



African volcanic emissions influencing atmospheric aerosol particles over the Amazon rain forest

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Abstract.

Long-range transport (LRT) plays an important role in the Amazon rain forest by bringing in different primary and secondary aerosol particles from distant sources. The atmospheric oxidation of dimethyl
30 sulfide (DMS), emitted from marine plankton, is considered an important sulfate source over the Amazon rain forest, with a lesser contribution from terrestrial soil and vegetation sulfur emissions. Volcanic sulfur emissions from Africa could be a source of particulate sulfate to the Amazonian atmosphere upon transatlantic transport but no observations have been published. By using satellite observations, together with ground-based and airborne aerosol particle observations, this paper provides
35 evidence of the influence that volcanic emissions have on the aerosol properties that have been observed



in central Amazonia. Under the volcanic influence, sulfate mass concentrations reached up to $3.6 \mu\text{g m}^{-3}$ (hourly mean) at ground level, the highest value ever reported in the Amazon region. The hygroscopicity parameter was higher than the characteristic dry-season average, reaching a maximum of 0.36 for accumulation mode aerosol particles. Airborne measurements and satellite data indicated the transport of two different volcanic plumes reaching the Amazon Basin in September 2014 with a sulfate-enhanced layer at an altitude between 4 and 5 km. These observations show that remote volcanic sources can episodically affect the aerosol cycling over the Amazon rain forest and perturb the background conditions. Further studies should address the long-term effect of volcanogenic aerosol particles over the Amazon Basin by running long-term and intensive field measurements in the Amazon region and by monitoring African emissions and their transatlantic transport.

1 Introduction

Sulfate aerosol particles are produced in the atmosphere by oxidation of sulfur dioxide (SO_2), emitted by fossil fuel (FF) combustion, volcanic emissions, and by oxidation of DMS (Andreae and Rosenfeld, 2008). These particles scatter solar radiation and act as efficient cloud condensation nuclei (CCN) (Stevens and Feingold, 2009). Anthropogenic SO_2 emissions have increased over the 20th century to a maximum around the year 1980 and declined somewhat thereafter to around 100 Tg SO_2 per year, but they are still the most important source of sulfur to the atmosphere (Boucher et al., 2013; <http://edgar.jrc.ec.europa.eu/overview.php?v=431>, last accessed 6 Sep 2017). Volcanic emissions are the predominant natural source of SO_2 and account for about 5 % of total SO_2 emissions (Yang et al., 2017). Sulfur dioxide is oxidized in the atmosphere to gaseous sulfuric acid, which is quickly converted to sulfate aerosol particles (Chin et al., 1996; Reiner and Arnold, 1994). Volcanic sulfur emissions can account for 20 – 40 % of sulfate aerosol particle mass concentrations in the middle troposphere at 650 hPa (Chin and Jacob, 1996). Volcanic eruptions can change the atmospheric composition (gas and particle phase) drastically in large areas (Mather et al., 2003). Two prominent examples are the Pinatubo eruption in 1991 (Kirchner et al., 1999) and the 2014 – 2015 eruption of the Holuhraun volcano in Iceland (Ilyinskaya et al., 2017), where the emissions affected the cloud-drop effective radius



(r_{eff}) while other cloud properties like the cloud optical depth and the cloud liquid water path remained unaffected (Malavelle et al., 2017; Yuan et al., 2011). Moreover, a connection between tropical volcanic explosive eruptions and El Niño-like events has been described recently (Khodri et al., 2017). Besides explosive-effusive eruptions, small eruptions and passive degassing account for relatively stable SO₂ fluxes ($23.0 \pm 2.3 \text{ Tg yr}^{-1}$, 2005–2015), and approximately one order of magnitude higher than explosive eruptive SO₂ fluxes (Carn et al., 2017). To what extent volcanic passive sulfur emissions can affect cloud properties is still debated (Ebmeier et al., 2014; Malavelle et al., 2017). There are several active volcanoes in Africa. The Nyamuragira-Nyiragongo neighboring volcanoes in the Democratic Republic of the Congo (DRC) were among the most persistent passively degassing volcanoes worldwide between 2004 and 2014, with around 150 days with detected degassing in 2014 and the highest average SO₂ index observed in the period 2004 – 2014 (Carn et al., 2016). The strong passive degassing activity of Nyamuragira starting in 2011 and culminating in a formation of a lava lake in late 2014 (Campion, 2014) lead to enhanced SO₂ emission from the Nyiragongo-Nyamuragira complex for several years (Bobrowski et al., 2017).

In the Amazon rain forest, biogenic sulfate aerosol is sustained by oceanic DMS emission, and to a lesser degree by hydrogen sulfide (H₂S), methanethiol (MeSH), and DMS emissions from soils and vegetation (Andreae and Andreae, 1988; Jardine et al., 2015; Martin et al., 2010). Occasional anthropogenic sulfur injections have been attributed to open biomass burning and fossil fuel combustion emissions, either from Brazil or, via LRT, from Africa (Andreae et al., 1990; Talbot et al., 1988). A recent study suggests that fossil fuel sources (e.g., ship traffic, power plants) do not significantly influence aerosol particles measured in the remote Amazon forest (Saturno et al., 2017) but can be important downwind of populated areas like Manaus, Brazil (Kuhn et al., 2010; Martin et al., 2016). Recent measurements during the South American Biomass Burning Analysis (SAMBBA) campaign, which focused on biomass burning (BB) emissions, found no correlation between sulfate aerosol and various kinds of BB aerosol particles (Brito et al., 2014). On the other hand, aircraft observations of haze layers at 2 – 4 km altitude over the Amazon rain forest have shown high sulfate enrichment in comparison to the boundary layer and the free troposphere concentrations and indicated these haze layers to be linked to LRT of aerosol particles from Africa (Andreae et al., 1988; Holanda et al., 2017).



