

Stratospheric aerosol formed by intense volcanism – sea interaction during the 2022 Hunga Ha’apai eruption

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Abstract. The Hunga Tonga eruption the 15 January 2022 (HT-22) induced vigorous volcano – sea interaction. Here we study the stratospheric aerosol and water vapor resulting from the eruption using satellite-based instruments: the CALIOP lidar and the Microwave Limb Sounder (MLS). We investigate the stratospheric relative humidity following the record-breaking water vapor injections from the HT-22 eruption, and the particle size of the aerosol. The HT-22 eruption injected its effluents into the deep Brewer-Dobson (BD) branch causing several years of stratospheric perturbation. The long duration, and aerosol concentration among the highest, makes the HT-22 eruption the strongest stratospheric aerosol event since the 1991 Mt. Pinatubo eruption despite a modest SO₂ injection explaining only ~30% of the AOD from the HT-22 eruption according to our estimates. The stratospheric AOD level was established after 2 weeks, or possibly even earlier, which is a short time compared with the usual 2 – 3 months required to reach the maximum AOD following volcanic eruptions. We discuss the sources of the aerosol from the HT-22 eruption in relation to the low emission of SO₂, its e-folding time and volcanological observations of strong interactions with the sea containing not only water but also high concentrations of dissolved substances.

1 Introduction

The stratospheric background conditions are frequently offset by injections of copious amounts of aerosol and gases from explosive volcanic eruptions (Kremser et al., 2016) and intense wildfires forming pyrocumulonimbus clouds (Fromm et al., 2010). These events cause variable stratospheric impact with durations of months to several years (Friberg et al., 2018), which are important to account for in climate models (Schmidt et al., 2018).

The Hunga Tonga – Hunga Ha’apai volcano erupted on 15 January 2022, with a volumetric flow rate an order of magnitude higher than that of the 1991 Mt Pinatubo eruption, and formed an umbrella cloud at 31 km and a second cloud at 17 km altitude (Gupta et al., 2022). Further, a record-breaking overshooting plume reached above 50 km (Carr et al., 2022, Proud et al., 2022, Taha et al., 2022). The volcanic explosivity index (VEI) was estimated to be 6, based on seismological observations (Poli and Shapiro, 2022). Despite the high VEI, ash could not be detected in the ice-rich stratospheric clouds from the HT-22 eruption (Gupta et al., 2022), and the UV aerosol index (UVAI) indicates low ash content (Carn et al., 2022). This is further supported by CALIOP (Cloud-Aerosol Lidar with Orthogonal Polarization) measurements finding very low depolarization ratios indicating dominance of spherical particles uncharacteristic of ash (Legras et al., 2022). Additionally, the volcanic layers in the stratosphere contained very low SO₂ amounts for such a strong eruption (Carn et al., 2022).

37 Widespread damage to the seafloor with runouts exceeding 100 km was caused by volcanoclastic density
38 currents, suggesting a collapsing eruption column entering the sea (Seabrook et al., 2023; Clare et al., 2023).
39 Such a sequence of events where hot volcanoclastic density currents form induces strong interaction with sea
40 water over vast areas, that can supply hot water vapor forming a plume that is buoyant at the base and
41 accelerates as it rises (Mastin et al., 2024). A relatively small eruption can in this way form umbrella clouds the
42 size and altitude of the HT-22 eruption, whereas entrainment of vapor from cold water does not (Mastin et al.,
43 2024). Other possible mechanisms include formation of an explosive steam from superheated water in contact
44 with the erupting magma (Millán et al., 2022).

45 The stratospheric background aerosol contains mainly sulfurous and carbonaceous components with some
46 extraterrestrial and tropospheric components (Murphy et al., 2007, Kremser et al., 2016, Martinsson et al.,
47 2019). Volcanic aerosol in the stratosphere normally contains large amounts of sulfuric acid formed from sulfur
48 dioxide (SO₂), water, carbonaceous material and ash (Martinsson et al., 2009; Andersson et al., 2013; Friberg et
49 al., 2014). Wildfires produce an aerosol initially dominated by organic and black carbon (Garofalo et al., 2019),
50 where the former component is rapidly removed by photolysis (half-life 10 days) in the stratosphere (Martinsson
51 et al., 2022; Friberg et al., 2023).

52 The volcanic and wildfire events also affect particle size distribution. During a long period with conditions close
53 to the background, spanning 1998 to 2004, the particle volume mode was 0.2 – 0.3 μm in diameter, whereas
54 approximately 1 μm in 1992 – 1993 after the Mt. Pinatubo eruption (Bauman et al., 2003; Wilson et al., 2008).
55 Measurements the second week after the 2017 Canadian wildfire showed particle diameter of 0.6 – 0.7 μm
56 (Haarig et al., 2018; Hu et al., 2019).

57 In this work we investigate the stratospheric aerosol resulting from the HT-22 eruption in relation to the
58 volcanological sequence of events during the eruption. We also investigate the interaction of the aerosol with the
59 large amounts of water vapor injected into the stratosphere. The global stratospheric aerosol optical depth
60 (AOD) is studied 1.5 years after the eruption, until the decommission of the NASA satellite CALIPSO (Cloud-
61 Aerosol Lidar and Infrared Pathfinder Satellite Observation) and its lidar sensor CALIOP. Our incrementally
62 developed evaluation software (Andersson et al., 2015; Friberg et al., 2018; Martinsson et al., 2022) based on
63 methodology presented in Vernier et al., (2011) was applied on CALIOP level 1B data. In contrast to limb-
64 oriented methodology, the nadir-oriented CALIOP provides viable results in dense aerosol layers from strong
65 volcanic eruptions and wildfires after correction for attenuation (Martinsson et al., 2022). We also use the
66 satellite Aura sensor MLS for measurements of water vapor and temperature. We find that the SO₂ emissions
67 from the HT-22 eruption cannot alone explain the high AOD level, nor can ash particles. We also find that the
68 aerosol went deep into the stratosphere and that the one-year AOD perturbation due to the HT-22 eruption is the
69 largest since that of Mt. Pinatubo in 1991.

70 **2 Methods**

71 Two satellite-based instruments were used to investigate the stratosphere following the HT-22 eruption. Aerosol
72 measurements were based on the CALIOP lidar aboard CALIPSO, whereas water vapor concentrations and
73 atmospheric temperature were obtained from MLS aboard Aura.

74 **2.1 CALIOP measurements**

75 CALIPSO orbits the globe 14 – 15 times per day between 82° S and 82° N. The vertical resolutions of CALIOP
76 are 30, 60, 180 and 300 m in the altitude ranges <8.2, 8.2 – 20.2, 20.2 – 30.1 and 30.1 – 40 km, respectively
77 (Winker et al., 2007, 2010). The average global stratospheric AOD from the tropopause (obtained from the
78 MERRA-2 reanalysis (Modern-Era Retrospective analysis for Research and Applications)) to 35 km altitude in
79 the stratosphere was computed from version 4-51 of CALIOP level 1B at the wavelength 532 nm using night-
80 time measurements. The stratospheric AOD was computed in three layers: the lowermost stratosphere (LMS,
81 tropopause to 380 K isentrope), the shallow BD branch (380 – 470 K isentropes) and deep BD branch (470 K
82 isentrope – 35 km altitude), where potential temperatures were obtained from MERRA-2 pressures and
83 temperatures. The effective lidar ratio was estimated based on single, intense volcanic layers day 1 – 28 after the
84 eruption. From initial high values (70 sr) the lidar ratio declined to 47.5 ± 10.2 sr. This is close to the commonly
85 used CALIOP effective lidar ratio of 50 sr, which we therefore applied in this study. The attenuated backscatter
86 CALIOP data were corrected by methods described in Martinsson et al. (2022). Based on measured parallel and
87 perpendicularly polarized scattering, the volume depolarization was obtained and converted to particle
88 depolarization ratios with methods described in Martinsson et al. (2022). Data were missing for a week from a
89 few days after the eruption, and a long gap appeared from 21 October to 7 December 2022. Several minor gaps
90 appeared during the first half-year of 2023 the last data produced by CALIOP.

91 **2.2 MLS measurements**

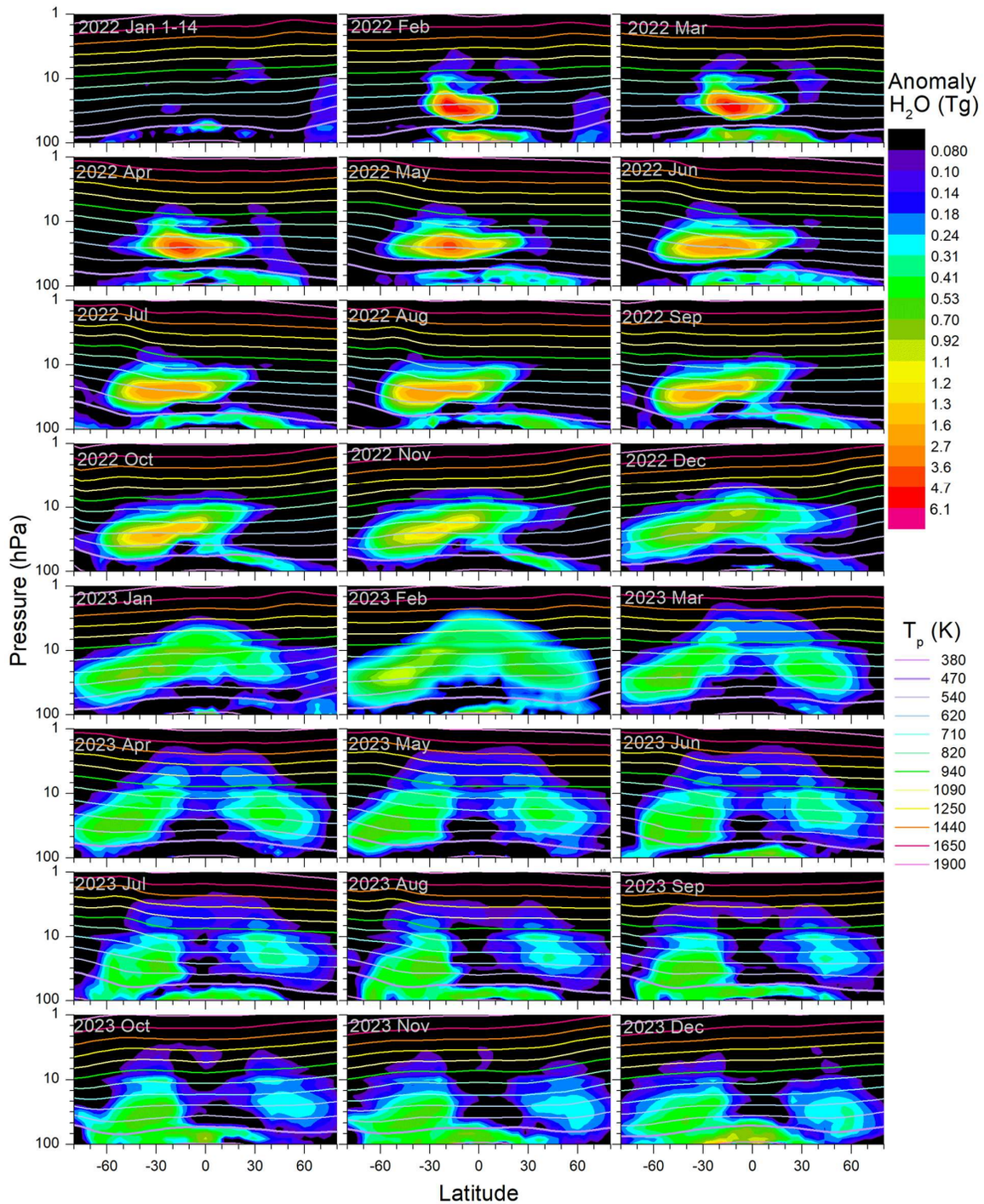
92 Water vapor concentrations were obtained in the 100 – 1 hPa range in 12 levels per decade from the MLS,
93 version 5.0-1.0a, level 2 (Waters et al., 2006). The vertical resolution is 1.3 – 3.6 km (Lambert et al., 2020;
94 Livesey et al., 2020). Data were screened based on error parameters supplied with the data, rendering a large
95 fraction of the volcanic data invalid the first two weeks after the eruption. From the beginning of February 2022,
96 when our evaluation starts, erroneous data became scarce.

