A novel pathway of atmospheric sulfate formation through carbonate radicals

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Abstract. Carbon dioxide is considered an inert gas that rarely participates in atmospheric chemical reactions. Nonetheless, we show here that CO2 is involved in some important photo-oxidation reactions in the atmosphere through the formation of carbonate radicals (CO3−3). This potentially active intermediate CO3−3 is routinely overlooked in atmospheric chemistry concerning its effect on sulfate formation. The present work demonstrates that the SO2 uptake coefficient is enhanced by 17 times on mineral dust particles driven by CO3−3. Importantly, upon irradiation, mineral dust particles are speculated to produce gas-phase carbonate radical ions when the atmospherically relevant concentration of CO2 presents, thereby potentially promoting external sulfate aerosol formation and oxidative potential in the atmosphere. Employing a suite of laboratory investigations of sulfate formation in the presence of carbonate radicals on the model and authentic dust particles, ground-based field measurements of sulfate and (bi)carbonate ions within ambient PM, together with density functional theory (DFT) calculations for single electron transfer processes in terms of CO3−3-initiated S(IV) oxidation, a novel role of carbonate radical in atmospheric chemistry is elucidated.

1 Introduction

Atmospheric composition changes are subjected to highly reactive light-induced radicals, such as hydroxyl (·OH), hydroperoxyl (HO2·), or nitrate radicals (NO3·), which are able to alter not only compositions but also the physical and chemical properties of particulate matter (T. Liu et al., 2022; Stevenson et al., 2020; Mahajan et al., 2021). However, when atmospheric chemical reactions occur over humified particles at ambient conditions, which creates a locally enriched aqueous medium of unique chemical activity, other radicals might likewise gain importance. The carbonate radical (CO3−3) is typically such an active radical. The lifetime of CO3−3 ranges from a microsecond to even a few milliseconds, and its concentration can be 2 orders of magnitude higher than that of hydroxyl radicals over the water surface (Chandrasekaran and Thomas, 1983; Goldstein et al., 2001; Shafirovich et al., 2001; Sulzberger et al., 1997). In addition, the one electron reduction potential of the E0(CO3−3 / CO2−3) coupling is 1.78 V vs. a normal hydrogen electrode (NHE) at neutral pH, leaving CO3−3 a strong oxidant in aquatic chemistry (Cope et al., 1973; Bisby et al., 1998; Merouani et al., 2010). Previous studies concerning carbonate radical in aqueous media demonstrate that it reacts rapidly with some organic compounds (Merouani et al., 2010), especially for those electron-rich amine compounds (Stenman et al., 2003; Yan et al., 2019). Also, it has been pointed out that the scavenging of hydroxyl radicals by (bi)carbonate species leads
to the formation of CO$_3^{2-}$ ions (Graedel and Weschler, 1981; Xiong et al., 2016). Besides, a high second-order rate constant, lying at 10$^7$ M$^{-1}$ s$^{-1}$, has been reported for the reaction of CO$_3^{2-}$ with porphyrins (Ferrer-Sueta et al., 2003), indicating that this radical ion has great oxidation capability that may trigger atmospherically relevant chemical reactions, i.e., secondary inorganic species formation. However, it is only regarded as a marginal intermediate in tropospheric anion chemistry so far (Lehtipalo et al., 2016; Beig and Brasseur, 2000; Herrmann et al., 2000; Ge et al., 2021), and its underlying role as an active oxidant for heterogeneous reaction in the atmosphere is barely explored. Very recently, our group observed the promotional effect of CO$_3^{2-}$ on atmospheric nitrate formation (Fang et al., 2021). Motivated by this finding, attempts were made to further explore its role in other important, atmospherically relevant reactions.

It is well documented that sulfate (SO$_4^{2-}$) is also a key constituent of aerosols in the atmosphere (Huang et al., 2015; Su et al., 2016). It is able to serve as the precursor of efficient cloud condensation nuclei, with optical properties leading to a cooling effect (Wang et al., 2011). As a consequence, the mechanism aspect of secondary sulfate formation was the focus of numerous studies over the past decades (Hung et al., 2018; Stone, 2002; Zheng et al., 2015). There is a consensus that high-valence sulfur (VI), produced from the oxidation of anthropogenic SO$_2$, is the dominant source of atmospheric secondary sulfate. However, a remarkable missing sulfate budget emerges for the atmospheric modeling (Huang et al., 2019; Liu et al., 2021; Itahashi et al., 2018), which significantly underpredicts SO$_4^{2-}$ with respect to observational results when heterogeneous aerosol chemistry is not considered (Zheng et al., 2015; Feng et al., 2018; Wu et al., 2021). This indicates that the heterogeneous sulfate production pathway is a crucial process, and exploring the unrecognized heterogeneous mechanism is very likely to narrow the gap between observations in lab studies, field measurements, and numerical modeling. Due to the missing chemical mechanism that initiated a fast SO$_2$ oxidation, atmospheric models fail to capture the key feature of atmospheric observations of high sulfate production during dust storm episodes in the troposphere (Yu et al., 2020; Dong et al., 2016; Huang et al., 2014), where an evident increase in Ca$^{2+}$ (Wang et al., 2005; Li et al., 2013), carbonate-containing particles with high alkalinity (Tang et al., 2016; Li et al., 2014; Abou-Ghanem et al., 2020), and photoactive mineral components (Nie et al., 2012; Ta et al., 2003) are prevalent. Air mass is usually in low relative humidity, reportedly being 25 %–35 % (Al-Salihi and Mohammed, 2015; Csavina et al., 2014; Najafpour et al., 2020) in these events, during which the photochemical process is able to alter atmospheric constituents (X. C. Liu et al., 2022). Consequently, there are unknown heterogeneous reaction pathways and previously unconsidered promoters that have great potential to accelerate sulfate formation in the dust-storm-relevant conditions.

Due to the high stability of CO$_2$ under ambient conditions (Hossain et al., 2020), there are rare studies concerning the influence of CO$_2$ in atmospheric chemical processes (Deng et al., 2020; Y. Liu et al., 2020; Xia et al., 2021). CO$_2$ is demonstrated to form (bi)carbonate species over humidified dust particles (Baltrusaitis et al., 2011; Nanayakkara et al., 2014) and is reduced to CO under solar illumination (Deng et al., 2020). Nonetheless, its impact on atmospheric heterogeneous reactions remains poorly characterized. Our early laboratory study illustrates that CO$_2$ decreases the sulfate formation on aluminum oxide particles in the dark (Y. Liu et al., 2020), while upon solar illumination its role in SO$_2$ oxidation over mineral dust surfaces is still an open question. In addition, carbonate salt is abundant in authentic dust particles (Cao et al., 2005) and is reported to reach over 10 wt % of Asian dust particles (McNaughton et al., 2009). It is generally accepted that CO$_3^{2-}$ affects atmospheric chemistry and aerosol characteristics mainly through its intrinsic alkalinity, which buffers aerosol acidity and increases SO$_2$ adsorption and corresponding sulfate production in the presence of oxidants (Bao et al., 2010; Kerminen et al., 2001; Yu et al., 2018; Li et al., 2007; Al-Hosney and Grassian, 2005). In fact, either CO$_2$ or carbonate salt is able to produce the active CO$_3^{2-}$ under the ambient circumstance (Graedel and Weschler, 1981; Ervens et al., 2003) and is prone to increase the oxidative capacity in the atmosphere. Our early study shows that carbonate radicals serve as an active oxidant to accelerate NO$_2$ oxidation over mineral dust particles (Fang et al., 2021), allowing us to consider the possibility that fast heterogeneous SO$_2$ oxidation can be triggered by this active intermediate as well. Nevertheless, to the best of our knowledge, no work has ever considered how and to what extent the carbonate radical influences SO$_2$ heterogeneous oxidation in the atmosphere.

In the current study, through laboratory studies, we present that carbon dioxide and calcium carbonate, working as the precursor of carbonate radicals, have the ability to accelerate sulfate formation over authentic particles in the atmosphere. Together with quantum chemistry calculations, a detailed molecular mechanism regarding a single electron transfer (SET) process between carbonate radical and sulfite ions is elucidated. Furthermore, ground-based observations validate some findings from the laboratory-based simulations.

2 Experimental methods

2.1 Laboratory studies

2.1.1 Methodology for uptake coefficient estimation

The reaction uptake coefficient was estimated by the following Eqs. (1)–(3), as suggested by the previous work (Kong et al., 2014):
where \( v \) is the mean molecular velocity of SO\(_2\), \( A_s \) is the effective sample surface, \( R \) is the gas constant, \( T \) is the temperature, \( M_{SO_2} \) is the molecular weight of SO\(_2\), and \( Z \) is a total number of surface collisions per unit of time. To be precise, the formation rates (\( \frac{d[SO_2^2^-]}{dt} \)) in the equation were determined by ion chromatography (IC) measurements, followed by a conversion factor calculation through linear regression analysis for the integrated absorbance of sulfate bands and corresponding sulfate concentrations. By employing this method, a conversion factor was established for various types of dust particles. For this purpose, we mixed the known proportion of K\(_2\)SO\(_4\) and dust particles of concern and thus obtained relationship curves between the integrated absorbance of DRIFTS (i.e., diffuse reflectance Fourier transformed infrared spectroscopy) sulfate bands and corresponding theoretical sulfate content through linear fitting (Fig. S4). In total, 10 time points during the heterogeneous reaction were applied for these kinetic calculations.

