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Measurements of gaseous H₂SO₄ by **AP-ID-CIMS during CAREBeijing 2008** Campaign

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As part of the 2008 Campaign of Air Quality Research in Beijing and Surrounding Regions (CAREBeijing 2008), measurements of gaseous sulfuric acid (H₂SO₄) have been conducted at an urban site in Beijing, China from 7 July to 25 September 2008 using atmospheric pressure ion drift – chemical ionization mass spectrometry (AP-ID-CIMS). This represents the first gaseous H₂SO₄ measurements in China. Diurnal profile of sulfuric acid is strongly dependent on the actinic flux, reaching a daily maximum around noontime and with an hourly average concentration of 5×10^6 molecule cm⁻³. Simulation of sulfuric acid on the basis of the measured sulfur dioxide concentration. photolysis rates of ozone and nitrogen dioxide, and aerosol surface areas captures the trend of the measured H₂SO₄ diurnal variation within the uncertainties, indicating that photochemical production and condensation onto preexisting particle surface dominate the observed diurnal H₂SO₄ profile. The frequency of the peak H₂SO₄ concentration exceeding 5×10^6 molecule cm⁻³ increases by 16% during the period of the summer Olympic Games (8-23 August 2008), because of the implementation of air quality control regulations. Using a multivariate statistical method, the critical nucleus during nucleation events is inferred, containing two H_2SO_4 molecules ($R^2 = 0.85$). When neither nucleation nor precipitation occurs, the condensation rate of H2SO4 correlates with the daytime sulfate mass concentration of the Aitken mode, but not with that of the accumulation mode aerosols.

1 Introduction

Gaseous sulfuric acid (GSA) is of critical atmospheric interest for its important role in various atmospheric processes, such as new particle formation (Seinfeld and Pandis, 1998; Finlayson-Pitts and Pitts, 1999) and modification of optical properties and hygroscopicity of preexisting aerosols (Khalizov et al., 2009a, b). GSA is predominantly formed through the gas-phase reaction between hydroxyl radical (OH) and sulfur dioxide (SO_2) in the presence of oxygen (O_2) and water (O_2),

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$$SO_2 + OH + M \rightarrow HSO_3 + M$$
 (R1)

$$HSO_3 + O_2 \rightarrow SO_3 + HO_2$$
 (R2)

$$SO_3 + H_2O + M \rightarrow H_2SO_4 + M \tag{R3}$$

GSA forms strong inter-molecular hydrogen bonds with H₂O, and H₂SO₄ has a low saturation vapor pressure at atmospherically relevant relative humidity (Zhang et al., 1993; Zhao et al., 2009). The fate of GSA in the atmosphere depends on the available surface area of preexisting aerosols. H₂SO₄ also forms molecular complexes with inorganic and organic species (Zhang et al., 2004). The size of these complexes is usually less than a few nanometers, below the detection limit of conventional instrumentation for nanoparticles. New particle formation includes nucleation to form the critical nucleus and subsequent growth of the critical nucleus to a detectable size of nanoparticles (L. Wang et al., 2010; Zhang, 2010). Although the detailed mechanisms of atmospheric aerosol nucleation and growth is still a subject of active research (Kulmala et al., 2004; Arnold, 2006; Nadykto and Yu, 2007; Young et al., 2008; Sipila et al., 2010; Zhang, 2010), H₂SO₄ is commonly considered as one of the dominant components of newly formed nanoparticles.

Atmospheric aerosols have important implications on the solar and terrestrial radiation budget (IPCC, 2007; Zhang et al., 2008), and multi-phase chemical processes (Molina et al., 1997). Freshly emitted soot contains mainly elementary carbon (EC) and to a lesser extent organic matter (OM). EC absorbs solar and terrestrial radiation and is also referred to as black carbon (BC). Fresh soot is largely hydrophobic and does not serve as cloud condensation nuclei (CCN) until a high supersaturation is reached (Zhang et al., 2008). However, after exposure to GSA, soot particles become hydrophilic, because of irreversibly uptake of GSA (Zhang and Zhang, 2005). Consequently, aged soot is more prone to wet deposition and has a smaller atmospheric residence time. Internally mixed soot particles with H₂SO₄ (or neutralized by ammonia to form ammonium sulfate) enhance light absorption and scattering, especially at