97 Stratospheric temperatures in the pressure range 100 – 1 hPa were obtained from the MLS, which were used
98 primarily to compute relative humidity and potential temperature. The latter allows analysis of transport in
99 relation to isentropic surfaces. The potential temperatures were also used as a common ground in comparisons
100 between MLS and CALIOP, where the native vertical scale of the former is atmospheric pressure and for the
101 latter geometric altitude.

102 **3 Results**

103 This work focuses on the stratospheric aerosol resulting from the HT-22 eruption. The altitude and latitude
104 distributions will be presented here together with the evolution of the stratospheric aerosol extinction
105 coefficients and AOD. However, we start by presenting stratospheric water vapor data from the HT-22 eruption

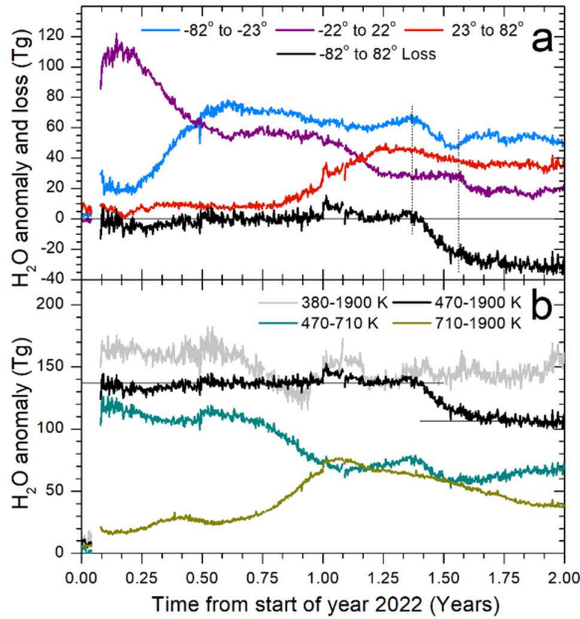
106 to highlight the contrasting evolution of the two volcanic components. Water vapor data are also used for
 107 computations on relative humidity and vertical air motions presented in the Discussion section.



108

109 **Figure 1.** Monthly averaged H₂O mass anomaly (Tg) against latitude and altitude with pixel size $(2.3 \pm 0.14) \times$
 110 $10^{16} \text{ m}^3 \text{ times } \cos(\Theta)$, where Θ is the latitude. Note that “2022 Jan 1 – 14” covers only the pre-eruption period 1
 111 – 14 January. Overlain isentropes in the range 380 – 1900 K are shown, where T_p is the potential temperature.

112 Note that the 380 K isentrope reaches below 100 hPa only in the tropics and that the 1900 K isentrope partly is
 113 found at pressures below 1 hPa. Vertical scale minor ticks: 1.5, 2.2, 3.2, 4.6, 6.8 and ten times these values.



114

115 **Figure 2.** Evolution of water vapor (H_2O) anomaly following the January 15, 2022, Hunga Tonga eruption. **a)**
 116 H_2O anomaly in three latitude intervals and loss of H_2O in a 4th latitude interval, all in the $470 < T_p < 1900$ K
 117 range (the deep BD branch). Vertical lines mark the main region of H_2O loss of the deep BD branch. **b)** H_2O
 118 anomaly in the latitude interval -82 to 82° in various potential temperature intervals (T_p). Horizontal lines show
 119 the average H_2O anomaly from end of January 2022 to mid-May 2023 (136.9 ± 0.2 (standard error) Tg) and from
 120 the beginning of October to the end of December 2023 (106.1 ± 0.3 Tg).

121 3.1 Water vapor

122 It has widely been reported about the record-breaking amounts of water vapor reaching the stratosphere
 123 following the HT-22 eruption (Millán et al., 2022; Schoeberl et al., 2022; Xu et al., 2022; Zhu et al., 2022;
 124 Nedoluha et al., 2024). Here we present the distribution related to isentropic surfaces in contrast to previous
 125 authors, in particular the fate of water that reaches the deep branch of the BD circulation, i.e., above the
 126 potential temperature (T_p) 470 K (Fueglistaler et al., 2009). Fig. 1 shows monthly mean water vapor mass
 127 anomalies for years 2022 and 2023, where the masses of year 2021 were subtracted, the exception being January
 128 2022 where only the days prior to the eruption are shown (January 1 – 14). The first two weeks after the
 129 eruption the MLS water vapor data from volcanic effluents frequently were erratic, probably due to high
 130 concentrations, and are not shown.

131 In February 2022 two layers appear, one minor in the shallow BD branch and the main layer in the deep BD
 132 branch, consistent with the reported eruption chronology (Gupta et al., 2022). The lower water vapor layer is
 133 spread rapidly latitudinally before it is transported below the lower atmospheric pressure limit used here (100
 134 hPa).

135 The first months after the eruption the water of the upper layer remains in the tropics, before a fraction clearly
136 visible in May 2022 is transported to the Southern extratropics (Figs. 1 and 2a). Towards the end of 2022
137 transport to the Northern extratropics starts, and in February 2023 the water from the HT-22 eruption covers
138 most of the globe. Later that year most of the water is found in the extratropics, whereas the water-rich air in the
139 tropics is replaced in the BD circulation by younger tropospheric air that is unaffected by the HT-22 eruption
140 (Figs. 1 and 2a). At the same time the water in the Southern extratropics of the deep BD branch approaches and
141 clearly passes-descends below the 470 K isentrope in May 2023 (Fig. 1 and 2a), consistent with the extratropical
142 downward motion of air.

143 The total amount of water vapor from the HT-22 eruption in the stratosphere at $T_p > 380$ K in the tropics and
144 100 hPa atmospheric pressure elsewhere, is 160 Tg. The mass in the deep BD branch, which is a part of the
145 previously mentioned layer, is 1378 Tg. After $\frac{3}{4}$ of a year these categories reach the same level (Fig. 2b),
146 implying that the lower water layer (injected below the deep BD branch) is transported down below the lower
147 limit in altitude (atmospheric pressure 100 hPa) of the data used here. The water vapor displays considerable
148 vertical transport in the deep BD branch. Dividing that branch into two T_p intervals (Fig. 2b) reveals a clear rise
149 in the amount of water in the upper interval in the last quarter of the year 2022. A small fraction of the water
150 vapor reached high altitudes in the tropics during the year 2023 (Fig. 1), and some even reached altitudes above
151 1 hPa atmospheric pressure (~ 48 km), i.e. the region of the stratopause, which can be seen in (supplementary
152 Fig. S1) but not in Fig. 1 because the much smaller integration volume increases the noise level.