2.1.2 Preparation for clay and dust membranes and investigation of sulfate formation on those authentic particles during the daytime and nighttime

Each particle suspension for Arizona test dust (ATD: 2.5 mg per 0.5 mL), Imt-2 (Ilite, Montana, USA; 10 mg per 0.5 mL) and K-GA (Kaolin, Georgia, USA; 10 mg per 0.5 mL) was first dispersed into water through an ultrasonic bath for 5 min. After that, sample suspensions were transformed onto the cleaned round quartz films (\( d = 2 \) cm) using a pipette and subsequently sent to the infrared drying oven for 10 min to prepare the dust membranes. Once taken out from the oven, samples were quickly sealed into a desiccator and cooled down to room temperature before starting the experiments. A membrane sample was then mounted at the center of the reaction chamber (the top half is made of quartz, and the bottom half is made of Teflon). Before each set of experiments, a gas flow (dry air) of 300 mL min\(^{-1}\) was introduced to the chamber for 5 min where a prepared membrane sample was installed. Afterward, samples were exposed to a 4.91 \( \times \) 10\(^{14} \) molec. cm\(^{-3} \) SO\(_2\) (+2.46 \( \times \) 10\(^{18} \) CO\(_2\) when necessary) / N\(_2\) + O\(_2\) mixture in the absence and presence of irradiation (light intensity (\( I \)) of 30 mW cm\(^{-2} \)) for 15 min before the sample was transferred to a beaker (scale of 10 mL), with 2 mL of 2% vol. isopropanol leaching solution, then ions were extracted in the ultrasonic tank for 5 min using 0.22 µm PTFE membrane filter, followed by sending the sample into IC. Noting that dust and clay particles possess a considerable volume of sulfate ion in the background, we thus measured the background ions for each batch of synthesized dust particle membranes following the procedures described above. All data demonstrated in Fig. 6 were obtained after the subtraction of background ions.

2.1.3 Determination of gas-phase reactive oxygen species (ROS) production in the flow cell reactor

To measure the concentration of ROS released from TiO\(_2\) particles in various reaction systems, an experimental approach using the probe molecule aniline was applied in this study. This is because compound aniline is reported to react rapidly with -OH radicals and CO\(_3^2^-\) radicals, which are also evidenced to be two major active ROS species that are responsible for the SO\(_2\) oxidation over mineral dust particles. The method applied in this study was almost implemented in the same way as that of the previous study (Behrman, 2018), with a slight modification. Briefly, the degradation rate of aniline in various reaction systems was monitored through high-performance liquid chromatography (HPLC; LC-10AD, Shimadzu, Japan). A Zorbax StableBond (SB) C18 (4.6 mm \( \times \) 150 mm, 5 µm) reverse-phase column at 25\(^\circ\) was used with a UV detector at 236 nm to measure the aniline concentration. The mobile phase consisted of an acetonitrile/water ratio of 55:45 (V/V), with a flow rate of 1 mL min\(^{-1}\).

The TiO\(_2\) suspension (5 mg TiO\(_2\) per 100 µL deionized water) was deposited onto the glass substrate (0.13–0.17 mm in thickness) using a pipette and then dried in an oven for 10 min to obtain a TiO\(_2\)-coated film. Dilute aniline solution, using a 67 mM phosphate buffer solution (pH of 7.0), was prepared and placed below the TiO\(_2\)-coated film, with an intervening gap between the TiO\(_2\) film and solution surface around 2 mm (Fig. 1). This short distance essentially guarantees that gaseous ROS (e.g., -OH radicals or CO\(_3^2^-\) radicals) will diffuse and react with aniline molecule (Rodriguez et al., 2013).

In the reaction system containing a TiO\(_2\) film upon irradiation (the UV wavelength of 310 nm) in the presence of humidified air (RH of 30%), when operated in a continuous mode, the overall degradation rate of the probe molecules in the presence of the TiO\(_2\) film can be described by Eq. (4), as follows (Wang et al., 2004):

\[
k_{obs} = \frac{d[An]}{dr} = r_A + r_U + r_{ROS} = r_A + r_U + k_{ROS, AN}[ROS][An].
\]
where \( k_{\text{obs}} \) is the observed degradation rate of aniline, [An] is the concentration of aniline, denoted as [An] hereafter, and \( r_A, \ r_U \), and \( r_{\text{ROS}} \) stand for aniline removal rates resulting from air stripping, UV photolysis, and ROS oxidation. \( k_{\text{ROS}}, \text{An} \) are the overall second-order reaction rate constants for An with ROS.

Reference experiments without the introduction of ROS were also conducted to measure \( r_A + r_U \) in each reaction system. Upon irradiation, the dust proxy TiO\(_2\) produces hole-electron pairs, further forming \( \cdot \text{OH} \) radicals and superoxide radicals (\( \text{O}_2^{-}\)) in the presence of absorbed water and oxygen molecules. Thus, an experiment using \( \text{N}_2 \) was adopted to investigate the role of \( \text{O}_2^{-} \) in consuming aniline. As illustrated in Fig. 7c, a slight change in the degradation rate of aniline after stripping oxygen from the air, indicating that \( \text{O}_2^{-} \) shows quite a smaller contribution than \( \cdot \text{OH} \). This result agrees well with the finding reported by Durán et al. (2019), where the removal of \( \text{O}_2^{-} \) by adding benzoquinone (BQ) into TiO\(_2\) suspension results in a negligible change in the An degradation rate.

Taken above, \( \cdot \text{OH} \) radicals are assumed to be the only active ROS that accounts for the An degradation. Hence, the maximum steady concentration of \( \cdot \text{OH} \) radicals can be given by the following equation:

\[
- \frac{d[\text{An}]}{dt} = k_{\exp}[\text{An}] = k_{\cdot \text{OH}, \text{An}}[\cdot \text{OH}]_{\text{ss-max}}[\text{An}].
\] (5)

The integration of Eq. (5) yields the following:

\[
\ln \left[ \frac{[\text{An}]_f}{[\text{An}]_0} \right] = k_{\exp} t
\] (6)

\[
k_{\exp} = k_{\cdot \text{OH}, \text{An}}[\cdot \text{OH}]_{\text{ss-max}}.
\] (7)

Together with the reported second-order rate constant \( (k_{\cdot \text{OH}, \text{An}} = 6.5 \times 10^8 \text{ M}^{-1} \text{s}^{-1}) \) (Samuni et al., 2002), the steady-state \( \cdot \text{OH} \) radical concentration \( [\cdot \text{OH}]_{\text{ss-max}} \) in a buffered An solution can be calculated from Eq. (7). The observed degradation rate constant of \( k_{\exp} \) can be obtained from the slope of the semi-log plot of an An degradation, as illustrated in Eq. (6). The maximum steady-state aqueous concentration of \( \cdot \text{OH} \) supplied by the partitioning process from gas-phase \( \cdot \text{OH} \) was thus estimated to be \( 2.15 \times 10^{-15} \text{ M} \) for the TiO\(_2\) + Air system.

When \( \text{CO}_2 \) (400 ppm, parts per million; atmospherically relevant concentration) is introduced into a flow cell chamber, an increased degradation rate of An is seen, which is very likely to be the generation of active carbonate radicals (Fig. 9). Similar to the method we adopted for the estimation of \( [\cdot \text{OH}]_{\text{ss-max}} \), reference experiments were conducted to determine the rates for air stripping and UV photolysis processes in the TiO\(_2\) + Air + \( \text{CO}_2 \) system. In the next step, we quenched the hydroxyl radicals by adding tertiary butanol (TBA). This is because it reacts rapidly with hydroxyl radicals (Li et al., 2020) \( k_{\cdot \text{OH}, \text{TBA}} = (6 \times 10^8 \text{ M}^{-1} \text{s}^{-1}) \), while showing a rather low reaction rate with carbonate radicals (Liu et al., 2015) \( k_{\cdot \text{CO}_3^{-}, \text{TBA}} < 1.6 \times 10^7 \text{ M}^{-1} \text{s}^{-1} \). Subsequently, we determined \( [\cdot \text{CO}_3^{-}]_{\text{ss}} \) using the previous protocol (Huang and Mabury, 2000) with known \( k_{\cdot \text{CO}_3^{-}, \text{AN}} \) \( (5.4 \times 10^8 \text{ M}^{-1} \text{s}^{-1}) \) (Wojnarovits et al., 2020). In the extreme case, assuming that all hydroxyl radical ions were fully trapped by absorbed and dissolved HCO\(_3^-\) / CO\(_3^{2-}\), the maximum steady-state \( \cdot \text{CO}_3^{-} \) concentration was determined to be \( 1.39 \times 10^{-13} \text{ M} \) for the TiO\(_2\) + Air + \( \text{CO}_2 \) system, matching well with an earlier study where the concentration of CO\(_3^{2-}\) is reported to be 2 orders of magnitudes more than \( \cdot \text{OH} \) over the water surface (Sulzberger et al., 1997).

