Evidence of New Particle Formation Within Etna and Stromboli Volcanic Plumes and Its Parameterization From Airborne In Situ Measurements

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Abstract

Volcanic emissions can significantly affect the Earth’s radiation budget by emitting aerosol particles and gas-phase species that can result in the new particle formation (NPF). These particles can scatter solar radiation or modify cloud properties, with consequences on health, weather, and climate. To our knowledge, this is the first dedicated study detailing how gas-phase precursors emitted from volcanic plumes can influence the NPF. A series of airborne measurements were performed around the Etna and Stromboli volcanoes within the framework of the CLerVolc and STRAP projects. The ATR-42 aircraft was equipped with a range of instrumentation allowing the measurement of particle number concentration in diameter range above 2.5 nm and gaseous species to investigate the aerosol dynamics and the processes governing the NPF and their growth within the volcanic plumes. We demonstrate that NPF occurs within the volcanic plumes in the free troposphere (FT) and boundary layer (BL). Typically, the NPF events were more pronounced in the FT, where the condensational sink was up to two orders of magnitude smaller and the temperature was ~20 °C lower than in the BL. Within the passive volcanic plume, the concentration of sulfur dioxide, sulfuric acid, and N2,5 were as high as 92 ppbV, 5.65 × 108 and 2.4 × 105 cm−3, respectively. Using these measurements, we propose a new parameterization for NPF rate (J2.5) within the passive volcanic plume in the FT. These results can be incorporated into mesoscale models to better assess the impact of the particle formed by natural processes, that is, volcanic plumes, on climate.

1. Introduction

Volcanic emissions are found to be one of the most abundant natural sources of particles and gases in the atmosphere (Bobrowski et al., 2007; Boulon et al., 2011; Haywood & Boucher, 2000; Oppenheimer et al., 2003, 2011; Robock, 2000; Tomasi & Lupi, 2016). Volcanos emit a wide range of different gases (SO2, CO2, H2O, H2S, HF, HBr, etc.) and particle types (ash and aerosol particles formed from condensable vapors and metals; Aiuppa et al., 2006; Bobrowski et al., 2007; Mather, 2015; Roberts et al., 2018; Simpson et al., 1999) into the atmosphere. Volcanic aerosols can scatter the solar radiation back to space contributing to a global cooling effect (direct effect; Albrecht, 1989; Haywood & Boucher, 2000; Robock, 2000) or modify the climatic impacts of clouds (indirect effect; Mather, 2015, and references within) by acting as cloud condensation nuclei (CCN; Gassó, 2008; Hobbs et al., 1982; Ilyinskaya et al., 2017; Mather, 2015; Mather et al., 2003) or ice nuclei (Hoyle et al., 2011). Moreover, volcanic emissions can have significant detrimental effects on human health, the impact of which depends on aerosol physical and chemical properties (Ilyinskaya et al., 2017; Schmidt et al., 2011, 2015).

The two main types of volcanic aerosols present in the atmosphere are either primarily emitted or secondarily formed (Mather, 2015; Mather et al., 2003; Petäjä et al., 2012; Roberts et al., 2018; Robock, 2000). The primary volcanic aerosols are mainly volcanic ash and can have diameters ranging from very fine ash (submicron) to 2 mm according to classic sedimentology. The very fine ash, which survives...
Sulfuric acid (SA), formed from the oxidation of SO2 through different channels, is known to be a key species in NPF processes (Kroll et al., 2015; Mauldin et al., 2003; Petäjä et al., 2011; Sipilä et al., 2010; Weber et al., 2012). While a large number of studies have investigated volcanic emissions through in situ ground-based and satellite/radar measurements (Carn et al., 2013; Galle et al., 2010; Kantzas & McGonigle, 2008; Mather, 2015; McCormick et al., 2016; McGonigle et al., 2017; McGonigle & Oppenheimer, 2003), airborne in situ measurements of volcanic emissions remain very scarce (Mauldin et al., 2003; Oppenheimer et al., 2010; Petäjä et al., 2012; Radke, 1982; Rose et al., 2006; Tulet et al., 2017; Vignelles et al., 2016; Weber et al., 2012). The limited number of volcanic plume airborne observations investigating NPF arises from challenges associated with restricted timescales and the impact of temporal and spatial plume's heterogeneities under typically harsh environments, besides the costly deployment of highly sophisticated instrumentation aboard an aircraft in such harsh conditions (Delmelle, 2003; Mauldin et al., 2003; Oppenheimer et al., 2003). In that context, the aim of this study is to investigate the aerosol dynamics and the processes governing aerosol formation and growth in different types of volcanic plumes. To the best of our knowledge, this is the first comprehensive dedicated study investigating how gas-phase precursors influence NPF events within different volcanic plumes over Etna and Stromboli using airborne measurements' platforms. Such investigation allows us to characterize the plume spatial extent, its properties, and its intensity and to derive a new parameterization of the rate of NPF. These will permit further improving the estimation of NPF from natural proximal sedimentation, usually ranges from submicron to a few microns, and they result from the fragmentation of the erupting magma into juvenile solid particles injected into a rising column and dispersed in the atmosphere (Allard et al., 2000; Rose & Durant, 2009). The secondary volcanic aerosol particles are produced from the gas-to-particle conversion (secondary formation) process, specifically from the oxidation of SO2 (Mather, 2015; Mather et al., 2004; Naughton et al., 1975; Schmidt et al., 2011), and this process is not yet well characterized within the volcanic plumes. This process is called new particle formation (NPF), where clusters are formed from the gaseous phase as a first step and, later on, grow to larger sizes (>100 nm) at which they can act as CCN (Hobbs et al., 1982; Mather et al., 2003) or ice nuclei (Hoyle et al., 2011) and impact the climate (Kerminen et al., 2012; Kulmala et al., 2001, 2004, 2014; Kulmala & Kerminen, 2008; Kulmala & Laaksonen, 1990; Makkonen et al., 2012). During active eruptions, both primary and secondary particles are present in different atmospheric vertical layers (Ilyinskaya et al., 2017; Mather & Pyle, 2015; Tulet et al., 2017). Conversely, during passive emissions, primary aerosols, with low concentrations, are often limited to the remobilization of accidental lithic (derived from the conduit and crater walls), while emissions of gaseous species may remain significant, likely to contribute to the formation of new particles. It is estimated that ~9 Tg/year of SO2 is emitted from degassing passive volcanoes worldwide (Allard et al., 1991; Mather et al., 2003; Mather & Pyle, 2015; Pyle & Mather, 2003), being roughly the same order of magnitude of continuously and sporadically eruptive volcanoes (Andres & Kasgnoc, 1998; Carn et al., 2016). Currently, volcanoes contribute to ~10% of the global budget of sulfur emission sources that are dominated by anthropogenic emissions (Allard et al., 1991; Smith et al., 2011). Past studies estimated that aerosol particles with diameters smaller than 0.1 μm contributed a total of 6% to 18% to the total aerosol volume in the passive plume from Etna in Italy (Watson & Oppenheimer, 2000), whereas in Stromboli (Italy), the contribution of particles in the nucleation and accumulation modes was estimated to be 66% of the total aerosol volume (Allard et al., 2000). Recent studies estimated that global NPF contributes up to 54% of CCN with a large uncertainty range of 38–66% in the present-day atmosphere (Gordon et al., 2017), which is higher than what has been estimated in past studies (Merikanto et al., 2009). In the preindustrial atmosphere simulations, NPF is shown to contribute up to 68% with an even larger range of uncertainty at 45–84% (Gordon et al., 2017). However, a large fraction of the uncertainty on the impact of aerosols on climate stems from the incomplete knowledge of the pre-industrial gas and aerosol concentrations and compositions (Carslaw et al., 2013; Gordon et al., 2016, 2017), therefore, further understanding of such natural processes is crucial.
sources, that is, volcanic degassing plumes, in models to evaluate more accurately the impact of those particles on climate.

