# The effect of ash, water vapor, and heterogeneous chemistry on the evolution of a Pinatubo-size volcanic cloud

Mohamed Abdelkader<sup>1</sup>, Georgiy Stenchikov<sup>1</sup>, Andrea Pozzer<sup>2</sup>, Holger Tost<sup>3</sup>, and Jos Lelieveld<sup>2</sup> <sup>1</sup>Division of Physical Sciences and Engineering, King Abdullah University of Science and Technology, Thuwal 23955-6900, Saudi Arabia <sup>2</sup>Air Chemistry Department, Max Planck Institute for Chemistry, Mainz, 55128, Germany <sup>3</sup>Institute for Atmospheric Physics, Johannes Gutenberg University of Mainz, Mainz, 55128, Germany

Correspondence: Mohamed Abdelkader (mohamed.ahmed@kaust.edu.sa)

Abstract. We employ the atmospheric chemistry general circulation model (EMAC) with gas phase, heterogeneous chemistry, and detailed aerosol microphysics to simulate that incorporates calculations of gas phase and heterogeneous chemistry coupled with the ozone cycle and aerosol formation, transport, and microphysics to calculate the 1991 Pinatubo volcanic cloud. We explicitly account for the interaction of simultaneously injected considered simultaneous injections of SO<sub>2</sub>, volcanic ash, and

- 5 water vaporand. We conducted multiple ensemble simulations with different injection configurations to test the simulated evolution of  $SO_2$ ,  $SO_4^{2-}$ , ash masses, stratospheric aerosol optical depth, surface area density (SAD), and the stratospheric temperature response against available observations. We find that the , masses and stratospheric aerosol optical depth (SAOD) are found that the volcanic cloud evolution is sensitive to the initial height of the altitude where volcanic debris is initially injected and the initial concentrations of the eruption products that affect radiative heating and lofting of the volcanic cloud. The
- 10 volcanic cloud interacts with tropopause and stratopause, and its composition is shaped by heterogeneous chemistry coupled with the ozone cycle. The height of the volcanic cloud in our simulations is also affected by dynamic processes within the cloud, i. e., heating and lofting of volcanic products. The mass of the injected water vapor has a moderate effect on the cloud evolution when volcanic numerical experiments with the injection of 12 Mt SO<sub>2</sub>, 75 Mt of volcanic ash, and 150 Mt of water vapor at 20 km show the best agreement with the observations aerosol optical depth and stratospheric temperature response. Volcanic
- 15 water injected by eruptive jet and/or intrudes through the tropopause accelerates  $SO_2$  oxidation. But the mass of volcanic water retained in the stratosphere is controlled by the stratospheric temperature at the injection level. E.g., if volcanic materials are released in the lower stratosphere because it cold point above the tropical tropopause, most of the injected water freezes and sediments as ice crystals. However, the injected water vapor at a higher altitude accelerates the oxidization of which is sensitive to the injected water vapor mass (via hydroxyl production and reaction rate) The water vapor directly injected into the
- 20 volcanic cloud increases the  $SO_4^{2-}$  mass and stratospheric aerosol optical depth by about 5%. The coarse ash comprises 98% of ash injection mass, which the ash injected mass. It sediments within a few days, but aged sub-micron ash could stay in the stratosphere for a few months providing SAD for heterogeneous chemistry. The presence of ash accelerates the SO<sub>2</sub> oxidation that leads to a faster by 10-20% due to heterogeneous chemistry, radiative heating, lofting, and faster dispersion of volcanic debris. Ash aging affects its lifetime and optical properties, almost doubling the ash radiative heating. The two-and-a-half-year
- 25 simulations show that the stratospheric temperature anomalies forced by radiative heating of volcanic debris in our experiments

with the 20 km injection height agree well with observations and reanalysis data. This indicates that the model captures the long-term evolution and climate effect of the Pinatubo volcanic cloud. The volcanic cloud's initial lofting, facilitated by ash particles' radiative heating, controls the oxidation rate of SO<sub>2</sub>. Ash accelerates the formation of the sulfate acrosol-layer in the first two months after the eruptionand has to . We also found that the interactive calculations of OH and heterogeneous

30 chemistry increase the volcanic cloud sensitivity to water vapor and ash injections. All those factors must be accounted for in modeling the impact of large-scale volcanic injections on climate and stratospheric chemistry.

#### 1 Introduction

#### Volcanic activity is a

(Azoulay et al., 2021) (Bittner et al., 2016) (Toohey et al., 2014) (Fujiwara et al., 2022) (Labitzke and McCormick, 1992) Driscoll et al

- 35 <u>Strong explosive volcanic eruptions are the</u> major natural cause of climate <u>variation-variability</u> on both global and regional scales (Robock, 2000). <u>Strong explosive volcanic eruptions-Volcanic explosions</u> inject a mixture of SO<sub>2</sub>, volcanic ash, water vapor, halogens, and other tracers into the <u>lower-stratosphere</u>. The injected volcanic materials scatter and absorb incoming solar and outgoing terrestrial radiation, warming the stratosphere and cooling the Earth's surface and the lower troposphere (Hansen et al., 1992; Stenchikov et al., 1998; Kirchner et al., 1999; Robock, 2000; Soden, 2002; Shindell et al., 2001). Stratospheric
- 40 warming (Stenchikov et al., 1998) and tropospheric cooling (Kirchner et al., 1999; Ramachandran et al., 2000) caused by the radiative impact of volcanic aerosols yield to changes in atmospheric circulation (Stenchikov et al., 1998; Kirchner et al., 1999; Ramachand affect El Nino Southern Oscillation (ENSO) (Predybaylo et al., 2017), and force a positive phase of the Arctic Oscillation (AO) (Graf et al., 1993; Kodera and Kuroda, 2000; Mao and Robock, 1998; Kodera and Kuroda, 2000; Stenchikov, 2002; Shindell, 2004; Stenchikov et al., 2006; Karpechko et al., 2010) causing boreal winter warming in middle and high latitudes over Eurasia
- 45 and North America (Stenchikov et al., 2004; Thomas et al., 2009; Poberaj et al., 2011).

Recent research shows that the mechanisms of stratosphere-troposphere dynamic interaction are complex, and large simulation ensembles are required to detect and attribute dynamic responses reliably. Stenchikov et al. (2006), Driscoll et al. (2012), Charlton-Perez et al. (2013) showed that the Intergovernmental Panel on Climate Change (IPCC) models have the problem of producing a stronger northern polar vortex in response to low-latitude volcanic eruptions. Conveying this signal to the

50 surface is even more problematic. E.g., Polvani et al. (2019) concluded that the positive AO phase after the Pinatubo eruption appeared only by chance. Toohey et al. (2014) further elaborated on the planetary-wave-based mechanism of winter warming after large low-latitude eruptions. Bittner et al. (2016) and Azoulay et al. (2021) showed that a stronger eruption could more reliably force a positive phase of the AO.

The eruption of Mount Pinatubo (Philippines, in the Philippines on 15 June 1991) with had an Explosivity Index of VEI=6 and caused the largest climate impact in the twentieth century. It is also by far was also the largest eruption to affect that affected a densely populated area. The observed global mean visible optical depth from the Pinatubo eruption reached 0.15. It was about two times higher than that of the second largest eruption in the 20<sup>th</sup> century, El Chichon in 1982 (Dutton and Christy, 1992). The 1991 Mt. Pinatubo eruption is also the best observed explosive event with a detected significant climate impact. It has been documented by satellite instruments (McCormick, 1987; Long and Stowe, 1994), ground-based LIDARs

- and sunphotometers (Antuna et al., 2002, 2003; Good and Pyle, 2004; Nagai et al., 2010; Dutton and Christy, 1992; Thomason, 1992), and airborne aerosol counters (McCormick et al., 1995; Pueschel et al., 1994; Borrmann et al., 1995; Deshler, 2003). Mount Pinatubo produced about five cubic kilometers of dacitic magma. Three magma. According to observations, three main volcanic explosions were reported to spread the on 15<sup>th</sup> June 1991 spread volcanic ash and gases over an area of 300,000 km<sup>2</sup>
- The SO<sub>2</sub> mass emitted by the Mount Pinatubo eruption was estimated using Stratospheric Aerosol and Gas Experiment (SAGE), TOVS, TOMS and ground-based LIDAR. The TIROS Operational Vertical Sounder (TOVS), and The Total Ozone Mapping Spectrometer (TOMS) retrievals (Guo et al., 2004a; Rose et al., 2006; Sheng et al., 2015; Krueger et al., 1995; Fisher et al., 2019). In addition to SO<sub>2</sub>, Pinatubo injected tens of megatons of water vapor and volcanic ash into the stratosphere (Guo et al., 2004a; Nedoluha et al., 1998; Joshi and Jones, 2009).
- In the stratosphere,  $SO_2$  is oxidized by the OH radical to form sulfuric acid, which then binary nucleates in the presence of water to form sulfate aerosol. The primary source of OH in the stratosphere is ozone photolysis by ultraviolet radiation. This reaction forms oxygen and atomic oxygen in the excitation state ( $O^{1d}$ ), which interacts with water vapor to form OH radicals. Thus, the  $SO_2$  oxidation is controlled by the abundance of OH, which depends on the concentration of stratospheric water vapor (Lovejoy et al., 1996). The co-injection of water vapor with  $SO_2$  therefore accelerates the formation of sulfuric acid (LeGrande
- 75 et al., 2016). The online calculation of OH is essential to correctly reproduce the dynamics of sulfate aerosol mass (Clyne et al., 2021; Stenchikov, 2021), and this has been neglected in many previous studies (Niemeier et al., 2009; Oman et al., 2006).
  previous studies (Marshall et al., 2018; Niemeier et al., 2009; Oman et al., 2006).

The sulfuric acid resulting results from  $SO_2$  oxidation nucleates to form long-lived sub-micron sulfate particles which interact droplets interacting with solar and terrestrial radiation. The radiative effect and lifetime of sulfate aerosols depends

- 80 depend on their size distribution, which is not definitively established. Therefore, different Pinatubo modeling studies report a wide range of visible (0.5-0.6 μm) Stratospheric Aerosol Optical Depth (SAOD) for the same amount of injected SO<sub>2</sub>. Brühl et al. (2015) obtained equatorial average SAOD=0.38 compared to SAOD=0.11 reported by LeGrande et al. (2016) for 17 Mt of injected SO<sub>2</sub>. Niemeier et al. (2021) and Stenchikov et al. (2021) obtained similar SAOD which is consistent with observations for 17 Mt of injected SO<sub>2</sub>. Dhomse et al. (2014), using a detailed aerosol microphysics model, found that in simulations of a
- Pinatubo-like eruption with a 10 Mt of  $SO_2$  injection, SAOD matches observations better than that with larger  $SO_2$  emission. Mills et al. (2016) also reported that in their model a 10 Mt  $SO_2$  injection produces the best fit to Pinatubo observations, while Sheng et al. (2015) and Sukhodolov et al. (2018) found that SAOD in their experiments with the emission of the 14 Mt of  $SO_2$ best fits SAGE observations. Timmreck et al. (2018) conducted ensembles of simulations with perturbed parameters, including the mass of injected  $SO_2$ - $SO_2$  and the injection height of volcanic debris, to quantify the uncertainties in the radiative forcing
- 90 of the 1991 Mt. Pinatubo eruption.

Volcanic ash (tephra) comprises silicate, and volcanic glass with traces of gas bubbles (Kremser et al., 2016). Ash particles have a wide range of sizes from sub-microns to millimeters (Rose and Durant, 2009) and highly irregular shapes. Large ash particles with radii r>1 µm sediment relatively quickly (Niemeier et al., 2021, 2009; Stenchikov et al., 2021), and are believed

to contribute little in the long-term evolution of a volcanic cloud. Fine ash particles with r<1 µm disperse over vast distances

- 95 and can survive in the stratosphere for several months (Pueschel et al., 1994; Zhu et al., 2020; Russell et al., 1996; Vernier et al., 2016), but. Still, their radiative effect is small because of their relatively smaller small mass. However, Stenchikov (2021) showed that despite the fact that although most ash mass sediments during the first week after an eruption, ash solar and IR heating and chemical/microphysical interactions with sulfate particles could affect the volcanic cloud formation and its long-term evolution. Ash particles could be coated by sulfate, becoming chemically aged (Muser et al., 2020; Zhu et al., 2020).
- 100 They also uptake SO<sub>2</sub>, thereby decreasing its abundance (Zhu et al., 2020). The coating and aging of ash particles increase their size, alter alter their optical properties, and increase their deposition velocities (Muser et al., 2020; Zhu et al., 2020). The enhanced ash sedimentation removes a portion of sulfate depending on the aging level. At the same time, stratospheric aerosol particles (ash and sulfate) provide surfaces for heterogeneous chemical reactions affecting stratospheric chemical composition (Muthers et al., 2015). Aerosol particles from volcanic eruptions increase the surface area density (SAD) and hence the rate
- 105 of heterogeneous reactions involving  $ClONO_2$  and  $N_2O_5$ . This damps the  $NO_x$  mixing ratios altering the OH stratospheric budget, which affects the rate of  $SO_2$  oxidation (Prather, 1992; Kilian et al., 2020). Fig. ?? summarizes the microphysical and chemical transformations of the erupted volcanic materials.

Like mineral dust, volcanic Volcanic ash absorbs and scatters solar shortwave (SW) and terrestrial longwave (LW) radiation. This has a significant impact on the chemistry significantly impacts the dynamics and radiation budget of the atmosphere in

- the first few days after an eruption, causing rapid lifting of volcanic debris (Stenchikov et al., 2021; Niemeier et al., 2009). The models with different physics that calculate the evolution of volcanic clouds and their impact on climate assume various SO<sub>2</sub> injection heights, initial plume composition (ash and water are often not injected), use different spatial-temporal resolutions , different treatment of ash-sulfate interaction, and ash chemical aging and treat differently ash-sulfate chemical interaction. The differences in physics and chemistry translate into the differences in volcanic cloud evolution and radiative
- 115 effect. For instance, those models which generate finer sulfate particles (Mills et al., 2017; Dhomse et al., 2020) overestimate the stratospheric sulfate lifetime, generating higher SAOD for the same injection mass.

Along with the  $SO_2$  mass, the injection height of volcanic debris is a critical parameter for correctly simulating the dispersion of a volcanic cloud, as it associates with the wind field that transports the volcanic plume. Different modeling studies assume different injection heights. The results show that the oxidation rate of  $SO_2$  depends strongly on the injection height according

- 120 to the availability of water vapor and OH radicals. Sheng et al. (2015) performed a sensitivity study for the initial mass and altitude of the injected  $SO_2$  for the Pinatubo eruption and showed that a mass of 17 Mt of  $SO_2$  or less gives the best agreement with the SAGE optical depth within a peak of the volcanic cloud between 18-21 km. The transient equilibrium height of the volcanic plume depends not only on the height of the initial injection but also on internal feedback mechanisms. Stenchikov et al. (2021) demonstrated that radiative heating by ash was lifting volcanic debris by 1 km per day during the first week
- 125 following the 1991 Pinatubo eruption. Muser et al. (2020) reported the lifting of a volcanic plume of Raikoke eruption that was the Raikoke eruption, an order of magnitude smaller than the 1991 Pinatubo eruption. Volcanic debris injections cause significantly different localized radiative heating and lofting when weaker lofting when their initial distribution is quasi-zonal, as in Brühl et al. (2015) and when than when it is localized, as in LeGrande et al. (2016). In contrast to addition to processes

considered in Stenchikov et al. (2021), we here explicitly calculate ash chemical aging, stratospheric ozone chemistry, and

130 aerosol microphysical processes-, accounting for the hygroscopic growth of sulfate/ash particles. However, we do not account for the heating by SO<sub>2</sub> because for the Pinatubo eruption, it is much weaker than radiative heating from ash and sulfate aerosols (Osipov et al., 2021; Stenchikov et al., 2021; Osipov et al., 2020).

The underlying dynamic and *for chemical mechanism chemical mechanisms* of the large sensitivity of SAOD to the injection height has not been recognized have not been tested yet in a fully interactive model. The effects of injected "volcanic" water and

- 135 chemical aging of volcanic ash on  $SO_2$  oxidation rate and  $SO_4^{2-}$  removal are not studied within the models with comprehensive gaseous and heterogeneous chemistry and detailed microphysics. Here we use the Atmospheric Chemistry and Aerosol General Circulation Model (EMAC ) EMAC model with multi-phase chemistry along with detailed aerosol microphysics to study the evolution of a Pinatubo-size volcanic cloud. We account for the entire range of dynamic, chemical, and microphysical complexity of the processe processes governing the development of volcanic cloud to address the following science questions:
- How does the initial spatial distribution and height of injected volcanic debris affect the evolution of a volcanic cloud?
  - What is the effect of heterogeneous chemistry on the SO<sub>2</sub> oxidation rate within a volcanic cloud?
  - How do co-injection of SO<sub>2</sub>, water vapor, and ash, affect volcanic cloud evolution?
  - How does the aging of co-injected ash affect volcanic cloud development?

#### 2 Data

- To constrain the simulations and evaluate the model results, we use the SAGE data set with partially filled gaps compiled by Stratospheric Processes and its Role in Climate (SPARC) and published in the Assessment of Stratospheric Aerosol Properties (ASAP) report (Thomason and Peter, 2006). This data set provides the aerosol effective radius and aerosol extinction in UV, visible (0.525 μm), and near IR (1.02 μm) wavelengths. The SAGE/ASAP SAOD is zonal mean and collected assembled on a monthly basis. It is available from 70°S to 70°N with 5° resolution in latitude between from 1984 and to 1999. We further
- 150 refer to this data as SAGE/ASAP SAOD or R<sub>eff</sub>. The SAGE observations of aerosol extinction contain multiple gaps at the initial stage of the volcanic cloud evolution because of the instrument's saturation (Thomason, 1992). The observations in near IR-near-IR observations are of better quality than in visible or UV-Ultraviolet (UV) (Stenchikov et al., 1998). Therefore, to obtain the visible SAGE/ASAP SAOD, we scale near IR SAOD using the angstrom exponent obtained from our simulations, similar to (Stenchikov et al., 1998). We also use the Advanced Very High-Resolution Radiometer (AVHRR) SAOD at 0.63
- 155 μm (Long and Stowe, 1994). The AVHRR observations are collected over the oceans at 0.1°x0.1° horizontal resolution for cloud-free conditions at daytimedaylight time. The AVHRR AOD is measured for the entire atmospheric column, including the troposphere. To obtain the AVHRR stratospheric AOD, we calculate the AVHRR AOD monthly climatology for the pre-Pinatubo period of 1985-1990 and subtract it from the total AOD for the Pinatubo period. Unfortunately, this can introduce some level of uncertainty due to the high variability of tropospheric AOD. We refer to the visible SAOD obtained from SAGE

160 the SAGE near IR observations as the scaled SAGE/ASAP SAOD. We compare scaled SAGE/ASAP and AVHRR SAOD at 0.63 to the model visible SAOD.

Krueger et al. (1995) estimated the mass of  $SO_2$  during the first 15 days after the 1991 Pinatubo eruption based on the Total Ozone Mapping Spectrometer (TOMS) observations. They concluded that the mass of the initially emitted  $SO_2$  was  $15\pm3$  Mt. Guo et al. (2004a) later estimated the emitted mass of  $SO_2$  to be 14-20 Mt. Recent estimates reduce the initial  $SO_2$  mass to 12

- 165 Mt (Fisher et al., 2019). Estimates of SO<sub>2</sub> mass using retrievals from the Optical Vertical Sounder/High-Resolution Infrared Radiation Sounder/2 (TOVS) on the Television Infrared Observation Satellite (TIROS) suggest that the initial SO<sub>2</sub> mass was  $19\pm4$  Mt (Guo et al., 2004a). However, the TOVS retrievals are less accurate than TOMS because they are affected by sulfate aerosol absorption in IR. SO<sub>4</sub><sup>2-</sup> mass has also been estimated using the High-Resolution Infrared Radiation Sounder(HIRS/2) (HIRS/2) (Guo et al., 2004a). However, the The estimated sulfate aerosol mass depends on the aerosol size distribution, which
- 170 is not well known, and this introduces uncertainties into adds uncertainties to the estimated  $SO_4^{2-}$  mass.

Volcanic ash mass is available was estimated for the first few days after the Pinatubo eruption from AVHRR and High-Resolution Infrared Radiation Sounder/2 (using the AVHRR and HIRS/2) observations (Guo et al., 2004b). HIRS/2 detected 80Mt of fine ash in the atmosphere on the first day after the eruption. AVHRR ash retrievals evaluate the spectral contrast of radiance (Aerosol Index) to distinguish between absorbing aerosols, such as volcanic ash, and non-absorbing aerosols, such as sulfate.
175 However, the retrieval algorithm does not consider particles smaller than 1 µm (Guo et al., 2004b).

We obtain the stratospheric temperature response to the 1991 Pinatubo eruption from the Modern-Era Retrospective analysis for Research and Applications (MERRA2) reanalysis data available on 0.5°x0.625° horizontal grid and 72 vertical levels from the surface to 0.01 hPa (Gelaro et al., 2017). To reproduce the effect of the 1991 Pinatubo eruption, MERRA2 assimilates observations from different satellite sensors such as TOVS and the Spinning Enhanced Visible and InfraRed

180 Imager (SEVIRI) on the Meteosat Second Generation (MSG) Satellite, as well as the Microwave Limb Sounder (MLS). The MERRA2 temperature anomalies fields are consistent with the observations reported by Labitzke and McCormick (1992). The reanalyses intercomparison S-RIR (Fujiwara et al., 2022) shows that the MERRA2 stratospheric temperature anomalies caused by the 1991 Pinatubo injection resemble the observations well, although the absolute stratospheric temperature is slightly underestimated. In this study, the MERRA2 temperature anomalies were calculated with respect to 1985-1990 climatology.

#### 185 3 Model

Here we employ the ECHAM5/MESSy2 atmospheric chemistry model, EMAC (Joeckel et al., 2005, 2006, 2010). EMAC is a modular model based on sub-models that describe processes in the stratosphere, the middle atmosphere, and the troposphere, accounting for anthropogenic emissions and interactions with oceans and land (Joeckel et al., 2010). EMAC has been used to study impacts of volcanic stratospheric aerosols on climate and stratospheric circulation (Brühl et al., 2012, 2015; Bingen et al., 2017; Löffl

as well as dust aging and dust-air pollution interaction interactions in the troposphere (Abdelkader et al., 2015, 2017; Klingmüller et al., 2019, 2020).

