O in July. But at no time during the year do the 2022 values stray dra-516 matically from the historical HNO_3/N_2O relationship, indicating that the observed vari-517 ations in HNO₃ in the SH midlatitudes are governed to a large extent by transport pro-518 cesses. The levels that show anomalously low values in the latter part of 2022 fall near 519 or below below the peak in the HNO_3 profile, which is generally situated at around 600 K 520 in the SH (e.g., Santee et al., 2004), while at potential temperatures above 700 K, HNO₃ 521 anomalies are slightly positive at that time (Figures 3c, 3j). These signatures are con-522

sistent with the markedly weaker diabatic descent and stronger poleward flow in the SH midlatitudes in the aftermath of HT-HH reported by Coy et al. (2022).

To help diagnose chemical effects, we approximate odd nitrogen, NO_y , as $2 \times N_2O_5$ 525 + NO_x + HNO₃ + ClONO₂, with HNO₃ taken from MLS and the others from ACE-FTS; 526 together, these species account for about 97% of the stratospheric NO_y budget (e.g., Berthet 527 et al., 2017). While the ACE-FTS (averages of sunrise and sunset occultations) and MLS 528 measurements of these diurnally varying species are obtained at different local times, com-529 plicating precise quantification of NO_y , our purpose here is merely to qualitatively com-530 pare the behavior observed in 2022 with that in prior years. The ratios of N_2O_5 and NO_x 531 to NO_y (Figures 2i, 2j) confirm that stratospheric nitrogen partitioning is shifted away 532 from those species in the latter half of 2022. In contrast, HNO_3 makes up an anomalously 533 large fraction of NO_{u} from June onward, redefining the previously observed range in early 534 October and November (Figure 2k). These results suggest that heterogeneous HNO_3 pro-535 duction does take place in this EqL band in the months following HT-HH, but the sig-536 nature of such processing in Figures 2h and 3c is overwhelmed by large countervailing 537 transport effects. 538

The evolution of HNO_3 in the SH midlatitudes in 2022 differs from that observed 539 by the predecessor MLS instrument on the Upper Atmosphere Research satellite (UARS). 540 launched in September 1991, a few months after the June eruption of Mt. Pinatubo. UARS 541 $MLS HNO_3$ retrievals in the equatorial region suffered from contamination by strongly 542 volcanically enhanced SO_2 for the first ~100 days of the mission, but retrievals at higher 543 latitudes were unaffected by this artifact. UARS MLS first observed middle and high south-544 ern latitudes in November, by which time HNO₃ abundances were elevated throughout 545 a broad swath of the hemisphere, with values at 585 K well outside the range measured 546 by UARS MLS at similar times and locations during the rest of the mission (Santee et 547 al., 2004). HNO₃ mixing ratios did not return to normal at the highest EqLs until the 548 following June (1992), and the enhancement lingered for several months longer at mid-549 latitudes, in line with residual volcanic aerosol there. Following Pinatubo, heating from 550 absorption of longwave radiation by the volcanic aerosol altered the radiative balance 551 and dynamics of the stratosphere, intensifying both the upwelling in the tropics and the 552 downwelling in the SH extratropics (Aquila et al., 2012, 2013). This aerosol-induced per-553 turbation in the mean circulation caused a positive anomaly in ozone in the SH midlat-554 itudes (Aquila et al., 2013) and presumably had a similar effect on HNO₃, adding to any 555 chemical enhancement that may have occurred. In contrast, as noted earlier, the radia-556 tive perturbation from the exceptional water vapor injection by HT-HH counteracted 557 that from aerosol, resulting in net longwave cooling and weaker extratropical descent. 558 The contrasting circulation changes likely account to a large extent for the apparently 559 different HNO₃ responses to the two eruptions. 560

The strong influence of dynamics on HNO_3 in the second half of 2022 calls into ques-561 tion a primary chemical provenance for the steep mid-year dive in HCl. As with HNO_3 , 562 the rapid decrease in HCl coincides with an increase in N_2O (cf. Figures 2n and 2d). Scat-563 ter plots for HCl also show a bimodal structure in the lower stratosphere in 2022, with 564 two discrete clumps linked by the points from July (Figure 4, middle column). In the 565 case of HCl, however, notable deviations from the overall climatological HCl/N_2O re-566 lationship are seen at some levels. The conspicuous departure from anticorrelation in the 567 grey dots at 460–620 K is a signature of heterogeneous chlorine activation on smoke par-568 ticles from ANY (Santee et al., 2022). While behavior as clearly anomalous as that in 569 2020 is not evident in 2022, points from July / August onward at 460-660 K (the domain 570 over which ClO displays enhancements) do fall in positions that were previously unoc-571 cupied in the distribution to a much greater extent than they did for HNO_3 . To help dis-572 entangle dynamical and chemical effects, we define inorganic chlorine, Cl_y , as $ClO + ClONO_2$ 573 + HCl, with ClO and HCl from MLS and ClONO₂ from ACE-FTS. Stratospheric trans-574 port is known to play a key role in controlling the Cl_{y} distribution (Strahan et al., 2014). 575

The ratios of ClO and ClONO₂ to Cl_y (Figures 20, 2p) confirm that stratospheric chlorine partitioning is shifted toward those species, especially ClO, in the second half of 2022. The HCl/Cl_y ratios, on the other hand, reveal an anomalously small contribution to inorganic chlorine from HCl in early June, July, and August (Figure 2q). We surmise that, although the pronounced reduction in SH midlatitude HCl in the latter part of 2022 is chiefly governed by transport, HCl abundances are also being suppressed by chemical processing.

To obtain a qualitative sense of the onset of anomalous post-HT-HH partitioning 583 via reactions R1–R5 at 500 K, where the observed perturbations in the chlorine species 584 in this EqL band are largest, Figure 5 compares their respective first-order rate constants 585 in 2022 to those in 2021. The rate constant for a heterogeneous reaction is proportional 586 to its reaction probability (γ value, Figure S1 in Supporting Information), SAD, and the 587 thermal velocity of the molecule. The calculations in Figure 5 are based on the water 588 vapor and temperature values shown in Figure 2, but they use constant abundances for 589 the other gas-phase species as well as a fixed SAD of $2\,\mu\text{m}^2\,\text{cm}^{-3}$, representing background 590 conditions. Thus, these results are intended to illuminate only the influence of water va-591 por and temperature on R1–R5; they take no account of the enhanced aerosol loading, 592 the true chlorine species mixing ratios, or the evolution of those quantities over the year. 593 While using an enhanced SAD value for 2022 would more accurately characterize post-594 HT-HH conditions, doing so would affect the rate constants for R1–R5 proportionally 595 and thus would reveal no additional information about the relative effectiveness of those 596 reactions beyond that provided in Figure 5. Calculation of actual reaction rates, and hence 597 full quantification of the individual contributions of R1–R5 to the observed repartition-598 ing, requires detailed chemical modeling beyond the scope of this analysis. Nevertheless, 599 Figure 5 provides useful insight. The rate constants for R2–R4 are slightly higher in 2022 600 than in the preceding year from late July onward, not dissimilar to the timing of the per-601 turbations in ClO and HCl (i.e., the HCl/Cl_y ratio). This might suggest a role for those 602 reactions. However, their rate constants are still several orders of magnitude smaller than 603 those for R1 and R5. Taking the reciprocal of the maximum 2022 rate constants (which 604 are in units of s^{-1}) in Figure 5 yields approximate chemical lifetimes for the gas-phase 605 reactants of 1 day for R1, 10 months for R2, 60 years for R3, 40 years for R4, and 5 hours 606 for R5. Our results are in line with the modeling work by Robrecht et al. (2019), who 607 demonstrated that R2 and R3 (and by implication also R4) do not become effective un-608 til temperatures fall below 205 K even for very high water vapor mixing ratios of 20 ppmy 609 and volcanic conditions of $10 \times$ background SAD. Although at its peak the EqL-mean 610 aerosol cross section in this region is enhanced by about a factor of 20, the maximum 611 water vapor values are just over 5 ppmv, and minimum temperatures are $\sim 210 \,\mathrm{K}$. We 612 conclude that, despite the heavy aerosol loading, excess moisture, and strong cooling in 613 the 2022 SH midlatitudes following HT-HH, ambient conditions remain unfavorable for 614 R2–R4, and only R1 and R5 are at play in producing the observed moderate chlorine 615 activation. This is in contrast to the situation following ANY in 2020, when the high sol-616 ubility of HCl in aged stratospheric smoke particles apparently triggered heterogeneous 617 chlorine activation, in particular via R3, under relatively warm conditions in the SH mid-618 latitudes (Solomon et al., 2023). 619

