Supporting Information for

Unprecedented ambient sulphur trioxide (SO₃) detection: possible formation mechanism and atmospheric implications

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**Text S1. Sampling site**

Our field measurements were conducted at BUCT (Beijing University of Chemical Technology) sampling site (39.94° N, 116.30° E), which was located on the west campus of BUCT. This site was located at the roof of a teaching building, which is approximately 15 m above the ground level. Around 130 m to the north and 550 m to the west are Zizhuyuan Road and West Third Ring Road, respectively. The “West Third Ring Road” is one of the main “Ring” roads in Beijing. Besides the influence of traffic, this site is also affected by local commercial and residential activities. Therefore, the BUCT monitoring site is representative of an urban site.

**Text S2. The nitrate-CI-API-LTOF mass spectrometer**

A nitrate-based Chemical Ionization - Atmospheric Pressure interface - Long-Time-of-Flight (CI-APi-LTOF, Aerodyne Research Inc, USA and Tofwerk AG, Switzerland) mass spectrometer was deployed to detect SO$_3$ and gas-phase sulfuric acid. The CI-APi-LTOF consists of an optimized inlet for chemical ionization (CI-inlet)$^{2,3}$ and an APi-LTOF mass spectrometer with the mass resolving power of ~10000 Th/Th. Nitrate ions (NO$_3^-$·(HNO$_3$)$_n$, $n=0,1$ and 2) were used as reagent ions. The working principle of nitrate-CI-APi-LTOF has been described in many previous studies.$^{2,4}$

In the charging part of CI-inlet, the nitrate ions are electrostatically pushed into ambient sample flow to react with SO$_3$ and H$_2$SO$_4$. In the CI-inlet, the ion-molecule reaction time was ~200 ms.$^4$ Pure air originated from a pure air generator (Aadco 737) was used as the sheath air. Ambient air was sampled into the CI-inlet through a ½ inch stainless steel tube. A 0.8 L min$^{-1}$ flow from the mixed flow entered the APi-LTOF. Data of CI-APi-LTOF were acquired at 5 s time resolution and analyzed with a MATLAB tofTools package.$^5$

**Text S3. Detection of sulfuric acid with nitrate reagent ions**

The ion-molecule reactions of sulfuric acid with nitrate reagent ions can be described by the following reaction$^{2,4}$:

$$\text{H}_2\text{SO}_4 + \text{NO}_3^-\cdot(\text{HNO}_3)_n \rightarrow \text{HSO}_4^-\cdot(\text{HNO}_3)_j + (n-j+1) \text{HNO}_3$$  \hspace{1cm} R (S1)

where $n = 0, 1$ or $2$ and $j = 0$ or $1$. Due to H$_2$SO$_4$ being a stronger acid, de-protonation occurs during its collision with nitrate ions. Thus, H$_2$SO$_4$ molecules can be detected as de-protonated monomer ions and cluster ions with HNO$_3$ in CI-API-LTOF.

To take the variation in the total reagent ions into account, neutral sulfuric acid was quantified according to the following equation:

$$[\text{H}_2\text{SO}_4] = C \times \frac{\sum_{n=0}^{\infty} (\text{HSO}_4^-\cdot(\text{HNO}_3)_j)}{\sum_{n=0}^{\infty} (\text{NO}_3^-\cdot(\text{HNO}_3)_n)}$$  \hspace{1cm} E (1)

where $C$ (in units of cm$^{-3}$) is a calibration coefficient from in-situ calibration.

