Chinese SO$_2$ pollution over Europe – Part 2: Simulation of aerosol and cloud condensation nuclei formation

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Received: 4 December 2008 – Accepted: 15 December 2008 – Published: 28 January 2009

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Published by Copernicus Publications on behalf of the European Geosciences Union.
Abstract

We report on sulfur dioxide (SO$_2$) induced formation of aerosols and cloud condensation nuclei in an SO$_2$ rich aged (9 days) pollution plume of Chinese origin, which we have detected at 5–7 km altitude during a research aircraft mission over the East Atlantic off the West coast of Ireland. Building on our measurements of SO$_2$ and other trace gases along with plume trajectory simulations, we have performed model simulations of SO$_2$ induced formation of gaseous sulfuric acid (GSA, H$_2$SO$_4$) followed by GSA induced formation and growth of aerosol particles. We find that efficient photochemical SO$_2$ conversion to GSA took place in the plume followed by efficient formation and growth of H$_2$SO$_4$-H$_2$O aerosol particles. Most particles reached sufficiently large sizes to act as cloud condensation nuclei whenever water vapor supersaturation exceeded 0.1–0.2%. As a consequence, smaller but more numerous cloud droplets are formed, which tend to increase the cloud albedo and to decrease the rainout efficiency. The detected plume represents an interesting example of the environmental impact of long range transport of fossil fuel combustion generated SO$_2$.

1 Introduction

Fossil fuel combustion represents the most important source of atmospheric sulfur dioxide (SO$_2$), a major air pollutant. Presently most atmospheric SO$_2$ is released in Europe and China from combustion of relatively sulfur rich coal (Lelieveld et al., 2001). Sulfur dioxide impacts the environment in several ways. It is toxic and after inhalation may cause severe adverse health effects (e.g. Sunyer et al., 2003; Venners et al., 2003; Longo et al., 2008). Moreover, it contributes to atmospheric acidity and acid-induced corrosion (e.g. Rodhe et al., 2002; Huang et al., 2008). In the atmosphere, SO$_2$ undergoes conversion to particulate sulfate and OH-induced gas-phase conversion to gaseous sulfuric acid (GSA, H$_2$SO$_4$), (e.g. Menon and Saxena, 1998; Reiner and Arnold, 1993, 1994; Fiedler et al., 2005). At low temperatures typical of the middle
and upper troposphere, GSA may undergo binary (H$_2$SO$_4$-H$_2$O) nucleation leading to new aerosol particles. These grow by condensation and coagulation and ultimately may become CCN (Seinfeld and Pandis, 2006). Consequently, atmospheric SO$_2$ promotes the formation of sulfuric acid-water aerosol particles which impact the radiation balance by absorbing and scattering sunlight and by prolonging the lifetime of clouds (e.g. Ramanathan et al., 2001; Harshvardhan et al., 2002; Garrett et al., 2002; Andreae et al., 2005).

After SO$_2$ release from ground level combustion sources substantial amounts of SO$_2$ may be transported to the middle and upper troposphere and thereby escape deposition at the surface and are less affected by removal via cloud processes. In the middle and upper troposphere, SO$_2$ is removed preferably by OH induced gas-phase conversion to GSA which leads to a SO$_2$ lifetime of about 7–14 days depending on the efficiency of photochemical OH formation (e.g. Finlayson-Pitts and Pitts, 2000). Hence, the SO$_2$ lifetime is sufficiently long to allow SO$_2$ long-range transport. Therefore, SO$_2$ may impact aerosols and clouds thousands of kilometers away from its emission source. An interesting question is to what extent the important SO$_2$ source China impacts regional aerosols and clouds. Here we report on model investigations of SO$_2$ induced formation of GSA, aerosols and CCN in a major SO$_2$ rich pollution plume of Chinese origin, which we have detected over the East Atlantic (Fiedler et al., 2008).

The model simulations are performed with an atmospheric chemistry and aerosol dynamics model AEROFOR (Pirjola, 1999; Pirjola and Kulmala, 2001).

Previous comparable AEROFOR model examples have been published e.g. by Pirjola et al. (1999) and by Laaksonen et al. (2000).

