Response to the editor

We are grateful to the editor for the careful review of our manuscript. We have corrected all of the grammar errors, typos, and format issues suggested by the editor and found by ourselves in the Main text and Supplement. Please see below the marked-up revised manuscript showing the changes.

Comments to the Author:

The authors have appropriately addressed the comments of the two anonymous referees and they have modified their manuscript accordingly. However, many alterations and corrections are needed for the Main text and Supplement before the manuscript can be published in ACP.

For the Main text:

Line 16: Replace "over years" by "over the years".

Line 23: Replace "in two" by "in the two".

Line 25: Replace "than in" by "compared to".

Line 30: Replace "with that" by "with those".

Lines 30-31: Replace "averagely contributed" by "contributed, on average,".

Line 32: Replace "76% of" by "76% of the".

Lines 50 and 51: Replace "Noziere" by "Nozière".

Line 50: Replace "organic matters in SOA are" by "organic matter in SOA is".

Lines 63, 272, and 540: Replace "Gomez-Gonzalez" by "Gómez-González".

Line 67: Replace "of OS" by "of the OS" and replace "in ambient" by "in the ambient".

Line 76: Replace ", which can" by " and can".

Line 79: Replace "In atmospheric" by "In the atmospheric".

Line 87: Replace "react with" by "react with the".

Line 88: Replace "in ambient" by "in the ambient".

Line 89: Abbreviations and acronyms, here OM, should be defined (written full-out) when first used.

Thus, replace "OM" by "organic mass (OM)".

Line 113: Replace "same or similar with" by "same as or similar to".

Line 117: Replace "Up to date" by "Up to now".

Line 122: Abbreviations and acronyms, here UPLC-ESI-QToF-MS, should be defined (written fullout) when first used in the Main text (outside of the Abstract). I suggest replacing "UPLC-ESI-QToF-MS" here by "an ultra-performance liquid chromatography quadrupole time-of-flight mass spectrometer equipped with an electrospray ionization (ESI) source (UPLC-ESI-QToF-MS)".

Line 125: Replace "were probed" by "were investigated".

Line 145: Replace "waters" by "Waters".

Line 154: Abbreviations and acronyms, here MS/MS, should be defined (written full-out) when first used. I suggest replacing "MS/MS" here by "tandem mass spectrometry (MS/MS)"; as a consequence, in lines 161-162, "tandem mass spectrometry (MS/MS)" should the be replaced by "MS/MS".

Line 158: Replace ")were" by ") were".

Line 167: Replace "by Shanghai" by "by the Shanghai".

Line 173: Replace "were analyzed" by "were determined".

Line 199: Replace "of OS" by "of the OS".

Line 201: Replace "from LC" by "from the LC".

Line 213: Replace "in two years" by "in the two years".

Line 232: Replace "Figure 3a, b show" by "Figure 3a and b show".

Line 234: Replace "than CHO" by "than the CHO".

Line 236: Replace "and a few" by "and for a few".

Line 240: Replace "to observed" by "to the observed".

Line 242: Replace "species were significantly" by "species was significantly".

Line 244: Replace "to biogenic" by "to that of biogenic".

Line 255: Replace "originate from" by "originates from".

Line 259: Replace "with that in" by "as in". Line 262: Replace "of identified" by "of the identified".

Line 263: Replace ", in polluted" by " in polluted".

Line 265: Replace "signals were not" by "signals was not".

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Line 285: Replace "in two" by "in the two".
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Line 295: Replace "Isoprene-derived OS" by "The isoprene-derived OS".

Line 313: Replace "Since 2-MT-OS" by "2-MT-OS".

Line 315: Replace "suggests that" by "suggest that".

Line 317: Replace "important SOA" by "important for SOA".

Line 327: Replace "expect for" by "except for".

Line 332: Replace "NOSs" by "NOS".

Line 334: Replace "Monoterpene-derived OS" by "The monoterpene-derived OS".

Line 344: Replace "in two" by "in the two".

Line 350: Replace "Among NOS species, the" by "Among the NOS species,".

Line 353: Replace "than observed" by "than that observed".

Lines 358-359: Replace "Although the" by "Although".

Line 363: Replace "NOS species" by "the NOS species".

Line 370: Replace "were quantified. The inter-annual variations of" by "was quantified. The inter-annual variation of".

Line 371: Replace "to biogenic OS" by "to the biogenic OS one".

Line 372: Replace "Among quantified OSa, the" by "Among the quantified OSa,".

Line 375: Replace "time with the" by "time as the".

Line 388: Replace "species averagely accounted for 19% of" by "species accounted, on average, for 19% of the".

Line 396: Replace "with those observed" by "with that observed".

Line 403: Replace "2019). The" by "2019).".

Line 403-404: Replace "most of" by "most of the".

Line 410: Replace "0.88). The" by "0.88).".

Line 411: Replace "2017), which was strongly correlated with the" by "2017); it was strongly correlated with".

Line 413: Replace "GA, where the" by "GA, where" and replace "most of" by "most of the".

Line 415: Replace "concentrations of" by "concentrations of the".

Line 432: Replace "seasons were also" by "seasons was also".

Line 435: Replace "that most" by "where most".

Line 437: Replace "that that the" by "that the".

Line 446: Replace "6, OS" by "6, the OS".

Line 458: Replace "of ion" by "of the ion".

Line 464: Replace "on average accounted" by "accounted, on average,".

Line 470: Replace "on average contributed to 19% of the concentration of quantified OS species.

The" by "contributed, on average, to 19% of the concentration of the quantified OS species.".

Line 472: Replace "76% of" by "76% of the".

Line 474: Replace "implies that" by "imply that".

Line 490: Replace "of CHOS signals that arose mainly from anthropogenic sources were" by "of the CHOS signals that arose mainly from anthropogenic sources was".

Lines 509-510, 647, and 757-759: Titles of journal articles should be in lower case, not in Title Case.

Lines 627-628: Replace "amp,apos, Dowd" by "O'Dowd".

Line 638: Replace "88," by "88, 3319-3322,".

Lines 683, 762, 764, and 806: For references with only 2 authors, there should be no comma before the "and" preceding the second author. Note that for references with 3 or more authors, there should be a comma before the "and" preceding the last author.

Line 709: Replace "117," by "117, D01207,".

Line 714: Replace "Kaberer" by "Kalberer".

Lines 714-719: "Noziere et al., 2015" should come after "Noziere et al., 2010".

Line 719: Replace "37," by "37, L05806,".

Line 796: Replace "8345" by "36, 8345".

Line 813: Replace "224," by "224, 117180,".

Line 818: Replace "724," by "724, 138134,".

Line 882: Replace "relative humid" by "relative humidity".

Line 889: Replace "The intensity fraction" by "Intensity fraction".

Line 890: Replace "The number of" by "Number of".

Line 896: Replace "The concentrations of different types of the quantified OS over four" by

"Concentrations of different types of the quantified OS over the four".

Line 899: Replace "The quantified" by "Quantified". Line 905: Replace "The quantified" by "Quantified".

Line 906: Replace "of circles" by "of the circles".

Line 907: Replace "with aerosol" by "with the aerosol".

Response: Thanks. We have corrected all of these issues in the Main text.

For the Supplement:

Table S3, first line of heading: Replace "The dates" by "Dates".