**Text S4. Calibration experiment for SO$_3$**

The calibration of SO$_3$ was implemented by introducing a known amount of gaseous SO$_3$ produced by the reaction of SO$_2$ and OH radicals formed by UV photolysis of water vapour, which is similar to the method for sulfuric acid calibration in the previous literature (i.e. Kürten et al., 2012)$^6$. During the calibration experiment, a 10 L·min$^{-1}$ N$_2$ flow, a 100 mL·min$^{-1}$ pure air flow, a 300 mL·min$^{-1}$ SO$_2$ flow and a set of 20 -
400 mL·min$^{-1}$ saturated water vapour flow were mixed together as the calibration sampling flow. Then, the mixed flow was exposed to 184.9 nm UV light to produce OH radicals which reacted with SO$_2$ to produce SO$_3$. The schematic of the experimental setup was shown in Figure S2. The UV lamp was turned on in an N$_2$ environment at least one hour before the actual calibration measurement in order to achieve a stable light intensity. During the calibration, the box was flushed with a 1 - 2 L·min$^{-1}$ dry N$_2$ flow to avoid the absorption of UV light by O$_2$. Different levels of SO$_3$ were achieved by adjusting the flow of saturated water vapour, that is, adjusting OH radical concentrations. The theoretical generated SO$_3$ concentrations were calculated by Numerical Model which has been introduced in Kürten et al., 2012$^6$. The time profiles of water content ([H$_2$O]) and normalized signals of sulfuric acid and SO$_3$ were exhibited in Figure S3. The correlation between normalized SO$_3$ signals measured by CI-APi-LTOF and SO$_3$ concentrations formed by photo-oxidation of SO$_2$ by OH radicals was depicted in Figure S4. After taking the diffusion loss of the sampling line into account, a calibration coefficient of $1.7 \times 10^{10}$ molecule cm$^{-3}$ was obtained. The diffusion loss was assumed as same as that of sulfuric acid.

Similar to the quantification of sulfuric acid$^2$, to regard the variation in the total reagent ions, SO$_3$ was quantified according to:

$$[SO_3] = C \times \frac{[SO_3 \cdot NO_3]}{\sum_{n=0}^{\infty} (NO_3) \cdot (HNO)_n}$$  \hspace{1cm} \text{(2)}$$

where $C$ (in units of cm$^{-3}$) is a calibration coefficient of SO$_3$.

**Text S5. Calculations for collision rate coefficients**

Using ion-molecule collision parametrizations from two previous studies (Su & Bowers, 1973 and the Su & Chesnavich, 1982; note that these are the standard approaches used to estimate ion-molecule collision rates also in CIMS studies)$^7,8$, with the dipole & polarisability of H$_2$SO$_4$ computed at the same level of theory (wb97xd/aug-cc-pVTZ; numerical values given below), the results for the collision rate coefficients (unit cm$^3$ molecule$^{-1}$ s$^{-1}$) are as follows:

H$_2$SO$_4$ (mass 98)
- Dipole moment 3.15093 Debye
- Polarisability 35.9657 bohr$^3$ = 5.330 Ånström$^3$

Collision rate at 298.15 K with HNO$_2$·NO$_3^-$ (mass 125) according to Su & Chesnavich 1982: 2.11E-09
Collision rate at 298.15 K with NO$_3^-$ (mass 62) according to Su & Chesnavich 1982: 2.54E-09

SO$_3$ (mass 80)
- Dipole moment 0 Debye
- Polarisability 28.4649 bohr$^3$ = 4.218 Ånström$^3$

Collision rate at 298.15 K with HNO$_2$·NO$_3^-$ (mass 125) according to Su & Chesnavich 1982: 6.88E-10
Collision rate at 298.15 K with NO$_3^-$ (mass 62) according to Su & Chesnavich 1982: 8.13E-10

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The main charging ions are HNO$_3$·NO$_3$· and NO$_3$·. The corresponding ratios (collision rate of H$_2$SO$_4$ / collision rate of SO$_3$) are 3.07, 3.12, 1.39 and 1.42, respectively. The newer ion-molecule collision rate parametrization (Su & Chesnavich, 1982) thus predicts a difference of a factor of 3, whereas the older one only predicts a difference of about a factor of 1.5. A non-polar molecule collides much slower with an ion than a strongly polar molecule. The product ions H$_2$SO$_4$·NO$_3$· (binding Gibbs free energy of -32.6 kcal/mol at the wB97xd/aug-cc-pVTZ level) and SO$_3$·NO$_3$· (binding Gibbs free energy -28.4 kcal/mol at the same level) are very strongly bound and stable. The sensitivity of SO$_3$ could be less than that of H$_2$SO$_4$ by a factor of 3. Thus, a factor of 3 difference would lead to an underestimation of SO$_3$ if the calibration factor for H$_2$SO$_4$ was used to quantify SO$_3$.