Finally, an obvious question is whether the mild enhancement in reactive chlorine 620 in 2022 had a discernible impact on ozone concentrations. As noted by Santee et al. (2022), 621 although the strong and sustained midlatitude chlorine activation that occurred in the 622 wake of ANY was unprecedented in the satellite record, it was still an order of magni-623 tude or more weaker than that in a typical winter polar vortex. They found no conclu-624 sive observational evidence of significant ANY-induced chemical ozone destruction; rather, 625 they attributed the anomalously low ozone observed in the SH midlatitudes in 2020 (and 626 until midway through 2021) largely to transport effects, as did Strahan et al. (2022). In 627 contrast, other studies argued that heterogeneous chemistry on ANY smoke particles did 628 give rise to detectable ozone loss (e.g., Rieger et al., 2021; Solomon et al., 2022, 2023; 629

Yu et al., 2021). Given that the degree of chlorine activation in 2022 is about half that following ANY and the large-scale dynamical perturbations are far greater, we expect the evidence of chemical ozone destruction from HT-HH to be even more equivocal.

Figure 2e shows that the behavior of ozone in the SH midlatitude lower stratosphere 633 in 2022 is not dissimilar to that in 2020, with ozone mixing ratios generally low but not 634 outside the range of previously observed variability. Below 600 K, the changes in ozone 635 track those in HNO_3 and HCl; above that level, ozone becomes positively correlated with 636 N_2O in this EqL band and increases in the second half of the year (Figures 1g, 3g, 3n). 637 Although the $2022 O_3/N_2O$ correlations show two distinct populations in the early and 638 late months of the year at lower potential temperatures, as they do also for HNO_3 and 639 HCl, for the most part the points do not lie outside the typical transport-controlled dis-640 tribution (Figure 4, right column). Thus MLS measurements indicate that the minor in-641 creases in ClO provoked by HT-HH do not result in appreciable chemical ozone loss in 642 the SH midlatitude lower stratosphere in 2022. 643

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4.2 Southern Hemisphere Subtropics (22°S–38°S EqL)

The evolution in the SH subtropics at 620 K is similar to that at midlatitudes and 645 is summarized only briefly. Aerosol increases rapidly almost immediately after the erup-646 tion, reaching maximum values between April (620-660 K) and August (460-500 K) that 647 are larger at a given level than those at more poleward EqLs (Figure 6a; Figures S3a, 648 S3h in Supporting Information). HNO_3 ticks upward in late January above 560 K (Fig-649 ure 6h; Figures S3c, S3j in Supporting Information). This positive anomaly, though still 650 small, starts earlier, achieves a larger magnitude (>0.6 ppbv at 580–660 K in April), and 651 extends to higher altitudes than seen in the $38^{\circ}S-54^{\circ}S$ EqL band. Maximum values pro-652 trude above the mission envelope (Figure 6h), albeit only marginally and for a short in-653 terval in some cases. HNO_3 values continue to slowly rise in subsequent months at the 654 highest altitudes (Figure S3j in Supporting Information). At lower levels, however, they 655 decline, with a particularly precipitous drop in May–June over 540–700 K, and then re-656 main anomalously low through the end of the year. The downturn starts sooner but is 657 less abrupt than that seen at midlatitudes, and for the most part HNO_3 values do not 658 dip below the MLS mission envelope. Although the sparser sampling of ACE-FTS in this 659 EqL band compromises identification of anomalies, slightly depressed N_2O_5 (Figures 6f, 660 6i) and NO_x (Figures 6g, 6j) in later months again suggest repartitioning within the odd 661 nitrogen family, and elevated HNO_3/NO_y ratios (Figure 6k) are consistent with some 662 chemical production of HNO₃. 663

HCl follows the climatological mean for the first few months of 2022 before falling 664 fairly steeply to values lower than any previously observed above $550 \,\mathrm{K}$ (Figure 6n); be-665 low that level, the record HCl deficits in 2020 arising from ANY still stand. The decrease 666 in HCl begins about a month earlier than in the $38^{\circ}S-54^{\circ}S$ EqL band. As before, scat-667 ter plots (Figure S4 in Supporting Information, left and middle columns) show two sep-668 arated clusters in 2022 in much of the lower stratosphere, connected by points with rapidly 669 decreasing HNO₃ and HCl but increasing N₂O, in this case during May–June. Here, how-670 ever, above 460 K the 2022 HCl values lie in previously unpopulated areas of the distri-671 bution to a much greater extent than at midlatitudes. In addition, the HCl/Cl_{y} ratios 672 are indicative of chlorine repartitioning from April onward (Figure 6q). ClO shows an 673 even more prominent signature of chemical processing, with significant positive anoma-674 lies appearing over 460–700 K as early as March and reaching maxima of as much as 30– 675 35 pptv in May–July, depending on the level, as the peak in the ClO anomaly profile grad-676 ually descends in concert with the downward progression of the sulfate aerosol (Figures S3d, 677 S3k in Supporting Information). At most levels, the 2022 ClO abundances sit near or 678 above the top of the mission envelope for a few months (e.g., Figure 61); at and above 679 620 K, they exceed the ANY enhancements at all times, while at lower levels they are 680 initially smaller than 2020 values but then stay elevated through the end of the year (Fig-681

⁶⁶² ure S3d in Supporting Information). The timing of the buildup, peak values, and decay
⁶⁶³ of the ClO anomaly coincides closely with that of the aerosol anomaly but generally pre⁶⁶⁴ cedes that of the water vapor anomaly (Figure S2 in Supporting Information). Although
⁶⁶⁵ the general picture is similar to that in the SH midlatitudes, the relationship between
⁶⁶⁶ active chlorine and aerosol cross section is clearer in this EqL band.

As with HNO₃ and HCl, ozone mixing ratios briefly dip below the climatological mean at 620 K in June–July, mirroring the bump up in N₂O; thereafter, ozone rises steadily until it skirts the top of the observed range from October onward (Figure 6e). None of the analyses, including the scatter plots in Figure S4 in Supporting Information (right column), show patterns indicative of significant chemical ozone loss in 2022.

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4.3 Southern Hemisphere Tropics (6°S–22°S EqL)

In the SH tropics, aerosol (Figures 7a, 8a, 8h) and water vapor (Figures 7b, 8b, 8i) are abruptly enhanced through much of the lower stratosphere immediately after the eruption. At the levels of interest here, the choice of aerosol dataset used makes no material difference to these results. A warming trend throughout the stratosphere in this EqL band prior to the event is reversed, with temperatures dropping by $\sim 2 \text{ K}$ during February (Figure 7c; see also Schoeberl et al., 2022).

