# **Enhanced Sulfate Formation in Mixed Biomass Burning and Sea-salt**

# **Interactions Mediated by Photosensitization: Effects of Chloride,**

# **Nitrogen-containing Compounds, and Atmospheric Aging**

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## **Abstract**

 Discrepancies persist between modeled simulations and measured sulfate concentrations in the marine boundary layer, especially when the marine air is influenced by biomass burning plumes. However, there is a notable dearth of research conducted on the interactions between sea-salt aerosol and biomass burning plumes, impeding a comprehensive understanding of the sulfate formation. This work studied sulfate formation by mixing real biomass burning (BB) extracts and NaCl, mimicking internal mixtures of BB and sea-salt particles. BB-NaCl particles had significantly higher sulfate formation rate than incense burning (IS)-NaCl particles. For fresh particles, the sulfate formation rate followed the trend of corn straw (CS)-NaCl>rice straw (RS)-NaCl>wheat straw (WS)-NaCl>IS-NaCl. The filter sample aging was achieved by exposure to OH• generated from UV irradiation. After aging, RS-NaCl particles exhibited the highest enhancement in sulfate formation rates among all the BB-NaCl particles, due to interactions between RS and NaCl. Bulk aqueous experiments spiked with NaCl using mixtures of model photosensitizers (PS) and nitrogen-containing organic compounds (NOCs), pyrazine (CHN) and 4-nitrocatechol (CHON), revealed positive effects of chloride in the PS-CHON system and negative effects in the PS-CHN system in sulfate formation. Our work suggests that BB reaching or near coastal areas can affect sulfate formation via photosensitizer-mediated reactions, potentially exacerbating air pollution.

**Keywords:** sulfate formation, biomass burning, photosensitization, sea-salt aerosol, chloride

## **1 Introduction**

 Recent fire outbreaks in areas like Canada, Amazonia, and Southeast Australia, together with the increased fire frequency and intensity reports in areas like western US have highlighted the risks of fire, especially biomass burning (BB), to human health and climate change (Bond et al., 2013; Andreae, 2019; Jones et al., 2022). As an agricultural powerhouse, China boasts immense agricultural crop yields, especially in rice, wheat, and corn throughout the country. These crop residues are frequently burned in rural areas for cooking and heating purposes, as well as for land preparation after harvest, resulting in the substantial production of light- absorbing species, such as brown carbon (BrC) (Chen et al., 2017). Recent studies have reported that specific BrC species from biomass burning, including vanillin (VL), acetovanillone, syringaldehyde (SyrAld), and naphthalene-derived secondary organic aerosol (Teich et al., 2016; Li et al., 2024; Liu et al., 2020; Wang et al., 2021b) can act as photosensitizers (PS) and 48 oxidize  $SO<sub>2</sub>$  to sulfate (Zhou et al., 2023; Liang et al., 2024). Atmospheric processes like aging or long-range transport, can alter the chemical compositions and optical properties of PS, and hence affect the sulfate formation potential (You et al., 2020; Li et al., 2019). Sea-salt aerosol (SSA), with its high particulate matter loadings and extensive surface area, plays a significant role in interfacial and multiphase reactions with reactive gases, thereby impacting global radiation balance and air quality in marine and coastal areas (Gantt and Meskhidze, 2013; Chi et al., 2015). Prior research has identified several secondary sulfate formation pathways in SSA, 55 e.g., multiphase SO<sub>2</sub> oxidation by O<sub>3</sub> (Alexander et al., 2012), coexistence of NO<sub>2</sub> (Zhang and Chan, 2023b), PS (Tang et al., 2023), chlorine-PS synergistic effects (Zhang and Chan, 2024), and Cl and OH radicals generated by chlorine photoactivation (Cao et al., 2024). The chlorine/chloride studies are particularly interesting as they highlight the importance of NaCl-based photochemistry in sulfate formation.

 SSA can frequently mix with organic matter through processes such as sea-to-air emission, photochemical oxidation and atmospheric transport (Liu et al., 2023b). Previous studies have observed elevated sulfate concentrations in coastal regions when air masses passed through inland areas due to intensive BB or other anthropogenic emissions, suggesting the possible interactions between the SSA (primarily sodium chloride) and anthropogenic emissions (Qiu et al., 2019; Huang et al., 2018; Wu et al., 2022). Van Pinxteren et al. (2015) observed an increase 66 in sulfate concentration (2.26  $\mu$ g m<sup>-3</sup>) during the RV MARIA S cruise as it approached the 67 African mainland, in contrast to the marine-origin aerosol (1.59  $\mu$ g m<sup>-3</sup>), showing significant influence of BB. Hence, mixing of sea-salt and biomass burning aerosols can lead to secondary aerosol production in coastal regions.

 Transmission electron microscopy (TEM) studies indicate that most coastal particles are internally mixed, showing a higher proportion of organic and salt mixtures in the presence of biomass burning aerosols, accompanied by an increase in sulfate (Dang et al., 2022; Li et al., 2003). However, discrepancies persist between modeled simulations and measured sulfate concentrations in MBL (Yu et al., 2023). The interactions of sea-salt and BB aerosols, especially in multiphase reactions, can potentially unravel the intricate chemistry of sulfate formation in BB affected MBL. Hence, internal mixtures of inorganic salt and water-soluble organic carbons

are often used in reaction studies (Tan et al., 2024).

 The two main atmospheric water systems in marine boundary layer (MBL) were wet aerosol (droplets in our case) and cloud/fog (bulk aqueous solutions), both droplet and aqueous reactions are relevant for studying multiphase sulfate formation within MBL (Ruiz-Lopez et al., 2020; Herrmann, 2003). Typically, droplet reactions were characterized by high ionic strength 82 (up to >10 M), low liquid water content  $(10^{-7} - 10^{-3} \text{ cm}^3 \text{ m}^{-3})$  and high surface-to-volume ratio whereas aqueous reactions exhibit the opposite characteristics. Additionally, droplet experiments can encompass certain interfacial reaction pathways that may occur in the atmosphere, especially in submicron particles (Ruiz-Lopez et al., 2020) .

 Previous studies have detected a significant proportion of NOCs, including nitroaromatics (CHON) and reduced nitrogen species (CHN) in biomass burning plumes, wildfires and ambient samples (Zhong et al., 2024; Wang et al., 2017b; Song et al., 2022; Cai et al., 2020). 89 These NOCs are considered ubiquitous contributors to BrC and can affect global climate and human health. Moreover, recent research has discovered aerosol pollution in marine background regions, with high levels of NOCs when air masses are transported from wildfires or biomass burning events in nearby (Zhong et al., 2024; Qin et al., 2024). These NOCs, combined with reactive gases, may mix with SSA and impact regional air quality in coastal zones. Therefore, it is essential to further investigate the interactions between NOCs, reactive gases, and SSA.