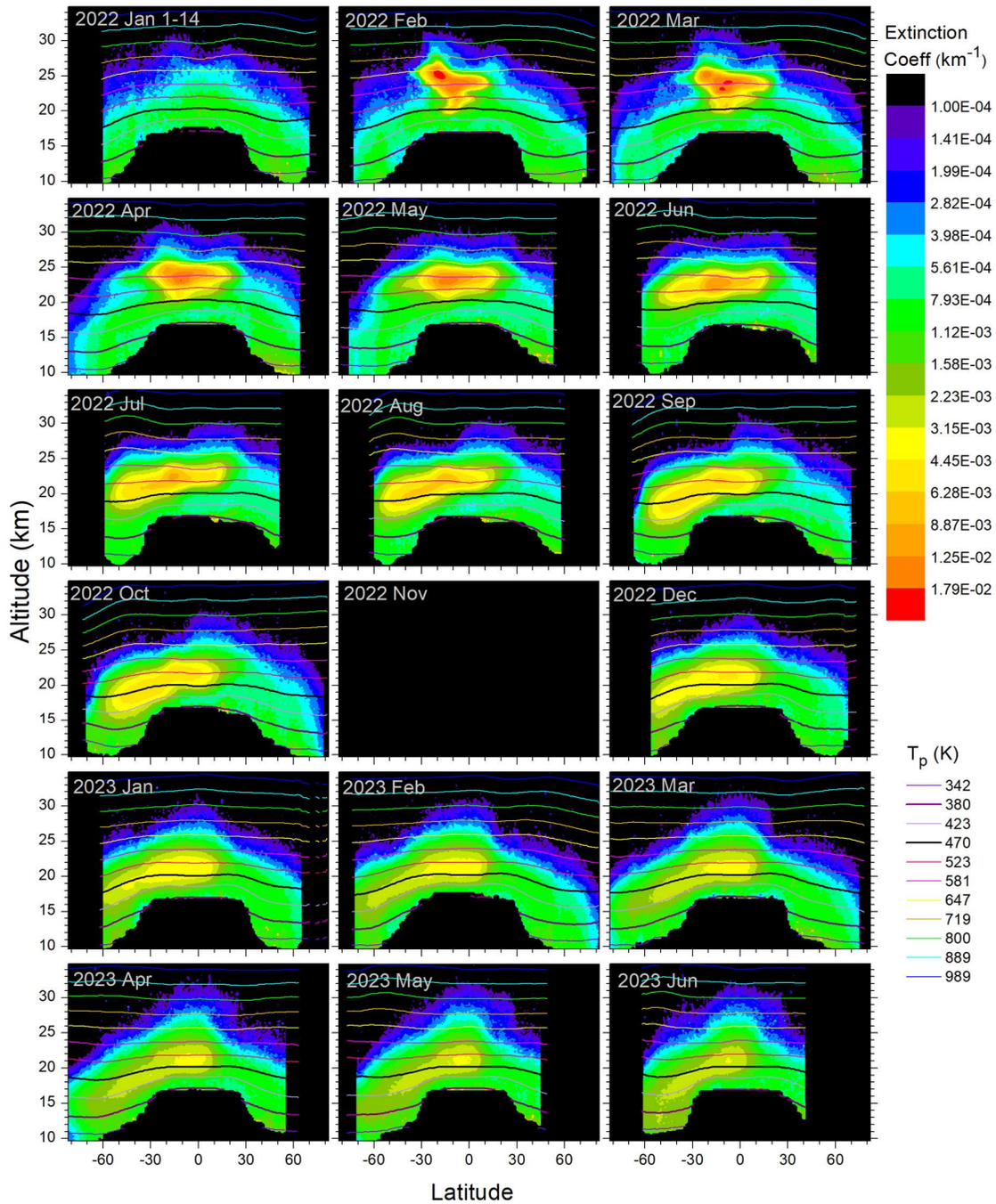
153 The water anomaly remained constant in the deep BD branch with only minor fluctuations from February 2022
154 to May 2023 (Fig. 2b), whereafter the anomaly is reduced by 23% due to transport to the shallow BD branch, a
155 level that remains until the end of 2023.

156 3.2 Aerosol

157 The evolution of the stratospheric AOD following the HT-22 eruption has been reported by several authors using
158 limb-viewing measurements (Bourassa et al., 2023; Sellitto et al., 2022; Taha et al., 2022) that suffer from event
159 termination (“saturation”) during the first months after strong volcanic or wildfire events (Fromm et al., 2014;
160 Chen et al., 2018; DeLand et al., 2019; Martinsson et al., 2022), and problems to measure the lower parts of the
161 stratosphere (Taha, 2020). Here we present results based on a nadir-viewing lidar technique (CALIOP) that is
162 better suited for measurements in dense aerosol layers because they do not suffer from saturation effects, and
163 attenuation of the lidar signal can be corrected for (Martinsson et al., 2022).

164 Just as for water vapor, we present monthly mean values of the aerosol distribution with overlaid isentropic
165 surfaces (Fig. 3). January 2022 aerosol data show conditions prior to the eruption. Initially (February – June
166 2022) almost all the HT-22 aerosol is found in the deep BD branch ($T_p > 470$ K). We identify downward motion
167 of the aerosol centroid in the tropics, the most intense part shifting from isentrope 581 to 523 K from March to
168 September 2022, despite the upward motion of air in the tropics as part of the BD circulation. This is caused by
169 gravitational settling, and the aerosol that reaches the Southern extratropics loses altitude even faster, aided by
170 downward air motion in the extratropics, leading to an increasing fraction of the aerosol in the shallow BD

171 branch from July 2022. The aerosol continues downwards, reaching the LMS (below 380 K) in December 2022
172 on its way out of the stratosphere.

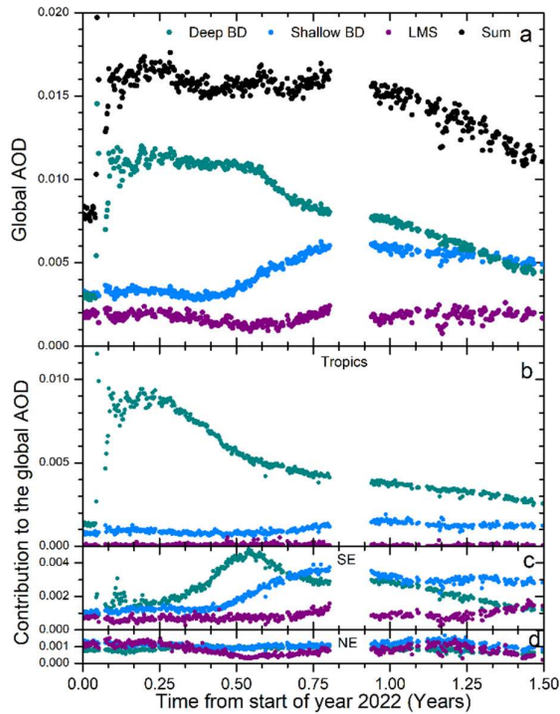


173

174 **Figure 3.** Monthly average extinction coefficients dependent on latitude and altitude with overlaid potential
175 temperature levels. Note that “2022 Jan 1 – 14” covers only the pre-eruption period 1 – 14 January.

176 Substantial amounts of aerosol entered the stratosphere because of the HT-22 eruption. The global average AOD
177 reached 0.016 (Fig. 4a), which is among the highest stratospheric aerosol loads since the 1991 Mt. Pinatubo

178 eruption. Already by the end of January, half a month after the eruption, the AOD level that remained for almost
 179 a year was reached. After that we see a decline where approximately half of the aerosol from the HT-22 eruption
 180 is removed during the first half-year of 2023. Almost the entire aerosol amount from HT-22 was found in the



181

182 **Figure 4. a)** Global average AOD of the stratosphere from the tropopause to 35 km altitude and -82 to 82° in
 183 latitude (Sum) with the sub layers: the tropopause to 380 K potential temperature (T_p) (LMS), 380 – 470 K T_p
 184 (shallow Brewer-Dobson (BD) branch) and T_p 470 K to 35 km altitude (deep BD). Latitude distributions of
 185 AOD **b)** tropics (-22 to 22°), **c)** Southern extratropics (SE) (-82 to -23°) and **d)** Northern extratropics (NE) (23
 186 to 82°). The AODs are related to the global scale, i.e. the sum of SE, tropics and NE graphs is the global AOD.

187 deep BD branch the first months after the eruption (Fig. 4a), in the tropics (Fig. 4b). We see transport to the
 188 Southern extratropics starting in April 2022 in the deep BD followed by downward motion to the shallow BD
 189 branch starting in June 2022 (Fig. 4c). Only a small fraction of the aerosol reached the Northern extratropics
 190 (Fig. 4d), in contrast to the transport of water vapor (Fig. 2a) that took place at a higher altitude (Fig. 1).

191 4 Discussion

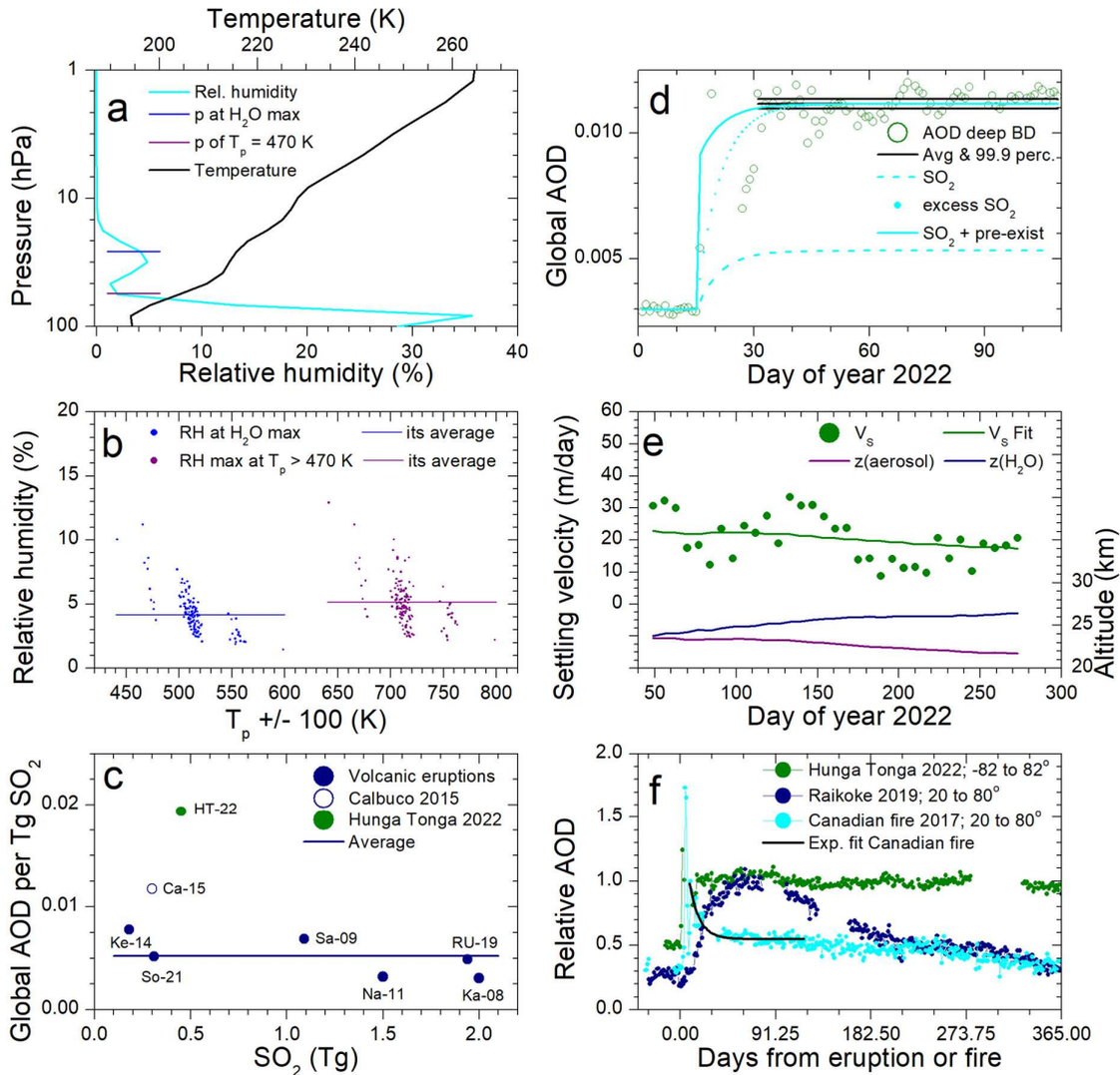
192 SO₂ emissions from HT-22 eruptions took place over a period from 19 December 2021 to 15 January 2022
 193 (Carn 2022). Most of these eruptions reached 15 – 18 km in altitude, whereas the main eruption’s umbrella
 194 cloud on 15 January 2022 reached 31 km with an overshooting plume reaching 55 – 58 km (Gupta et al., 2022).
 195 Based on several methods the total SO₂ emissions during this period is estimated to 0.6 – 0.7 Tg, and that of the
 196 main umbrella cloud, reaching deep into the stratosphere, contained 0.4 – 0.5 Tg SO₂ (Carn et al., 2022).
 197 Altitude-resolved SO₂ measurements from MLS find a similar SO₂ amount deep into the stratosphere (Millán
 198 2022). Compared with the SO₂ emissions, the stratospheric AOD generated by the HT-22 eruption is

199 unexpectedly high. Here we will discuss reasons for this seeming discrepancy, and we start by examining water
200 uptake as an explanation.