**Text S6. Quantum chemical calculations**

Quantum chemical calculations demonstrate that the SO$_3$·(NO$_3$·) cluster is very strongly bound compared to the HNO$_3$·(NO$_3$·) cluster (Table S1). The difference in binding is over 10 kcal/mol both in electronic and free energies, with the more rigorous coupled-cluster methods predicting a larger difference than the density functional theory method used here. SO$_3$ molecules will thus be very efficiently charged by nitrate ions in a nitrate-CI-API-LTOF instrument, as the charge transfer reaction HNO$_3$·(NO$_3$·) + SO$_3$ → SO$_3$·(NO$_3$·) + HNO$_3$ is highly favourable. Furthermore, the thermal evaporation rate of SO$_3$·(NO$_3$·) clusters in the CI-inlet will be negligible, and also the (non-thermal) fragmentation of the cluster in the ion optics of the instrument will be considerably smaller than for example that of the (H$_2$SO$_4$)·HSO$_4$· cluster, which has binding energy comparable to HNO$_3$·(NO$_3$·)$^\text{9}$. All of this supports the hypothesis that the instrument sensitivity toward SO$_3$ will be very high. The optimized structure of the SO$_3$·(NO$_3$·) cluster is shown in Figure S5. The strength of the O$_3$S…ONO$_2$ interaction is reflected in the relatively short S…O distance.

The thermodynamics of the SO$_3$·H$_2$O + NO$_3$·(HNO$_3$) → SO$_3$·(NO$_3$·)·H$_2$O + HNO$_3$ reaction (R5) was assessed at the wB97X-D/aug-cc-pVTZ level. Three different hydrogen bonding patterns (conformers) for SO$_3$·(NO$_3$·)·H$_2$O were assessed, with the H$_2$O molecule placed either close to the SO$_3$ moiety, the NO$_3$· moiety, or in a bridging position between the two. The latter structure, where H$_2$O H-bonds to both O-S and O-N oxygen atoms, was found to be the lowest in free energy (at 298 K) (see Figure S6), though the differences between conformers were fairly small (below 2 kcal/mol). By comparison to the results in Table S1, it is likely that higher-level energy corrections (omitted here for computational reasons) would lead to an even more negative (favourable) reaction free energy.

**Text S7. Computational details**

Molecular and cluster geometries were optimized, and harmonic vibrational frequencies were calculated, with the ωB97X-D density functional$^{10}$ and the aug-cc-pVTZ basis set$^{11}$, using the Gaussian 16 program suite (Gaussian 16, Revision A.03)$^{12}$. Special care was taken in the optimizations to ensure that SO$_3$ and NO$_3$· had the correct D$_3$h symmetry point group (with a rotational symmetry number of 6). Thermal and vibrational zero-point contributions to enthalpies and entropies were calculated using the standard rigid rotor - harmonic oscillator approximations. Single-point energy corrections were performed on top of the ωB97X-D/aug-cc-pVTZ structures using explicitly correlated coupled cluster theory; specifically RHF-RCCSD(T)-F12$^{13-15}$ with the VDZ-F12 and VTZ-F12 basis sets$^{16}$. The Molpro 2015.1 program was used for these calculations (MOLPRO, version 2019.2, a package of ab initio programs, see https://www.molpro.net)$^{17}$. With the smaller basis set, the “F12a” variant was used, based on recommendations in the program manual. With the larger basis set, both “F12a” and “F12b” energies were calculated. The differences between binding energies
computed with the two variants were minimal (around 0.01 kcal/mol or less), and even the difference between
the basis sets was less than 0.3 kcal/mol for both HNO$_3$·(NO$_3^-$) and SO$_3$·(NO$_3^-$). Final results are presented
using the RHF-RCCSD(T)-F12b/VTZ-F12 values.

**Text S8. PM$_{2.5}$, black carbon (BC), particulate sulfate, trace gases, meteorological parameters and
UVB measurements.**

Mass concentrations of PM$_{2.5}$ were recorded by a TEOM (tapered element oscillating microbalances) monitor.
Non-haze and haze days were categorized as daily mean PM$_{2.5}$ mass concentrations of <100 μg m$^{-3}$, and ≥
100 μg m$^{-3}$, respectively.

Mass concentrations of BC in PM$_{2.5}$ were measured by the aethalometer (Magee AE33). PM$_{2.5}$ particles were
continuously collected through the filter tape. Then the transmission of light through the filter tape containing
the sample was obtained at seven different wavelengths. AE33 calculates the instantaneous concentration of
optically absorbing aerosols from the rate of change of the attenuation of light transmitted through the
particle-laden filter $^{18}$. The determination of black carbon concentration is based on the measurement of light
absorption on a filter loaded. In this study, aerosol particles are continually sampled on the filter, and the
optical attenuation is measured with high time resolution 1s. Optical attenuation (ATN) is measured on two
spots with different sample flows and the reference spot without the flow calculated as Equation 3:

$$\text{ATN} = -100 \ln \left( \frac{I}{I_0} \right) \quad \text{E(3)}$$

where $I$ and $I_0$ represent the detector intensity signal for the measurement spot and reference signal,
respectively. The factor 100 is there for convenience only $^{17}$.