 In this study, we performed in-situ droplet and bulk aqueous solution reaction experiments using BB extracts-NaCl mixtures to explore the possible interplay between biomass burning and marine aerosols in coastal areas. BB was derived from the burning of rice straw (RS), wheat straw (WS), and corn straw (CS) as well as incense burning (IS). The aims of this study are to: (i) compare the differences in sulfate formation among different kinds of BB-NaCl particles and BB extracts; (ii) examine the impacts of OH• aging on sulfate formation across different BB-NaCl particles and BB extracts; (iii) investigate the role of NOCs and chloride ions in BB extracts mediated sulfate formation.

**2 Material and methods**

### **2.1 Burning experiments**

 Three types of commonly used biomass (RS, WS and CS) were cut into small, uniform pieces 107 ( $\sim$ 10 cm in length) and dried. About 100 g of the dried biomass materials ( $\sim$ 10% moisture content) was then introduced into a traditional iron stove commonly used in rural areas (Figure S1). The stove was covered with a hood and the biomass was ignited using a propane lighter. 110 The generated BB smoke was collected onto 90-mm quartz filters at  $0.9 \text{ m}^3 \text{ min}^{-1}$  for 10 minutes by a custom-made aerosol sampler under mixed combustion conditions (include flaming and 112 smoldering, modified combustion efficiency MCE,  $0.85 \le \Delta [CO_2]/(\Delta [CO_2] + \Delta [CO]) \le 0.95)$ 113 (Ting et al., 2018). The sampler was placed 1 meter above the ground and connected to a PM<sub>2.5</sub> sampling head through a sampling pump. For incense burning (IS), laboratory-generated 115 smoldering smoke was collected on 47-mm quartz filters at a flow rate of  $\sim 6.0$  L min<sup>-1</sup> for 80 min using a stainless-steel combustion chamber. Note that the different combustion modes of  IS and BB are intentionally used to represent real-world combustion conditions. Our previous study demonstrated that IS was representative of BB based on GC×GC chromatograms and pixel-based partial least squares discriminant analysis (Tang et al., 2023). Hereafter, we will use BB to represent both the real BB materials and the surrogate materials (IS) unless otherwise specified. After sampling, the collected BB samples (fresh BB) were wrapped by pre-baked

122 aluminum foil (550 °C for 6 h) and stored at -20 °C until further analysis.

 To achieve atmospheric OH• aging, the collected fresh BB filter samples were placed in a pre- flushedchamber (zero air, more than 24 h) and illuminated with UV lamps for 40 min. We used lamps of 185 nm and 254 nm, the combination of which has been widely used in oxidation flow 126 reactor design and experiments for mimicking atmospheric OH• aging conditions (Peng and Jimenez, 2020; Rowe et al., 2020; Tkacik et al., 2014; Hu et al., 2022). The estimated OH 128 exposure was  $\sim 2.0 \times 10^{12}$  molecules cm<sup>-3</sup> s, equivalent to an atmospheric aging period of 15 129 days (assuming an average atmospheric OH concentration of  $1.5 \times 10^6$  molecules cm<sup>-3</sup>) (Mao et al., 2009). Detailed characterization of the OH exposure can be found in our previous study (Tang et al., 2023). We will discuss the sulfate formation of these fresh and OH• aged samples later.

### **2.2 Materials and instrumentation**

 Aqueous stock solutions of BB samples were prepared by dissolving the collected filters in ultrapure water and subjecting them to ultrasonication in a cooled-water bath three times, each for 20 minutes. The resulting water extracts of the BB were then filtered through 0.22 μm PTFE filters and stored in brown vials at 4℃ in a refrigerator. The anions, i.e., chloride, sulfate and nitrate of the BB extracts were analyzed by Dionex ion chromatography (ICS 1100, CA). An 139 aliquot (~0.5 ml) of the BB or IS extracts was used for water-soluble organics detection by ultra-high performance liquid chromatography (Thermo Scientific Dionex UltiMate 3000 UHPLC) coupled with high-resolution Orbitrap Fusion Lumos Tribrid mass spectrometry (Orbitrap HRMS, Thermo Fisher Scientific, USA). The particulate organic matter was characterized by directly heating the filter samples using a thermal desorption module (TDS3, Gerstel) coupled to comprehensive two-dimensional gas chromatography-mass spectrometer (GCMS-TQ™8050 NX, Shimadzu, Japan). UV-Vis spectrometry (UV-3600, Shimadzu, Japan) was employed to examine the absorbance of BB extracts. Total organic carbon (TOC) was measured by total carbon analyzer (TOC-L CPH, Shimadzu, Japan). Metal concentrations were measured by inductively coupled plasma-mass spectrometry (ICP-MS, Agilent 7800). Detailed analysis can be found in Text S1. Aqueous stock solution of sodium chloride (≥99.8%, Unichem) was prepared by dissolving the corresponding salt in ultrapure water to obtain a concentration of 1M. The study utilized high purity grade synthetic air and nitrogen supplied by the Linde HKO Ltd., while sulfur dioxide was obtained from the Scientific Gas Engineering Co., Ltd.

#### **2.3 Multiphase and aqueous-phase reactions of S(IV)**

 In SO<sub>2</sub> uptake experiments, the stock solution of BB (fresh and aged) extracts was premixed 155 with sodium chloride solution (1M) at a volume ratio of 1:1 and the solutions had pH a of 4-6. A droplet generator (Model 201, Uni-Photon Inc.) was then utilized to deposit droplets onto a 157 hydrophobic substrate (model 5793, YSI Inc.) for SO<sub>2</sub> uptake experiments. Reactive SO<sub>2</sub> uptake  experiments were performed via a flow cell/in-situ Raman system at controlled room temperature (23-25℃). The top and bottom quartz windows of the flow cell were used for Raman analysis and UV irradiation, respectively. The light experiment was performed using a 161 xenon lamp (model 6258, ozone free, 300W, Newport), with photon flux of  $9.8 \times 10^{15}$  photons 162 cm<sup>-2</sup> s<sup>-1</sup> in 280-420 nm received by particles in the flow cell (Zhang and Chan, 2023b). Identical experiments were conducted in the dark, with the lights off and the experimental area kept in complete darkness. The relative humidity (RH) inside the flow cell was adjusted to 80% by mixing dry and wet synthetic air or nitrogen. The particles were then equilibrated at 80% RH 166 for over 60 min and remained liquid throughout the experiment period.  $SO<sub>2</sub>$  was introduced into the system to reach a concentration of 8.0 ppm. The prescribed size used in our in-situ 168 Raman research was  $60 \pm 5$  µm. Despite using particles for droplet experiments that were larger 169 than ambient fine particles, we employed the SO<sub>2</sub> uptake coefficient ( $\gamma_{SO_2}$ ) as a kinetic 170 parameter to account for the particle size effects. Comprehensive calculation of  $\gamma_{SO2}$  can be found in our previous studies (Gen et al., 2019a, b; Tang et al., 2023; Zhang et al., 2020a).