201 The temperature is rising with altitude in the stratosphere, making the air very dry after passing the tropical cold
202 point tropopause. The amount of water vapor injected by the HT-22 eruption is unprecedented in the modern
203 satellite era (Zhu et al., 2022). It has been suggested that hygroscopic growth could be an important process that
204 affects the aerosol particle size and light scattering (Legras et al., 2022; Sellitto et al., 2022). Here we investigate
205 the relative humidity by examining the five highest daily water vapor concentrations measured by the MLS
206 during February 2022 (140 MLS profiles), when the volcanic effluents were concentrated to a relatively small
207 volume. Based on MLS water vapor and temperature measurements the relative humidity was computed, where
208 the saturation water vapor pressures were obtained from Murphy and Koop (2005). Fig. 5a shows the average
209 relative humidity of the profiles (140) from February 2022. At the lowest altitudes, close to 100 hPa, the relative
210 humidity reaches 35% because of the low temperature (Fig. 5a, upper scale), and, to a smaller degree, the lower
211 volcanic layer (Fig. 1, February 2022). At higher altitude, the relative humidity rapidly declines as the
212 temperature increases, becoming close to zero at altitudes above 10 hPa. However, a peak appears at 30 hPa
213 caused by the main volcanic layer (above 470 K potential temperature) containing most of the stratospheric
214 water vapor from the HT-22 eruption (Fig. 2b). In the following discussion we concentrate on that layer. The
215 average positions of the 470 K isentrope and the peak water vapor concentration are shown in Fig. 5a, where the
216 shift of the maximum relative humidity from the peak water vapor concentration is caused by the temperature
217 gradient. The relative humidity at the peak water vapor concentration as well as the maximum relative humidity
218 of all the 140 measurements are shown in Fig. 5b (note the shift of ± 100 K in potential temperature to separate
219 the two categories). The measurements of each of the two categories appear in groups depending on the altitude
220 (or pressure level) of the water vapor layer. The maximum relative humidity above the 470 K isentrope is 13%,
221 and that of the peak water vapor is 11%, whereas the averages are 5.1 and 4.2%. Such low relative humidities
222 causes no or modest hygroscopic growth (Winkler, 1973) that affects particle size or light scattering only to a
223 small degree.

224 Several authors regard the aerosol from the HT-22 eruption as a sulfate aerosol (Khaykin et al., 2022; Legras et
225 al., 2022; Sellitto et al., 2022; Taha et al., 2022; Zhu et al., 2022; Bernath et al., 2023; Duchamp et al., 2023;
226 Kahn et al., 2024; Sellitto et al., 2024), although with questions on the relatively small amount of SO₂ emitted in
227 relation to the AOD level (Carn et al., 2022). Here we will investigate this relation in more detail by forming the
228 ratio of the maximum global stratospheric AOD rise above the pre-eruption AOD to the amount of SO₂ emitted
229 by eight recent volcanic eruptions (Table 1 and Fig. 5c). This ratio is approximately 0.005 Tg⁻¹ for most of the
230 eruptions, whereas the Calbuco (Ca-15) and HT-22 deviate by having higher AOD per SO₂ mass emitted. Most
231 of these volcanic eruptions showed depolarization ratio less than 0.05 (Hoffmann et al., 2010; O'Neill et al.,
232 2012; Zhuang & Yi 2016; Voudouri et al., 2023) typical of aerosol dominated by spherical sulfuric acid
233 particles. Volcanic ash settles rapidly by gravitation, but a fraction can remain for months in the stratosphere
234 (Andersson et al., 2013). Vernier et al. (2016) found that this can affect stratospheric AOD, detecting elevated
235 depolarization ratio (0.05) a month after the Kelut eruption (Ke-14). The depolarization ratio of the aerosol from
236 the Ca-15 eruption was much higher (0.18) a month after the eruption (Klekociuk et al., 2020) thus indicating a
237 strong influence from ash on the AOD that likely explains the strong deviation in AOD-to-SO₂ ratio from the

238 other eruptions. Ca-15 was therefore not included in the average AOD-to-SO₂ ratio calculated here. The HT-22
 239 eruption has the highest AOD-to-SO₂ ratio but low depolarization ratio (supplementary Fig. S2), thus high ash
 240 concentration is not a valid explanation (Gupta et al., 2022; Carn et al., 2022; Legras et al., 2022).



241

242 **Figure 5.** Stratospheric characteristics after the HT-22 eruption. **a)** Average relative humidity (RH) and
 243 temperature of the five daily H₂O profiles with the highest concentration during February 2022. **b)** RH at the
 244 maximum H₂O concentration and maximum RH at potential temperatures > 470 K of all the profiles mentioned
 245 in (a) with average RH of 4.2 and 5.1%, respectively. The potential temperature (T_p) was shifted ± 100 K to
 246 separate the two groups of data. **c)** Global AOD per Tg SO₂ emitted by recent volcanic eruptions related to SO₂,
 247 the average being 0.0052 global AOD per Tg SO₂ (see Table 1). **d)** AOD in the upper BD branch with 99.9
 248 percentile of the average marked and reported SO₂ of 0.45 Tg (Carn et al., 2022) converted to AOD according to
 249 (c) (broken line), and the dotted line tests the evolution using an excess of 1.1 Tg SO₂ to reach the measured
 250 AOD. The full cyan line displays the SO₂ AOD (broken line) added by an assumed AOD from pre-existing non-
 251 sulfate aerosol from the eruption to reach the measured AOD. **e)** Aerosol gravitational settling velocity (V_s) and

252 fit (equivalent aerodynamic diameter 1.1 μm) and average altitudes (z ; right scale) of the HT-22 aerosol and
 253 water vapor at latitudes -14 to -6°. **f**) Normalized stratospheric AOD evolution during one year for one wildfire
 254 event (Martinsson et al., 2022) and two volcanic eruptions.

255 **Table 1.** Recent volcanic eruptions with SO₂ emissions, global stratospheric optical depths (AOD) and literature
 256 references.

Date	Eruption	Short name	SO ₂ (Tg)	SO ₂ references	Global AOD ^a	Depolarization Ratio references
2008-08-07	Kasatochi	Ka-08	2	Yang et al., 2010	0.0061	Hoffmann et al., 2010
2009-06-12	Sarychev	Sa-09	1.09	Sandvik et al., 2021	0.0075	O'Neill et al., 2012
2011-06-12	Nabro	Na-11	1.5	Clarisse et al., 2012	0.0048	Zhuang & Yi 2016
2014-02-14	Kelut	Ke-14	0.18	Li et al., 2017	0.0014	Vernier et al., 2016
2015-04-23	Calbuco	Ca-15	0.3	Pardini et al., 2018	0.0035	Klekociuk et al., 2020
2019-06-22	Raikoke		1.5			
2019-06-26	Ulawun	RU-19	0.14	Kloss et al., 2021	0.0095	Voudouri et al., 2023
2019-08-03	Ulawun		0.3			
2021-04-10	Soufriere	So-21	0.31	Taylor et al., 2023	0.0016	^b Lidar browse images
2022-01-15	Hunga Tonga	HT-22	0.45	Carn et al., 2022	0.0087	This work

257 ^a) Global stratospheric AOD maximum increase due to the eruptions. References: Friberg et al., 2018 and this
 258 work (2019 – 2023)

259 ^b) Lidar Level 1 Browse Images - 2021-04-26 09:42:19Z - Section 1 ([https://www-
 260 calipso.larc.nasa.gov/products/lidar/browse_images/std_v451_index.php](https://www-calipso.larc.nasa.gov/products/lidar/browse_images/std_v451_index.php))

261 We adopt the central estimate of Carn et al. (2022), i.e., 0.45 Tg SO₂ with an e-folding time of ~6 days. The e-
 262 folding time is unusually short for stratospheric conditions, probably due to elevated water vapor concentrations
 263 (Carn et al., 2022). Fig. 5d shows the AOD, with double-sided 99.9% confidence interval of the mean in the
 264 deep BD branch, where all the aerosol from the HT-22 eruption was injected (Fig. 4a). Using the AOD-to-SO₂
 265 ratio based on six volcanic eruptions (Fig. 5c) to estimate the AOD based on the SO₂ emissions, we end up with
 266 far too low AOD (Fig. 5d, broken line). To investigate the timing, we added 1.1 Tg excess SO₂ to reach the
 267 measured AOD while preserving the measured e-folding time (dotted line). The excess SO₂ reaches into the
 268 99.9% confidence interval of the average AOD after approximately 50% longer time from the eruption
 269 compared to the time required for CALIOP to record a stable AOD. It is thus unlikely that the aerosol from the
 270 HT-22 eruption was formed from SO₂ conversion alone, mainly because of the low SO₂ emissions, but also
 271 because of the timing. Other material must have been present already the first days after the eruption. Making
 272 use of the AOD-to-SO₂ ratio from Fig. 5c, adding ~~pre-existing non-sulfate~~ aerosol from the HT-22 eruption
 273 ~~adjusted~~ to obtain the measured AOD and using the measured SO₂ mass and e-folding time, results in the cyan
 274 full line in Fig. 5d. Such a combination of ~~pre-existing non-sulfate~~ aerosol from the eruption and SO₂ conversion
 275 is consistent with the 99.9% confidence interval of the AOD average.