The Modular Earth Submodel System (MESSy) links the various submodels. The submodels comprise AEROPT, CLOUD, CONVECT. CVTRANS, DDEP, GMXE, JVAL, LNOX, MECCA, OFFEMIS, ONEMIS, RAD4ALL, SCAV, SEDI, TNUDGE. and TROPOP. Table 1 shows the submodels used in this study indicating their functionality, while the detailed description of all

195 EMAC submodels can be found in Joeckel et al. (2010) (Joeckel et al., 2010). We configure EMAC using MESSy version 2.52 with the 5<sup>th</sup> generation European Centre Hamburg Atmospheric general circulation Model version 5.3.02, ECHAM5 (Roeckner et al., 2006) - and employ the same chemistry and aerosol microphysics setup as in (Brühl et al., 2012, 2015). For vertical approximation we employ, we use 90 sigma-hybrid levels from the earth surface up to 0.01 hPa, and T42 spectral approximation horizontally that corresponding to  $2.8^{\circ}$  grid spacing at the equator both in longitude and latitude. Varying monthly sea surface temperature and sea ice are prescribed from AMIPII dataset (Taylor et al., 2000).

200

We apply the guasi-biennial oscillation (OBO) submodel to capture the observed phase of OBO and account for its effect on the stratospheric circulation, similar to Stenchikov et al. (2004) (Stenchikov et al., 2004). No other constraints are imposed on the model dynamics, e. g., we do not nudge tropospheric winds.

- The emission inventory comprises the sources of greenhouse gases, NO<sub>x</sub>, CO, NMVOCs, NH<sub>3</sub>, SO<sub>2</sub>, black carbon (BC), 205 and organic carbon (OC). We also account for the background emission of CFCs, Halogens, and Halons. The emissions are monthly mean and geographically distributed according to the EDGAR4-2009 EDGAR4 emission inventory and the Global Fire Emissions Database (GFED) version 3 (van der Werf et al., 2010). We also-account for the DMS and OCS emissions similar to Brühl et al. (2015). To calculate atmospheric composition, we We employ 230 gas-phase chemical reactions, 76 photolytic reactions, and 12 heterogeneous reactions for 159 species to calculate atmospheric composition. The photolysis
- 210 rates are calculated within the model for the spectral range 178.6 nm  $\leq \lambda \leq$  752.5 nm accounting for gaseous absorption (O<sub>3</sub>) and  $O_2$ ), Rayleigh scattering, absorption, and scattering by aerosols and clouds (Landgraf and Crutzen, 1998; Sander et al., 2011). In this setup, the photolysis rates are not coupled to volcanic aerosol. The model calculates the instantaneous radiative forcing using double radiation calls, with and without aerosols. Aerosol microphysics and chemistry are called every model time step, while the radiation sub-model is called every third-time step.

#### 215 3.1 Stratospheric sulfate chemistry

Volcanic sulfate results from the oxidation of  $SO_2$  by OH in the presence of water vapor. OH is produced by ozone photolysis by UV radiation with wavelengths less than 0.242  $\mu$ m. This reaction forms O<sub>2</sub> and excited oxygen O(<sup>1d</sup>)(Eq. ??). The excited oxygen radical interacts with water to form the hydroxyl radical OH(Eq. ??), which oxidizes  $SO_2$  in two steps to form sulfate. At the first step, OH oxidizes SO<sub>2</sub> to form SO<sub>3</sub> and HO<sub>2</sub>(Eq. ??). At the second step, SO<sub>3</sub> interacts with water molecules to

- 220 form sulphuric acid (Eq. ??). (Seinfeld and Pandis, 2006). The rate of reaction in Eq. ?? depends on the concentration of water molecules that are also in the reactants (Burkholder et al., 2015). Therefore, higher water vapor concentrations significantly increase concentration significantly increases the formation rate of sulphuric acid. New sulfate particles are generated by the binary nucleation of sulfuric acid and water molecules. Thus, the formation of sulfate particles in a volcanic cloud depends strongly on water vapor concentration. The models that do not parameterize nucleation explicitly are less sensitive to the
- abundance of water vapor in a volcanic cloud than those that do (LeGrande et al., 2016). 225

The aerosol setup in EMAC has been described in detail in (Pringle et-, 2010; Pringle et al., 2010; Pringle et al., 2012; Pringle et al., 2015; Pringle et al., 2005, 2011, 2019). Aerosol size distributions in the model are approximated by seven lognormal modes: four soluble modes (nucleation, Aitken, accumulation, coarse) and three insoluble modes (Aitken, accumulation, coarse). In our simulations, sulfate represents by the particles are represented by soluble modes, and while ash is initially considered insoluble until it ages, i.e., five monolayers of sulfate particles coat the ash particle. The modes' median radii change in time during aerosol microphysical transformations, but the widths of the modes remain fixed. The median radii for three insoluble modes and dry cores of four soluble modes initially are equal

to 0.0015, 0.025, 0.25, and 2.5 µm for nucleation, Aitken, accumulation, and coarse modes, respectively. The widths geometric standard deviations of the lognormal distributions for the above modes are 1.59, 1.59, 1.49, and 1.70, respectively (Brühl et al., 2015).

Aerosols in soluble modes evolve by uptake or loss of water and  $SO_4^{2-}$  molecules , and coagulation. The hygroscopic growth of ash is only allowed in a soluble mode (Abdelkader et al., 2015). The mass of large or fine aerosol particles in the size

240 distribution tails is assigned to a corresponding neighboring mode when the mode's median radius reaches a certain threshold. The aerosol modes and the thresholds are schematically shown in Fig. 1. In our simulations, we choose threshold radii equal to 0.0005, 0.006, 0.07, 1.6 µm for the nucleation, Aiken, accumulation, and coarse modes respectively, as in (Brühl et al., 2015).

#### 3.3 Volcanic ash - Model Implementation

We introduce introduced a new "ash" tracer to account for volcanic ash in the EMAC setting. We assume the ash density to be
 2400 kg m<sup>-3</sup>, similar to that of . Ash is similar to mineral dust, as ash comprises mainly silicate comprising mainly silicates (SiO<sub>2</sub>). Therefore, for calculating chemical aging, we assume that ash particles have the same water uptake and accommodation coefficients as dust particles (Abdelkader et al., 2015).

High-density In our simulations, high-density ash particles sediment faster than pumice assumed in (Zhu et al., 2020). Zhu et al. (2020) considered the Kelud eruption that emitted 100 times less volcanic material than the 1991 Pinatubo eruption; therefore, even the long-lived pumice ash produced negligible radiative heating. Stenchikov et al. (2021) showed that applying the assumption about long-lived pumice ash for the larger volcanic explosions like the 1991 Pinatubo eruption could cause unrealistic overheating of the stratosphere.

For a full representation of chemical aging, we use a comprehensive chemistry scheme that enables the production of the primary inorganic acids which contribute to the chemical aging of ash particles (Metzger et al., 2016).

255

250

<sup>5</sup> Volcanic ash is removed from the stratosphere mainly by gravitational sedimentation. Sedimentation parameterization in EMAC utilizes the Walcek scheme (Walcek, 2000; Kerkweg et al., 2006a). Ash scavenging in the troposphere is implemented in EMAC by Tost et al. (2006a) and is fully coupled with the aerosol and gas-phase chemistry.

To calculate the optical properties of volcanic ash, we choose its complex refractive index to be equal to that of dust assuming ash particles. For a complete representation of chemical aging, we use a comprehensive chemistry scheme that enables the production of the primary inorganic acids which contribute to the chemical aging of ash particles (Metzger et al., 2016).

Ash particles scatter and absorb solar and terrestrial radiation(Pollack et al., 1973; Vogel et al., 2017; Stenchikov et al., 2021). In visible light. To calculate their optical properties we choose the volcanic ash's complex refractive index according to (Pollack et al., 1973; Vogel et al., 2017; Stenchikov et al., 2021). For visible wavelengths, the ash refractive index RI=1.53 + 0.004*i*. Ash is more absorbing in UV, near-infrared (NIR), and Infrared (IR) near-IR, and IR than in visible. Table 1–S1 in the supplement shows the volcanic ash refractive index as a function of wavelength. Fig. 8–S8 shows the refractive indices used in

265

260

## the EMAC model for different aerosols as a function of wavelength.

#### 3.4 Aerosol Radiative Effect

We use the AEROPT submodel to calculate extinction, single-scattering albedo, and asymmetry parameter, the aerosol optical properties required for the radiative transfer calculations. It is assumed that different types of aerosols are mixed internally so
that the refractive index of the mixture is calculated from the volume fractions of the aerosol components. The sensitivity to this assumption is discussed in detail by Klingmüller et al. (2014). The optical properties are calculated for each aerosol mode independently. The RAD submodel calculates radiative transfer (Roeckner and Coauthors, 2003). The Fouquart and Bonnel seheme (Fouquart and Bonnel, 1980) Fouquart and Bonnel (1980) scheme is used for calculating shortwave radiation, while longwave radiation is calculated using RRTM-a Rapid Radiative Transfer Model (RRTM) (Iacono et al., 2008). Scattering of the infrared-IR radiation by aerosols is neglected. RAD accounts for shortwave and longwave absorption of water vapor, clouds, O<sub>3</sub>, CH<sub>4</sub>, N2O, CO2, CFCs, and aerosols, including sulfate and volcanic ash implemented in this study. Table 2 shows

the shortwave and longwave bands used in the radiative transfer calculations in EMAC. For comparison with observations, we consider the first two SW bands in Table 2 as visible and near-infrarednear-IR.

#### 4 Experimental Setup

The Table 3 shows the complete set of experiments is listed in Table 3numerical experiments conducted in this study. The control experiment (ctrl) describes the state of the atmosphere from 1990 to 2000when unperturbed by volcanic eruption. All, assuming there were no volcanic eruptions. In the perturbed simulations (those with volcanic aerosols present)were conducted from June 1, 1991, to December 31 with the injection of volcanic aerosols), 1994. We we emit 17 Mt of sulfur dioxide in all perturbed simulations, with the exception of one experimentas in (Stenchikov et al., 2021), except in one experiment, specifically marked and used to study sensitivity to SO<sub>2</sub> emission mass. The perturbed simulations were calculated from June

285 specifically marked and used to study sensitivity to SO<sub>2</sub> emission mass. The perturbed simulations were calculated from 1, 1991, to December 31, 1994, with a one-year spin-up not included in the analysis.

Along with  $SO_2$ , we consider co-injections of water vapor and ash. For ash, we adopt the same initial size distribution as in (Niemeier et al., 2009) and (Stenchikov et al., 2021). We redistribute the total emitted fine ash mass of 75 Mt (Guo et al., 2004b) between two insoluble modes, accumulation and coarse (Fig. 1). The accumulation mode comprises 1.5 Mt of ash, and

- 290 the coarse mode comprises 73.5 Mt of ash. The ash massDespite its small mass, ash in the accumulation mode is important since it has a much longer lifetime than ash in the coarse mode. We use the standard EMAC *import* and *offemis* IMPORT and OFFEMIS submodels to initialize the SO<sub>2</sub>, water vapor, and ash tracers (Kerkweg et al., 2006b). Section 1 in the supplement explains the implementation of SO<sub>2</sub>, water vapor, and volcanic ash injection mechanism mechanisms in EMAC.
- In the main set of experiments, we release volcanic products in the specified model grid box centered at the <u>altitude of 17</u> km, 20 km, or 25 km <u>height</u> at the geographical coordinates of Mt. Pinatubo (15.1429 °N, 120.3496 °E) with <u>pre-calculated</u> the <u>precalculated</u> emission rates (in molecules m<sup>-3</sup> s<sup>-1</sup>) during 24 h. <u>Here we refer to these as We call it a one-grid-box</u> emission scheme <u>- See (see</u> Table 3 for details). In the 1s1-17km, 1s1-20km, and 1s1-25km experiments, we assume that only SO<sub>2</sub> is injected at 17 km, 20 km or 25 km, respectively. In the 1w1-20km experiment we release SO<sub>2</sub> and water vapor (Nedoluha et al., 1998; Joshi and Jones, 2009) at 20 km. The va0 experiments employ the same settings as 1w1 but
- 300 assume injection of 75 Mt of ash. The va0 experiments do not account for the chemical aging of ash. The va1 experiments are similar to va0 but account for ash aging. In contrast to the one-grid-box emission scheme, in experiment 3s10-25km we inject SO<sub>2</sub> in the 3000 km wide latitude belt centered at the latitude of the eruption mimicking the setting in Brühl et al. (2015)(Brühl et al., 2015). The injected layer is 10 boxes thick (5-from 22.5km to 27.5km) and is centered at the altitude of 25 km. The experiments are listed in Table 3. We When this does not confuse, we refer to the clusters of experiments
- 305 with the same physics using a generic name without specifying injection altitudealtitudes, such as 1s1, 1w1, va0, and va1, in instances when this will not cause confusion. Experiments 1s1 are used to study the sensitivity to the height of the injection of volcanic SO<sub>2</sub>. The 1w1 experiments with 150 Mt and 15 Mt injected water allow allows us to quantify the dependence of sensitivity to the mass of injected water vapor. Experiments va0 and va1 are designed to quantify the effect of ash and ash aging, respectively. Experiment 3s10-25km mimics the quasi-zonal injection from Brühl et al. (2015). Experiment.
- 310 va1-20km-12Mt is designed to study dependence on the amount of the sensitivity to the amount injected SO<sub>2</sub>. All simulations are conducted with a one-year spin-up not included in the analysis. To reduce the effect of internal model variability in each experiment, we calculate five ensemble members using different atmospheric initial conditions. The analysis in this study is performed and presented for the ensemble means. We show the "ensemble" variability for the selected variables.

#### 315 5 Results

First, we compare the model results with observations, focusing on spatial-temporal distributions of  $SO_2$ ,  $SO_4^{2-}$  and other related chemicals. We also compare the test the simulated stratospheric AOD (SAOD) which defines volcanic radiative effect, and the stratospheric temperature response which measures volcanic climate impact against available observations. In addition, we compare the Surface Area Density (SAD) that controls heterogeneous chemistry within the volcanic cloud, and aerosol effective radius ( $R_{eff}$ ) that characterizes aerosol size distribution (see Fig. 2-5). The spatially averaged  $R_{eff}$  is calculated as a

effective radius ( $R_{eff}$ ) that characterizes aerosol size distribution (see Fig. 2-5). The spatially averaged  $R_{eff}$  is calculated as a ratio of the third  $M3_m$  and the second  $M2_m$  moments of each aerosol mode *m* integrated over the entire domain (Eq. 1 and

Eq. 2). The effective radii for individual modes and for the entire aerosol size distribution are given by (Eq. 3) and (Eq. 4), respectively.

$$M2_m = \iiint_v N_m R_m^2 exp^{(2ln^2\sigma_m)} dxdydz \tag{1}$$

325

$$M3_m = \iiint N_m R_m^3 exp^{(\frac{9}{2}ln^2\sigma_m)} dxdydz \tag{2}$$

$$R_{eff}^m = \frac{M3_m}{M2_m} \tag{3}$$

$$R_{eff} \underbrace{\overset{total}{\underline{\phantom{a}}}}_{m=1} = \frac{\sum_{m=1}^{m=Nmodes} M3_m}{\sum_{m=1}^{m=Nmodes} M2_m}$$
(4)

330

Where  $N_m$  is the number density for aerosol mode m,  $R_m$  is the median radius, and  $\sigma_m$  is the width of the aerosol mode m.  $N_{modes}$  is the number of aerosol modes.

Figures 2-4 show-compare various parameters in 3s10-25km, and 1s1 experiments with the different injection heights , as well as-with the AVHRR, and SAGE/ASAP observations. The AVHRR zonal mean visible SAOD is largely consistent in spatial-temporal behavior with the scaled SAGE/ASAP SAOD (Fig. 3). The original SAGE/ASAP visible SAOD is almost half of the AVHRR SAOD , because the SAGE II sensor was saturated during the few first-first few weeks after the eruption, therefore , Therefore data at the initial stage of eruption are sparse. The AVHRR continuously sensed the entire atmospheric columnineluding , including the troposphere, the effect of which could be estimated only approximately (Thomason, 1992; Russell et al., 1996; Kremser et al., 2016). The consistency between the scaled SAGE/ASAP and AVHRR visible SAODs lessens in the late fall of 1991 ,-when scaled SAGE/ASAP SAOD begins overestimating AVHRR SAOD. Discrepancies between different data sets are discussed in (Bingen et al., 2004). Despite sparse observations at the initial stage of volcanic cloud development, SAGE/ASAP is the only global satellite observation that recorded the vertical structure of the Pinatubo cloud.

For example, Fig. 4 demonstrates aerosol SAD at different altitudes as reported by SAGE/ASAP and simulated in the model.

Below we study the sensitivity of volcanic cloud evolution to all the main factors: injection height, amount of injected water, injection of ash, and ash aging. We start from sensitivity to injection height using the simplest 1s1 experiments with  $SO_2$ 

only injections. The cloud height is essential because it defines the wind field that drives eloud dispersion the dispersion of volcanic debris. The  $O_3$  mixing ratio and abundance of water vapor and water vapor mixing ratios, which affect chemical and microphysics transformations reactions and aerosol microphysics within the plume, are also height dependent.

### 5.1 Sensitivity to Injection Height

Figure Figures 2a,b,c compares compare the observed and simulated SAOD, SO<sub>2</sub> and SO<sub>4</sub><sup>2-</sup> masses, and R<sub>eff</sub> in the 1s1 exper-

- 350 iments with different injection heights, respectively. The altitudes where volcanic debris resides depend not only on the initial injection height but <u>on</u> upward stratospheric motion and lofting driven by radiative heating of volcanic debris (Stenchikov, 2021; Niemeier et al., 2009; Kinnison et al., 1994; Aquila et al., 2012). The latter process and the rate of chemical transformations within a volcanic cloud are sensitive to the initial concentrations of optically and chemically active materials within a fresh volcanic cloud, i.e., in terms of our simulation settings, from to the volume that a cloud initially occupies.
- Experiment 3s10-25km assumes a zonally uniform  $SO_2$  release at 25 km altitude within a latitude belt centered at the latitude of the eruption (15.1429°N). The visible SAOD from this experiment compares well with observations. In experiment 1s1-25km, we release  $SO_2$  centered at the same height -as in the 3s10-25km experiment, but within one model grid box at the geographic coordinates of the Pinatubo eruption (15.1429°N, 120.3496°E). This causes initially higher  $SO_2$  concentrations compared to the 3s10-25km experiment. The volcanic debris is cruptive materials are released with a constant mass emission
- 360 rate and spread for more than 1000 km during the 24 hours of emission. Despite that  $SO_2$  was released at the same altitude, these two experiments exhibit remarkable differences in the globally averaged SAODs (see Fig. 2a),  $SO_4^{2-}$  masses (Fig. 2b), and spatial distributions of SAOD and SAD (Fig. 3 and Fig. 4). To understand the mechanism of the strong sensitivity of the volcanic cloud evolution to its initial stage, below-we test the 3s10-25km experiment and the one-grid-cell -only injection experiments-1s1 with experiments with the 17 km, 20 km, and 25 km injection heights against observations.

#### 365 5.1.1 SAOD

Contrary to the 1s1-25km experiment, the visible tropical SAOD in experiment 1s1-20km compares well with that <u>SAOD</u> from the scaled SAGE/ASAP and AVHRR observations (Fig. 2a). The visible SAOD from the Sixth Coupled Model Intercomparison Project (CMIP6) (Eyring et al., 2016; Zanchettin et al., 2016) which mimics the original visible SAGE/ASAP extinctions <u>SAOD</u> develops slowly and is half of the scaled SAGE/ASAP and AVHRR.

- The equatorial average (20S-20N) SAOD in 1s1-17km is half the size of the 1s1-20km and 3s10-25km SAODs. The 1s1-25km SAOD is even smaller (Fig. 2a). All SAODs except that in the 1s1-25km experiment are bigger than the CMIP6 SAOD. The SAOD in the 3s10-25m experiment grows faster than in the SAOD in the 1s1 experiments, reaching 0.33 in August 1991. At a given chemical composition of sulfate aerosol particles, the transient SAOD depends both on the  $SO_4^{2-}$  mass, i.e., the rate of oxidation of  $SO_2$  to  $SO_4^{2-}$ , and on aerosol size distribution, i.e.,  $R_{eff}$ . The smaller sulfate aerosol particles have a bigger
- 375 collective cross-section per unit mass than larger ones. So a bigger mass of large sulfate particles might have a smaller SAOD than a smaller mass of smaller sulfate particles. This must be considered when evaluating the mass of  $SO_4^{2-}$  and the sulfate aerosol SAOD in observations and model experiments.

#### 5.1.2 Oxidation of SO<sub>2</sub>

Figure 2b shows the globally integrated SO<sub>2</sub> and SO<sub>4</sub><sup>2-</sup> masses in the 3s10-3s10-25km and 1s1 experiments with the different
emission heights as functions of time. The SO<sub>2</sub> mass in the 1s1-20km experiment decreases more slowly than in all other experiments. Furthermore, the The SO<sub>4</sub><sup>2-</sup> mass in the 3s10-25km grows faster than in the other experiments in Fig.Figure 2b.

This is because the  $SO_2$  oxidation rate depends on the abundance of OH radicals. The OH production depends on  $O_3$  concentration and incoming UV radiation. Because in the 3s10-25km experiment, SO<sub>2</sub> is distributed zonally over the entire latitude beltin the 3s10-25km experiment, its concentration in a volcanic cloud is lower than in all one-grid-box-injection experiments.

385 Hence, the SO<sub>2</sub> oxidation is more efficient in the 3s10-25km experiment because than in the 1s1 runs because initially, there are more OH radicals available per  $\underline{an}$  SO<sub>2</sub> molecule in the latitude belt than in  $\underline{a}$  smaller volcanic cloud volume as in the one-grid-boxexperiments. Furthermore, is less depleted by in a larger volume. All 1s1 experiments underestimate  $SO_4^{2-}$  mass in the first few days in comparison with the available observations (Fig. 2b). The presence of  $SO_4^{2-}$  in a fresh volcanic plume detected in observations is confusing puzzling as the models usually do not account for the physical mechanisms that could produce it in such a short time. To explain this discrepancy, Guo et al. (2004a) suggested that 1-2 Mt of  $SO_4^{2-}$  was injected at

the initial stage of the eruption. However, we do not account for the initial  $SO_4^{2-}$  release in this study.