This paper consists of the following sections: Sect. 2 presents a short review of the SO$_2$ measurements off the Irish west coast. Section 3 describes the AEROFOR model simulation inputs, the model results and discusses the sensitivity of the results to several input parameters. Section 4 interprets the model simulations further concerning the SO$_2$ impact on CCN formation.
2 Measurements

The measurements are described in detail in the accompanying paper "Chinese SO$_2$ pollution over Europe – Part 1" (Fiedler et al., 2008). The measurement flight started from Brest in Northern France on 3 May 2006 and took place over the Atlantic Ocean south and west of Ireland. The SO$_2$ measurement method was chemical ionization mass spectrometry (CIMS) using a permanent calibration with isotopically labeled SO$_2$. The objective was to probe an Asian pollution plume, which had been predicted before by the lagrangian particle dispersion model FLEXPART (Stohl et al., 2002, 2005). Between 10:00 and 11:00 UTC several pollution plumes were detected as particularly indicated by very markedly elevated SO$_2$. A comprehensive description of the measurements especially in the light of other simultaneously measured trace gases can be found in Fiedler et al. (2008).

Figure 1 shows the vertical distribution of the SO$_2$ mole fraction measured during that flight. Apart from a SO$_2$ pollution layer of American origin at approximately 1.5 km altitude, a deep layer of SO$_2$ has been detected in the altitude range from 5 to 7.5 km. The atmospheric SO$_2$ background in this layer is about 150 pmol/mol with a single peak of 900 pmol/mol which is discussed in detail in the accompanying paper Fiedler et al. (2008).

FLEXPART particle dispersion model analyses identified a region in North East China as the source region of the SO$_2$ emissions.

3 Aerosol model simulations

3.1 Initialization of the model

The air mass with the highest SO$_2$ mole fraction (around 10:40 UTC, e.g. in Fig. 3 of the paper Fiedler et al., 2008) was investigated using simulations by the atmospheric chemistry and aerosol dynamics model AEROFOR along the trajectory. This model
has previously been described (e.g. Pirjola, 1999; Pirjola and Kulmala, 2001). AERO-
FOR is a sectional box model which treats aerosol formation by homogeneous binary
nucleation of H$_2$SO$_4$ and H$_2$O (Vehkamäki et al., 2002) as well as aerosol growth by
H$_2$SO$_4$-H$_2$O condensation (Fuchs and Sutugin, 1971) and aerosol coagulation (Fuchs,
1964). In this work a Lagrangian approach is used. From a prescribed SO$_2$ and OH
concentration the H$_2$SO$_4$ concentration is calculated. Further model input needed is
an initial particle concentration, pressure, relative humidity and temperature along the
trajectory. The model then delivers the homogeneous binary nucleation rate $J_{\text{honu}}$
condensation sink $CS$, which is principally the inverse H$_2$SO$_4$ lifetime, and particle number
size distributions between 0.86 nm and 1 µm. In this work we used 54 size bins.

The particle concentration calculation in the model moreover builds on a bimodal
initial particle size distribution, possessing the lognormal parameters

$$N_1 = 200 \text{ cm}^{-3}$$  \hspace{1cm} (1)
$$d_1 = 130 \text{ nm}$$  \hspace{1cm} (2)
$$\sigma_1 = 1.45$$  \hspace{1cm} (3)

and

$$N_2 = 50 \text{ cm}^{-3}$$  \hspace{1cm} (4)
$$d_2 = 250 \text{ nm}$$  \hspace{1cm} (5)
$$\sigma_2 = 1.9$$  \hspace{1cm} (6)

with $N_1$ the initial particle number concentration in mode 1, $d_1$ the geometric mean
diameter of that mode and $\sigma_1$ the geometric standard deviation of the lognormal distri-
bution, $N_2$, $d_2$ and $\sigma_2$ respectively in mode 2. Here, four different scenarios (A, B, C,
D) will be investigated for the highest SO$_2$ peak. Since no measurements existed for
the preexisting particle size distribution, the initial particle concentrations are varied by
multiplying the above values by 0, 1, 2 and 4, resulting in total initial particle concen-
trations of 0, 250, 500 and 1000 cm$^{-3}$.  