Table S3, first line of footnote: Replace "in the parentheses are the concentrations (ppm) of OS" by "in parentheses are the concentrations (ppm) of the OS".

Table S4, first line of heading: Replace "The molecular" by "Molecular". Figure S1, first line of caption: Remove "The " after "Figure S1".

Response: Thanks. We have corrected all of these issues in the Supplement.

- Organosulfates in atmospheric aerosols in Shanghai, China: seasonal and interannual variability,
- 2 origin, and formation mechanisms
- 3 Yao Wang¹, Yue Zhao^{1,2,*}, Yuchen Wang³, Jian-Zhen Yu^{3,4}, Jingyuan Shao⁵, Ping Liu¹, Wenfei Zhu¹, Zhen Cheng¹, Ziyue
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Abstract

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42 43 Organosulfates (OS) are ubiquitous in the atmosphere and serve as important tracers for secondary organic aerosols (SOA). Despite intense research over the years, the abundance, origin, and formation mechanisms of OS in ambient aerosols, in particular in regions with severe anthropogenic pollution, are still not well understood. In this study, we collected filter samples of ambient fine particulate matter (PM_{2.5}) over four seasons in both 2015/2016 and 2018/2019 at an urban site in Shanghai, China, and comprehensively characterized the OS species in these PM_{2.5} samples using an ultra-performance liquid chromatography quadrupole time-of-flight mass spectrometer equipped with an electrospray ionization (ESI) source (UPLC-ESI-QToF-MS). We find that while the concentration of organic aerosol (OA) decreased by 29% in 2018/2019, compared to that in 2015/2016, mainly a result of the reduction in anthropogenic pollutant emissions in eastern China, the annually averaged concentrations of 35 quantified OS were overall similar in the two years (65.5±77.5 ng m⁻³, 0.57%±0.56% of OA in 2015/2016 versus 59.4±79.7 ng m⁻³, 0.66%±0.56% of OA in 2018/2019), suggesting an increased contribution of SOA to OA in 2018/2019 compared to 2015/2016. Isoprene- and monoterpene-derived OS were the two most abundant OS families, on average accounting for 36.3% and 31.0% of the quantified OS concentrations, respectively, during the two sampling years, suggesting an important contribution of biogenic emissions to the production of OS and SOA in Shanghai. The abundance of biogenic OS, particularly those arising from isoprene, exhibited strong seasonality (peaked in summer) but no significant interannual variability. In contrast, the quantified anthropogenic OS had little seasonal variability and declined in 2018/2019 compared with those in 2015/2016. The C₂/C₃ OS species that have both biogenic and anthropogenic origins contributed, on average, to 19.0% of the quantified OS, with C₂H₃O₆S⁻, C₃H₅O₅S⁻, and C₃H₅O₆S⁻ being the most abundant ones, together accounting for 76% of the C₂/C₃ OS concentrations in 2015/2016 and 2018/2019. 2-Methyltetrol sulfate (2-MT-OS, C₅H₁₁O₇S²) and monoterpene-derived C₁₀H₁₆NO₇S⁻ were the most abundant OS and nitrooxy-OS in summer, on average contributing to 31% and 5% of the quantified OS, respectively, during the summertime of the sampling years. The substantially larger concentration ratio of 2-MT-OS to 2-methylglyceric acid sulfate (2-MA-OS, C₄H₇O₇S⁻) in summer (6.8-7.8) than in other seasons (0.31-0.78) implies that low-NO_x oxidation pathways played a dominant role in isoprene-derived SOA formation in summer, while high-NO_x reaction pathways were more important in other seasons. We further find that the production of OS was largely controlled by the level of O_x ($O_x = O_3 + NO_2$), namely, the photochemistry of OS precursors, in particular in summer, though sulfate concentration, aerosol acidity, as well as aerosol liquid water content (ALWC) that could affect the heterogeneous chemistry of reactive intermediates leading to OS formation also played a role. Our study provides valuable insights into the characteristics and mechanisms of OS formation in a typical Chinese megacity and implies that mitigation of O_x pollution can effectively reduce the production of OS and SOA in eastern China.

1 Introduction

Secondary organic aerosol (SOA) accounts for a significant fraction of atmospheric fine particulate matter (PM_{2.5}) (Jimenez et al., 2009; Huang et al., 2014) and contributes significantly to deteriorated air quality and Earth's climate forcing (Ramanathan et al., 2001; Mahowald, 2011; Shrivastava et al., 2017; Huang et al., 2014). SOA consists of thousands of organic compounds that are diverse in molecular properties. Identification and quantification of the composition of SOA are essential for understanding the composition, the chemistry of formation and evolution, properties, and climate and health impacts of SOA (Hoffmann et al., 2011; Nozière et al., 2015). However, currently only a small portion of organic matter in SOA is identified as specific compounds (Hoffmann et al., 2011; Nozière et al., 2015; Johnston and Kerecman, 2019). Organosulfates (OS) are important constituents of SOA and have been frequently detected in both polluted and clean environments (Iinuma et al., 2007a; Surratt et al., 2008; Claeys et al., 2010; Froyd et al., 2010; Hawkins et al., 2010; Hatch et al., 2011; Lin et al., 2012a; Stone et al., 2012; Hansen et al., 2014; He et al., 2014; Ma et al., 2014; Tao et al., 2014; Liao et al., 2015; Shakya and Peltier, 2015; Kourtchev et al., 2016; Meade et al., 2016; Wang et al., 2016b; Hettiyadura et al., 2017; Huang et al., 2018; Le Breton et al., 2018; Wang et al., 2018; Hettiyadura et al., 2019; Wang et al., 2019a; Brüggemann et al., 2020). It has been estimated that OS accounted for 6-12% of total sulfur in a rural area in K-puszta, Hungary (Lukacs et al., 2009), 1.3% of fine particulate organic mass (FPOM) in Fairbanks, Alaska (Shakya and Peltier, 2013), and 1-13% of FPOM across the continental United States (Tolocka and Turpin, 2012; Shakya and Peltier, 2015). Studies have also shown that OS can affect aerosol properties such as acidity, viscosity, hygroscopicity, and light-absorbing properties (Nguyen et al., 2012; Hansen et al., 2015; Estillore et al., 2016; DeRieux et al., 2018; Fleming et al., 2019; Riva et al., 2019; Olson et al., 2019).

Chamber studies have revealed that OS can originate from the (photo)oxidation of both biogenic precursors such as isoprene (Gómez-González et al., 2008; Surratt et al., 2007b; Surratt et al., 2007a), monoterpenes (Iinuma et al., 2007a; Surratt et al., 2007a; Surratt et al., 2007b; Iinuma et al., 2009), sesquiterpenes (Chan et al., 2011), and 2-methyl-3-buten-2-ol (Zhang et al., 2012), as well as anthropogenic precursors such as polycyclic aromatic hydrocarbons, long-chain alkanes, naphthenes (Riva et al., 2015; Riva et al., 2016b), and diesel and biodiesel fuel vapors (Blair et al., 2017) in the presence of sulfate aerosol or SO₂. Many of the OS observed in these chamber studies have also been detected in the ambient atmosphere, among which isoprene- and monoterpene-derived OS are usually most abundant in forested, rural, and even urban areas (Stone et al., 2012; Ma et al., 2014; Meade et al., 2016; Hettiyadura et al., 2019; Kourtchev et al., 2016; Kristensen and Glasius, 2011; Wang et al., 2019a; He et al., 2014; Hatch et al., 2011; Surratt et al., 2008; Hettiyadura et al., 2017).