2.2 Quantum chemical calculation

We employed density functional theory (DFT) calculations in the term of the single electron transfer (SET) process using the Gaussian 09 package to investigate this novel route (detailed in Texts S11 and S20 in the Supplement).

2.3 Field observations

2.3.1 Sampling

Our sampling for atmospheric particle matter was launched on the roof of the Department of Environmental Science and Engineering, Fudan University (Jiangwan Campus, 31.340661° N, 121.506747° E, 16 km away from the city downtown). More geographical information for sampling has been described in detail elsewhere (Y. Liu et al., 2020). Observations for the water-soluble ionic components of particulate matter were performed using an eight-stage non-viable cascade-impactor-type sampler (Tisch Environmental Inc., USA), the size gradients of which are in the sequence of \(<0.43, 0.43–0.65, 0.65–1.1, 1.1–2.1, 2.1–3, 3.3–4.7, 4.7–5.8, 5.8–9.0, \) and \(~9.0 \mu\text{m} \). These sizes represent the effective cutoff diameter at each level for unit density spherical particles. In our sampling, the estimations for (bi)carbontate ions in atmospheric particulate matter is conducted for the initial four stages \(~3.3 \mu\text{m} \). Atmospheric airflow from the head was maintained at the constant rate of 28.3 L min\(^{-1}\) to meet the operation criterion required for the Anderson-type sampler. Quartz filters (81 mm in diameter, Whatman, GE Healthcare, UK) were applied for the sampling, and the
membranes were rinsed with ultrapure water (electrical resistivity of 18.2 MΩ) no fewer than three times; that is, they were kept in the ultrasonic cleaning tank for 40 min, then rinsed with ultrapure water twice, before being sent into the infrared drying oven, followed by packing them in aluminum foil prior to the field sampling. We carried out this procedure to eliminate the water-soluble background ions as much as possible and to ensure the balance calculation for (bi)carbonate ions. Additionally, following the same aforementioned pretreatments, we measured the background concentrations of ions in blank membranes. We separated our daily samplings into two periods of 11 h each to give an insight into the influence of potential photoinduced reactions on secondary sulfate formation in the atmosphere.

### 2.3.2 (Bi)carbonate estimation

The concentration of bi(carbonate) ions were estimated following the protocol reported in the early works (Zhang et al., 2011; Palmer and Cherry, 1984; Fang et al., 2021; Y. Liu et al., 2020). The following two assumptions were made to simplify the estimation for bi(carbonate) concentrations within the system: (a) dominating cations in each system are H⁺, Li⁺, Na⁺, NH₄⁺, K⁺, Mg²⁺, and Ca²⁺, whereas those transition and heavy metal ions were out of consideration, considering their limited contents of dissolve cations in the atmospheric particulate matter. (b) There were three typical organic acid ions (CH₃COO⁻, COOH⁻, and C₂O₄²⁻, which are major soluble organic acid ions in the atmosphere) taken into account for the ionization balance and the rest of the charge gap in each system was assumed to originate from (bi)carbonate ions. Then we established balance equations (Eqs. 8–11) for each sample on the basis of charge conservation and ionization equilibrium constant of carbonic acid (K₁ = 4.47 × 10⁻⁷ and K₂ = 4.69 × 10⁻¹¹ at 273 K).

\[
\begin{align*}
[H⁺] + [Li⁺] + [Na⁺] + [NH₄⁺] + [K⁺] + 2[Cr²⁺] \\
+ 2[Ca²⁺] = & [OH⁻] + [F⁻] + [CH₃COO⁻] + [COOH⁻] \\
+ [NO₂⁻] + [Cl⁻] + [NO₃⁻] + 3[PO₄³⁻] \\
+ 2[SO₄²⁻] + 2[C₂O₄²⁻] + [HCO₃⁻] + 2[CO₃²⁻]
\end{align*}
\] (8)

\[
[HCO₃⁻] = \frac{K₁[H₂CO₃]}{[H⁺]}
\] (9)

\[
[CO₃²⁻] = \frac{K₁K₂[H₂CO₃]}{[H⁺]²}
\] (10)

It is worth mentioning that [X] is referring to the charge concentration for ions (coulomb M). Additionally, the temperature factor was also considered to correct the equilibrium constant for (bi)carbonate ions using the following equation:

\[
\ln \frac{K′}{K} = \frac{-ΔH}{RT} \left(\frac{1}{T′} - \frac{1}{T}ight),
\] (11)

where ΔH is the temperature variation (K), R is the ideal gas constant (8.31451 J mol⁻¹ K⁻¹), and T is the temperature (K) during pH measurements. We then solved those equations to obtain a series of [HCO₃⁻], which were eventually corrected by subtracting blank values.

### 2.4 Other measurements and analysis

In addition to the above descriptions of measurements and calculation methodologies, more information, including the heterogenous reaction setup, kinetics reaction order determination, pretreatment of mineral dust, DRIFTS, IC, Raman measurements and analysis, etc., is available in Texts S1–S13 in the Supplement.

### 3 Results and discussion

#### 3.1 Accelerated sulfate production in the presence of carbonate

The physicochemical properties of the employed mineral dust proxies, including Brunauer–Emmett–Teller (BET) surface area, crystal phase, and structure, were first characterized (Fig. S1), consistent with early studies (Su et al., 2008; Balachandran and Eror, 1982; Shang et al., 2010), with an additional discussion in Text S14. The spectral irradiance of the solar simulator applied in the present study is well covered by natural sunlight (Fig. S2), and we tried as much as possible to have the experimental results from the lab simulate the real atmosphere. Upon solar irradiation under RH of 30 % SO₂ and 50 wt % TiO₂ and 50 wt % CaCO₃, measured by IC, is significantly enhanced by 7 times and 23 times compared to that of pristine TiO₂ and CaCO₃ (Fig. 2a), respectively. In stark contrast, there is a negligible increase in sulfate production detected in the TiO₂ + CaCO₃ mixture relative to that of pristine TiO₂ and CaCO₃ in dark experiments (Fig. S3). Great discrepancies in sulfate production over TiO₂ + CaCO₃ particles between dark and light experiments suggest that carbonate salt may play a different role in these two scenarios. However, the alkalinity of carbonate salt favors SO₂ adsorption (Al-Hosney and Grassian, 2005; Yu et al., 2018), and the photo-oxidation process assisted by TiO₂ particles is able to strengthen the oxidation efficiency of adsorbed SO₂ (Shang et al., 2010; Chen et al., 2012), which is a plausible explanation for the increased sulfate production over TiO₂ + CaCO₃ particles. Following this speculation, two types of mixtures, TiO₂ + CaCO₃ and TiO₂ + CaO, were employed. In the dark experiments (Fig. 2b), both TiO₂ + CaO and TiO₂ + CaCO₃
almost yield an identical concentration of sulfite and sulfate. On the contrary, TiO$_2$ + CaCO$_3$ particles produce nearly 2 times more sulfate than TiO$_2$ + CaO particles once irradiated, along with a sharp decrease in S(IV) species on the surface of TiO$_2$ + CaCO$_3$ surfaces. Besides, CaCO$_3$ tends to show relatively humble physical properties, including BET surface area, surface pH, and solubility, etc., relative to CaO (see the detailed discussion in Text S15). The above results allow us to assert that the carbonate-containing system contains an alternative important mechanism for sulfate formation beyond the production of an alkaline environment (additional discussion available in Text S16). Figure 2c and d illustrate that the DRIFTS features of S(IV) and S(VI) species (Nanayakkara et al., 2014; Wu et al., 2011) increase beyond the production of an alkaline environment (ad- ditional discussion available in Text S16). Figure 2c and d illustrate that the DRIFTS features of S(IV) and S(VI) species (Nanayakkara et al., 2014; Wu et al., 2011) increase beyond the production of an alkaline environment (additional discussion available in Text S16). Figure 2c and d illustrate that the DRIFTS features of S(IV) and S(VI) species (Nanayakkara et al., 2014; Wu et al., 2011) increase beyond the production of an alkaline environment (additional discussion available in Text S16). Figure 2c and d illustrate that the DRIFTS features of S(IV) and S(VI) species (Nanayakkara et al., 2014; Wu et al., 2011) increase beyond the production of an alkaline environment (additional discussion available in Text S16). Figure 2c and d illustrate that the DRIFTS features of S(IV) and S(VI) species (Nanayakkara et al., 2014; Wu et al., 2011) increase beyond the production of an alkaline environment (additional discussion available in Text S16).
Figure 2. (a) Sulfate or sulfite concentration quantified by IC on TiO$_2$, CaCO$_3$, and TiO$_2$ + CaCO$_3$ particles (wt/wt = 50/50) after exposure to gaseous SO$_2$ under irradiation for 60 min. (b) Sulfate and sulfite concentration quantified by IC on mineral dust particles of concern after exposure to SO$_2$ under dark or irradiation for 20 min. In situ DRIFTS of S(IV) and S(VI) production on theoretical and experimental TiO$_2$ + CaCO$_3$ mixtures (wt/wt = 50/50) upon irradiation for 90 min. All spectra were processed by the Kubelka–Munk (K–M) algorithm. Note that the production of sulfur species in theoretical TiO$_2$ + CaCO$_3$ mixtures refers to 0.5 × K–M bands of sulfur species of TiO$_2$ − 0.5 × K–M bands of sulfur species of CaCO$_3$, while those for experimental TiO$_2$ + CaCO$_3$ mixtures refer to 1 × K–M bands of sulfur species of TiO$_2$ + CaCO$_3$ mixtures (wt/wt = 50/50). Reaction conditions are RH of 30%, light intensity (I) of 30 mW cm$^{-2}$, total flow rate of 52.5 mL min$^{-1}$, and SO$_2$ of 2.21 × 10$^{14}$ molec. cm$^{-3}$.

