Modeled and Observed Volcanic Aerosol Control on Stratospheric NOy and Cly

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Abstract Decreases in stratospheric NOy associated with enhanced aerosol have been observed after large volcanic eruptions, for example, after the eruption of Mount Pinatubo in 1991. While the 1991 Mount Pinatubo eruption was the last large explosive eruption, recent studies have shed light on the impacts of moderate-sized eruptions since the year 2000 on the global stratospheric aerosol budget. We use an ensemble of simulations from a coupled climate-chemistry model to quantify and analyze changes in NO and NO2 (NOy), N2O5, HNO3, ClO, and ClONO2 during periods of increased stratospheric volcanic aerosol concentrations since 2000. By using an ensemble approach, we are able to distinguish forced responses from internal variability. We also compare the model ensemble results to satellite measurements of these changes in atmospheric composition, including measurements from the Optical Spectrograph and Infrared Imaging Spectrometer on the Odin satellite and the Aura Microwave Limb Sounder. We find decreases in stratospheric NOy concentrations up to 20 hPa, consistent with increases in stratospheric HNO3 concentrations. The HNO3 perturbations also extend higher, up to 5 hPa, associated with periods of increased volcanic aerosol concentrations in both model simulations and observations, though correlations with volcanic aerosol are considerably higher in the model simulations. The model simulates increases in ClO at altitudes and magnitudes similar to the NOy reductions, but this response is below the detectable limit in the available observations (100 pptv). We also demonstrate the value of accounting for transport-related anomalies of atmospheric trace gases by regression onto N2O anomalies.

1 Introduction

Heterogeneous chemistry on the surfaces of sulfate aerosol injected by volcanic eruptions can affect NOy (e.g., NO2, N2O5, and HNO3) and Cly (e.g., Cl, ClO, ClONO2, and HCl) partitioning in the stratosphere (e.g., Coffey, 1996; Hanson et al., 1994). After large sulfur injections from the El Chichón (1982) and Pinatubo (1991) eruptions, extensive research was carried out to identify and quantify these responses. Johnston et al. (1992) and Mills et al. (1993) found large NO2 reductions over New Zealand and Colorado, respectively, after Pinatubo; Rinsland et al. (1994) found decreases in N2O5 and increases in HNO3 on the basis of on-ground and stratospheric aircraft measurements, Coffey (1996) found decreases in NO and NO2 (hereafter NOy) and corresponding increases in HNO3 column abundances after both eruptions; Mickley et al. (1997) found reductions in NOy as high as 17 hPa after Pinatubo, and Avallone et al. (1993) found increases in ClO near 20–30°N after Pinatubo.

The heterogeneous reactions that are thought to control the NOy/Cl response to sulfate aerosol are as follows (Hanson et al., 1994):

\[
\text{ClONO}_2 + \text{HCl} \rightarrow \text{HNO}_3 + \text{Cl}_2 \quad \text{(R1)}
\]
\[
\text{HOCl} + \text{HCl} \rightarrow \text{H}_2\text{O} + \text{Cl}_2 \quad \text{(R2)}
\]
\[
\text{N}_2\text{O}_5 + \text{H}_2\text{O} \rightarrow 2\text{HNO}_3 \quad \text{(R3)}
\]
As such, the expected responses to an increase in volcanic aerosol concentrations include a decrease in NO, NO\textsubscript{2}, and N\textsubscript{2}O\textsubscript{4} mainly by reaction (R3) (N\textsubscript{2}O\textsubscript{4} is a reservoir species for NO and NO\textsubscript{2}, and therefore, hydrolysis of N\textsubscript{2}O\textsubscript{4} depletes the reservoir) and a corresponding increase in HNO\textsubscript{3} by (R3), as well as (R1), (R4), and (R5). Similarly, anomalies of opposite signs are possible in ClONO\textsubscript{2} (negative; (R1) and (R4)) and ClO (positive; (R1) and (R2)), though (R4) is likely most significant inside the polar vortex (Wegner et al., 2012).

There has been no eruption nearly as large in magnitude as Mount Pinatubo (18 Tg SO\textsubscript{2}) since it erupted in 1991; however, on average about 0.8 Tg SO\textsubscript{2}/year was emitted into the stratosphere from volcanic eruptions over 2005–2013, with injections as high as 21 km (Table S1 in the supporting information). This constitutes a substantial increase from the 0.06 Tg SO\textsubscript{2}/year emitted, on average, over 1999–2004, and there has been an increasing focus recently on understanding the role of these smaller but still significant injections of volcanic sulfur into the stratosphere on, for example, the global energy budget and climate change (Schmidt et al., 2018; Solomon et al., 2011) and atmospheric chemistry (Adams et al., 2017; Berthet et al., 2017). For the latter, Berthet et al. (2017) found reductions of NO\textsubscript{2} and increased HNO\textsubscript{3} below 19 km after the 2009 Sarychev Peak eruption, while Adams et al. (2017) showed NO\textsubscript{2} reductions up to 25 km after several eruptions from 2005 to 2014, as measured by the Optical Spectrograph and Infrared Imaging System (OSIRIS).

Though the research on the topic is extensive, previous studies largely looked at the lowermost stratosphere, and only a few have examined the impacts of volcanic eruptions since 2000. The goal of this study is to probe not only the lower but also the upper layers of the stratosphere to investigate whether there are any significant observable responses in NO\textsubscript{y} or Cl\textsubscript{y} to the multiple moderate-magnitude volcanic eruptions in the post-Pinatubo era. Using satellite data from OSIRIS and the Microwave Limb Sounder (MLS), and an ensemble of simulations from the Community Earth System Model version 1 with the Whole Atmosphere Community Climate Model component (CESM1|WACCM4; Hurrell et al., 2013; Mills et al., 2016), we examine the effect of recent volcanic eruptions on stratospheric composition up to 5 hPa.