In addition to the precursors, detailed formation mechanisms of OS have also been widely studied (Brüggemann et al., 2020). The acid-catalyzed ring-opening reaction of epoxides was established to be an important mechanism for the formation of OS (Surratt et al., 2010; Lin et al., 2012b; Iinuma et al., 2009; Zhang et al., 2014), in particular for isoprene-derived OS (Surratt et al., 2010; Lin et al., 2012b; Hatch et al., 2011). 2-Methyltetrol sulfate (2-MT-OS, C₅H₁₁O₇S⁻), formed via reactive uptake of isoprene epoxide (IEPOX) on sulfate, is one of the most abundant OS in atmospheric aerosol (Chan et al., 2010; Liao et al., 2015) and can contribute up to 12.6% of the organic carbon mass in Atlanta, GA (Hettiyadura et al., 2019). Another OS formation pathway is the nucleophilic substitution of tertiary organonitrates by inorganic sulfate. Darer et al. (2011) found that tertiary organonitrates are thermodynamically unstable and can undergo nucleophilic substitution with sulfate to generate

OS rapidly. This mechanism can also explain the formation of some nitrooxy-OS (NOS). In the atmospheric aqueous phase, sulfate radicals that can be produced by oxidation of S(IV) species in the presence of transition metal ions (TMI) (Grgic et al., 1998; Herrmann, 2003) or by OH radical reaction with bisulfate (Herrmann, 2003; Jiang et al., 1992) can also react with unsaturated organic compounds to form OS. Laboratory studies have shown that a large number of OS were produced by the bulk aqueous-phase oxidation of aromatic compounds, isoprene or its oxidation products, methyl vinyl ketone (MVK) and methacrolein (MACR), in the presence of inorganic sulfate or peroxodisulfate under irradiation (Nozière et al., 2010; Schindelka et al., 2013; Huang et al., 2020), or in the presence of S(IV) and TMI under dark conditions (Huang et al., 2019). However, field observational evidence for this mechanism is still lacking. In addition, reactive uptake of SO₂ on organic aerosol can also result in the production of OS. Laboratory studies have found that sulfur dioxide (SO₂) could react with the C=C bond in unsaturated fatty acids under dark conditions to form OS (Shang et al., 2016). Such OS have been detected in the ambient atmosphere with an estimated contribution of 0.3%-0.9% to organic mass (OM) in PM_{2.5} in southern China (Zhu et al., 2019). Recent studies have also shown efficient production of OS from heterogeneous/aqueous-phase reactions of SO₂ with organic peroxide-containing aerosol and SOA (Wang et al., 2019b; Yao et al., 2019). Such OS production was found to be mainly a result of the direct reaction between SO₂ and peroxides, rather than acid-catalyzed reaction involving inorganic sulfate (Wang et al., 2019b). Currently, the acid-catalyzed ring-opening reaction of IEPOX has been the most well-studied mechanism and proved to be important in atmospheric OS formation by both field and modelling studies (Surratt et al., 2010; Kourtchev et al., 2016; Hettiyadura et al., 2019; Hatch et al., 2011; Chan et al., 2010; McNeill et al., 2012; Worton et al., 2013; He et al., 2018; Pye et al., 2013). However, atmospheric importance of other OS formation mechanisms remains to be evaluated.

The OS formation pathways aforementioned can be affected by aerosol properties such as acidity, aerosol liquid water content (ALWC), and sulfate concentration. There is ample evidence from laboratory studies that increased aerosol acidity significantly enhances the production of OS from acid-catalyzed reactions (Iinuma et al., 2007b; Chan et al., 2011; Surratt et al., 2007a; Surratt et al., 2007b; Zhang et al., 2012), while field studies have shown that the abundance of OS was not or only weakly correlated with aerosol acidity in some locations (Nguyen et al., 2014; Budisulistiorini et al., 2015; Brüggemann et al., 2017; Rattanavaraha et al., 2017), suggesting the existence of other factors (e.g., ALWC, sulfate content, etc.) that control OS formation in these areas. ALWC has dual effects on OS formation. On one hand, elevated ALWC can reduce the viscosity and/or inhibit the liquid-liquid phase separation of aerosol, which would favor the dissolution and mixing of reactive intermediates such as IEPOX and multifunctional aldehydes in aqueous sulfate aerosol (Shiraiwa et al., 2011; Liao et al., 2015; McNeill et al., 2012) or SO₂ in organic aerosol (Shang et al., 2016; Passananti et al., 2016; Yao et al., 2019), thereby enhancing OS formation. On the other hand, high ALWC would decrease aerosol acidity via dilution, hence inhibiting the acid-catalyzed OS formation. High ALWC may also promote the hydrolysis of OS in aqueous aerosol (Darer et al., 2011).

Quantification of OS is important for understanding their abundance and the chemistry of formation and evolution in the atmosphere. Owing to the lack of authentic standards, the quantification of OS remains a challenging task. Recently, several research groups have synthesized a series of authentic standards (e.g., glycolic acid sulfate, lactic acid sulfate, hydroxyacetone sulfate, 2-methyltetrol sulfate, benzyl sulfate, α -pinene sulfate, β -pinene sulfate, and limonene sulfate) that are structurally the same as or similar to the OS found in atmospheric aerosols (Kundu et al., 2013; Staudt et al., 2014; Hettiyadura et al.,

2015; Wang et al., 2017; Olson et al., 2011; Huang et al., 2018; Budisulistiorini et al., 2015). They used these authentic standards to quantify OS in ambient aerosols and provided important constraints on the abundance, origin, and chemistry of OS in the atmosphere.

Up to now, there are few studies characterizing atmospheric OS in areas with severe anthropogenic pollution. Situated in the eastern Yangtze River Delta (YRD) of China, Shanghai has a population of more than 24 million and is plagued by air pollution (Behera et al., 2015; Wang et al., 2016a). Here we conducted a comprehensive investigation of the molecular composition, abundance, sources, and formation processes of OS in ambient aerosols in Shanghai. More than 150 ambient PM_{2.5} samples collected over four seasons during both 2015/2016 and 2018/2019 in urban Shanghai were analyzed using an ultra-performance liquid chromatography quadrupole time-of-flight mass spectrometer equipped with an electrospray ionization (ESI) source (UPLC-ESI-QToF-MS), and 35 OS were quantified using 7 synthesized and commercially purchased OS standards. Seasonal and interannual variations of OS, in response to the changes in emissions, meteorological conditions, and PM_{2.5} chemical compositions, were comprehensively characterized, and the influencing factors such as aerosol acidity, ALWC, sulfate content, oxidant level and so on for OS formation were investigated. This study would help to understand the characteristics and mechanisms of OS and SOA production under the strong influence of anthropogenic pollution in Chinese megacities.