390

#### 5.1.3 Spatial-temporal Evolution of SAOD and SAD

The spatial-temporal patterns of visible SAOD in the 1s1-20km experiment compare well with AVHRR and scaled SAGE/ASAP observations (Fig. 3), although the aerosol poleward. However, the poleward aerosol transport in the model is too fast. This 395 is a known deficiency of global models which simulate subtropical barriers which are too transparent, which simulate too transparent subtropical barriers due to coarse spatial resolution -(Oman et al., 2006). The 1s1-25km visible SAOD is smaller than the scaled SAGE/ASAP and AVHRR SAODs - and exhibits qualitatively incorrect evolution of the volcanic cloud, which moves too far north, similar to that reported by Stenchikov et al. (2021) for volcanic injection at 24 km altitude. The 3s10-25km SAOD has a realistic spatial-temporal structure but substantially overestimates observed SAODs. SAOD in the 1s1-17km experiment (Fig. 3) exhibits even faster poleward transport than in the 1s1-20km run due to stronger more vigorous wave activity 400 at lower altitudes in the stratosphere. In this experiment, the equatorial aerosol reservoir dissipates too quickly because of its proximity to the tropopause and too-intensive poleward transport.

Figure 4 compares the SAD in the 1s1-17km, 1s1-20km, and 1s1-25km experiments with the SAGE/ASAP observations (Thomason et al., 1997). SAD facilitates heterogeneous reactions in the volcanic cloud. Both sulfate aerosols and volcanic

- ash contribute to SAD, but in 1s1 experiments, we only account for sulfate aerosol surfaces. Thereforeit is expected that, the 405 simulated SAD will is expected to be smaller than the observed one, especially at the very beginning after the eruption. Only the 1s1-20km experiment shows SAD distributions consistent with observations at all three levels; 20 km, 25 km, and 30 km. Both the model and SAGE/ASAP show that the peak SAD is at 20. At higher altitudes (25 and 30), SAD is smaller than at 20 altitude. This suggests that volcanic material in the simulations has been lifted by at least 5 above the injection level. In the
- 410 1s1-17km experiment the model underestimates SAD at 25 km and 30 km, while the volcanic cloud remains at and below 20 level. In the 1s1-25km experiment, the volcanic cloud resides climbs unrealistically high, at and above 30 km. At that height, sulfate droplets tend to evaporate and the. The sulfuric acid photolyzes back to SO<sub>2</sub>, and is eventually transported to which reaches the mesosphere (Rinsland et al., 1995).

#### 5.1.4 Aerosol Size Distribution

415 Figure 2c compares  $R_{eff}$  from SAGE/ASAP averaged over the tropical belt, the 3S10-25km, and the 1s1 experiments with the 17, 20, and 25 km injection heights. In the control case, the model  $R_{eff}$ =0.14 µm is lower than the observed unperturbed value of 0.17-0.19 µm (Russell et al., 1993), as the model underestimates the effect of anthropogenic sulfur emissions on the stratospheric Junge layer (Marandino et al., 2013; Brühl et al., 2015).

SAGE-II observations suggest that aerosol extinction increases, and its maximum shifts from 0.385 µm to 525-0.525 µm
soon after the Pinatubo eruption, indicating the sudden increase of sizes of aerosol particles (Thomason, 1992; Thomason and Peter, 2006; Kremser et al., 2016). The observed effective radius increases from the background level to about 0.5 µm in six months (Russell et al., 1996).

In Figure 2c,  $R_{eff}$  in the 1s1-20km experiment increases gradually, reaching maximum  $R_{eff} = 0.4 \ \mu\text{m}$  in September of 1991, and then decreases due to settling of larger particles. In the 1s1-25km experiment,  $R_{eff}$  is the largest - when compared

- 425 with other experiments, as it generates the largest concentration. It among all experiments.  $R_{eff}$  in this experiment initially grows faster than in all other runs, reaching maximum  $R_{eff} = 0.45 \ \mu m$  in August of 1991, and then decreases, merging with all other experiments in December of 1991. In experiment the 1s1-17km experiment,  $R_{eff}$  is the smallest when compared to other experiments, as it looses loses SO<sub>2</sub> mass through the tropopause. All the simulations are predicting quite similar temporal evolution of  $R_{eff}$ .
- 430 The tropical visible SAODs in Fig. 2a are consistent with mass and  $R_{eff}$ . That is, the mass in the 1s1-25km experiment is larger than in the 1s1-17km experiment, but SAOD is smaller because 1s1-25km  $R_{eff}$  is bigger. The When testing our simulated  $R_{eff}$  in 1s1-25km and 3s10-25km have maximums above that in SAGE/ASAP. The  $R_{eff}$  in the 1s1-17km and 1s1-20km runs is always below the against observations, we have to mention that the SAGE/ASAP  $R_{eff}$ . However the SAGE/ASAP  $R_{eff}$  is itself is quite uncertain (Ansmann et al., 1997).

#### 435 5.1.5 Impact on Chemical Composition

Figure 5 shows vertical cross-sections of the mixing ratios or concentrations of  $SO_2$ ,  $SO_4^{2-}$ , OH,  $H_2SO_4$ ,  $NO_x$ ,  $NO_{y,2}O_3$  in the 1s1 experiments with the 17km, 20km, and 25km injection heights (see Eq. ?? - Eq. ??) averaged over the equatorial belt (20°S - 20°N). We do not account here for the  $SO_2$  radiative effect (Stenchikov, 2021), and there is no ash injection in these experiments. Therefore it is , only sulfate aerosols that cause heating and lofting of the volcanic cloud. After the

- 440 injectionBefore a significant amount of  $SO_4^{2-}$  develops, lifting is caused by regional upward motion in the Brewer–Dobson circulationbefore develops, being reproduced by EMAC Brewer-Dobson circulation, which EMAC captures. The SO<sub>2</sub> and  $SO_4^{2-}$  clouds separate due to gravitational settling of sulfate aerosols (Fig. 5a-f). This initiates multi-layer distributions of all other tracers. Stratospheric vertical uplift depends on the altitude which is getting stronger at higher altitude altitudes (at least in EMAC). This is well seen in the 1s1-25km run in comparison with the 1s1-17km and 1s1-20km runs (see Fig. 5a-c).
- 445 The volcanic cloud in the 1s1-25km experiment rises to 30 km, significantly higher than in all other experiments. This affects the development of the volcanic cloud. The SO<sub>2</sub> oxidation rate slows down as the temperature rises. The Therefore, the SO<sub>4</sub><sup>2-</sup>

mass is therefore smaller than in the other experiments. In addition, in the 1s1-25km experiment,  $R_{eff}$  is higher than in other experiments. This factor tends to lower the SAOD, since larger particles in the 1s1-25km experiment are less optically efficient per unit mass, and have a lower lifetime with respect to gravitational settling. Therefore, SAOD and SAD in this experiment are smaller than in the others in Fig. other experiments in Figure 2 and Fig. 4Figure 4, respectively.

450

465

480

The experiments with different emission heights result in differences in the SAOD in generate different SAOD (Fig. 2and SAD in ) and SAD (Fig. 4). This partially results from different SO<sub>2</sub> oxidation rates that are defined by the abundance of OH radicals at different altitudes. Oxidation of volcanic in the stratosphere also perturbs the Chapman cycle and reduces the ozone mixing ratio in the stratosphere.

Three weeks after the eruption, OH is reduced around the injection height because of stratospheric water consumption by the oxidation of SO<sub>2</sub> in all three experiments in Fig.Figures 5g, h, i. OH remains depleted above the  $SO_4^{2-}$  cloud, where SO<sub>2</sub> mixing ratio is high. The change in OH is generally the largest in the 1s1-25km experiment.

The increase of  $H_2SO_4$  is also-more pronounced in the 1s1-25km experiment (Fig. 5j,k,l). Initially, the  $H_2SO_4$  increase develops increases at the emission level. This is seen until November of 1991. Then a secondary plume of  $H_2SO_4$  is formed at

460 a higher altitude, above 29km. Oxidation of volcanic  $SO_2$  in the stratosphere also perturbs the Chapman cycle and reduces the ozone mixing ratio in the stratosphere (Fig. 5s,t,u).

We account for twelve heterogeneous reactions. Heterogeneous reactions on aerosols explain the re-partitioning between NO<sub>x</sub> and the reactive nitrogen reservoir NO<sub>y</sub>. Following Danilin et al. (1999), we evaluate the effect of the heterogeneous reactions heterogeneous chemistry by the abundance of NO<sub>x</sub> (NO+NO<sub>2</sub>) and total inorganic nitrogen, NO<sub>y</sub> (NO<sub>x</sub> + NO<sub>3</sub> + HNO<sub>3</sub> +  $2*N_2O_5$  + HONO + HNO<sub>4</sub> + ClONO<sub>2</sub> + BrONO<sub>2</sub>).

The In addition, the heterogeneous chemistry might also influence the oxidation capacity be influenced by the halogen reactions by chlorine and bromine activation; however, as no additional halogen emissions from the eruption are considered, this effect. However, we do not account for the volcanic halogen injections in this study; therefore, their impact might be minor. Here, the main pathway in this transformation is the oxidation of NO<sub>x</sub> to form  $N_2O_5$ , which interacts with water vapor

- to form HNO<sub>3</sub>. N<sub>2</sub>O<sub>5</sub> can also interact with halogens on the surface of aerosols (sulfate or ash). The heterogeneous reaction of N<sub>2</sub>O<sub>5</sub> and water on the surface of aerosols effectively depletes NO<sub>2</sub> from the active reaction system depending on SAD (Fig. 4). During the daytime, HNO<sub>3</sub> can photo-dissolve and form OH, interacting with NO<sub>x</sub> to form N<sub>2</sub>O<sub>5</sub> and then HNO<sub>3</sub>. While at night time, the formation of HNO<sub>3</sub> is one-way via oxidation of NO<sub>x</sub> and N<sub>2</sub>O<sub>5</sub>. N<sub>2</sub>O<sub>5</sub> can decompose back to NO<sub>3</sub> and NO<sub>2</sub> either photochemically or thermally, depending on the overhead column of ozone. As altitude increases, temperature
- 475 increases, and the production rate of HNO<sub>3</sub> increases. The limiting factors in the heterogeneous formation of HNO<sub>3</sub> are that of NO<sub>2</sub>, O<sub>3</sub> column, and SAD (Seinfeld and Pandis, 2006).

Figure 5 m-r shows the strong dependence of  $NO_x$  and  $NO_y$  on injection height within the aerosol cloud. The  $NO_x$  mixing ratio decreases, and the  $NO_y$  mixing ratio increases, along with the increase of the injection height. The changes in  $NO_x$  and  $NO_y$  affect the ozone cycle (Seinfeld and Pandis, 2006). The dependence of the background ozone concatenations concentrations on altitude adds to the sensitivity of the cloud evolution to injection height. Furthermore, the modified ozone

concentrations feed concentration (Fig. 5 s.t.u) feeds back to the OH OH production and hence the sulfur oxidation . SO<sub>2</sub>

oxidation rate. For the 1s1-17km experiment, the depletion of  $NO_x$  (Fig. 5 m) is lower than for 25km injection (Fig. 5 o), while the production of  $NO_y$  at 25km (Fig. 5 r) injection is higher than for 17km experiment (Fig. 5 p). At higher altitude, the ozone concentration and SAD (Fig. 4 c) is higher, and hence the formation of HNO<sub>3</sub> is higher for the 1s1-25km experiment (see

- 485 Fig. S9 in the supplement). Although the change in  $NO_y$  for 1s1 experiments at 17km, 20km, and 25km injection is insignificant (see Fig. S9b in the supplement), the heterogeneous transformation from  $N_2O_5$  to HNO<sub>3</sub> is efficient. The transformation is enhanced (Fig. S9b,c,d in the supplement) by the injection of ash particles due to the additional SAD and heating by ash and the associated stronger lofting of the volcanic plume.
- Along with the chemical processes, the interaction of volcanic debris with the tropopause and the stratopause , adds in adds to
  the sensitivity of the SO<sub>4</sub><sup>2-</sup> mass to the height of the injection. In the 17 km injection height experiments experiment, the cloud loses part of the mass through the tropopause, but in the 25 km injection height experiment, part of the sulfur is transported to the mesosphere and gets lost for immediate sulfate formation. It descends to the stratosphere again in high latitudes in winter. The volcanic debris injected at 20 km stabilizes in the middle of the stratosphere. Henceit, in this case, the volcanic cloud is less affected by interaction with the tropopause and the tropopause and stratopause.

#### 495 5.2 Water Vapor Intrusiondue to Tropopause Heating

As expected, warming of the tropical tropopause layer by radiative heating of volcanic debris facilitates the cross-tropopause troposphere-to-stratosphere transport of water vapor (Oltmans and Hofmann, 1995; Nedoluha et al., 1998; Joshi and Jones, 2009). The presence of extra water vapor in the stratosphere intensifies OH production and accelerates  $SO_2$  oxidation to form sulfate particles (LeGrande et al., 2016).

- For the 1s1-17km experiment, the stratospheric (i.e., above 100 hPa) water vapor mass integrated over the tropical belt increases by about 30 Mt (Fig. 6a)at the equatorial belt. However, changes an increase of water vapor just above the tropopause do does not affect volcanic cloud evolution much because the bulk of this water vapor is well below the altitude where the core of the volcanic cloud resides. Cross-tropopause water transport decreases as injection height increases. For example, the 1s1-25km injection experiment shows no cross-tropopause water transport. In three Three weeks after the injection, the aerosol water associated with sulfate aerosols in the 1s1-17km experiment (Fig. 6c) is higher compared with other experiments ,
- because than in other experiments because, in the 1s1-17km experiment more water vapor penetrates the stratosphere through the tropopause . However, the volcanic cloud is closer to the tropopause than in the other experiments. In the 1s1-20km run, the mass of  $SO_4^{2-}$  in the 1s1-20km run continues to increase continues increasing during August and September 1991 (Fig. 2b), and the associated aerosol water also increases to 3.5 Mt as shown in Fig. 6c. Little ice is accumulated in the stratosphere in
- 510 all experiments (Fig. 6b), since it is quickly removed by gravitational sedimentation. In the 1s1-20km experiment liquid water mass peaks at 3 Mt; in the 1s1-17km at 2Mt; and in 1s1-25km at 1 Mt (Fig. 6d).

#### 5.3 Volcanic Water Injection

Water vapor injected into the stratosphere with a volcanic plume could directly affect the initial evolution of a volcanic cloud since it is concentrated within it. Most of this water is brought by the entrainment of tropospheric water in an explosive jet

- 515 or co-ignimbrite convective updrafts; nevertheless, the term "volcanic" water is used here. A wide range (75 Mt 150 Mt) of volcanic water vapor injection for the Pinatubo eruption was reported (Joshi and Jones, 2009; Nedoluha et al., 1998). However, the amount of volcanic water retained in the stratosphere depends on the height of the injection. That is, almost injection height. Almost all water vapor injected at a low temperature just above the tropopause forms ice and quickly sediments sedimented (Stenchikov et al., 2021). A larger fraction of water vapor injected at higher altitudes, where stratospheric temperaturesare
- 520 higher with higher stratospheric temperatures, could remain in the stratosphere. To test the sensitivity of volcanic clouds to the amount of volcanic water vapor, we conduct the 1w1 simulations injecting simultaneously  $SO_2$  and  $\frac{15 \text{ or } 150 \text{ different amounts}}{150 \text{ of water vapor}}$  of water vapor at 20 and 25k heights.

species comprise water vapor, ice, and aerosol water. The aerosol water accumulates in sulfate and over ash particles.

Figure 7 compares the time series of the equatorial SAODs, and changes in the globally integrated masses of sulfate and water species in the stratosphere (above 100 hPa) in the 1w1 experiments, with the simultaneous injection of  $SO_2$  and 15 Mt or 150 Mt of volcanic water vapor at 20 km and 25 km with respect to the corresponding 1s1 experiments (see Table 3). Water

525

The effect of volcanic water on the generation of the  $SO_4^{2-}$  mass and SAOD is dependent depends on the amount of water vapor retained in the stratosphere after the eruption. The sensitivity of SAOD and  $SO_4^{2-}$  mass to the injected volcanic water vapor is higher in the 1w1-25km experiment compared to the 1s1-20km experiment (Fig. 7c,d). The increase in sulfate mass

530 results from the acceleration of SO<sub>2</sub> oxidation facilitated by the higher water vapor concentration (see Eq. ??). Water vapor emission in the (Seinfeld and Pandis, 2006). The sensitivity to the amount of water vapor in the experiments with the 20 km injection experiment has a weaker effect altitude is weaker than in the experiments with the 25 km injection experiment, height because most of the water vapor injected at 20 km condenses and deposits from the stratosphere , since the temperature is lower at 20 km is lower than at 25 km (Fig. 7i). Because more injected water remains in the stratosphere in the 1w1-25km experiment, its effect is more significant than in the 1w1-20km experiment.

### 5.4 Volcanic Ash Injection

In the va0 and va1 experiments, we inject 75 Mt of ash together with SO<sub>2</sub> and water vapor. The va1 experiment accounts for ash aging, but the va0 does not. In both experiments, we assume that the volcanic ash is initially hydrophobic. Therefore, we inject it into the insoluble (dry) accumulation and coarse modes (Fig. 1 and Fig. 8). In the va1 experiments, volcanic ash ages quickly, populating the soluble (wet) modes (Fig. 8c,d), while ash particles in the va0 experiments remain in the dry modes (accumulation and coarse). In the va1 experiments, ash particles increase in size due to the aging and associated water and SO<sub>4</sub><sup>2-</sup> uptake, which tends to transfer particles from the accumulation to the coarse mode.

In the va0 experiments, ash in the coarse mode (see Fig. 8b) sediments from the stratosphere in for two days, but ash particles in the accumulation mode remain in the stratosphere for a week (Fig. 8a). In the va1 experiments, the ash mass in the 545 wet modes increases quickly due to dry-to-wet particle conversion shown by the arrow in Fig. 8. The aging of ash particles slows the decrease of ash mass in both accumulation and coarse modes.

In the experiments with a 25 km injection height, it takes longer for ash to reach the tropopause and leave the stratosphere in comparison to the 20m than in the experiments with the 20 km experimentinjection height. For instance, after the first day

of injection, 60 Mt of insoluble coarse mode ash mass ash remains in the stratosphere (not shown) for the va0 experiment with

550 25 km injection height compared to 1.7 Mt when ash is injected at 20 km (see Fig. 8b).

Figure 9a shows the evolution of the stratospheric ash masses in the va0 and va1 experiments compared to the AVHRR and HIRS/2 retrievals (Guo et al., 2004b). The mass of volcanic ash in va0 is smaller than that in va1. In the va1 experiment, the model ash mass is higher than in the AVHRR and HIRS/2 observations, while in the va0 experiment, the ash mass is underestimated when compared with observations. However, the uncertainties in the AVHRR derived ash mass are  $\pm 53\%$  (Gu

et al., 2003) and  $\pm 85\%$  in HIRS observations (Yu and Rose, 2000). retrievals (Yu and Rose, 2000). The larger ash mass in the val experiment (compared to va0) on the first day after the eruption causes stronger heating and lofting of ash in the val experiment than in va0, which prolongs its lifetime in the stratosphere (Fig. 12).

The injection of volcanic ash significantly increases the stratospheric optical depth and  $R_{eff}$  during the few days after injection the eruption. This is shown in Fig.Figures 10a,b which compares compare the time series of SAOD and the effective

- 560 radius averaged within 20S 20N latitude belt above 100 hPa in the 1s1-20km, 1w1-20km, va0-20km, and va1-20km experiments, with available observations. The AVHRR and scaled SAGE/ASAP SAODs are consistent for at least 4-5 months after the eruption. The CMIP6 SAOD appears to be half the size when compared with to them. During the 4-5 months following the eruption, the simulated SAOD (Fig. 10a) is slightly larger than in observations but decreases more quickly when compared to than in observations later on in all experiments except va0-20km. The va1-20km and va0-SAODs va0-20km
- 565 experiments grow more rapidly during the first two months than in all experiments without ash injection. the 1s1-20km and 1w1-20km experiments (Fig. 10a). The effective radii in the val and va0 experiments spike to about 0.6-0.8 µm during the first week after the eruption, when a significant amount of ash is present in the volcanic cloud.

Figure 10c,d shows the evolution of the  $SO_4^{2-}$  mass in the coarse and accumulation modes in the same experiments integrated over the 20S - 20N latitude belt. In the va0 and va1 experiments, the stratospheric sulfate mass increases more rapidly than in the 1w1 and 1s1 experiments. This is consistent with SAOD in Fig. 10a and with the more rapid depletion of  $SO_2$  mass in

570 in the 1w1 and 1s1 experiments. This is consistent with SAOD in Fig. 10a and with the more rapid depletion of  $SO_2$  mass in Fig. 9b, which demonstrates a better agreement with  $SO_2$  mass observations. We relate the faster  $SO_2$  oxidation in the va0 and va1 experiments with the effect of heterogeneous reactions on ash particles , and more intensive volcanic cloud dispersion facilitated by ash radiative heating.

Two months after the eruption, in the 1s1 and 1w1 experiments, the sulfate mass in accumulation and coarse modes reaches maximums of 9Mt and 0.7 Mt, respectively. Thus, the sulfate formation rate increases in the va1 and va0 experiments compared to experiments without ash in both accumulation and coarse modes. The  $SO_4^{2-}$  mass reaches the maximum two weeks earlier in experiments with ash than in experiments without ash (Fig. 10c,d).