2. Methodology and Measurements Conditions

2.1. The Volcanoes

The airborne measurements were conducted around the Etna and Stromboli volcanoes (Italy). Etna is located on the East coast of Sicily in the Mediterranean Sea (37.75°N, 14.99°E). The vent is located at 3,330 m above sea level (a.s.l.), typically in the free troposphere (FT). Mount Etna exhibits basaltic eruptions ranging from weakly explosive low-volume activity, such as Stromboli, to more powerful explosive activity leading to fire fountains, which feed columns of scoria, bombs, and ash as jets to heights of tens to hundreds of meters (Calvari et al., 2011). Occasionally, Mount Etna exhibits even more powerful eruptions and produces subplinian plumes injecting large amounts of ash and gas, although limited to the troposphere. Intereruptive periods are usually characterized by significant emissions of gas, making Etna volcano one of the most important SO2 emitter (Calvari et al., 2011). During the eruptive activity, the average flux of SO2 emitted at Etna is typically in the range 10–25 kt/day (Caltabiano et al., 1994) and decreases to 0.6–2 kt/day (Aiuppa et al., 2008; Roberts et al., 2018) during passive emissions. Stromboli is one of the Eolian Islands in the Mediterranean Sea located in the north coast of Sicily (38.79°N, 15.21°E), and the vent is at 924 m a.s.l., estimated to be in the boundary layer (BL) during our measurements in summer daytime (Seidel et al., 2012). Stromboli volcano is known to exhibit short-lived low-explosive activity with explosions occurring at a time interval of a few tens of minutes on average (Blackburn et al., 1976). The average flux of SO2 emitted during a standard level of activity lies in the range of 0.15–0.6 kt/day (Burton et al., 2008).

During the time of our campaign, Etna was not erupting, and only products of passive emissions could be recorded. On the contrary, at Stromboli volcano, the Northeast craters exhibited the typical Strombolian activity with small gas bursts accompanied by the ejection of ballistics every 5–10 min, while the Southwest crater produced less frequent ash-rich explosions. The threshold of the SO2 flux rate is up to ~5,000 t/day for Etna volcano and ~200–300 t/day for Stromboli volcano as reported by the National Institute of Geophysics and Volcanology in Italy for the week between 13 and 20 June 2016, Report. no. 25/2016 on 21 June 2016 (National Institute of Geophysics and Volcanology, 2016a, 2016b).

2.2. Research Flights

In 2016, as part of the CLerVolc and STRAP projects (Centre Clermontois de Recherche sur le Volcanisme and Trans-disciplinary collaboration to investigate volcano plumes risks), a series of airborne-based (French research aircraft, ATR-42) measurements were performed around Etna and Stromboli volcanoes on the 15 and 16 June 2016. The ATR-42, operated by the French SAFIRE Facility (Service des Avions Français Instrumentés pour la Recherche en Environnement), intercepted the volcanic plume close to the vent (~2.5 to 5 km) and tracked its evolution for up to 120 km. During this campaign, four flights were conducted: two around Etna (called herein ETNA13 and ETNA14) and two around the Stromboli (called herein STRO14 and STRO15; Table 1).

2.3. Airborne Instrumentations

The characterization of the aerosols and gases in the volcanic plumes involved installing a number of instruments in the ATR-42, including the following: (1) an ultrafine water-based condensation particle counters (TSI 3788; Kucp et al., 2013) to measure the total number of particle concentration at a cut-off size of >2.5 nm in diameter; 2) the CONDensation PArticle System (COPAS) condensation particle counters (Weigel et al., 2009), which is specifically dedicated to aircraft measurements, to measure particles number concentration at a cut-off size of >10 nm in diameter; (3) an optical particle counter (Sky OPC, Grimm, # 1.129) to measure the particle size distribution and number concentration in each size bin in the range distributed between 250 nm and 2.5 μm (according to the works of Pirjola et al., 1999, the optical particle counter data in that size range were used to calculate the condensational sink (CS); thus, it represents the lower limit of CS calculated for this study); (4) a UV Fluorescence SO2 Analyzer Teledyne API (T100 V) to measure the SO2 concentration with 10-s time resolution; and (5) a newly designed ambient ionization (AI) inlet coupled with the Atmospheric Pressure interface – Time Of Flight (AI-API-ToF) mass
spectrometer (MS), developed at the Institute for Atmospheric and Earth System Research of the University of Helsinki (Finland; Junninen et al., 2010), to measure SA concentration (more details about SA calibration are further given in section 2.4).

This combination of different instruments, all having a time resolution of 1 s, covered a wide particle size range (2.5 up to ~2,500 nm in diameter), allowing the measurements of aerosol physical properties and the detection of both nanoparticle nucleation and growth processes. The variables defined from the in situ measurements of total particle number concentrations (cm$^{-3}$) together with the SA and SO$_2$ are summarized in Table 2.

Differents meteorological variables, such as temperature ($T$), relative humidity (RH), wind speed ($W$), dew point temperature, pressure, and turbulence, were also measured aboard the aircraft with a 1‐s time resolution. The statistics and the variation range of the $T$, RH, and $W$ are shown in Table 3. The time series of the abovementioned meteorological parameters are displayed in Figure S1 (supporting information) for each flight.