90 A modeled global sulfate source attribution study showed that southern Africa peak sulfate concentrations occur between June and August. For this three-month period, estimated emissions were 0.81 Tg S and decreased in the following three months to 0.66 Tg S (Yang et al., 2017).

Northeasterly and southeasterly trade winds (north and south of the inter-tropical convergence zone (ITCZ), respectively) are able to transport aerosols over large distances given the typically weak wet
95 deposition in this latitude band (Wang et al., 2016). During the Amazonian dry season (August – November), the transport of African smoke from southern Africa savanna and shrubland fires is an important source of aerosol in addition to the emissions by regional fires in South America (Andreae et al., 1994). When the ITCZ shifts north in the dry season, south-east trade winds originating from southern Africa are more likely to reach the central Amazon rain forest. Even though the potential
100 impact of transatlantic transport of volcanic sulfur emission has been suggested (Yang et al., 2017), no ground-based evidence has been reported previously in the literature concerning the impact of African volcanic sources.

In this paper, we present satellite observations that show volcanic SO₂ emissions in central Africa that have been transported over the South Atlantic Ocean and reach the Amazon rain forest after being
105 oxidized to particulate sulfate. Satellite, airborne, and ground-based observations are used to show that volcanogenic sulfate can significantly affect the aerosol physical and optical properties over the Amazon Basin during the dry season.

2 Data and methods

2.1 Ground-based instrumentation

110 The ground-based aerosol data presented here have been collected at the Amazon Tall Tower Observatory (ATTO) site (called *T0a* in the GoAmazon2014/5 experiment, (Martin et al., 2016)), located in the Uatumã Sustainable Development Reserve, Amazonas, Brazil. Details about the ATTO site infrastructure, instrumentation and an overview of running measurements can be found elsewhere (Andreae et al., 2015). Figure 1 shows the ATTO site location and the place of the Nyamuragira



115 volcano in the DRC located at 1.41° S, 29.2° E, 3058 m a.s.l. The long-term measurements at ATTO
started in 2012. A systematic backward trajectory (BT) analysis of air masses arriving at ATTO can be
found in Pöhlker et al. (2017a). The ATTO aerosol measurements were taken on a triangular mast (02°
08.602' S, 59° 00.033' W, 130 m above sea level, a.s.l.) using a 25 mm diameter, 60 m high stainless
steel tube with a laminar sampling flow rate of 30 L min⁻¹. The instruments were installed inside an
120 air-conditioned container where the temperature was kept between 29 and 31 °C.

Equivalent black carbon (BC_e) mass concentrations, M_{BC_e} , were calculated from absorption
measurements by a multi-angle absorption photometer, MAAP (Model 5012, Thermo Electron Group,
Waltham, USA). The details of the instrument are described elsewhere (Petzold and Schönlinner, 2004).
The BC mass absorption cross-section (MAC) was retrieved from fitting MAAP absorption coefficients
125 at 637 nm wavelength and refractory black carbon (rBC) mass concentrations measured by using a
single-particle soot photometer (SP2) revision D (Droplet Measurement Technologies, Boulder, USA).
Details of the technique can be found in Stephens et al. (2003). The MAC calculations are described in
Saturno et al. (2017). Light scattering coefficients were measured using a three-wavelength integrating
nephelometer (Aurora 3000, Ecotech Pty Ltd., Knoxfield, Australia). For details of the instrument, see
130 Müller et al. (2011). Absorption and interpolated scattering measurements at 637 nm wavelength were
used to calculate the single scattering albedo of dry aerosol particles, ω_0 , at this wavelength, which is
defined as the ratio of scattering to extinction coefficients (extinction = scattering + absorption). A
detailed study of aerosol optical properties measured at the ATTO site can be found in Saturno et al.
(2017).

135 An aerosol chemical speciation monitor (ACSM) (Aerodyne Research Inc., Billerica, USA) was used to
measure online non-refractory aerosol chemical composition (Carbone et al., 2017). These
measurements started in February 2014 at the ATTO site. The technique resolves the sub-micron
aerosol chemical species in the following categories: Organics, sulfate, nitrate, ammonium and chloride
(Ng et al., 2011). In this paper, we only use organics and sulfate mass concentration data, M_{org} and
140 $M_{sulfate}$, respectively.




Cloud condensation nuclei (CCN) number concentrations, N_{CCN} , were measured with a CCN counter (CCNC, model CCN-100, Droplet Measurement Technologies, Boulder, USA), which was deployed at the ATTO site starting in March 2014. The instrument scanned over a range of different supersaturations and particle diameters; more details can be found elsewhere (Pöhlker et al., 2016). The
145 hygroscopicity parameter, κ , retrieved for a CCN activation ratio of 50 % is used in this study. Condensation nuclei number concentrations (> 10 nm), N_{CN} , were measured with a condensation particle counter (CPC, model 3022A, TSI, USA).

2.2 Airborne *in-situ* measurements

Chemical species of sub-micron aerosol particles were measured using a compact time-of-flight aerosol
150 mass spectrometer (C-ToF-AMS) installed on board of the German High-Altitude and Long Range Research Aircraft (HALO, <http://www.halo.dlr.de>, last visited 13 September 2017), a modified business jet G550 (Gulfstream, Savannah, USA). The C-ToF-MS details are presented elsewhere (Drewnick et al., 2005). The measurements took place between 6 September and 1 October 2014, during the
155 “Aerosol, Cloud, Precipitation, and Radiation Interactions and Dynamics of Convective Cloud Systems” (ACRIDICON) - “Cloud Processes of the Main Precipitation Systems in Brazil: A Contribution to Cloud Resolving Modeling and to the GPM (Global Precipitation Measurement)” (CHUVA) campaign over the Amazon rain forest (Machado et al., 2014). More details on the flight trajectories and instrumentation can be found elsewhere (Wendisch et al., 2016). In this study, only data up to 7 km altitude have been used.