Airflow ($F$) is measured after the air passes the filter (Eq.3), lateral airflow in the optical chamber $\zeta$ has to be
taken into account:

$$F_{\text{in}} = F_{\text{out}} (1 - \zeta) \quad \text{E(4)}$$

where the value of $\zeta$ is determined by measuring input and output flow and was found to be in the 0.02–0.07
range.

Attenuation coefficient ($b_{\text{ATN}}$) and Absorption coefficient ($b_{\text{ABS}}$) can be calculated as Equations 5 and 6

$$b_{\text{ATN}} = \frac{S \cdot \Delta \text{ATN}}{F \cdot \Delta t} \quad \text{E(5)}$$

$$b_{\text{ABS}} = \frac{b_{\text{ATN}}}{C} \quad \text{E(6)}$$

where $F$ is the volumetric flow and $S$ is the filter surface area on which the aerosol particles are deposited, $t$
represents the time, and $C$ is multiple scattering parameter $^{20}$.

BC concentration can be calculated using Equation 7:

$$BC = \frac{b_{\text{ABS}}}{\sigma_{\text{air}}} \quad \text{E(7)}$$

where $\sigma_{\text{air}}$ is the mass absorption cross-section, here we obtain the mass absorption cross-section value is 7.19
m$^2$ g$^{-1}$ measured at 970 nm to convert the observed light attenuation to the mass concentration of BC $^{21}$.

Combining Eqs. 5, 6 and 7, we obtain Eq.8 to calculate the concentration of BC:

$$BC = \frac{b_{\text{ABS}}}{\sigma_{\text{air}}} \quad \text{E(8)}$$
where $k$ is a loading effect parameter.

The mass concentrations of non-refractory PM$_{2.5}$ including sulfate concentration were measured by an online Time-of-Flight Aerosol Chemical Speciation Monitor (ToF-ACSM, Aerodyne Research Inc., USA) equipped with a cyclone to select PM$_{2.5}$ particles. Trace gases including SO$_2$, O$_3$ and NO$_x$ were recorded by Thermo analyzers (Model 43i, 49i and 42i, Thermo Scientific, USA). A weather station (Vaisala Inc., Finland) was deployed to measure meteorological parameters (ambient relative humidity (RH), temperature, wind speed, visibility). The UVB (280-315 nm) intensity was measured using CUV3 radiometer (Kipp & Zonen, USA).

The mixing layer height (MLH) was determined with the enhanced single-lens ceilometers CL-51 (Vaisala Inc., Finland), which utilized the strobe laser lidar technique (910 nm) to measure the attenuated backscattering coefficient profiles. The Vaisala software product BL-VIEW was used to determine the mixing layer height by finding the position with the maximum negative gradient (-dβ/dx) in the attenuated backscattering coefficient profiles as the top of the mixing layer.$^{22}$

Text S9. Sub-3 nm particles measurement

The number size distributions of sub-3 nm clusters/particles were measured using an Airmodus A10 particle size magnifier (PSM) coupled with an Airmodus A20 condensation particle counter (CPC)$^{4,23}$. The measured mobility diameter was from $\sim$1.2 to 2.5 nm. Diethylene glycol and n-butanol were used as working fluids of the PSM and CPC, respectively. The sub-3 nm clusters/particles can grow up to $\sim$90 nm by condensation of diethylene glycol vapour inside the PSM and then detected by the CPC$^4$. To minimize the loss of clusters/particles and increase the sampling efficiency, a core sampling system and high flow rate (7.5 L min$^{-1}$) were utilized$^1$. The time resolution of a full scan from 0.1 L min$^{-1}$ to 1.3 L min$^{-1}$ was 240 s.

Text S10. Calculation of condensation sink

Condensation sink (CS) describes the condensing vapour sink caused by the particle population$^{24}$:

$$\text{CS} = 2\pi D \sum_{d_p} \beta_{m,d_p} d_p N_{d_p}$$  \hspace{1cm} E(9)

where $D$ is the diffusion coefficient of the condensing vapour (usually assumed to be sulfuric acid), and $\beta_{m,d_p}$ is the transitional regime correction factor.