### Table 1

<table>
<thead>
<tr>
<th>Date</th>
<th>Flight name and number</th>
<th>Takeoff-landing time (UTC)</th>
<th>Latitude range</th>
<th>Longitude range</th>
<th>Altitude range (m)</th>
<th>Pressure (hPa)</th>
<th>FT or BL</th>
</tr>
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</table>

Note. UTC = local time—2 hr.

### Table 2

<table>
<thead>
<tr>
<th>Variable</th>
<th>Name</th>
<th>Unit</th>
<th>Instrument/calculation method</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total number concentration for particles (N) at diameter &gt; 2.5 nm</td>
<td>$N_{2.5}$</td>
<td>cm$^{-3}$</td>
<td>Water CPC (TSI 3788)</td>
<td>Kupc et al. (2013)</td>
</tr>
<tr>
<td>N at diameter &gt; 10 nm</td>
<td>$N_{10}$</td>
<td>cm$^{-3}$</td>
<td>CPC</td>
<td>Weigel et al. (2009)</td>
</tr>
<tr>
<td>N at diameter &gt; 250 nm</td>
<td>$N_{250}$</td>
<td>cm$^{-3}$</td>
<td>Sky OPC GRIMM, # 1.129</td>
<td>Manual (GRIMM, 2008)</td>
</tr>
<tr>
<td>N at diameter range between 2.5 and 10 nm</td>
<td>$N_{2.5-10}$</td>
<td>cm$^{-3}$</td>
<td>Derived in this study</td>
<td></td>
</tr>
<tr>
<td>N between 10 and 250 nm</td>
<td>$N_{10-250}$</td>
<td>cm$^{-3}$</td>
<td>$N_{10} - N_{250}$</td>
<td>Derived in this study</td>
</tr>
<tr>
<td>Sulfuric acid concentration</td>
<td>SA</td>
<td>cm$^{-3}$</td>
<td>CI-API-ToF</td>
<td>Junninen et al. (2010)</td>
</tr>
<tr>
<td>Sulfur dioxide mixing ratio</td>
<td>SO$_2$</td>
<td>ppbV</td>
<td>UV Fluorescence Analyzer Teledyne API</td>
<td>Manual (Model T100U Trace Level Sulfur Dioxide Analyzer, 2011)</td>
</tr>
</tbody>
</table>

Note. CPC = condensation particle counters, OPC = optical particle counter
For flights ETNA13 and ETNA14 that took place in the FT, the temperature was above 5 °C, reaching a maximum of 17.5 °C at ~2 km in altitude. In STRO14 and STRO15 that took place in the BL, the temperature was always detected over 20 °C at lower altitudes, and it was 10 to 20 °C higher than the temperatures observed in the FT. The RH did not exceed 62.5% and 82% for flights ETNA13 and ETNA14, and it was even lower in the Stromboli plumes (Figure S1 and Table 3). The measurements of the cloud droplet probe and fast cloud droplet probe together with the RH measurements confirm that flights were undertaken in cloud-free conditions.

2.4. SA Calibration From AI-API-ToF

Following the strict aircraft instrumentation regulation concerning chemicals, a new AI inlet has been developed for the field campaign avoiding the use of chemical reagents. The AI-API-ToF is used for the first time on board an aircraft to perform such measurements and provides a 1-s time resolution of a lower estimate of SA concentration. The system utilizes a soft X-ray source (Hamamatsu L9490) to ionize directly the sampled air and increase the overall signal for fast (1 s) measurement. The instrument was also operated in ion mode, like a classical APi-ToF MS mode, where only natural ions are sampled. However, in the ion mode, a long integration time (minimum 10 min) is necessary to obtain the correct signal. The X-ray source was periodically switched ON and OFF (for these flights, 10 s ON and 10 s OFF), allowing both sampling of natural ions and forced ionized ions. The ATR-42 was flying at an average speed of 360 km/hr; hence, we use the X-ray mode (with 1-s resolution) to first identify the different air masses (i.e., inside the plume and outside the plume) and in a second step to analyze the average natural ions spectrum within the volcanic plume.

After the flight campaign, a calibration campaign took place during CLOUD11 at the CLOUD chamber similarly to previous CLOUD experiments (Kirkby et al., 2011; Rondo et al., 2016). In the CLOUD chamber, various atmospheric systems were studied with a wide range of species, that is, SO2 (0–2.6 ppb), NOx (0–33 ppb), and organic vapors (alpha-pinene (0–4.5 ppb), isoprene (0–5 ppb), trimethylbenzene (0–9 ppb), allowing the characterization of the new flying AI in different atmospheric systems. During this calibration campaign, the new flying AI inlet worked mainly in O2− chemical ionization modes; however, when NOx was at high concentrations in the chamber, NO3− ionization could also contribute. Estimates of SA concentration are conducted from the signal that is produced from the ionization of O2− ions (HSO4−) and NO3− (H2SO4·NO3), obtaining a good correlation between well-characterized nitrate API-ToF MS systems and the instrument used here (supporting information Figure S3).

2.5. Backward Trajectories

The 72-h air mass backward trajectories were calculated first at the vent of the volcano at the beginning of the flight and then at the flight track every 10 min along the path of each flight trajectory using the HYSPPLIT model (Draxler, 2003; Stein et al., 2016; Figure S2). According to back-trajectory calculations, the air mass that reached Etna originated mostly from the Atlantic Ocean passing through Spain and the Mediterranean Sea. Similar back-trajectory was observed for STRO14; however, STRO15 back-trajectory suggests an origin crossing above the Saharan desert. During the latter event, a significant aerosol surface area in the background was observed and could be explained by the presence of Saharan dust, in contrast to the other three flights.
2.6. Background and Plume Conditions

We measured the SO2 concentration over the Mediterranean in both the FT and BL outside of the volcanic plumes of Etna and Stromboli, and the values were ranging from 1.4 to 1.9 ppbV. From the air mass characterization upwind and downwind of Etna (being almost all the time in the FT) and Stromboli (in the BL), a background plume threshold value (PTV) of 2 ppbV of SO2 is used. This threshold value is used to exclude any contributions from ship emissions or other anthropogenic sources, and so on. Although the PTV could be considered relatively high when compared to anthropogenic emission levels (usually on average below 1 ppbV) or to other studies (Mauldin et al., 2003), these values were chosen to ensure that our data analysis focused only on measurements within the different volcanic plumes and to ensure that we do not consider contributions from other sources. The background (outside plume conditions) was also characterized in terms of particle concentrations, being defined as median of all the measures (defined in Table 2) when SO2 < 2 ppbV for Etna (i.e., 1.61 ppbV, 1,781 cm$^{-3}$, 3.2 cm$^{-3}$, and 0.85 × 10$^8$ cm$^{-3}$ for SO$_2$, $N_{2,5}$, $N_{250}$, and SA, respectively) and Stromboli (i.e., 1.26 ppbV, 2,100 cm$^{-3}$, 10.75 cm$^{-3}$, and 0.87 × 10$^8$ cm$^{-3}$ for SO$_2$, $N_{2,5}$, $N_{250}$, and SA, respectively). The SO$_2$, SA, and particle number concentrations were corrected considering that their backgrounds were subtracted from the plume concentrations in order to quantify the volcanic plume increment. Thus, all the analysis presented in the following sections is under plume conditions.