160 2.3 Air mass trajectories

To investigate the probability of the volcanic sulfate plume reaching the ATTO site, trajectories were calculated using the National Oceanic and Atmospheric Administration (NOAA) hybrid single-particle Lagrangian integrated trajectory HYSPLIT model (Draxler and Hess, 1997, 1998; Stein et al., 2015). NOAA Global Data Assimilation System (GDAS) (Kleist et al., 2009) data at  1° resolution were
165 used as meteorological input for HYSPLIT.



2.4 Satellite SO₂ VCD data

As one of the most abundant gases in a volcanic plume, SO₂ is often used as a tracer for volcanic emissions by a variety of spectroscopic remote sensing techniques. The strong characteristic absorption features in the UV spectral range allow the quantification of SO₂ using differential optical absorption spectroscopy (DOAS, see e.g. Platt and Stutz, 2008; Richter and Wagner, 2011), both from the ground (e.g. Bobrowski and Platt, 2007; Galle et al., 2003) and from space (e.g., Eisinger and Burrows, 1998; Khokhar et al., 2005; Krueger and J., 1985).

The ozone monitoring instrument (OMI) on board of the National Aeronautics and Space Administration (NASA) Aura satellite, launched in 2004, detects backscattered solar radiation in the UV-visible range (Levelt et al., 2006). The polar-orbiting instrument crosses the equator at 13:30 local time. DOAS analysis of OMI spectra yields column densities of trace gases such as NO₂, SO₂, and HCHO with a spatial resolution of about 13 × 24 km² away from the swath edges. OMI's wide swath of 2600 km allowed daily global coverage until the first occurrence of the so-called row anomaly in June 2007, an instrumental problem that causes grievous radiance errors in up to half of OMI's ground pixels (Van Hoek and Claas, 2010). The row anomaly strongly affects the reliability of observations; therefore all affected pixels were removed from the data set prior to analysis.

The OMI SO₂ vertical column density (VCD) data presented in this paper were retrieved using the NASA's principal-component based algorithm with an a-priori vertical profile representative of a volcanic plume in the middle troposphere (TRM, Li et al., 2013, 2017). The assumption that the volcanic plume is located in the mid-troposphere is justified by the elevation of the volcano (3700 m), the strength of the eruption, and, particularly, the HYSPLIT trajectory analysis presented later in this paper. It is, however, important to note that the sensitivity of the satellite measurements depends systematically on plume altitude. Thus, the absolute values of the SO₂ VCD derived from the satellite observations over-estimate the true values if the plume is located at a higher or lower altitude, respectively. Fortunately, this does not influence our study, as the focus of this paper is on the spatial pattern of the SO₂ plumes, and not on SO₂ amount.



The level-2 data were downloaded from: <https://mirador.gsfc.nasa.gov/> (last visited 27 October 2017) and gridded to a regular, $0.1^\circ \times 0.1^\circ$ grid for easily handling.

Multi-year OMI SO₂ VCD daily averages from 11° S to 17° N are summarized in Fig. S1 as a function
195 of time and longitude. The figure shows observations during the ATTO measurement period (March 2012 to July 2017) and a snapshot of September 2014. Given that the Nyamuragira and Nyiragongo volcanoes are so close to each other (within ~15 km), their emissions detected by remote sensors are often treated as a paired source (Carn et al., 2017). Hereafter, the term “Nyamuragira” refers to the couple Nyamuragira-Nyiragongo in this text. Time series of area-averaged OMI SO₂ observations are
200 shown in Fig. 2a. Nyamuragira produced frequent intensive SO₂ emission events especially from 2012 to the end of 2015. The area where the average was calculated (Fig. 2b) corresponds to approximately 200,000 km² around Nyamuragira. Emissions from Nyamuragira were often transported westward, as can be observed in HYSPLIT forward trajectories calculated for 2014 (Fig. 2).

3 Results and discussion

205 3.1 Satellite measurements and trajectory analysis of the volcanic plume

The most important activity at the Nyamuragira volcano since 2011 occurred in September 2014 (Global Volcanism Program, 2017), coinciding with ground and airborne measurement campaigns in the Amazon Basin (Andreae et al., 2015; Wendisch et al., 2016; Martin et al., 2017). Satellite SO₂ VCD observations over central Africa and the Atlantic Ocean were examined during this period in order to
210 precisely identify the volcanic eruptive period and the plume trajectory. A map of gridded OMI SO₂ TRM VCD observations from 7 to 17 September 2014, is available as supplementary material (Fig. S3). Two important emission events were observed at the Nyamuragira location, one on 7 September and another on 12 September. The latter one exhibits a clear westward transport of the SO₂ plume starting from 13 to 17 September. Figure 3 shows SO₂ VCD observations during this period within the region
215 between 20° W – 30° E, and 15° S – 5° N with SO₂ VCD larger than 2.5×10^{16} molecules cm⁻². Several sets of trajectory calculations were performed. First, three to seven starting locations were selected within