The aerosol water mass increases when sulfate mass increases, both for in the accumulation and the coarse modes (Fig. 10e,f). A sulfate mass of 9Mt is associated with aerosol water mass of 3 Mt in the accumulation mode (Fig. 10c,e). This is consistent with the 75% sulfuric acid solution assumed by Stenchikov et al. (1998). For the coarse mode, the aerosol water mass of 0.5 Mt is associated with roughly 0.8 Mt of sulfate (Fig. 10d,f). Both sulfate and wet ash particles accumulate aerosol water. In the long runhowever, due to the shorter lifetime of the ash particles, aerosol water is associated mainly with mainly retained in sulfate aerosols. FigureFigures 10g,h show SO<sub>4</sub><sup>2-</sup> mass in the coarse and accumulation modes in the troposphere (integrated below

100 hPa) globally for the same experiments. The tropospheric  $SO_4^{2-}$  mass of volcanic origin comprises  $SO_4^{2-}$  sedimented 585 from the stratosphere. Because of the rapid wet removal, little sulfate (not exceeding 0.4 Mt in each mode) is accumulated in the troposphere both in accumulation and coarse modes. This is more than an order of magnitude less than the  $SO_4^{2-}$  mass in the stratosphere. The tropospheric mass of volcanic origin comprises sedimented from the stratosphere. More sulfate mass sediments into the troposphere in the va0 and va1 experiments than in the 1s1 and 1w1 runs (Fig. 10e). This is, because in the va1 and va0 experiments, the stratospheric sulfate mass is bigger-larger than in the 1s1 and 1w1 experiments, as is deposition.

#### 590 5.5 Ash Aging

595

600

Ash particles provide surface areas, enhancing the heterogeneous reactions and leading to significant changes in stratospheric chemistry (Danilin et al., 1999). Ash SAD is especially important in the first week after the eruption when a few sulfate aerosols form. Fig.formed. Figures 11a-d show the 20S-20N mean SO<sub>2</sub> mixing ratio and  $SO_4^{2-}$  concentration as a function of time and height for the va0-20km and va1-20km experiments. Ash radiative heating causes lofting of the SO<sub>2</sub> plume by about 1 km per day in both experiments, similar to that found in (Stenchikov et al., 2021), although ash in our simulations is more absorbing than in (Stenchikov et al., 2021).

In both cases, the  $SO_4^{2-}$  cloud is below 35 km, but  $SO_2$  reaches the stratopause. Therefore some  $SO_2$  penetrates the mesosphere. This effect is more significant in the va0 experiment because of slower  $SO_2$  oxidation compared with the va1 experiment. The enhanced mixing ratio of  $SO_2$  in the mesosphere above 45 km km was detected in ATMOS observations (Rinsland et al., 1995) and simulated in Brühl et al. (2015) by Brühl et al. (2015) after the 1991 Pinatubo eruption.

- FigureFigures 11e,g ,i,k show the change show the changes in mixing ratio of  $H_2SO_4$ , aerosol water, and OH, and concentrations in the va0-20km experiment with respect to 1w1-20km, in order to demonstrate demonstrating the effect of ash injection. Fig.Figures 11f,h ,j,l show changes of the same characteristic characteristics, except in the va1-20km experiment with respect to the va0-20km experiment to demonstrate the effect of aging. If aging is turned on,  $H_2SO_4$  condenses on volcanic ash, decreas-
- 605 ing sulfuric acid concentration. At the same time, the presence of ash facilitates the ash facilitates heterogeneous reactions. The Moreover, the stratospheric water vapor mass is larger for the val experiment than va0 because of the more substantial heating (longwave and shortwave) in the val experiment (Fig. 11b,c). The more significant heating of the val experiment increases the tropospheric water vapor transport into the stratosphere. As a result, the stratospheric water vapor mass in val is 15Mt higher than in va0 after four months of the eruption. The increased stratospheric water vapor facilitates OH formation (11h). The
- 610 combined effect of ash aging radiative heating, aging, and heterogeneous chemistry in our setting resulted in an increase of the mass of sulphuric acid and sulfate mass by about 10%-20%, compared to those experiments without ash injections.

Figure 12 shows the averaged over the tropical belt (20S-20N) shortwave and longwave heating rates caused by volcanic cloud for the va0-20km (left column) and va1-20km (right column) experiments, as <u>a</u> function of time and height. The contour lines show the ash concentrations for the accumulation (top row) and coarse (bottom row) modes. <u>There are Both experiments</u>

615 <u>have</u> two distinguished periods of shortwave heating in both experiments increased radiative heating (Fig. 12a,b). The first period is time the heating peaks just after the eruption, and the second is time - ten days later. The first period peak is associated with ash solar absorption, and the second period - with sulfate aerosol absorption. In both cases the shortwave SW heating by sulfate peaks at 25 km due to lofting of and ash plumes (see Fig. 11a,b). The average ash  $\underline{SW}$  heating rates is about 0.4 Kday<sup>-1</sup> in experiment va1-20km, and 0.15 Kday<sup>-1</sup> - in experiment va0-20km (Fig. 12a,b). The shortwave heating caused by sulfate in the va1-20km experiment (Fig. 12a) is higher than in the va0-20km experiment (Fig. 12b).

620

The thermal absorption of the volcanic ash layer IR radiation cools the top of the volcanic cloud during the first few days after the eruption. Still, the absorption of upward IR radiation heats the bottom of the volcanic cloud. Heating caused by absorption of IR radiation by sulfate aerosols is seen in about ten days when enough  $SO_4^{2-}$  is generated. Fig.Figure 12d shows that in the va1-20km experiment, the longwave heating rate reaches 0.2 Kday<sup>-1</sup>. We observe much weaker Radiative heating in the va0-20km experiment is weaker than in the va1-20km experiment (Fig. 12c). To summarize, we can say that ash Thus, aging

625

630

significantly enhances the radiative effect of ash for about a week after the eruption.

# 5.6 Long-term climate response to volcanic forcing

In section 5.1, we showed that during the first six months after the eruption, the the model demonstrates strong SAOD sensitivity to the injection height during the first six months after the eruption. We also found that the spatial-temporal SAOD development in simulations with the volcanic emission of 17 Mt  $SO_2$  at 20 km best fit the observations during the fits the

- observations six months after the eruption but overestimate SAODs, but SAOD is overestimated. Here we further test the volcanic cloud evolution and stratospheric temperature response for the entire post-eruption\_post-Pinatubo period of 2.5 years against observations. We take advantage of the fact that the climate response provides another constrain to SAOD, since it defines stratospheric warming and tropospheric cooling constraint to SAOD (Stenchikov et al., 1998; Kirchner et al., 1999).
- 635 We also quantify the sensitivity of volcanic cloud evolution to the amount of injected  $SO_2$  considering the 12 Mt  $SO_2$  emission at 20 km height.

Figure 13 compares the post-eruption evolution of SAODs (visible and near IR) and  $SO_4^{2-}$  mass in the val experiments with the 20km injection height and the 17 Mt and 12 Mt SO<sub>2</sub> emission with the observations from CMIP6, AVHRR, SAGE/ASAP (scaled visible and original NIRnear IR) SAODs for 2.5 years. The SAOD in the val-25km experiment with In the val-20km

- 640 experiment with the 17 Mt SO<sub>2</sub> injection<del>SAOD overestimates</del>, the simulated SAOD overestimated the AVHRR and scaled SAGE/ASAP SAODs both in visible and near-IR (Fig. 14a-d) in July-August 1991. In the experiment with the 12 Mt of emitted SO<sub>2</sub>, the peak of SAOD is reduced and overestimates the observed SAOD maximum only slightly in both visible and near IRemission, the SAOD reduces and fits the observations in visible and near-IR. It is important to that the initial rate of development of visible and NIR-near IR SAODs are similar in the model and observations both in the tropics and globally. It
- suggests the model on the stage when the aerosol cloud is still confined in the tropics captures the  $SO_2$  oxidation process and  $SO_4^{2-}$  development quite well.

Starting from September 1991, the exaggerated excessive speed of poleward transport of aerosols in the model causes a faster decrease of SAOD in the simulations (both in the tropics and globally) than in the observations (Fig. 13e-h). This is because sulfur is sulfate aerosols are mainly deposited in the mid-latitude storm tracks through tropopause faults and in the

650 polar regions in the downward branch of B-D circulation (Gao et al., 2007)<del>, and the \_. The</del> faster poleward aerosol transport makes both of these processeswork more effectivelyfacilitates both these processes.

The CMIP6 visible SAOD is half the size during the first three months when compared to the scaled SAGE/ASAP and AVHRR. This is primarily due to missing data in the original visible SAGE/ASAP data set.

We further evaluate the long-term model stratospheric temperature response to test the consistency of val simulations with observations. Fig.Figure 14 shows the temperature anomalies for in the 1s1, val-20km, val-20km-12Mt experiments, and MERRA2 reanalysis. The left column of Fig.14 depicts the hovemoller diagrams of zonal mean temperature anomaly at 50 hPa, and the right column is the temporal evolution of the global mean (70S-70N) temperature anomaly as a function of height or pressure(or pressure) and time. All experiments in Fig. 14 resemble the spatial-temporal structure of the stratospheric temperature response well. They reproduce stratospheric heating by the volcanic plume volcanic heating in the first year after the eruption - and the additional heating associated with the change of the QBO phase in 1993. The simulations resemble the

MERRA2 geographical temperate response and its vertical distribution well.

In the val-20km-val-20km-12Mt experiment, the peak of temperature response is higher than in the val-20km-12Mt experiment, reaching SAOD is about 30% lower than in val-20km, causing lower radiative heating. In the val-20km experiment, the peak of temperature response reaches 4 K at 50 hPa. In the val-20km-12Mt experiment, the temperature peak is half the

- 665 size at about 2.5 K (Fig. 14a,c), which better agrees with observations (Labitzke and McCormick, 1992) and the MERRA2 temperature anomalies reanalisis (Fig. 14e). Fig.Figures 14b,d,f show a peak temperature anomaly at 30 hPa in the model simulations and the reanalysis. Again, the temperature response in the va1-20km-12Mt experiment fits the MERRA2 temperature anomalies better than the va1-20km experiment (Fig. 14b,d,f). Thus, reducing the injected SO<sub>2</sub> to 12Mt from 17 mass to 12 Mt shows a better agreement with the observations (Labitzke and McCormick, 1992), and the MERRA2 temperature response.
- 670 It results anomalies, resulting in more realistic heating at 50 hPa in both tropics and subtropics. In the val-20km-12Mt experiment, the SAOD is about 30% lower than in val-20km. The lower SAOD causes weaker aerosol radiative heating and a less vigorous temperature response.

#### 6 Conclusions

In this study, we use the EMAC model with well-developed stratospheric chemistry (including heterogeneous chemistry) and

- 675 detailed aerosol microphysics to explore the evolution of the volcanic cloud from the 1991 Pinatubo eruption, the largest and best observed volcanic event in the 20<sup>th</sup> century. We tested the model results with available observations of volcanic clouds and their radiative effect. We conducted ensemble simulations to study the impact of volcanic cloud sensitivity to the injection height and its initial volume the initial volume where volcanic materials were released (one grid-box versus a latitude belt), as well as considered the effects of co-injection of water vapor, ash, and ash aging on the formation of the volcanic cloud.
- The model simulations with 20 km injection height exhibit the best agreement of , , and ash masses, the spatial-temporal evolution of SAOD with the AVHRR SAOD and SAGE/ASAP SAOD and SADat different altitudes. In the 20 km injection experiments, the volcanic cloud is afterwards lifted to an altitude lifted to a height of 25 km by radiative heating, while in the experiments with volcanic materials injection at 25 km overshoots volcanic cloud overshoots the 30 km level. The vertical distribution of SAOD and SAD in the observations and the model experiments with a the 20 km injection height, show that the

685 aerosol-cloud aerosol cloud stabilizes in the middle of the stratosphere at 25 km. In the experiments with the 17 km and 25 km injection heights, the volcanic cloud interacts with the tropopause and the stratopause, respectively, causing some aerosol mass to be lost too quicklyescape from the stratosphere. The stratospheric oxidation capacity and wind fields are different at different altitudes, strengthening the sensitivity to the injection height. In the experiments with the zonally uniform SO<sub>2</sub> injection in a latitude belt at a the height of 25 km, the SAOD is significantly lofting is weak, so the volcanic cloud remains at the same

Because of the coarse spatial resolution (T42L90), similar to other global models, EMAC simulates a too fast aerosol poleward transport with a too quick escape of the volcanic materials from the tropical stratosphere. This process accelerates accelerating the loss of the aerosol mass to deposition at the polesand in the , in troppause folds, and storm tracks.

altitude. The SAOD in the tropical belt in this experiment is overestimated due to the higher oxidization oxidation rate.

- The increase of water vapor in the stratosphere leads to an increase of the oxidization the rise of the oxidation rate of  $SO_2$  to  $SO_4^{2-}$ . The water vapor could be brought into the stratosphere by an eruptive jet, co-ignimbrite convection, and/or intruded through the tropopause heated by absorption of solar and IR radiation by volcanic debris. The cross-tropopause water vapor intrusion does not affect the volcanic cloud evolution much, as most of the water penetrating through the tropopause accumulates below the volcanic cloud. The water vapor directly injected in into the volcanic cloud in the 1s1-20km experiment increases the  $SO_4^{2-}$  mass and SAOD by about 5%. The sensitivity of the  $SO_4^{2-}$  mass to the amount of injected water in this
- 700 experiment the experiment with the 20 km injection height is low because most of the water vapor freezes and is quickly removed quickly removes from the stratosphere in agreement with (Stenchikov et al., 2021). So the masses of remaining stratospheric water vapor in the 1w1-20km experiments with 15 Mt and 150 Mt of water vapor injections do not differ much. A significant acceleration of SO<sub>2</sub> oxidation due to injection of water vapor (LeGrande et al., 2016) is only reproduced in the experiments with the 25 km injection height , where temperature is higher than at 20 , and when a significant mass of injected
  700 water vapor is rate ined in the stratosphere.
- 705 water vapor is retained in the stratosphere.

690

710

The Our experiments' simulated mass of ash in our experiments is within the estimates of AVHRR and HIRS estimates, but observations themselves are uncertain. Volcanic ash provides SAD for heterogeneous chemistry. This is most important during the first few weeks after the eruption when ash is still abundant, but sulfate aerosol is not yet developed. The simultaneous injection of water vapor and non-aging ash in the va0-20km experiment increases the maximum SAOD and  $SO_4^{2-}$  mass by 10%.

In the val-20km experiment, ash particles in the accumulation and coarse modes are entirely aged within a day after the injection. Aging increases the mass of ash particles. They continue up-taking water and  $SO_4^{2-}$  molecules until removed by transport or sedimentation. The coarse ash particles deposit within a week, while it takes six months to reduce the mass of the ash accumulation mode from 1.2 Mt to 0.3 Mt. Overall, aging increases the SAOD by 20% and the  $SO_4^{2-}$  mass by 10%. Aging

715 doubles increases the radiative effect of ash both in SW and IR. The injections of volcanic water vapor and ash significantly accelerate the formation of the sulfate aerosols during the first two months after the eruption in the val-20km and va0-20km experiments.

The simulated maximum SAOD and stratospheric temperature anomalies in the va1-20km-12Mt experiment with the 12 Mt  $SO_2$  injection quite closely resemble the temperature anomalies obtained from the reanalysis both in latitude and height.

- 720 The inclusion of volcanic ash adds to the radiative heating of the volcanic debris during the first week after the eruption in agreement with (Stenchikov et al., 2021), showing that the initial local heating results in the lofting of the aerosol cloud. Our simulations show that the interactive calculations of OH and heterogeneous chemistry increase the volcanic cloud sensitivity to water vapor and ash injections and have to be accounted for in simulations of volcanic impacts on climate and stratospheric chemistry.
- 725 *Code and data availability.* The EMAC code modifications, including all initialization data sets, and selected simulation results, are available at the KAUST repository site at https://repository.kaust.edu.sa/handle/10754/675509, DOI 10.25781/KAUST-0W317

*Author contributions.* MA performed the calculations and prepared all the figures. MA and GS wrote the manuscript. GS planned the analysis and calculations, led the discussion, and reviewed and improved the manuscript. AP, HT, and JL advised on EMAC modifications, discussed the results, reviewed and improved the manuscript.

730 Competing interests. The authors declare that they have no conflict of interest.

*Acknowledgements.* This research has been supported by the KAUST Competitive Research Grant (URF/1/2180-01-01) 1000Combined Radiative and Air Quality Effects of Anthropogenic Air Pollution and Dust over the Arabian Peninsula,1000and the KAUST Base Research Grant (BAS/1/1309-01-01). The authors thank the KAUST Supercomputing Laboratory for providing computer resources. We are thankful to Christoph Brühl for valuable discussion and help in EMAC setting, and Linda and Mark Everett for proofreading this manuscript. HT acknowledges funding from the Carl-Zeiss foundation and from the Deutsche Forschungsgemeinschaft (DFG, German Research Foundation)

-TRR 301 - Project-ID - TRR 301-Project-ID 428312742.

#### References

- Abdelkader, M., Metzger, S., Mamouri, R. E., Astitha, M., Barrie, L., Levin, Z., and Lelieveld, J.: Dust-air pollution dynamics over the eastern Mediterranean, Atmospheric Chemistry and Physics, 15, 9173–9189, https://doi.org/10.5194/acp-15-9173-2015, 2015.
- 740 Abdelkader, M., Metzger, S., Steil, B., Klingmüller, K., Tost, H., Pozzer, A., Stenchikov, G., Barrie, L., and Lelieveld, J.: Sensitivity of transatlantic dust transport to chemical aging and related atmospheric processes, Atmos. Chem. Phys., 17, 3799–3821, https://doi.org/10.5194/acp-17-3799-2017, 2017.
  - Ansmann, A., Mattis, I., Wandinger, U., Wagner, F., Reichardt, J., and Deshler, T.: Evolution of the Pinatubo Aerosol: Raman Lidar Observations of Particle Optical Depth, Effective Radius, Mass, and Surface Area over Central Europe at 53.4°N, Journal of the Atmospheric
- 745 Sciences, 54, 2630–2641, https://doi.org/10.1175/1520-0469(1997)054<2630:EOTPAR>2.0.CO;2, publisher: American Meteorological Society Section: Journal of the Atmospheric Sciences, 1997.
  - Antuna, J. C., Robock, A., Stenchikov, G. L., Thomason, L. W., and Barnes, J. E.: Lidar validation of SAGE II aerosol measurements after the 1991 Mount Pinatubo eruption, Journal of Geophysical Research-Atmospheres, 107, 4194, https://doi.org/10.1029/2001JD001441, wOS:000178977300035, 2002.
- 750 Antuna, J. C., Robock, A., Stenchikov, G., Zhou, J., David, C., Barnes, J., and Thomason, L.: Spatial and temporal variability of the stratospheric aerosol cloud produced by the 1991 Mount Pinatubo eruption, Journal of Geophysical Research-Atmospheres, 108, 4624, https://doi.org/10.1029/2003JD003722, wOS:000186088200008, 2003.
  - Aquila, V., Oman, L. D., Stolarski, R. S., Colarco, P. R., and Newman, P. A.: Dispersion of the volcanic sulfate cloud from a Mount Pinatubo–like eruption, Journal of Geophysical Research: Atmospheres, 117, https://doi.org/10.1029/2011JD016968, \_eprint: https://agupubs.onlinelibrary.wiley.com/doi/pdf/10.1029/2011JD016968, 2012.
- Azoulay, A., Schmidt, H., and Timmreck, C.: The Arctic Polar Vortex Response to Volcanic Forcing of Different Strengths, Journal of Geophysical Research: Atmospheres, 126, https://doi.org/10.1029/2020JD034450, 2021.
  - Banda, N., Krol, M., van Weele, M., van Noije, T., and Rockmann, T.: Analysis of global methane changes after the 1991 Pinatubo volcanic eruption, Atmospheric Chemistry and Physics, 13, 2267–2281, https://doi.org/10.5194/acp-13-2267-2013, wOS:000315406600033, 2013.
- 760 Bândă, N., Krol, M., Noije, T. v., Weele, M. v., Williams, J. E., Sager, P. L., Niemeier, U., Thomason, L., and Röckmann, T.: The effect of stratospheric sulfur from Mount Pinatubo on tropospheric oxidizing capacity and methane, Journal of Geophysical Research: Atmospheres, 120, 1202–1220, https://doi.org/10.1002/2014JD022137, 2015.
  - Bingen, C., Fussen, D., and Vanhellemont, F.: A global climatology of stratospheric aerosol size distribution parameters derived from SAGE II data over the period 1984–2000: 2. Reference data, Journal of Geophysical Research: Atmospheres, 109, https://doi.org/10.1020/20021D002511.2004
- 765 https://doi.org/10.1029/2003JD003511, 2004.
  - Bingen, C., Robert, C. E., Stebel, K., Brühl, C., Schallock, J., Vanhellemont, F., Mateshvili, N., Höpfner, M., Trickl, T., Barnes, J. E., Jumelet, J., Vernier, J.-P., Popp, T., de Leeuw, G., and Pinnock, S.: Stratospheric aerosol data records for the climate change initiative: Development, validation and application to chemistry-climate modelling, Remote Sensing of Environment, 203, 296–321, https://doi.org/10.1016/j.rse.2017.06.002, 2017.
- 770 Bittner, M., Schmidt, H., Timmreck, C., and Sienz, F.: Using a large ensemble of simulations to assess the Northern Hemisphere stratospheric dynamical response to tropical volcanic eruptions and its uncertainty: LARGE ENSEMBLES FOR VOLCANIC ERUPTIONS, Geophysical Research Letters, 43, 9324–9332, https://doi.org/10.1002/2016GL070587, 2016.

- Borrmann, S., Dye, J. E., Baumgardner, D., Proffitt, M. H., Margitan, J. J., Wilson, J. C., Jonsson, H. H., Brock, C. A., Loewenstein, M., Podolske, J. R., and Ferry, G. V.: Aerosols as dynamical tracers in the lower stratosphere: Ozone versus aerosol correlation after
- 775 the Mount Pinatubo eruption, Journal of Geophysical Research: Atmospheres, 100, 11 147–11 156, https://doi.org/10.1029/95JD00016, publisher: John Wiley & Sons, Ltd, 1995.
  - Brühl, C., Lelieveld, J., Crutzen, P. J., and Tost, H.: The role of carbonyl sulphide as a source of stratospheric sulphate aerosol and its impact on climate, Atmospheric Chemistry and Physics, 12, 1239–1253, https://doi.org/10.5194/acp-12-1239-2012, 2012.
  - Brühl, C., Lelieveld, J., Tost, H., Höpfner, M., and Glatthor, N.: Stratospheric sulfur and its implications for radiative forc-
- 780 ing simulated by the chemistry climate model EMAC, Journal of Geophysical Research: Atmospheres, 120, 2103–2118, https://doi.org/10.1002/2014JD022430, 2015.