2 Materials and methods

2.1 Ambient Sample collection

In total 156 ambient PM_{2.5} samples were collected from 8 April 2015 to 16 January 2016 and from 23 October 2018 to 5 August 2019 in Shanghai, China. The sampling site is located on the rooftop of a 20-meter-tall teaching building on the Xuhui Campus of Shanghai Jiao Tong University at 31.201°N, 121.429°E, which is in downtown and surrounded by residential and commercial areas (see Fig. 1a, b). There is a main street 230 m east to the sampling site. The PM_{2.5} samples were collected on pre-baked (550 °C, 8 h) quartz-fiber filters (Whatman) from 9:00 am to 8:00 am of the next day using a high-volume sampler (HiVol 3000, Ecotech) at a flow rate of 67.8 m³ h⁻¹. The collected samples were wrapped in pre-baked (550 °C, 8 h) aluminum foil and stored at -20 °C before analysis.

2.2 Organosulfate measurement with UPLC-ESI-ToF-MS

An aliquot of ~17 cm² was removed from each filter sample and extracted in 3 mL of methanol (LC-MS grade, CNW Technologies GmbH) twice under sonication in an ice bath at $4 \, {}^{\circ}\!\underline{C}$ for 30 min. The extracts derived each time were combined and filtered through a 0.45 μ m poly tetra fluoroethylene (PTFE) syringe filter (CNW Technologies GmbH) to remove insoluble materials, and subsequently concentrated to 250 μ L with a gentle stream of ultra-high-purity nitrogen (Shanghai Likang Gas Co., Ltd). The resulting extracts were mixed with ultrapure water (milliQ, 18.2 M Ω ·cm) of the same volume and centrifugated at 12000 rpm and $4 \, {}^{\circ}\!\underline{C}$ for 20 min using a centrifuge (Cence, TGL-16M) to get supernatant for analysis.

The resulting solutions were analyzed using an Acquity UPLC (Waters) coupled to a Xevo G2-XS QToF-MS (Waters) having a mass resolving power of \geq 40000 and equipped with an ESI source. The analytes were separated by an ethylene bridged hybrid (BEH) C_{18} column (2.1×100 mm, 1.7 μ m particle size, Waters) at 50 °C. A gradient elution procedure was performed

using water (A) and methanol (B) both containing 0.1% acetic acid (v/v) as the eluents: A was maintained at 99% for 1.5 min, decreased to 46% in 6.5 min and to 5% in 3 min, then decreased to 1% in 1 min and held for 2 min, finally returned to 99% in 0.5 min and held for 1.5 min to equilibrate the column. The total eluent flow rate was 0.33 mL min⁻¹ and the sample injection volume was 2.0 μ L. The ESI source was operated in the negative ion mode under optimum conditions as follows: capillary voltage 2.0 kV, sampling cone voltage 40 V, source offset voltage 80 V, source temperature 115°C, desolvation gas temperature 450°C, cone gas 50 L h⁻¹, desolvation gas 900 L h⁻¹.

2.3 Auxiliary measurements

 Meteorological parameters, including temperature, relative humidity (RH), and wind speed (WS) were continuously monitored by the Shanghai Hongqiao international airport station, which is 9 km west to the sampling site (Fig. 1c). The concentrations of SO₂, nitrogen dioxide (NO₂), O₃ and PM_{2.5} were measured by a state-controlled air quality monitoring station on the Xuhui Campus of Shanghai Normal University, which is 4.5 km southwest to the sampling site for the PM_{2.5} filter samples (Fig. 1c). Organic carbon (OC) and elemental carbon (EC) in PM_{2.5} filter samples were measured by a thermal-optical multiwavelength carbon analyzer (DRI Model 2015). The concentration of OM was derived by multiplying the OC by 1.6 (Tao et al., 2017). Water-soluble inorganic compounds including sulfate, nitrate, chloride, ammonium, potassium, and calcium were determined with an ion chromatograph (Metrohm MIC). The seasonal and annual average values of meteorological parameters and concentrations of trace gases, PM_{2.5} and its major components in 2015/2016 and 2018/2019 are listed in Table S1.

2.4 Estimation of aerosol liquid water content and pH

The ISORROPIA-II thermodynamic model (Fountoukis and Nenes, 2007) was employed to predict ALWC and aerosol pH.

The aerosol water-soluble inorganic ion concentrations, as well as temperature and RH were used as the model input. The

model was run in the forward model for metastable aerosol, which was shown to give a more accurate representation of aerosol pH than using the reverse-mode calculations when with only aerosol data input (Guo et al., 2015; Hennigan et al., 2015). ISORROPIA-II calculates the equilibrium concentration of aerosol hydronium ions (H^{+}_{air}) per volume of air ($\mu g m^{-3}$), along with ALWC ($\mu g m^{-3}$). The aerosol pH was then derived by

$$pH = -\log_{10}(H_{aq}^{+}) = -\log_{10}\frac{1000H_{air}^{+}}{ALWC},$$
 (1)

where H^+_{aq} is the concentration of hydronium ions in aqueous aerosol (mol L^{-1}). In this study, ALWC associated with organic aerosol and its influences on aerosol pH were not considered. However, previous studies showed that water uptake by organic aerosol only contributed to a minor fraction (5%) of total ALWC and had a negligible influence on aerosol pH in haze events in China (Liu et al., 2017). The seasonally and annually averaged ALWC and aerosol pH levels in 2015/2016 and 2018/2019 are also given in Table S1.

2.5 Quality control

The extraction efficiency of OS species in filter samples was evaluated by measuring the recovery of ten different OS standards (see Table S2). The synthesized and commercially purchased OS standards were spiked into blank and pre-baked quartz filters, followed by extraction and analysis with the same procedures for ambient samples. The recoveries of OS standards were about 84-94% except for Δ -Carene OS, Lactic acid sulfate, and Glycolic acid sulfate, the recovery of which was 66%, 72.5%, and 77.8%, respectively (see Table S2). This result suggests a fairly high extraction efficiency for the majority of OS species in this study.

In addition, we evaluated the matrix effect on the signal response of OS by comparing the measured signal intensity of OS standards added to the extracts of ambient $PM_{2.5}$ filter samples with that of pure OS standard solutions. Table S3 gives the ratios of measured signal intensity of OS standards in filter sample extracts to that in pure solutions. As for the standards that were already present in the samples, we subtracted the response in the sample from the total (sample + standard) before calculating the ratio. Most of the OS standards had a ratio around 1, suggesting no obvious matrix effect on the measurement of the majority of OS species. However, the two smallest OS standards, methyl sulfate and glycolic acid sulfate that were the very first species eluted from the LC column, had a ratio significantly smaller than 1, suggesting the inhibited ionization of these two OS likely by the highly soluble and polar species in the filter samples that were co-eluted with these two OS. We note that the matrix effect for these two OS is dependent on the $PM_{2.5}$ mass loading. For example, the signal ratio of glycolic acid sulfate standard measured in filter sample extracts versus in pure solutions ranged from 0.17-0.31 (Exps. 1-2) for very polluted days to 0.45-0.53 for clean days (Exps. 3-4). This implies that the abundance of glycolic acid sulfate in ambient aerosols reported here may be underestimated by a factor of 2-6 due to the matrix effect.