The model ensemble approach is a powerful approach for separating forced responses from natural or internal variability (e.g., Deser et al., 2014; Kay et al., 2015; Otto-Bliessner et al., 2016). This approach is especially important for elucidating the responses to forcings, such as volcanic eruptions, which can alter not only the mean state but also the internal variability (e.g., Bittner et al., 2016). While the ensemble approach has been shown to be an effective tool in dynamical studies, it has seen less application for chemistry. Here we show the importance of making use of statistical information from ensemble simulations to ensure appropriate identification of chemical phenomena, especially when comparing data and models.

We also demonstrate the value of accounting for transport-related anomalies of atmospheric trace gases by regressing onto N\textsubscript{2}O anomalies. This method has seen limited application in prior studies (e.g., Rinsland et al., 2009; Stolarski et al., 2018), but this work suggests that it can provide an improved means of accounting for quasi-biennial oscillation (QBO) perturbations to chemically active species.

2. Data and Model Description
2.1. CESM1(WACCM)

CESM1 is a fully coupled climate model with atmosphere, ocean, land, and sea ice components (Hurrell et al., 2013). WACCM version 4 is the high-top atmospheric component of CESM1 that has a horizontal resolution of 1.9° latitude by 2.5° longitude; the free running version of WACCM used here has 66 vertical levels with a model top near 140 km in altitude (Garcia et al., 2017; Marsh et al., 2013; Solomon et al., 2015). The WACCM chemical scheme used in the study includes a representation of tropospheric, stratospheric, mesospheric, and lower thermospheric chemistry (Kinnison et al., 2007; Tilmes et al., 2016). The species included within this mechanism are contained within the O\textsubscript{x}, NO\textsubscript{x}, HO\textsubscript{x}, ClO\textsubscript{x}, and BrO\textsubscript{x} chemical families, along with CH\textsubscript{4} and its degradation products. In addition, 20 primary nonmethane hydrocarbons and related oxygenated organic compounds are represented along with their surface emissions. There is a total of 183 different species, 341 gas phase reactions, 114 photolytic processes, and 17 heterogeneous reactions on aerosols (i.e., sulfate, nitric acid trihydrate, and water ice).
Using this model, a fully coupled ensemble of simulations is generated. The ensemble uses a free running atmosphere-ocean chemistry climate general circulation model whose 10 members are examined over the 1999–2014 period, allowing a better distinction between the forced response and internal variability, thereby allowing us to identify to what extent any volcanic signals should be expected in the observations. These simulations have a repeated cyclic 28-month QBO based on rocketsonde data (Matthes et al., 2010) and no solar cycle. Sulfate aerosol surface area densities (SAD) include volcanic and nonvolcanic sources and are specified based on calculations from Mills et al. (2016), which used volcanic SO$_2$ injections from Neely and Schmidt (2016) for the period spanning 1999–2014. The calculated aerosol properties were shown to compare well to lidar-based stratospheric aerosol extinction data (Mills et al., 2016) and the Coupled Model Intercomparison Project phase 6 stratospheric aerosol optical depth data set (Schmidt et al., 2018; Thomason et al., 2018). Ensemble members are branched from a single Chemistry-Climate Model Initiative (Eyring et al., 2013) REF-C2 simulation in 1995, and distinct ensemble members are generated by randomly perturbing the initial temperature fields by roundoff magnitude (order $10^{-14}$ K). The SAD calculations are based on simulations nudged using meteorological fields from the NASA Global Modeling and Assimilation Office Modern-Era Retrospective Analysis for Research and Applications (Rienecker et al., 2011), including stratospheric winds, so the transport of aerosols is not affected by the cyclic QBO used in these simulations. Greenhouse gas concentrations in the simulations evolve according to the Representative Concentration Pathway 6.0 (Fujino et al., 2006; Hijioka et al., 2008), and concentrations of ozone-depleting substances are prescribed according to the Chemistry-Climate Model Initiative (Eyring et al., 2013).

### 2.2. MLS

The Earth Observing System MLS on board the Aura satellite launched 15 July 2004 observes thermal emission through the Earth’s atmospheric limb, which enables daily near-global (82°S to 82°N) measurements of temperature, pressure, and atmospheric gases from the troposphere to the mesosphere (Waters et al., 2006). The data used in this study are MLS v4.2 “Level 2” products generated from the calibrated microwave radiance observations; improvements in the v4.2 data include better handling of cloudy regions (Livesey et al., 2017).

The standard MLS ClO product is taken from the 640-GHz retrievals. The vertical resolution of the v4.2 ClO is 3–4.5 km. Precision on individual profiles is ±0.1 ppbv over most of the stratosphere, increasing to ±0.3 ppbv at 147 and 1 hPa. The systematic uncertainty in the measurements ranges from ±0.02 to ±0.4 ppbv.

There exists a latitude-dependent bias in the ClO profiles in the 68 to 147 hPa range, which is corrected by subtracting the bias estimates from individual measurements at these levels. V2.2 ClO measurements are validated in Santee et al. (2008); for a similar comparison of different observing platforms with v3 ClO products, to which the v4.2 ClO product is most similar, see Livesey et al. (2013).

For pressures greater than or equal to 22 hPa, the MLS HNO$_3$ product is derived from the 240-GHz radiances; for pressures less than 22 hPa, retrievals from the 190-GHz radiometer are used. The vertical resolution of the v4.2 HNO$_3$ is 3–5 km. Precision on individual profiles is ±0.6 ppbv over most of the stratosphere, increasing to ±0.8 and ±1.2 ppbv at 2.1 and 1.5 hPa, respectively, and systematic uncertainties are between ±0.1 and ±2.4 ppbv at the various levels (Livesey et al., 2017). The detailed validation of the v2.2 HNO$_3$ products is presented by Santee et al. (2007), with analysis of the more comparable v3 HNO$_3$ measurements in Livesey et al. (2013).