276 The next question is what is the source of the ~~non-sulfate~~ aerosol ~~that existing was present~~ before the conversion
 277 of SO₂? We have no measurements of the aerosol composition to aid in this respect. From the depolarization
 278 ratio (supplementary Fig. S2) we can rule out significant fractions of volcanic ash, which is also supported by
 279 other measurements (Gupta et al., 2022; Carn et al., 2022). To find another plausible source of the ~~pre-
 280 existing non-sulfate~~ aerosol we consider the intense sea – volcanism interaction during the HT-22 eruption
 281 (Seabrook et al., 2023; Clare et al., 2023, Mastin et al., 2024, Millán et al., 2022) causing enhanced bubble
 282 bursting (Keene et al., 2007) and/or explosive superheated water. Such events are not only sources of water

283 vapor but also releases the entire sea water substance to the atmosphere that includes sea salts. High
284 concentrations of sea salt in volcanic ash fallout from the HT-22 eruption has been documented (Colombier et
285 al., 2023). Sea salt particles enter the volcanic column together with the water vapor. As the particles are
286 hygroscopic, they readily serve as condensation nuclei in cloud formations as the air cools on the way up to the
287 stratosphere. In the competition for water, preferentially large particles are scavenged in cloud formations prior
288 to the formation of precipitation. This leaves the smaller particles as an interstitial aerosol (Martinsson et al.,
289 1999). The amount of aerosol from the eruption existing-present before the SO₂ conversion (Fig. 5d) would
290 correspond to aerosol formation from 1.1 Tg SO₂ based on the AOD-to-SO₂ ratio (Fig. 5c). Using this number as
291 a coarse estimate we can compare it with the amount of water injected into the deep BD branch (137 Tg; Fig.
292 2b). With the typical salinity of sea water (35 g/kg) that amount of water corresponds to 4.8 Tg of sea salt,
293 which is four times the coarse estimate of pre-existing-non-sulfate aerosol mass.

294 Besides the water from enhanced bubble bursting induced by volcanoclastic density currents or explosive
295 superheated water, water evaporates directly from a heated ocean without sea salt emissions. Additional
296 quantitative uncertainties pertain to the relative losses of water and sea salt to precipitation. Given the orders of
297 magnitude of these estimates we can from this standpoint conclude that aerosol formation from strong sea –
298 volcanism interaction is a plausible source of a large fraction of the stratospheric aerosol from the HT-22
299 eruption. However, we also need to consider the low depolarization ratio of the HT-22 aerosol. Cubic sodium
300 chloride particles can according to modeling show depolarization ratios in the range 0 to approximately 0.25
301 with strong dependence on the particle size, being close to 0 for particle volume mean diameters less than 0.7 –
302 0.8 μm before it gradually increases (Murayama et al., 1999; Haerig et al., 2017). The ageing of sea salt particles
303 in the atmosphere tends to round the particles (Adachi and Buseck, 2015) thus reducing depolarizations. To
304 further investigate this matter, we need to consider the particle size distribution.

305 Several authors have reported on the stratospheric aerosol particle size following the HT-22 eruption, i.e., 0.6 –
306 1 μm diameter (Boichu et al., 2023), 0.8 μm (Duchamp et al., 2023) and 2 – 3 μm (Legras et al., 2022). Whereas
307 the former two estimates show good agreement, the latter, based on estimating the gravitational settling velocity,
308 stands out by finding the particles to be larger than the other estimates. We used the same method as Legras et
309 al., (2022) to estimate the settling velocity: $V(\text{sedimentation}) = V(\text{aerosol}) - V(\text{air})$, where V is the vertical
310 velocity, $V(\text{aerosol})$ the observed weekly change in the aerosol centroid altitude and $V(\text{air})$ is estimated from the
311 weekly change in the altitude of the water vapor centroid. Applying a 3-week moving average dampened
312 variations in settling velocity leading to Fig. 5e. The gravitational settling velocity varies around the value 20
313 m/day, agreeing well with the results of Legras et al. (2022) whereas the conversion to particle size differs. The
314 settling velocity of a given particle depends on the pressure and temperature because of the air viscosity and the
315 Cunningham slip correction factor's dependence on the mean free path of the air. We computed the particle size
316 that best fits the weekly settling velocity observations. Fig. 5e shows decreased settling velocity as the aerosol
317 falls to lower altitude. We found that the equivalent aerodynamic diameter was 1.1 μm, which is based on the
318 assumptions of a spherical particle shape and particle density of 1 g/cm³. The low depolarization ratio
319 (supplementary Fig. S2) validates the first assumption. The density of the particles is not known a priori.
320 However, the low relative humidity (Figs. 5a and b) results in concentrated solution drops of sulfuric acid and
321 sea salts, having density clearly exceeding 1 g/cm³, e.g., a 76.5% sulfuric acid – water solution has a density of

322 1.75 g/cm³ at stratospheric conditions (Myhre et al., 1998). Applying that density results in 0.70 μm geometric
323 diameter and changing the density to 1.5 and 2 g/cm³ results 0.81 and 0.62 μm diameter, respectively, which is
324 in good agreement with estimates based on other methods. Based on our results and others (Boichu et al., 2023;
325 Duchamp et al., 2023) we conclude that the HT-22 aerosol is submicron in diameter, in between stratospheric
326 background and Mt. Pinatubo particle sizes (Bauman et al., 2003; Wilson et al., 2008). The depolarization ratio
327 was low already the first days after the eruption when only a small fraction of the SO₂ conversion was
328 completed. However, the particle size of the HT-22 aerosol falls in the region where the depolarization ratio for
329 cubic sodium chloride particles is small, thus not contradicting that sea salt from volcanism – sea interaction
330 was a strong source of the HT-22 aerosol.

331 The water vapor injected into the deep BD branch remained in the stratosphere for the full two years of this
332 study, although 23% was transported from the deep BD branch to the shallow one 1.5 years after the eruption
333 (Fig. 2b). The stratospheric AOD remained almost constant for one year before starting to decline (Fig. 4a).
334 Because of gravitational settling aerosol remains in the stratosphere for a shorter time than gases with low
335 chemical reactivity. The combined effect of the 2019 Raikoke and Ulawun eruptions on the maximum global
336 stratospheric AOD is the highest observed for recent eruptions (Table 1) when also the lowest part of the
337 stratosphere are accounted for. The peak AOD from HT-22 eruption is slightly lower. However, the long
338 duration of the AOD from the HT-22 eruption, caused by the powerful eruption placing the effluents in the deep
339 BD branch in the tropics, makes it the most important in terms of stratospheric AOD since the 1991 eruption of
340 Mt. Pinatubo (Fig. 5f). The first year after the eruption the AOD was 0.016. Subtracting average background
341 AOD (Friberg et al., 2018) the stratospheric global mean AOD from the HT-22 eruption becomes 0.010. This
342 corresponds to -0.24 W/m² in global stratospheric total volcanic effective radiative forcing during the first year
343 after the eruption, according to results based on volcanic activity years 1979 to 2015 (Schmidt et al., 2018).

344 The HT-22 was the last major volcanic eruption to be studied based on data from the CALIOP lidar aboard the
345 CALIPSO satellite that ended its mission in June 2023. This is by far the most efficient method for studies of the
346 initial months of stratospheric aerosol formation following volcanic eruptions and wildfires, because of its
347 brilliant vertical resolution and optically short vertical path. Limb-viewing techniques suffer from event
348 termination (saturation) during 2 – 3 months after a major stratospheric aerosol event (Martinsson et al., 2022;
349 Fromm et al., 2014; Chen et al., 2018; DeLand, 2019). Fig. 5f illustrates the importance of CALIOP by showing
350 the AOD of two volcanic eruptions and one wildfire. Conversion of SO₂ formed the Raikoke aerosol, resulting
351 in 2 – 3 months delay before the AOD peaked which is the case for most volcanic eruptions (Friberg et al.,
352 2018). In contrast, sea salt aerosol from HT-22 ~~existing-present~~ before the SO₂ conversion dominated its AOD
353 and we observed the maximum already after two weeks. That was the time required for the aerosol to become
354 dispersed enough to allow approximately ten CALIOP measurements per day in the volcanic effluents, thereby
355 reducing the uncertainty in the daily average. Another special case was the 2017 Canadian wildfire where we
356 observed a strong and rapid decline of the stratospheric AOD (Fig. 5f) indicative of photolytic loss of organic
357 aerosol (Martinsson et al., 2022). A study of the 2019/2020 Australian wildfire showed similar losses, where
358 also a complex feed of wildfire aerosol from the upper troposphere during 1 – 2 weeks after the fire was
359 identified (Friberg et al., 2023), thanks to the mentioned special properties of the CALIOP instrument. The

360 decommissioning of the ageing CALIOP in June 2023 severely diminishes future studies of aerosol formation
361 and losses in the stratosphere, prompting the need for new satellite-based lidar systems.

362 **4 Conclusions**

363 Aerosol and water vapor in the stratosphere emanating from the 15 January 2022 eruption in Hunga Tonga (HT-
364 22) is investigated using satellite-based instruments CALIOP and MLS. Most of its effluents were injected into
365 the deep branch of the stratospheric Brewer-Dobson (BD) circulation.

366 A small fraction of the record-breaking water vapor injections into the deep BD branch reached up to the
367 stratopause after 1.25 years in the stratosphere, whereas 23% was transported down to the shallow BD branch as
368 the water vapor spread vertically. The water vapor injected into the deep BD branch remained in the stratosphere
369 for the full two years of this study. The water vapor from the HT-22 eruption in the southern tropics steadily
370 increased its latitudinal coverage, first to the southern midlatitudes. After a year most of the global stratosphere
371 was covered with water vapor from the HT-22 eruption, before a reduction of the tropical stratospheric
372 concentration appeared as the BD circulation brought tropospheric air that was unaffected by the HT-22
373 eruption.

374 The aerosol and its precursor gases were initially at the same altitude as the water vapor from the HT-22
375 eruption, but gravitational settling of the aerosol particles gradually opened a gap in altitude which resulted in
376 the aerosol from the HT-22 eruption mainly appearing in the tropics and the southern hemisphere. The
377 stratospheric aerosol optical depth (AOD) remained constant for a year after the eruption, before transport out of
378 the stratosphere started. At the time of the decommission of the CALIOP instrument in June 2023, 50% of the
379 aerosol from the HT-22 eruption had been removed from the stratosphere.