3. Results and Discussion

3.1. Plume Spatial Extent and Total Particle Number Concentrations

The spatial extent (vertical and horizontal) of the different volcanic plumes, represented by the SO$_2$, is displayed in Figure 1. Once the aircraft arrived at the volcano, a total vertical profile ranging from 0.2 up to 4 km for both plumes (Figure 1) was performed. The Etna plume appeared to be located at altitudes between 2 and 3.6 km, whereas the Stromboli plume was centered around 0.8 km during STRO14 and as low as 0.2 km.
during STRO15. We, therefore, assume that both volcanic plumes investigated around Stromboli were in the BL. Once the plume vertical distribution was located, a series of horizontal transects took place up to distances of 120 km.

For ETNA13 and ETNA14 and within the plume conditions, the vertical sounding shows that the air mass containing the plume moved toward downwind above 2 km in altitude confirming the presence of the plume in the FT. The median SO2 concentrations were 13.7 and 13.55 ppbV and reached maximum values at 92.3 and ~77 ppbV for ETNA13 and ETNA14, respectively. Since we did not sample the exact center of the plume at the vent where SO2 should peak, the maximum of SO2 was observed at altitudes above 3 and ~2.8 km at ~13 and 10 km away from the vent for ETNA 13 and 14, respectively (Figures 1a and 1b). Subsequently, SO2 concentrations decreased with distance from the vent and decreased in altitude while travelling downwind below 2.8 km (Figures 1a and 1b). The plume can still be observed above the altitude of 2.8 km with relatively high SO2 values above 40 ppbV at ~22-km distance from the vent (Figures 1a and 1b).

For STRO14 and STRO15 in BL, the SO2 median concentrations were 28.33 and 7.3 ppbV and reached the maxima of 83.3 and 78.3 ppbV, respectively (Figures 1c and 1d). The median concentration of SO2 observed in STRO15 is a factor of ~2 less than what was observed in both cases in the FT, whereas, in STRO14, the median was a factor of ~2 larger than in the FT. The differences in the median concentrations of the two BL flights can be explained by the relatively low wind speed measured in the case of STRO14 compared to the other flights (Table 3), resulting in the plume being less spread in both horizontal and vertical directions and more concentrated over a shorter range of distance (Figure 1). Since the aircraft missed the core of the plume at the vent, the maximum values were observed at distances of ~10 to 12 km from the vent similarly to Etna plumes and at heights of 860 and 120 m (Figures 1c and 1d), respectively, indicating a downward transport of the plume. The SO2 concentration decreases significantly (<15 ppbV; Figure 1d) with distance from the vent.

The concentration of SO2 measured during all flights is on the same order of magnitude as those observed in previously reported airborne measurement (larger than 10 ppbV) over the Miyaka volcano in the Pacific Ocean in Japan (Mauldin et al., 2003).

The maps of the aircraft trajectories for the four flights are shown in Figure 2, where the color bar represents the concentration of SO2, N10−250, and N250. In the FT, N10−250 increased along the plume with median concentrations of ~2,500 and 14,000 cm−3 for ETNA13 and ETNA14, respectively (Table 4). The maximum concentrations of N10−250 were measured to be 34,100 and 25,400 cm−3 at ~26- and ~47-km distance from the vent. Thus, the N10−250 maxima positions were at ~13 and 37 km farther than the areas where SO2 maxima were detected (Figures 2a, 2b, 2e, and 2f). The N10−250 increases, on average, from a few thousands at ~5 km to a few ten thousands at a distance of >25 km from the vent, for Etna flights (Figures 2e and 2f), and the rate of this increase roughly was ~1,500 cm−3·km−1. This suggests the occurrence of NPF and growth along the volcanic plume. For larger particles, the median of N250 were 18.8 and 11 cm−3 for ETNA14 and ETNA14, respectively (Table 4). During ETNA flights, the N250 was relatively high (~70 cm−3) close to the vent (~7 km) but decreased significantly (<20 cm−3) with the plume dilution (Figures 2i, 2j, and S4). This is opposite to what was measured for smaller particles (N10−250) where a higher concentration was detected along the volcanic plume in the FT especially in the diluted plume (>25 km) of both Etna flights (Figures 2e and 2f).

Considering that the volcanic emissions in ETNA13 and ETNA14 during the time of the campaign were passive, the presence of large particles can mainly be interpreted as the rapid growth of freshly nucleated particles. This hypothesis is confirmed by the correlation between N100, N250, and N10 (Figures S5a and S5c). We do not exclude that some fraction of these particles may also be due to the presence of very fine primary particles (accidental lithic) remobilized from the previous deposits within the conduit and volcanic crater walls, but we estimate that their contribution to the total aerosol concentration was minimal in our observation in the FT (Figures S5a and S5c).

For STRO14 and STRO15, N10−250 was observed with relatively large values (>104 cm−3) close to the vent at ~4 km and then decreased significantly with distance from the vent (Figures 2g and 2h), except the area where N10−250 peaks at ~12-km distance from the vent for STRO14 (Figure 2g). The N10−250 maxima were 18,809 and 19,700 cm−3 at ~12 and ~7 km for STRO14 and STRO15, respectively, where SO2 was also
observed to be relatively large (>60 ppbV). This is in contrast to what we observed for Etna plumes, where the N$_{10-250}$ maxima were located farther downwind (i.e., at 26- and 47-km distance from the vent). The median of N$_{10-250}$ concentrations were 2,470 and 1,300 cm$^{-3}$ for both Stromboli flights, that is, close to the N$_{10-250}$ median concentration measured within the ETNA13 but one order of magnitude less than the median N$_{10-250}$ observed for ETNA14.