Concomitant with these changes, HNO₃ increases rapidly above 500 K (Figures 8c, 699 8j), overshooting the previously observed range by early February, with values then near 700 or above the top of the envelope through the first half of the year (Figure 7h). Large anoma-701 lies of $\sim 0.7-0.8$ ppbv are seen from 580–800 K (Figures 8j). The HNO₃/N₂O scatter plots 702 (Figure 9, left column) show that the March–July points lie at the outer edge of or slightly 703 apart from the main distribution at most levels, suggesting that transport is not the sole 704 factor controlling HNO_3 abundances at this time; rather, the observed behavior is in-705 dicative of heterogeneous processing. This determination is substantiated by the very 706 low N_2O_5 (Figures 7f, 7i) and NO_x (Figures 7g, 7j) values, as well as the greatly elevated 707 HNO_3/NO_4 ratios (Figure 7k) throughout 2022. Although the picture is unfortunately 708 fragmentary because of the poor ACE-FTS sampling at low latitudes, the 2022 values 709 of all of these quantities are clearly highly anomalous. 710

After rising steadily for a few weeks, however, HNO₃ mixing ratios then level off 711 and remain approximately flat until June, when they start to increase again, largely fol-712 lowing the typical seasonal pattern (Figure 7h). We interpret the March–May plateau 713 in HNO₃ as evidence of the saturation of N_2O_5 hydrolysis. The amount of N_2O_5 avail-714 able to participate in R1 is limited by its formation (mainly from the recombination of 715 NO_2 and NO_3 at night, whose rate depends quadratically on NO_x). When aerosol SAD 716 exceeds a certain threshold, N_2O_5 is hydrolyzed as fast as it can be formed, and further 717 increases in SAD do not affect the rate of conversion of NO_x to HNO_3 by R1 (e.g., Fa-718 hey et al., 1993; Tie et al., 1994). Although ACE-FTS coverage in the tropics is too sparse 719 to permit diagnosis of saturation in the NO_x data, this mechanism may explain why HNO_3 720 production appears to stall by early March at many levels (Figure 8j), while aerosol (Fig-721 ure 8h) and water vapor (Figure 8i) are still increasing. It may also account for the con-722 tinuous gradual increase in HNO_3 in April at higher potential temperatures (660–850 K), 723 as well as the lack of strong enhancement at lower potential temperatures $(460, 500 \, \text{K})$ 724 despite the presence of substantial aerosol there, since saturation occurs for relatively 725 lower aerosol amounts at lower altitudes (e.g., Mills et al., 1993). Similarly, Santee et al. 726 (2004) reported considerably greater impact from Pinatubo on HNO₃ abundances at higher 727 (585 K) than at lower (420, 465 K) potential temperatures based on UARS MLS mea-728 surements and attributed the differences to R1 saturation. 729

The evolution of ClO is also consistent with substantial sustained heterogeneous processing in this EqL band in 2022. Significant enhancements appear by mid-January from 500 to 700 K, with maxima as large as 40–45 pptv at 580–660 K in March (Figure 8k),

far exceeding the previously observed range of behavior (Figures 71, 8d; see also Figure 1d). 733 The enhancement slowly decays after April, but mixing ratios remain elevated over much 734 of the domain through the end of the year. A secondary MLS ClO product retrieved from 735 radiances in a different spectral band than that used in generating the standard ClO data 736 indicates a similar (albeit noisier) persistent strong enhancement (not shown). $CIONO_2$ 737 is also high during the brief windows of ACE-FTS sampling through most of the year 738 (Figures 7m, 7p). As with HNO₃, after an initial steep rise, ClO values then remain more 739 or less steady for the next several months (Figures 7l, 8k). A similar nonlinear response 740 to increasing aerosol SAD seen in airborne in situ ClO measurements after the eruption 741 of Mt. Pinatubo was attributed to the saturation of N_2O_5 hydrolysis on sulfate aerosols 742 (Avallone et al., 1993; Fahey et al., 1993). The aircraft campaigns also found the largest 743 ClO enhancements at the lowest latitudes sampled (Avallone et al., 1993). Thus, our re-744 sults are qualitatively in accord with the earlier in situ findings. In addition, the MLS-745 based CIO enhancements are comparable to post-eruption values simulated in several 746 modeling studies (e.g., Kinnison et al., 1994; Tie et al., 1994; Zambri et al., 2019). On 747 the other hand, Zambri et al. (2019) also examined MLS ClO measurements after mul-748 tiple moderate-magnitude eruptions in the post-Pinatubo era (see Section 3.2) and found 749 no enhancements matching those modeled, possibly because the HT-HH impact on strato-750 spheric chlorine activation in the SH tropics greatly surpasses that of any prior event. 751

In the immediate aftermath of the HT-HH eruption, as HNO_3 and ClO shoot up, 752 HCl displays a modest (<0.08 ppbv) but fairly sharp decrease over a limited altitude range 753 (540–660 K; Figure 81). This dip in January, which is not seen at higher latitudes, places 754 the 2022 HCl values at or near the bottom of the envelope at those altitudes in Febru-755 ary and March (Figure 7n). Given that N_2O is generally low at this level at this time 756 in 2022 (Figures 7d, 8f), correspondingly high HCl mixing ratios would have been ex-757 pected. After its initial decrease, HCl follows climatological patterns (i.e., deviations from 758 the mean remain essentially flat) for a few months, echoing the plateau in ClO (Figures 7n, 759 81) until dynamically driven changes—negative anomalies below 660 K, positive anoma-760 lies above (Figures 8e, 8l)—develop in the latter part of 2022. Although the HCl/N_2O 761 correlations are not especially unusual for the first half of the year, from about July on-762 ward they stand apart from the typical relationships at and above 620 K (Figure 9, mid-763 dle column). This behavior likely signifies continuing mid-year chemical suppression of 764 HCl, in consonance with the evolution of ClO. The extremely low HCl/Cl_{y} ratios seen 765 during the intervals of ACE-FTS coverage (Figure 7q) confirm the depletion of HCl through 766 heterogeneous chemical processing. 767

In the 620–660 K layer, close to the altitude of the maximum HT-HH water vapor 768 injection (Figure 8i; Millán et al., 2022), the steep increases in HNO₃ and ClO and the 769 decrease in HCl in the first weeks after the eruption occur nearly simultaneously with 770 the pulses in both water vapor and aerosol (Figure S2 in Supporting Information). Be-771 low that layer, however, HNO_3 , CIO, and HCl change more or less in unison with the 772 increase in water vapor, while the aerosol curves slope upward more gently and peak weeks 773 to months later. This behavior in the SH tropics differs from that in the subtropics and 774 midlatitudes, where, as discussed in the previous section, ClO enhancement generally 775 tracks the evolution of aerosol more closely than that of water vapor at altitudes below 776 600 K. Climate model simulations have shown that the extraordinary magnitude of the 777 HT-HH water vapor anomaly halved the SO₂ lifetime, leading to more rapid coagula-778 tion of larger sulfate particles than is typical (Zhu et al., 2022). The slower buildup in 779 aerosol at lower potential temperatures in the tropics may reflect the longer time required 780 for sulfate formation at less water-rich levels and/or the timescale for sedimenting par-781 ticles from above to reach those levels. The observed latitudinal differences in the rela-782 tionships between aerosol, water vapor, and reactive chlorine may arise because, by the 783 time a substantial portion of the HT-HH plume disperses to the SH subtropics, the con-784 version to sulfate aerosol is well underway. Nevertheless, sufficient sulfate aerosol to en-785 able heterogeneous processing is clearly present soon after the eruption even at lower al-786

titudes in the tropics, as evidenced by the rapid rise in HNO₃ and ClO there (Figures 8j,
8k; Figure S2 in Supporting Information).