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higher mass fraction of sulfate (Cheng et al., 2008; Khalizov et al., 2009a). GSA significantly influences the soot-ageing process by modifying the morphology, optical properties, and hygroscopicity of soot particles (Zhang et al., 2008). Measurements of GSA provide a valuable input for evaluation of the soot aging process under atmospheric conditions. Furthermore, sulfuric acid can efficiently catalyze aqueous reactions of organic compounds, formed from photo-oxidation of volatile organic compounds (Suh et al., 2001; Zhang et al., 2002b), contributing to the formation of secondary organic aerosols (Zhao et al., 2005, 2006).

Because of rapid economic development and urbanization, Beijing has become one of the most populated cities in China. Increasing demand for energy in both industrial and domestic sectors leads to large fossil fuel consumption, with annual emissions of 2.39×10^5 tons of nitrogen oxides (NO_x = NO + NO₂), 2.4×10^5 tons of volatile organic compounds (VOCs), 1.9×10^5 tons of SO₂, and 5.8×10^4 tons of soot (Chan and Yao, 2008). Consequently, Beijing is frequently experiencing severe air pollution events, which are characterized by high concentrations of ozone (O₃) and particulate matter (PM) and severe deterioration in visibility. Rapid soot aging has been reported in the Beijing area, when the single scattering albedo of the aerosols increases to values above 0.9 within several hours due to secondary aerosol formation and condensation growth (Cheng et al., 2009).

Because of concerns on air quality during the 2008 Olympic Games period, the Beijing Municipal Environmental Protection Bureau (EPB) adopted a series of air quality control measures (Wang et al., 2009). On 20 July, a rule for odd/even numbers of the registration plate of automobiles was applied, aiming a reduction of 50% of the traffic volume from the private car sector. During the Olympic Games period, extra 20% of state owned vehicles were off the road, but extra diesel buses were added to public transportation fleet to accommodate the increasing demand for public traffic among different stadiums. After the Paralympics ended on 17 September, most of the regulations were lifted and on-road traffic increased. The Campaign of Air Quality Research in Beijing and Surrounding Regions (CAREBeijing 2008) was one of a series

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of comprehensive field studies (e.g., CAREBeijing 2006 and 2007) targeting the understanding of the chemical and physical processes responsible for the air pollution episodes. During the CAREBeijing 2008 campaign, a suite of state-of-the-art instruments were deployed to measure trace gases, meteorological parameters, and aerosol

This paper reports measurements of GSA during the CAREBeijing 2008 field study. In a separate publication, the role of GSA in new particle formation and growth has been investigated by Yue et al. (2010). Data validation of the GSA measurements is performed using a normalized sequential difference analysis (NSD). Meteorological effects on GSA are also discussed. The GSA diurnal features are presented and the influences from the administrative regulations on GSA measurements are discussed. Moreover, the fate of GSA in the Beijing urban environment is evaluated, focusing on its photochemical production and condensation sink. By applying a multivariate statistical method, the number of H_2SO_4 molecules in the critical nuclei is estimated during nucleation events. The contribution of H_2SO_4 to particulate sulfate is assessed.

5 number size distribution and chemical compositions (PM₁).

2 Experimental

2.1 GSA measurements

The observation site was on the rooftop of a six-story building on the campus of the Peking University (39°59′21″ N and 116°18′26″ E), located in the NW part of the urban Beijing area. GSA was measured with atmospheric pressure ion-drift chemical ionization mass spectrometry (AP-ID-CIMS). The detailed instrument description has been provided elsewhere (Fortner et al., 2004; Zheng et al., 2010). Briefly, the AP-ID-CIMS consisted of an inlet (10 cm ID and 60 cm long), an Am-241 ion source, an atmospheric pressure ion-drift tube, and a quadrupole mass spectrometer (QMS). The AP-ID-CIMS was housed inside an air-conditioned observatory. To minimize wall loss, the inlet was machined into a scooped-shape and the sampling rate was kept at about