Text S11. Source identification of SO$_2$ in winter

In winter, the median concentrations of SO$_2$ exhibited similar diurnal trends as SO$_3$ (Figure 2A). A similar diurnal variation of SO$_2$ with an early morning peak has already been reported from another site in urban Beijing$^{25}$. We also studied the evaluation of median mixing layer height (MLH) together with the diurnal trend of the median concentration of SO$_2$, SO$_3$ and UVB (Figure S8). The median MLH was merely 200-300 m, and stable in the morning (~05:00 to ~08:30). Many studies have reported regional SO$_2$ is tightly linked with the combustion of sulfur-containing fuels and the smelting of sulfur-containing ores$^{26-32}$. Together with stable weather conditions (Figure S8), the elevated SO$_2$ concentration during the early morning could mainly be attributed to local emissions (e.g. residential and industry emission) and transportation$^{33}$.  

$$BC = \frac{S \times (\text{ATN} \times 100)}{F(1 - 0.5 \times \sigma_{\text{air}} \times \Delta \text{ATN} \times (1 - k \times \text{ATN}) \times \Delta t} \hspace{1cm} E(8)$$
Figure S1. High-resolution peak fitting of the peak $^{32}$SO$_3$·NO$_3$ and its main isotope peak $^{34}$SO$_3$·NO$_3$.

Figure S2. The schematic of the calibration experiment setup.
Figure S3. Time series of normalized signals of H$_2$SO$_4$ and SO$_3$, and [H$_2$O] in the calibration experiment.

Figure S4. The correlation between normalized SO$_3$ signals measured by CI-API-LTOF and SO$_3$ concentrations formed by the photo-oxidation of SO$_2$ by OH radicals. Our calibration experiment yielded a calibration coefficient of $1.7 \times 10^{10}$ cm$^{-3}$ for SO$_3$. This factor has taken diffusion loss of the sampling line of CI-API-LTOF into account.
Figure S5. Optimized structure of the SO$_3$·(NO$_3^-$) cluster, at the ωB97X-D/aug-cc-pVTZ level. The S…O interaction between SO$_3$ and NO$_3^-$ is shown as a covalent bond based on the relatively short interatomic distance, as well as the very strong binding energy. Key bond lengths are given in Ångström. Color coding: yellow=S, blue=N, red=O.

Figure S6. Lowest free energy (at 298 K) structure found for SO$_3$·(NO$_3^-$)·H$_2$O, at the wB97X-D/aug-cc-pVTZ level. Hydrogen bonds are indicated with dotted lines, and the corresponding distances are given in Ångström. Covalent bond lengths differ by less than 0.02 Å from those shown in Figure S5. Color coding: yellow=S, blue=N, red=O.
**Figure S7.** The averaged mass spectra of atmospheric naturally charged ions for one whole day (10 November 2018).

**Figure S8.** Median diurnal variation of concentrations of SO$_3$ and SO$_2$, the mixing layer heights (MLH), intensities of UVB, and wind speeds during winter.
Figure S9. Time profile of SO$_3$ concentration and mass concentration of sulfate in PM$_{2.5}$ from 7 February to 11 February 2019 (A) and median diel variation of SO$_3$ and sulfate for all non-haze days during the winter measurement period (B).
Figure S10. The relationship between the atmospheric ion signals of HSO₄⁻ and SO₃⁻NO₃⁻ during night time (18:00-5:00 next day) and early morning (5:00-8:00) from 9 to 22 November 2018.

Table S1. Comparison of the binding thermodynamics of HNO₃(NO₃⁻) and SO₃⁻(NO₃⁻) ion-molecule clusters, in kcal/mol. ΔE_{DFT} and ΔE_{F12} correspond to the electronic energies (not including vibrational zero-point corrections) computed at the oB97X-D/aug-cc-pVTZ and RHF-RCCSD(T)-F12b/VTZ-F12 levels, respectively, both at the optimized geometry corresponding to the former method. ΔG_{DFT,F12} corresponds to the Gibbs free energy (at 298.15 K and 1 atm reference pressure) obtained by combining the RHF-RCCSD(T)-F12b/VTZ-F12 electronic energy with the oB97X-D/aug-cc-pVTZ thermal and vibrational zero-point contributions.

|          | ΔE_{DFT} | ΔE_{F12} | ΔG_{DFT,F12} |
|----------|----------|----------|--------------|
| HNO₃(NO₃⁻) | -29.1    | -29.2    | -21.5        |
| SO₄(NO₃⁻)  | -40.1    | -44.4    | -32.7        |
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