Table 1. Chemical compositions of mineral dust simulants.

<table>
<thead>
<tr>
<th></th>
<th>SiO$_2$ (wt %)$^b$</th>
<th>Al$_2$O$_3$ (wt %)$^b$</th>
<th>CaCO$_3$ (wt %)$^{a,b}$</th>
<th>TiO$_2$ (wt %)$^b$</th>
<th>Ca : Al (Fe : Al)$^c$</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO$_2$ + Al$_2$O$_3$</td>
<td>89.46</td>
<td>10.54</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>SiO$_2$ + Al$_2$O$_3$ + CaCO$_3$</td>
<td>82.47</td>
<td>9.72</td>
<td>7.81</td>
<td>–</td>
<td>0.73 (-)</td>
</tr>
<tr>
<td>SiO$_2$ + Al$_2$O$_3$ + TiO$_2$</td>
<td>88.42</td>
<td>10.42</td>
<td>–</td>
<td>1.15</td>
<td>–</td>
</tr>
<tr>
<td>SiO$_2$ + Al$_2$O$_3$ + TiO$_2$ + CaCO$_3$</td>
<td>81.59</td>
<td>9.62</td>
<td>7.73</td>
<td>1.06</td>
<td>0.73 (-)</td>
</tr>
<tr>
<td>ATD</td>
<td>71.27</td>
<td>8.4</td>
<td>–</td>
<td>0.93</td>
<td>0.73 (0.17)</td>
</tr>
</tbody>
</table>

$^a$ In the present study, CaCO$_3$ was taken as representative of alkaline earth metal oxide in our proxies for the authentic dust.
$^b$ The mass ratio of four components (if any) in the simulants were controlled in the ratio of SiO$_2$:Al$_2$O$_3$:CaCO$_3$:TiO$_2$ = 81.59:9.62:7.73:1.06. For instance, the mass ratio of SiO$_2$ to Al$_2$O$_3$ is 81.59 : 9.62 in the SiO$_2$ + Al$_2$O$_3$ simulant, whereas it is 81.59 : 9.62 : 7.73 in the SiO$_2$ + Al$_2$O$_3$ + CaCO$_3$ mixture. The ratios of each component are derived from the EDS mapping analysis of ATD dust particles. $^c$ This column refers to the molar ratio.
Figure 3. (a) Energy dispersive spectroscopy (EDS) mapping of sulfur. (b) A selected HRTEM (high-resolution transmission electron microscopy) region containing a high density of sulfur for further observation is shown, and the red rectangle refers to the region shown in panel (c). (c) The HRTEM image in high resolution with lattice fringes and (d) corresponding fast Fourier transform (FFT) power spectra, lattice indexing, and (1–6) the inverse FFT analysis of lattice signal is shown in panel (d). In panel (e), the term C-SO$_4^{2-}$ stands for crystalline SO$_2^-$, i.e., CaSO$_4$ and Ti(SO$_4$)$_2$. Particles employed for the HRTEM measurement refer to TiO$_2$ (50 wt %) + CaCO$_3$ (50 wt %) mixture particles upon exposure to the 4.42 × 10$^{14}$ molec. cm$^{-3}$ SO$_2$/N$_2$+O$_2$ for 60 min, while other reaction conditions are as same as those of the above sulfate quantification experiments in Fig. 1. Reaction conditions are RH of 30 %, light intensity (I) of 30 mW cm$^{-2}$, and total flow rate of 52.5 mL min$^{-1}$.

Figure 4. Sulfate concentration quantified by IC. Sulfate concentration was measured by IC on mineral dust simulants after exposure to gaseous SO$_2$ (2.46 × 10$^{14}$ molec. cm$^{-3}$) under irradiation. Noting that SiO$_2$ : Al$_2$O$_3$ : CaCO$_3$ : TiO$_2$ refers to the mass fraction ratios of the components in simulants. Experiments were all conducted at an RH of 30 % and light intensity (I) of 30 mW cm$^{-2}$.

reported to produce electron–hole pairs under solar irradiation (Li et al., 2019), thus likely involving the reaction channel considered in this work (see a detailed discussion in a later section). Similar to the protocol applied for synthesizing the TiO$_2$ + CaCO$_3$ mixture, αFe$_2$O$_3$ + CaCO$_3$ are prepared accordingly. In Fig. S9a, our results show that αFe$_2$O$_3$ cannot trigger fast SO$_2$ oxidation in the presence of carbonate ions upon irradiation, which is distinguished from the results we derived from the TiO$_2$ + CaCO$_3$ mixture. This can be explained by the fact that Fe$_2$O$_3$ shows a lower redox activity.
relative to TiO$_2$ (Fig. S9b), where its strong redox capability essentially enables photoinduced electrons and holes to produce O$_2^-$ and -OH radical ions. In stark contrast, the valence band and conduct band of Fe$_2$O$_3$ lie at $-0.18$ and at 1.68 V vs. NHE (pH = 7), lower than the redox potential required for generating an O$_2^-$, -OH and carbonate-containing ROS (Li et al., 2016). Hence, no promoted sulfate production is seen for $\alpha$Fe$_2$O$_3$ + CaCO$_3$ particles under irradiation. We also note the inconsistency between our study and the previous literature with regard to the response of SO$_2$ oxidation to solar irradiation over $\alpha$Fe$_2$O$_3$ particles, which has been interpreted in Text S17.

Overall, we show that, upon irradiation, the atmospherically relevant content of TiO$_2$ (nearly 1%) in mineral dust simulants is able to interact with carbonate ions to launch an increased sulfate production, which is beyond the conventional regime of alkaline neutralization of H$_2$SO$_4$. Unlike TiO$_2$, $\alpha$Fe$_2$O$_3$ lacks the ability to initiate fast SO$_2$ oxidation by generating carbonate-containing ROS due to its limited photochemical activity, although ferric chemistry is important in secondary sulfate formation in the atmosphere (Sullivan et al., 2007; Yermakov and Purmal, 2003).

### 3.2 Accelerated sulfate production in the presence of CO$_2$

Atmospheric CO$_2$ is also an important source of (bi)carbonate. Its influence on photochemical SO$_2$ uptake on mineral dust was thus studied. Distinguishing S(VI) from S(IV) species over TiO$_2$ particles relies upon the position of the infrared (IR) bands, according to the assignment of the previous literature (Nanayakkara et al., 2014), and S(VI) and S(IV) build up as heterogeneous reactions occur in all cases (Fig. 5). In the presence of atmospherically relevant CO$_2$ ($9.83 \times 10^{15}$ molec. cm$^{-3}$), the sulfate yield was increased under irradiation as compared to the CO$_2$-free case (Fig. 5a and b). We cautiously examined the net effect of formed (bi)carbonate on sulfate production by time-resolved DRIFTS spectra (Fig. 5c and d) using dark experiments. CO$_2$ suppresses both S(IV) and S(VI) products in the dark probably because of the competitive adsorption effect, as we observed over Al$_2$O$_3$ particles (Y. Liu et al., 2020).

The above observation leads us to speculate that the active intermediate derived from (bi)carbonate species upon irradiation is a plausibe force to drive rapid sulfate formation. Besides, a nearly 50% increase in the SO$_2$ uptake coefficient is observed for the mineral dust proxy TiO$_2$ after being exposed to the 9.83 $\times$ 10$^{15}$ molec. cm$^{-3}$ (400 ppm) CO$_2$ + SO$_2$/N$_2$ + O$_2$ mixture (Table S1). The pseudo first reaction order (1.13) was also determined in the selected concentration range of 400–20 000 ppm (Fig. S5b), which satisfies the prerequisite for the uptake coefficients derived from laboratory chambers potentially being generalized to the atmosphere condition, as we expounded in the early context.