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1200 standard liters per minute (slpm). About 55 slpm of the total air sample was introduced into the drift tube, where ambient H₂SO₄ underwent proton-transfer reactions with nitrate anion (NO₃·HNO₃) (m/e 125), generated from HNO₃ vapor by α particle radiation, to form HSO₄·HNO₃ (*m/e* 160). All reagent and product ions were sequentially monitored by the QMS in selective ion monitor (SIM) mode for about 12 s. Instrument calibrations were conducted twice a week with a custom-made water photolysis GSA primary source adjustable in the concentration range of 10⁷ to 10⁸ molecules cm⁻³. For a 5 min average time, the detection limit for GSA was about 10⁵ molecules cm⁻³. From in-situ calibrations, the sensitivity of AP-ID-CIMS (counts per second (Hz) per unit H₂SO₄ molecule cm⁻³) varied within 36% of the mean value and was reported as the uncertainty in the present work.

Aerosol and trace gas measurements

Aerosol number size distribution (3 nm to 10 µm) was measured by a twin differential mobility particle sizer (TDMPS) and an aerodynamic particle sizer (APS) (TSI Inc., model 3321). The TDMPS consists of two Hauke-type differential mobility analyzers and two condensation particle counters (TSI Inc., model 3010 and 2025) and are used to measure the particle size distributions from 3 to 900 nm (Stokes diameter). The APS is used to measure aerosols with aerodynamic diameter between 500 nm and 10 µm. A detailed description and operation conditions of the TDMPS and APS are provided by Yue et al. (2010). The concentration of particle surface area is integrated from the particle number distribution, by assuming a spherical geometry for the particle.

An Aerodyne High-Resolution Time-of-Flight Aerosol Mass Spectrometer (HR-ToF-AMS) was deployed to characterize particle chemical compositions. The detailed instrumental description of HR-ToF-AMS has been given by DeCarlo et al. (2006). The operation procedure of the HR-ToF-AMS during the 2008 CAREBeijing campaign has been described by Huang et al. (2010) in a companion paper of this special issue. Briefly, the HR-ToF-AMS was operated sequentially in a cycle of 5 modes every 10 min, including a 2 min V-mode to obtain the mass concentrations of the non-refractory

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species, a 2 min W-mode to obtain high resolution mass spectral data, a 4 min separate PToF (particle time-of-flight) mode to determine size distributions of species under the V-mode, and a 2 min Soft-EI mode using a lower EI voltage (~13 eV).

The photolysis frequencies of O_3 (J_{O^1D}) and NO_2 (J_{NO_2}) were measured by specifically designed filter radiometers provided by Forschungszentrum Jülich, Germany (FZJ). The instruments were calibrated at FZJ with a reference spectroradiometer before and after the campaign. The accuracy of the photolysis frequency measurements was estimated to be 10% at solar zenith angles smaller than 80° (Bohn et al., 2008).

Measurements of meteorology parameters including wind direction, wind speed, temperature, pressure, relative humidity, precipitation, UVA and UVB were conducted with a LASTEM auto meteorology station (LSI-LASTEM, M7115). SO₂ was measured with a commercial fluorescence sulfur dioxide analyzer (ECTECH, E9850). Both instruments were maintained and calibrated routinely.

3 Results and discussion

3.1 Instrument performance and meteorological effect

Accurate measurements of GSA represent a great challenge because of its low concentration. A normalized sequential difference (NSD) analysis of the GSA data is performed to evaluate the performance of the AP-ID-CIMS instrument. NSD at time t is calculated from (Arnold et al., 2007)

$$NSD_{t} = \frac{C_{t} - C_{t-1}}{\sqrt[2]{C_{t} \times C_{t-1}}}$$
 (1)

where C_t and C_{t-1} correspond to two consecutive GSA data points at time t and t-1, respectively. NSD represents the consistency of the instrument performance. Since all the factors affecting the observation results are random in nature, NSD_t is expected to exhibit a normal distribution with a mean of zero if there is not a systematic bias

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associated with the instrument. Figure 1a shows the frequency histogram of the NSD plotted in bins of 0.023 width. The histogram appears to be nearly a Gaussian distribution with a slight positive skewness. A p-value of 0.24 (larger than 0.05) is obtained from a standard t-test between the NSD and a randomly generated normal distribution with the same standard deviation as the calculated NSD but a mean value of zero, indicating that the positive skewness is statistically insignificant. Therefore, the GSA data shows no systematic instrument bias in either direction. Similarly, by plotting NSD with GSA concentration (Fig. 1b), it is demonstrated that within the observed GSA concentration range, the NSD is also normally distributed, i.e., the AP-ID-CIMS does not show a systematic bias within the range of observed GSA level.