In contrast to R1, reactions R2 and R5 are not subject to saturation in the lower 789 stratosphere. Thus, the fact that HNO_3 and CIO do not continue to increase in tandem 790 with aerosol implies that R1 is playing a dominant role in their production. Moreover, 791 a larger decrease in HCl, as well as a decrease (rather than an increase) in $CIONO_2$, would 792 be expected if direct heterogeneous chlorine activation were proceeding. The results of 793 our rate constant calculations (Figure 10) support this supposition. Although the ambient (temperature, water vapor) conditions in 2022 increase the rate constants for R2– 795 R4 considerably over their respective 2021 values starting in late January / early Febru-796 ary, they remain several orders of magnitude smaller than those for R1 and R5. There-797 fore, we conclude that, as exceptional as the aerosol and water vapor enhancements in 798 this EqL band are, they are not sufficient to promote R2–R4 to any significant extent 799 under the colder than normal but still relatively (compared to polar winter) warm con-800 ditions that prevail in the first half of 2022 (Figure 7c). 801

Figures 5 and 10 show that the rate constant for R5 is as much as 5-6 times faster 802 than that for R1. Photolysis of the HOBr and HNO₃ resulting from BrONO₂ hydrol-803 ysis is an important source of OH, rivalling that from nitrogen chemistry under condi-804 tions of volcanically enhanced aerosol; indeed, about half of the simulated changes in strato-805 spheric chlorine species after the Sarychev eruption were attributed to elevated OH abun-806 dances from R5 (Berthet et al., 2017). In addition to $ClONO_2$ and $BrONO_2$ hydroly-807 sis as a source of OH, the excess water vapor from HT-HH likely amplifies direct OH for-808 mation from H_2O oxidation; model simulations by Zhu et al. (2022) show increases in 809 OH in the SH tropics immediately following the eruption compared to a run with no wa-810 ter vapor injection. Unfortunately, MLS no longer measures OH itself, but measurements 811 of HO₂ (the other main member of the HO_x family) are available from MLS in the mid-812 dle stratosphere; those data indicate some enhancement in early 2022 (Figure S5 in Sup-813 porting Information). As noted above, the small decreasing tendency in HCl levels off 814 at approximately the same time as the increases in HNO_3 and ClO; this behavior may 815 point to saturation and hence the influence of enhanced OH arising from R1, rather than 816 from R5 or from the additional water vapor from HT-HH. Again, full quantification of 817 the effectiveness of the different processes contributing to OH production (and hence chlo-818 rine repartitioning) requires detailed chemical modeling beyond the scope of this study. 819

Finally, although heterogeneously driven chemical loss typically plays little or no 820 role in determining ozone abundances in the tropics, we investigate whether the unprece-821 dented degree of chlorine activation in the 500–700 K layer following HT-HH leads to per-822 ceptible reductions in ozone. At 620 K, ozone displays substantial variability through-823 out 2022, rising and falling fairly rapidly at various times and occasionally redefining the 824 top of the mission envelope (Figure 7e). Of particular note is the steep decline in March 825 and early April, which coincides with the maximum enhancement in ClO (Figure 71) and 826 thus could be a sign of chemical loss. However, ozone mixing ratios at this level barely 827 fall below the climatological mean before beginning to increase again at the end of May, 828 and surrounding levels also see only small decreases (Figure 8n). Ozone remains largely 829 anticorrelated with N_2O in the tropics at this altitude (cf. Figures 8f and 8g), and the 830 O_3/N_2O scatter plots show no hint of abnormally low O_3 values suggestive of depletion 831 at any point during the year (Figure 9, right column). Therefore, any signature of chlorine-832 catalyzed ozone loss that may be taking place in the lower stratosphere is masked by much 833 larger changes due to transport. On the other hand, points from the latter half of 2022 834 fall well outside the previously observed O_3/N_2O distribution at 660 K (and higher lev-835 els; not shown). These anomalously high ozone values may reflect diminished efficiency 836 of the NO_x catalytic loss cycle, which dominates ozone destruction in the middle strato-837 sphere (see Section 3.4). 838

5 Summary and Conclusions

The eruption of HT-HH in January 2022 caused a substantial perturbation in strato-840 spheric aerosol loading, which, although not unprecedented in the modern instrumen-841 tal era, was nevertheless the largest in the last several decades. Even more noteworthy 842 was the uniquely extreme enhancement in stratospheric water vapor. The submarine vent 843 erupted at a "Goldilocks" depth—sufficiently shallow that the pressure of the overlying 844 water did not dampen the power of the blast, sufficiently deep to allow extensive magma-845 seawater interaction (Witze, 2022). In addition, the plume was injected at an advanta-846 geous location, as lofting into lower altitudes or higher latitudes where air is colder would 847 have resulted in smaller water vapor anomalies (Schoeberl et al., 2022). The excess mois-848 ture led to strong stratospheric cooling that in turn induced anomalous circulation pat-849 terns. 850

In this study, we use constituent profile measurements from Aura MLS and ACE-FTS, 851 aerosol extinction from OMPS-LP, and meteorological parameters from MERRA-2 to 852 investigate the extent to which the dense aerosol, high humidity, and strong cooling brought 853 about by HT-HH prompted heterogeneous chemical reactions to occur in regions where 854 they are not normally active to a significant degree. We look for signatures of chlorine 855 and nitrogen repartitioning in the SH midlatitudes, subtropics, and tropics over a range 856 of altitudes in the lower and middle stratosphere. Calculated rate constants for the het-857 erogeneous reactions R1–R5 provide insights into their relative importance in altering 858 trace gas distributions. Ozone is also examined for evidence of chemical loss as a result 859 of the eruption. 860

In the SH midlatitudes ($38^{\circ}S-54^{\circ}S$ EqL), N₂O₅ and NO_x exhibit considerable de-861 creases at 500 K in the second half of 2022 that are consistent with heterogeneous pro-862 cessing on HT-HH aerosol. Although enhancements in HNO_3 would be expected in association with the reductions in N_2O_5 and NO_x , strong dynamical effects instead lead 864 to unusually low HNO_3 values that in some cases redefine the previously observed range, 865 and HNO₃ production only becomes apparent in the HNO_3/NO_y ratios. Signs of het-866 erogeneous chlorine activation are more compelling, with a positive anomaly in ClO reach-867 ing 35–40 pptv at its peak near 500 K from June through August, accompanied by an 868 increase in ClONO₂. Although significant at the $>2\sigma$ level, however, the ClO anomaly 869 is only marginally outside the envelope of previous behavior over a narrow layer for a 870 few months. Transport effects dominate the evolution of HCl in the latter part of 2022, 871 but chemically induced deficits in HCl that mirror the enhancements in ClO and ClONO₂ 872 are evident in the HCl/Cl_{y} ratios. Perturbations in all chlorine species are considerably 873 weaker than those measured at these levels in this EqL band in 2020 in the aftermath 874 of the ANY fires. Rate constant calculations suggest that, despite the unusually moist 875 and cool conditions in the SH midlatitude stratosphere following HT-HH, heterogeneous 876 chlorine activation reactions R2–R4 are not at work to any significant extent, and R1 877 and R5 (hydrolysis of N_2O_5 and BrONO₂, respectively) are primarily responsible for the 878 observed composition changes. Evidence of widespread and persistent HT-HH-induced 879 chlorine and nitrogen repartitioning is even clearer in the SH subtropics $(22^{\circ}S-38^{\circ}S EqL)$, 880 but the signatures of heterogeneous processing in HNO_3 and HCl are again obscured by 881 dynamical influences. 882

In the SH tropics (6°S–22°S EqL), unambiguous signatures of substantial hetero-883 geneous processing appear over a broad vertical domain almost immediately after the 884 eruption. Substantial repartitioning of both the nitrogen and the chlorine families oc-885 curs, as evidenced by depletion of N_2O_5 and NO_x together with enhancement of HNO_3 , 886 ClO, and $ClONO_2$. Unlike at higher latitudes, in this EqL band HCl also decreases rapidly 887 right after the eruption, and, in the second half of 2022, it displays extremely low HCl/Cl_{u} 888 ratios, as well as notable departures from the typical HCl/N_2O correlations. These pat-889 terns all point to significant chemical suppression of HCl. To our knowledge, volcanically 890 induced HCl depletion has not been reported previously. After initially rising steeply, 891