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The simulation time is 8.5 days starting at 00:00 UTC on 25th of April 2006, so ending around noon on the 3rd of May 2006, which was the measurement flight day and time. The initial \(\text{SO}_2\) concentration of \(4.4 \times 10^{10}\ \text{cm}^{-3}\) was chosen in such a way that the final modeled \(\text{SO}_2\) concentration (\(1.3 \times 10^{10}\ \text{cm}^{-3}\), which corresponds at that temperature and pressure level to 900 pmol/mol) matches the measured \(\text{SO}_2\). At the beginning of the simulation the air parcel under consideration was lifted to 8 km (\(\approx 335\ \text{hPa}\)). Table 1 summarizes the model input data of the four model cases.

For the OH calculation a clear sky assumption was made. This is in accordance with a rough satellite cloud top temperature data analysis. Latitude, longitude and time in UTC have been taken into account as well as the length of the day. Typical maximum OH concentrations were adopted from Logan et al. (1981). In Fig. 2 6 h mean values of temperature and relative humidity, RH, calculated from the water vapor concentration along the trajectory are plotted. The temperature stays rather constant around 240 K the whole time, relative humidity varies between 5% and 68% with two maxima at the beginning and on day 6 of the simulation. We also calculated the saturation vapor pressure over ice based on the Magnus equation (Pruppacher and Klett, 2000) and the relative humidity over ice, RHice, also shown in Fig. 2. Since for temperatures below 0°C the water vapor saturation pressure over liquid water is always higher than over ice, RHice is higher than RH for the same atmospheric water vapor concentration. However, water vapor supersaturation does not occur in the 8.5 simulation days.

3.2 Model results

Figures 3–8 describe the AEROFOR model results. All times are in UTC. Figures 3 and 4 show the model results for model scenario A with an initial particle concentration of 0 cm\(^{-3}\). Plotted versus time are: \(\text{SO}_2\) concentration, GSA and prescribed OH, rate \(J\) of \(\text{H}_2\text{SO}_4\cdot\text{H}_2\text{O}\) nucleation, and number concentrations \(N_d\) of aerosol particles with diameters larger than \(d\) nm. The OH (Fig. 3, upper panel) undergoes a semi-sinusoidal diurnal variation with maximum local noontime OH concentrations of about 2–4\(\times\)10\(^6\) cm\(^{-3}\) depending mostly on geographic latitude. The OH converts \(\text{SO}_2\) to GSA and causes,
within the 8.5 days of simulation, SO₂ to decrease from $4.4 \times 10^{10} \text{cm}^{-3}$ (corresponding to a mole fraction of almost 3700 pmol/mol) to $1.3 \times 10^{10} \text{cm}^{-3}$ (corresponding to the measured maximum mole fraction of around 900 pmol/mol). In comparison, the FLEXPART particle dispersion model predicted a SO₂ mole fraction of 2500–3500 pmol/mol at the location and time of our measurement (paper 1, Fig. 9, Fiedler et al., 2008).

Considering that FLEXPART treats SO₂ as an inert tracer, this compares well to the assumption of 3700 pmol/mol as initial SO₂ mole fraction input for the AEROFOR model.

After 8.5 days, about 70% of the initially available SO₂ has undergone photochemical conversion to GSA, which in turn has been incorporated into aerosol particles. The H₂SO₄ concentration (Fig. 3, upper panel) shows a diurnal variation, following the diurnal variation of OH. Right in the beginning, H₂SO₄ local noontime concentrations of $3 \times 10^7 \text{cm}^{-3}$ are reached. The maximum varies slightly each day between $1 \times 10^7$ and $3 \times 10^7 \text{cm}^{-3}$. The homogeneous nucleation rate $J_{\text{honu}}$ (Fig. 3, lower panel) shows two strong maxima around noon on day 0 and day 1 (7000 and 20 000 cm$^{-3}$ s$^{-1}$) and three smaller peaks on day 2 and 6, which corresponds to the local minima in temperature accompanied by local maxima in relative humidity. Low temperatures and high relative humidities favor new particle formation. The condensation sink CS (Fig. 4, lower panel) starts at 0 s$^{-1}$ as no initial particles exist, but increases immediately to 0.01 s$^{-1}$ simultaneously to the occurring nucleation. During nighttime the CS decreases caused by the decrease in the total particle surface, which results from coagulation and growth of the existing particles. The surface to volume ratio decreases with increasing radius of the particles. So, if small particles coagulate to form bigger ones, the total aerosol surface will decrease. This means that the total aerosol surface available for gaseous sulfuric acid scavenging decreases during night, if no new particles are formed. After the new nucleation on simulation day 1 the CS reaches its maximum of 0.013 s$^{-1}$.