- Burkholder, J. B., Sander, S. P., Abbatt, J. P. D., Barker, J. R., Huie, R. E., Kolb, C. E., Kurylo, M. J., Orkin, V. L., Wilmouth, D. M., and Wine, P. H.: Chemical kinetics and photochemical data for use in atmospheric studies: evaluation number 18, Tech. Rep. JPL Pub 15-10, Pasadena, CA : Jet Propulsion Laboratory, National Aeronautics and Space Administration, https://trs.jpl.nasa.gov/bitstream/handle/2014/ 45510/JPL%20Pub%2015-10.pdf?sequence=1&isAllowed=y, 2015.
- Charlton-Perez, A. J., Baldwin, M. P., Birner, T., Black, R. X., Butler, A. H., Calvo, N., Davis, N. A., Gerber, E. P., Gillett, N., Hardiman, S., Kim, J., Krüger, K., Lee, Y.-Y., Manzini, E., McDaniel, B. A., Polvani, L., Reichler, T., Shaw, T. A., Sigmond, M., Son, S.-W., Toohey, M., Wilcox, L., Yoden, S., Christiansen, B., Lott, F., Shindell, D., Yukimoto, S., and Watanabe, S.: On the lack of stratospheric dynamical variability in low-top versions of the CMIP5 models: STRATOSPHERE IN CMIP5 MODELS, Journal of Geophysical Research: Atmospheres, 118, 2494–2505, https://doi.org/10.1002/jgrd.50125, 2013.
  - Clyne, M., Lamarque, J.-F., Mills, M. J., Khodri, M., Ball, W., Bekki, S., Dhomse, S. S., Lebas, N., Mann, G., Marshall, L., Niemeier, U., Poulain, V., Robock, A., Rozanov, E., Schmidt, A., Stenke, A., Sukhodolov, T., Timmreck, C., Toohey, M., Tummon, F., Zanchettin, D., Zhu, Y., and Toon, O. B.: Model physics and chemistry causing intermodel disagreement within the VolMIP-Tambora Interactive Stratospheric Aerosol ensemble, Atmospheric Chemistry and Physics, 21, 3317–3343, https://doi.org/10.5194/acp-21-3317-2021, 2021.
- 795 Danilin, M. Y., Rodriguez, J. M., Hu, W., Ko, M. K. W., Weisenstein, D. K., Kumer, J. B., Mergenthaler, J. L., Russell, J. M., Koike, M., Yue, G. K., Jones, N. B., and Johnston, P. V.: Nitrogen species in the post-Pinatubo stratosphere: Model analysis utilizing UARS measurements, Journal of Geophysical Research: Atmospheres, 104, 8247–8262, https://doi.org/https://doi.org/10.1029/1999JD900024, \_\_eprint: https://agupubs.onlinelibrary.wiley.com/doi/pdf/10.1029/1999JD900024, 1999.
- de Meij, A., Pozzer, A., Pringle, K., Tost, H., and Lelieveld, J.: EMAC model evaluation and analysis of atmospheric
   800 aerosol properties and distribution with a focus on the Mediterranean region, Atmospheric Research, 114-115, 38–69, https://doi.org/10.1016/j.atmosres.2012.05.014, 2012.
  - Deshler, T.: Thirty years of in situ stratospheric aerosol size distribution measurements from Laramie, Wyoming (41°N), using balloon-borne instruments, Journal of Geophysical Research, 108, 4167, https://doi.org/10.1029/2002JD002514, 2003.
  - Dhomse, S. S., Emmerson, K. M., Mann, G. W., Bellouin, N., Carslaw, K. S., Chipperfield, M. P., Hommel, R., Abraham, N. L., Telford, P.,
- 805 Braesicke, P., Dalvi, M., Johnson, C. E., O'Connor, F., Morgenstern, O., Pyle, J. A., Deshler, T., Zawodny, J. M., and Thomason, L. W.: Aerosol microphysics simulations of the Mt. Pinatubo eruption with the UM-UKCA composition-climate model, Atmospheric Chemistry and Physics, 14, 11 221–11 246, https://doi.org/10.5194/acp-14-11221-2014, wOS:000344165800017, 2014.
  - Dhomse, S. S., Mann, G. W., Antuña Marrero, J. C., Shallcross, S. E., Chipperfield, M. P., Carslaw, K. S., Marshall, L., Abraham, N. L., and Johnson, C. E.: Evaluating the simulated radiative forcings, aerosol properties and stratospheric warmings from the 1963 Agung, 1982

- 810 El Chichón and 1991 Mt Pinatubo volcanic aerosol clouds, preprint, Aerosols/Atmospheric Modelling/Stratosphere/Chemistry (chemical composition and reactions), https://doi.org/10.5194/acp-2020-344, 2020.
  - Driscoll, S., Bozzo, A., Gray, L. J., Robock, A., and Stenchikov, G.: Coupled Model Intercomparison Project 5 (CMIP5) simulations of climate following volcanic eruptions, Journal of Geophysical Research: Atmospheres, 117, https://doi.org/10.1029/2012JD017607, publisher: John Wiley & Sons, Ltd, 2012.
- 815 Dutton, E. G. and Christy, J. R.: Solar radiative forcing at selected locations and evidence for global lower tropospheric cooling following the eruptions of El Chichón and Pinatubo, Geophysical Research Letters, 19, 2313–2316, https://doi.org/10.1029/92GL02495, 1992.
  - English, J. M., Toon, O. B., and Mills, M. J.: Microphysical simulations of large volcanic eruptions: Pinatubo and Toba, Journal of Geophysical Research-Atmospheres, 118, 1880–1895, https://doi.org/10.1002/jgrd.50196, wOS:000317841000022, 2013.
- Eyring, V., Bony, S., Meehl, G. A., Senior, C. A., Stevens, B., Stouffer, R. J., and Taylor, K. E.: Overview of the Coupled Model
  Intercomparison Project Phase 6 (CMIP6) experimental design and organization, Geoscientific Model Development, 9, 1937–1958, https://doi.org/10.5194/gmd-9-1937-2016, 2016.
  - Fisher, B. L., Krotkov, N. A., Bhartia, P. K., Li, C., Carn, S. A., Hughes, E., and Leonard, P. J. T.: A new discrete wavelength backscattered ultraviolet algorithm for consistent volcanic SO<sub&gt;2&lt;/sub&gt; retrievals from multiple satellite missions, Atmospheric Measurement Techniques, 12, 5137–5153, https://doi.org/10.5194/amt-12-5137-2019, 2019.
- Fountoukis, C. and Nenes, A.: ISORROPIA II: a computationally efficient thermodynamic equilibrium model for  $K^+$   $Ca^{2+}$   $Mg^{2+}$   $NH_4^+$ -  $Na^+$  -  $SO_4^{2-}$  -  $NO_3^-$  -  $Cl^-$  -  $H_2O$  aerosols, Atmos. Chem. Phys., 7, 4639–4659, https://doi.org/10.5194/acp-7-4639-2007, 2007.
  - Fouquart, Y. and Bonnel, B.: Computations of solar heating of the earth's atmosphere: a new parameterization, Beitrage zur Physik der Atmosphare, 53, 35–62, 1980.
- Fujiwara, M., Gloria L. Manney, Lesley J. Gray, and onathon S. Wright: SPARC Reanalysis Intercomparison Project (S-RIP) Final Report,
   Tech. Rep. SPARC Report No. 10, WCRP-6/2021, https://doi.org/10.17874/800dee57d13, 2022.
  - Gao, C., Oman, L., Robock, A., and Stenchikov, G. L.: Atmospheric volcanic loading derived from bipolar ice cores: Accounting for the spatial distribution of volcanic deposition, Journal of Geophysical Research, 112, D09 109, https://doi.org/10.1029/2006JD007461, 2007.
  - Gelaro, R., McCarty, W., Suárez, M. J., Todling, R., Molod, A., Takacs, L., Randles, C. A., Darmenov, A., Bosilovich, M. G., Reichle, R., Wargan, K., Coy, L., Cullather, R., Draper, C., Akella, S., Buchard, V., Conaty, A., da Silva, A. M., Gu, W., Kim, G.-K., Koster, R.,
- Lucchesi, R., Merkova, D., Nielsen, J. E., Partyka, G., Pawson, S., Putman, W., Rienecker, M., Schubert, S. D., Sienkiewicz, M., and Zhao,
   B.: The Modern-Era Retrospective Analysis for Research and Applications, Version 2 (MERRA-2), Journal of Climate, 30, 5419–5454, https://doi.org/10.1175/JCLI-D-16-0758.1, 2017.
  - Good, P. and Pyle, J.: Refinements in the use of equivalent latitude for assimilating sporadic inhomogeneous stratospheric tracer observations, 1: Detecting transport of Pinatubo aerosol across a strong vortex edge, Atmospheric Chemistry and Physics, 4, 1823–1836, wOS:000223829000003, 2004.

- Graft, H.-F., Kirchner, I., Robock, A., and Schult, I.: Pinatubo eruption winter climate effects: model versus observations, Climate Dynamics, 9, 81–93, https://doi.org/10.1007/BF00210011, 1993.
- Gu, Y., Rose, W. I., and Bluth, G. J. S.: Retrieval of mass and sizes of particles in sandstorms using two MODIS IR bands: A case study of April 7, 2001 sandstorm in China, Geophysical Research Letters, 30, https://doi.org/10.1029/2003GL017405, 2003.
- 845 Guo, S., Bluth, G. J. S., Rose, W. I., Watson, I. M., and Prata, A. J.: Re-evaluation of SO2 release of the 15 June 1991 Pinatubo eruption using ultraviolet and infrared satellite sensors, Geochemistry, Geophysics, Geosystems, 5, https://doi.org/10.1029/2003GC000654, 2004a.

- Guo, S., Rose, W. I., Bluth, G. J. S., and Watson, I. M.: Particles in the great Pinatubo volcanic cloud of June 1991: The role of ice: JUNE 1991 PINATUBO VOLCANIC CLOUDS, Geochemistry, Geophysics, Geosystems, 5, n/a–n/a, https://doi.org/10.1029/2003GC000655, 2004b.
- 850 Hansen, J., Lacis, A., Ruedy, R., and Sato, M.: Potential climate impact of Mount Pinatubo eruption, Geophysical Research Letters, 19, 215–218, https://doi.org/10.1029/91GL02788, 1992.
  - Iacono, M. J., Delamere, J. S., Mlawer, E. J., Shephard, M. W., Clough, S. A., and Collins, W. D.: Radiative forcing by longlived greenhouse gases: Calculations with the AER radiative transfer models, Journal of Geophysical Research, 113, D13103, https://doi.org/10.1029/2008JD009944, 2008.
- Joeckel, P., Sander, R., Kerkweg, A., Tost, H., and Lelieveld, J.: Technical Note: The Modular Earth Submodel System (MESSy) a new approach towards Earth System Modeling, Atmospheric Chemistry and Physics, 5, 433–444, https://doi.org/10.5194/acp-5-433-2005, 2005.
  - Joeckel, P., Tost, H., Pozzer, A., Bruehl, C., Buchholz, J., Ganzeveld, L., Hoor, P., Kerkweg, A., Lawrence, M., Sander, R., Steil, B., Stiller, G., Tanarhte, M., Taraborrelli, D., van Aardenne, J., and Lelieveld, J.: The atmospheric chemistry general circulation model
- 860 ECHAM5/MESSy1: consistent simulation of ozone from the surface to the mesosphere, Atmospheric Chemistry and Physics, 6, 5067– 5104, https://doi.org/10.5194/acp-6-5067-2006, 2006.
  - Joeckel, P., Kerkweg, A., Pozzer, A., Sander, R., Tost, H., Riede, H., Baumgaertner, A., Gromov, S., and Kern, B.: Development cycle 2 of the Modular Earth Submodel System (MESSy2), Geoscientific Model Development, 3, 717–752, https://doi.org/10.5194/gmd-3-717-2010, 2010.
- 865 Joshi, M. M. and Jones, G. S.: The climatic effects of the direct injection of water vapour into the stratosphere by large volcanic eruptions, Atmos. Chem. Phys., 9, 6109–6118, https://doi.org/10.5194/acp-9-6109-2009, 2009.
  - Karpechko, A. Y., Gillett, N. P., Dall'Amico, M., and Gray, L. J.: Southern Hemisphere atmospheric circulation response to the El Chichon and Pinatubo eruptions in coupled climate models, Quarterly Journal of the Royal Meteorological Society, 136, 1813–1822, https://doi.org/10.1002/qj.683, wOS:000284039900013, 2010.
- 870 Kerkweg, A., Buchholz, J., Ganzeveld, L., Pozzer, A., Tost, H., and Jöckel, P.: Technical Note: An implementation of the dry removal processes DRY DEPosition and SEDImentation in the Modular Earth Submodel System (MESSy), Atmos. Chem. Phys., 6, 4617–4632, https://doi.org/10.5194/acp-6-4617-2006, 2006a.
  - Kerkweg, A., Sander, R., Tost, H., and Jöckel, P.: Technical note: Implementation of prescribed (OFFLEM), calculated (ONLEM), and pseudo-emissions (TNUDGE) of chemical species in the Modular Earth Submodel System (MESSy), Atmospheric Chemistry and Physics,
- 6, 3603–3609, https://doi.org/10.5194/acp-6-3603-2006, 2006b.
  - Kilian, M., Brinkop, S., and Jöckel, P.: Impact of the eruption of Mt Pinatubo on the chemical composition of the stratosphere, Atmospheric Chemistry and Physics, 20, 11 697–11 715, https://doi.org/10.5194/acp-20-11697-2020, 2020.
    - Kinnison, D. E., Grant, K. E., Connell, P. S., Rotman, D. A., and Wuebbles, D. J.: The chemical and radiative effects of the Mount Pinatubo eruption, Journal of Geophysical Research: Atmospheres, 99, 25705–25731, https://doi.org/10.1029/94JD02318, 1994.
- 880 Kirchner, I., Stenchikov, G. L., Graf, H.-F., Robock, A., and Antuña, J. C.: Climate model simulation of winter warming and summer cooling following the 1991 Mount Pinatubo volcanic eruption, Journal of Geophysical Research: Atmospheres, 104, 19039–19055, https://doi.org/10.1029/1999JD900213, 1999.

- Klingmüller, K., Steil, B., Brühl, C., Tost, H., and Lelieveld, J.: Sensitivity of aerosol radiative effects to different mixing assumptions in the AEROPT 1.0 submodel of the EMAC atmospheric-chemistry–climate model, Geoscientific Model Development, 7, 2503–2516, https://doi.org/10.5194/gmd-7-2503-2014, 2014.
- Klingmüller, K., Lelieveld, J., Karydis, V. A., and Stenchikov, G. L.: Direct radiative effect of dust-pollution interactions, Atmospheric Chemistry and Physics, 19, 7397–7408, https://doi.org/10.5194/acp-19-7397-2019, 2019.
  - Klingmüller, K., Karydis, V. A., Bacer, S., Stenchikov, G. L., and Lelieveld, J.: Weaker cooling by aerosols due to dust–pollution interactions, Atmospheric Chemistry and Physics, 20, 15285–15295, https://doi.org/10.5194/acp-20-15285-2020, 2020.
- 890 Kodera, K. and Kuroda, Y.: Tropospheric and stratospheric aspects of the Arctic oscillation, Geophysical Research Letters, 27, 3349–3352, https://doi.org/10.1029/2000GL012017, 2000.
  - Kremser, S., Thomason, L. W., von Hobe, M., Hermann, M., Deshler, T., Timmreck, C., Toohey, M., Stenke, A., Schwarz, J. P., Weigel, R., Fueglistaler, S., Prata, F. J., Vernier, J.-P., Schlager, H., Barnes, J. E., Antuña-Marrero, J.-C., Fairlie, D., Palm, M., Mahieu, E., Notholt, J., Rex, M., Bingen, C., Vanhellemont, F., Bourassa, A., Plane, J. M. C., Klocke, D., Carn, S. A., Clarisse, L., Trickl, T., Neely, R., James,
- A. D., Rieger, L., Wilson, J. C., and Meland, B.: Stratospheric aerosol-Observations, processes, and impact on climate: Stratospheric Aerosol, Reviews of Geophysics, 54, 278–335, https://doi.org/10.1002/2015RG000511, 2016.
  - Krueger, A. J., Walter, L. S., Bhartia, P. K., Schnetzler, C. C., Krotkov, N. A., Sprod, I., and Bluth, G. J. S.: Volcanic sulfur dioxide measurements from the total ozone mapping spectrometer instruments, Journal of Geophysical Research: Atmospheres, 100, 14057–14076, https://doi.org/10.1029/95JD01222, 1995.
- 900 Labitzke, K. and McCormick, M. P.: Stratospheric temperature increases due to Pinatubo aerosols, Geophysical Research Letters, 19, 207–210, https://doi.org/10.1029/91GL02940, 1992.
  - Landgraf, J. and Crutzen, P. J.: An Efficient Method for Online Calculations of Photolysis and Heating Rates, Journal of the Atmospheric Sciences, 55, 863–878, https://doi.org/10.1175/1520-0469(1998)055<0863:AEMFOC>2.0.CO;2, 1998.

LeGrande, A. N., Tsigaridis, K., and Bauer, S. E.: Role of atmospheric chemistry in the climate impacts of stratospheric volcanic injections,

905 Nature Geoscience, 9, 652–655, https://doi.org/10.1038/ngeo2771, 2016.

885

Löffler, M., Brinkop, S., and Jöckel, P.: Impact of major volcanic eruptions on stratospheric water vapour, Atmospheric Chemistry and Physics, 16, 6547–6562, https://doi.org/10.5194/acp-16-6547-2016, 2016.

- Long, C. S. and Stowe, L. L.: using the NOAA/AVHRR to study stratospheric aerosol optical thicknesses following the Mt. Pinatubo Eruption, Geophysical Research Letters, 21, 2215–2218, https://doi.org/10.1029/94GL01322, publisher: John Wiley & Sons, Ltd, 1994.
- 910 Lovejoy, E. R., Hanson, D. R., and Huey, L. G.: Kinetics and Products of the Gas-Phase Reaction of SO3 with Water, The Journal of Physical Chemistry, 100, 19911–19916, https://doi.org/10.1021/jp962414d, publisher: American Chemical Society, 1996.
  - Mao, J. and Robock, A.: Surface Air Temperature Simulations by AMIP General Circulation Models: Volcanic and ENSO Signals and Systematic Errors, Journal of Climate, 11, 1538–1552, https://doi.org/10.1175/1520-0442(1998)011<1538:SATSBA>2.0.CO;2, 1998.
- Marandino, C. A., Tegtmeier, S., Krüger, K., Zindler, C., Atlas, E. L., Moore, F., and Bange, H. W.: Dimethylsulphide (DMS) emissions
   from the western Pacific Ocean: a potential marine source for stratospheric sulphur?, Atmospheric Chemistry and Physics, 13, 8427– 8437, https://doi.org/https://doi.org/10.5194/acp-13-8427-2013, 2013.
  - Marshall, L., Schmidt, A., Toohey, M., Carslaw, K. S., Mann, G. W., Sigl, M., Khodri, M., Timmreck, C., Zanchettin, D., Ball, W. T.,
     Bekki, S., Brooke, J. S. A., Dhomse, S., Johnson, C., Lamarque, J.-F., LeGrande, A. N., Mills, M. J., Niemeier, U., Pope, J. O.,
     Poulain, V., Robock, A., Rozanov, E., Stenke, A., Sukhodolov, T., Tilmes, S., Tsigaridis, K., and Tummon, F.: Multi-model comparison

920 of the volcanic sulfate deposition from the 1815 eruption of Mt. Tambora, Atmospheric Chemistry and Physics, 18, 2307–2328, https://doi.org/10.5194/acp-18-2307-2018, 2018.

McCormick, M. P.: Sage II: An overview, Advances in Space Research, 7, 219–226, https://doi.org/10.1016/0273-1177(87)90151-7, 1987.
McCormick, M. P., Thomason, L. W., and Trepte, C. R.: Atmospheric effects of the Mt Pinatubo eruption, Nature, 373, 399, http://dx.doi.org/10.1038/373399a0, 1995.

- 925 Metzger, S., Steil, B., Abdelkader, M., Klingmüller, K., Xu, L., Penner, J. E., Fountoukis, C., Nenes, A., and Lelieveld, J.: Aerosol water parameterisation: a single parameter framework, Atmospheric Chemistry and Physics, 16, 7213–7237, https://doi.org/10.5194/acp-16-7213-2016, 2016.
  - Mills, M. J., Schmidt, A., Easter, R., Solomon, S., Kinnison, D. E., Ghan, S. J., Neely, R. R., Marsh, D. R., Conley, A., Bardeen, C. G., and Gettelman, A.: Global volcanic aerosol properties derived from emissions, 1990–2014, using CESM1(WACCM), Journal of Geophysical Research: Atmospheres, 121, 2015JD024 290, https://doi.org/10.1002/2015JD024290, 2016.
- Mills, M. J., Richter, J. H., Tilmes, S., Kravitz, B., MacMartin, D. G., Glanville, A. A., Tribbia, J. J., Lamarque, J.-F., Vitt, F., Schmidt, A., Gettelman, A., Hannay, C., Bacmeister, J. T., and Kinnison, D. E.: Radiative and Chemical Response to Interactive Stratospheric Sulfate Aerosols in Fully Coupled CESM1(WACCM), Journal of Geophysical Research: Atmospheres, 122, https://doi.org/10.1002/2017JD027006, 2017.