Previous versions of MLS N$_2$O were derived from the 640-GHz retrievals. However, due to deterioration of the derived N$_2$O product, the standard product for MLS v4.2 N$_2$O is taken from the 190-GHz radiances. The vertical resolution of the 190-GHz N$_2$O product is 4–8 km, and the estimated precision varies from ±15 to ±19 ppbv; these values are slightly worse than those of the 640-GHz retrievals, except at 68 hPa (Livesey et al., 2017). Compared to v2.2 N$_2$O, which is validated in Lambert et al. (2007), average v4.2 N$_2$O values are up to 10% smaller at 100 and 68 hPa and within a few percent at other levels.

After screening individual profiles as described in Livesey et al. (2017), MLS ClO, HNO$_3$, and N$_2$O retrievals are averaged to create daily means on 7.5° latitude × 15° longitude grids; daily means are then averaged zonally and in time to construct monthly zonal averages on each pressure level. Because the focus of this
analysis is partially on the effectiveness of the N$_2$O regression fit, analysis of MLS products is performed on the intersection of the N$_2$O and ClO/HNO$_3$ vertical grids: 68–1 hPa for ClO and 68–1.5 hPa for HNO$_3$.

2.3. OSIRIS

OSIRIS is a Canadian instrument on board the Swedish Odin satellite (Llewellyn et al., 2004; McLinden et al., 2012). Launched on 20 February 2001, the orbit of OSIRIS is Sun-synchronous, at an altitude of approximately 600 km. Similar to MLS retrievals, limb-scattering measurements from OSIRIS have near-global, 82°S to 82°N, spatial coverage. The OSIRIS data used in this study are based on the v6 NO$_2$, which are described and validated in Sioris et al. (2017). The v6 NO$_2$ data are based on an improvement of previously developed retrieval algorithms, and measurement uncertainties are estimated to be about 15–20% above ~20 km (Bourassa et al., 2011; Sioris et al., 2003, 2004, 2007).

Based on OSIRIS measurements during descending portions of the orbit, with an equatorial crossing time near 06:30 local solar time (LST), NO$_2$ number density profiles are derived. NO$_2$ (and NO) is in rapid photochemical equilibrium and exhibits a strong diurnal cycle (Brasseur & Solomon, 2005; Park et al., 2017, their Figure 1). In order to facilitate the comparison of the OSIRIS NO$_2$ observations with model output, Park et al. (2017) derived daily-mean NO$_2$ (NO + NO$_2$) concentrations using a photochemical box model (Adams et al., 2017; McLinden et al., 2000; Prather, 1992). After adjusting the OSIRIS NO$_2$ profiles to represent NO$_2$ concentrations at 06:30 a.m. LST (some of the profiles are measured before or after 6:30 LST), the box model was used to derive a scaling factor based on the ratio of daily-mean NO$_2$ concentrations to NO$_2$ concentrations at 6:30 LST; the scaling is then applied to estimate the daily average NO$_2$ concentrations.

As in Park et al. (2017), we convert NO$_2$ number densities to mixing ratios using temperature (T) and pressure (p) from the European Centre for Medium-Range Weather Forecasting (ECMWF) interim reanalysis (ERA-Interim; Dee et al., 2011). The box model includes input of OSIRIS O$_3$ and aerosol extinction, as well as the retrieved surface reflectance and T and p from ERA-Interim.

Gridded OSIRIS NO$_2$ profiles are computed similarly to the MLS profiles: daily NO$_2$ profiles are used to construct daily gridded data on the same horizontal grid as the MLS data, and the daily gridded data are averaged to create monthly zonal means. On the other hand, the OSIRIS NO$_2$ product is available on a vertical grid of 1 km from 10.5 to 39.5 km, which we interpolate vertically to the MLS N$_2$O pressure grid using log-pressure coordinates.

3. Methods

We calculate the climatological monthly mean and standard deviation of the WACCM SAD, $\overline{SAD}$, and $\sigma_{SAD}$, respectively, at each latitude and pressure level for the period 1999–2013. We define the “volcanically active” periods at each latitude $\phi$ and pressure level $p$ as the set of all times $t$ such that

$$SAD(t, \phi, p) > \overline{SAD}(\phi, p) + \sigma_{SAD}(\phi, p)$$  (1)

In other words, we define a volcanically active period as one where the SAD is greater than one standard deviation above the mean for that month, latitude, and pressure level or approximately greater than the 84th percentile. The background climatologies, from which all anomalies are calculated, are then calculated as the means of the “volcanically clean” periods, that is, the times for which equation (1) is not satisfied. The background values are subtracted at all points, and the linear trend is removed. Finally, we divide by the background values, so that unless otherwise noted, anomalies are given as deseasonalized and detrended percent changes.

Figure 1 shows the deseasonalized and detrended WACCM NO$_2$ anomalies for a single ensemble member (5) at several levels in the stratosphere. Black contour lines in the figure highlight the periods of enhanced stratospheric aerosol concentrations as defined by equation (1), based upon the volcanic inputs and modal aerosol model presented in Mills et al. (2016). The figure suggests suppression of NO$_2$ during periods of high aerosol concentrations at the lowest levels (72 and 51 hPa; Figures 1d and 1e) but with considerable noise. Overall, the most prominent feature is the 28-month periodic QBO signal, which can be seen at all levels in the tropics and midlatitudes in Figure 1. The effect of the QBO on stratospheric composition has been
Figure 1. WACCM NO\textsubscript{x} (NO + NO\textsubscript{2}) anomalies (\%; a–e) and QBO1 time series (f) for ensemble member 5. Black line contours indicate where sulfate aerosol surface area density is greater than 1\(\sigma\) above the mean, and triangles indicate the timing and latitudes of the volcanic eruptions listed in Table S1. The number in the top right corner of panel (f) is the percent variance of the zonal wind explained by QBO1. WACCM = Whole Atmosphere Community Climate Model; QBO = quasi-biennial oscillation.
widely studied (e.g., Park et al., 2017; Randel & Thompson, 2011; Randel & Wu, 1996). We first present a fit to QBO-related anomalies obtained using a multiple linear regression, regressing the first two principal components (PCs) of the QBO (QBO1 and QBO2, respectively) onto the WACCM NO\textsubscript{x} anomalies (Adams et al., 2017; Bourassa et al., 2014; Park et al., 2017; Randel & Wu, 1996); the results of the fit are plotted in Figure 2, and the relative amplitudes of the QBO regression fits are given in Figures 3a and 3b. After regressing out the first two PCs of QBO, altitudes above 50 hPa still show a clear quasi-biennial signal on WACCM NO\textsubscript{x} anomalies (period ~22 months; Figures 2a–2c). We chose to show WACCM NO\textsubscript{x} anomalies