For BL flights, the N$_{250}$ is observed to be high, peaking at 1,081 and 718.5 cm$^{-3}$ near the vent at ~7 and 5.2 km and continue to possess high values downwind along the plume dilution reaching 225 cm$^{-3}$ at 23-km distance from the vent (Figures 2c, 2d, 2k, and 2l). In general, the N$_{250}$ in the BL was observed to be

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**Figure 2.** Maps of the trajectory of the different flights in the free troposphere and boundary layer. The color-coded bars represent the corrected (background subtracted) observation of SO$_2$ (a–d), N$_{10-250}$ (e–h), and N$_{250}$ (i–l).
higher than in the FT; the medians are factor of 3 to 6 higher in the FT (Table 4), and the absolute values are two orders of magnitude higher close to the vent and all along the plume dilution (Figures 2i–2l). The presence of those large particles in the BL flights is due to the quick growth of newly formed particles at distances less than 10 km close to the vent (Figures S5b and S5d; blue points marked by a blue circle) and distant from the vent with a different growth rate (Figures S5b and S5d; light blue to red points marked by a red circle). Besides, close to the vent in STRO14, there is likely a contribution from very fine juvenile ash ejected during the short explosions at Stromboli (Figures S5b and S5d; blue points marked by a green circle), which can be a significant contribution to the total aerosol surface area. Moreover, in STRO15 and based on the backward trajectory analysis, there may also be a contribution from the Saharan dust particles that arrive from North Africa to the measured areas (Figure S2). Due to the lack of aerosol composition measurements, we are unable to accurately quantify these contributions. The higher concentrations of large particles during BL flights would eventually contribute to a larger aerosol surface area along the volcanic plumes compared to FT flights within Etna’s volcanic plumes.

In the following section, we will investigate the NPF and their potential growth processes occurring within these two different volcanic plumes.

### 3.2. Observation of NPF Within Different Volcanic Plumes

The total number concentrations of particles between 2.5 and 10 nm ($N_{2.5-10}$), the SA concentrations (defined in Table 2), and CS are illustrated in Figure 3. For FT flights, $N_{2.5-10}$ concentrations were observed above the detection limit throughout both plumes and up to ~45 km away from the vent. For both Etna flights, the $N_{2.5-10}$ is a factor of ~2 to 10 larger than the $N_{10-250}$ concentrations. This suggests that nucleation is taking place within the volcanic passive plume in the FT (Figures 2e, 2f, 3a, and 3b). Linked to this nucleation process occurring along the volcanic passive plume, a high concentration of SA ($>10^8$ cm$^{-3}$) is continuously observed, being produced from the oxidation of SO$_2$ (Figures 3e and 3f). The maximum of $N_{2.5-10}$ was observed where the SA concentration was greater than $2 \times 10^8$ cm$^{-3}$, and the CS was minimum (Figures 3a, 3b, 3e, 3f, 3i, and 3j).

During BL flights, NPF was also observed, but high concentrations of $N_{2.5-10}$ were mostly located close to the vent (<10 km). However, the median concentration of $N_{2.5-10}$ was up to two orders of magnitude less than in the FT flights. These show the evidence that new particles were formed within the different volcanic plumes close to the vent (~3 km) for both Etna and Stromboli volcanic plumes with different rates but also far from the vent (>35 km) along the dilution of the passive plume from Etna in the FT. The higher concentrations of large particles (>250 nm) in the BL result in a higher CS than in the FT (Figures 1i–1l, 2i–2l, S4, and S5).

### Table 4

<table>
<thead>
<tr>
<th>Variable</th>
<th>ETNA13</th>
<th>ETNA14</th>
<th>STRO14</th>
<th>STRO15</th>
</tr>
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<tbody>
<tr>
<td>$N_{2.5-10}$ (cm$^{-3}$)</td>
<td>[0.99 2.4] $\times 10^4$</td>
<td>[0.41 1.1] $\times 10^5$</td>
<td>[0.4 1.3] $\times 10^3$</td>
<td>[171 741]</td>
</tr>
<tr>
<td>$N_{10-250}$ (cm$^{-3}$)</td>
<td>[0.053 1.9] $\times 10^4$</td>
<td>[0.7 1.81] $\times 10^4$</td>
<td>[1.1 3.7] $\times 10^3$</td>
<td>[428 3.1 $\times 10^3$]</td>
</tr>
<tr>
<td>$N_{250}$ (cm$^{-3}$)</td>
<td>[11.2 27.6]</td>
<td>[5.4 16.8]</td>
<td>[16.6 150.5]</td>
<td>[2.5 72.5]</td>
</tr>
<tr>
<td>SA (cm$^{-3}$)</td>
<td>[0.85 3.9] $\times 10^8$</td>
<td>[3.35 3.94] $\times 10^8$</td>
<td>[2.34 2.94] $\times 10^8$</td>
<td>[3.35 $\times 10^8$]</td>
</tr>
<tr>
<td>SO$_2$ (ppbV)</td>
<td>[6.2 25]</td>
<td>[7.5 21.6]</td>
<td>[13 37.6]</td>
<td>[4.5 9.6]</td>
</tr>
<tr>
<td>CS (s$^{-1}$)</td>
<td>[0.5 3.1] $\times 10^{-4}$</td>
<td>[0.23] $\times 10^{-4}$</td>
<td>[0.65 9.8] $\times 10^{-3}$</td>
<td>[0.45 $\times 10^{-3}$]</td>
</tr>
</tbody>
</table>

Note. The 25th and 75th percentiles are in the first line in square brackets, and medians are in the second line.
the BL flights than in the FT flights, are likely to explain the weaker NPF events within the volcanic plumes in the BL (Tables 2 and 4; Figures 3 and S1).

Moreover, the growth behavior of the newly formed particles was distinct between ETNA and STRO plumes (Figures S4 and S5). The Correlations between newly formed particles and larger particles are observed along the volcanic plumes in both the FT and the BL (Figures S4 and S5). The growth rate, represented by the different slopes, varied according to the distance from the vent and the SO$_2$ abundance (Figures S4 and S5). The growth is observed to increase at areas closer to the vent (<20 km) than at farther areas than 20 km (Figures S4 and S5). Since SA was abundant during all flights (Figures 3e–3h), it likely played a

Figure 3. Observed N$_{2.5-10}$ (a–d), sulfuric acid (SA; e–h), and condensational sink (CS; i–l) along the flight trajectory within the plumes of Etna and Stromboli. Gaps visible in the different trajectories are attributed to instrumental data nonavailability.
key role in NPF and growth processes (CCN active size) within the volcanic plumes in FT and BL. It should be stressed that in the absence of the chemical analysis of the grown particles in our observation, we do not exclude the contribution of other condensable vapors to the growth of the freshly formed particles in the volcanic plumes. The growth of the newly formed particles to CCN active diameters illustrates that those particles within the volcanic plumes can contribute to cloud formation, thus impacting the weather and climate. Such an observation might be useful for further modeling studies to investigate the contribution of NPF to the CCN and their impact on climate and reduce the associated uncertainty.