the SO₂ plumes detected by OMI on 12–17 September 2014. At each location, 15-day (360 hours) forward trajectories were started at the time of the satellite overpass (11 – 14 UTC) at seven altitudes spaced equally between 1 and 7 km. The resulting trajectories initialized at 4 km altitude on
220 13 September are in best agreement with the satellite data and are shown in Fig. 3. All starting parameters were systematically varied and very consistent patterns were found. The trajectories are superimposed on the map presenting all SO₂ plumes detected by OMI between 12 and 17 September. Trajectories started within the leading edge of the plume are in good agreement with the OMI data, as after 24 hours (second marker) both trajectories are located within the plume detected on 14 September,
225 and after 48 hours (third marker) both trajectories are located within the plume detected on 15 September. The two southernmost trajectories make a sharp turn after 15 September, which is in agreement with the observed pattern, although there is no longer an exact match with the respective OMI observations (in red and maroon). This discrepancy may be due to inaccuracy of the individual trajectories, or the SO₂ plumes might have been below OMI's detection limit. The southernmost
230 trajectories reach South America and come within several hundreds of kilometers of ATTO within 15 days. One of them reached ATTO on 25 September at 1.8 km altitude, whereas the other one passed at an altitude of 1.5 km at the point nearest to ATTO on 24 September.

In addition to the plume forward trajectory analysis, backward trajectories initiated at the ATTO site at an altitude of 300 m (approximately 170 m above ground) were calculated for 360 hours. These
235 trajectories were initiated every hour from 20 September 0:00 UTC up to 30 September 23:00 UTC. The results, presented as a trajectory density plot in Fig. 4, indicate that although during this time period essentially all air masses come from southern Africa, they usually come from further south. Nevertheless, a significant number of trajectories originates close to the volcano and its plume.


3.2 Airborne observation of the volcanogenic aerosol particles

240 Enhanced sulfate aerosol mass concentrations were observed above 3 km height over the Amazon Basin during the ACRIDICON-CHUVA campaign compared to lower altitudes. A map including all airborne observations on the different flights can be found in the supplementary material (Fig. S4). However, given the multiple sulfate aerosol sources, sulfate itself can not be used as a tracer of volcanic




emissions. In order to distinguish the volcanogenic sulfate from additional aerosol sources like BB,
245 which is important during this time of the year, we examined the M_{sulfate} vertical profiles together with
their sulfate-to-OA mass ratio ($M_{\text{sulfate}} / M_{\text{org}}$). A list of the ACRIDICON-CHUVA flights and M_{sulfate}
vertical profiles are presented as supplementary information in Table S1 and Fig. S5, respectively. From
the different airborne observations, flight AC14 showed the highest sulfate-to-OA mass ratio, indicating
the strongest volcanogenic influence. The M_{sulfate} vertical profile measured on 21 September 2014
250 (AC14) is presented in Fig. 5. The observations show a sulfate-enhanced layer between 4 and 5 km
height. The average M_{sulfate} observed during flight AC14 was $1.1 \pm 0.5 \mu\text{g m}^{-3}$ between 3 and 6 km
height. This sulfate-enhanced layer exhibits a sulfate-to-OA ratio generally larger than 1. It can be
distinguished from lower layers, below 3 km height, which are characteristically rich in OA due to BB
and biogenic emissions. Usually, BB aerosol particles have been shown to have higher OA mass
255 concentrations than other aerosol particles (McNaughton et al., 2011; Saturno et al., 2017). Therefore,
the high sulfate-to-OA ratio is an indication of the volcanic origin of the probed aerosol. Backward
trajectories were calculated from several points along the flight paths. Figure 6 shows backward
trajectories started at nine points along the AC14 flight track where sulfate-to-OA ratios larger (colored
lines) or smaller than 1 (gray lines) were detected; the flight and aerosol data measured at each point are
260 presented in Table 1. Figure 6 clearly shows that the (colored) trajectories, with one exception,
initialized within the sulfate plume originate from central Africa, whereas the gray trajectories, started
outside of the sulfate plume, appear to originate from South America or from more southern regions
over or across the Atlantic Ocean. The air mass trajectory analysis indicates that the AC14 observations
were likely the result of probing the volcanic plume emitted on 7 September, the first one detected by
265 OMI (see Fig. S3). For flight AC17 a similar pattern is observed, with three out of four (colored)
trajectories started within the sulfate plume originating from central Africa and half of the other
trajectories clearly coming from regions more to the South (see supplement, Fig. S6 and Table S2).
Figures 3, 4, and 6 show that the trajectories agree well, but not perfectly with the ground-based,
airborne, and satellite measurements, which is mainly caused by the uncertainty of such long
270 trajectories. Nevertheless, the fact that forward and backward trajectories calculated from various
starting points and times agree on the sulfate source is a strong indication that the sulfate plumes



observed at and near ATTO originate from the Nya  agira volcano. Combined with the westward transport pattern derived from SO₂ satellite data and the lack of an alternative strong sulfate source makes this a quite convincing case.

275 3.3 Volcanic emission effects on the aerosol particle properties

The arrival of the African volcanic emissions over the Amazon rain forest affected the aerosol physical and chemical properties measured at the ATTO site. The most evident effect was the significant increase in M_{sulfate} . The 90th percentile of M_{sulfate} measured at the ATTO site during the dry season 2014 was used as a threshold to define the volcanic influence event (Nya2014) as the time when this
280 threshold was exceeded. By this criterion, the Nya2014 event spanned from 21 September 2014 at 02:00 UTC to 1 October 2014 at 01:00 UTC. Figure 7 shows different aerosol parameters measured before, during and after the Nya2014 event. The N_{CN}  shown in Fig. 7a, did not vary greatly from the values typical of the season (Pöhlker et al., 2016). The average N_{CN} during the dry season in 2014 was
285 $(1.3 \pm 0.6) \times 10^3$ particles cm⁻³, whereas during the Nya2014 event, there were three peaks lasting for few hours with particle number concentrations higher than 3.0×10^3 particles cm⁻³ on 27, 29, and 30 September 2014.