 Aqueous-phase photochemical reactions were performed using a quartz photo reactor (Mabato et al., 2023; 2022). Specifically, a 500 mL solution containing 100 ppm bisulfite and 1 ppm BB TOC extracts were continuously mixed using a magnetic stirrer throughout the experiments. Note that the 1 ppm BB TOC and 100 ppm bisulfite align well with the atmospheric-relevant ranges in aqueous aerosols, fogs and clouds, where PS concentration can reach hundreds of micromolar and total sulfur concentration can exceed several millimolar (Anastasio et al., 1997; Guo et al., 2012; Shen et al., 2012; Rao and Collett, 1995). To achieve air-saturated conditions, 179 synthetic air was continuously introduced to the solutions at a flow rate 0.5 L min<sup>-1</sup> throughout the experiments. The above mixed solutions were then exposed to radiation via the same xenon 181 lamp (light intensity of 1318 mW/cm<sup>2</sup>) as in the droplet experiments. Samples were collected at 1h interval for a total of 8 h for sulfate and bisulfite analysis using ion chromatography. Unlike droplet experiments, NaCl was not added to the bulk solution in most studies, unless specified otherwise.

## **3 Results and Discussion**

### **3.1 Enhanced sulfate production in BB-NaCl droplets over IS-NaCl droplets.**

 As no sulfate was detected under dark conditions for any of the experiments, we have focused on the light experiments. Figure 1 depicts the sulfate production by (a) fresh BB-NaCl; (b) aged 189 BB-NaCl droplets as a function of time in the presence of light, air, and  $SO_2$  at 80% RH. Our previous study (Tang et al., 2023) has found significantly higher sulfate formation of IS-NaCl droplets over NaCl droplets. Here, we only focus on the comparison of sulfate formation between different kinds of BB-NaCl droplets and IS-NaCl droplets. Regardless of whether the extracts were fresh or aged, the sulfate production by BB-NaCl droplets was higher than IS- NaCl droplets. Specifically, sulfate formed by fresh (F) BB-NaCl droplets followed the trends 195 of CS<sub>F</sub>-NaCl (16.8  $\pm$  2.6 mM ppmC<sup>-1</sup>) >RS<sub>F</sub>-NaCl (9.8  $\pm$  0.1mM ppmC<sup>-1</sup>) >WS<sub>F</sub>-NaCl (4.2  $\pm$ 196 0.2mM ppmC<sup>-1</sup>) >IS<sub>F</sub>-NaCl (0.8 mM ppmC<sup>-1</sup>) after illumination for 1080 min. In aged (A) 197 samples, while  $BB_A-NaCl$  is more efficient than  $IS_A-NaCl$  in sulfate formation, the order of 198 sulfate formation was different from the fresh samples:  $RS_A$ -NaCl (35.2  $\pm$  0.6 mM ppmC<sup>-1</sup>) >  $CS_A-NaCl (13.0 \pm 0.1 \text{ mM ppmC}^{-1}) > WS_A-NaCl (6.0 \pm 1.6 \text{ mM ppmC}^{-1}) > IS_A-NaCl (0.6 \text{ mM}$ 

200 ppmC<sup>-1</sup>). The sulfate enhancement factors of  $RS_F$ -NaCl, WS<sub>F</sub>-NaCl, and CS<sub>F</sub>-NaCl over IS<sub>F</sub>-201 NaCl after 18 h SO<sub>2</sub> uptake (Sulfate<sub>BB<sub>F</sub>-NaCl/IS<sub>F</sub>-NaCl) were 11.7, 5.0 and 20.0, respectively.</sub> The enhancement of sulfate can also be observed in aged BB samples, with values of 54.3, 9.2 203 and 20.1 for  $RS_A-NaCl$ ,  $WS_A-NaCl$ , and  $CS_A-NaCl$ , respectively. The lower sulfate formation of IS-NaCl droplets than BB-NaCl droplets can be explained by the significantly higher TOC concentration of IS due to the incomplete and smoldering combustion (Table S1). The TOC 206 concentration of the IS extracts ( $>550$  mg L<sup>-1</sup>) was nearly an order of magnitude higher than 207 that of the BB extracts (34.0-69.9 mg L<sup>-1</sup>), and ∑anions/WSOC exhibited a more than tenfold increase in BB extracts than in IS extracts. Previous studies have confirmed that the smoldering condition of BB results in significantly more organic compounds and less ions than flaming condition (Wang et al., 2020b; Fushimi et al., 2017; Kalogridis et al., 2018; Kim et al., 2018). Additionally, significantly higher polycyclic aromatic hydrocarbons (PAHs) proportion 212 (12.2%-16.6% by intensity) in BB than IS  $(\sim 5.0\%)$  were observed by GC×GC-MS. Huang et 213 al. (2022a) reported higher PAHs in BB particles (CS, WS, RS,  $>262.5$  mg kg<sup>-1</sup>,  $>3.7\%$  of 214 organic matter) than in IS particles  $(3.3 \text{ mg kg}^{-1}, 0.9\%$  of organic matter) (Song et al., 2023). Fushimi et al. (2017) and Kim et al. (2021) demonstrated that more PAHs would be emitted under flaming than smoldering conditions. PAHs like pyrene, fluoranthene, and phenanthrene have been recognized as PS (Jiang et al., 2021; Yang et al., 2021) and are mainly from combustion processes, e.g., pyrosynthesis from aliphatic and aromatic precursors in biomass burning processes and the constituents vary with temperatures and oxygen contents (Pozzoli et al., 2004). The higher percentage of PAHs in BB together with the collection procedure (mixed combustion and higher temperature for real BB and smoldering and lower temperature for IS) suggested that the BB materials would generate more PAHs at high temperatures and may contribute to sulfate formation.