3 Results and Discussion

3.1 Overview of pollution characteristics during sampling periods

Figure 2 shows the time series of meteorological parameters, O₃, NO₂, SO₂, PM_{2.5} and its major components, as well as H⁺_{aq}

and ALWC during the sampling periods. The average values (concentrations) of each parameter (species) were given in Table S1. The meteorological conditions (wind speed, temperature, and RH) were overall similar in 2015/2016 and 2018/2019. While the NO₂ concentration decreased from 27.0±13.0 ppb in 2015/2016 to 21.3±10.3 ppb in 2018/2019, the O₃ level had no obvious difference in the two years (29.8±15.2 ppb in 2015/2016 versus 29.6±13.9 ppb in 2018/2019), consistent with the nonlinear response of O₃ production to precursor emissions (Liu and Wang, 2020). The annual average mass loading of PM_{2.5} declined by 34.5% in 2018/2019 (38.6±24.0 µg m⁻³) compared to that 2015/2016 (59.0±37.9 µg m⁻³), largely driven by the strong decrease in the abundance of OM (29.1%) and sulfate (37.4%). The decrease of PM_{2.5}, OM, and sulfate concentrations from 2015/2016 to 2018/2019 reflects a significant reduction in anthropogenic pollutant emissions in eastern China in recent years. In contrast to OM and sulfate, the concentration of nitrate had little change between 2015/2016 (8.8±8.9 μg m⁻³) and 2018/2019 (8.4±7.8 μg m⁻³), despite an obvious decrease in NO₂ concentration. This is at least partly a result of reduced aerosol acidity (H⁺_{ao}, see Fig. 2 and Table S1) and thereby enhanced partitioning of HNO₃ into the particle phase. Furthermore, the nitrate concentration showed a strong seasonality, ranging from 1.0±1.1 and 3.4±3.2 µg m⁻³ in summer to 16.6±10.0 and 14.1±10.0 μg m⁻³ in winter in 2015/2016 and 2018/2019, respectively, partly owing to the seasonal variation of temperature and aerosol acidity that modulates the gas/particle partitioning of nitrate (Fisseha et al., 2006; Guo et al., 2015; Griffith et al., 2015; Guo et al., 2016). A similar strong reduction in PM_{2.5} concentration and variations in aerosol composition over the past several years were observed in different regions in China (Tao et al., 2017; Wang et al., 2020b; Ding et al., 2019a; Wen et al., 2018). As a result of strong reductions in inorganic ion concentrations, ALWC decreased dramatically in 2018/2019 (14.8±20.4 μg m⁻³), compared to that in 2015/2016 (24.4±27.0 μg m⁻³). In short, anthropogenic pollutant emissions, as well as aerosol concentration and composition varied significantly between 2015/2016 and 2018/2019 in Shanghai, which, as will be discussed below, has important implications for the production of OS in ambient aerosols.

3.2 Molecular composition of sulfur-containing organic compounds

The organic compounds in ambient PM_{2.5} identified using UPLC-ESI(-)-QToF-MS were classified into four groups based on their elemental composition, i.e., CHO, CHON, CHOS, and CHONS. Figure 3a and b show the average mass spectra of organic compounds in PM_{2.5} over a typical winter (21-26 January 2019) and summer (23-28 July 2019) pollution episode. The S-containing compounds were overall larger in molecular size than the CHO and CHON compounds, likely because of the addition of a sulfate group to the molecule. The molecular weight (MW) of most S-containing compounds was between 100-400 Da, and for a few between 400-700 Da. The high-MW CHOS species (400-700 Da, see Table S4) showed a larger contribution in winter than in summer, suggesting that they are more likely to arise from anthropogenic sources than biogenic emissions. Figure 3c shows the signal contribution of different compound categories as well as concentrations of sulfate, OM, and quantified OS, and Fig. 3d, e shows the number of identified organic compounds in each category during two pollution episodes. The CHOS compounds contributed most by signal and number to the observed organic compounds in both winter and summer. The signal contributions and number of unquantified CHOS and CHONS did not vary significantly from winter to summer, whereas the signal contribution of quantified CHOS and CHONS species was significantly larger in summer than in winter (on average 15% vs. 7% for CHOS and 11% vs. 7% for CHONS). As will be discussed later, the abundance of quantified anthropogenic OS was fairly constant across different seasons, in striking contrast to that of biogenic OS that showed strong seasonal variability. Therefore, lack of seasonal variability for unquantified CHOS and CHONS implies that

they may originate mainly from anthropogenic sources. In addition, both signal intensity and the number of CHO species increased significantly in summer, compared to those in winter. In contrast, CHON compounds contributed substantially more to the observed signals in winter than in summer (on average 25% vs. 7%), though their numbers are quite similar during the two periods. This suggests an enhanced production and/or suppressed depletion of nitrogen-containing organic species in winter.

The CHOS compounds with an O/S ratio of \geq 4 were assigned as potential OS species. Similarly, the CHONS compounds with an O/(N+S) ratio of \geq 7 could be assigned as potential NOS species (Lin et al., 2012a). The $C_8H_{17}O_4S^-$, $C_8H_{15}O_4S^-$, and $C_5H_{11}O_4S^-$ were the highest OS peaks observed in the pollution episode in winter. The $C_8H_{17}O_4S^-$ and $C_8H_{15}O_4S^-$ species may be derived from the photooxidation of diesel fuel vapors according to previous chamber studies (Blair et al., 2017). The $C_5H_{11}O_4S^-$ species was correlated well with $C_8H_{17}O_4S^-$ in 2015/2016 (r=0.76) and 2018/2019 (r=0.84), suggesting it may also be derived from diesel fuel vapors. The highest NOS peak in winter was $C_{10}H_{16}NO_7S^-$, which likely originates from monoterpene oxidation (Surratt et al., 2008). The $C_5H_{11}O_7S^-$, $C_{15}H_{29}O_5S^-$, and $C_{13}H_{25}O_5S^-$ were observed among the highest OS peaks in the summer pollution episode. The $C_5H_{11}O_7S^-$ is an IEPOX-derived OS species (Surratt et al., 2010), while $C_{15}H_{29}O_5S^-$ and $C_{13}H_{25}O_5S^-$ may be derived from the oxidation of diesel fuel vapors (Blair et al., 2017). The highest NOS peak in summer is monoterpene-derived $C_{10}H_{16}NO_7S^-$, the same as in winter.

3.3 Quantified organosulfates

In this study, we quantified twenty-nine OS and six NOS compounds using a variety of authentic and surrogate OS standards (Table 1). The quantified OS and NOS accounted for 14-18% and 47-67% by intensity of the identified CHOS and CHONS in polluted winter days and 15-37% and 58-87% in polluted summer days (Fig. 3c), respectively. Increased contribution of the quantified OS and NOS in summer is because they are mainly derived from biogenic VOCs, which have greater emissions in summer than in other seasons (Guenther et al., 1995). We note that a large fraction of OS signals was not quantified owing to the lack of proper standards in this study. As discussed above, these unquantified OS mainly originated from anthropogenic sources. Future studies of their abundances and formation mechanisms are warranted.