Eventually, this results in the particle concentrations depicted in Fig. 4, upper panel. A strong increase in the total particle concentration (up to $6 \times 10^6 \text{cm}^{-3}$), caused by the two nucleation bursts on the first and second day of the simulation, is followed by a slow decrease of the total particles to a final value of $\approx 1600 \text{cm}^{-3}$. Condensational
and coagulational growth of the freshly formed particles forms particles with diameters larger than 30 nm already on the first day. Particles of these size classes may act as cloud condensation nuclei (CCN) and are therefore available for cloud formation.

Similar graphs for model scenario B with an initial particle concentration of 250 cm$^{-3}$ are not shown as no major changes occurred compared to scenario A. In model case B, H$_2$SO$_4$ concentration and homogeneous nucleation rate show nearly the same behavior as in model case A. The CS starts at 0.002 s$^{-1}$ instead of 0 s$^{-1}$, because of the initial particles that are available for condensation already when the simulation starts. However, the CS maximum value is 0.013 s$^{-1}$ as in scenario A. Freshly nucleated particles are formed in the same amount as in scenario A, so there is obviously enough H$_2$SO$_4$ available for both, growth of the initial particles and nucleation of new ones. The formation of CCN after 1 day is still enhanced and a final total particle concentration of 1300 cm$^{-3}$ at the measurement site can be expected.

The plots for model scenario C (initial particle concentration 500 cm$^{-3}$) are shown in Figs. 5 and 6. The higher initial particle concentration again has no substantial influence on the H$_2$SO$_4$ concentration and the nucleation rate, but the higher CS (starting at 0.005 s$^{-1}$ and 0.0135 s$^{-1}$ in maximum) slows down the growth of the particles particularly in the size classes N50, N100 and N200. The growth starts one day later than in scenario A and the total increase in the number concentration of CCN is less developed. Nevertheless the final concentration of CCN reaches in the sum of initial and freshly formed particles 1200 cm$^{-3}$.

Model scenario D, depicted in Figs. 7 and 8, eventually starts with the assumption of 1000 cm$^{-3}$ as initial particle concentration, which is almost the final value in scenario C. The H$_2$SO$_4$ concentration development stays nearly the same, but the nucleation rate J$_{honu}$ is slightly lowered. The condensation sink is further increasing and is all the time above 0.01 s$^{-1}$ with a maximum of 0.0143 s$^{-1}$. The increase in the total particle concentration on the first two simulation days is still appreciable, but the growth in all size classes is now markedly reduced. So the H$_2$SO$_4$ concentration seems still to be high enough for new particle formation, but a large amount of H$_2$SO$_4$ will be
already consumed by condensation onto preexisting particles, so that not much H₂SO₄ is left for condensation on newly formed particles. The final concentration of CCN is approximately 700 cm⁻³.

Typical previously measured particle concentrations for background aerosol are around 500 cm⁻³ and in very polluted cases also several 1000 cm⁻³ can be reached (e.g., Minikin et al., 2003). So at least model scenario C seems to be quite realistic.

With even higher initial particle concentrations (e.g. 2000 cm⁻³) nucleation would not occur at all due to a complete removal of the condensable gases by condensation onto the preexisting aerosol.