224 Table 1 and Figure S2 present the reactive  $(\gamma_{SO_2})$  and normalized reactive SO<sub>2</sub> uptake 225 coefficients  $(n\gamma_{SO_2})$  of different BB-NaCl droplets. The  $\gamma_{SO_2}$  obtained in our study are 0.9 -226 6.6×10<sup>-6</sup>, which are consistent but fall on the low side of the reported heterogeneous SO<sub>2</sub> 227 oxidation processes, including nitrate photolysis (10<sup>-6</sup>-10<sup>-5</sup>) (Gen et al., 2019a), TMI-catalyzed 228 oxidation  $(10^{-6} - 10^{-4})$  (Zhang et al., 2024), NO<sub>2</sub>/O<sub>3</sub> oxidation  $(10^{-6} - 10^{-4})$  (Zhang et al., 2021a; 229 Zhang and Chan, 2023a) and peroxide oxidation  $(10^{-6} - 10^{-1})$  (Wang et al., 2021a; Ye et al., 2018; 230 Yao et al., 2019). Additionally, the reported  $\gamma_{SO_2}$  in our study aligns well with the results 231 obtained from ambient samples in Beijing (Zhang et al., 2020b). The large discrepancy of the 232 reported  $\gamma_{SO_2}$  can be attributed to the differences in aerosol components, particle size, RH, SO<sub>2</sub> 233 and oxidants concentrations. Higher  $n\gamma_{SO_2}$  were found for fresh and aged real BB-NaCl than IS-NaCl droplets, following the trend of :CS<sub>F</sub>-NaCl  $(8.8\times10^{-8} \text{ ppmC}^{-1})$ >RS<sub>F</sub>-NaCl  $(6.2\times10^{-8} \text{ mm}^{-1})$ 234 ppmC<sup>-1</sup>)>WS<sub>F</sub>-NaCl (2.0×10<sup>-8</sup> ppmC<sup>-1</sup>)>IS<sub>F</sub>-NaCl (0.61×10<sup>-8</sup> ppmC<sup>-1</sup>) and RS<sub>A</sub>-NaCl (2.2×10<sup>-1</sup>) 235 <sup>7</sup> ppmC<sup>-1</sup>)>CS<sub>A</sub>-NaCl (6.2×10<sup>-8</sup> ppmC<sup>-1</sup>)>WS<sub>A</sub>-NaCl (3.5×10<sup>-8</sup> ppmC<sup>-1</sup>)>IS<sub>A</sub>-NaCl (0.46×10<sup>-8</sup> 236 237 ppm $C^{-1}$ ), respectively.

 In our previous study, we observed a significant increase in sulfate formation in IS-NaCl droplets than NaCl droplets, which we attributed to PS present in the IS (Tang et al., 2023). Considering the fact that BB-NaCl droplets produced sulfate more efficiently than IS-NaCl droplets and NaCl droplets, we explore the underlying mechanisms driving this phenomenon. 242 Possible reasons include nitrate (from BB extracts or newly formed) photolysis, [Cl<sup>-</sup>H<sub>3</sub>O<sup>+</sup>-O<sub>2</sub>]

243 photoexcitation (Cl from BB extracts),  $H_2O_2$  oxidation, BC-catalyzed oxidation, reactive 244 nitrogen species oxidation, and organics-driven pathways e.g., HCHO, photosensitizing 245 components, organic peroxide, and TMI-organic oxidation (Ye et al., 2023).

 Since there was no nitrate peak in our Raman spectra in all experiments, the potential impact 247 from nitrate photolysis was excluded. Besides, the significantly low Cl concentration (0.0002- 0.001M) in the original BB extracts (compared to 1M NaCl, Table S1) has minimized the 249 influence of chloride photoexcitation of  $\left[Cl - H_3O^+ - O_2\right]$  (Cl from BB extracts) on the sulfate 250 formation. Reactive nitrogen species e.g.,  $NO<sub>x</sub>$ , HONO and  $NH<sub>3</sub>$  were neither introduced nor detected in our system, indicating that the oxidation pathway involving reactive nitrogen species was insignificant. Additionally, the water extraction process has excluded the possibility of BC-catalyzed oxidation. The absence of sulfate formation in dark conditions ruled out the 254 involvement of direct  $H_2O_2$  oxidation and organic peroxide oxidation pathways. The concentrations of TMI did not exhibit a consistent relationship with the sulfate formation 256 observed in both  $BB_F-NaCl$  and  $BB_A-NaCl$  droplets (Figure S3), suggesting that the TMI- catalyzed oxidation pathway may not be responsible for the observed phenomenon. Therefore, the most probable reason for the enhancement of sulfate formation by BB-NaCl droplets over NaCl droplets would be the photosensitizing components. Given the complexity and the lack of a method to quantify PS in BB aerosols, using the total TOC concentration as an upper limit 261 for estimating PS concentration is considered a compromise that allows systematic comparison. The sulfate formation reported here can be considered as the lower limit of photosensitizing capacity. Our goal is to compare the photosensitizing ability in different chemical systems, but not to quantify their absolute values.

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266 State-of-the-art mass spectrometry analysis including UHPLC-Orbitrap-MS and GC×GC-MS 267 showed the existence of possible PS such as PAHs (e.g., fluoranthene, pyrene, 268 cyclopenta[cd]pyrene, 4-methylphenanthrene, benzo[a]pyrene, perylene, Table S2) and 269 aromatic carbonyls (SyrAld, VL, 3,4-dimethoxybenzaldehyde, acetophenone, acetosyringone, 270 Table S2). Photosensitizing components can directly or indirectly (by forming secondary 271 oxidants in the presence of oxygen) oxidize S(IV) to S(VI). Wang et al. (2020a) proposed a 272 direct oxidation process of  $S(IV)$  to sulfate by excited triplet states of photosensitizers ( ${}^{3}PS*$ ). 273 To explore the contribution of the direct  ${}^{3}P S^{*}$  oxidation on sulfate formation, we performed the 274 same sets of experiments in N<sub>2</sub>-saturated condition, shown in Figure S4. Under N<sub>2</sub>-saturated 275 conditions, secondary oxidants such as  $HO<sub>2</sub>•$ ,  $OH<sub>•</sub>$  oxidation pathway can be ruled out due to 276 the lack of oxygen. Chlorine radicals from droplets can also react with dissolved  $SO<sub>2</sub>$  to 277 generate sulfite radicals, but  $O_2$  is required to form sulfate. Despite initial molecular oxygen in 278 the droplets may also participate in sulfate formation under  $N_2$ -saturated conditions, its  $279$  contributions are likely minimal. Consequently, the sulfate formed under N<sub>2</sub>-saturated condition 280 can be considered as the upper limit of direct <sup>3</sup>PS\* oxidation. The BB-NaCl droplets showed 281 only direct PS\* oxidation contribution of 3.6% to 22.7%, highlighting the predominant role of 282 secondary oxidants (Tang et al., 2023). For BB<sub>F</sub>-NaCl droplets, the contribution of direct <sup>3</sup>PS\* 283 followed the trend of  $W\text{S}_\text{F}\text{-}\text{NaCl}$  (22.7%) >  $\text{RS}_\text{F}\text{-}\text{NaCl}$  (15.7%) >  $\text{CS}_\text{F}\text{-}\text{NaCl}$  (7.0%), while for 284 BB<sub>A</sub>-NaCl droplets,  $WS_A-NaCl(10.2\%) > CS_A-NaCl(6.7\%) > RS_A-NaCl(3.6\%)$  was observed. 285 In summary, regardless of whether fresh or aged, the secondary oxidants triggered by indirect 286 <sup>3</sup>PS\* oxidation were the main reason for sulfate formation, highlighting the importance of O<sub>2</sub>





289 Figure 1. Sulfate production under different droplet compositions as a function of time: (a) fresh 290 BB-NaCl droplets; (b) aged BB-NaCl droplets in air at 80% RH. RS, WS, CS and IS represent 291 rice straw, wheat straw, corn straw and incense burning, respectively. The subscripts F and A 292 represent fresh and aged, respectively.