Table 2 summarizes the seasonally and annually averaged concentrations of the quantified OS, as well as their contributions to OM in 2015/2016 and 2018/2019. The average concentration of quantified OS was 65.5±77.5 ng m⁻³ in 2015/2016 and 59.4±79.7 ng m⁻³ in 2018/2019. Although there was little change in OS concentration in these two years, the contribution of OS to OM was larger in 2018/2019 (0.66%±0.56%) than in 2015/2016 (0.57%±0.56%), mainly due to a significant reduction of OM in 2018/2019. Since OS species are important tracers for SOA (Surratt et al., 2007b; Gómez-González et al., 2008; Surratt et al., 2008; McNeill et al., 2012; Zhang et al., 2012; Surratt et al., 2010; Lin et al., 2013), an increase of OS/OM ratios in 2018/2019 implies an enhanced contribution of SOA to OA in Shanghai. A previous study by Ma et al. (2014) reported an average OS concentration in urban Shanghai in 2012/2013 of about 8.6 ng m⁻³, substantially smaller than the concentration reported here. This is likely due to a different number of OS species quantified (17 vs. 35) and different OS standards used (octyl and benzyl sulfates vs. seven authentic/surrogate standards) in Ma et al. (2014) and the present study. As can be seen in Fig. 2e and Table 2, the OS concentration and OS/OM ratio both showed a strong seasonal variation and peaked in summer. The concentration of OS and its contribution to OM in summertime Shanghai (on average, 114.1 ng m⁻³ and 1.13% in July

2015 and 102.1 ng m⁻³ and 1.18% in July 2019) were larger or comparable to those observed in Beijing (55.2 ng m⁻³, 0.42%) (Wang et al., 2018) and Birmingham, Alabama (205.4 ng m⁻³, 2% of OC) (Rattanavaraha et al., 2017), but significantly lower than those observed in Atlanta, GA (2366.4 ng m⁻³, 16.5% of OC) and Centreville, AL (812 ng m⁻³, 7.3% of OC) (Hettiyadura et al., 2019) where the production of OS and SOA is dominated by the oxidation of biogenic emissions. The contribution of OS to OM in wintertime Shanghai (on average, 0.32% in January 2016 and 0.36% in January 2019) was larger than that observed in Xi'an (~0.2%) (Huang et al., 2018), though the quantified OS concentrations in the two regions were comparable. This may suggest a stronger secondary formation of OA in Shanghai than in Xi'an, consistent with independent measurements by Huang et al. (2014).

To further characterize the seasonality and interannual variability of OS, as well as their origin and formation mechanisms, the quantified OS were assigned to four different source categories based on their molecular composition and literature data (Surratt et al., 2008; Surratt et al., 2007a; Nozière et al., 2010; Surratt et al., 2010; Schindelka et al., 2013; Zhang et al., 2014; Riva et al., 2015; Riva et al., 2016b; Blair et al., 2017; Nestorowicz et al., 2018). The OS species for each OS source category are listed in Table 1 and the seasonal and interannual variations in the abundance of grouped and individual OS are shown in Fig. 4 and Table S5, respectively.

3.3.1 Isoprene-derived organosulfates

The isoprene-derived OS (hereafter referred to as OS_i) include ten $C_{4.5}$ species and one dimeric species ($C_7H_9O_7S^-$). The average concentration of OS_i in summer was 76.5 ± 93.4 ng m⁻³ for 2015/2016 and 68.4 ± 102.2 ng m⁻³ for 2018/2019, significantly larger than the concentrations (10.4-17.1 ng m⁻³) in other seasons (Fig. 4a). Similar strong seasonality of OS_i was also observed in suburban areas in the Mid-Atlantic United States (Meade et al., 2016) and the Pearl River Delta in Southern China (He et al., 2014) The significantly increased production of OS_i in summer is mainly a result of enhanced isoprene emissions (Guenther et al., 1995) and photochemistry due to strong solar radiation and high temperatures in this warmer season.

The most abundant species among OS_i was 2-MT-OS ($C_5H_{11}O_7S^-$), produced by reactive uptake of IEPOX on sulfate during the photooxidation of isoprene under low-NO_x conditions (Surratt et al., 2010). The average concentration of 2-MT-OS was ~31 ng m⁻³ in summer, contributing to about 45% of OS_i , whereas it decreased to 0.4-1.3 ng m⁻³ in other seasons, accounting for only 4-10% of OS_i in 2015/2016 and 2018/2019 (see Table S5). In addition, 2-MA-OS ($C_4H_7O_7S^-$) was also abundantly detected with an average concentration of 4.5 ng m⁻³ in summer and 1.0-2.2 ng m⁻³ in other seasons during the two sampling years. 2-MA-OS is formed from reactive uptake of methacrylic acid epoxide (MAE) (Lin et al., 2013) and hydroxymethylmethyl-lactone (HMML) (Nguyen et al., 2015) on sulfate aerosol during isoprene photooxidation under high-NO_x conditions. It is worthwhile noting that the concentration ratio of 2-MT-OS/2-MA-OS in summer (6.8-7.8) is substantially larger than that in other seasons (0.31-0.78). This is consistent with a dramatic reduction of NO_x level (e.g., NO_2) in summer (~13 ppb) compared to that in other seasons (~24-34 ppb) (see Table S1). Furthermore, the NO_x -influenced oxidation pathways may not be conducive to 2-MA-OS formation in summer, given that the formation of reactive intermediates such as MAE is unfavorable at high temperatures owing to enhanced thermal decomposition of its precursor methacryloylperoxynitrate (MPAN) (Worton et al., 2013), 2-MT-OS and 2-MA-OS are key tracers for isoprene-derived SOA under low- and high-NO_x

conditions, respectively (Surratt et al., 2010; Lin et al., 2013; Nguyen et al., 2015). The dramatically larger ratios of 2-MT-OS/2-MA-OS in summer than in other seasons therefore strongly suggest that the low-NO $_x$ oxidation pathways dominated the production of isoprene-derived SOA in summer, while the processes favorable under high-NO $_x$ conditions were important for SOA formation in other seasons. We note that the 2-MT-OS/2-MA-OS ratios observed in summertime Shanghai are smaller than those (17.0-33.8) observed in less polluted environments such as the southeastern United States (Hettiyadura et al., 2019; Budisulistiorini et al., 2015; Riva et al., 2019), but significantly larger than those (0.55-1.57) observed in Beijing (Wang et al., 2018; Bryant et al., 2020) and the Pearl River Delta (PRD) region of China (He et al., 2018).

Other abundant OS_i species include $C_5H_7O_7S^-$, $C_5H_9O_7S^-$, and $C_4H_7O_6S^-$. The $C_5H_7O_7S^-$ and $C_5H_9O_7S^-$ species can be produced by photooxidation of isoprene (Surratt et al., 2008; Nestorowicz et al., 2018) and/or the oxidative aging of 2-MT-OS (Hettiyadura et al., 2015; Chen et al., 2020). The $C_4H_7O_6S^-$ can be generated both from sulfate radical reaction with MACR/MVK (Nozière et al., 2010; Schindelka et al., 2013; Wach et al., 2019) and isoprene photooxidation (Lin et al., 2013; Surratt et al., 2007a; Nestorowicz et al., 2018). The $C_5H_7O_7S^-$ and $C_4H_7O_6S^-$ are also consistent in molecular formula with the OS species formed from the photooxidation of diesel fuel vapors (Blair et al., 2017). However, these two species had moderate to strong correlations with MT-OS and $C_5H_9O_7S^-$ in different seasons except for autumn ($C_5H_7O_7S^-$: r=0.68-0.96, $C_4H_7O_6S^-$: r=0.62-0.96), indicating that they are mainly derived from isoprene oxidation. We note that the five most abundant OS_i species as discussed above were moderately correlated with EC and CO in winter (r=0.5-0.67), suggesting that there might be anthropogenic sources of isoprene in winter. Borbon et al. (2001) measured the hourly isoprene concentration at an urban site in Lille, France for two years and found that isoprene was largely derived from vehicle emissions in winter. In addition to OS species, two isoprene-derived NOS ($C_5H_{10}NO_9S^-$ and $C_5H_8NO_{10}S^-$) were also observed, in particular in summer.