GSA is highly sticky to almost all kind of surfaces. In order to minimize the inlet losses, a short inlet and a high sampling flow rate are used for the GSA measurement. The inlet is made of a 10 cm ID aluminum tube, which significantly reduces diffusion. Turbulence induced wind gust may impact GSA measurements. Figure 2 presents (a) a polar plot of the NSD with wind direction (North = 0°) and (b) a scatter plot of NDS with wind direction. Overall, NSD scatters in all wind direction with a slight bias under NW wind, but NSD is symmetrically distributed at any wind speed. This result is explained by the orientation of the inlet, i.e., when wind blows from perpendicular to the inlet or at a higher speed, more wall loss is expected as the air flow changes direction for both cases, but this bias is statistically insignificant.

H₂SO₄ measurements

The measurement period spans from 7 July to 25 September, covering the entire period of the XXIX Olympic Games and the Paralympics. Because GSA is photochemically produced and readily lost to aerosol surfaces, its concentration is expected to follow solar radiation intensity and is anti-correlated with the aerosol loading. Figure 3 shows the time series of GSA concentration measured during the 2008 CAREBeijing field campaign. The gaps within the series are due to interruption of power supply or instrument calibration. Diurnally, GSA typically remains a minimum value from late night

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to early morning and reaches a daily peak at noontime. The maximum peak value of GSA is measured on 10 September, with a concentration of 1.9×10^7 molecules cm⁻³.

The reaction between SO₂ and OH (Reactions R1 to R3) represents the primary pathway to form GSA (Finlayson-Pitts and Pitts, 1999), whereas GSA is readily removed by condensing onto aerosol surfaces. Simultaneous measurements of GSA and aerosol surface area present a unique opportunity to evaluate the competition between production and removal of GSA. The steady-state GSA concentration is calculated from

$$P_{\rm GSA} = L_{\rm GSA} \tag{2}$$

where $P_{\rm GSA} = k_{\rm OH+SO_2}[{\rm OH}][{\rm SO_2}]$ and $L_{\rm GSA} = \frac{\gamma \cdot S \cdot \bar{\nu}}{4}[{\rm H_2SO_4}]$ (Freiberg and Schwartz, 1981), $P_{\rm GSA}$ and $L_{\rm GSA}$ denote GSA production and loss, respectively, $k_{\rm OH+SO_2}$ is the rate constant of Reaction (R1) (JPL, 2006), $\bar{\nu}$ is the mean velocity of GSA molecules, S is the aerosol surface concentration (μ m² cm⁻³), and γ is the uptake coefficient of GSA on aerosol surface. A value of 0.73 (Jefferson et al., 1997) is adopted for γ . Because OH is not measured in this work, the OH concentration is estimated according to Ehhalt and Rohrer (2000),

$$[OH] = a(J_{O^1D})^{\alpha} (J_{NO_2})^{\beta} \frac{bNO_2 + 1}{cNO_2^2 + dNO_2 + 1}$$
(3)

where $J_{\text{O}^1\text{D}}$ and J_{NO_2} are observed photolysis rates of O_3 and NO_2 , respectively. The values of the constants are taken from Ehhalt and Rohrer (2000), i.e., $a=4.1\times10^9$, b=140, c=0.41, d=1.7, $\alpha=0.83$, and $\beta=0.19$. It has been suggested that Eq. (3) is valid under high NO_{X} conditions, which is typically the case in the megacity Beijing with large NO_{X} emissions (Liu and Shao, 2007). Because only dry deposition is considered in Eq. (2), GSA measurements when precipitation occurs are excluded in this investigation. Figure 4a shows hourly average diurnal plots of the measured aerosol surface area, SO_2 , and calculated OH concentration. Figure 4b displays both the hourly average GSA measurements with one σ error bar and calculated GSA from