The volcanic SA was also observed to be abundant in other volcanic plumes of Etna (Roberts et al., 2018) and in other locations, that is, Miyaka and Kilauea (Kroll et al., 2015; Mauldin et al., 2003). This abundance was found to be variable according to several factors related to SO2 concentrations and its oxidation rates and meteorological variables, that is, wind speed, temperature, and relative humidity (Kroll et al., 2015; Roberts et al., 2018). In comparison to other studies, the SA observed within ETNA and STRO flights were almost of the same order of magnitude to what has been reported in the Pacific BL volcanic plume from the Miyaka volcano (Mauldin et al., 2003; Weber et al., 2003). Conversely, the median of N_{2.5-10} measured in Stromboli plumes in MBL is almost one order of magnitude larger than the upper limit (100 cm^{-3}) reported in the MBL volcanic plume in the Pacific from Miyaka volcano for the similar size range (3–4 and 3–8 nm; Mauldin et al., 2003). The presence of large aerosol surface area may explain our observation of relatively low N_{3.8} in the BL, in comparison to N_{2.5-10} observed in the FT. Similarly, in the Pacific BL volcanic plume from Miyaka volcano (Mauldin et al., 2003), the presence of preexisting particles is given by the authors as an explanation of the low N_{3.4} and N_{5.8} concentrations.

The spatial distributions of the plumes were analyzed in more details, through plume latitudinal transects at different distances from the vent in both ETNA13 and STRO15 (Figure 4).

In each transect (highlighted in blue, red, green, and purple in Figures 4c and 4f), we determined the center of the plume by the peak of SO2 (not shown). The plume widths are of the order of 6–8 km for the two transects nearest to the volcanoes (respectively at ~7 and ~10 km from the vent) for both volcanoes (Figures 4a, 4b, 4d, and 4e). Further downwind, the STRO15 plume width increases to about 30 km at ~80-km distance from the vent (Figures 4g and 4h). For the ETNA13 and STRO15 flights, their corresponding N_{2.5-10} maxima were not observed at the center of the plume but at the edges of the plumes (Figures 4a and 4d). Conversely, N_{10} possesses the maxima exactly at the center of the plume in the FT (Figure 4b), whereas in the BL it was also shifted to the plume’s border (Figure 4e). These results support that the presence of large CS at the center of the plume is likely to explain the peaks shifting of N_{2.5-10} (and N_{10} in the BL), inhibiting the nucleation process (despite SA continuously produced by oxidation of SO2) compared to more favorable conditions at the plume periphery, where the CS is lower. The relationships between the particle concentrations in the smallest size bin and their gas-phase precursors, as a function of the distance from the volcanoes’ vents, are further investigated in the following section.

### 3.3. Derivation of NPF Parameterization

The data from the in situ measurements are used to derive a parameterization that can be useful to describe the rate of NPF as a function of SA concentration under natural conditions. One hypothesis in our derivation of the nucleation rate is that losses of newly formed particles due to coagulation are negligible compared to the strength of the nucleation rate. Thus, we believe that our calculation is the lower estimate of the nucleation rate. Nucleation rates were derived when the nucleation mode particle concentrations (N_{2.5}) was increasing with processing time (t) for several periods and locations, where NPF was observed to occur (Figures 5a and 5d; areas are highlighted by blue). The parameter t is the time needed for an air mass originating from the vent to reach the point where it was sampled by the aircraft. This time is estimated here by integrating the wind speed along the plume with the distance from the vent as follows:

$$
t = \frac{x_0}{\bar{V}_x} + \sum_{i=x_0+1}^{L} \frac{\Delta x}{V(i)}
$$

x_0 is the distance between the closest point from the flight trajectory inside the volcanic plume and the vent, \(\bar{V}_x\) is the mean wind speed of all the trajectory points, L is the farthest point from the vent of flight trajectory.
within the plume, $\Delta x$ is the distance traveled by the aircraft from the source point (vent), and $V(i)$ is the corresponding wind speed of each point of the flight trajectory. Since the nucleation events were found to be more pronounced in the FT than in the BL, a nucleation rate could only be calculated along the plumes in the FT, and hence, the parameterization is solely based on Etna emissions. We plot the particle concentration increase as a function of the processing time by taking into account the dilution of the plume with transport within the plume. It is important to note that the volcanic passive plume at Etna contains a low concentration of CO (upper limit was 110 ppbV), a typical gas used as a dilution factor ($d_i$). Therefore, during this study, SO$_2$ is used as a dilution factor, while its concentration is strongly enhanced in the plume and has a typical tropospheric average lifetime of 1–2 days (Beirle et al., 2014), and thus, is partially consumed during the plume evolution. A dilution factor, defined as the SO$_2$ concentration normalized by its maximum value for each flight, was hence applied to the particle concentration to calculate normalized particle number concentrations. Figures 5b, 5c, 5e, and 5f show the normalized $N_{2.5}$ concentrations as a function of $t$ to evaluate the correlation between these two variables. For ETNA13, we divided the flight into two periods: (a) from 10:40 to 10:55 when the aircraft only crossed the plume at different distances from the vent, and (b) from 11:00 to 11:15 when the aircraft was flying within the center of the plume. For ETNA14, we furthermore chose two periods where the NPF events were observed to occur at two different altitudes in the FT: (c) from 14:28 to 14:32 at ~2.8 km and (d) from

---

**Figure 4.** $N_{2.5-10}$ and $N_{10}$ as a function of distance from the center of the plume for ETNA13 (a and b) and STRO15 (d, e, g, and h). The plots (g) and (h) are similar to (d) and (e) but with a larger horizontal extent from the center of the plume in STRO15. The considered transects are highlighted by the corresponding colors in the right panel (c for Etna and f for Stromboli).
14:34 to 14:38 at ~3.3 km. In period (a), we calculated the mean value of each particle peak, each corresponding to a single processing time (Figure 5b). The fit of those points against $t$ was linear (Figure 5b), yielding a slope of $11 \text{ cm}^{-3}\text{s}^{-1}$. In period (b), the function between normalized $N_{2.5}$ concentrations and the $t$ was exponential (Figure 5c). The rate of NPF ($J_{2.5}$) for each nucleation event is then the derivative of each of the abovementioned regressions for the two periods for flight ETNA13.