3.3.2 Monoterpene-derived organosulfates

The monoterpene-derived OS (hereafter referred to as OS_m) include seven $C_{7\text{-}10}$ OS species and three $C_{9\text{-}10}$ NOS species. Compared to the OS_n , the OS_m showed a weaker seasonal variation and a relatively larger abundance except in summer (Fig. 4b). This is consistent with the fact that isoprene emissions have stronger seasonal variability than monoterpene emissions (Guenther et al., 1995). The seasonally averaged concentrations of OS_m were higher in spring and summer, but lower in autumn and winter. This is different from previous observations in 2012/2013 in Shanghai by Ma et al (2014). They found that the OS_m were most abundant in summer, followed by autumn, winter, and spring. The differences in seasonal variations of OS_m observed by the two studies may be attributed to different meteorological and chemical conditions that affected precursor emissions and chemistry of OS_m formation over the sampling periods. Given that the OS_m concentration had an obvious daily variation, the number of samples collected would significantly affect the seasonally averaged concentration. As such, the difference in the number of samples collected each season (18-20 samples in this study versus 6 samples within three days in Ma et al. (2014)) may also contribute to the different seasonality observed in the two studies.

The NOS species such as $C_{10}H_{16}NO_7S^-$, $C_9H_{14}NO_8S^-$, and $C_{10}H_{16}NO_{10}S^-$ were the most abundant OS_m species, which arises mainly from monoterpenes photooxidation in the presence of NO_x or nighttime NO_3 chemistry (Surratt et al., 2008; Iinuma et al., 2007a). The concentrations of these three NOS were all lower in summer than in spring and autumn (Table S5), consistent with the seasonal trend of NO_x concentrations (Fig. 2 and Table S1). Similar seasonal variations for these NOS

species were also observed in the PRD region of China (He et al., 2014) and the Mid-Atlantic United States (Meade et al., 2016). Among the NOS species, the $C_{10}H_{16}NO_7S^-$ was most abundant, contributing to 22-48% of OS_m . This species had an annual average concentration of 6.2 ± 6.5 ng m⁻³ in 2015/2016 and 5.5 ± 6.2 ng m⁻³ in 2018/2019, which is comparable to the concentrations observed in Beijing (12 ng m⁻³) (Wang et al., 2018) and Atlanta, GA (9.0 ng m⁻³) (Hettiyadura et al., 2019), but much lower than that observed in the PRD region of China (52.4 ng m⁻³ in summer and 151 ng m⁻³ in autumn) (He et al., 2014). The prevalence of monoterpene-derived NOS in Shanghai as observed in this study is consistent with recent observations that monoterpenes accounted for up to 60% of nighttime NO_3 radical loss in the YRD region of China (Wang et al., 2020a).

The most abundant nitrogen-free OS_m species was $C_{10}H_{15}O_7S^-$ (m/z 279.0538), which was shown to be produced from the photooxidation of monoterpenes (Surratt et al., 2008) or sulfate radical reaction with α -pinene (Nozière et al., 2010). Although $C_{10}H_{15}O_7S^-$ is consistent in molecular composition with the OS species formed by photooxidation of cyclodecane in the presence of sulfate aerosol (Riva et al., 2016b), its moderate to strong correlation (r=0.51-0.93) with the three monoterpenederived NOS in all seasons except for winter suggests that it is mainly derived from monoterpene oxidation. The concentration of $C_{10}H_{15}O_7S^-$ was on average 3.0 ± 3.0 ng m⁻³ in 2015/2016, lower than that $(4.0 \pm 3.4$ ng m⁻³) in 2018/2019. In contrast to the NOS species, the $C_{10}H_{15}O_7S^-$ species was most abundant in summer in both years, again suggesting a strong contribution of low-NO_x chemistry in OS and SOA formation in summer.

3.3.3 Anthropogenic organosulfates

The quantified anthropogenic OS (hereafter referred to as OS_a) in this study include phenyl sulfate (C₆H₅O₄S⁻), benzyl sulfate $(C_7H_7O_4S^-)$, $C_8H_{17}O_4S^-$, as well as $C_4H_7O_4S^-$, $C_5H_7O_6S^-$, and $C_6H_9O_6S^-$. The annual average concentrations of OS_a in 2015/2016 and 2018/2019 were 5.6±2.8 ng m⁻³ and 3.8±3.3 ng m⁻³, respectively. Although the concentration of quantified OS_a decreased in 2018/2019, we are not sure whether the total concentration of OS_a decreased because only a small fraction of OS_a was quantified. The inter-annual variation of OS_a warrants further studies. As shown in Fig. 4c, the quantified OS_a concentration was substantially smaller compared to the biogenic OS one. In addition, the OS_a concentration had little seasonal variations in both 2015/2016 and 2018/2019. Among the quantified OS_a, C₄H₇O₄S⁻ was most abundant with an annual average concentration of 2.0±1.5 ng m⁻³ in 2015/2016 and 1.8±2.6 ng m⁻³ in 2018/2019, which is comparable to the concentrations in Atlanta, GA (Hettiyadura et al., 2019). Blair et al. (2017) found that photooxidation of diesel vapors in the presence of SO₂ can form $C_4H_7O_4S^-$, $C_5H_7O_6S^-$, and $C_6H_9O_6S^-$ species. The $C_8H_{17}O_4S^-$ species had the same retention time as the octyl sulfate standard in the LC column, suggesting it is a long-chain aliphatic OS. This OS species was correlated with C₅H₇O₆S⁻ and C₆H₉O₆S⁻ that were potential diesel vapor-derived OS. Phenyl sulfate and benzyl sulfate may be produced by photooxidation of naphthalene and 2-methylnaphthalene (Riva et al., 2015) and/or sulfate radical reaction with aromatic compounds such as benzoic acid and toluene in the aqueous phase (Huang et al., 2020), but phenyl sulfate was only detected in 42 out of 75 samples in 2015/2016 and 8 out of 81 samples in 2018/2019, primarily in winter. The benzyl sulfate concentrations in 2015/2016 and 2018/2019 were 0.4±0.1 ng m⁻³ and 0.2±0.10 ng m⁻³, respectively, which were higher than the observations in springtime Lahore, Pakistan (Staudt et al., 2014) and in wintertime Xi'an, China (Huang et al., 2018). Benzyl sulfate had a strong correlation with phenyl sulfate and was also correlated with C₆H₉O₆S⁻ and C₈H₁₇O₄S⁻.