380 The AOD level of the stratosphere was established already 2 weeks after the eruption and was unexpectedly
381 high for a modest injection of 0.4 – 0.5 Tg SO₂. Given the exceptional water vapor amounts from the HT-22
382 eruption, we investigated if hygroscopic growth affected the aerosol optical properties. Despite the record-
383 breaking water vapor emissions, the average relative humidity remained below 5% in the dry stratosphere,
384 causing no or limited hygroscopic growth.

385 The gravitational settling velocity of the aerosol is estimated from the altitude evolution to ~20 m/day,
386 corresponding to an equivalent aerodynamic diameter of 1.1 μm at the altitude of the aerosol layer. Assuming
387 density of concentrated solution drops of 1.5 – 2 g/cm³ the geometrical diameter becomes 0.6 – 0.8 μm.

388 Comparing eight recent volcanic eruptions we find that the global AOD per mass of SO₂ emitted from the HT-22
389 eruption is 4 times that of most other eruptions. The amount of SO₂ and ash emitted to the stratosphere was
390 unusually small for an eruption with volcanic explosivity index (VEI) of 6. Aerosol formation from intense
391 volcano – sea interaction provides sea salt aerosol as a plausible explanation for the unexpectedly high AOD.

392 The maximum global stratospheric AOD following the HT-22 eruption is among the highest observed in more
393 than 30 years. The injection in the deep branch of DB circulation prolonged the perturbation of the stratospheric

394 aerosol, making the HT-22 eruption the largest aerosol event since that of Mt. Pinatubo in 1991. The 1-year
395 average global AOD of 0.01 from the HT-22 eruption can be estimated to -0.24 W/m² in global stratospheric
396 total volcanic effective radiative forcing.

397 **References**

- 398 Adachi K. and P.R. Buseck, Changes in shape and composition of sea-salt particles upon aging in an urban
399 atmosphere. *Atmos. Environ.* 100, 1-9, <http://dx.doi.org/10.1016/j.atmosenv.2014.10.036>,
400 2015.
- 401 Andersson, S. M., Martinsson, B. G., Friberg, J., Brenninkmeijer, C. A. M., Rauthe-Schöch, A., Hermann,
402 M., van Velthoven, P. F. J., and Zahn, A.: Composition and evolution of volcanic aerosol from
403 eruptions of Kasatochi, Sarychev and Eyjafjallajökull in 2008–2010 based on CARIBIC observations,
404 *Atmos. Chem. Phys.*, 13, 1781–1796, <https://doi.org/10.5194/acp13-1781-2013>, 2013.
- 405 Andersson, S. M., Martinsson, B. G., Vernier, J. P., Friberg, J., Brenninkmeijer, C. A. M., Hermann, M., van
406 Velthoven, P. F. J., and Zahn, A.: Significant radiative impact of volcanic aerosol in the lowermost
407 stratosphere, *Nat. Commun.*, 6, 1–8, <https://doi.org/10.1038/ncomms8692>, 2015.
- 408 Bauman J. J., P. B. Russell, M. A. Geller, and P. Hamill, A stratospheric aerosol climatology from SAGE II
409 and CLAES measurements: 2. Results and comparisons, 1984–1999, *J. Geophys. Res.*, 108(D13),
410 4383, doi:10.1029/2002JD002993, 2003.
- 411 Bernath P., C. Boone, A. Pastorek, D. Cameron and M. Lecours, Satellite characterization of global
412 stratospheric sulfate aerosols released by Tonga volcano. *J. Quant. Spectrosc. Rad. Transf.* 299,
413 108520, 2023.
- 414 Boichu M., R. Grandin, L. Blarel, B. Torres, Y. Derimian, P. Goloub, C. Brogniez, I. Chiapello, O. Dubovic,
415 T. Mathurin, N. Pascal, M. Patou and J. Riedi, Growth and Global Persistence of Stratospheric Sulfate
416 Aerosols From the 2022 Hunga Tonga–Hunga Ha'apai Volcanic Eruption. *J. Geophys. Res. Atmos.* 128,
417 e2023JD039010. <https://doi.org/10.1029/2023JD039010>, 2023.
- 418 Bourassa, A. E., Zawada, D. J., Rieger, L. A., Warnock, T. W., Toohey, M., & Degenstein, D. A.,
419 Tomographic retrievals of Hunga Tonga-Hunga Ha'apai volcanic aerosol. *Geophysical Research*
420 *Letters*, 50, e2022GL101978. <https://doi.org/10.1029/2022GL101978>, 2023.
- 421 Carn S.A., N.A. Krotkov, B.L. Fisher and C. Li, Out of the blue: Volcanic SO₂ emissions during the 2021–
422 2022 eruptions of Hunga Tonga—Hunga Ha'apai (Tonga). *Front. Earth Sci.* 10:976962. doi:
423 10.3389/feart.2022.976962, 2022.
- 424 Carr J.L., A. Horváth, D.L. Wu, and M.D. Friberg, Stereo plume height and motion retrievals for the
425 record-setting Hunga TongaHunga Ha'apai eruption of 15 January 2022. *Geophysical Research Letters*
426 49, e2022GL098131, <https://doi.org/10.1029/2022GL098131>, 2022.
- 427 Chen, Z., Bhartia, P. K., Loughman, R., Colarco, P., and De Land, M.: Improvement of stratospheric
428 aerosol extinction re trieval from OMPS/LP using a new aerosol model, *Atmos. Meas. Tech.*, 11, 6495–
429 6509, <https://doi.org/10.5194/amt-11-6495-2018>, 2018.
- 430 Clare M.A., I.A. Yeo, S. Watson, R. Wysoczanski, S. Seabrook, K. Mackay, J.E. Hunt, E. Lane, P.J. Talling,
431 E. Pope, S. Cronin, M. Ribó, T. Kula, D. Tappin, S. Henrys, C. de Ronde, M. Urlaub, S. Kutterolf, S.

432 Fonua, S. Panuve, D. Veverka, R. Rapp, V. Kamalov and M. Williams, Fast and destructive density
433 currents created by ocean-entering volcanic eruptions. *Science* 381, 1085–1092, 2023.

434 Clarisse L., D. Hurtmans, C. Clerbaux, J. Hadji-Lazaro, Y. Ngadi, and P.-F. Coheur, Retrieval of sulphur
435 dioxide from the infrared atmospheric sounding interferometer (IASI). *Atmos. Meas. Tech.*, 5, 581–
436 594, 2012.

437 Colombier M., I. A. Uktins, S. Tegtmeier, B. Scheu, S. J. Cronin, S. Thivet, J. Paredes-Mariño, C.
438 Cimarelli, K.-U. Hess, T. Kula, F.H. Latu'ila and D. B. Dingwell, Atmosphere injection of sea salts
439 during large explosive submarine volcanic eruptions. *Scientific Reports* 13:14435,
440 <https://doi.org/10.1038/s41598-023-41639-8>, 2023.

441 DeLand, M.: Readme document for the Soumi-NPP OPMS LP L2 AER675 Daily product, Goddard Earth
442 Sciences Data and Information Services Center (GES DISC), <http://disc.gsfc.nasa.gov> (last access:
443 October 2021), 2019.

444 Duchamp C., F. Wrana, B. Legras, P. Sellitto, R. Belhadji and C. von Savigny, Observation of the Aerosol
445 Plume From the 2022 Hunga Tonga—Hunga Ha'apai Eruption With SAGE III/ISS. *Geophys. Res. Lett.*
446 50, e2023GL105076, <https://doi.org/10.1029/2023GL105076>, 2023.

447 Friberg J., Martinsson B. G., Andersson S. M., Brenninkmeijer C. A. M., Hermann M., Van Velthoven P. F.
448 J., and Zahn A., Sources of increase in lowermost stratospheric sulphurous and carbonaceous aerosol
449 background concentrations during 1999–2008 derived from CARIBIC flights, *Tellus B*, 66, 23428,
450 <https://doi.org/10.3402/tellusb.v66.23428>, 2014.

451 Friberg, J., Martinsson, B. G., Andersson, S. M., and Sandvik, O. S.: Volcanic impact on the climate– the
452 stratospheric aerosol load in the period 2006–2015, *Atmos. Chem. Phys.*, 18, 11149–11169,
453 <https://doi.org/10.5194/acp-18-11149-2018>, 2018.

454 Friberg J., B.G. Martinsson, and M.K. Sporre, Short- and long-term stratospheric impact of smoke from the
455 2019–2020 Australian wildfires. *Atmos. Chem. Phys.*, 23, 12557–12570, 2023
456 <https://doi.org/10.5194/acp-23-12557-2023>, 2023.

457 Fromm M., Lindsey D. T., Servranckx, R., Yue G., Trickl T., Sica R., Doucet P., and Godin-Beekmann S.,
458 The untold story of pyrocumulonimbus, *B. Am. Meteorol. Soc.*, 91, 1193–1209, 2010.

459 Fromm, M., Kablick III, G., Nedoluha, G., Carboni, E., Grainger, R., Campbell, J., and Lewis, L.:
460 Correcting the record of volcanic stratospheric aerosol impact: Nabro and Sarychev Peak, *J. Geophys.*
461 *Res.-Atmos.*, 119, 1–22, <https://doi.org/10.1002/2014JD021507>, 2014.

462 Fueglistaler S., Dessler A. E., Dunkerton T. J., Folkins I., Fu Q., and Ote P. W., Tropical tropopause layer,
463 *Rev. Geophys.*, 47, RG1004, <https://doi.org/10.1029/2008RG000267>, 2009.

464 Garofalo, L. A., Levin, E. J. T., Campos, T., Kreidenweis, S. N., and Farmer, D. K.: Emission and evolution
465 of submicron organic aerosol in smoke from wildfires in the western United States. *ACS Space Chem.*,
466 3, 1237–1247, 2019.

467 Gupta A.K., R. Bennartz, K.E. Fauria and T. Mittal, Eruption chronology of the December 2021 to January
468 2022 Hunga Tonga-Hunga Ha'apai eruption sequence. *Comm. Earth Environm.* 3:314,
469 <https://doi.org/10.1038/s43247-022-00606-3>, 2022.

470 Haarig M., A. Ansmann, J. Gasteiger, K. Kandler, D. Althausen, H. Baars, M. Radenz, and D.A. Farrell,
471 Dry versus wet marine particle optical properties: RH dependence of depolarization ratio, backscatter,

472 and extinction from multiwavelength lidar measurements during SALTRACE. *Atmos. Chem. Phys.*,
473 17, 14199–14217, <https://doi.org/10.5194/acp-17-14199-2017>, 2017.