<table>
<thead>
<tr>
<th>Minerals</th>
<th>ATD (%)$^a$</th>
<th>IMt-2 (%)$^a$</th>
<th>K-Ga-2 (%)$^a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO$_2$</td>
<td>78.11</td>
<td>59.57</td>
<td>56.93</td>
</tr>
<tr>
<td>Al$_2$O$_3$</td>
<td>7.19</td>
<td>19.47</td>
<td>37.49</td>
</tr>
<tr>
<td>Fe$_2$O$_3$</td>
<td>2.57</td>
<td>7.95</td>
<td>1.81</td>
</tr>
<tr>
<td>FeO</td>
<td>n.d.$^b$</td>
<td>0.05</td>
<td>0.15</td>
</tr>
<tr>
<td>MgO</td>
<td>1.22</td>
<td>2.42</td>
<td>0.03</td>
</tr>
<tr>
<td>CaO</td>
<td>3.03</td>
<td>0.37</td>
<td>0.01</td>
</tr>
<tr>
<td>Na$_2$O</td>
<td>1.39</td>
<td>0.08</td>
<td>n.d.$^b$</td>
</tr>
<tr>
<td>K$_2$O</td>
<td>2.06</td>
<td>8.72</td>
<td>0.06</td>
</tr>
<tr>
<td>TiO$_2$</td>
<td>0.46</td>
<td>0.99</td>
<td>3.43</td>
</tr>
<tr>
<td>P$_2$O$_5$</td>
<td>0.10</td>
<td>0.07</td>
<td>0.05</td>
</tr>
<tr>
<td>MnO</td>
<td>0.04</td>
<td>0.03</td>
<td>n.d.$^b$</td>
</tr>
<tr>
<td>S</td>
<td>n.d.$^b$</td>
<td>0.03</td>
<td>0.02</td>
</tr>
<tr>
<td>Total</td>
<td>99.21</td>
<td>99.75</td>
<td>99.98</td>
</tr>
<tr>
<td>Total AE$^c$</td>
<td>7.70</td>
<td>11.59</td>
<td>0.10</td>
</tr>
</tbody>
</table>

$^a$ Chemical compositions of the dust and clays were determined by XRF (X-ray fluorescence) results. Note that $^b$ n.d. refers to not detected and that $^c$ AE refers to alkaline earth metal oxide.

As another step toward a real scenario in the atmosphere, experimental trials employing authentic mineral dust particles, i.e., K-Ga-2 (Kaolin, Georgia, USA), Arizona test dust (ATD), and IMt-2 (Illite, Montana, USA) clays were implemented, with the component analysis results shown in Table 2. In Fig. 6, the pronounced increase in sulfate yield (a nearly 100% increase in sulfate production in the CO$_2$-involved case under irradiation) is best seen in K-Ga-2 clay (Fig. 6a). The promotional effect of CO$_2$ on sulfate formation under irradiation, nonetheless, is less evident for IMt-2 (the content of TiO$_2$ $\approx$ 0.99%) and ATD (the content of TiO$_2$ $\approx$ 0.46%) as compared to K-Ga-2 particles. This may correlate to their higher mass fraction of alkaline earth metal oxide (denoted as AE), which enables dust particles to possess a substantial number of (bi)carbonate species within the natural environment where they have experienced long-term exposure to atmospheric CO$_2$ during regional transport. Therefore, the aforementioned synergetic effect takes effect over IMt-2 and ATD particles even without exposure to CO$_2$, presumably due to the presence of the abundant alkaline carbonate formed, and a relatively moderate increase in sulfate production was thus observed. On the other hand, TiO$_2$ content is not necessarily an accurate predictor of photoactivity; the content and proportion of the active phase of TiO$_2$ in K-Ga-2 altogether contribute to a more pronounced increase in sulfate production relative to the other two clays (see the detailed discussion in Text S18).

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3.3 Reaction mechanism

The heterogeneous reaction of SO$_2$ on dust particles in the atmosphere is a complicated process, covering a series of reactions taking place in both homogeneous and heterogeneous ways. At a sufficiently low RH condition (normally below 10 % RH), water readily dissociates on the surface of metal oxide under ambient atmospheric conditions, where the metal oxide surface is terminated by hydroxyl groups that hydrogen bond to adsorbed water molecules (Cwiertny et al., 2008). In this case, SO$_2$ oxidation over dust particles is dominated by the reaction regime where the resulting hydroxyl groups react with gaseous SO$_2$ to form adsorbed S(IV)$_{ad}$ species. Afterward, S(IV)$_{ad}$ will be oxidized by oxidants in the atmosphere or photoinduced active intermediates produced from the dust surface upon irradiation. As the RH increases beyond 10 %–15 %, multilayer water coverage occurs, reaching approximately two monolayers at RH of 30 % (Mogili et al., 2006). Under these circumstances, the amount of water adsorbed onto the surface of the dust particles is be-
Mineral dust + hv → h⁺ + e⁻  

(R1)

h⁺ + CO₃²⁻ → CO₃⁻.  

(R2)

In the second pathway, carbonate radicals evolve through the reaction of (bi)carbonate anion with hydroxyl radicals ·OH forming over mineral dust surfaces (Zhang et al., 2015; Reaction R3 and R4).

h⁺ + H₂O_ads → ·OH + H⁺  

(R3)

·OH + HCO₃⁻/CO₃²⁻ → CO₃⁻ + H₂O/OH⁻.  

(R4)

The above assumptions are supported by nanosecond transient absorption spectra (NTAS), in which a signal (∆OD) of a carbonate radical CO₃⁻ at 600 nm (Bhattacharya et al., 1998) only emerges for dust suspension containing (bi)carbonate species (Fig. 7a). An increased degradation rate of aniline observed in TiO₂ suspension due to the presence of carbonate ions produces additional evidence of the formation of active CO₃⁻ ions and the strengthened oxidation capability of TiO₂ (Fig. 7b; see the additional discussion in Text S19). The CO₃⁻-induced chemistry was further evidenced by ·OH scavenging experiments using tertiary butyl alcohol (TBA) and isopropanol (i-PrOH), as they show lower reaction rates with CO₃⁻ (kₐ₉₋,TBA = 4.0 × 10⁴ M⁻¹ s⁻¹ and kₐ₈₋,TBA = 1.6 × 10² M⁻¹ s⁻¹) relative to those with ·OH (k₁₋,PrOH→OH < 1.9 × 10⁹ M⁻¹ s⁻¹ and k₁₋,OH,TBA = 6 × 10⁸ M⁻¹ s⁻¹; Buxton et al., 2009; Liu et al., 2015; Li et al., 2020). TBA dramatically decreases the yield of sulfate on the TiO₂ surface by nearly 70%, with sulfite ions being the dominant sulfur species (Fig. 7c). Meanwhile, a great loss of sulfate yield when TiO₂ suspension was added with i-PrOH (Fig. 7d). This is in strong contrast to the result of a carbonate-involved system, where the reactivity is sustained as carbonate radicals offer an alternative reaction pathway for SO₂ oxidation. This is plausible since the carbonate ions are an excellent ·OH scavenger, and CO₃⁻ becomes the predominant species in a relatively strong alkaline aqueous-like environment in the presence of carbonate salt. This is supported by a previous work (Sun et al., 2016) in which adding 0.1 M of bicarbonate salt into the UV/H₂O₂ system (H₂O₂ = 0.3 mM) was sufficient to suppress ·OH concentration to around 10⁻¹⁵ M, creating a carbonate-radical-dominated regime ([CO₃⁻] = 8.64 × 10⁻¹² M). In our experiments (Fig. 7b), 0.2 M of carbonate salt was employed, and the reaction rate of CO₃⁻ with ·OH is nearly 2 orders of magnitude higher than that of HCO₃⁻, thus giving rise to carbonate radical being the substitute for the hydroxyl radical in the reaction. The above results suggest that ·OH is a major contributor to the sulfate yield on TiO₂ particles in the absence of carbonate ions, while CO₃⁻ ions dominate the SO₂ oxidation over humidified carbonate-containing TiO₂ particles upon irradiation. In addition to experimental investigations, the carbonate radical formation process is proved to be thermodynamically favorable and supported by density functional theory (DFT) calculations (Fig. S10 and Text S20).