Figure 9 gives an overview of the modeled final particle concentrations N₃₀, N₁₀₀ and N₂₀₀ (after 8.5 days). For comparison, the corresponding initial particle concentrations are also given. The right panel of Fig. 9 shows model predictions with H₂SO₄ formation switched off. Here new aerosol formation does not occur and preexisting particles grow only by coagulation. For increasing preexisting particle concentrations coagulation becomes more efficient resulting in a decrease of N₃₀ and N₁₀₀ with respect to the corresponding concentrations at time 0. In contrast, N₂₀₀ increases but the increase is only very small. When photochemical SO₂ conversion to H₂SO₄ is switched on, the situation changes drastically (Fig. 9, left panel). Now, for each of the 4 scenarios A, B, C and D, the final concentrations of N₃₀, N₁₀₀ and N₂₀₀ are much larger. Hence, the AEROFOR model simulations indicate very substantial SO₂-mediated growth of preexisting and newly formed aerosol particles.

3.3 Sensitivity analyses

The simulation of the model case C was repeated to test the sensitivity of the concentration of CCN after 8.5 days against the initial SO₂ concentration, OH concentration, nucleation rate and initial particle size distribution.

When multiplying the nucleation rate by a factor of 100 and keeping the OH and SO₂ concentrations as in case C the final N₅₀, N₁₀₀ and N₂₀₀ increased only 4%, 7% and 2%, respectively. The total particle concentration became threefold but the major part
of the particles remained smaller than 4 nm in diameter size. Thus we can conclude that the CCN production is much more sensitive to particle growth, i.e. condensable vapor concentration, than to the actual nucleation rate.

Since the GSA formation rate is proportional to OH×SO$_2$, the simulations were repeated by multiplying first the OH by a factor of 2 (SO$_2$ as in case C) and then the initial SO$_2$ by a factor of 2 (OH as in case C). The final SO$_2$ concentration was $4.5 \times 10^9$ cm$^{-3}$ ($\approx 305$ pmol/mol) and $2.8 \times 10^{10}$ cm$^{-3}$ ($\approx 1900$ pmol/mol), respectively. The increases in N50, N100 and N200 were 16%, 41% and 35%, and 25%, 67% and 92%, respectively.

The other factor which affects the growth of nucleated particles is the condensation sink CS of the preexisting particles. The CS is mainly determined by the number concentration and the size distribution of the preexistent particles. Therefore, a CS sensitivity study is already included in the 4 simulation cases. The CS of case B (initial particle concentration of 250 cm$^{-3}$) e.g. corresponds to the CS of case C (initial particle concentration of 500 cm$^{-3}$) if the diameter of the particles in mode 1 of case C (N1=400 cm$^{-3}$) is changed to 62 nm and the diameter in mode 2 of case C (N2=100 cm$^{-3}$) is changed to 124 nm (see also Table 1 for comparison). On the other hand, if the diameters of the particles in mode 1 and 2 of case D are changed to 100 nm and 200 nm, respectively, the CS of case B is the same CS as in case D.

4 Implications for CCN

Our above findings have interesting implications for CCN and cloud droplet formation. In the atmosphere, water vapor supersaturation (WSS) occasionally occurs and aerosol particles with sufficiently large diameters and hygroscopicity may act as CCN and induce water cloud droplet formation. Mostly, WSS is induced by adiabatic cooling or mixing of air masses with different temperatures and humidities (e.g. Seinfeld and Pandis, 2006; Gettelman et al., 2006). Typical atmospheric water vapor supersaturation rarely surpasses 2%, the median of observed supersaturation even is 0.1% (Pruppacher and Klett, 2000).
Sulfuric acid is very hygroscopic and therefore \( \text{H}_2\text{SO}_4 \) represents an ideal CCN material. \( \text{H}_2\text{SO}_4\)-\( \text{H}_2\text{O} \) aerosols are liquid in the temperature and RH conditions encountered in the troposphere. While \( \text{H}_2\text{SO}_4 \) contained in an aerosol droplet may promote water vapor uptake, it may at least initially hinder droplet freezing depending on the \( \text{H}_2\text{SO}_4 \) fraction of the droplet mass (Carleton et al., 1997; Ettner et al., 2004). As water vapor uptake proceeds and the droplet size has increased sufficiently and the \( \text{H}_2\text{SO}_4 \) mass fraction has decreased sufficiently, the droplet may ultimately freeze even quasi-homogeneously, depending on RH. However, a pure water droplet only freezes at Temperatures \(<235\text{ K} \text{ and } \text{RHi}>145\%\). Those values have not been reached during the 8.5 simulation days (Fig. 2). Figure 10 depicts the minimum diameter a \( \text{H}_2\text{SO}_4\)-\( \text{H}_2\text{O} \) aerosol particle needs to possess in order to become a water vapor condensation nucleus (activation diameter). For example for WSS=0.5\%, the critical diameter is about 56 nm. As suggested by our above AEROFOR model simulations, in the Chinese pollution plume even newly formed particles grow to diameters larger than 50 nm within only a few days and therefore will act as CCN at WSS=0.5\%. For lower WSS the minimum diameter increases very steeply (Fig. 10), which is due to the Kelvin effect.