HNO₃ and ClO essentially plateau, maintaining fairly constant anomalies (of as much 892 as 0.7–0.8 ppbv and 40–45 pptv, respectively) for several months. This behavior is con-893 sistent with the saturation of N_2O_5 hydrolysis, suggesting that R1 is the primary mech-894 anism for the production of HNO_3 and, after associated gas-phase chemistry, the increases 895 in ClO and ClONO₂. Enhancements in OH brought about by the excess water vapor from 896 HT-HH may also play a role in the chlorine repartitioning. As at midlatitudes, rate con-897 stant calculations imply that reactions R2-R4 play no significant role in the observed 898 composition changes. Quantification of the relative contributions of the different pro-899 cesses perturbing stratospheric composition in the wake of HT-HH will require detailed 900 modeling efforts. 901

The moderate enhancements in reactive chlorine seen throughout the SH mid- and 902 low-latitude lower stratosphere following HT-HH do not lead to appreciable chemical loss 903 of ozone; rather, lower-stratospheric ozone in those regions remains primarily controlled 904 by transport processes. The lack of chemical ozone depletion in 2022 substantial enough 905 over an area extensive enough to be perceptible in the EqL means examined here is not 906 surprising. Although the 2022 ClO anomalies are unprecedented in the tropics and sub-907 tropics and are exceeded only by those induced by ANY in the midlatitudes, they are 908 still only about half as large as those arising from that event. Moreover, the ClO per-909 turbations caused by ANY were themselves more than an order of magnitude weaker than 910 the enhancements in typical winter polar vortices. In the middle stratosphere, by con-911 trast, reduced NO_x may have decreased the efficiency of that loss cycle, leading to mild 912 increases in ozone at and above 660 K. 913

The strong positive stratospheric aerosol anomaly from HT-HH has already sub-914 stantially abated, and, assuming no further major perturbations, stratospheric aerosol 915 loading is expected to return to pre-eruption levels by around mid-2023 (Khaykin et al., 916 2022). In contrast, stratospheric water vapor abundances remain historically high, with 917 the extreme enhancement likely to endure for several more years (Khavkin et al., 2022) 918 Millán et al., 2022). The HT-HH water vapor plume was effectively excluded from the 919 2022 Antarctic polar vortex and thus had little or no impact on that year's ozone hole; 920 similarly, it did not reach northern high latitudes in time to influence Arctic ozone loss 921 in boreal spring 2023 (Manney et al., 2023). However, the sustained HT-HH water va-922 por enhancement is expected to significantly affect Antarctic and Arctic vortex chem-923 istry and ozone loss during their upcoming 2023 and 2023/2024 winter/spring seasons. 924 Our ability to assess the impact of HT-HH on stratospheric chemistry and polar ozone 925 depletion over the next few years will depend strongly on the continuing availability of 926 satellite observations from Aura MLS and ACE-FTS. 927

6 Data Availability Statement

⁹²⁹ The data sets used here are publicly available, as follows:

930 •	Aura MLS Level 2 data (Lambert, Read, & Livesey, 2020; Manney et al., 2020;
931	Santee et al., 2020; Froidevaux et al., 2020; Lambert, Livesey, & Read, 2020; Schwartz
932	et al., 2020; Wang et al., 2020): https://disc.gsfc.nasa.gov/datasets?page=
933	1&keywords=AURA%20MLS
934 •	Aura MLS Derived Meteorological Products (DMPs; Manney & Millán, 2007–present):
935	https://mls.jpl.nasa.gov/eos-aura-mls/dmp (registration required)
936 •	ACE-FTS 4.1/4.2 data: http://www.ace.uwaterloo.ca (registration required
937	at https://databace.scisat.ca/l2signup.php)
938 •	ACE-FTS v4.1/4.2 error flags (Sheese & Walker, 2020): https://doi.org/10.5683/
939	SP2/BC4ATC
940 •	OMPS-LP L2 Aerosol Extinction Vertical Profile swath multi-wavelength daily
941	3slit Collection 2 V2.0 data (Taha, 2020): https://doi.org/10.5067/CX2B9NW6FI27

 MERRA-2 (Global Modeling and Assimilation Office (GMAO), 2015): https:// disc.sci.gsfc.nasa.gov/uui/datasets?keywords=%22MERRA-2%22

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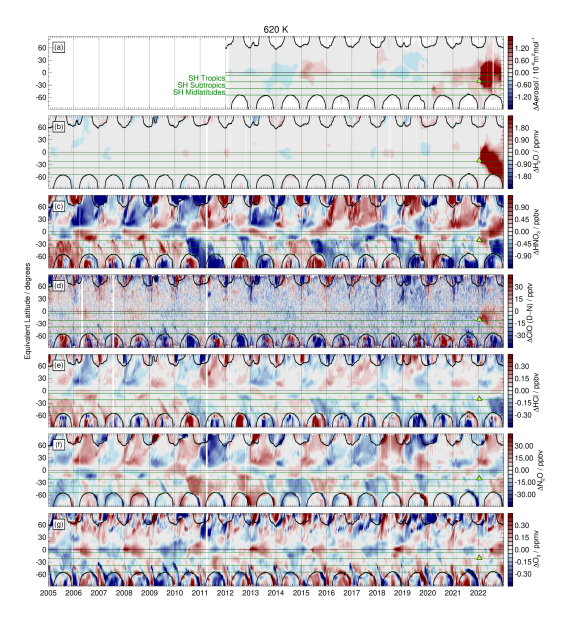


Figure 1. Equivalent latitude (EqL) / time series at 620 K of anomalies (calculated by subtracting from the daily data in each EqL bin the corresponding climatological value over 2005– 2021 for MLS and 2012–2021 for OMPS-LP for that day of the year) in (a) NASA OMPS-LP aerosol cross section at 869 nm and MLS (b) H_2O , (c) HNO_3 , (d) CIO (day minus night), (e) HCl, (f) N₂O, and (g) O₃. To ameliorate a negative drift in the MLS N₂O measurements (see Section 2), the N₂O anomalies (panel (f)) have been "detrended" by removing a linear fit to the daily values across the Aura mission at each EqL/potential temperature gridpoint. Blank spaces represent data gaps. Black overlays identify the approximate boundary of the winter polar vortices (as defined by Lawrence et al., 2018). The black horizontal line marks the equator; green horizontal lines mark the EqLs defining the regions examined in this study: the southern tropics (6°S–22°S), subtropics (22°S–38°S), and midlatitudes (38°S–54°S). The bright green triangle marks the time and latitude of the main HT-HH eruption.