On the other hand, early studies (Das, 2001; Neta and Huie, 1985; Chameides and Davis, 1982) agree with the key role of the sulfite radical (SO₃⁻) in rapid sulfate production in an aqueous medium, and the present reaction system creates a localized environment in which the SO₃⁻ can be readily produced from the TiO₂ and S(IV) species upon solar illumination (Salama et al., 1995). Consequently, a probe light of NTAS at wavelength 255 nm (ascribed to the sulfite radical) and 600 nm (ascribed to carbonate radical) were simultaneously monitored (Hayon et al., 1972; Ghalei et al., 2016; Goldstein et al., 2001). A weak signal of sulfite radical was observed in the system of the TiO₂ + Na₂SO₃ suspension under irradiation (Fig. 7e). On the contrary, the sulfite radical signal is strengthened after the introduction of carbonate ions into the TiO₂ + Na₂SO₃ suspension, along with a significant decrease in the signal for carbonate radical. Electron spin resonance (ESR) data (Fig. 7f) further confirm the increase in the SO₃²⁻ after 2 min UV irradiation in the presence of a carbonate ion. Based on the above results, one may deduce that the interplay between carbonate radical and sulfite ions is a crucial step giving rise to the increased SO₃²⁻, which is reported to account for rapid atmospheric sulfate formation through chain propagation reactions that involve SO₄⁻ and SO₂⁻ intermediates (Hung and Hoffmann, 2015; Hung et al., 2018). Additionally, this sulfite radical ion chemistry is believed to drive fast sulfate formation over mineral dust particles as well (Gankanda et al., 2016; Rubasinghe et al., 2010). Nevertheless, there are two possibilities that might explain the aforementioned interaction. One way is through the
Figure 7. (a) Single-wavelength transient absorption spectra of various aqueous solutions. (b) The removal rate of aniline after exposure to airflow under irradiation in the absence and presence of mineral dust particles for 300 s. Reaction conditions are RH of 30 %, light intensity (I) of 30 mW cm\(^{-2}\), and a total flow rate of 52.5 mL min\(^{-1}\). Note that an adequate amount of NaOH was introduced into the TiO\(_2\) suspension system to achieve a pH environment condition identical to that of the TiO\(_2\)+Na\(_2\)CO\(_3\) suspension system. (c) Determination of sulfite and sulfate concentration after exposure to airflow under irradiation in the absence and presence of mineral dust particles for 20 min. Reaction conditions are RH of 30 %, light intensity (I) of 30 mW cm\(^{-2}\), and a total flow rate of 52.5 mL min\(^{-1}\). (d) Sulfate formation change \((\text{SO}_3^{2-})\) is determined by different sulfate concentrations with and without the addition of isopropanol as hydroxyl radical scavenger. (e) The difference in transient absorption kinetics of the sulfite radical and carbonate radical at the various aqueous solutions and their corresponding growth–decay fit curves. A \(A_1\) signal was recorded at 255 and 600 nm after a pulsed 355 nm laser excitation. (f) Electron spin resonance (ESR) spectrometry of \([\text{DMPO–SO}_3^{-}]\) intermediary formed in a solution of TiO\(_2\) (3 mg per 4 mL) + 0.1 M Na\(_2\)SO\(_3\) and TiO\(_2\) (3 mg per 4 mL) + 0.5 M Na\(_2\)CO\(_3\) + 0.1 M Na\(_2\)SO\(_3\). For clarity, the integrated areas of the ESR profiles were also presented for direct comparison. Note that Exp. and Sti. stand for experimental results and corresponding fitting results using isotropic radicals software.

We first examined the oxygen transfer path through \(^{18}\)O isotope labeling experiments. TiO\(_2\) particles were initially exposed to C\(^{16}\)O\(_2\)/N\(_2\) and C\(^{18}\)O\(_2\)/N\(_2\), followed by the exposure of SO\(_2\)/N\(_2\)+O\(_2\) under irradiation (Fig. 8a). A bidentate carbonate band centered at 1573 cm\(^{-1}\) appears after the introduction of C\(^{16}\)O\(_2\)/N\(_2\), while this band shifts to 1558 cm\(^{-1}\) when C\(^{18}\)O\(_2\)/N\(_2\) is introduced, indicating the incorporation of \(^{18}\)O into bidentate carbonate species, in accordance with a previous report (Liao et al., 2002). However, no shift of IR features at 1269, 1219, and 1159 cm\(^{-1}\), assigned to bisulfate species on TiO\(_2\) particles, was observed throughout the reaction. This implies that the oxygen transfer path does not account for the rapid SO\(_2\) oxidation on particles of concern.

In light of the above analysis, the electron transfer might be a plausible pathway to explain the fast oxidation within the reaction system. DFT calculations provide an accessible approach to studying the electron transfer pathway. The result in Fig. 8b illustrates SO\(_3^{2-}\) formation is a SET process of CO\(_3^{2-}\) and SO\(_2^{2-}\), where the O atom in SO\(_2^{2-}\) transfers an electron to the O atom in CO\(_3^{2-}\) to form SO\(_3^{2-}\) and CO\(_2^{-}\). This SET reaction is a thermodynamically favorable process, with the difference in the Gibbs free energy between reactant and product lying at \(-24.09\) kcal mol\(^{-1}\). We note that the insufficient O\(_2\) supply in aqueous media may be an underlying constraint to the proposed CO\(_3^{2-}\)-initiated SO\(_2\) oxida-
Figure 8. (a) In situ DRIFTS of heterogeneous reaction of SO₂ on the TiO₂ particles for 2 and 60 min after being exposed to C₁₆(¹⁸)O₂ / N₂ for 20 min under irradiation. (b) Reaction pathway of the interaction between the carbonate radical (CO₃²⁻) and sulfite (SO₃²⁻) and the (c) interaction between the hydroxyl radical (·OH) and sulfite (SO₃²⁻) through the SET process at the coupled cluster calculations with all single and double excitations (CCSDs) (T)-F12/cc-PVDZ-F12//M06-2X/6-311++G (3df, 3pd) level (where polarized valence double zeta is PVDZ), and ΔG_{SET}^r represents the difference in the Gibbs free energy between the reactant and product. The white, black, yellow, and red spheres represent H, C, S, and O atoms, respectively. In order to visualize the variation in the surface products in oxygen isotope experiments (a), DRIFTS features of these concerned species are highlighted in dark colors.

In this SET process, the electron donor SO₃²⁻ reacts spontaneously with electron acceptor ·OH (Fig. 8c), and the calculated activation free energy barrier ΔG_{SET}^r for this SET reaction is 2.50 kcal mol⁻¹. Hence, the reaction process of ·OH with SO₃²⁻ is diffusion controlled, and the total rate constant k_{SET−2} was calculated to be 7.12 × 10⁹ M⁻¹ s⁻¹. In comparison, the rate constant k_{SET−1} of the diffusion-controlled SET process for CO₃²⁻ and SO₃²⁻ was estimated to be 7.42 × 10¹⁰ M⁻¹ s⁻¹. Despite a slight net increase in the rate, the distinguishable concentration of CO₃²⁻ and ·OH should also be taken into account for the rate comparison in varied reaction paths. To visualize the difference, relative rates were calculated according to Eq. (12), as follows:

\[ r = \frac{v_{CO_3^{2-} + SO_3^{2-}}}{v_{OH + SO_3^{2-}}} = \frac{k_{SET−1} [CO_3^{2-}] [SO_3^{2-}]}{k_{SET−2} [·OH] [SO_3^{2-}]}, \]  

where \( r \) is the ratio of two reaction rates, and \([CO_3^{2-}]\), \([SO_3^{2-}]\), and \([·OH]\) refer to the concentration of corresponding reactants. In the literature, it is suggested that the concentration of carbonate radicals is able to show 2 orders of magnitude higher than that of hydroxyl radicals at the surface of the water under solar irradiation (Chandrasekaran and Thomas, 1983; Goldstein et al., 2001; Sulzberger et al., 1997), which is consistent with the concentration gap between carbonate radicals and hydroxyl radicals through a partitioning process from the gas phase determined in our reaction system (Fig. 9). While the net concentrations of carbonate and hydroxyl radicals in the water layers of humified particles are very likely to be different from those found in the bulk aqueous media, the concentration inputs of two radicals with the gap of 2 orders of magnitude could somehow reflect the relative contribution of carbonate radicals and hydroxyl radicals to sulfate production, based on the literature results and our experimental trails. The concentrations of CO₃²⁻ and ·OH were set in the range from 1.0 × 10⁻¹⁰ to 1.0 × 10⁻¹² mol L⁻¹ and from 1.0 × 10⁻¹¹ to 1.0 × 10⁻¹⁴ mol L⁻¹ (Sulzberger et al., 1997), and the \( r \) value could thus reach to 1.04 × 10³ at most (Fig. S11). As a result, we speculate that the formation pathway of SO₃²⁻ via an interaction between CO₃²⁻ and SO₃²⁻ is a more efficient route that corresponds well with our experimental results.