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294 Table 1. Sulfate formation rate  $(k_{so_4^2})$ , reactive  $(\gamma_{SO_2})$  and normalized SO<sub>2</sub> uptake coefficient 295 ( $n\gamma_{SO_2}$ ) at various particle compositions at 80% RH as well as sulfate formation rate ( $k_{SO_4^{2-}}$ ) 296 using different BB extracts and model compounds in bulk aqueous reactions. 1, 10, 100 and 297 200 represent the concentration of different compounds (in ppm).

<b>Droplet Experiments</b>			
Particle Composition	$k_{so_4^{2-}}$	$\gamma_{SO_2}$	$n\gamma_{SO_2}$ a
	$(\mu M \text{ min}^{-1} \text{ ppmC}^{-1})$		$ppmC^{-1}$
$RS_F-NaCl$	$9.4 \pm 0.10$	$(2.2 \pm 0.023) \times 10^{-6}$	$(6.2 \pm 0.066) \times 10^{-8}$
WS <sub>F</sub> -NaCl	$3.7 \pm 0.048$	$(0.66 \pm 0.0086) \times 10^{-6}$	$(2.0 \pm 0.027) \times 10^{-8}$
$CS_F-NaCl$	$15.6 \pm 0.11$	$(2.0 \pm 0.015) \times 10^{-6}$	$(8.8 \pm 0.065) \times 10^{-8}$
$IS_F-NaCl$	$0.83 \pm 0.011$	$(1.7 \pm 0.034) \times 10^{-6}$	$(0.61 \pm 0.012) \times 10^{-8}$
$RS_A-NaCl$	$33.5 \pm 0.38$	$(6.6 \pm 0.074) \times 10^{-6}$	$(21.5 \pm 0.24) \times 10^{-8}$
$WS_A-NaCl$	$6.2 \pm 0.18$	$(0.92 \pm 0.027) \times 10^{-6}$	$(3.5 \pm 0.10) \times 10^{-8}$
$CS_A-NaCl$	$10.6 \pm 0.23$	$(1.0 \pm 0.023) \times 10^{-6}$	$(6.2 \pm 0.13) \times 10^{-8}$
$IS_A-NaCl$	$0.72 \pm 0.026$	$(1.3 \pm 0.052) \times 10^{-6}$	$(0.46 \pm 0.017) \times 10^{-8}$
		<b>Bulk Aqueous Experiments</b>	
Aqueous Reactions	Concentration	$k_{so_4^{2-}}$	$k_{so_4^{2-}}$
	(ppm)		





<sup>a</sup>The  $n\gamma_{SO_2}$  was calculated by normalizing the  $\gamma_{SO_2}$  with the TOC concentration in the BB extracts, i.e.,  $n\gamma_{SO_2} = \gamma_{SO_2}/TOC$ 

#### 298

## 299 **3.2 Aging effects on sulfate formation across various BB materials**

 To investigate the aging effects across various BB materials on sulfate formation, we subjected the collected BB filters to OH radical aging by irradiating them with UV lights at wavelengths of 185 nm and 254 nm. This combination effectively generate OH radicals (Tang et al., 2023). Figure S5 exhibits the differences in sulfate formation rates in droplets of different fresh and aged BB materials. RS and WS show sulfate formation enhancement, while CS and IS show reduction after aging. Figure 2(a) shows that the 18h sulfate enhancement factor 306 (Sulfate<sub>A</sub>/Sulfate<sub>F</sub>) followed the trend of RS-NaCl (3.6) >WS-NaCl (1.4) > CS-NaCl (0.8)  $\approx$  IS- NaCl (0.8), which is neither consistent with the trends of sulfate formation for BB<sub>F</sub>-NaCl nor BBA-NaCl, indicating the varying effects of aging of BB materials. A similar trend was found 309 for  $n\gamma_{SO_2}$ , showing the highest and lowest sulfate enhancement for RS-NaCl (3.5) and IS-NaCl (0.7), respectively.





312 Figure 2. Enhancement factor of (a) sulfate and (b) normalized  $SO_2$  uptake coefficient  $n\gamma_{SO_2}$ 313 between fresh and aged BB-NaCl droplets.

 Bulk aqueous reactions using fresh/aged BB extracts without NaCl were performed to investigate the aging effects on the sulfate formation in the cloud phase (Figure S6). As the experiment proceeded, sulfate concentrations accumulated while bisulfite concentrations decreased. Concurrently, the pH of the aqueous solution decreased from approximately 5.0 to 3.0, reflecting enhanced acidity. Bulk reactions showed lower sulfate formation rates than droplets reactions, which may be attributed to the accelerated reactions induced by PS at the air-water interface (Wang et al., 2024c; Martins-Costa et al., 2022), as well as the differences in concentrations of S(IV) and NaCl between bulk and droplet surfaces. However, given that interfacial reactions are closely linked to particle size (Wei et al., 2020; Chen et al., 2022b), and additional research is needed to better understand its influence. Our experiments involve large droplets of the size of 60 μm. The interfacial effects of such large droplets may not be evident. Future work to examine the interfacial effects of submicron and nanometer size particles is needed.