Figure 11 shows the modeled number concentration of CCN sized \( \text{H}_2\text{SO}_4\)-\( \text{H}_2\text{O} \) aerosol particles versus WSS. CCN concentrations refer to the end of the simulation period and to the four model cases A (\( N_{\text{ini}}=0 \)), B (\( N_{\text{ini}}=250 \)) etc. The right panel is the model with gaseous sulfuric acid formation switched off. Here particles grow only due to coagulation. In this case for models B, C and D the CCN concentrations increase initially steeply with WSS which means that an increasing fraction of the particles can act as CCN. For WSS>0.4\% (critical diameter: 65 nm), CCN concentrations remain nearly constant, which means that all particles present at the end of the simulation period act as CCN. Due to coagulation, during the simulation period, their total number concentration has decreased and their diameters have increased. The left panel of Fig. 11 shows the model results with gaseous sulfuric acid formation switched on. Now, \( \text{H}_2\text{SO}_4\)-\( \text{H}_2\text{O} \) nucleation generates new particles and both, new and preexisting particles grow by \( \text{H}_2\text{SO}_4\)-\( \text{H}_2\text{O} \) condensation and coagulation. Now, for WSS>0.1\% (critical
diameter=163 nm), CCN concentrations become mostly (with exception of model case D) much larger than for the model without GSA formation since numerous newly formed particles become CCN. For WSS<0.1%, CCN concentrations are largest for case D and smallest for case A, which is similar to the case without GSA formation (bottom panel). For WSS>0.1%, CCN concentrations are largest for case A and smallest for case D. For WSS>0.5%, CCN concentrations become nearly constant. This means that all particles, including even the newly formed ones, become CCN.

Our findings suggest that, already on day 6, the Chinese pollution plume had developed its full CCN potential (maximum N100) and was primed for the formation of large concentrations of relatively small water droplets whenever WSS of about 0.2% built up (activation diameter 100 nm for 0.2% WSS, view Fig. 10). Condensation of a given mass of supersaturated water vapor on more CCN leads to smaller and more numerous cloud droplets. This tends to increase the albedo of the cloud and to increase the cloud lifetime with respect to rainout, since the droplet sedimentation velocity is approximately proportional to the square of the droplet diameter. In other words, after 9 days and at a distance of about 20 000 km from its birthplace, the plume had very suitable conditions for the formation of whiter clouds possibly possessing a reduced tendency for rainout. The Chinese pollution plume reported here represents a striking example of an environmental impact of long range transport of fossil fuel combustion generated SO$_2$.

5 Conclusions

We have investigated aerosol and CCN formation in a SO$_2$ rich pollution plume of Chinese origin which had traveled within 9 days from China to Europe where we have detected the plume at 5–7 km altitude during an aircraft mission by measuring SO$_2$. During most of the plume travel clouds were absent inside and above the plume and therefore efficient photochemical conversion of SO$_2$ to GSA took place, followed by GSA induced formation and growth of H$_2$SO$_4$-H$_2$O aerosol particles. Our AEROFOR
model simulations of aerosol and CCN formation lead to the following conclusions: in the plume, high SO\textsubscript{2} induced a strong increase of the concentration of H\textsubscript{2}SO\textsubscript{4}-H\textsubscript{2}O aerosol particles sufficiently large to act as cloud condensation nuclei already at a water vapor supersaturation exceeding only 0.1%. This implies that whenever WSS exceeded 0.1–0.2% in the 9 days aged plume, smaller and more numerous cloud droplets were formed. This tends to increase the cloud albedo and to decrease the tendency for rainout. The investigated Chinese pollution plume represents an interesting example of a potential environmental impact of long range transport of fossil fuel combustion generated SO\textsubscript{2}.