Figure 2. (a–f) WACCM NO\textsubscript{x} anomalies (%) for ensemble member 5 with QBO1 + QBO2 fits removed. Black line contours indicate where sulfate aerosol surface area density is greater than 1σ above the mean, and triangles indicate the timing and latitudes of the volcanic eruptions listed in Table S1. WACCM = Whole Atmosphere Community Climate Model; QBO = quasi-biennial oscillation.
as an illustrative example here, but this pattern is consistent across the different gases analyzed (and between model and observations). We also investigated the impacts of the semiannual oscillation (SAO) of zonal winds in the tropical upper stratosphere but found little correlation between the SAO and NO\textsubscript{y} concentrations, so SAO regression is excluded from results shown here. Instead, the residual quasi-biennial signal at upper levels appears to be a tape recorder-like effect (Mote et al., 1996), where QBO-induced anomalies at lower levels are advected upward, with variable time lags relative to lower levels.

Park et al. (2017) demonstrated the strong anticorrelation between NO\textsubscript{y} and HNO\textsubscript{3} and source gas N\textsubscript{2}O in the tropical stratosphere, where N\textsubscript{2}O is oxidized to form NO\textsubscript{y}. We next explore this relationship and its value for accounting for dynamical variability including the QBO. In Figure 4, scatter plots of deseasonalized and detrended N\textsubscript{2}O versus NO\textsubscript{y} (NO + NO\textsubscript{2} + NO\textsubscript{3} + 2N\textsubscript{2}O\textsubscript{3} + HNO\textsubscript{3} + ClONO\textsubscript{2}) anomalies at various levels and latitude bands demonstrate this relationship, which extends outside the tropical latitudes studied by Park et al. (2017). At all pressure levels above and including 72 hPa (i.e., at pressures less than or equal to 72 hPa), NO\textsubscript{y} shows strong anticorrelations with N\textsubscript{2}O, except at high latitudes in the upper stratosphere (poleward of 30° in both hemispheres at 5 hPa; Figure 4a), where NO\textsubscript{y} is positively correlated with N\textsubscript{2}O, with Pearson correlation coefficients greater than 0.5. Correlation coefficients between N\textsubscript{2}O and NO\textsubscript{y} are below −0.9 at all levels in the 10°S to 10°N band, with the strongest anticorrelation ~0.99 at 5 and 10 hPa (Figures 4a and 4b). At and below 20 hPa (Figures 4c–4e), correlation coefficients are below (i.e., stronger than) −0.9 at all latitudes in the 60°S to 60°N band considered here.

Park et al. (2017) showed the dependence of N\textsubscript{2}O on the QBO; therefore, regressing N\textsubscript{2}O anomalies onto the NO\textsubscript{y} anomalies should also capture QBO variability. Because of the compact relationship between NO\textsubscript{y} species and N\textsubscript{2}O seen in Figure 4, this method should better account for sources of dynamical variability that regressing out QBO alone might not capture. In Figure 5, we show the WACCM NO\textsubscript{x} time series obtained when detrended model N\textsubscript{2}O anomalies are regressed onto the NO\textsubscript{x} anomalies instead of using the QBO PCs; the relative amplitude of the N\textsubscript{2}O fit is shown in Figure 3c. The tight relationship between N\textsubscript{2}O and