# **3.3 Chemical characterization of BB extracts and sulfate formation in bulk aqueous solutions**

329 In bulk experiments, all BB extracts have higher  $k_{so4}^2$  after aging. The increased sulfate formation of BB extracts after aging may be due to changes in their chemical compositions. 331 Compared to  $RS_F (28.3\%$  for CHON- and 67.3% for CHN+ in total intensity),  $RS_A$  has higher CHON- (36.1%) and CHN+ (88.3%) percentages (Figs. S7-S8). Zhao et al. (2022) observed a slight increase in CHON percentage for RS from 53.4% to 56.2% after aging. Similar trend was observed for CS extracts, where CHON- and CHN+ percentage increases from 26.7% and 65.2% to 31.5% and 68.8%, respectively, after aging. To semi-qualitatively distinguish BrC chromophores from the rest of dissolved organic carbon, we used the DBE value to the range 337 of 0.5C≤DBE≤0.9C as a criterion (Lin et al., 2018). For example,  $RS<sub>A-BrC</sub>$  denotes the BrC chromophores in RSA, according to the above. Larger intensity fractions of CHON- species 339 were found in  $RS_{A-BrC}$  (41.9%) and  $CS_{A-BrC}$  (35.5%) than  $RS_{F-BrC}$  (32.3%) and  $CS_{F-BrC}$  (34.7%). One of the key categories of CHON- is nitrated aromatics, which have been widely identified in lab-generated BB smoke (Huang et al., 2022b; Wang et al., 2017a; Zhang et al., 2022; Xie et al., 2019) and field campaigns (Salvador et al., 2020; Mohr et al., 2013; Chen et al., 2022a). A 343 series of CHON- species, e.g.,  $C_6H_5NO_3$ ,  $C_6H_5NO_4$ ,  $C_7H_7NO_3$ , and  $C_8H_9NO_3$ , which were tentatively identified as nitrophenol, nitrocatechol, methyl-nitrophenol, and dimethyl- nitrophenol, have been detected in our BB extracts. Nitrophenols photolysis has been found to be a potential source of OH radicals (Sangwan and Zhu, 2018; Guo and Li, 2023; Cheng et al., 2009; Sangwan and Zhu, 2016) and it may partially contribute to the increase in sulfate 348 formation by  $RS_A$  and  $CS_A$ .

 Approximately 80% of the CHN+ species identified exhibited a diatomic nitrogen composition in their molecular formula. The precise determination of the molecular structures of these compounds solely based on elemental composition is challenging due to the presence of stable isomers. However, the N-bases, which contain two nitrogen atoms, can be attributed to various 353 N-heterocyclic alkaloids (Figure S9). For example, homologs of  $C_5H_6N_2(CH_2)_n$  were likely pyrazine, pyrimidine or amino pyridine, which were composed of six-membered heterocyclic 355 rings with N atoms and alkyl side chains (Lin et al., 2012; Laskin et al., 2009).  $C_5H_8N_2(CH_2)_n$   were likely alkyl-substituted imidazole compounds, featuring a five-membered heterocyclic ring with two nitrogen atoms as the core structure and alkyl side chains (Lin et al., 2012; Laskin 358 et al., 2009). For  $C_7H_6N_2(CH_2)_n$  homologs, the core skeleton was  $C_7H_6N_2$ , with an AI<sub>mod</sub> of 0.8, indicating its distinctive characteristics of compounds containing fused five-membered and six- membered rings, such as benzimidazole or indazole (Wang et al., 2017b). Redox-inactive heterocyclic nitrogen-containing bases, e.g., pyridine, imidazole, and their derivatives, have been shown to enhance the redox activity of humic-like substances (HULIS) fraction by hydrogen-atom transfer, with the degree of enhancement directly correlated to their concentration (Dou et al., 2015; Kipp et al., 2004). Thus, the increased CHN+ percentage may 365 also contribute to the enhanced sulfate formation of  $RS_A$  and  $CS_A$  by acting as a H-bond 366 acceptor to facilitate the  ${}^{3}P S^*$ -mediated oxidation by generating more oxidants.

367 However, the CHON- and CHN+ percentages in  $WS_A$  were lower than  $WS_F$ , indicating that the sulfate enhancement in WS<sup>A</sup> was not due to the CHON and CHN species. Instead, CHO-369 accounted for higher proportion in  $WS_A$  (68.5%) and  $WS_{A-BrC}$  (68.9%) than  $WS_F$  (65.0%) and WSF-BrC (64.8%).This aligns with a prior AMS study, showing increased CHO proportions in aged wheat burning emissions (Fang et al., 2017). We suppose that CHO- compounds, particularly photosensitizing compounds with carbonyl groups, would explain the difference of sulfate formation in WS extracts (Gómez Alvarez et al., 2012; Mabato et al., 2023; Felber et al., 2020; Fu et al., 2015). Therefore, we filtered the chemical formula of CHO- species from UHPLC-Orbitrap-HRMS by applying the maximum carbonyl ratio (MCR) (Zhang et al., 2021b; Wang et al., 2024a; Calderon-Arrieta et al., 2024; Liu et al., 2023a), H/C, O/C as well as 377 modified aromaticity index  $(AI_{mod})$  to focus on potential PS (Zherebker et al., 2022; Koch and Dittmar, 2006). In short, molecular formula were classified into six groups, namely, condensed 379 aromatics ( $AI_{mod} \ge 0.67$ ), polyphenolics (0.50< $AI_{mod}$ <0.67), highly unsaturated and phenolic compounds (AImod≤0.5, H/C<1.5), aliphatics (H/C≥1.5, O/C≤0.9, N=0), peptide-like compounds (H/C≥1.5, O/C≤0.9, N>0) and sugar-like compounds (H/C≥1.5, O/C>0.9), details can be found in Text S1. As aliphatics, peptide-like compounds and sugar-like compounds are unlikely to be PS, we exclude them as potential PS. By applying further data filtration involving CHO-, condensed aromatics, polyphenolics, highly unsaturated and phenolic compounds based on the aforementioned criteria, as well as MCR≥0.9 (which includes oxidized unsaturated and highly unsaturated compounds such as PS like imidazole-carboxaldehyde and PAHs) (Zhang et al., 2021b), 52.6% and 49.7% of the compounds (by intensity) can be considered as potential PS in WS<sub>A</sub> and WS<sub>F</sub>, respectively. The main compositional difference lies in polyphenolics, comprising 26.3% and 21.8% of WS<sup>A</sup> and WS<sup>F</sup> respectively. Therefore, the higher sulfate 390 formation in  $WS_A$  may be related to the higher contributions of the polyphenolics, e.g.,  $C_8H_8O_3$ .

 To summarize, we propose that in aqueous reactions, the enhanced sulfate formation in CS<sup>A</sup> and  $RS_A$  was likely due to the increased proportions (by intensity) of CHON and CHN species, potentially nitrophenols and N-heterocyclic compounds. Conversely, the increased sulfate 394 formation in  $WS_A$  appears to be linked to a higher percentage of CHO species. However, the association between detailed chemical characteristics and sulfate formation is not available in this study due to the complexity of the interactions between different chemical categories and difficulties in the interpretation of the coefficients. Future studies are needed to elucidate the relationships between sulfate formation and the chemical characteristics.