Similar to period (b), the regression for the two chosen periods for flight ETNA14 was found to be exponential as well (Figures 5e and 5f).

Table 5 shows all the derived relationships between the normalized $N_{2.5}$ versus $t$ with their fitting parameters $A$ and $B$ to estimate the nucleation rate $J_{2.5}$. The relationship has either a linear form as in period (a; illustrated in Figure 5b) or an exponential form as in periods (b, c, and d; illustrated in Figures 5c, 5e, and 5f):

$$J_{2.5} = d(N_{2.5}/dt) = A e^{Bt} \quad (1)$$

Figure 6 shows all estimated $J_{2.5}$ for all Etna points in the FT versus SA together with their averages (black dots) every $0.25 \times 10^8 \text{ cm}^{-3}$. We derive the parameterization of NPF rate $J_{2.5}$ by fitting the averages of all
The black dots in panel (b) are the mean values of the normalized N2.5 and corresponding processing time shown in Figures 5b, 5c, 5e, and 5f with their fitting parameters for the different periods in the case of ETNA13 and ETNA14 in the free troposphere. The calculated regression of N2.5 versus sulfuric acid (SA) for the different periods in the free troposphere is shown in Table 5.

Table 5
Summary of the Derivative of the Correlation Between the Normalized N2.5 and Corresponding Processing Time Shown in Figures 5b, 5c, 5e, and 5f With Their Fitting Parameters for the Different Periods in the Case of ETNA13 and ETNA14 in the Free Troposphere

<table>
<thead>
<tr>
<th>Period</th>
<th>(a)</th>
<th>(b)</th>
<th>(c)</th>
<th>(d)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regression of N2.5/dt versus t</td>
<td>10:40 to 10:55</td>
<td>11:00 to 11:15</td>
<td>14:28 to 14:32</td>
<td>14:34 to 14:38</td>
</tr>
<tr>
<td>J2.5 = d(N2.5/dt)/dt</td>
<td>11 × t + 17,000</td>
<td>4.505 × 10^3 • 9.5 × 10^-15s⁻¹</td>
<td>3.280 × 10^3 • 7.1 × 10^-15s⁻¹</td>
<td>3.799 × 10^3 • 8.4 × 10^-15s⁻¹</td>
</tr>
<tr>
<td>R²</td>
<td>0.99</td>
<td>0.8814</td>
<td>0.9123</td>
<td>0.8124</td>
</tr>
<tr>
<td>Adjusted R²</td>
<td>0.98</td>
<td>0.881</td>
<td>0.9119</td>
<td>0.8116</td>
</tr>
</tbody>
</table>

Note. The confidence bounds for all regressions were 95%.

Figure 6. The calculated J2.5 versus sulfuric acid (SA) for the different periods of all ETNA flights in the free troposphere. The black dots and the associated error bars on the panel (a) are the mean and the standard deviation (STD) of the J2.5 obtained within 0.25 × 10^8 cm⁻³ SA equal bins. The black dots in panel (b) are the mean J2.5 as a function of SA for the specific SA range defined by the light red shaded area on the panel (a) when SA is smaller than 4.6 × 10^8 cm⁻³. In the panel (b), the correlation between J2.5 and SA is a power fit (red line), which represents the parameterization of the new particle formation within the volcanic plume J2.5 = 1.844 × 10^-3 • [SA]^{1.12}, with R-square = 0.85.

estimated J2.5 to the corresponding SA using the bi-squares method, with 95% confidence bounds, of the simple power model that has the form

\[ J_{2.5} = K[SA]^P \]  \hspace{1cm} (2)

where the prefactor K and exponent P are the fitting parameters of the power function and, estimated to be 1.844 × 10⁻⁸ s⁻¹ and 1.12 (95% confidence interval 0.76 and 1.47), respectively. The exponent P, found in the current study, is closer to 1 (associated with activation-type nucleation; Kulmala et al., 2006) rather than kinetic-type nucleation (2; McMurry & Friedlander, 1979), in agreement with what has previously been hypothesized (Kuang et al., 2008; Sihto et al., 2006). According to our derivation and by substituting the values of K, P, and the medians of SA from Table 4 for ETNA13 and ETNA14, the average J2.5 is 68.6 ± 39.9 and 59.23 ± 29.8 cm⁻³ s⁻¹, respectively, and equal to 63.23 ± 34.8 cm⁻³ s⁻¹ for all ETNA points.