During the Nya2014 period, M_{sulfate} averaged $1.7 \pm 0.6 \mu\text{g m}^{-3}$, which was significantly above the dry season 2014 average of $0.7 \pm 0.3 \mu\text{g m}^{-3}$, see Fig. 7b. The highest M_{sulfate} value observed at the ATTO site was $3.6 \mu\text{g m}^{-3}$ (hourly mean) on 26 September 2014. For comparison, during the SAMBBA
290 campaign in southern Amazonia, M_{sulfate} barely exceeded $1.0 \mu\text{g m}^{-3}$, despite organics nearly reaching $100 \mu\text{g m}^{-3}$, M_{BCe} of $5 \mu\text{g m}^{-3}$ and N_{CN} above 25×10^3 particles cm⁻³ during the peak of biomass burning (Brito et al., 2014). A long-term measurement study, also conducted in southern Amazonia, reported
295 M_{sulfate} of $1.1 \pm 0.7 \mu\text{g m}^{-3}$ during the dry season, with a maximum of $3.3 \mu\text{g m}^{-3}$ for aerosol particles with mobility diameters smaller than $2 \mu\text{m}$ (Artaxo et al., 2002). It is important to note that Artaxo et al.'s (2002) sulfate observations were done under strong BB influence with average elemental carbon (EC) mass concentrations, M_{EC} , of $3.8 \pm 4.2 \mu\text{g m}^{-3}$, with a maximum of $25 \mu\text{g m}^{-3}$. In contrast, the BC_e measurements at the ATTO site during the Nya2014 event had an average of $0.4 \pm 0.1 \mu\text{g m}^{-3}$, with a maximum of $0.8 \mu\text{g m}^{-3}$, indicating that the BB influence was relatively weak during the period of



interest, with some short (few hours) spikes due to the influence of near-by fire events, see Fig. 7b. At a
300 sampling site impacted by Manaus emissions, the sub-micron M_{sulfate} was about $0.2 \mu\text{g m}^{-3}$ during the
wet season, rarely exceeding $0.6 \mu\text{g m}^{-3}$ (de Sá et al., 2017). Therefore, even considering a range of
pollution sources, our measurements at ATTO during the Nya2014 event are the highest sub-micron
sulfate concentration ever reported in the Amazon Basin; see Martin et al. (2010) for a summary of wet
and dry season aerosol observations. For comparison, the ACRIDICON-CHUVA airborne
305 measurements are also included in Fig. 7b. The M_{sulfate} measured on flight AC14 was significantly
enhanced between 3 to 6 km altitude, reaching a median of $1.0 \mu\text{g m}^{-3}$ and a 75th percentile of
 $1.6 \mu\text{g m}^{-3}$. Previous aircraft measurements during the SAMBBA campaign reported a M_{sulfate} flight
average of $0.48 \mu\text{g m}^{-3}$ (Allan et al., 2014).

The increased M_{sulfate} period was accompanied by an enhanced sulfate-to-OA mass ratio, according to
310 the ATTO observations (Fig. 7c). The Ny2014 sulfate-to-OA average over about 10 days was
 0.38 ± 0.09 , significantly higher than the dry season average of 0.18 ± 0.15 . During some BB pulses,
decreased sulfate-to-OA ratios were observed, but the whole Nya2014 period was exceptionally high
compared to the typical dry season conditions. The sulfate-to-OA values measured at ground level were
usually lower than the airborne values observed between 3 and 6 km height because the OA sources
315 (BB and biogenic emissions) are located at ground level and the LRT sulfate that arrives at higher
altitudes is diluted upon vertical mixing. The possibility of fossil-fuel (FF) burning was ruled out as an
important sulfur source during the event discussed here because of the particularly high dry-aerosol ω_0
measured during the event (0.89 ± 0.04), as can be observed in the color code data in Fig. 7c. Usually
FF emissions, rich in BC, present characteristically low ω_0 (0.2 – 0.3) (Bond and Bergstrom, 2006) and
320 its addition would have lowered the value of ω_0 . Instead, an increase in ω_0 was observed to values
higher than 0.90 during the period of maximum M_{sulfate} (26 – 27 September 2014).

The effects of the volcanic sulfur plume on the aerosol hygroscopicity was explored by analyzing the κ
values measured at different supersaturations. Higher κ values were measured for the accumulation
mode aerosol (particles with diameter greater than 100 nm) (Fig. 7d; note the color-coded particle
325 activation diameter, D_a). During the Nya2014 event, the κ values increased significantly, especially
when the maximum M_{sulfate} was observed. For example, the average κ for a supersaturation of 0.10 %



($D_a = 167 - 179$ nm, 25th and 75th percentile, respectively) was 0.26 ± 0.04 during the Nya2014 event, with a maximum of 0.36. The Nya2014 κ average was slightly higher than the 2014 dry season average of 0.21 ± 0.03 for 0.10 % supersaturation (excluding the volcanic event), and significantly higher than a strong BB event average of 0.18 ± 0.01 for 0.10 % supersaturation, whose high OA content (sulfate-to-OA < 0.04) caused a significant drop in the hygroscopicity parameter (Pöhlker et al., 2017b).