**Acknowledgements.** The authors gratefully acknowledge A. Dörnbrack (DLR) for providing ECMWF temperature and humidity data along the backward trajectories as model input data. The DWD is acknowledged to provide access to the ECMWF analysis data. We are grateful to the crew of the DLR Flight Department for their commitment and support to collect this data set. We also thank Bernd Kärcher (DLR) for the fruitful discussions and comments. This work was funded by DLR, MPI-K and Metropolia University of Applied Sciences, Helsinki. The data analysis was also supported in part by the Deutsche Forschungsgemeinschaft (DFG) within the Priority Program HALO under contract SPP1294.

This Open Access Publication is financed by the Max Planck Society.

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Table 1. Number concentration of initial particles $N_1$ (130 nm diameter) and $N_2$ (250 nm diameter) in the four model cases A–D. The initial SO$_2$ concentration was in all cases the same.

| Model Case | A  | B  | C  | D  |
|------------|----|----|----|----|
| Initial Particles $N_1$ [cm$^{-3}$] | 0  | 200 | 400 | 800 |
| Initial Particles $N_2$ [cm$^{-3}$] | 0  | 50  | 100 | 200 |
| Initial SO$_2$ concentration [cm$^{-3}$] | 4.4$\times 10^{10}$ |
Fig. 1. SO$_2$ mole fraction altitude profile of flight 20060503, smoothed with a running mean averaging 30 data points.
Fig. 2. AEROFOR model simulation input: temperature (blue) and relative humidity over water (green) and over ice (red) along the air mass back trajectory.
Fig. 3. AEROFOR simulation scenario A: no initial particle concentration. Upper panel: modeled OH and H$_2$SO$_4$ concentrations. Lower panel: homogeneous nucleation rate $J_{\text{honu}}$ and SO$_2$ concentration.
Fig. 4. AEROFOR simulation scenario A: no initial particle concentration. Upper panel: particle concentrations of particles of different size classes. Lower panel: condensation sink CS.
**Fig. 5.** AEROFOR simulation scenario C: initial particle concentration 500 cm$^{-3}$. Upper panel: modeled OH and H$_2$SO$_4$ concentrations. Lower panel: homogeneous nucleation rate $J_{\text{honu}}$ and SO$_2$ concentration.
**Fig. 6.** AEROFOR simulation scenario C: initial particle concentration 500 cm$^{-3}$. Upper panel: particle concentrations of particles of different size classes. Lower panel: condensation sink CS.
Fig. 7. AEROFOR simulation scenario D: initial particle concentration 1000 cm$^{-3}$. Upper panel: modeled OH and H$_2$SO$_4$ concentrations. Lower panel: homogeneous nucleation rate $J_{\text{honu}}$ and SO$_2$ concentration.
Fig. 8. AEROFOR simulation scenario D: initial particle concentration 1000 cm$^{-3}$. Upper panel: particle concentrations of particles of different size classes. Lower panel: condensation sink CS.
Fig. 9. Right panel: Particle concentrations for the four different model simulation cases initially (crosses) and after 8.5 simulation days (dots) if only coagulation is considered. Left panel: Particle concentrations initially (crosses) and after 8.5 simulation days (squares) if photochemical SO$_2$ conversion to H$_2$SO$_4$ is switched on.
Fig. 10. The critical activation diameter of a H$_2$SO$_4$-H$_2$O CCN as a function of water vapor supersaturation WSS.
Fig. 11. Modeled number concentration of CCN sized H$_2$SO$_4$-H$_2$O aerosol particles versus water vapor supersaturation WSS. CCN concentrations refer to the end of the simulation period. The four model cases (Nini=0, 250, 500, 1000 cm$^{-3}$) are considered. Right panel is the model with gaseous sulfuric acid formation switched off. Mind the different y-axis scaling.