930

- 935 Muser, L. O., Hoshyaripour, G. A., Bruckert, J., Horváth, Á., Malinina, E., Wallis, S., Prata, F. J., Rozanov, A., von Savigny, C., Vogel, H., and Vogel, B.: Particle aging and aerosol–radiation interaction affect volcanic plume dispersion: evidence from the Raikoke 2019 eruption, Atmospheric Chemistry and Physics, 20, 15015–15036, https://doi.org/10.5194/acp-20-15015-2020, 2020.
  - Muthers, S., Arfeuille, F., Raible, C. C., and Rozanov, E.: The impacts of volcanic aerosol on stratospheric ozone and the Northern Hemisphere polar vortex: separating radiative-dynamical changes from direct effects due to enhanced aerosol heterogeneous chemistry, Atmospheric Chemistry and Physics, 15, 11 461–11 476, https://doi.org/10.5194/acp-15-11461-2015, 2015.
- Nagai, T., Liley, B., Sakai, T., Shibata, T., and Uchino, O.: Post-Pinatubo Evolution and Subsequent Trend of the Stratospheric Aerosol Layer Observed by Mid-Latitude Lidars in Both Hemispheres, Sola, 6, 69–72, https://doi.org/10.2151/sola.2010-018, wOS:000278466500003, 2010.
- Nedoluha, G. E., Bevilacqua, R. M., Gomez, R. M., Siskind, D. E., Hicks, B. C., Russell, J. M., and Connor, B. J.: Increases in middle
   atmospheric water vapor as observed by the Halogen Occultation Experiment and the ground-based Water Vapor Millimeter-Wave Spec
  - trometer from 1991 to 1997, Journal of Geophysical Research: Atmospheres, 103, 3531–3543, https://doi.org/10.1029/97JD03282, 1998. Niemeier, U., Timmreck, C., Graf, H.-F., Kinne, S., Rast, S., and Self, S.: Initial fate of fine ash and sulfur from large volcanic eruptions, Atmospheric Chemistry and Physics, 9, 9043–9057, https://doi.org/10.5194/acp-9-9043-2009, 2009.
- Niemeier, U., Riede, F., and Timmreck, C.: Simulation of ash clouds after a Laacher See-type eruption, Climate of the Past, 17, 633–652, https://doi.org/10.5194/cp-17-633-2021, 2021.
  - Oltmans, S. J. and Hofmann, D. J.: Increase in lower-stratospheric water vapour at a mid-latitude Northern Hemisphere site from 1981 to 1994, Nature, 374, 146–149, https://doi.org/10.1038/374146a0, 1995.
- Oman, L., Robock, A., Stenchikov, G. L., Thordarson, T., Koch, D., Shindell, D. T., and Gao, C.: Modeling the distribution of the volcanic aerosol cloud from the 1783–1784 Laki eruption, Journal of Geophysical Research: Atmospheres, 111, D12 209, https://doi.org/10.1029/2005JD006899, 2006.
  - 29

- Osipov, S., Stenchikov, G., Tsigaridis, K., LeGrande, A. N., and Bauer, S. E.: The Role of the SO Radiative Effect in Sustaining the Volcanic Winter and Soothing the Toba Impact on Climate, Journal of Geophysical Research: Atmospheres, 125, https://doi.org/10.1029/2019JD031726, 2020.
- Osipov, S., Stenchikov, G., Tsigaridis, K., LeGrande, A. N., Bauer, S. E., Fnais, M., and Lelieveld, J.: The Toba supervolcano eruption caused
   severe tropical stratospheric ozone depletion, Communications Earth & Environment, 2, 71, https://doi.org/10.1038/s43247-021-00141-7, 2021.
  - Poberaj, C. S., Staehelin, J., and Brunner, D.: Missing Stratospheric Ozone Decrease at Southern Hemisphere Middle Latitudes after Mt. Pinatubo: A Dynamical Perspective, Journal of the Atmospheric Sciences, 68, 1922–1945, https://doi.org/10.1175/JAS-D-10-05004.1, 2011.
- 965 Pollack, J. B., Toon, O. B., and Khare, B. N.: Optical properties of some terrestrial rocks and glasses, Icarus, 19, 372–389, https://doi.org/10.1016/0019-1035(73)90115-2, 1973.
  - Polvani, L. M., Banerjee, A., and Schmidt, A.: Northern Hemisphere continental winter warming following the 1991 Mt. Pinatubo eruption: reconciling models and observations, Atmospheric Chemistry and Physics, 19, 6351–6366, https://doi.org/10.5194/acp-19-6351-2019, 2019.
- 970 Pozzer, A., Jöckel, P., Sander, R., Williams, J., Ganzeveld, L., and Lelieveld, J.: Technical Note: The MESSy-submodel AIRSEA calculating the air-sea exchange of chemical species, Atmospheric Chemistry and Physics, 6, 5435–5444, https://doi.org/10.5194/acp-6-5435-2006, 2006.
  - Pozzer, A., de Meij, A., Pringle, K. J., Tost, H., Doering, U. M., van Aardenne, J., and Lelieveld, J.: Distributions and regional budgets of aerosols and their precursors simulated with the EMAC chemistry-climate model, Atmospheric Chemistry and Physics, 12, 961–987,
- 975 https://doi.org/10.5194/acp-12-961-2012, 2012.
  - Prather, M.: Catastrophic loss of stratospheric ozone in dense volcanic clouds, Journal of Geophysical Research: Atmospheres, 97, 10187– 10191, https://doi.org/10.1029/92JD00845, 1992.
  - Predybaylo, E., Stenchikov, G. L., Wittenberg, A. T., and Zeng, F.: Impacts of a Pinatubo-size volcanic eruption on ENSO: VOLCANIC IMPACTS ON ENSO, Journal of Geophysical Research: Atmospheres, 122, 925–947, https://doi.org/10.1002/2016JD025796, 2017.
- 980 Pringle, K. J., Tost, H., Metzger, S., Steil, B., Giannadaki, D., Nenes, A., Fountoukis, C., Stier, P., Vignati, E., and Lelieveld, J.: Description and evaluation of GMXe: a new aerosol submodel for global simulations (v1), Geoscientific Model Development, 3, 391–412, https://doi.org/10.5194/gmd-3-391-2010, 2010.
  - Pueschel, R. F., Russell, P. B., Allen, D. A., Ferry, G. V., Snetsinger, K. G., Livingston, J. M., and Verma, S.: Physical and optical properties of the Pinatubo volcanic aerosol: Aircraft observations with impactors and a Sun-tracking
- 985 photometer, Journal of Geophysical Research: Atmospheres, 99, 12915–12922, https://doi.org/10.1029/94JD00621, \_eprint: https://agupubs.onlinelibrary.wiley.com/doi/pdf/10.1029/94JD00621, 1994.
  - Ramachandran, S., Ramaswamy, V., Stenchikov, G. L., and Robock, A.: Radiative impact of the Mount Pinatubo volcanic eruption: Lower stratospheric response, Journal of Geophysical Research: Atmospheres, 105, 24409–24429, https://doi.org/10.1029/2000JD900355, 2000.
- 990 Rinsland, C. P., Gunson, M. R., Ko, M. K. W., Weisenstein, D. W., Zander, R., Abrams, M. C., Goldman, A., Sze, N. D., and Yue, G. K.: H<sub>2</sub> SO 4 photolysis: A source of sulfur dioxide in the upper stratosphere, Geophysical Research Letters, 22, 1109–1112, https://doi.org/10.1029/95GL00917, 1995.
  - Robock, A.: Volcanic eruptions and climate, Reviews of Geophysics, 38, 191-219, https://doi.org/10.1029/1998RG000054, 2000.

Roeckner, E. and Coauthors: The atmospheric general circulation model ECHAM5. Part I: Model description. Max Planck Institute for

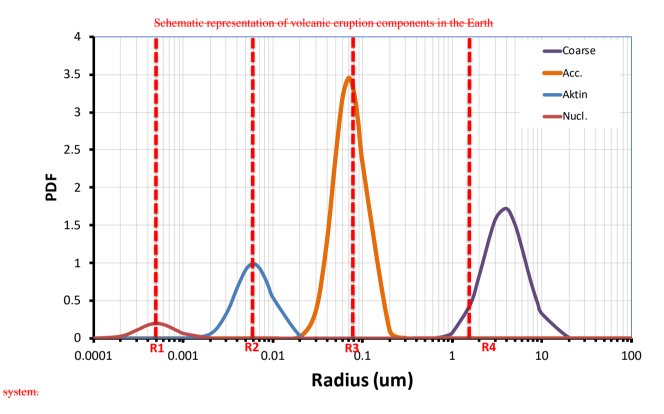
- 995 Meteorology, Tech. Rep. 349, MPI for Meteorology, Hamburg, Germany, https://pure.mpg.de/rest/items/item\_995269\_6/component/file\_ 3192562/content, 2003.
  - Roeckner, E., Brokopf, R., Esch, M., Giorgetta, M., Hagemann, S., Kornblueh, L., Manzini, E., Schlese, U., and Schulzweida, U.: Sensitivity of Simulated Climate to Horizontal and Vertical Resolution in the ECHAM5 Atmosphere Model, Journal of Climate, 19, 3771–3791, https://doi.org/10.1175/JCLI3824.1, 2006.
- 1000 Rose and Durant, A.: Fine ash content of explosive eruptions, Journal of Volcanology and Geothermal Research, 186, 32–39, https://doi.org/10.1016/j.jvolgeores.2009.01.010, 2009.
  - Rose, W. I., Millard, G. A., Mather, T. A., Hunton, D. E., Anderson, B., Oppenheimer, C., Thornton, B. F., Gerlach, T. M., Viggiano, A. A., Kondo, Y., Miller, T. M., and Ballenthin, J. O.: Atmospheric chemistry of a 33–34 hour old volcanic cloud from Hekla Volcano (Iceland): Insights from direct sampling and the application of chemical box modeling, Journal of Geophysical Research: Atmospheres,
- 1005 111, https://doi.org/10.1029/2005JD006872, 2006.
- Russell, P. B., Livingston, J. M., Dutton, E. G., Pueschel, R. F., Reagan, J. A., DeFoor, T. E., Box, M. A., Allen, D., Pilewskie, P., Herman, B. M., Kinne, S. A., and Hofmann, D. J.: Pinatubo and pre-Pinatubo optical-depth spectra: Mauna Loa measurements, comparisons, inferred particle size distributions, radiative effects, and relationship to lidar data, Journal of Geophysical Research: Atmospheres, 98, 22 969–22 985, https://doi.org/10.1029/93JD02308, 1993.
- 1010 Russell, P. B., Livingston, J. M., Pueschel, R. F., Bauman, J. J., Pollack, J. B., Brooks, S. L., Hamill, P., Thomason, L. W., Stowe, L. L., Deshler, T., Dutton, E. G., and Bergstrom, R. W.: Global to microscale evolution of the Pinatubo volcanic aerosol derived from diverse measurements and analyses, Journal of Geophysical Research: Atmospheres, 101, 18745–18763, https://doi.org/10.1029/96JD01162, 1996.
- Sander, R., Kerkweg, A., Jöckel, P., and Lelieveld, J.: Technical note: The new comprehensive atmospheric chemistry module MECCA,
   Atmospheric Chemistry and Physics, 5, 445–450, https://doi.org/10.5194/acp-5-445-2005, 2005.
- Sander, R., Baumgaertner, A., Gromov, S., Harder, H., Jöckel, P., Kerkweg, A., Kubistin, D., Regelin, E., Riede, H., Sandu, A., Taraborrelli, D., Tost, H., and Xie, Z.-Q.: The atmospheric chemistry box model CAABA/MECCA-3.0, Geoscientific Model Development, 4, 373–380, https://doi.org/10.5194/gmd-4-373-2011, publisher: Copernicus GmbH, 2011.
- Sander, R., Jöckel, P., Kirner, O., Kunert, A. T., Landgraf, J., and Pozzer, A.: The photolysis module JVAL-14, compatible with the MESSy
   standard, and the JVal PreProcessor (JVPP), Geoscientific Model Development, 7, 2653–2662, https://doi.org/10.5194/gmd-7-2653-2014, 2014.
  - Sander, R., Baumgaertner, A., Cabrera-Perez, D., Frank, F., Gromov, S., Grooß, J.-U., Harder, H., Huijnen, V., Jöckel, P., Karydis, V. A., Niemeyer, K. E., Pozzer, A., Riede, H., Schultz, M. G., Taraborrelli, D., and Tauer, S.: The community atmospheric chemistry box model CAABA/MECCA-4.0, Geoscientific Model Development, 12, 1365–1385, https://doi.org/10.5194/gmd-12-1365-2019, 2019.
- 1025 Seinfeld, J. H. and Pandis, S. N.: Atmospheric chemistry and physics : from air pollution to climate change, J. Wiley, Hoboken, N.J., 2006. Sheng, J.-X., Weisenstein, D. K., Luo, B.-P., Rozanov, E., Arfeuille, F., and Peter, T.: A perturbed parameter model ensemble to investigate Mt. Pinatubo's 1991 initial sulfur mass emission, Atmospheric Chemistry and Physics, 15, 11501–11512, https://doi.org/10.5194/acp-15-11501-2015, wOS:000364316800004, 2015.

Shindell, D. T.: Dynamic winter climate response to large tropical volcanic eruptions since 1600, Journal of Geophysical Research, 109, https://doi.org/10.1029/2003JD004151, 2004.

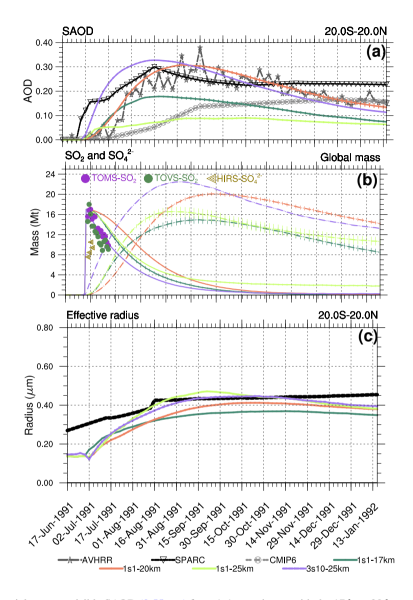
- Shindell, D. T., Schmidt, G. A., Miller, R. L., and Rind, D.: Northern hemisphere winter climate response to greenhouse gas, ozone, solar, and volcanic forcing, Journal of Geophysical Research: Atmospheres, 106, 7193–7210, https://doi.org/10.1029/2000JD900547, 2001.
- Soden, B. J.: Global Cooling After the Eruption of Mount Pinatubo: A Test of Climate Feedback by Water Vapor, Science, 296, 727–730, https://doi.org/10.1126/science.296.5568.727, 2002.
- 1035 Stenchikov, G.: Arctic Oscillation response to the 1991 Mount Pinatubo eruption: Effects of volcanic aerosols and ozone depletion, Journal of Geophysical Research, 107, https://doi.org/10.1029/2002JD002090, 2002.
  - Stenchikov, G.: The role of volcanic activity in climate and global changes, in: Climate Change, pp. 607–643, Elsevier, https://doi.org/10.1016/B978-0-12-821575-3.00029-3, 2021.
- Stenchikov, G., Kirchner, I., Robock, A., Graf, H.-F., Antuña, J. C., Grainger, R. G., Lambert, A., and Thomason, L.: Radiative
  forcing from the 1991 Mount Pinatubo volcanic eruption, Journal of Geophysical Research: Atmospheres, 103, 13837–13857, https://doi.org/10.1029/98JD00693, publisher: John Wiley & Sons, Ltd, 1998.
  - Stenchikov, G., Hamilton, K., Robock, A., Ramaswamy, V., and Schwarzkopf, M. D.: Arctic oscillation response to the 1991 Pinatubo eruption in the SKYHI general circulation model with a realistic quasi-biennial oscillation, Journal of Geophysical Research-Atmospheres, 109, D03 112, https://doi.org/10.1029/2003JD003699, wOS:000189052800001, 2004.
- 1045 Stenchikov, G., Hamilton, K., Stouffer, R. J., Robock, A., Ramaswamy, V., Santer, B., and Graf, H.-F.: Arctic Oscillation response to volcanic eruptions in the IPCC AR4 climate models, Journal of Geophysical Research, 111, https://doi.org/10.1029/2005JD006286, 2006.
  - Stenchikov, G., Ukhov, A., Osipov, S., Ahmadov, R., Grell, G., Cady-Pereira, K., Mlawer, E., and Iacono, M.: How does a Pinatubo-size Volcanic Cloud Reach the Middle Stratosphere?, Journal of Geophysical Research: Atmospheres, https://doi.org/10.1029/2020JD033829, 2021.
- 1050 Sukhodolov, T., Sheng, J.-X., Feinberg, A., Luo, B.-P., Peter, T., Revell, L., Stenke, A., Weisenstein, D. K., and Rozanov, E.: Stratospheric aerosol evolution after Pinatubo simulated with a coupled size-resolved aerosol–chemistry–climate model, SOCOL-AERv1.0, Geoscientific Model Development, 11, 2633–2647, https://doi.org/10.5194/gmd-11-2633-2018, 2018.
  - Taylor, K., Williamson, D., and Zwiers, F.: The Sea Surface Temperature And Sea-Ice Concentration Boundary Conditions For AMIP II Simulations, Tech. Rep. 60, National Center for Atmospheric Research, Boulder, CO, USA, 2000.
- 1055 Thomas, M. A., Giorgetta, M. A., Timmreck, C., Graf, H.-F., and Stenchikov, G.: Simulation of the climate impact of Mt. Pinatubo eruption using ECHAM5-Part 2: Sensitivity to the phase of the QBO and ENSO, Atmospheric Chemistry and Physics, 9, 3001–3009, wOS:000266189700008, 2009.
  - Thomason: Observations of a new SAGE II aerosol extinction mode following the eruption of Mt. Pinatubo, Geophysical Research Letters, 19, 2179–2182, https://doi.org/10.1029/92GL02185, \_eprint: https://agupubs.onlinelibrary.wiley.com/doi/pdf/10.1029/92GL02185, 1992.
- 1060 Thomason and Peter, T.: SPARC Assessment of Stratospheric Aerosol Properties (ASAP), Tech. rep., SPARC Office, http://www. sparc-climate.org/publications/sparc-reports/, 2006.
  - Thomason, Poole, L. R., and Deshler, T.: A global climatology of stratospheric aerosol surface area density deduced from Stratospheric Aerosol and Gas Experiment II measurements: 1984-1994, Journal of Geophysical Research: Atmospheres, 102, 8967–8976, https://doi.org/10.1029/96JD02962, 1997.
- 1065 Timmreck, C., Mann, G. W., Aquila, V., Hommel, R., Lee, L. A., Schmidt, A., Brühl, C., Carn, S., Chin, M., Dhomse, S. S., Diehl, T., English, J. M., Mills, M. J., Neely, R., Sheng, J., Toohey, M., and Weisenstein, D.(2018)...; The Interactive Stratospheric Aerosol Model Intercomparison Project (ISA-MIP): motivation and experimental design... Geoscientific Model Development, 11(7): 2581–2608. https://doi.org/10.5194/gmd-11-2581-2018, 2018.

Toohey, M., Krüger, K., Bittner, M., Timmreck, C., and Schmidt, H.: The impact of volcanic aerosol on the Northern Hemisphere

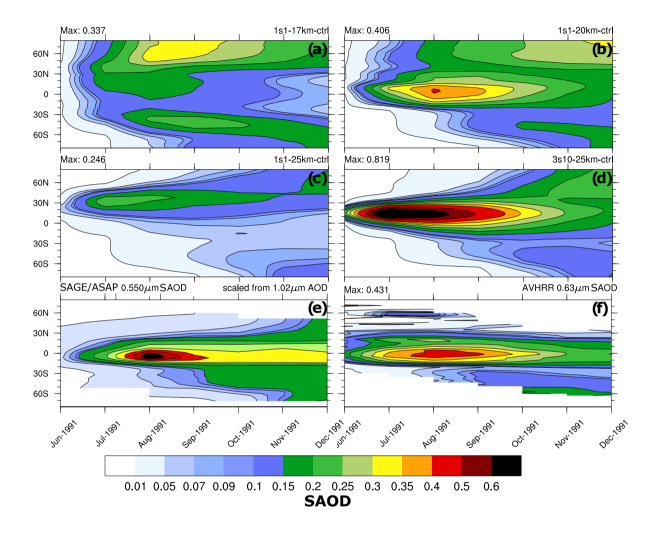
- 1070 stratospheric polar vortex: mechanisms and sensitivity to forcing structure, Atmospheric Chemistry and Physics, 14, 13063–13079, https://doi.org/10.5194/acp-14-13063-2014, 2014.
  - Tost, H., Jöckel, P., Kerkweg, A., Sander, R., and Lelieveld, J.: Technical note: A new comprehensive SCAVenging submodel for global atmospheric chemistry modelling, Atmos. Chem. Phys., 6, 565–574, https://doi.org/10.5194/acp-6-565-2006, publisher: Copernicus Publications, 2006a.
- 1075 Tost, H., Jöckel, P., and Lelieveld, J.: Influence of different convection parameterisations in a GCM, Atmospheric Chemistry and Physics, 6, 5475–5493, https://doi.org/10.5194/acp-6-5475-2006, 2006b.
  - Tost, H., Jöckel, P., and Lelieveld, J.: Lightning and convection parameterisations & amp;ndash; uncertainties in global modelling, Atmospheric Chemistry and Physics, 7, 4553–4568, https://doi.org/10.5194/acp-7-4553-2007, 2007.
- Tost, H., Lawrence, M. G., Brühl, C., Jöckel, P., Team, T. G., and Team, T. S.-O.-D.: Uncertainties in atmospheric chemistry modelling due
   to convection parameterisations and subsequent scavenging, Atmos. Chem. Phys., 10, 1931–1951, https://doi.org/10.5194/acp-10-1931-2010, 2010.
  - van der Werf, G. R., Randerson, J. T., Giglio, L., Collatz, G. J., Mu, M., Kasibhatla, P. S., Morton, D. C., DeFries, R. S., Jin, Y., and van Leeuwen, T. T.: Global fire emissions and the contribution of deforestation, savanna, forest, agricultural, and peat fires (1997–2009), Atmospheric Chemistry and Physics, 10, 11707–11735, https://doi.org/10.5194/acp-10-11707-2010, 2010.
- 1085 Vernier, J.-P., Fairlie, T. D., Deshler, T., Natarajan, M., Knepp, T., Foster, K., Wienhold, F. G., Bedka, K. M., Thomason, L., and Trepte, C.: In situ and space-based observations of the Kelud volcanic plume: The persistence of ash in the lower stratosphere, Journal of Geophysical Research: Atmospheres, 121, 11,104–11,118, https://doi.org/10.1002/2016JD025344, publisher: John Wiley & Sons, Ltd, 2016.
- Vogel, A., Diplas, S., Durant, A. J., Azar, A. S., Sunding, M. F., Rose, W. I., Sytchkova, A., Bonadonna, C., Krüger, K., and Stohl, A.: Reference data set of volcanic ash physicochemical and optical properties, Journal of Geophysical Research: Atmospheres, 122, 9485–9514, https://doi.org/10.1002/2016JD026328, 2017.
  - Walcek, C. J.: Minor flux adjustment near mixing ratio extremes for simplified yet highly accurate monotonic calculation of tracer advection, Journal of Geophysical Research: Atmospheres, 105, 9335–9348, https://doi.org/10.1029/1999JD901142, \_eprint: https://agupubs.onlinelibrary.wiley.com/doi/pdf/10.1029/1999JD901142, 2000.
- Yu, T. and Rose, W. I.: Retrieval of Sulfate and Silicate Ash Masses in Young (1 to 4 Days Old) Eruption Clouds Using
   Multiband Infrared HIRS/2 Data, in: Remote Sensing of Active Volcanism, pp. 87–100, American Geophysical Union (AGU), https://doi.org/10.1029/GM116p0087, \_eprint: https://agupubs.onlinelibrary.wiley.com/doi/pdf/10.1029/GM116p0087, 2000.
  - Zanchettin, D., Khodri, M., Timmreck, C., Toohey, M., Schmidt, A., Gerber, E. P., Hegerl, G., Robock, A., Pausata, F. S. R., Ball, W. T.,
    Bauer, S. E., Bekki, S., Dhomse, S. S., LeGrande, A. N., Mann, G. W., Marshall, L., Mills, M., Marchand, M., Niemeier, U., Poulain,
    V., Rozanov, E., Rubino, A., Stenke, A., Tsigaridis, K., and Tummon, F.: The Model Intercomparison Project on the climatic response
- 1100 toVolcanic forcing (VolMIP): experimental design and forcing input data for CMIP6, Geoscientific Model Development, 9, 2701–2719, https://doi.org/10.5194/gmd-9-2701-2016, 2016.
  - Zhu, Y., Toon, O. B., Jensen, E. J., Bardeen, C. G., Mills, M. J., Tolbert, M. A., Yu, P., and Woods, S.: Persisting volcanic ash particles impact stratospheric SO<sub>2</sub> lifetime and aerosol optical properties, Nature Communications, 11, 4526, https://doi.org/10.1038/s41467-020-18352-5, 2020.