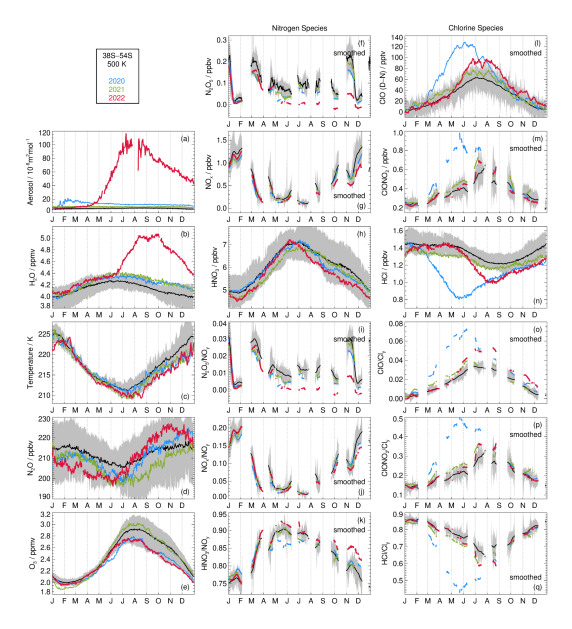


Figure 2. Averages over $38^{\circ}S-54^{\circ}S$ EqL (southern midlatitudes) at 500 K of (a) NASA OMPS-LP aerosol cross section at 869 nm, (b) MLS H₂O, (c) MERRA-2 temperature, (d) MLS N₂O, (e) MLS O₃, (f) ACE-FTS N₂O₅, (g) ACE-FTS NO_x (NO+NO₂), (h) MLS HNO₃, (i) N₂O₅/NO_y (where NO_y=N₂O₅+NO_x+HNO₃+ClONO₂), (j) NO_x/NO_y, (k) HNO₃/NO_y, (l) MLS ClO (day minus night), (m) ACE-FTS ClONO₂, (n) MLS HCl, (o) ClO/Cl_y (where Cl_y=ClO+ClONO₂+HCl), (p) ClONO₂/Cl_y, and (q) HCl/Cl_y. Values for 2022 are shown in red, for 2021 in green, and for 2020 in blue. Grey shading depicts the envelope of behavior and black lines the mean over 2005–2019 (thus excluding the highlighted years). ClO is one of the noisier MLS products; in addition, for this analysis we are taking day minus night differences, so fewer individual points contribute to these EqL-band averages than for the other species measured by MLS. Consequently, the ClO field displays a fairly large degree of day-to-day variability. To enhance clarity, the ClO results (highlighted years and mean, but not the envelope) have been smoothed using a boxcar of width 10 days. Although such smoothing attenuates extreme values, it facilitates detection of persistent anomalies. Similarly, all ACE-FTS fields, as well as all quantities derived using them, have also been smoothed with a 10-day boxcar average.

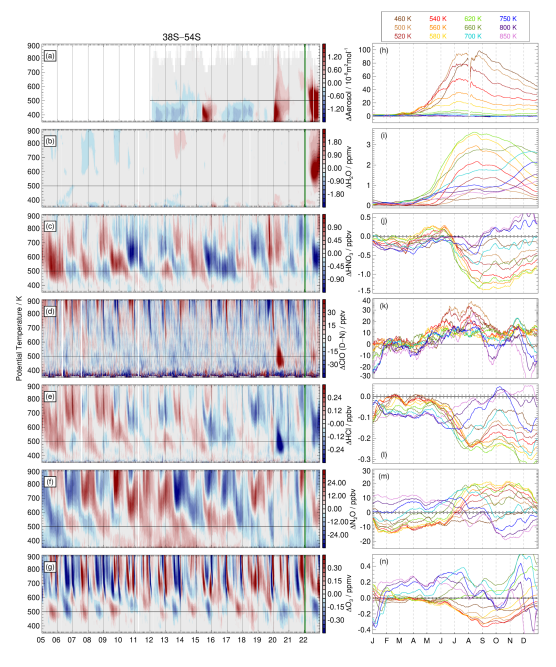


Figure 3. Potential temperature / time sections of anomalies (calculated as in Figure 1) in the 38°S–54°S EqL band of (a) NASA OMPS-LP aerosol cross section at 869 nm and MLS (b) H₂O, (c) HNO₃, (d) ClO (day minus night), (e) HCl, (f) N₂O, and (g) O₃. The vertical green line marks the date of the main HT-HH eruption. Panels (h)–(n) show the 2022 anomalies for each species at multiple potential temperatures (see legend). To enhance clarity, all curves in (h)–(n) except ClO have been smoothed using a boxcar of width 10 days; for ClO the smoothing window is 20 days. This smoothing removes some of the small-scale structure seen in the unsmoothed EqL-band averages shown in Figure 2. Solid black horizontal lines mark zero. Dashed black horizontal lines on the bottom five panels denote the 2σ contributions (± values) of MLS measurement noise to the smoothed EqL-band average anomalies, taken to indicate likely statistical significance for individual points in the anomaly timeseries.

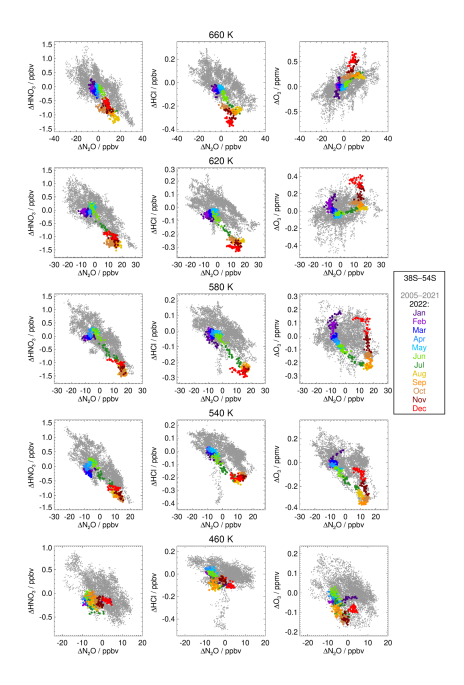


Figure 4. Scatter plots of daily HNO₃ (left), HCl (middle), and O₃ (right) vs. N₂O anomalies in the $38^{\circ}S-54^{\circ}S$ EqL band at various potential temperature levels. Grey dots depict data from 2005–2021; 2022 data are color-coded by month (see legend).

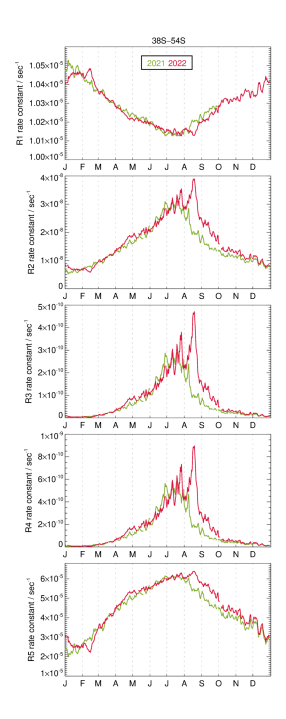


Figure 5. First-order rate constants for reactions R1–R5 in the $38^{\circ}S-54^{\circ}S$ EqL band at 500 K, based on the daily mean temperature and water vapor abundances observed in this region and the reactive uptake coefficients in Figure S1 in Supporting Information, assuming fixed abundances of 2 ppbv HCl, 0.1 ppbv ClONO₂, 0.1 ppbv HOCl, and 0.1 ppbv H₂SO₄ for a mean particle radius of 0.1μ m and an SAD of $2\,\mu$ m² cm⁻³. Values for 2022 are shown in red and for 2021 in green.

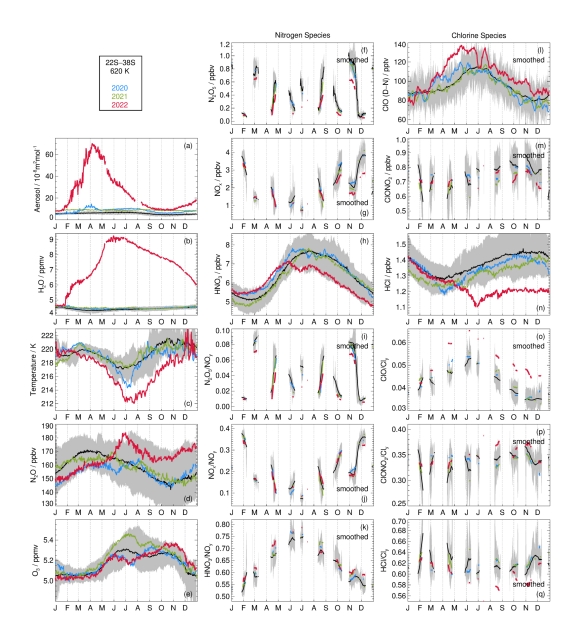


Figure 6. As Figure 2, but for 22°S–38°S EqL (southern subtropics) at 620 K.