#### **3.4 Effects of Chloride and Nitrogen-containing Species on Sulfate Formation**

 Unlike the droplet experiments where RS-NaCl has the highest sulfate enhancement factor after aging, aqueous reaction results (without NaCl) show a sulfate enhancement trend of WS>CS>RS>IS, suggesting that chloride may take effect in the droplet experiments, especially in RS-NaCl system. To evaluate the effects of chloride on sulfate formation, we conducted bulk reaction experiments using RS extracts as an example with 100-200 ppm NaCl additions, where the NaCl to TOC ratio ranged from 100:1 to 200:1 to match the 100:1 to 1000:1 range in droplet 406 experiments. Interestingly, incorporating NaCl yielded contrasting results for  $RS_F$  and  $RS_A$ 407 (Figure 3). While the addition of NaCl enhanced sulfate formation in  $RS_A$ , it showed the 408 opposite trend in  $RS_F$ . The nature of the cations and ionic strength may affect the sulfate formation rate; however, previous studies have indicated that their effects are negligible (Zhang 410 and Chan, 2024; Parker and Mitch, 2016). The opposite effect of the NaCl addition on  $RS_F$  and RS<sub>A</sub>, to some extent, explains the significantly higher sulfate and SO<sub>2</sub> uptake coefficient enhancement factor for RS-NaCl in Fig. 2. Compared to the RS-based system, NaCl control experiment showed minimum (but non-zero) sulfate formation (Table 1 and Figure 3). On one hand, it supported the findings that chloride participated in the sulfate formation under light but no sulfate formation under dark (Cao et al., 2024; Tang et al., 2023; Zhang and Chan, 2024). 416 On the other hand, the opposite trend of Cl effects on RSF and RSA reflects its complex interactions with BB extracts under light and air. While direct reaction between S(IV) species 418 and  $3PS*$  may occur (Wang et al., 2020a), other pathways, i.e., interactions among halide ions, PS and oxygen should also be considered. Detailed mechanisms will be discussed later.

 Statistical analysis using the Spearman correlation coefficients, as guided by the Shapiro-Wilk test (Table S4), revealed that the CHO, CHON, and CHN species exhibited significant 422 correlations ( $|R|>0.7$ ) with the sulfate formation rate ( $p < 0.01$ , Figure S10). While PS can be the main CHO species contributing to sulfate formation, N-containing organic compounds (NOCs), i.e., CHN and CHON species, may affect the chloride contribution to the sulfate formation rate. Therefore, we selected SyrAld and VL as model CHO (PS), pyrazine (Pyz) as a model CHN, and 4-nitrocatechol (4-NC) as a model CHON to elucidate how the N containing 427 species can alter the effects of chloride on sulfate formation rate by studying the CHO+Cl, 428 CHO+CHN+Cl, and CHO+CHON+Cl systems. For SyrAld and VL, as the [Cl]o/[PS]o 429 increases,  $k_{so_4^2}$ - initially decreases and then increases. The initial decrease of  $k_{so_4^2}$ - may be 430 attributed to the quenching of  ${}^{3}P S^{*}$  by electron transfer from Cl<sup>-</sup> or loss of OH radials by 431 forming ClOH• through reaction of OH•+Cl<sup>-</sup> ↔ ClOH• (Anastasio and Newberg, 2007). Excessive chloride (e.g. 100 and 200 ppm) may generate Cl and OH radicals through 433 photoexcitation in the presence of air and water and compensate for the loss of  ${}^{3}P S^*$  or OH radicals. Previous studies have shown contradicting influence of halides on the photosensitized oxidation of organic compounds or bisulfite. Parker and Mitch (2016) and Zhang et al. (2023) attributed the significantly higher photodegradation of dienes, thioethers and acetaminophen to the formation of reactive halogen species generated by the reactions of PS and halides. Zhang 438 and Chan (2024) reported that  $\left[Cl/PS\right]_0$  in the range of 1:2 to 4:1 did not lead to significant difference in sulfate formation. The differences between the current results and the 440 aforementioned study might be attributed to the higher [Cl<sup>-</sup>/PS]<sub>0</sub> used in this study (up to 200:1) which may have been sufficient to initiate the relevant reactions, as well as the difference in

- 442 photosensitizing capacities of the PS studied (triplet quantum yield of  $0.86 \pm 0.05$  for 2-IC and  $443$  0.21  $\pm$  0.01 for VL) (Felber et al., 2021; 2020). Safiarian et al. (2023) reported that increasing 444 chloride concentrations facilitated anthracene photosensitization by producing high-level 445 reactive oxygen species (ROS). Wang et al. (2023a) found that the effects of chloride on sulfate 446 formation depended on the specific PS: enhancing sulfate production for benzophenone (BP) 447 and 3,4-dimethoxybenzaldehyde (DMB), but decreasing it for 1,4-naphthoquinone.
- 148 Incorporating CHN species yielded a 2-3-fold increase in  $k_{so_4^{2-}}$ , due to the enhanced H transfer 449 by CHN acting as H-bond acceptor (Dou et al., 2015). With the addition of NaCl, the enhanced 450 H-transfer effect by CHN was inhibited, possibly due to the consumption of  ${}^{3}P S^{*}$  by Cl. On 451 the other hand, the addition of model CHON species into PS decreased  $k_{so_4^{2-}}$ , due to the 452 consumption of  ${}^{3}P S^{*}$  by CHON species, in agreement with Wang et al. (2023b) who reported 453 increased effective quantum yield of 4-NC under co-photolysis with VL. Further addition of 454 NaCl increased the  $k_{so_4^{2-}}$ , possibly due to the consumption of 4-NC by RCS (Wang et al., 455 2024b), which, to some extent, reduced the loss of  ${}^{3}PS*$ . Generally, the addition of chloride 456 increased  $k_{so_4^{2-}}$  of PS-CHON but decreased  $k_{so_4^{2-}}$  of PS-CHN. However, the ambient air is 457 characterized by the presence of tens of thousands of chemical compounds. As a result, the 458 interplay among this diverse array of species may occur in ways that exceed current 459 understanding, necessitating additional research to investigate the interactions between 460 different organic compounds more thoroughly.



462 Figure 3. (a) Sulfate formation rate and (b) (c) bisulfite decay in RS-NaCl aqueous reactions. 1-0,

463 1-100, and 1-200 refer to the concentration ratios of TOC<sub>RS</sub> and NaCl, in which 1, 100, 200 represent

464 1 ppm, 100 ppm and 200 ppm, respectively.