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Eq. (2). Daytime GSA reaches a daily maximum around noontime with a value of \sim 5 x 10⁶ molecules cm⁻³. The calculated GSA captures the measured trend in general, indicating that photochemistry dominates the GSA production. However, since photochemistry is the only source considered, nighttime GSA cannot be accounted for. Given that the lifetime of GSA relative to aerosol uptake is less than a minute, the nighttime concentration of GSA ($\sim 1 \times 10^6$ molecules cm⁻³) clearly indicates that other non-photochemical mechanism is operative to produce OH radicals, such as the ozonolysis of alkenes (Zhang et al., 2002a; Fan and Zhang, 2004). The planetary boundary layer height typically reaches the minimum in the early morning and grows rapidly within a few hours after sunrise. As a result, surface air pollutants are vented upward and air masses of the regional sources are mixed down, explaining the variations in the particle surface area concentration and SO₂ concentration. Both OH and GSA are photochemically driven and their concentrations show clear variations with the solar diurnal cycle.

Another interesting feature in Fig. 3 is that the measured GSA peak value exhibits a periodic variation of about one week. Such a variation is related to the aerosol loading, which is regulated by the large-scale weather pattern in this area. A change in the weather pattern, such as the wind direction (from northwest) or precipitation, leads to a cleaner air condition, allowing GSA to reach a higher concentration.

In order to improve air quality during the Olympic Games, control measures have been implemented to reduce primary air pollutant emissions, especially, from industries, on-road vehicles and construction sites. Considerable reductions in both gasphase and aerosol-phase pollutants have been observed by both mobile laboratory (Wang et al., 2009) and surface stations (T. Wang et al., 2010). Moreover, more precipitation and cleaner conditions associated with the northerly wind were experienced during the period. Consequently, more "Blue-Sky" days were observed during the Olympic Games in the Beijing region (Zhang et al., 2010). Since GSA production is enhanced by stronger actinic flux and uptake by preexisting aerosol surfaces represents the dominant sink for GSA, an increasing trend in the daily GSA concentration is

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evident in Fig. 3, especially during the Olympics and Paralympics period, when more stringent traffic control measures were adopted. For example, the frequency of the peak H_2SO_4 concentration exceeding 5×10^6 molecule cm⁻³ increases by 16% during 8–23 August, compared to the time period prior to 8 August. This trend is consistent with that of the frequency of new particle formation events (Yue et al., 2010).

3.3 H₂SO₄ in new particle formation

During the CAREBeijing 2008 campaign, fourteen nucleation events were observed, which has been discussed previously in a companion paper in this special issue (Yue et al., 2010). In this work, a multivariate statistical method (MSM) (McGraw and Zhang, 2008) is employed to analyze the molecular components of the critical nuclei during eleven nucleation events. Table 1 summarizes new particle formation rate (FR) and GSA measurements.

To perform the MSM analysis, a multicomponent extension form of the kinetic nucleation theorem (Ford, 1997) is deduced by treating the critical clusters as the transition state during the nucleation process, i.e., the nucleation rate is determined from the sum of flux of each condensable gas that contributes to the growth of clusters over the critical size. Details of the MSM analysis for aerosol nucleation are described by (McGraw and Zhang, 2008). Applying the data listed in Table 1, we find that the most likely number of $\rm H_2SO_4$ molecules in the critical clusters is 2. This is consistent with the result by Anderson et al. (2008), who suggest that at least two sulfuric acid molecules are required to form a stable critical cluster. It should be pointed out that other species, including organics, can also be presence in the critical nucleus (Zhang et al., 2004, 2009; Metzger et al., 2010), but cannot be concluded from our present analysis.