Figure 3. Relative amplitude of regression fits (%/σ) for WACCM NO\textsubscript{x} for QBO only (a, b), N\textsubscript{2}O only (c), and QBO + N\textsubscript{2}O (d–f). Results are shown for the mean of the regression fits of the individual ensemble members. WACCM = Whole Atmosphere Community Climate Model; QBO = quasi-biennial oscillation.
NO\textsubscript{x} is further demonstrated in Figure 3c, which shows up to 20% decreases in NO\textsubscript{x} for a 1\(\sigma\) change in N\textsubscript{2}O in the tropical middle stratosphere (e.g., near 0° and 10 hPa). At levels above 50 hPa (Figures 5a–5c), the periodic oscillations are now well removed by regressing out N\textsubscript{2}O anomalies, and novel information is obtained about the relationship between aerosol anomalies and NO\textsubscript{x}. Specifically, at 10 and 20 hPa notable decreases in NO\textsubscript{x} concentrations exceeding 10% are simulated in times of increased volcanic aerosol (Figures 5b and 5c). On the other hand, upon comparison of Figures 2d and 5d, there is now a QBO signal at 51 hPa (Figure 5d) which was not evident after removing the dynamical influence associated with QBO1 and QBO2 (Figure 2d). This is likely to be related in part to weak gradients in N\textsubscript{2}O at lower altitudes (higher pressures). In light of the effectiveness of the N\textsubscript{2}O regression above 50 hPa and the apparent utility of the QBO regression at greater pressures, we next use a multiple linear regression to remove dynamical anomalies associated with both the QBO and N\textsubscript{2}O; the results are shown in Figure S1, and Figure 3 shows the relative amplitudes of the regression fits for each of the three methods ([a–b] QBO, [c] N\textsubscript{2}O, and [d–f] QBO + N\textsubscript{2}O). Figure 3 demonstrates that the N\textsubscript{2}O anomalies dominate the QBO + N\textsubscript{2}O regression; this should be expected, since N\textsubscript{2}O anomalies will capture some of the QBO variability. However, Figures 3d and 3e show that retaining the QBO fit as a separate function remains important for optimal fitting at pressures \(\geq\) 30 hPa. We note that the QBO also impacts the temperature structure of the stratosphere, which affects temperature-dependent chemistry; Figure S1 shows that using both the QBO and N\textsubscript{2}O anomalies in the regression provides a remarkably effective means for accounting for both chemical and dynamical influences on NO\textsubscript{x}, demonstrating the value of this approach for studies of chemical perturbations. Figure S2 shows the relative regression amplitudes for N\textsubscript{2}O and the QBO. Stratospheric N\textsubscript{2}O is highly correlated with the QBO above 30 hPa; on the other hand, variability in the equatorial stratospheric zonal winds has little influence on N\textsubscript{2}O concentrations at and below 30 hPa. In
addition, Figure S3 shows the WACCM ensemble‐mean NOx anomalies subtracted from those after N\textsubscript{2}O regression only, which is approximately equal to the effect of including the QBO regression. From Figure S3, it can be concluded that the QBO influence for the QBO+N\textsubscript{2}O fit is limited to exactly those levels (pressure ≥ 30 hPa) where QBO and N\textsubscript{2}O are uncorrelated. Therefore, the potential nonorthogonality of the QBO and N\textsubscript{2}O regressors is not an issue for this application.

Figure 5. (a–e) WACCM NO\textsubscript{x} anomalies (%) for ensemble member 5 with the dynamical variability identified by the N\textsubscript{2}O fit removed. Black line contours indicate where sulfate aerosol surface area density is greater than 1\(\sigma\) above the mean, and triangles indicate the timing and latitudes of the volcanic eruptions listed in Table S1. WACCM = Whole Atmosphere Community Climate Model.

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We also make use of the ensemble of coupled climate-chemistry model simulations by analyzing the signal-to-noise ratio (SNR; e.g., Deser et al., 2014). We define the SNR as the ensemble-mean anomaly—the “forced” response—divided by the standard deviation of the response across ensemble members, which quantifies the internal variability. The SNR analysis provides a quantitative comparison of the respective magnitudes of the forced response and the internal variability; this is useful for comparing the model ensemble to the observations in that it can help to identify times and locations in the observations at which one might best be able to see a forced response that is distinguishable from the natural variability.

4. Results

After removing the dynamics-related anomalies, there is a clear volcanic signal in the WACCM ensemble mean NO$_x$ time series in Figure 6, with anomalies related to increased sulfate aerosol concentrations (contours show SAD) at levels up to 10 hPa but most prevalent at and below 20 hPa (i.e., at pressures ≥20 hPa). Specifically, at 10 hPa (Figure 6b), decreased NO$_x$ concentrations are simulated poleward of 30°S in 2002–2004 during periods of increased volcanic aerosol concentrations; however, the timing of the response in 2002, which begins in July, indicates that the reduction is not related to the Ruang eruption, which occurred in September 2002. Reductions in NO$_x$ are also simulated at NH midlatitudes in 2003, 2007, and 2008 in association with higher volcanic aerosol concentrations. The times and locations of these responses are consistent with volcanic aerosol concentration increases due to the eruptions listed in Table S1. The SNR in Figure 8b indicates that the most consistent forced anomaly at 10 hPa occurs in the NH tropics in 2007, where the SNR is above 4, indicating robustness of the reduction. In addition, Figure S4, which shows the absolute NO$_x$ anomalies, indicates that the apparent correlation between aerosol concentrations and NO$_x$ prior to 2005 is less robust, with both positive and negative anomalies simulated in times of increased aerosol concentrations. It is likely that the previously discussed reductions at midlatitudes in Figure 6b are partly related to the apparent seasonal cycle of sulfate aerosol SAD at this altitude. However, removal of the seasonal variations in the gaseous species demonstrates a robust aerosol chemical effect: comparing, for example, years 2002–2005, where this seasonal cycle is apparent, to 2010–2014, when the aerosol SAD is reduced at this level, it can be seen that these reductions in NO$_x$, though occurring seasonally, also vanish when the SAD is reduced.