474 Haarig, M., Ansmann, A., Baars, H., Jimenez, C., Veselovskii, I., Engelmann, R., and Althausen, D.:
475 Depolarization and lidar ratios at 355, 532, and 1064nm and microphysical properties of aged
476 tropospheric and stratospheric Canadian wildfire smoke, *Atmos. Chem. Phys.*, 18, 11847–11861,
477 <https://doi.org/10.5194/acp-18-11847-2018>, 2018.

478 Hoffmann A., C. Ritter, M. Stock, M. Maturilli, S. Eckhardt, A. Herber, and R. Neuber, Lidar
479 measurements of the Kasatochi aerosol plume in August and September 2008 in Ny-Ålesund,
480 Spitsbergen. *J. Geophys. Res.*, 115, D00L12, doi:10.1029/2009JD013039, 2010.

481 Hu, Q., Goloub, P., Veselovskii, I., Bravo-Aranda, J.-A., Popovici, I. E., Podvin, T., Haefelin, M., Lopatin,
482 A., Dubovik, O., Pietras, C., Huang, X., Torres, B., and Chen, C.: Long-range transported Canadian
483 smoke plumes in the lower stratosphere over northern France, *Atmos. Chem. Phys.*, 19, 1173–1193,
484 <https://doi.org/10.5194/acp-19-1173-2019>, 2019.

485 Kahn, R. A., Limbacher, J. A., Junghenn Noyes, K. T., Flower, V. J. B., Zamora, L. M., & McKee, K. F.,
486 Evolving particles in the 2022 Hunga Tonga—Hunga Ha'apai volcano eruption plume. *J. Geophys.*
487 *Res.*, 129, e2023JD039963. <https://doi.org/10.1029/2023JD039963>, 2024.

488 Keene W.C., H. Maring, J.R. Maben, D.J. Kieber, A.A.P. Pszenny, E.E. Dahl, M.A. Izaguirre, A.J. Davis,
489 M.S. Long, X. Zhou, L. Smoydzin, and R. Sander, Chemical and physical characteristics of nascent
490 aerosols produced by bursting bubbles at a model air-sea interface, *J. Geophys. Res.*, 112, D21202,
491 doi:10.1029/2007JD008464, 2007.

492 Khaykin S., A. Podglajen, F. Ploeger, J.-U. Groöf, F. Tence, S. Bekki, K. Khlopenkov, K. Bedka, L.
493 Rieger,, A. Baron, S. Godin-Beekmann, B. Legras, P. Sellitto, T. Sakai, J. Barnes, O. Uchino, I. Morino,
494 T. Nagai, R. Wing, G. Baumgarten, M. Gerding, V. Dufлот, G. Payen, J. Jumelet, R. Querel, B. Liley, A.
495 Bourassa, B. Clouser, A. Feofilov, A. Hauchecorne and F. Ravetta, Global perturbation of stratospheric
496 water and aerosol burden by Hunga eruption. *Comm. Earth Environm.*, 3:316,
497 <https://doi.org/10.1038/s43247-022-00652-x|www.nature.com/commsenv>, 2022.

498 Klekociuk A.R., D.J. Ottaway, A.D. MacKinnon, I.M. Reid, L.M. Twigger and S.P. Alexander, Australian
499 Lidar Measurements of Aerosol Layers Associated with the 2015 Calbuco Eruption. *Atmosphere* 11,
500 124, 2020.

501 Kloss C., G. Berthet, P. Sellitto, F. Ploeger, G. Taha, M. Tidiga, M. Eremenko, A. Bossolasco, F. Jégou, J.-
502 B. Renard, and B. Legras, Stratospheric aerosol layer perturbation caused by the 2019 Raikoke and
503 Ulawun eruptions and their radiative forcing. *Atmos. Chem. Phys.*, 21, 535–560, 2021.

504 Kremser, S., Thomason, L. W., von Hobe, M., Hermann, M., Deshler, T., Timmreck, C., Toohey, M.,
505 Stenke, A., Schwarz, J. P., Weigel, R., Fueglistaler, S., Prata, F. J., Vernier, J. P., Schlager, H., Barnes, J.
506 E., Antuña-Marrero, J. C., Fairlie, D., Palm, M., Mahieu, E., Notholt, J., Rex, M., Bingen, C.,
507 Vanhellemont, F., Bourassa, A., Plane, J. M. C., Klocke, D., Carn, S. A., Clarisse, L., Trickl, T., Neely,
508 R., James, A. D., Rieger, L., Wilson, J. C., and Meland, B.: Stratospheric aerosol– Observations,
509 processes, and impact on climate, *Rev. Geophys.*, 54, 278–335,
510 <https://doi.org/10.1002/2015RG000511>, 2016.

511 Lambert, A., Read, W., and Livesey, N.: MLS/Aura Level 2 Water Vapor (H₂O) Mixing Ratio V005,
512 Greenbelt, MD, USA, Goddard Earth Sciences Data and Information Services Center (GES DISC),
513 <https://doi.org/10.5067/Aura/MLS/DATA2508>, 2020.

514 Legras B., C. Duchamp, P. Sellitto, A. Podglajen, E. Carboni, R. Siddans, J.-U. Grooß, S. Khaykin, and F.
515 Ploeger, The evolution and dynamics of the Hunga Tonga–Hunga Ha’apai sulfate aerosol plume in the
516 stratosphere. *Atmos. Chem. Phys.*, 22, 14957–14970, 2022.

517 Li C., N.A. Krotkov, S. Carn, Y. Zhang, R.J.D. Spurr, and J. Joiner, New-generation NASA Aura Ozone
518 Monitoring Instrument (OMI) volcanic SO₂ dataset: algorithm description, initial results, and
519 continuation with the Suomi-NPP Ozone Mapping and Profiler Suite (OMPS). *Atmos. Meas. Tech.*, 10,
520 445–458, 2017.

521 Livesey, N. J., Read, W. G., Wagner, P. A., Froidevaux, L., Santee, M. L., Schwartz, M. J., Lambert, A.,
522 Manney, G. L., Valle, L. F. M., Pumphrey, H. C., Fuller, R. A., Jarnot, R. F., Knosp, B. W., and Lay,
523 R.R.: EOS MLS Version 5.0x Level2 and 3 data quality and description document, Tech. rep., Jet
524 Propulsion Laboratory D734 105336 Rev. A, <https://mls.jpl.nasa.gov/publications>, 2020.

525 Martinsson B.G., G. Frank, S.-I. Cederfelt, E. Swietlicki, O.H. Berg, J. Zhou, K.N. Bower, C. Bradbury, W.
526 Birmili, F. Stratmann, M. Wendisch, A. Wiedensohler, B.A. Yuskiewicz, Droplet nucleation and growth
527 in orographic clouds in relation to the aerosol population. *Atmos. Res.* 50, 289–315, 1999.

528 Martinsson, B. G., Brenninkmeijer, C. A. M., Cam, S. A., Hermann, M., Heue, K.P., van Velthoven, P. F. J.,
529 and Zahn, A.: Influence of the 2008 Kasatochi volcanic eruption on sulfurous and carbonaceous
530 aerosol constituents in the lower stratosphere, *Geophys. Res. Lett.*, 36, 1–5,
531 <https://doi.org/10.1029/2009GL038735>, 2009.

532 Martinsson, B. G., Friberg, J., Sandvik, O. S., Hermann, M., van Velthoven, P. F. J., and Zahn, A.:
533 Formation and composition of the UTLS aerosol, *Npj Climate and Atmospheric Science*, 2, 1–6,
534 <https://doi.org/10.1038/s41612-019-0097-1>, 2019.

535 Martinsson, B. G., Friberg, J., Sandvik, O. S., and Sporre, M. K.: Five-satellite-sensor study of the rapid
536 decline of wildfire smoke in the stratosphere, *Atmos. Chem. Phys.*, 22, 3967–3984,
537 <https://doi.org/10.5194/acp-22-3967-2022>, 2022.

538 Mastin L.G., A.R. Van Eaton and S.J. Cronin, Did steam boost the height and growth rate of the giant
539 Hunga eruption plume? *Bull. Volcanology* 86:64, <https://doi.org/10.1007/s00445-024-01749-1>, 2024.

540 Millán, L., Santee, M. L., Lambert, A., Livesey, N. J., Werner, F., Schwartz, M. J., Pumphrey H.C., Manney
541 G.L., Wang Y., Su H., Read W.G. and Froidevaux H.C., The Hunga Tonga-Hunga Ha'apai Hydration of
542 the Stratosphere. *Geophysical Research Letters*, 49, e2022GL099381. <https://doi.org/10.1029/2022GL099381>, 2022.

544 Murayama T., H. Okamoto, N. Kaneyasu, H. Kamataki, and K. Miura, Application of lidar depolarization
545 measurement in the atmospheric boundary layer: Effects of dust and sea-salt particles. *J. Geophys. Res.*
546 104, 31781-31792, 1999.

547 Murphy, D. M. and Koop, T., Review of the vapour pressures of ice and supercooled water for atmospheric
548 applications, *Q. J. Roy. Meteor. Soc.*, 131, 1539–1565, <https://doi.org/10.1256/qj.04.94>, 2005.

549 Murphy D. M., Cziczo D. J., Hudson P. K., and Thomson D. S., Carbonaceous material in aerosol particles
550 in the lower stratosphere and tropopause region, *J. Geophys. Res.*, 112, D04203,
551 <https://doi.org/10.1029/2006JD007297>, 2007.

552 Myhre C.E.L., C.J. Nielsen, and O.W. Saastad, Density and Surface Tension of Aqueous H₂SO₄ at Low
553 Temperature. *J. Chem. Eng. Data* 43, 617-622, 1998.

554 Nedoluha G.E., Gomez R.M., Boyd I., Neal H., Allen D.R. and Lambert A., The spread of the Hunga Tonga
555 H₂O plume in the middle atmosphere over the first two years since eruption. *J. Geophys. Res. Atmos.*
556 129, e2024JD040907. <https://doi.org/10.1029/2024JD040907>, 2024.

557 O'Neill N.T., C. Perro, A. Saha, G. Lesins, T. J. Duck, E. W. Eloranta, G. J. Nott, A. Hoffman, M. L.
558 Karumudi, C. Ritter, A. Bourassa, I. Abboud, S. A. Carn, and V. Savastiouk, Properties of Sarychev
559 sulphate aerosols over the Arctic, *J. Geophys. Res.*, 117, D04203, doi:10.1029/2011JD016838, 2012.