3.4 Contribution of GSA to particulate sulfate

To evaluate the likely contribution of GSA to particulate sulfate, we compare the calculated condensation rates to the measured sulfate mass concentrations by the

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concentration (Fig. 4b). For the Aitken mode aerosols, the calculated GSA condensation rate exhibits a close correlation with the measured concentration of sulfate mass from the HR-ToF-AMS. This may imply that condensation of GSA (followed by neutralization with ammonia) dominates particle growth in the mode. In contrast, for the case of accumulation mode aerosols, the increase in the concentration of sulfate mass from the HR-ToF-AMS is noticeably delayed than that of the calculated GSA condensation rate, suggesting that GSA condensation hardly contributes to the growth in this mode. Instead, the growth in accumulation mode aerosols is more likely to be explained by inter-modal coagulation. It has been suggested by Guo et al. (2010) that fine mode sulfate in the Beijing area can arise from secondary sources, since biomass burning and automobile exhausts also contribute to fine particles. Also, according to He et al. (2010), sulfate production through the agueous phase reaction between hydrogen peroxides (H₂O₂) and SO₂

HR-ToF-AMS. Figure 5 shows the diurnal profiles of the calculated GSA condensa-

tion rates for the Aitken and accumulation modes, which are evaluated using Eq. (2). The aerosol surface area of each mode is integrated from TDMPS number size dis-

tribution measurement. Also shown in Fig. 5 for comparison are diurnal profiles of the corresponding aerosol sulfate mass concentrations for the Aitken and accumulation modes measured by the HR-ToF-AMS. Limited by the sampling efficiency of the

HR-ToF-AMS, we define Aitken and accumulation modes as 40 to 100 nm and 100 to 400 nm in Stokes diameters, respectively. Both modes of aerosol sulfate show sig-

nature of predominant secondary formation, i.e., the lowest concentration in the early

morning and the highest concentration in either the afternoon or early evening. The calculated GSA condensation rate in both modes is strongly influenced by the GSA

 $SO_2 + H_2O \leftrightarrow HSO_3^- + H^+$ (R4)

$$HSO_3^- + H_2O_2 \longrightarrow HSO_4^- + H_2O$$
 (R5)

in aqueous aerosols is not significant during the CAREBeijing 2008 campaign. Other possible sulfate production mechanisms include heterogeneous reactions on the 5030

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As part of the CAREBeijing 2008 field study, GSA measurement by AP-ID-CIMS was conducted at an urban site in Beijing City from 7 July 2008 to 25 September 2008. This represents the first GSA measurements in China. Data quality is examined using a normalized sequential difference procedure. GSA data set shows no statistically significant systematic errors. Correlations between GSA and meteorology parameters are also analyzed. Only a minor bias is identified when the wind direction is perpendicular to the sample inlet, likely due to turbulence-induced wall losses. Diurnally, GSA shows strong dependence on actinic flux and reaches daily maximum around noon, ranging from a few 10^6 to 1.9×10^7 molecules cm⁻³. Nighttime GSA ($\sim 1 \times 10^6$ molecules cm⁻³) is also observed, indicating that non-photochemical OH sources are operative in Beijing, such as the ozonolysis of alkenes. Traffic control and other administrative measures pertaining to the Olympic Games period appear to be effectively improving air quality in Beijing with significant reduction in aerosol loading, which is consistent with the observation that GSA daily maximum is noticeably increased from the beginning to the later part of the campaign. On the basis of the assumption that GSA concentration is governed by photochemical production and aerosol surface condensation loss, a steady state calculation is performed to evaluate GSA chemistry in Beijing. In general, simulated GSA captures the diurnal variations of measured GSA, indicating that condensation on preexisting aerosol surface corresponds to the dominant loss mechanism of GSA. Hence, GSA can potentially contribute to the aerosol aging process, considerably altering aerosol optical properties and hygroscopicity. GSA time series show a

aerosol surface or in-cloud processing (Seinfeld and Pandis, 1998). Future work, especially a detailed aerosol dynamic model along other aerosol and gaseous mea-

surements, is required to quantitatively assess the contributions of GSA condensation,

inter-modal coagulation, and other sulfate production mechanisms to the measured

sulfate mass concentrations in the Aitken and accumulation modes.

wave shape pattern characterized by accumulation of aerosol followed by a washout. Analysis of the critical cluster composition using a multivariate statistical method indicates the presence of two sulfuric acid molecules. The calculated condensation rate of H₂SO₄ correlates with the daytime sulfate mass concentration of the Aitken mode, but not that of the accumulation mode aerosols. The results imply that condensation of GSA is likely responsible for the particle growth in the Aitken mode, but the growth in accumulation mode aerosols is more likely to be explained by inter-modal coagulation.