Panels c–e in Figures 6 and 7 show reductions in NO$_x$ that are both greater in magnitude and more directly related to identifiable volcanic injections. At 20 hPa (Figure 6c), similar decreases related to the seasonal occurrence of increased volcanic aerosol concentrations are simulated robustly in 2002 (SH) and 2003 (NH). On the other hand, NO$_x$ reductions in the second half of 2003 and early 2004 are simulated from 30°N and southward and appear to be related to the 2003 Anatahan and/or Soufrière Hills eruptions (Table S1, based on Table S6 of Mills et al., 2016; eruption latitudes 16°N and 17°N, respectively). Again, the reductions there are less than 10%, but the SNR (Figure 8c) is above 4, indicating a chemical response to the increased volcanic aerosol concentrations that is robust and consistent across the model ensemble and therefore should be detectable in the single realization available in observations. The largest NO$_x$ reductions at 20 hPa are simulated in 2005 (Manam, 4°S, Sierra Negra, 1°S; reductions >10%) and 2006 (Manam, 4°S, Soufrière Hills, 17°N, Tungurahua, 1°S; reductions >15%), with both periods simulating SNR above 4 (Figure 8c), again indicating that the signal should be observable. At 51 and 72 hPa (panels d and e of Figures 7 and 8), the volcanic aerosol effect is much more apparent, with WACCM model ensemble mean NO$_x$ decreases of more than 20% in the tropics after Manam (4°S) and Sierra Negra (1°S; 2005); after Manam (4°S), Soufrière Hills (17°N), and Rabaul (4°S; 2006), with decreases lasting through 2007; and after Merapi (15°N) and Nabro (13°N; 2010–2011). Outside the tropics, large decreases are simulated in the SH after Manam (4°S, 2006), Rabaul (4°S; 2006; NO$_x$ reduction in 2007), Chaitén (43°S; 2008), and Merapi (8°S; 2010; NO$_x$ reduction in 2011). At NH midlatitudes, the largest reductions follow the eruptions of Soufrière Hills (17°N, 2006), Okmok (53°N), Kasatochi (52°N, 2008), Redoubt (60°N), Sarychev Peak (48°N; 2009), and Nabro (13°N; 2011). Figure S4 shows the absolute changes in NO$_x$ in parts per billion by volume (ppbv), which are consistent with the percent changes in that the largest decreases are associated with increased volcanic aerosol, especially at and below 20 hPa. N$_2$O$_5$ anomalies (Figure S5) show similar decreases in response to the volcanic aerosol, with decreases exceeding 25% at ~50 hPa and below (Figures S5d–S5e).
To further emphasize the added value of using $N_2O$ as a tracer to remove anomalies related to dynamics, the WACCM NO$_x$ SNR are shown after regressing out QBO only (Figure S6) and N$_2$O only (Figure S7).

Comparison of SNR after QBO fit (Figure S6) with SNR after QBO + N$_2$O fit (Figure 7) shows that the full fitting process is much more effective in isolating the volcanic signal from dynamical variability. When using the QBO-only approach, Figure S6 shows SNR frequently above 4 at all levels, with the same spatiotemporal patterns as the residual periodic oscillations seen in Figure 2. The high SNR and therefore low variability

Figure 6. (a–e) WACCM NO$_x$ anomalies (%) for the ensemble mean with QBO + N$_2$O fits removed. Black line contours indicate where sulfate aerosol surface area density is greater than 1σ above the mean, and triangles indicate the timing and latitudes of the volcanic eruptions listed in Table S1. WACCM = Whole Atmosphere Community Climate Model; QBO = quasi-biennial oscillation.
indicates a continued dynamical control on the simulated response, even after regression of the QBO influence. In contrast, Figure 7 shows that after the more robust removal of dynamics-related anomalies, SNR is either generally very low (e.g., Figures 7a and 7b) or that the high SNR is associated with increases in volcanic aerosol concentrations (Figures 7c–7e). A similar comparison between Figure 7 and S7 shows that the N$_2$O fit appears to be sufficient at higher altitudes but does not account for all dynamical

**Figure 7.** (a–e) WACCM NO$_x$ signal-to-noise ratio (SNR), defined as the ensemble mean divided by the standard deviation across ensemble members. Black line contours indicate where sulfate aerosol surface area density is greater than 1σ above the mean, and triangles indicate the timing and latitudes of the volcanic eruptions listed in Table S1. WACCM = Whole Atmosphere Community Climate Model; QBO = quasi-biennial oscillation.
variability below 20 hPa, where high SNR remain, unrelated to volcanic aerosol concentrations (Figures S7c–S7d).

Figure 8 shows the OSIRIS NOx observed anomalies in percent change, and absolute anomalies (ppbv) are given in Figure S8. For all observations (i.e., OSIRIS NOx and MLS HNO3 and ClO), we perform the same regression fit as was done for the model output, using MLS N2O for the N2O regression and using the equatorial zonal wind observations over Singapore (Naujokat, 1986) to compute the QBO components. The WACCM ensemble mean shows good agreement with the OSIRIS NOx time series in Figure 8, with both

Figure 8. (a–e) OSIRIS derived NOx anomalies (%) with QBO + N2O fits removed. Black line contours indicate where sulfate aerosol surface area density is greater than 1σ above the mean, and triangles indicate the timing and latitudes of the volcanic eruptions listed in Table S1. OSIRIS = Optical Spectrograph and Infrared Imaging System.
showing the largest reductions at 46 and 68 hPa, though the observations match better when compared with a single ensemble member, for example, Figure S1. Comparing the observations to a single ensemble member results in generally better agreement because much of the internal variability is averaged out when computing the ensemble mean response. In comparing WACCM simulation #5 (Figure S1) with the OSIRIS data (Figure 8), the two show very good agreement at all latitudes from July 2006 to 2008, with long-lasting reductions in NOx in response to the sustained increase in volcanic aerosol during these years. In addition, the responses to Okmok (53°N) and Kasatochi (52°N; 2008), Sarychev Peak (48°N; 2009), Meraapi (8°N; 2010), and Nabro (13°N; 2011) as seen in WACCM are also captured in the OSIRIS observations. On the other hand, a few key differences can also be seen. First, at 46 and 68 hPa, there is no observable reduction due to the increased volcanic aerosol concentrations after the Manam (4°S) eruption in 2005; this is consistent with Adams et al. (2017), who also showed minimal observed changes in response to this eruption. For the eruptions after 2008, the response in OSIRIS NOx appears only at 68 hPa, as opposed to the WACCM simulations, which show reductions at 50 hPa in 2008, 2009, and 2011; this suggests potential differences between the observed and modeled transport of aerosol to the higher altitudes. The response to Okmok (53°N) and Kasatochi (52°N; 2008) is much larger and has greater latitudinal extent in the WACCM simulations than in the OSIRIS data.