Summary and conclusions

Satellite SO₂ observations showed two explosive events at the Nyamuragira volcano on 7 and 12 September 2014. These emissions were observed to be transported over the South Atlantic Ocean. Air mass trajectory modeling from the plume location showed that the plume was transported towards South America, likely over the ATTO site and its surroundings in central Amazonia. Airborne observations during the ACRIDICON-CHUVA campaign showed a sulfate-enhanced layer between 4 and 5 km height on 21 September 2014 (flight AC14). Additionally, this layer exhibited an increased sulfate-to-OA mass ratio with medians higher than 1 for measurements between 3 and 6 km height. The ground-based M_{sulfate} measured at the ATTO site reached an hourly mean of $3.6 \mu\text{g m}^{-3}$ on 26 September 2014, the highest values ever reported in the Amazon Basin. The sulfate-to-OA mass ratio increased from a dry-season average of 0.18 ± 0.15 to 0.38 ± 0.09 during the volcanic influence event, which spanned for a period of about 10 days. Increased sulfate-to-OA and single scattering albedo (ω_0) were assumed as an indication of the low influence of BB and FF sources, respectively. In terms of aerosol hygroscopicity, the values of κ (for 0.10 % supersaturation) measured during the volcanic event reached an average of 0.26 ± 0.04 (and a maximum of 0.36), slightly higher than the dry season average of 0.21 ± 0.03 .

The evidence presented here shows one particular event of volcanic SO₂ emissions in Africa influencing the aerosol particles' chemical composition, hygroscopicity, and optical properties in the Amazon Basin. Therefore, our study indicates that these emissions and their transatlantic transport could potentially affect the Amazonian cloud microphysical properties. However, the extent and relevance of the episodic volcanic influence on the Amazonian atmosphere would require more extensive studies. Beyond the



effects and implications of this particular event, the results of our study represent a reference case of the dynamics and conditions of transatlantic aerosol transport from southern Africa to South America. This could help to understand the inter-continental advection of other aerosol species, such as combustion aerosol particles that are more difficult to trace.

Data availability. The data presented in this paper can be accessed via e-mail request to Jorge Saturno (j.saturno@mpic.de) or Christopher Pöhlker (c.pohlker@mpic.de). OMI data are available online at <https://disc.gsfc.nasa.gov/>.

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

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Table 1. Measurements at points along track of the flight AC14 (21 September 2014) selected as starting points for backward trajectories presented in Fig. 6. Data points with sulfate-to-OA > 1 are emphasized by bold font.

Time (UTC)	Latitude [°N]	Longitude [°E]	Altitude [km]	M_{sulfate} [$\mu\text{g m}^{-3}$]	Sulfate-to-OA	Color in Fig. 6
15:14	-2.75	-60.34	1.61	1.2	0.5	Gray
16:19	-3.19	-60.21	4.50	1.0	1.1	Blue
16:36	-4.00	-59.50	4.50	3.0	2.9	Green
16:54	-5.20	-59.25	4.49	3.1	7.8	Red
17:11	-6.45	-58.98	6.45	0.6	0.8	Gray
17:46	-4.52	-59.40	6.43	0.5	0.8	Gray
19:40	-4.44	-59.41	7.68	1.2	1.0	Gray
21:41	-3.79	-59.55	4.79	1.8	2.2	Light blue
21:59	-3.04	-60.20	0.90	1.2	0.3	Gray

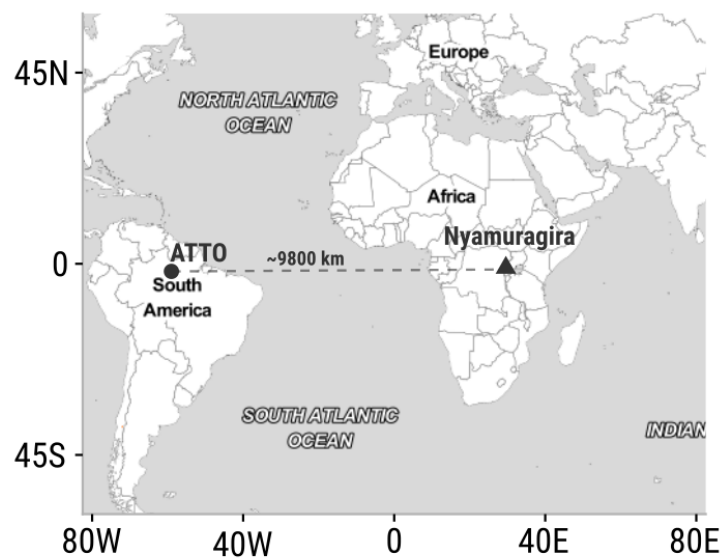


Figure 1. ATTO site and Nyamuragira volcano locations.