560 Pardini F., M. Burton, F. Arzilli, G. La Spina, M. Polacci, SO₂ emissions, plume heights and magmatic
561 processes inferred from satellite data: The 2015 Calbuco eruptions. *J. Volcanol. Geotherm. Res.* 361,
562 12–24, 2018.

563 Poli P. and N.M. Shapiro, Rapid characterization of large volcanic eruptions: measuring the impulse of the
564 Hunga Tonga Ha'apai explosion from teleseismic waves. *Geophys. Res. Letters* 49, e2022GL098123.
565 <https://doi.org/10.1029/2022GL098123>, 2022.

566 Proud S.R., A.T. Prata and S. Schmauss, The January 2022 eruption of Hunga Tonga-Hunga Ha'apai
567 volcano reached the mesosphere. *Science* 378, 554–557, 2022.

568 Sandvik O.S., J. Friberg, M.K. Sporre, and B.G. Martinsson, Methodology to obtain highly resolved SO₂
569 vertical profiles for representation of volcanic emissions in climate models. *Atmos. Meas. Tech.*, 14,
570 7153–7165, 2021.

571 Schmidt, A., Mills M. J., Ghan S., Gregory J. M., Allan R. P., Andrews T., Bardeen C.G., Conley A., Forster
572 P.M., Gettelman A., Portmann R.W., Solomon S. And Toon O.B., Volcanic radiative forcing from 1979
573 to 2015. *J. Geophys. Res. Atmos.* 123, 12,491–12,508. <https://doi.org/10.1029/2018JD028776>, 2018.

574 Schoeberl, M. R., Wang, Y., Ueyama, R., Taha, G., Jensen, E., and Yu, W., Analysis and impact of the
575 Hunga Tonga-Hunga Ha'apai stratospheric water vapor plume. *Geophysical Research Letters*, 49,
576 e2022GL100248. <https://doi.org/10.1029/2022GL100248>, 2022.

577 Seabrook S., K. Mackay, S.J. Watson, M.A. Clare, J.E. Hunt, I.A. Yeo, E.M. Lane, M.R. Clark, R.
578 Wysoczanski, A.A. Rowden, T. Kula, L.J. Hoffmann, E. Armstrong and M.J.M. Williams,
579 Volcaniclastic density currents explain widespread and diverse seafloor impacts of the 2022 Hunga
580 Volcano eruption. *Nat. Commun.*, 14:7881, <https://doi.org/10.1038/s41467-023-43607-2>, 2023.

581 Sellitto P., A. Podglajen, R. Belhadj, M. Boichu, E. Carboni, J. Cuesta, C. Duchamp, C. Kloss, R. Siddans,
582 N. Bègue, L. Blarel, F. Jegou, S. Khaykin, J.-B. Renard and B. Legras, The unexpected radiative
583 impact of the Hunga Tonga eruption of 15th January 2022. *Comm. Earth Environm.* 3:288
584 [|https://doi.org/10.1038/s43247-022-00618-z](https://doi.org/10.1038/s43247-022-00618-z), 2022.

585 Sellitto P., R. Siddans, R. Belhadj, E. Carboni, B. Legras, A. Podglajen, C. Duchamp, and B. Kerridge,
586 Observing the SO₂ and Sulfate Aerosol Plumes From the 2022 Hunga Eruption With the Infrared
587 Atmospheric Sounding Interferometer (IASI). *Geophys. Res. Lett.* 51, e2023GL105565.
588 <https://doi.org/10.1029/2023GL105565>, 2024.

589 Taha, G., OMPS-NPP L2 LP Aerosol Extinction Vertical Profile swath daily 3slit V2, Greenbelt, MD,
590 USA, Goddard Earth Sciences Data and Information Services Center (GES DISC),
591 <https://doi.org/10.5067/CX2B9NW6FI27>, 2020.

592 Taha, G., Loughman, R., Colarco, P. R., Zhu, T., Thomason, L. W., and Jaross, G., Tracking the 2022
593 Hunga Tonga-Hunga Ha'apai aerosol cloud in the upper and middle stratosphere using space-based
594 observations. *Geophysical Research Letters*, 49, e2022GL100091.
595 <https://doi.org/10.1029/2022GL100091>, 2022.

596 Taylor I.A., R.G. Grainger, A.T. Prata, S.R. Proud, T.A. Mather, and D.M. Pyle, Asatellite chronology of
597 plumes from the April 2021 eruption of La Soufrière, St Vincent. *Atmos. Chem. Phys.*, 23, 15209–
598 15234, 2023.

599 Vernier J.-P., Thomason L. W., Pommereau J. P., Bourassa A., Pelon J., Garnier A., Hauchecorne A., Blanot
600 L., Trepte C., Degenstein D., and Vargas F., Major influence of tropical volcanic eruptions on the
601 stratospheric aerosol layer during the last decade, *Geophys. Res. Lett.*, 38, 1–8,
602 <https://doi.org/10.1029/2011GL047563>, 2011.

603 Vernier J.-P., D. Farlie, T. Deshler, M. Natarajan, T. Knepp, K. Foster, F.G. Weingold, K.M. Bedka, L.
604 Thomason and C. Trepte, In situ and space-based observations of the Kelud volcanic plume: The
605 persistence of ash in the lower stratosphere. *J. Geophys. Res. Atmos.* 121, 11,104–11,118, doi:10.1002/
606 2016JD025344, 2016.

607 Voudouri K.A., K. Michailidis, M.-E. Koukouli, S. Rémy, A. Inness, G. Taha, G. Peletidou, N. Siomos, D.
608 Balis and M. Parrington, Investigating a Persistent Stratospheric Aerosol Layer Observed over
609 Southern Europe during 2019. *Remote Sens.* 15, 5394. <https://doi.org/10.3390/rs15225394>, 2023.

610 Waters, J. W., Froidevaux, L., Harwood, R., Jarnot, R., Pickett, H., Read, W., Siegel, P., Cofield, R.,
611 Filipiak, M., Flower, D., Holden, J., Lau, G., Livesey, N., Manney, G., Pumphrey, H., Santee, M., Wu,
612 D., Cuddy, D., Lay, R., Loo, M., Perun, V., Schwartz, M., Stek, P., Thurstans, R., Boyles, M., Chandra,
613 S., Chavez, M., Chen, G.-S., Chudasama, B., Dodge, R., Fuller, R., Girard, M., Jiang, J., Jiang, Y.,
614 Knosp, B., LaBelle, R., Lam, J., Lee, K., Miller, D., Oswald, J., Patel, N., Pukala, D., Quintero, O.,
615 Scaff, D., Snyder, W., Tope, M., Wagner, P., and Walch, M.: The earth observing system microwave
616 limb sounder (EOS MLS) on the Aura satellite, *IEEE T. Geosci. Remote*, 44, 1106–1121, 2006

617 Wilson J.C., S.-H. Lee, J. M. Reeves, C. A. Brock, H. H. Jonsson, B. G. Lafleur, M. Loewenstein, J.
618 Podolske, E. Atlas, K. Boering, G. Toon, D. Fahey, T. P. Bui, G. Diskin, and F. Moore, Steady-state
619 aerosol distributions in the extra-tropical, lower stratosphere and the processes that maintain them.
620 *Atmos. Chem. Phys.*, 8, 6617–6626, 2008.

621 Winker, D. M., Hunt, W. H., and McGill, M. J.: Initial performance assessment of CALIOP, *Geophys. Res.*
622 *Lett.*, 34, 1–5, <https://doi.org/10.1029/2007GL030135>, 2007.

623 Winker, D. M., Pelon, J., Coakley, J. A., Ackerman, S. A., Charlson, R. J., Colarco, P. R., Flamant, P., Fu,
624 Q., Hoff, R. M., Kittaka, C., Kubar, T. L., Le Treut, H., McCormick, M. P., Mégie, G., Poole, L.,
625 Powell, K., Trepte, K., Vaughan, M. A., and Wielicki, B. A.: The CALIPSO mission– A global 3D view
626 of aerosols and clouds, *B. Am. Meteorol. Soc.*, 91, 1211–1229,
627 <https://doi.org/10.1175/2010BAMS3009.1>, 2010.

628 Winkler P., The growth of atmospheric aerosol particles as a function of the relative humidity – II Improved
629 concept of mixed nuclei. *Aerosol Sci.* 4, 373-387, 1973.

630 Xu J., D. Li D., Z. Bai, M. Tao and J. Bian, Large Amounts of Water Vapor Were Injected into the
631 Stratosphere by the Hunga Tonga–Hunga Ha’apai Volcano Eruption. *Atmosphere* 13, 912. <https://doi.org/10.3390/atmos13060912>, 2022.

632
633 Yang, K., X. Liu, P. K. Bhartia, N. A. Krotkov, S. A. Carn, E. J. Hughes, A. J. Krueger, R. J. D. Spurr, and
634 S. G. Trahan, Direct retrieval of sulfur dioxide amount and altitude from spaceborne hyperspectral UV
635 measurements: Theory and application. *J. Geophys. Res.*, 115, D00L09, doi:10.1029/2010JD013982,
636 2010.

637 Zhu Y., C.G. Bardeen, S. Tilmes, M.J. Mills, X. Wang, V. Lynn Harvey G. Taha, D. Kinnison, R.W.
638 Portmann, P. Yu, K.H. Rosenlof, M. Avery, C. Kloss, C. Li,10, A.S. Glanville, L. Millán, T. Deshler, N.
639 Krotkov and O.B. Toon, Perturbations in stratospheric aerosol evolution due to the water-rich plume of
640 the 2022 Hunga-Tonga eruption. *Comm. Earth Environm.* 3:248, <https://doi.org/10.1038/s43247-022-00580-w>, 2022.

641
642 Zhuang J. and F. Yi, Nabro aerosol evolution observed jointly by lidars at a mid-latitude site and CALIPSO.
643 *Atmos. Environm.* 140, 106-116, 2016.

644 *Data availability.* The data used are publicly available: CALIOP V4.51 lidar data (<https://search.earthdata.nasa.gov/search?fp=CALIPSO>), and MLS data (version 5.0-1.0a, level 2) from https://disc.gsfc.nasa.gov/datasets?page=1&keywords=ML2H2O_005.

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647 *Author contributions.* BGM planned the study, undertook most of the data analysis and wrote the paper. JF
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