WACCM ensemble mean HNOx anomalies (%) were analyzed in the same manner as those for NOx presented above and are plotted in Figure 9. In contrast to NOx, which shows the largest effect of aerosol below 20 hPa, Figure 9 shows that WACCM simulates impacts on HNOx throughout the stratosphere, with large percentage increases at 5 hPa during times of increased sulfate aerosol (Figure 9a). At 10 hPa (Figure 9b), WACCM simulates increases in HNOx following Manam (4°S; 2005 and 2006), Soufrière Hills (17°N), Rabaul (4°S; 2006; HNO3 response in 2007), Okmok (53°N), and Kasatochi (52°N; 2008; HNO3 response in 2009). At 20 and 72 hPa (Figures 9c and 9e), the aerosol-induced increases in HNO3 are similar in magnitude to the associated decreases in NOx (e.g., Figures 6c and 6e); at 51 hPa (Figure 9d), percent changes in HNO3 are lower than those for NOx (Figure 6d) and mostly confined to the tropics. WACCM HNO3 SNR analysis (Figure S9) is consistent with Figure 9, although SNR at volcanically active times is hard to distinguish from the background SNR at 5 hPa (Figure S9a). The responses in Figure 9 at 10–72 hPa appear to be robust, as SNR at these altitudes tends to be highest during times of increased volcanic aerosol concentrations (Figures S9b–S9e). The model results compare well to HNO3 observations from MLS (Figure 10), which show increases at times of increased volcanic aerosol concentrations at all levels but 46 hPa. Figures S10 and S11 show the absolute changes in HNO3 for WACCM and MLS, respectively; the time series in these two figures agree quite well at all levels except near 50 hPa, where any concurrence of increased volcanic aerosol concentrations and MLS HNO3 appears to be spurious.

WACCM ensemble mean ClO anomalies are plotted in Figure 11. Because of its presence at very low mixing ratios in the stratosphere, we present ClO anomalies as absolute anomalies in parts per trillion by volume (pptv). WACCM simulates increases in ClO at 10 hPa and below, with anomalies of similar magnitudes to the WACCM NOx reductions. SNR analysis for the ClO response (Figure S12) indicates that the anomalies at 10 hPa, though consistent in terms of their coincidence with increased volcanic aerosol concentrations, may be difficult to distinguish from natural variability in the observations. The most distinct volcanic responses, as determined by high SNR relative to the background periods, are at 51 hPa in 2006–2007 and at 51 and 72 hPa in 2008, 2009, and 2011. WACCM ClONO2 anomalies (Figure S13) show no discernible response to the increased volcanic aerosol, which indicates that the ClO increase at these levels is simply a result of the reductions in NOx and not related to chlorine heterogeneous chemistry (e.g., (R1) and (R4)).

MLS ClO show consistent increases associated with volcanic aerosol at 4 hPa only (Figure S14a), which is the opposite of the vertical extent of the response simulated in WACCM. Anomalies at 21 hPa are generally positive at times of increased volcanic aerosol concentrations as well, specifically in 2007, at NH midlatitudes in 2009 and in the tropics in 2011 after Nabro (13°N; though this does not meet the 1σ aerosol criterion). However, anomalies in 2005 at the same level do not appear to be affected by increased volcanic aerosol concentrations. Overall, positive MLS ClO anomalies at times of increased volcanic aerosol concentrations do not stand out when compared to anomalies during volcanically clean periods. However, the changes in ClO simulated by WACCM are well below 100 pptv, which is the reported precision of the MLS product at these levels; therefore, it is probably the case that this response is simply undetectable in the observations.
5. Discussion

Figure 12 shows the Pearson correlation coefficients between sulfate aerosol SAD and the simulated and observed changes in NO\textsubscript{x}, HNO\textsubscript{3}, and ClO. For WACCM (Figures 12a–12c), the mean of the correlations from the ensemble members is shown; the correlations from individual ensemble members are in general very similar. WACCM simulates high anticorrelations between NO\textsubscript{x} and SAD (Figure 12a), with correlation coefficients less than \textasciitilde0.8 in the 20°S to 20°N band and from about 50–20 hPa. High

Figure 9. (a–e) WACCM HNO\textsubscript{3} anomalies (%) for the ensemble mean with QBO + N\textsubscript{2}O fits removed. Black line contours indicate where sulfate aerosol surface area density is greater than 1σ above the mean, and triangles indicate the timing and latitudes of the volcanic eruptions listed in Table S1. WACCM = Whole Atmosphere Community Climate Model; QBO = quasi-biennial oscillation.
Correlations between HNO\(_3\) and SAD (Figure 12b) are also simulated in WACCM, with correlations as high as 0.5 extending upward past 10 hPa, in agreement with the WACCM HNO\(_3\) time series discussed above. ClO correlations with SAD (Figure 12c) show a similar spatial pattern to the NO\(_x\) correlations but are generally lower in magnitude. In all three species, the magnitudes of the correlations generally decrease with increased distance from the equator. It is likely that higher aerosol concentrations and longer aerosol lifetimes in the tropics contribute to the generally higher correlations at these latitudes.

Figure 10. (a–e) MLS HNO\(_3\) anomalies (%) with quasi-biennial oscillation + N\(_2\)O fits removed. Black line contours indicate where sulfate aerosol surface area density is greater than 1σ above the mean, and triangles indicate the timing and latitudes of the volcanic eruptions listed in Table S1. MLS = Microwave Limb Sounder.