## 465 **3.5 Proposed mechanism for sulfate formation**

466 A conceptual diagram of PS and chloride mediated ROS and RCS production in the oxidation 467 of S (IV) to S (VI) was shown in Fig. 4. Initially, the PS (e.g., SyrAld and VL) absorb solar 468 radiation and produce the singlet state  ${}^{1}PS*$ , which then undergoes a spin conversion through 469 intersystem crossing, leading to the formation of the triplet state  ${}^{3}P S^{*}$ . The  ${}^{3}P S^{*}$  can react with 470 molecular oxygen through energy transfer and generate singlet state  ${}^{1}O_{2}$ \*, while the  ${}^{3}PS*$  returns 471 to ground state. The  ${}^{1}O_{2}$ \* can then transform to  $O_{2}$ \* via electron transfer. The  ${}^{3}PS*$ can also react 472 with an H donor (RH, e.g., organic acids, syringol, guaiacol, Table S3), leading to the formation 473 of alkyl or phenoxy radical ( $\mathbb{R}^{\bullet}$ ) and a ketyl radical (PSH $\bullet$ ).  $\mathbb{R}^{\bullet}$  can react with  $O_2$  and form  $RO_2$ 474 radicals while PSH• can transfer an H atom to  $O_2$  and form  $HO_2^*$ , returning to its ground state 475 PS. Additionally,  ${}^{3}P S^{*}$  can react with an electron donor, e.g., Cl, and form chlorine radicals 476 and PS<sup>\*</sup>. The formed PS<sup>\*</sup> then reacts with  $O_2$  and form  $O_2$ <sup>\*</sup>, which undergoes a series of 477 reactions and form  $HO_2\bullet$ ,  $H_2O_2$  and  $OH\bullet$ . The above-mentioned reactions are the main processes 478 in the ROS pathway. Recently, Zhang and Chan(2024) have proposed that the reactive chlorine 479 species (RCS) would contribute to sulfate formation. Cao et al. (2024) proposed a mechanism 480 of OH and Cl radicals formation by  $\left[Cl - H_3O^+ - O_2\right]$  under light irradiation through an electron 481 transfer process. Our results also demonstrate that the addition of Cl will affect the oxidation 482 process of S(VI) (Figures 3, S11-S13). Combining the above, the RCS pathway was shown in yellow arrows in Figure 4. The Cl• can be formed in two pathways, photoexcitation of the [Cl-483  $- H_3 O^{\dagger} - O_2$  complex that generates Cl radicals in deliquescent BB-NaCl droplets or aqueous 485 BB-NaCl solution (Cao et al., 2024), and PS\* mediated Cl• formation via electron transfer by 486 Cl<sup>-</sup> (Corral Arroyo et al., 2019). The formed Cl• can then react with each other through radical-487 radical reactions and produce molecular Cl<sub>2</sub>. The Cl• can also react with Cl<sup>-</sup> or Cl<sub>2</sub>•, forming 488 Cl<sub>2</sub>• or Cl<sub>2</sub>. Cl• and Cl<sub>2</sub>• can also react with OH and form HOCl. <sup>3</sup>PS<sup>\*</sup> itself can also oxidize 489 the S(IV) (e.g., dissolved  $SO_2$  or bisulfite) to S(VI). However, significantly lower sulfate 490 formation was found in the presence of  $N_2$  than air (Figure S4), highlighting the importance of 491 secondary oxidants compared to direct PS\* oxidation. As a consequence, these reactive species, 492 e.g., OH•/HO<sub>2</sub>•/O<sub>2</sub>• and Cl•/Cl<sub>2</sub>• may all participate in the oxidation of S(IV) to S(VI). In 493 addition, nitrogen-containing heterocyclic compounds such as pyrazine can act as H-bonding 494 acceptor and facilitate the H transfer, which then generates more ROS (Dou et al., 2015). Note 495 that although ROS and RCS pathways both contribute to the oxidation from S(IV) to S(IV), 496 they may act as competitive relationships due to the co-consumption of PS\*. Therefore, 497 different Cl effects may occur regarding various combinations of reactants (Figure 3, promoting 498 effect in  $RS_A$ , inhibiting effects on  $RS_F$ ).

499



 Figure 4. Conceptual diagram of PS and chloride-mediated ROS and RCS production, in the oxidation processes from S(IV) to S(VI)

## **4 Atmospheric Implication**

 This study provided laboratory evidence that the PS in biomass burning extracts can enhance the sulfate formation in NaCl particles, primarily by triggering the formation of secondary oxidants under light and air, with less contribution of direct photosensitization via triplets 507 (evidenced by  $N_2$  atmosphere, Figure S4). The sulfate formation rates of  $BB_F-NaCl$  particles 508 were ~10 folds higher than that of  $IS_F-NaCl$ , following the trends of  $CS_F-NaCl > RS_F$ -509 NaCl>WS<sub>F</sub>-NaCl>IS<sub>F</sub>-NaCl. Upon UV exposure, the sulfate formation trends shifted to  $RS_A$ - NaCl>CSA-NaCl>WSA-NaCl>ISA-NaCl, which might be explained by the effects of chloride (evidenced by aqueous reactions, Figure 3and Table 1). Interestingly, the incorporation of Cl- 512 into bulk solutions increased the sulfate formation rate in  $RS_A$ , while decreased it in  $RS_F$ . This seems to be different from our group's previous work where no significant sulfate formation 514 rate was found with the addition of Cl (Zhang and Chan, 2024). The difference can be explained 515 by the following reasons: 1) differences in PS/Cl<sup>-</sup>, the prior study might have used an

516 insufficient PS/Cl<sup>-</sup> ratio (2:1-1:4) while the current one significantly expanded it to 1:200. 2) differences in photosensitizing capacity: the former study used a strong PS, while the current study focused on the real BB (using TOC as a metric, with only a small portion of TOC considered as PS). 3) the complexity of the reaction system, the former study focused on mixing two individual species, while in real BB extracts, more complicated reactions may occur. Furthermore, our results using model PS show that although additional model CHN species would increase the sulfate formation by expedited H transfer via acting as H-bond acceptor, the addition of chloride could inhibit the sulfate formation rate, suggesting that the RCS pathway was less efficient in sulfate formation compared to ROS pathway in PS-CHN bulk system (Figure S11 and S12).

 While our prior study has examined the potential interplay between chloride and PS at limited mixing ratios (up to 4:1 in bulk solution) (Zhang and Chan, 2024), this work expanded the Cl- /PS ratio to a broader range (200:1) and systematically identified the interactions among different organics, including PS, NOCs, and chloride, using sulfate formation as a compass. This highlights the importance of studying secondary aerosol formation in mixed experimental systems under air pollution complex. Our work suggests that in coastal regions heavily influenced by anthropogenic emissions like biomass burning, especially those near the rice- growing regions or affected by transported wildfire smoke, such as Guangdong, Fujian and Taiwan, the transported BB plumes together with the high RH (Cheung et al., 2015) and abundant reactive gases, would play an important role in sulfate and potentially secondary organic aerosol formation.

### **Data availability**

 Datasets are available upon request to the corresponding author, Chak K. Chan ([chak.chan@kaust.edu.sa\)](mailto:chak.chan@kaust.edu.sa).

### **Author contributions**

 RT and CC conceptualized and designed the study. YQ and YC collected the samples. RT performed the experiments, data analysis and wrote the draft. JM provided assistance in data

processing. All the authors reviewed, edited and contributed to the scientific discussions.

## **Competing interests**

The authors declare no conflicts of interest.

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