In our derivation, the coagulation process was neglected in comparison to the strength of the nucleation process; therefore, the J2.5 values derived here are considered as the lower limits of nucleation. Based on Quantum Chemistry-normalized Classical Nucleation Theory (QC-CNT) and CLOUD measurements presented in Duplissy et al. (2016), the nucleation of new particles is minimized when SA was below 10^8 cm⁻³ at temperatures above 10 °C. This indicates that the SA background (up to 0.85 × 10^8 cm⁻³), which was subtracted from the data, would not have a significant impact on our derivation of the NPF rate. Moreover, this parameterization is valid when SA is less than 4.6 × 10^8 cm⁻³ (Figure 6a). For values of SA larger than 4.6 × 10^8, J2.5 has been observed to significantly decrease with increasing SA (Figure 6a) due to the large CS estimated at distances close to the vent (less than 5 km) or in the center of the plume, where growth is observed to be stronger than NPF. For the SA values greater than 4.6 × 10^8 cm⁻³, we believe that the approximation of negligible coagulation is no longer valid. In comparison to more cleaner environments, our lower limits estimations of J2.5 are one to almost two orders of magnitude higher than what has been previously measured in Hyytiälä (Finland) during the QUEST2 campaign in 2003 by Sihto et al. (2006) for J3. The formation rate J2.5 is a factor of ~2.5 to 4 higher than the upper limit of J1 found for the same campaign (Kuang et al., 2008; Sihto et al., 2006). These suggest a quicker occurrence of NPF within the harsh environment of the volcanic passive plume in comparison to what
has been found in other cleaner environments (Sihto et al., 2006). In comparison to the controlled CLOUD3 and CLOUD5 experiments of SA-water binary particle formation for the same range of SA, our estimation of \( J_{2.5} \) is comparable or an order of magnitude higher than the \( J \) that resulted from the exposure to different beams in the CLOUD chamber (Figure 9 in Duplissy et al., 2016). This may indicate that condensable vapors other than SA could be contributing to the NPF events. Indeed, Kirkby et al. (2011) showed that ~100 pptV of ammonia may increase nucleation up to a factor of 1,000 more than what binary SA-water nucleation can produce. However, in the absence of chemical characterization, our observations were not able to confirm the contribution of the other species than SA to the NPF process. Our estimation of \( J_{2.5} \) is found to be more than one order of magnitude higher than what was estimated, on average, within the aged volcanic plume that reached the puy de Dôme station, Massif Central (France; \( 4.76 \pm 2.63 \text{ cm}^{-3} \text{ s}^{-1} \)) in May 2010 from the Eyjafjallajökull eruption event (Boulon et al., 2011). This is expected since the plume in that study traveled several thousands of kilometers before reaching the station, whereas our measurements are occurring directly within the passive plume. This indicates how efficient the volcanic passive plume can be especially in the FT where NPF is favored. Substituting the average estimated value of SA (3.67 \( \pm 0.78 \times 10^7 \) molecules/cm\(^3\)) from Boulon et al. (2011) in our parameterization formula (equation (2)) would give an NPF rate of 5.02 cm\(^{-3}\) s\(^{-1}\), which is close to the average nucleation rate actually calculated from the Eyjafjallajökull plume (Boulon et al., 2011). This indicates that our parameterization would be able to reproduce the average nucleation rate \( J_5 \) estimated for the volcanic plumes even with SA less than 10\(^8\) cm\(^{-3}\) at locations far from the eruptive point. Yet the binary homogeneous H\(_2\)SO\(_4\)-H\(_2\)O nucleation scheme (Kulmala et al., 1998) and activation nucleation (Sihto et al., 2006) have been used in a previous modeling study that investigated the impact of volcanic aerosols on climate (Schmidt et al., 2012). The binary homogeneous H\(_2\)SO\(_4\)-H\(_2\)O nucleation scheme was found to underestimate the climatic impact of freshly formed particles in the volcanic degassing plumes (Boulon et al., 2011; Schmidt et al., 2012). Therefore, our current analysis together with the parameterization can effectively contribute to better understand and quantify the climatic impacts of aerosol nucleation and their evolution within volcanic plumes near the volcanic source and in the diluted volcanic plumes.

### 4. Conclusions

This study presents a comprehensive investigation of NPF and growth within volcanic plumes located in the FT and in the BL. This was conducted by performing airborne in situ measurements within the plumes of Etna and Stromboli, Italy.

We evidenced the occurrence of NPF and growth of these newly formed particles within the different plumes from Etna in the FT and Stromboli in the BL. In the FT, the NPF events were measured in the volcanic passive plume near the vent with a rapid growth rate and continue to occur efficiently along the plume at distances farther than 35 km. The rapid growth of the newly formed particles was observed close to the vent results in a relatively high number concentration of large particles (N\(_{250} > 55 \text{ cm}^{-3}\)) and hence a relatively significant CS (up to \( 10^{-2} \text{ s}^{-1} \)) in absolute value. However, the concentration of these large particles is diluted with distance, and therefore, the threshold ratio between the condensable gases and the condensational sink is overcome by the presence of sufficient SA from SO\(_2\) oxidation to allow for further nucleation events. In the BL, the NPF events were also observed close to the vent of the volcano and with smaller concentrations of ultrafine particles than in the FT. The SO\(_2\) fluxes at Stromboli were reported to be weak (0.15–0.6 kt/day; Burton et al., 2008), but they remain comparable with those emitted at Etna during passive degassing (0.6–2 kt/day (Aiuppa et al., 2008; Roberts et al., 2018). This has been supported by our observations, where we found that SO\(_2\) and SA concentrations were relatively comparable, and the differences in their values in different volcanic plumes of Etna (being passive) and Stromboli are small. Thus, these small differences in SO\(_2\) and SA concentrations between Etna and Stromboli do not explain solely the NPF being more dominant in Etna (in the FT) than in Stromboli (in the BL). Therefore, the occurrence of the NPF events in the different volcanic plumes seems to be largely influenced by the presence of large particles leading to large CS at the very proximity to the vent. Thus, the weaker NPF events in the BL is a result of a larger aerosol surface along the plume, where CS was up to two orders of magnitude higher than in the FT and with temperatures reaching 23 and 30 °C in both BL flights, which is up to 20 °C higher than in the FT. This detailed analysis of the growth of freshly nucleated
particles to the CCN sizes (Figures S4 and S5) is beneficial for further modeling studies to investigate the contribution of NPF to the CCN and their impact on climate.

To the authors’ knowledge, this is the first dedicated study that addresses the relationship between the newly formed nanoparticles and their gas-phase precursors in the vicinity of different volcanic plumes over Etna and Stromboli. The in situ airborne measurements performed as part of this study within the ETNA passive plume were used to derive NPF rate parameterizations $J_{NPF}$ that can eventually be incorporated into the models. The NPF rate was an exponential function of the processing time in most of the observed individual nucleation events along the flight trajectories and implicitly includes the information about the plume’s dilution. The NPF rate parameterization was a power law function of $SA$, with an exponent value of 1.12, which is accepted within the range of what has previously been reported (Kuang et al., 2008; Sihto et al., 2006). The latter exponent value implies that the nucleation within the studied plumes is a natural process that is a mixture of both activation (Kulmala et al., 2006) and kinetic (McMurry & Friedlander, 1979) nucleation modes, but more close to the activation nucleation mode. We believe that our calculation is the lower estimate of the nucleation rate within the volcanic plume since the losses due to coagulation are neglected compared to the strength of the nucleation rate. This new parameterization has a simple formula and is able to reproduce the same average nucleation rate for the volcanic plumes observed in locations thousands of kilometers distant from the erupted event (Boulon et al., 2011). Therefore, this parametrization of particle formation rate, based on actual measurements, is a more representative of the nucleation process that occurred under largely uncharacterized volcanic degassing plumes conditions. The new parametrization should further be tested in mesoscale models coupled with chemistry transport scheme and compared with preexisting parameterizations for NPF within volcanic plumes. It should be noted that although SA is the key factor for the NPF events within the different volcanic plumes, we cannot exclude that condensable vapors other than SA, for example, halogens and organic vapors, could be participating to the nucleation and growth processes in the FT and BL, and the latter should be investigated by deeper chemical characterization in future studies. Finally, this study contributes to better understand and quantify the natural process of the gas to particle conversion within volcanic plumes and how this process with the resulted aerosol concentrations evolves temporally and spatially in the atmosphere aiming to reduce the uncertainty of the aerosol’s impact on climate.

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References