correlations between HNO\(_3\) and SAD (Figure 12b) are also simulated in WACCM, with correlations as high as 0.5 extending upward past 10 hPa, in agreement with the WACCM HNO\(_3\) time series discussed above. ClO correlations with SAD (Figure 12c) show a similar spatial pattern to the NO\(_x\) correlations but are generally lower in magnitude. In all three species, the magnitudes of the correlations generally decrease with increased distance from the equator. It is likely that higher aerosol concentrations and longer aerosol lifetimes in the tropics contribute to the generally higher correlations at these latitudes.
Observations in Figures 12d–12f show overall good agreement on the sign of the correlation, and the top and bottom rows appear to be spatially coherent, with Pearson correlation coefficients of 0.57, 0.56, and 0.28 below 10 hPa for NO\textsubscript{x}, HNO\textsubscript{3}, and ClO, respectively. On the other hand, the magnitudes of the observed correlations are generally much lower than those simulated in WACCM. In addition, there are several noticeable differences in Figure 12 between the model (top row) and observations (bottom row). OSIRIS NO\textsubscript{x}
aerosol correlations (Figure 12d) appear to be the lowest in the 20-hPa range and are actually larger than WACCM correlations (Figure 12a) at 10 hPa and above. The spatial patterns of HNO$_3$-aerosol correlations agree quite well between WACCM and MLS at and below 10 hPa (Figures 12b and 12e, respectively) and at 5 hPa in the NH. Correlations between MLS ClO and volcanic aerosol are mostly positive (Figure 12f), in agreement with the WACCM simulations (Figure 12c), though as previously mentioned, Figure S14 indicates that these correlations are purely coincidental.

Though the chemical response to volcanic aerosol was identified robustly in both the model and the observations, we have also identified at some points considerable differences in the timing, location, and magnitude of the responses. In particular, the model responses are in general stronger than the observed responses. While these differences may be in part attributed to potential differences in the timing and locations of the modeled and observed aerosol, the aerosol properties in the WACCM SAD data set have been validated extensively in Mills et al. (2016) and Schmidt et al. (2018). Among other factors that could affect correlations found in observations versus the model are model errors in chemistry, noise in the observations, and interference between the aerosols and the chemical signals needed for the satellite retrievals. Changes in solar irradiance were not included in the model simulations; such changes can affect the dynamics (Kodera & Kuroda, 2002) and chemistry (Matthes et al., 2010; Shindell et al., 1999) of the stratosphere, though the difference in, for example, ozone abundance was found to be less than 5% between solar maxima and minima. In addition, a 28-month, cyclic QBO was imposed in the WACCM ensemble. The repeated 28-month cycle does not capture the full variability of the QBO, including potential aerosol effects on the QBO. In this way, simplifications in the model setup certainly play a role in the more consistent chemical response in the model. However, tests suggest that the time intervals considered may be the most important driver of the differences and are constrained by data availability. The 2005–2014 period for analysis of observations was...
used purely because of the availability of MLS measurements, while 1999–2013 was used for WACCM because years outside this interval were forced with repeating year 2000 aerosol loading in these simulations. Relative to the 1999–2004 period, 2005–2013 was much more volcanically active. In terms of sulfur dioxide (SO₂) loading, the eruptions in Table S1 spanning 1999–2004 account for an average of 0.06 Tg SO₂/year, while the average for the eruptions over 2005–2013 is 0.8 Tg SO₂/year. We demonstrate the impact that the choice of base period and background concentrations has on the response in Figures S15–S18, where we use the same period as in the observations (2005–2013) to compute the WACCM anomalies. This has a very large effect on the results. Figures S15–S17 show that NO₂/ClO-SAD correlations are about 20% lower in magnitude when excluding 1999–2004. This is further illustrated in the WACCM SAD correlation coefficients in Figures S18a–S18c, which are reduced by up to half when considering only 2005–2013, and then show much better agreement with the observations (Figures S18d–S18f) in terms of magnitudes of the correlations.

Park et al. (2017), among others, showed the strong dependence of variability of stratospheric trace gases and the QBO, as well as the compact relationship between N₂O and NOₓ, we have shown that the latter can be exploited to remove anomalies related to dynamical variability, the most prevalent form of which is the QBO (e.g., Randel & Wu, 1996). Application of the N₂O regression in conjunction with a traditional QBO regression has been shown to be a more effective method for removing a majority of transport-related anomalies than either one alone, which is essential for isolating the chemical response to volcanic aerosol.

### 6. Conclusions

In this study, we sought to probe the stratospheric NOy/Cly response to recent increases in stratospheric volcanic aerosol concentrations. Related to the occurrence of a series of moderate-magnitude volcanic eruptions between 2005 and 2014, we found a clear response in stratospheric NOₓ concentrations in both the WACCM ensemble and the observations, with some distinct differences. While OSIRIS NOₓ shows decreases during times of increased volcanic aerosol concentrations at and below 10 hPa, the response is much more consistent in WACCM, with respect to both the mean and individual ensemble members; this is evident in the much higher correlation coefficients for WACCM NOₓ. In general, WACCM appears to simulate the chemical response much more consistently. When hydrolysis of N₂O₃ increases, depleting the NOₓ reservoir, HNO₃ is enhanced. This response is also seen in the WACCM ensemble and MLS observations, though there are some differences in the timing and location as compared to the NOₓ response. In particular, HNO₃ enhancements are more prevalent at higher altitudes than NOₓ reductions. In addition, compared to lower altitudes, some of the largest perturbations from the mean in the WACCM SAD data set at ~20–5 hPa occur before 2005; this undoubtedly has an impact on the somewhat reduced response in MLS HNO₃ for which the observations begin in 2005. For ClO, WACCM simulates robust increases at and below 20 hPa as a result of the NOₓ depletion. Similar ClO enhancements are absent from the MLS observations but are likely below the detection threshold for the instrument.

In summary, our results demonstrate that recent moderate volcanic eruptions have affected NOₓ, HNO₃, and ClO concentrations in the stratosphere. The use of N₂O data along with QBO winds as a means of accounting for dynamical and chemical variability has also been shown to allow improved identification of such chemical signatures as distinct from dynamical noise. Finally, ensemble simulations are an important tool in ensuring a robust understanding of the SNR ratios of such responses and hence interpreting the single realization available in observations. The results presented here further highlight the impact of moderate volcanic activity, can help to improve predictability following a future volcanic eruption, and underline the importance and utility of N₂O observations for distinguishing the effects of heterogeneous chemistry from natural variability on stratospheric composition.

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