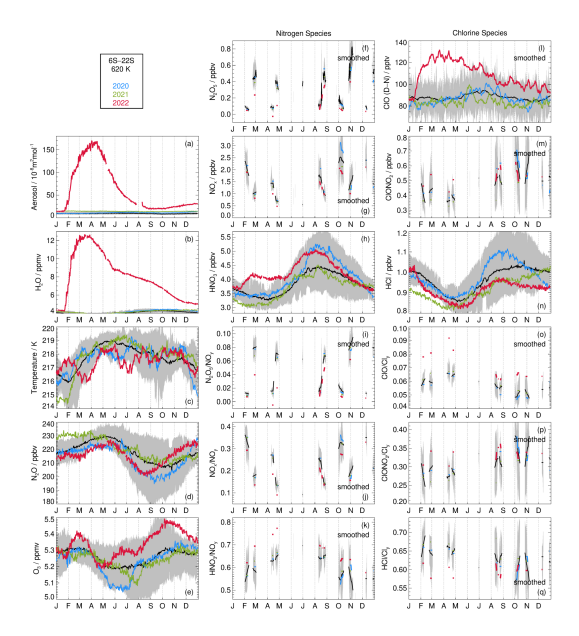


Figure 7. As Figure 2, but for 6°S–22°S EqL (southern tropics) at 620 K.

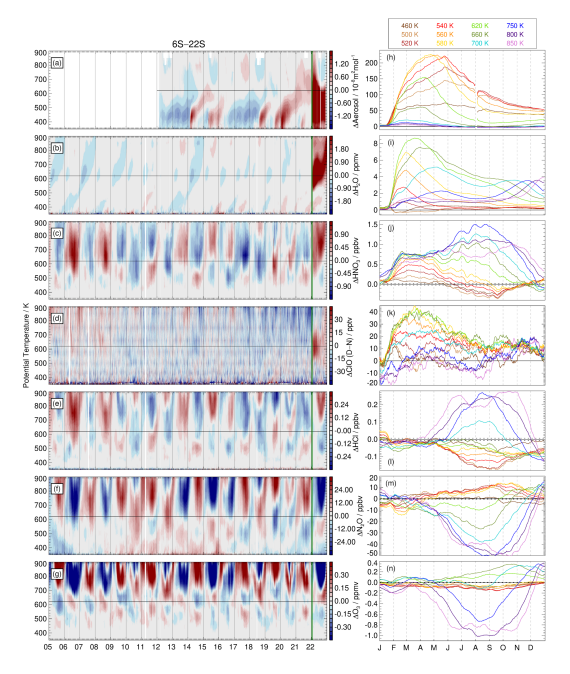


Figure 8. As Figure 3, but for 6°S–22°S EqL.

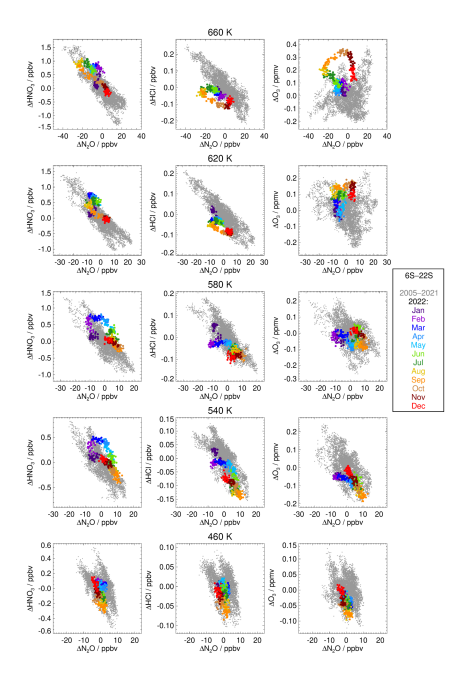


Figure 9. As Figure 4, but for $6^{\circ}S-22^{\circ}S$ EqL.

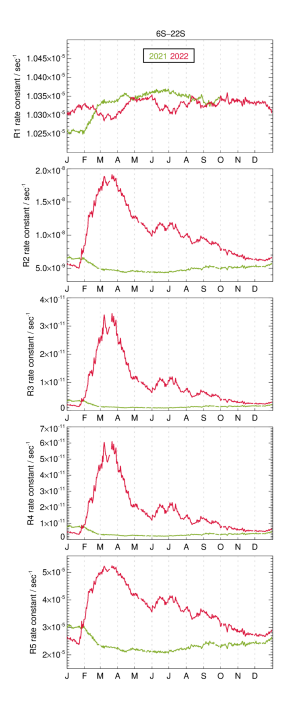


Figure 10. As Figure 5, but for $6^{\circ}S-22^{\circ}S$ EqL at 620 K.

- Supporting Information for "Strong Evidence of
- ² Heterogeneous Processing on Stratospheric Sulfate
- Aerosol in the Extrapolar Southern Hemisphere
- Following the 2022 Hunga Tonga-Hunga Ha'apai

⁵ Eruption"

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⁹ Contents of this file

10 1. Figures S1 to S5

11 Introduction

¹² Supporting Information contains additional figures supplementing the discussion in the ¹³ main text. Figure S1 shows the reactive uptake coefficients as a function of temperature ¹⁴ for the heterogeneous reactions R1–R5 discussed in the main text. The reactive uptake ¹⁵ coefficient (or surface reaction probability, γ) is the fraction of collisions between a gas-

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X - 2SANTEE ET AL.: HETEROGENEOUS PROCESSING ON HT-HH STRATOSPHERIC SULFATE AEROSOL phase reactant and a particle surface that lead to reactive uptake of the trace gas species 16 on/in the particle. Figure S2 shows anomalies of aerosol cross section from the Suomi-17 NPP Ozone Mapping and Profiler Suite Limb Profiler (OMPS-LP) and of water vapor, 18 HNO₃, ClO, HCl, N₂O and ozone from the Aura Microwave Limb Sounder (MLS) similar 19 to those shown in the main text. Here, however, to facilitate comparison of changes across 20 different quantities and equivalent latitude bands, all anomalies have been normalized by 21 dividing the raw magnitudes by their respective maximum absolute values. Results are 22 shown for the Southern Hemisphere (SH) midlatitudes, subtropics, and tropics. Figure S3, 23 similar to Figures 3 and 8 in the main text, gives the vertical view of the anomalies in 24 various quantities for the SH subtropics. Figure S4 shows MLS HNO₃/N₂O, HCl/N₂O, 25 and O_3/N_2O correlations as in Figures 4 and 9 in the main text, but for the SH subtropics. 26 Figure S5 shows a potential temperature / time section similar to those in Figure S3, but 27 for the anomaly in MLS monthly HO_2 in the SH tropics; in this case, the lowest potential 28 temperature shown is $670 \,\mathrm{K}$, as the MLS HO_2 data are not recommended for scientific 29 study at retrieval levels below (i.e., pressures greater than) 10 hPa (Livesey et al., 2020). 30

References

31	Burkholder, J. B., et al. (2019). Chemical kinetics and photochemical data for use in
32	atmospheric studies, evaluation number 19 (Tech. Rep. No. JPL Publication 19-5).
33	Pasadena: Jet Propulsion Laboratory. (Available at http://jpldataeval.jpl.nasa.gov)
34	Livesey, N. J., et al. (2020). Version 5.0x Level 2 and 3 data quality and description docu-
35	ment (Tech. Rep. No. JPL D-105336 Rev. A). Pasadena: Jet Propulsion Laboratory.
36	(Available at http://mls.jpl.nasa.gov.)