In addition to the pathway launched by photo-generated holes, the sink of photo-generated electrons is also considered. In our reaction system, O₂ is thought to be an electron...
Figure 9. The degradation rate of aniline after exposure to air flow under irradiation in the absence (a) and presence (b) of CO₂ over mineral dust proxy particles TiO₂ as a function of the reaction time. Reaction conditions are RH of 30 %, light intensity (I) of 30 mW cm⁻², and a total flow rate of 52.5 mL min⁻¹.

trap and to produce the superoxide radical ions (O₂⁻), which are reported to play a non-negligible role in sulfate formation (Shang et al., 2010) and should be taken into account to give a whole picture of reaction scheme in triggering sulfate formation on the surface of TiO₂-containing mineral dust particles. p-benzoquinone is a commonly used O₂⁻ scavenger for trapping the O₂⁻ radical ions (Yan et al., 2018). Our data show that adding an excess amount of p-benzoquinone into TiO₂ particles reduces the sulfate yield by 32 %, along with the appearance of sulfite ions over TiO₂ particles upon exposure to SO₂ (Fig. 7c). Notably, the decrease in sulfate yield by around 30 % in the presence of O₂⁻ scavenger p-benzoquinone is almost complementary to that added with -OH scavenger using TBA (70 %), pointing toward a minor sulfate formation pathway contributed by O₂⁻ relative to the major pathway by CO₃²⁻ when carbonate ions are presented to efficiently capture -OH ions. Following the work of Shang et al. (2010), O₂⁻-involved SO₂ oxidation can be given as Reactions (R10)–(R12), as follows:

\[ e^- + O_2 \rightarrow O_2^- \]  \quad (R10)

\[ SO_2 + O_2^- \rightarrow SO_3 + O^-^- \]  \quad (R11)

\[ SO_3 + H_2O \rightarrow H_2SO_4, \]  \quad (R12)

where intermediate SO₃, formed via the interaction between SO₂ and O₂⁻, subsequently couples with water molecules to produce sulfate species as a final product. pH is an important factor within aqueous chemical reaction processes and is preferred to alter the dominant regime for sulfate production. Yet, so far, adjusting the pH of particle surfaces is quite tough, and exploring the role of dust surface pH in the reactivity of CO₃²⁻ is not easily achieved. Notwithstanding, the increase in pH in TiO₂ suspension was observed to promote the production of CO₃²⁻, further strengthening the oxidation capability of dust particles (Fig. 7b). In contrast, decreasing pH is expected to reduce the yield of CO₃²⁻ since the reaction rate of CO₃²⁻ with -OH is nearly 2 orders of magnitude higher than that with HCO₃⁻. On this basis, a question arises as to whether the surface pH of mineral dust can be sustained to maintain the fast SO₂ oxidation triggered by CO₃²⁻ in the typical lifespan of mineral dust.

Considering that the SO₂ concentration employed in this work is higher than that in the real atmosphere, the concept of equivalent exposure time is introduced to evaluate the influence of pH on the SO₂ oxidation pathway initiated by CO₃²⁻ (see a more detailed discussion on determining equivalent exposure time in Text S22). The heterogeneous sulfate production over TiO₂ and TiO₂ + CaCO₃ particles versus equivalent exposure time were plotted (Fig. S12). Clearly, the sulfate yield builds up steadily during the 2-week equivalent exposure time, suggesting that the regime of CO₃²⁻-initiated SO₂ oxidation over TiO₂ and TiO₂ + CaCO₃ particles is slightly affected by the possible decrease in surface pH because of the accumulation of sulfate production over the entire reaction course. In the atmosphere, the lifetime of mineral dust particles ranges from several days to weeks (Bauer and Koch, 2005), and the equivalent exposure time considered in this study (nearly 2 weeks) falls right within the characteristic lifespan range of mineral dust particles. Besides, 20 ppb is assumed to be an atmospherically relevant concentration to calculate equivalent exposure time in this study, whereas even low SO₂ concentrations (several or a few tens ppb of SO₂) were monitored in the field observation (He et al., 2014; Watanabe et al., 2020). Therefore, the reduction of dust surfaces pH would be more moderate than what we now consider, and even a little influence of surface pH on our proposed reaction scheme would have an impact. Therefore, persistent growth of sulfate shows a negligible effect on the CO₃²⁻-initiated SO₂ oxidation scheme proposed in this work.
Additionally, dust particles are reported to eject the radical ions from the surface under solar light irradiation, severing as an underlying pathway for sulfate aerosol formation in the atmosphere (Chen et al., 2021; Dupart et al., 2012) and is described as follows:

\[
\text{Mineral Dust} + h\nu \rightarrow \text{ROS (g)} \quad \text{(R13)}
\]
\[
\text{ROS (g) + humidified Air + SO}_2 \rightarrow \text{Sulfate(g)}, \quad \text{(R14)}
\]

where ROS (g) stands for the active intermediates in the gas phase. Over 400 ppm of CO$_2$ is universal in the atmosphere, and it is expected to form (bi)carbonate ions once enters the quasi-liquid layer of humified particles. Bi(carbonate) ions are then prone to reacting with hydroxyl radical ions to form carbonate radicals. Following this line of reasoning, we attempt to monitor the plausible gas ROS species that are formed in the presence of CO$_2$ (see the detailed discussion in Sect. 2).

When CO$_2$ (atmospherically relevant concentration) is introduced into the homemade flow chamber with the intervening gap between the TiO$_2$-coated film and probe molecule solution fixing at nearly 2 mm, the short distance guarantees that gaseous ROS will diffuse and react with aniline (Rodriguez et al., 2013). An increased degradation rate of this probe molecule was seen, which can be speculated to be the generation of active carbonate radical ions (Fig. 9). Aqueous CO$_3^{2-}$ is believed to be supplied by the partitioning processes from CO$_3^{3-}$ (g) that come from humified dust particles, and its maximum steady-state concentration was determined to be $1.39 \times 10^{-13}$ M for TiO$_2$ + Air + CO$_2$ system, which is over 1.8 orders of magnitudes higher than that of $\cdot$OH for the TiO$_2$ + Air system ($2.15 \times 10^{-15}$ M). This observation matches with an earlier study, where the concentration of a carbonate radical can be nearly 2 orders of magnitudes greater than $\cdot$OH over the water surface (Sulzberger et al., 1997).

Overall, the above results suggest that the photochemistry that involves carbonate ions, more precisely CO$_3^{3-}$ radicals, increases sulfate production. This finding broadens the prevailing view that the acceleration of SO$_2$ oxidation over the carbonate salt is merely due to the favorable neutralization of H$_2$SO$_4$ over an alkaline surface. Worth noting is that upon irradiation the active component TiO$_2$ in mineral dust produces carbonate radicals in the gas phase when CO$_2$ presents, therefore potentially promoting sulfate aerosol formation in the atmosphere. Overall, it could be speculated that (bi)carbonate species strengthen the oxidative capacity of TiO$_2$-containing dust particles with regard to SO$_2$ oxidation.

### 3.4 Field measurements of sulfate and (bi)carbonate ions

Complementary field sampling and analysis were further conducted to examine our hypothesis that intermediate CO$_3^{3-}$ may play role in secondary sulfate formation in the atmosphere. We first considered the meteorological condition of wind speed, which is an important parameter determining whether the local chemical process gains importance in affecting secondary sulfate formation. Meteorological information was collected from an open-access database (https://www.aqistudy.cn/, last access: 7 September 2021). During the sampling period, the wind scale mainly varies from 0 to 1, corresponding to the wind speed ranging from 0 to 1.5 m s$^{-1}$ (Fig. S13). All plots shown in Fig. S13 give rise to a statistical wind speed of 0.76 ± 0.73, which represents the weak dispersion of pollutants at a low wind speed (not exceeding 2.5 m s$^{-1}$) (Witkowska et al., 2016; P. Liu et al., 2020), indicating that a local source is a dominant contributor to the air pollution.

Under stagnant meteorological conditions (wind speed < 1.5 m s$^{-1}$), for the coarse mode (2.5–10 μm) of sulfate, the heterogeneous reaction of SO$_2$ on the dust surfaces is thought to be a major contributor (Liu et al., 2017). This correlates to the fact that a large mass fraction of mineral dust is abundant in the coarse-mode particulate matter (PM; Fang et al., 2017; Miller-Schulze et al., 2015) in which TiO$_2$ was found at mass mixing ratios ranging from 0.1 % to 10 % depending on the exact location where particles were lifted up (Hanisch and Crowley, 2003; Chen et al., 2012). Therefore, PM with relatively larger size dimensions are expected to contribute to secondary sulfate formation via heterogeneous reactions, which is supported by the recent field study in which the carbonate fraction of coarse PM is evidenced to promote secondary sulfate production (Song et al., 2018). Considering this, rather than determining the concentration of water-soluble ions in all stages, special attention is focused on PM collected in stages 1–4 (particles with their dimension of ≥ 3.3 μm). As (bi)carbonate ions are known as key precursors in producing CO$_3^{3-}$ and accelerating sulfate formation, quantifications of those relevant water-soluble ions were thus conducted (see details in Sect. 2 and Text S13).