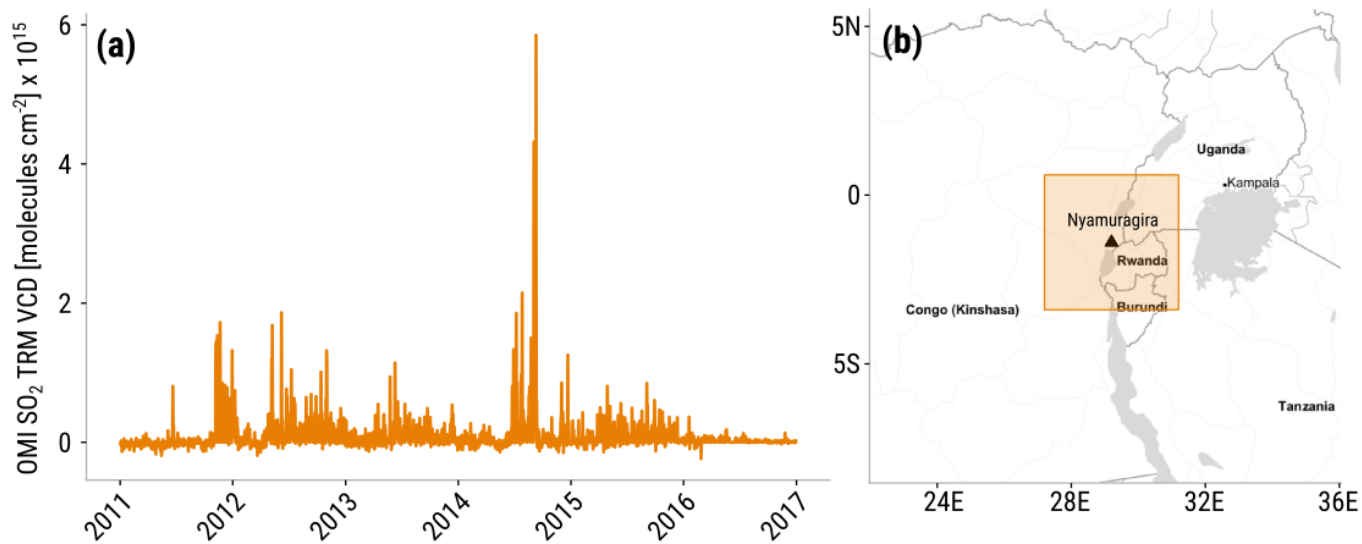


Figure 2. (a) Time series of daily-averaged OMI SO₂ TRM VCD observations corresponding to the averages over the area delimited by 27.2° E, 3.4° S, 31.2° E, and 0.6° N (b). Map of eastern Africa showing the averaging area (orange square). The location of Nyamuragira is represented by a black triangle.

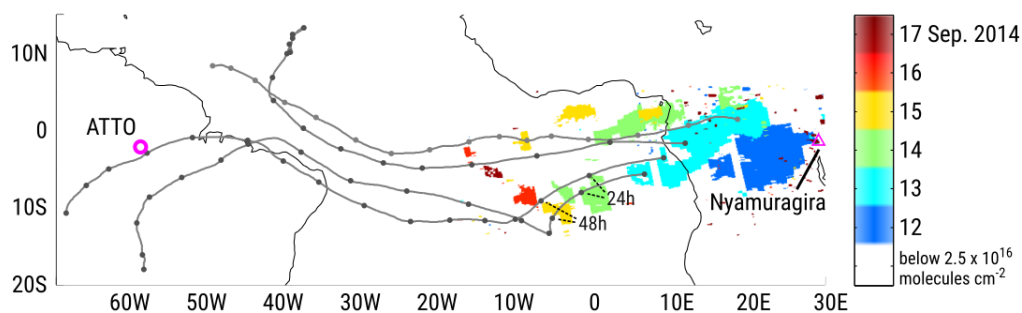


Figure 3. Map of SO₂ plumes with VCD > 2.5 × 10¹⁶ molecules cm⁻² color-coded by date of observation. Forward trajectories started at 4 km (above mean sea level, a.m.s.l.) at four locations within the plume detected on 13 September 2014 (light blue) are indicated by black lines with markers at 24-hour intervals. The ATTO site is marked by a pink circle; the location of Nyamuragira is indicated by a pink triangle.

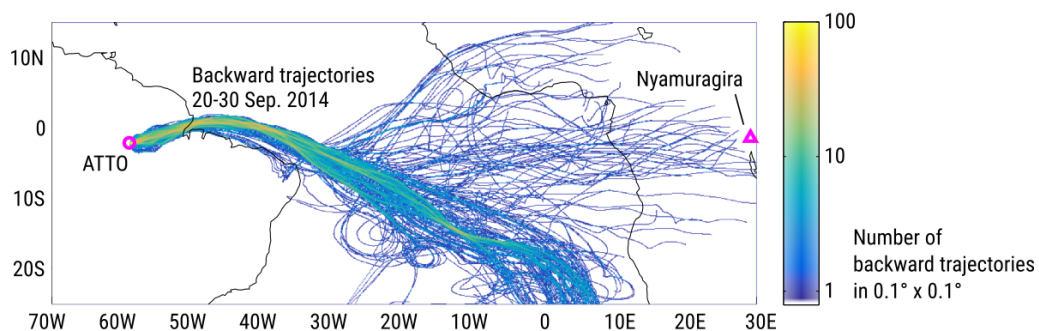


Figure 4. Density of backward trajectories started from ATTO at an altitude of 300 m a.m.s.l. on every hour starting at 0:00 UTC on 20 September 2014 up to 23:00 UTC on 30 September 2014. The locations of the ATTO site and Nyamuragira are shown on the map.