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Table 1. Summary of nucleation events during the CAREBeijing 2008 field study including aerosol formation rate (FR), GSA concentration, and relative humidity (RH%) that used in the multivariate analysis to evaluate the number of sulfuric acid molecule in the critical cluster.

Date	$FR (cm^{-3} s^{-1})$	$H_2SO_4 (10^6 cm^{-3})$	RH %
12-Jul	7.3	5.0	27.8
17-Jul	13.2	7.1	40.8
1-Aug	6.9	4.8	58.9
12-Aug	13.2	7.4	63.4
15-Aug	8.7	5.5	51.5
18-Aug	2.1	3.2	50.5
23-Aug	7.3	6.3	41.2
31-Aug	5.8	4.5	47.3
1-Sep	4.2	5.3	40.8
18-Sep	2.4	4.0	44.3

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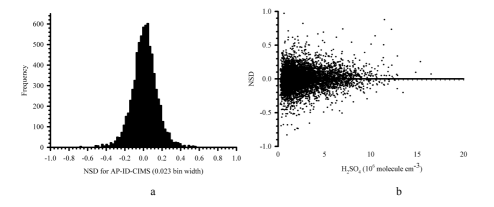


Fig. 1. (a) Frequency histogram of normalized sequential difference (NSD) for the AP-ID-CIMS; **(b)** scatter plot of NSD with GSA concentration.

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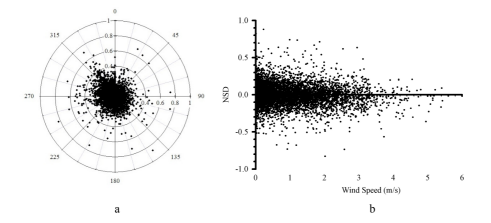


Fig. 2. (a) A plot of NSD with wind direction (North represented by 0 degree). **(b)** A scatter plot of NSD with wind speed.

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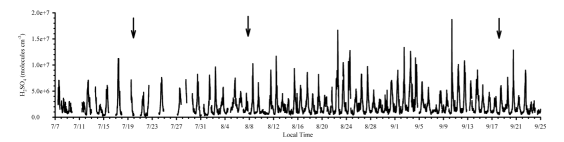


Fig. 3. Time series of 10 min averaged GSA concentration observed from 7 July to 25 September during the CAREBeijing 2008 Campaign. The arrows indicate the date of implementation of some major emission control measures, i.e., 20 July when odd/even plate number rule was applied, 8 August when full scale control was applied, and 18 September when the regulations were loosened.

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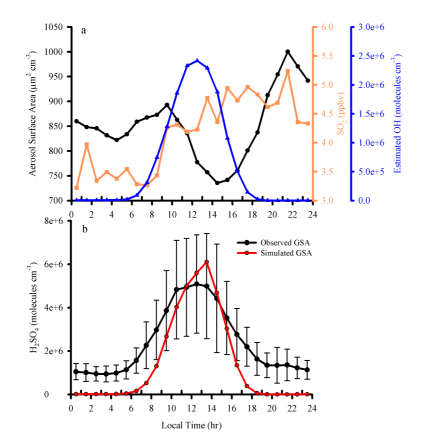
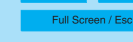


Fig. 4. (a) Hourly average diurnal profiles of aerosol surface area, SO₂, and modeled OH from 7 July to 25 September; (b) calculated (red) and measured (black) diurnal profiles of GSA.





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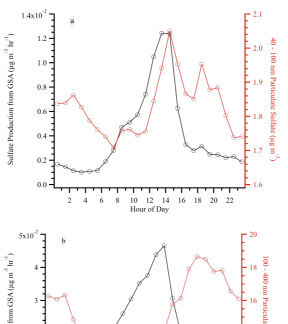
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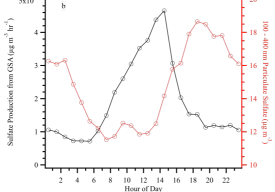


Fig. 5. (a) Diurnal profiles of condensation rates of GSA onto Aitken mode aerosols (40-100 nm) and sulfate concentrations in Aitken mode; (b) diurnal profiles of condensation rates of GSA onto accumulation mode particles (100-400 nm) and sulfate concentrations in accumulation mode.