We further consider the relationships between sulfate ions and (bi)carbonate ions by means of linear regression analysis. However, under the low wind speed (0.76 ± 0.73), correlation coefficients $R^2$ obtained for the relationship between bi(carbonate) and sulfate ions are not promising, with 0.56 (sulfate vs. carbonate) and 0.61 (sulfate vs. bicarbonate) for PM$_{3.3}$–PM$_{9.0}$ during daytime hours (Fig. 10). A plausible explanation is that in spite of having little significance, a local primary emission source also brings bias and uncertainty to the correlation results. Shanghai is a coastal city, and sulfate species such as K$_2$SO$_4$ and Na$_2$SO$_4$ from the sea salt contribute to the local sulfate emission as well (Long et al., 2014). On the other hand, this novel SO$_2$ oxidation chan-
nel is yet to be in the infant stage, and only active mineral dust components have been considered in this work, whereas other components found in the coarse mode of PM such as organic matter, elemental carbon, and sea salt (Cheung et al., 2011) are likely to involve this mechanism and alter the response of sulfate yield to SO$_2$ heterogeneous uptake. In addition, the water-soluble ions determined in these samples may not come from the net contribution of heterogeneous reaction processes in absolute daytime and nighttime periods. In other words, some of the collected samples, experiencing the heterogeneous reaction that occurs during the day/night-night/day shifts periods, will inevitably be assigned to the sulfate ions measured in separate sampling hours, thus reducing the correlation coefficients.

For those large particles (LPs), which refer to the particles with a diameter larger than 9 µm in this work, sulfate ions show a rather weak or even no correlation to (bi)carbonate ions measured in separate sampling hours, thus reducing the correlation coefficients. Likewise, higher correlation coefficients are also observed for PM$_{3.3}$–PM$_{9.0}$ than PM$_{>9.0}$ in the sample collected during the nighttime periods.

While we note that the correlations between sulfate and (bi)carbonate are not high in this work, ground-based field measurements of sulfate and (bi)carbonate ions shed light on their distinct correlations during the daytime and nighttime hours. In Figs. 10 and S14, the negative correlations between the mass concentrations of sulfate ions and (bi)carbonate ions are observed in the nighttime hours, consistent with the suppression of sulfate formation by CO$_2$ in the dark experiments. This is also supported by lab-based observations where CO$_2$-derived (bi)carbonate species are demonstrated to suppress sulfate production over two dominant mineral dust components, i.e., aluminum oxide (Y. Liu et al., 2020) and silicon dioxide (Fig. S15 and Text S23). Alternatively, while CO$_2$-derived (bi)carbonate may slightly affect sulfate accumulation over PM with high water content in the dark scenario, fresh PM is usually dry when emitted into the atmosphere. Due to the competitive adsorption, the occurrence of the suppression of SO$_2$ adsorption and subsequent sulfate formation is possible in the early emission stage before PM becomes wet, thus contributing to the overall negative correlation.

Instead, positive correlations were seen for those ions within PM sampled during the daytime hours, regardless of the size ranges and carbonate types (HCO$_3^-$ / CO$_3^{2-}$). This matches with the scenarios in which sulfate production, upon irradiation in the presence of (bi)carbonate ions, is increased over both model and authentic dust particles. Except for this case (nighttime period, size larger than 9 µm), most of the significance $P$ values for their correlations were smaller than 0.1, specifically with significant $P$ values below 0.5 determined for bicarbonate vs. sulfate, implying the plausible underlying connection between sulfate and (bi)carbonate ions. In fact, preceding ground-based observations of a highly correlated relationship between Ca$^{2+}$ and SO$_4^{2-}$ water-soluble
ions (Wu et al., 2020) during the carbonate-enriched dust storm episodes, together with persistent reports on the significant role of photochemical channels in elevating the sulfate concentration level during the daytime hours (Wei et al., 2019; Kim et al., 2018; Wu et al., 2017), indirectly reflects the possibility of accelerated SO$_2$ oxidation triggered by photo-generated active intermediates associated with carbonate species.

Overall, this is the first time that relationships between those ions have been explored separately in these two periods. Taken together, carbonate radicals are likely to promote sulfate production in the atmosphere during daytime hours. A detailed and systematic SO$_2$ oxidation channel triggered by CO$_3^{2-}$ needs further investigation to enable a better interpretation of the correlations between these inorganic ions at the given meteorological conditions of sampling and physicochemical properties of PM.

4 Conclusion

On the basis of the experimental and theoretical results derived from this work, we, for the first time, propose a novel reaction channel for fast SO$_2$ oxidation over mineral dust particles due to the formation of carbonate radical ions. A schematic chart for the sulfate formation in the presence of carbonate radicals upon solar light or bi(carbonate) ions under dark conditions is summarized and elucidated in Fig. 11. During the nighttime hours at 298 K (ambient temperature) CO$_2$-derived (bi)carbonate species are prone to having a slightly negative effect on sulfate formation, presumably due to the competitive adsorption between CO$_2$ and SO$_2$. For alkaline carbonate salt, it favors sulfate formation through the neutralization process. On the other hand, in the daytime, both CO$_2$-derived (bi)carbonate species and carbonate salt work as the precursors of CO$_3^{2-}$, which promotes sulfate formation. In particular, uptake coefficients for carbonate salt containing mineral dust can be increased 17 times, which is more pronounced than the increase due to the neutralization regime in the dark condition. Consistent with the findings reported in the early studies (Chen et al., 2021; Dupart et al., 2012), we speculate the production of gas-phase CO$_3^+$ when mineral dust particles are irradiated in the presence of CO$_2$ (atmospherically relevant concentration 400 ppm). This observation potentially implies that the increased sulfate yield in part comes from increased external secondary sulfate aerosol triggered by CO$_3^{2-}$ (g).

By means of ROS scavenger experiments, direct observation of carbonate radicals using NTAS analysis, oxygen isotope assay, ESR measurement, and DFT calculations, CO$_3^{2-}$-initiated S(IV) oxidation involving single electron transfer process is elucidated. While carbonate radical ions are mainly responsible for rapid sulfate formation, superoxide radical ions are likely to serve as a minor pathway over TiO$_2$-containing mineral dust particles (not shown in the schematic chart to direct readers’ focus on CO$_3^{2-}$). In addition, a weak correlation between sulfate ions and (bi)carbonate ions observed for PM$_{3.3-9.0}$ in this work reasonably correlates to non-chemical primary emission and the complicated nature of CO$_3^{2-}$ regime of sulfate production in the atmosphere. Nonetheless, complementary field sampling of ambient PM and analysis of sulfate and (bi)carbonate ions in this study unfold their distinct correlations during the daytime and nighttime hours; these two tendencies agree with the experimental observations.

In this work, only atmospheric secondary sulfate formation was considered, whereas the oxidation of primary organic species has not yet been investigated. In fact, carbonate radical ions are prone to reacting rapidly with electron-rich organic amines (Stenman et al., 2003; Yan et al., 2019) and phenol (Xiong et al., 2016; Busset et al., 2007), and they may potentially serve as the key oxidants that drive the fast formation of secondary organic aerosol in the atmosphere. Besides, an observation of the strengthened photochemistry launched by carbonate radicals suggests that such chemistry may be amplified on atmospherically relevant reactions that occur in cloud droplets and fog water, where they often contain hydroxyl radicals and water-soluble (bi)carbonate ions.

Importantly, gas-phase carbonate radical ions are speculated to be formed in the atmospherically relevant CO$_2$ concentration (400 ppm) when mineral dust is irradiated. This will help in the formation of external sulfate aerosol formation. Since both sulfate aerosol and CO$_2$ are known to regulate the radiation budget and solar energy balance on the Earth (Cheung et al., 2011; Möller, 1964), coupled with the CO$_2$-initiated promoted sulfate pathway found here, their

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overall influence on the global climate needs further investigation. Therefore, our study highlights the necessity for a comprehensive understanding of the CO$_3^{2-}$-relevant chemistry in the underlying impacts of fine PM concentration, human health, and climate. All these assumptions need to be investigated in further detail. This study provides the indication that carbonate radicals not only play a role as a marginal intermediate in tropospheric anion chemistry but also as a strong oxidant for the surface processing of trace gas in the atmosphere.

**Data availability.** The data that support the results are available from the corresponding author upon request.

**Supplement.** The supplement related to this article is available online at: https://doi.org/10.5194/acp-22-9175-2022-supplement.

**Author contributions.** YL, YD, and LZ initially proposed the idea. YL and YD designed the experiments together, and YL performed the most of the experiments. JL performed DFT calculations. YL, XF, and TW contributed to field samplings and data analysis. KL, KG, AUB, IN, QG, XZ, CG, and LZ provided suggestions on the experiments and writing of the draft. All authors wrote the paper.

**Competing interests.** The contact author has declared that none of the authors has any competing interests.

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