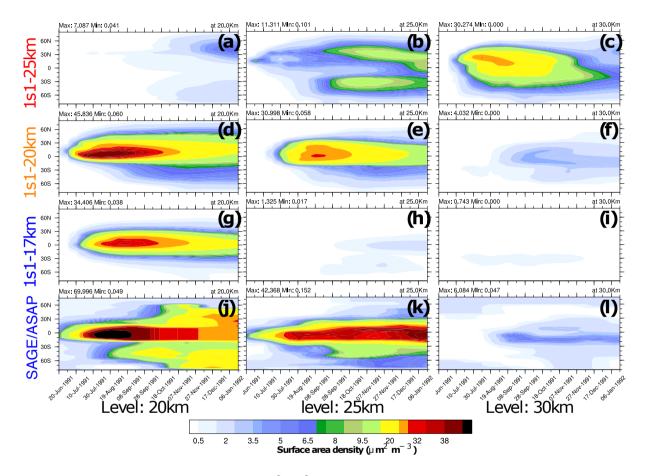
**Figure 1.** Schematic representation of the initial size distribution of aerosol modes in EMAC (nucleation, Aikten, accumulation and coarse). The threshold radii R1, R2, R3 and R4 control particle exchange between the modes. Initially 1.5 Mt of volcanic ash was injected in accumulation mode, and 73.5 Mt in coarse mode.



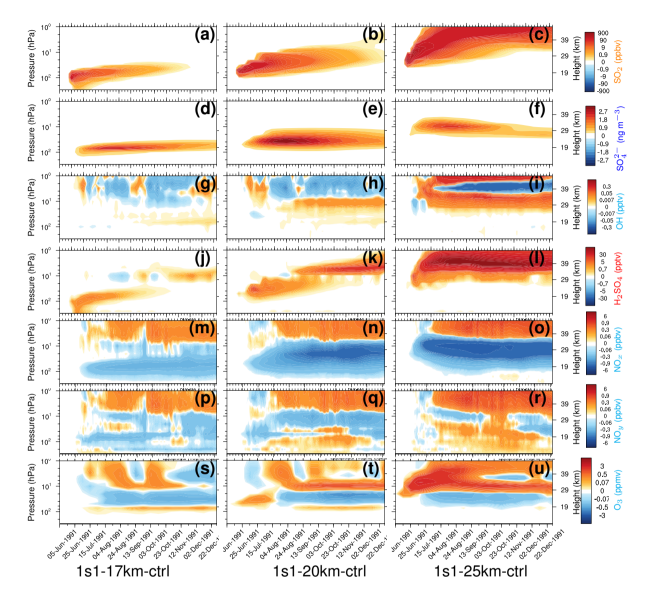
**Figure 2.** a) 20S-20N equatorial average visible SAOD ( $0.55 \mu$ m) from 1s1 experiment with the 17 km, 20 km and 25 km emission heights, AVHRR 0.630 µm, scaled visible SAGE/ASAP, and 0.525 µm CMIP6; b) SO<sub>2</sub> (solid lines) and SO<sub>4</sub><sup>2-</sup> (dashed lines) globally integrated masses calculated using output from 1s1 experiment with 17 km, 20 km and 25 km emission heights, the observed Guo et al. (2004b) SO<sub>2</sub> and SO<sub>4</sub><sup>2-</sup> masses are shown by markers; c) Equatorial average effective radius from 1s1 experiment with 17 km, 20 km and 25 km emission heights, and SAGE/ASAP (Stratospheric Processes and its Role in Climate published in the Assessment of Stratospheric Aerosol Properties) retrievals (solid black). Error bars represent one sigma from the ensemble mean.



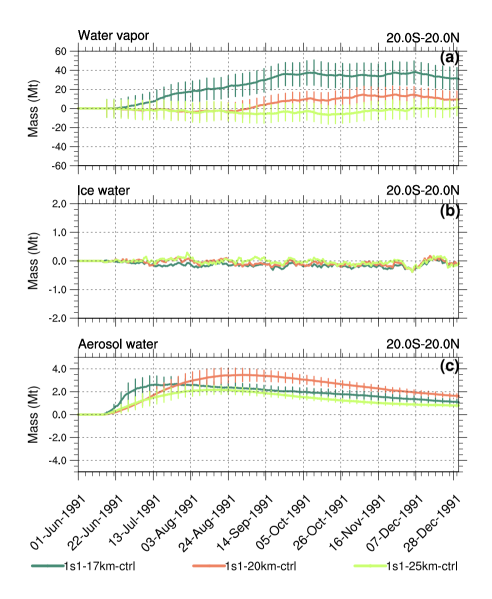
**Figure 3.** Zonally average visible SAOD ( $0.55 \mu$ m) as a function of latitude and time. a) 1s1-17km, b)1s1-20km, c) 1s1-25km, d) 3s10-25km, e) scaled visible SAGE/ASAP, f) 0.630  $\mu$ m AVHRR (Advanced Very High-Resolution Radiometer).



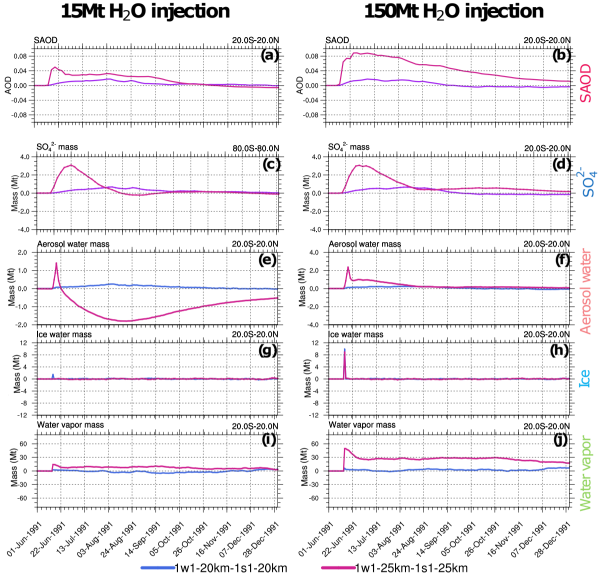
**Figure 4.** Zonally average Surface Area density (SAD,  $\mu m^{-2} cm^{-3}$ ) as a function of latitude and time at 20 km (left, a,d,g,j), 25 km (middle, b,e,h,k), and 30 km (right, c,f,i,l). For experiments: a-c) 1s1-25km, d-f) 1s1-20km, g-i) 1s1-17km, and observed j-l) SAGE/ASAP



**Figure 5.** Perturbations (with respect to control) of 20S-20N equatorial average chemical constituents as a function of pressure (from 300 hPa to 1 hPa) and time in 1s1-17km (left, a,d,g,j,m,p,s), 1s1-20km (middle, b,e,h,k,n,q,t), and 1s1-25km (right, c,f,i,l,o,r,u). a-c) SO<sub>2</sub> (ppbv), d-f) SO<sub>4</sub><sup>2-</sup> (ngm<sup>-3</sup>), g-i) OH (pptv), j-l) H<sub>2</sub>SO<sub>4</sub> (pptv), m-o) NO<sub>x</sub> (ppbv), p-r) NO<sub>y</sub> (ppbv), NO<sub>x</sub> = (NO+NO<sub>2</sub>), NO<sub>y</sub> = (NO<sub>x</sub> + NO<sub>3</sub> + HNO<sub>3</sub> + 2\*N<sub>2</sub>O<sub>5</sub> + HONO + HNO<sub>4</sub> + ClONO<sub>2</sub> + BrONO<sub>2</sub>), *S*-u) O<sub>3</sub>(ppmv).

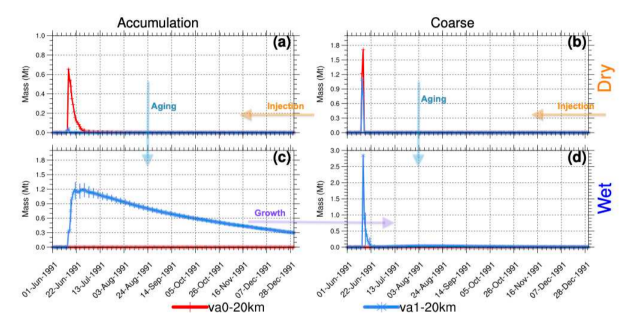


**Figure 6.** Perturbations (with respect to control) of 20S-20N stratospheric (above 100 hPa) integrated masses (Mt) in the 1s1 experiments with injection heights 17 km, 20 km and 25 km, as a function of time. a) water vapor, b) ice, c) aerosol water. Error bars represent one sigma from the ensemble mean.

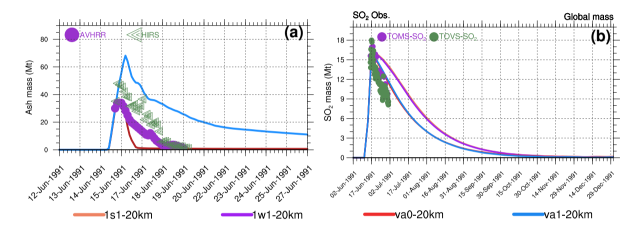


15Mt H<sub>2</sub>O injection

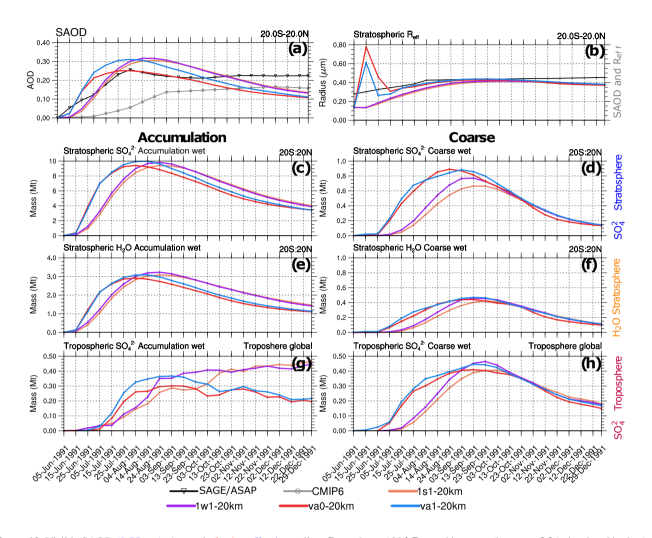
Figure 7. Perturbations (with respect to control) of 20S-20N stratospheric (above 100 hPa) SAOD and integrated masses (Mt) in the 1w1-20km and 1w1-25km experiments with respect to, correspondingly, 1s1-20km and 1s1-25km experiments as a function of time. Left column (a,c,e,g,f,ki) - 1w1 experiments with 15 Mt of volcanic water vapor injection and right column (b,d,f,h,j,+) - 1w1 experiments with 150 Mt water vapor injection. a-b) SAOD, c-d)  $SO_4^{2-}$ , e-f) aerosol water, g-h) ice, i-j) water vapor.



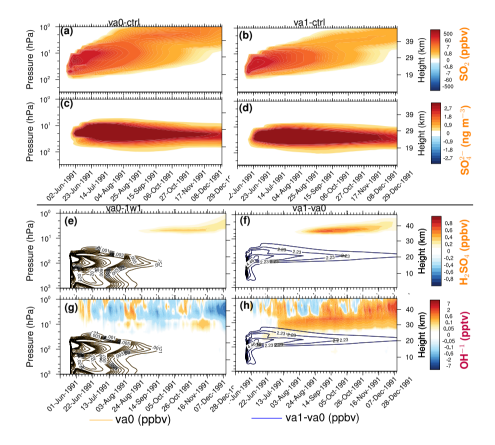
**Figure 8.** Globally integrated stratospheric volcanic ash mass (Mt) above 70 hPa as a function of time in the va0-20km and va1-20km experiments. a) Dry accumulation mode, b) Dry coarse mode, c) Wet accumulation mode, d) Wet coarse mode. Error bars represent one sigma from the ensemble mean.



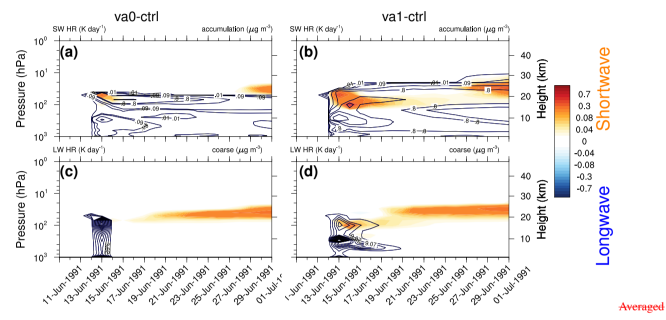
**Figure 9.** Globally integrated stratospheric masses (Mt) as a function of time. a) Volcanic ash in va0-20km, va1-20km, as well as in AVHRR, and HIRS (<u>High-Resolution Infrared Radiation Sounder</u>) retrievals Guo et al. (2004b), b) SO<sub>2</sub> in the 1s1-20km, 1w1-20km, va0-20km and va1-20km experiments, as well as in TOMS (<u>Total Ozone Mapping Spectrometer</u>) and TOVS (<u>TIROS Operational Vertical Sounder</u>) observations.



**Figure 10.** Visible SAOD (0.55 µm), Aerosol effective effective radius,  $R_{eff}$  above 100 hPa, and integrated masses (Mt) simulated in the 1s1-20km, 1w1-20km, va0-20km, va1-20km experiments as a function of time. a) simulated, as well as observed AVHRR, scaled SAGE/ASAP, and CMIP6 20S-20N SAODs, b) Simulated stratospheric (above 100 hPa)  $R_{eff}$ , as well as observed in SAGE/ASAP in 20S-20N, c) Stratospheric SO<sub>4</sub><sup>2-</sup> in accumulation mode in 20S-20N, d) Stratospheric SO<sub>4</sub><sup>2-</sup> in coarse mode in 20S-20N, e) Stratospheric aerosol water in accumulation mode in 20S-20N, f) Stratospheric aerosol water in coarse mode in 20S-20N, e) Tropospheric (below 100 hPa) SO<sub>4</sub><sup>2-</sup> in accumulation mode integrated globally, f) Tropospheric SO<sub>4</sub><sup>2-</sup> in coarse mode integrated globally.



**Figure 11.** 20S-20N average perturbations of chemical constituents as a function of pressure (from 300 hPa to 1 hPa) and time in va0-20km and va1-20km experiments. a) SO<sub>2</sub> in va0-ctr (ppbv), b) SO<sub>2</sub> in va1-ctr (ppbv), c) SO<sub>4</sub><sup>2-</sup> in va0-ctr (ngm<sup>-3</sup>), d) SO<sub>4</sub><sup>2-</sup> in va1-ctr (ngm<sup>-3</sup>), e) H<sub>2</sub>SO<sub>4</sub> in va0-1w1 (ppbv), f) H<sub>2</sub>SO<sub>4</sub> in va1-va0 (ppbv), g) OH in va0-1w1 (pptv), h) OH in va1-va0 (pptv). The contour lines shows the accumulation mode ash mixing ratio (ppbv); orange contour lines for va0 and blue contour lines for va1-va0 in the (e-h) panels.



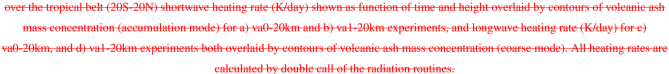
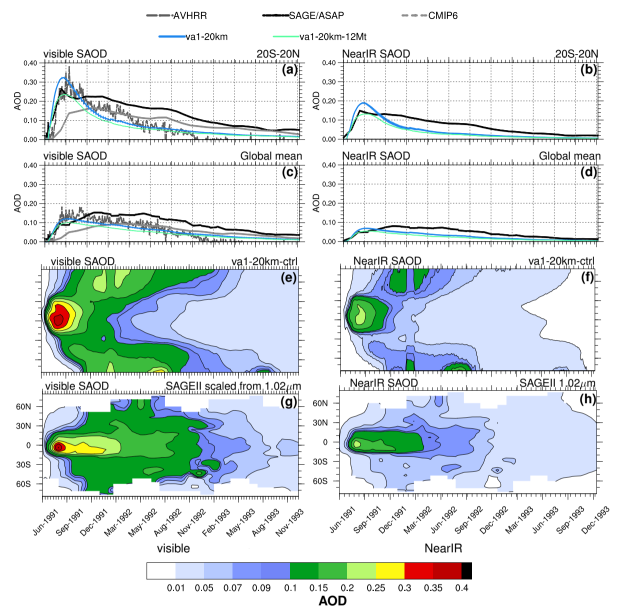
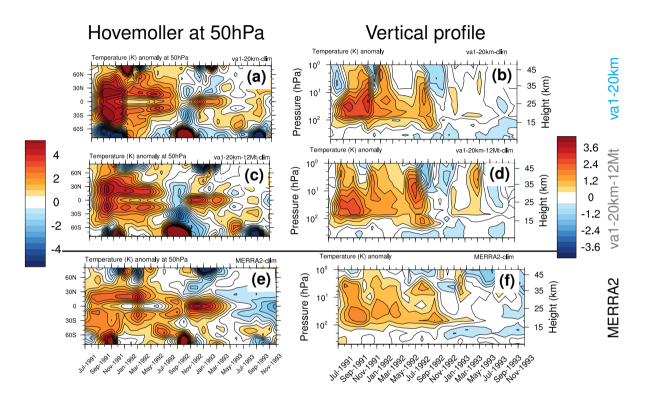


Figure 12. Averaged over the tropical belt (20S-20N) shortwave heating rate (K/day) shown as function of time and height overlaid by contours of volcanic ash mass concentration (accumulation mode) for a) va0-20km and b) va1-20km experiments, and longwave heating rate (K/day) for c) va0-20km, and d) va1-20km experiments both overlaid by contours of volcanic ash mass concentration (coarse mode). All heating rates are calculated by double call of the radiation routines.



**Figure 13.** Visible and NearIR SAODs in the va1-20km and va1-20km-12Mt experiments, as well as in AVHRR, scaled SAGE/ASAP, and CMIP6. a) Visible 20S-20N SAODs as function of time, b) NearIR 20S-20N SAODs as function of time, c) Visible globally averaged SAODs as function of time, d) NearIR globally averaged SAODs as function of time, e) Simulated visible zonally average SAOD in va1-20km as a function of time and latitude, f) Simulated NearIR zonally average SAOD in va1-20km as a function of time and latitude, g) SAGE/ASAP scaled visible zonally average SAOD as a function of time and latitude, h) SAGE/ASAP NearIR zonally average SAOD as a function of time and latitude, b) SAGE/ASAP NearIR zonally average SAOD as a function of time and latitude, h) SAGE/ASAP NearIR zonally average SAOD as a function of time and latitude. Error bars represent one sigma from the ensemble mean.



**Figure 14.** Atmospheric temperature anomalies (K) for the post-Pinatubo period with respect to the 1990-2000 climatology from the val-20km (a,b), val-20km-12Mt (c,d), and MERRA-2 <u>reranalysis reanalysis</u> (e,f). The left column depicts zonally average anomalies at the 50 hPa pressure level as a function of time and latitude, and the right column depicts globally (70S-70N) averaged anomalies as a function of time and height/pressure.

Submode	Description	
AEROPT	calculation of aerosol optical properties.	
AIRSEA	air-sea exchange of trace gases	
CLOUD	ECHAM5 cloud scheme as MESSy submodel	
CONVECT	convection parameterisations	
CVTRANS	convective tracer transport	
DDEP	dry deposition of trace gases and aerosols	
GMXE	aerosol microphysics and gas aerosol partitioning	
JVAL	photolysis rates	Landgraf and Crutzen (1998); Sander et al. (2014) Landgraf and Crutzen (1
LNOX	production of $NO_x$ from lightining	
MECCA	atmospheric chemistry computations	
OFFEMIS	prescribed emissions of trace gases and aerosols	
ONEMIS	on-line calculated emissions of trace gases and aerosols	
RAD	ECHAM5 radiative transfer as EMAC submodel	Roeckner et
SCAV	scavenging and wet deposition of trace gases and aerosol	
SEDI	sedimentation of aerosol particles	
TNUDGE	Newtonian relaxation of species as pseudo-emissions	
TROPOP	calculation of tropopause height	

Table 1. List or EMAC submodels used in this study. A complete list of all EMAC submodels can be found in Joeckel et al. (2010)

No.	Shortwave $(\mu m)$	longwave(µm)
1	0.25-0.69 (Visible)	3.3,3.8
2	0.69-1.19 (NearIR)	3.8-4.2
3	1.19-2.38	4.2-4.4
4	2.38-4.00	4.4-4.8
5		4.8-5.6
6		5.6-6.8
7		6.8-7.2
8		7.2-8.5
9		8.5-9.3
10		9.3-10.2
11		10.2-12.2
12		12.2-14.3
13		14.3-15.9
14		15.9-20.0
15		20.0-40.0
16		40.0-1000

Table 2. Shortwave and longwave bands used in the raditaion transfere calculations

**Table 3.** Description of experiments. The experiments are labeled according to the initial injection size and constituents of the injected plume. All experiment with "1x1" format represents injection in one grid box, 3s10 represents zonal injection with 10 grid points in latitude direction, the letter "s" denotes that only  $SO_2$ -SO<sub>2</sub> in injected (dry injection), letter "w" denotes that  $SO_2$ -SO<sub>2</sub> and water vapor and injected (wet injection), and va0 injection of volcanic ash with no aging and va1 is aging case. for the 1w1,va0, va1 experiments is injected with 15Mt 15 Mt and 150Mt-150 Mt of water vapor each has 5 ensemble members.