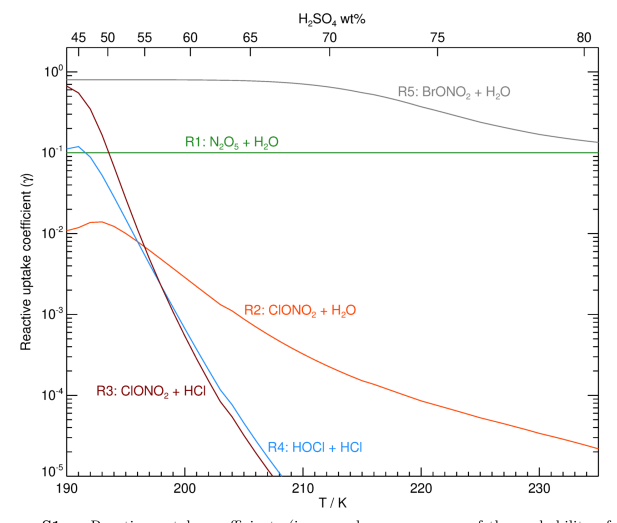


Figure S1. Reactive uptake coefficients (i.e., γ values, a measure of the probability of occurrence of a given reaction) for R1–R5 on stratospheric sulfuric acid aerosols as a function of temperature. Calculations follow the recommendations given by Burkholder et al. (2019) and assume fixed abundances of 5 ppmv H₂O, 1 ppbv HCl, 0.6 ppbv ClONO₂, 0.1 ppbv HOCl, and 0.1 ppbv H₂SO₄ for a mean particle radius of 0.1 μ m at 50 hPa.

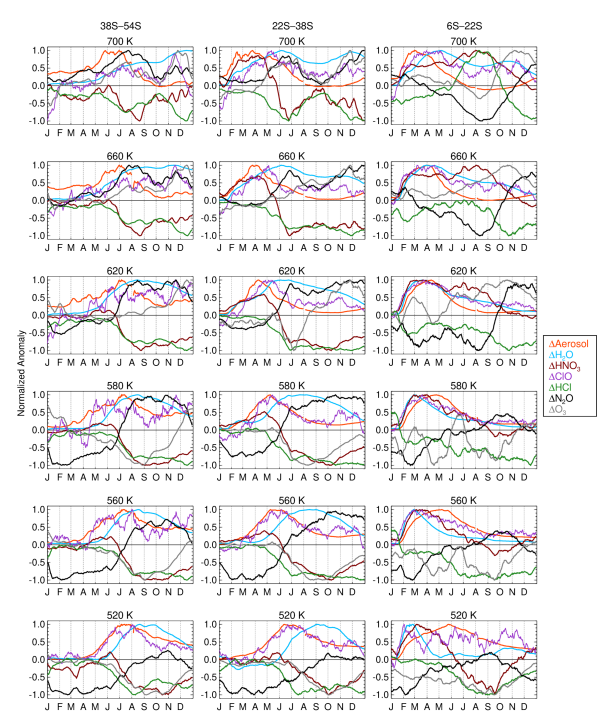


Figure S2. Normalized anomalies of NASA OMPS-LP aerosol cross section at 869 nm and MLS H_2O , HNO_3 , ClO (day minus night), HCl, N_2O , and O_3 (see legend) at six selected potential temperatures for the (left) 38°S–54°S, (middle) 22°S–38°S, and (right) 6°S–22°S EqL bands. Anomalies are normalized by dividing the raw magnitudes by their respective maximum absolute values. Solid black horizontal lines mark zero.

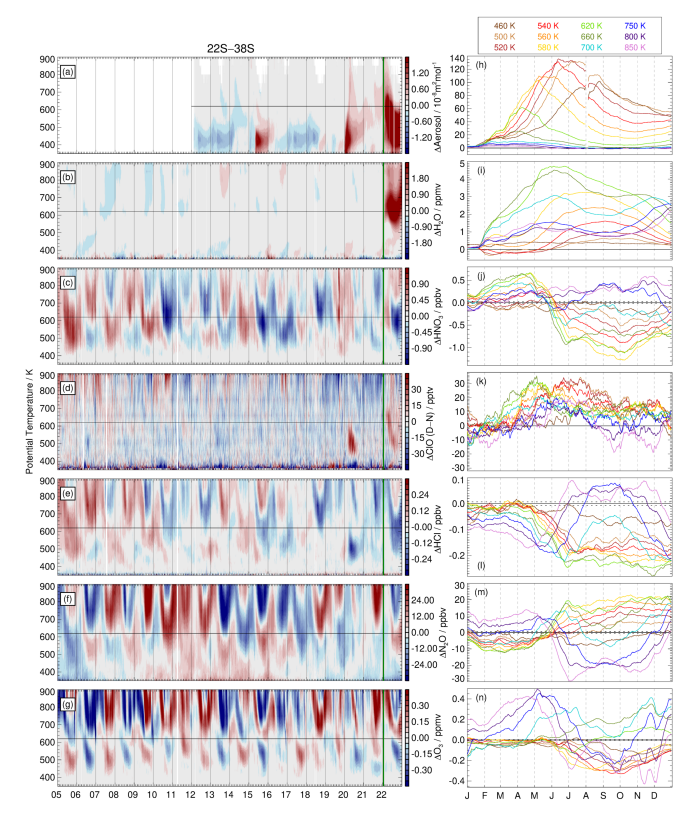


Figure S3. As Figure 3 in the main text, but for the 22°S–38°S EqL band.

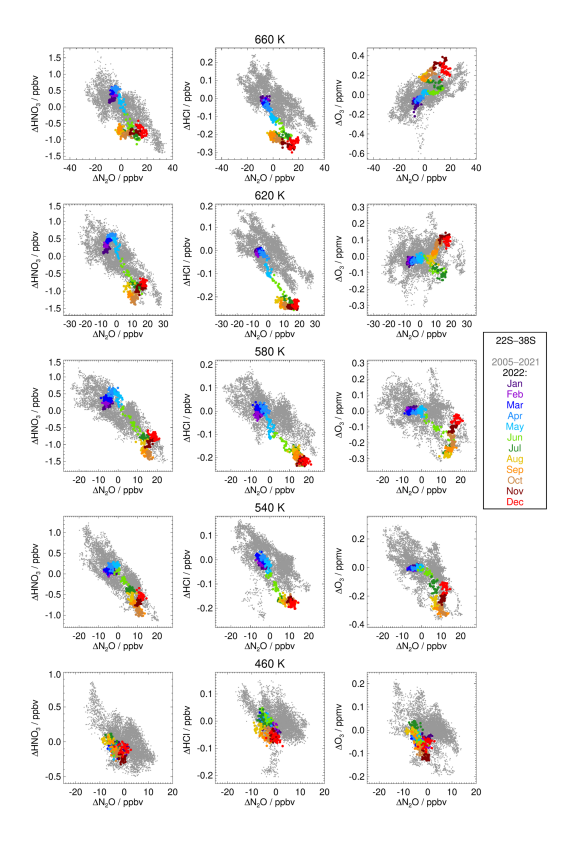


Figure S4. As Figure 4 in the main text, but for the 22°S–38°S EqL band.

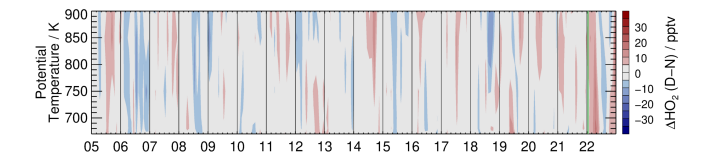


Figure S5. Potential temperature / time section, as in Figures 3 and 8 in the main text, but for the anomaly in MLS monthly HO_2 (day minus night) in the 6°S–22°S EqL band.