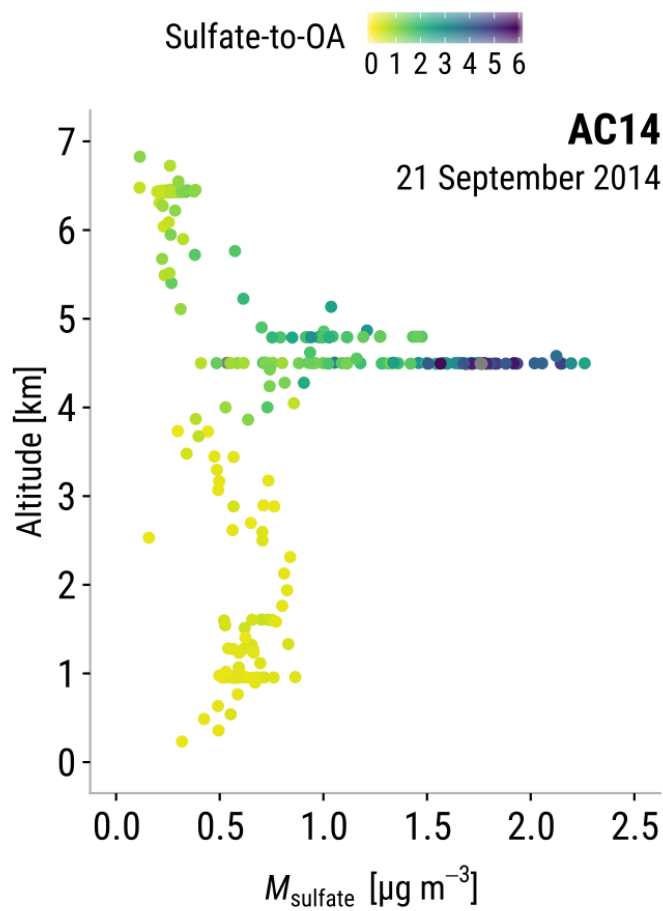


Figure 5. Sulfate layer observations over the Amazon rain forest. The figure shows the M_{sulfate} vertical profile observed during flight AC14 (21 September 2014). Color-coded sulfate-to-OA values are truncated at a maximum of 6.

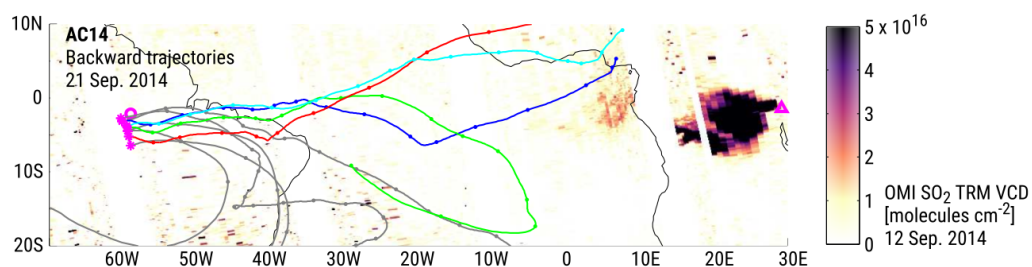


Figure 6. Map of gridded OMI SO₂ VCD, observed on 12 September 2014. Backward trajectories were started at several points along track of the flight AC14 (21 September 2014) at flight altitude. Trajectories starting at points where sulfate-to-OA > 1 are shown in color (see Table 1 for details), all other trajectories are shown in gray; dots are placed at 24-hour intervals. The path of flight AC14 is marked in pink, with stars denoting the starting points of the backward trajectories. The locations of the ATTO site and Nyamuragira are marked with a pink circle and triangle, respectively.

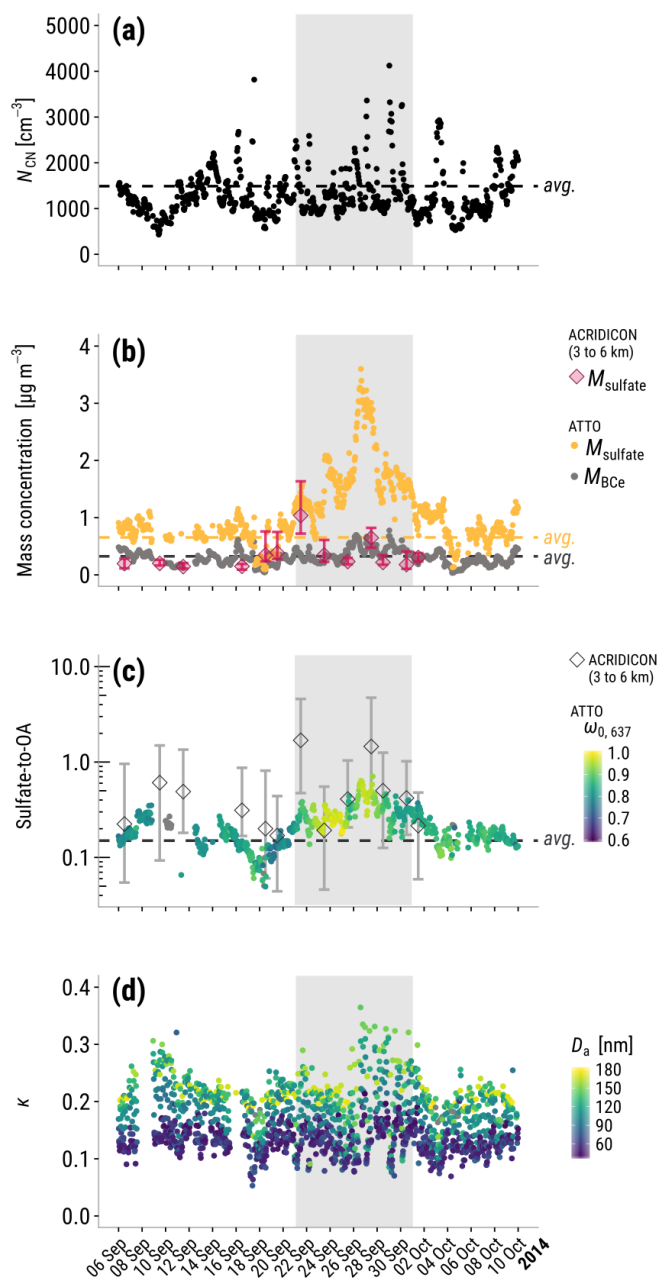


Figure 7. Different aerosol properties measured during the Nya2014 event (gray shaded area), These time series include ATTO hourly averages of (a) aerosol particle number concentration, N_{CN} , (b) sulfate and BC_e mass concentration, (c) sulfate-to-OA mass ratio with color-coded aerosol particle single scattering albedo at 637 nm wavelength, and (d) hygroscopicity parameter, κ , original time resolution, with color-coded particle activation diameter. Dry season averages are shown as dashed lines. ACRIDICON-CHUVA flight medians and inter-quartile ranges from 3 to 6 km altitude are shown in (b) and (c).