Case name	SO <sub>2</sub> mass (Mt)	Water vapor mass(Mt)	Ash mass (Mt)	Injection height (km)	Number of ensembles	Emission volume*
ctrl	-	-	-	-	5	-
1s1-17km	17	-	-	17	5	1 grid box*
1s1-20km	17	-	-	20	5	1 grid box*
1s1-25km	17	-	-	25	5	1 grid box*
1w1-20km	17	150 or 15	-	20	5x2	1 grid box*
1w1-25km	17	150 or 15	-	25	5x2	1 grid box*
va0-20km	17	150 or 15	75	20	5x2	1 grid box*
va0-25km	17	150 or 15	75	25	5x2	1 grid box*
va1-20km	17	150 or 15	75	20	5x2	1 grid box*
va1-25km	17	150 or 15	75	25	5x2	1 grid box*
3s10-25km	17	-	-	25	5	Zonal**
va1-20km-12Mt	12	150	75	17	1	1 grid box*

 $^{*}$  1 grid box - 280x280  $\rm km^{2}$  with thickness of 1 km at 17 km and 20 km altitude and 0.5 km at 25 km altitude

\*\* 10 grid box in latitude and 10 grid boxes in height ( 5km thickness)

**Table 4.** List of the studies that simulated interactive chemistry for Pinatubo case and the injected  $\frac{SO_2}{SO_2}SO_2$  height, maximum AOD and the time (in months) for the maximum AOD.

Ref.	Altitude range (km)	Initial thickness (km)	Max. AOD	Time of Max. AOD (months)	SO <sub>2</sub> Mass (Mt)
Aquila et al. (2012)	16-18	2	<del>2</del> -0.2	10	20
English et al. (2013)	15.1-28.5	13.4	0.24	7	20
Banda et al. (2013)	15-30	15	0.15	6	18.5
Dhomse et al. (2014)	19 - 27	8	0.35	2	10
Bândă et al. (2015)	17-21	4	NA	2	18.5
Sheng et al. (2015)	17-30	7-12	NA	3	14-20
Mills et al. (2016)	18-20	2	0.15	2	12

## The effect of ash, water vapor, and heterogeneous chemistry on the evolution of a Pinatubo-size volcanic cloud

Mohamed Abdelkader<sup>1</sup>, Georgiy Stenchikov<sup>1</sup>, Andrea Pozzer<sup>2</sup>, Holger Tost<sup>3</sup>, and Jos Lelieveld<sup>2</sup>

<sup>1</sup>Division of Physical Sciences and Engineering, King Abdullah University of Science and Technology, Thuwal 23955-6900, Saudi Arabia

<sup>2</sup>Air Chemistry Department, Max Planck Institute for Chemistry, Mainz, 55128, Germany

<sup>3</sup>Institute for Atmospheric Physics, Johannes Gutenberg University of Mainz, Mainz, 55128, Germany

**Correspondence:** Mohamed Abdelkader (mohamed.ahmed@kaust.edu.sa)

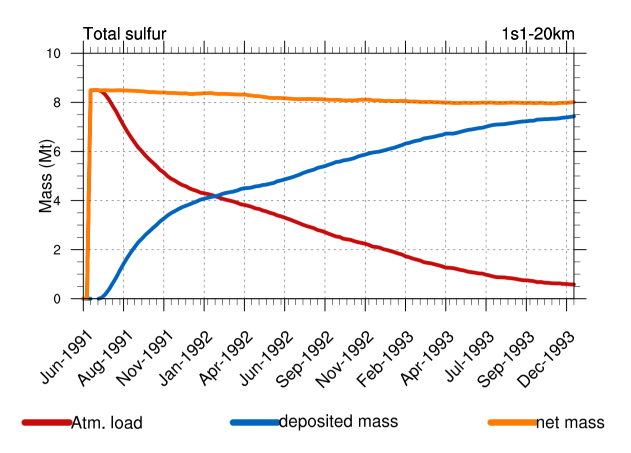
## 1 Implementation of injection

We use the EMAC import and offemis submodels to initialize the  $SO_2$ , water vapor, and volcanic ash. The emission rate is provided to the model via a prepared file that contains a four-dimensional field in space and time based on the model grid. The grid cells that correspond to the injection location are filled with the required emission rate, and the other cells are kept

- 5 with zero values. For instance, for the 1s1-20km experiment, the grid cell that corresponds to the location of Pinatubo at 20km altitude is filled with the daily emission rate for  $SO_2$ . We use the same file to initialize water vapor and volcanic ash. We use different emission rates for water vapor and volcanic ash. Then, the file is read by import submodel, and the tracers are associated with offemiss submodel.
- Figure S1 shows the sulfur mass content in the atmosphere, accumulated deposited sulfur mass, and their sum we below ref ere as a net sulfur mass. The net sulfur mass was initially 8.5Mt for 17 Mt injection of SO<sub>2</sub>. However, by the end of 1993, the net sulfur mass decreases to 8Mt, which means the model lost only 5.8% of mass to numerical errors in two and half years of simulations. This model quality is essential to accurately calculate the aerosol, and especially ash, mass balance.
- Figure S2b,c shows that the SAOD response to the water vapor mass and aging of volcanic ash. The increase of SAOD due to injection of water vapor is small compared to the increase due to injection of volcanic ash. In the val experiment, the SAOD increases (compared to 1s1) by 0.14 independent of the amount of injected water vapor. Although in va0, the sulfate mass increases by 5.5Mt (Fig. S2c), the rise in SAOD is lower than in the case with aging volcanic ash (Fig. S2d). The effect of ash is seen for about two months, and then the SAOD relaxes to that we see in the 1s1 simulation. The sensitivity to the water vapor is low because the oxidation rate of SO<sub>2</sub> depends on the water remaining in volcanic cloud, which is limited by the temperature at the injection height. It happens that in the experiments with an injection of 15Mt and 150 Mt of water vapor at 20km, about the same amount of extra water retains in the stratosphere (Fig. S2e-j). In case of 150Mt injection at 25km, about 70Mt of water vapor is retained above 50hPa level(Fig. S2e,g,i). In the va1-20km experiment only 20Mt of extra water vapor is retained above 50hPa. The difference in water vapor mass in these experiments (20Mt compared to 70Mt) resulted from the different immersion freezing at the injection level defined by the ambient temperature. The remaining water vapor mass in all cases affects the formation rate of SO<sup>2</sup>/<sub>4</sub><sup>-</sup> and the associated aerosol (Fig. S2).

Figure S3a shows the hovmoller diagram of  $SO_4^{2-}$  accumulated removal mass by sedimentation, scavenging, and and turbulent deposition in the val-20km experiment. Sulfate aerosols are removed mostly in Intertropical Convergence Zone (ITCZ)

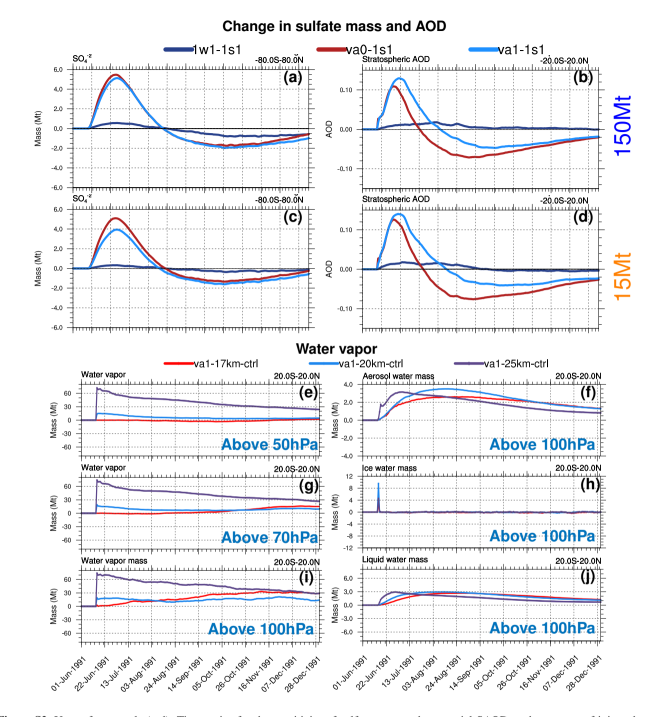
30 and in the storm tracks, in agreement with **?**. Fig. S3b shows the global accumulated sulfur deposition mass in different experiments. The sulfate removal in the 1s1 experiment is significantly lower than in other cases. Injection of water vapor increases the total removal of sulfate in the 1w1 experiment mainly because of larger sulfate mass (see Fig. 11 in the main text). In the va0 experiment, deposition increases due to the enhanced production of sulfate in the presence of ash particles. The aging of volcanic ash increases sulfate removal because aged particles are bigger and have larger deposition velocity. Also, in this 35 experiment, sulfate mass grows faster because the oxidation rate is enhanced by the heterogeneous reactions, which leads to the increase of the deposition mass.



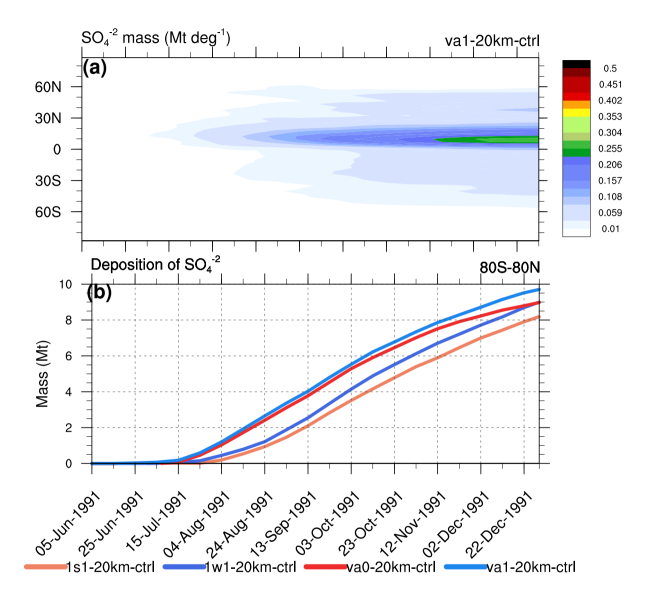
**Figure S1.** Time series for the total sulfur mass closure for 1s1-20km experiment *Atm. load* is the atmospheric sulfur mass, *Deposited mass* is the total deposition of sulfur, and *net mass* is the net atmospheric mass (Atmo. load+Deposited mass).

wave length (µm).	real	imaginary
0.31	1.53	0.02
0.42	1.53	0.01
0.52	1.53	0.01
0.63	1.53	0.00
0.75	1.53	0.00
0.88	1.53	0.00
1.00	1.53	0.00
1.13	1.53	0.00
1.34	1.53	0.01
1.64	1.53	0.01
1.93	1.53	0.01
2.23	1.53	0.01
2.58	1.48	0.02
2.99	1.47	0.02

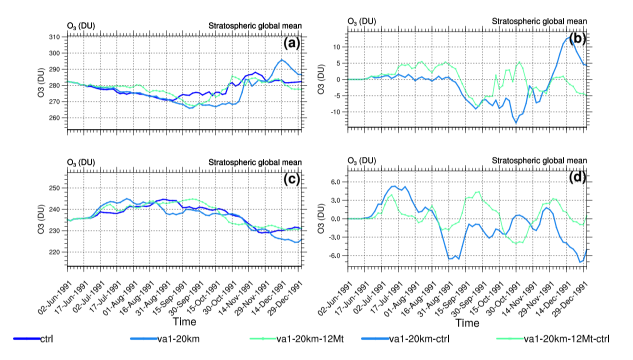
Table S1. Real and imaginary radiative indices for the volcanic ash as a function of wavelength



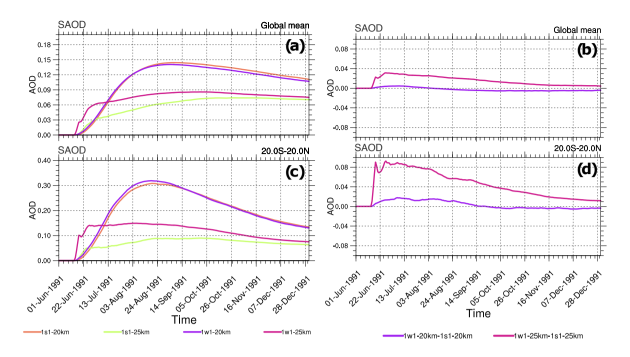
**Figure S2.** Upper four panels (a-d); Time series for the sensitivity of sulfate mass and equatorial SAOD to the amount of injected water vapor at 20km height in the presence of aged and non-aged volcanic ash: a) global sulfate mass for 150 Mt injected water vapor, b) SAOD for 150Mt injected water vapor, c) same as a using 15Mt water vapor, and d) same as b using 15 Mt water vapor. Lower six panels (e-j) time series for the change in the stratospheric water mass due to different injection heights for val experiment: e,g,i) stratospheric water vapor mass above 50 hPa, 70 hPa and 100 hPa respectively, f) aerosol water, g) ice, h) cloud liquid water above 100 hPa.



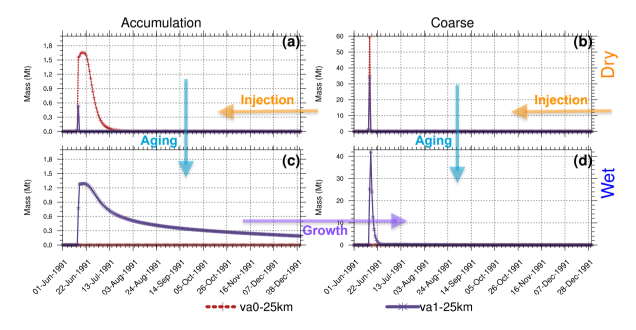
**Figure S3.** Total accumulated sulfate removal (scavenging +sedimentation) for va1-20km experiment; a) hovemoller diagram (total removal per unit degree of latitude), b) time series for the total integrated removal of  $SO_4^{2-}$  for different experiments.



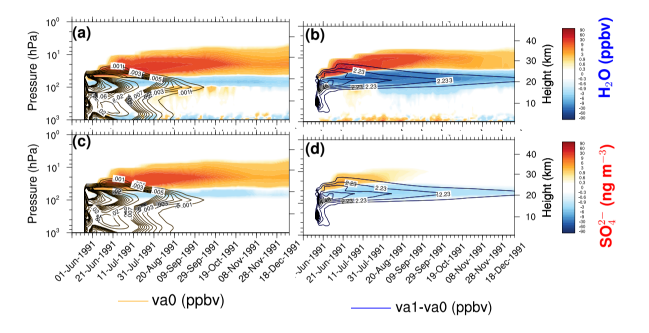
**Figure S4.** Time series for <u>stratospheric</u> ozone in Dobson unit: a) the global mean, b) the change in the global mass w.r.t control simulation, c) equatorial mean (20S-20N), d) the change in equatorial mean w.r.t control simulation for the va1-20km and va1-20km-12Mt simulations.



**Figure S5.** Time series for SAOD: a) the global mean, b) global mean of the change due to water vapor injection, c) equatorial mean (20S-20N), d) similar to (b) for equatorial mean. The left column is the SAOD for the difference w.r.t control simulation, right column is w.r.t 1w1 simulations at different injection heights.



**Figure S6.** Time series for the stratospheric (above 70 hPa) volcanic ash mass for 25 km injection heights: left column) accumulation mode, right column) coarse mode for the non-aged (dry) and aged (wet) particles for va0 and va1 experiments.



**Figure S7.** 20S-20N average perturbations of chemical constituents as function of pressure (from 300 hPa to 1 hPa) and time in va0-20km and va1-20km experiments. a)  $H_2O$  in va0-1w1 (ppbv), b)  $H_2O$  in va1-va0 (ppbv), c)  $SO_4^{2-}$  in va0-1w1 (ngm<sup>-3</sup>), d)  $SO_4^{2-}$  in va1-va0 (ng m<sup>-3</sup>). The contour lines shows the accumulation mode ash mixing ratio (ppbv); orange contour lines for va0 and blue contour lines for va1-va0 in the (a-d) panels.

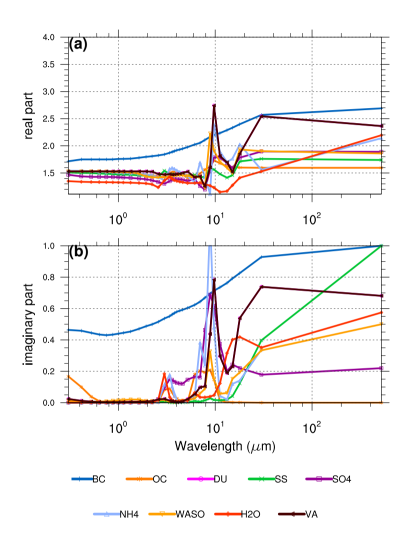


Figure S8. Radiative indices for different aerosols used in EMAC: a)real part, b) imaginary part.

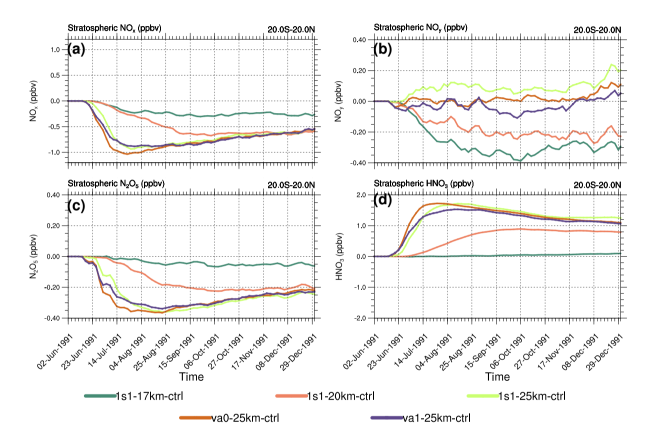


Figure S9. Time series for the stratospheric equatorial mixing ratio of: a)  $NO_x$ , b)  $NO_y$ , c)  $N_2O5$ , d)  $HNO_3$ .

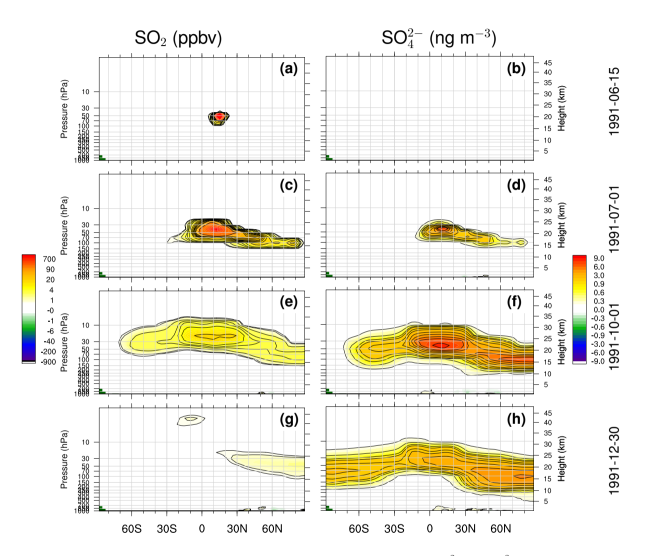
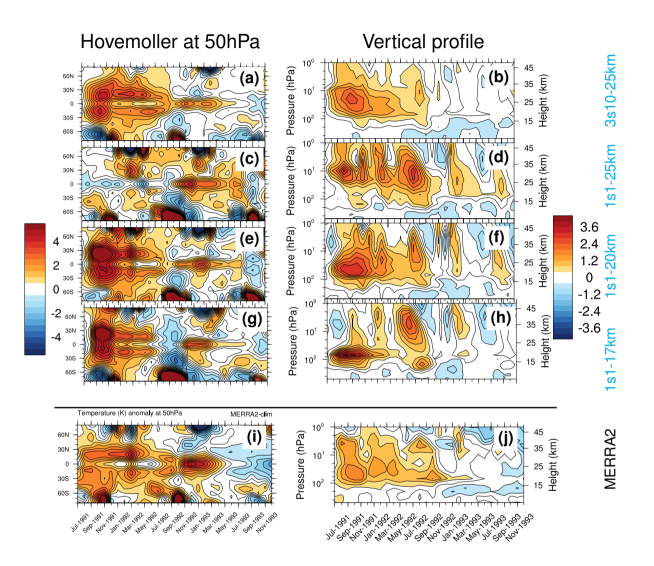


Figure S10. Zonal cross section for 1s1-20km experiment: a.c.e.g) SO<sub>2</sub> (ppbv) and b.d.f.h)  $SO_4^{2-}$  (ng m<sup>-3</sup>) at selected times from top to bottom 15<sup>th</sup> June 1991, 1<sup>st</sup> July 1991, 1<sup>st</sup> October 1991, 30<sup>th</sup> December 1991.



**Figure S11.** Atmospheric temperature anomalies (K) for the post-Pinatubo period with respect to the 1990-2000 climatology from the 3s10-25km (a,b), 1s1-25km (c,d), 1s1-20km (e,f), 1s1-17km (g,h), and MERRA-2 reanalysis (i,j). The left column depicts zonally average anomalies at the 50 hPa pressure level as a function of time and latitude, and the right column depicts globally (70S-70N) averaged anomalies as a function of time and height/pressure.