3.3.4 C₂/C₃ organosulfates

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The OS species with two or three carbon atoms are grouped together since many of them are considered to have both biogenic and anthropogenic origins. The C_2/C_3 OS quantified in this study include $C_2H_3O_6S^-$ (m/z 154.9650), $C_3H_5O_5S^-$ (m/z 152.9858), $C_3H_5O_6S^-$ (m/z 168.9807), $C_2H_3O_5S^-$ (m/z 138.9701), $C_3H_5O_4S^-$ (m/z 136.9909), and $C_3H_7O_5S^-$ (m/z 155.0014). The C_2/C_3 OS species accounted, on average, for 19% of the quantified OS concentrations and they were overall more abundant in summer than in other seasons in 2015/2016 and 2018/2019 (Fig. 4d). The C₂H₃O₆S⁻, C₃H₅O₅S⁻, and C₃H₅O₆S⁻ species, which were previously assigned to glycolic acid sulfate (GAS), hydroxyacetone sulfate (HAS), and lactic acid sulfate (LAS), respectively (Olson et al., 2011; Hettiyadura et al., 2017; Huang et al., 2018; Wang et al., 2018; Hettiyadura et al., 2019), were among the most abundant C_2/C_3 OS species, together contributing to 76% of C_2/C_3 OS concentrations during the two sampling years. The concentration of C₂H₃O₆S⁻ (GAS) was on average 2.9±2.2 ng m⁻³ in 2015/2016 and 2.3±1.7 ng m⁻³ in 2018/2019, which was lower than the concentrations measured in Beijing (19.5 ng m⁻³) (Wang et al., 2018), Xi'an (77.3 ng m⁻³) (Huang et al., 2018), Atlanta, GA (58.5 ng m⁻³) (Hettiyadura et al., 2019), Centreville, AL (20.6 ng m⁻³) (Hettiyadura et al., 2017), Lahore, Pakistan (11.3 ng m⁻³), and Bakersfield, CA (4.5-5.4 ng m⁻³) (Olson et al., 2011), and similar with that observed in Riverside, CA (3.3 ng m⁻³) (Olson et al., 2011). We note that if accounting for the underestimation (2-6 times) in concentration due to matrix effects (see Sect. 2.5), the GAS concentration measured in Shanghai would be comparable to that in most of the regions mentioned above. The concentrations of C₃H₅O₅S⁻ (HAS) and C₃H₅O₆S⁻ (LAS) were quite similar, on average 2.3 and 2.2 ng m⁻³ in 2015/2016 and 1.8 and 1.9 ng m⁻³ in 2018/2019, respectively, which were comparable to the concentrations observed in Beijing (2.2 ng m⁻³ and 4.4 ng m⁻³) (Wang et al., 2018), and Xi'an (1.3 ng m⁻³ for HAS) (Huang et al., 2018), but lower than those measured in Centreville, AL (5.8 ng m⁻³ and 16.5 ng m⁻³) (Hettiyadura et al., 2017) and Atlanta, GA (10.1 ng m⁻³ and 38.4 ng m⁻³) (Hettiyadura et al., 2019), C₂H₃O₆S⁻, C₃H₅O₆S⁻, C₃H₅O₆S⁻, and C₂H₃O₅S⁻ were strongly correlated with most of the OS_i species (r=0.52-0.96 for $C_2H_3O_6S^-$, r=0.53-0.99 for $C_3H_5O_5S^-$, r=0.53-0.90 for $C_3H_5O_6S^-$, and r=0.53-0.94 for C₂H₃O₅S⁻), indicating that they originated mainly from isoprene chemistry. This is in line with recent findings that a series of C₂/C₃ OS species, including C₂H₃O₆S⁻, C₃H₅O₆S⁻, and C₂H₃O₅S⁻, can be produced by heterogeneous OH oxidation of particulate 2-MT-OS (Chen et al., 2020). The C₃H₅O₄S⁻ species, proposed to contain an allyl group (Hettiyadura et al., 2017), was previously found to be produced by diesel photooxidation (Blair et al., 2017), and was correlated with anthropogenic OS such as the potential diesel vapor-derived OS (C₈H₁₇O₄S⁻, C₄H₇O₄S⁻, C₅H₇O₆S⁻, and C₆H₉O₆S⁻, r=0.53-0.87) and benzyl sulfate (C₇H₇O₄S⁻, r=0.49-0.88). C₃H₇O₅S⁻ is likely an OS species containing one hydroxyl group (Hettiyadura et al., 2017); it was strongly correlated with C₃H₅O₄S⁻ in all seasons and correlated with the diesel vapor-derived OS_a (C₆H₉O₆S⁻ and C₈H₁₇O₄S⁻) in spring and autumn, suggesting that it may be largely derived from the photooxidation of diesel vapors. This result is different from the observations in Atlanta, GA, where C₃H₇O₅S⁻ was correlated with most of the OS_i, leading to the suggestion that it was derived from the oxidation of isoprene (Hettiyadura et al., 2019). We note that the concentrations of the C₃H₅O₄S⁻ and C₃H₇O₅S⁻ species decreased significantly from 2015/2016 to 2018/2019 (except for summer, see Table S5), overall consistent with the interannual variations of OS_a species. This further supports that these two OS species mainly originated from anthropogenic sources.

3.4 Factors influencing organosulfate formation

Laboratory and field studies have shown that aerosol properties such as acidity, sulfate concentration, and ALWC play important roles in the formation of OS (Iinuma et al., 2007b; Chan et al., 2011; Surratt et al., 2007a; Surratt et al., 2007b; Liao et al., 2015; Hettiyadura et al., 2019; Riva et al., 2019). Here we examined the influences of these factors, as well as the level of oxidants and temperature on OS formation in ambient aerosols in Shanghai. Aerosol pH and ALWC here were calculated using ISORROPIA-II (see Sect. 2.4). Figure 5 shows the OS concentration versus the O_x level, sulfate concentration, aerosol pH, and ALWC observed in the spring, autumn, and winter of 2015-2016 and 2018-2019. Since the OS concentrations in summer were significantly greater than in other seasons, they were plotted separately in Fig. 6. As shown in Figs. 5 and 6, the aerosol pH in Shanghai ranged between 1.5 and 5.3 in summer and between 2.5 and 6.1 in other seasons, overall within the pH range (2-6) reported for ambient aerosols in northern China (Liu et al., 2017; Shi et al., 2017; Ding et al., 2019b; Song et al., 2019; Wang et al., 2018). A recent study by Zheng et al. (2020) has suggested that aerosol pH levels in populated continental regions including eastern and northern China are widely buffered by ammonium/ammonia, where the variation in aerosol pH is mainly driven by the variation in ALWC and temperature. Therefore, we infer that the lower aerosol pH in summer than in other seasons in Shanghai was mainly a result of decreased ALWC (Figs. 5 and 6c, d) and enhanced temperature (Fig. 2a). Decreased aerosol pH in summer compared to other seasons was also observed in Beijing (Ding et al., 2019b) and the southeastern United States (Guo et al., 2015; Nah et al., 2018).

As can be seen in Fig. 5, the OS concentration in spring, autumn, and winter increased obviously with increasing O_x level, sulfate concentration, and aerosol acidity (Fig. 5a, b). A similar result was also found in Beijing where most OS species were correlated strongly with the product of ozone and particulate sulfate ($[O_3] \cdot [SO_4^{2-}]$) (Bryant et al., 2020). In addition, an overall positive correlation was observed between the OS concentration and ALWC (Fig. 5c, d). Therefore, it is likely that the OS species were mainly produced by acid-catalyzed heterogeneous/aqueous-phase reactions of oxidized organic compounds with sulfate in these seasons. Previous studies have shown that elevated ALWC could inhibit OS production by decreasing aerosol acidity through dilution (Lewandowski et al., 2015; Nestorowicz et al., 2018). However, as the increase of ALWC was accompanied by elevated sulfate concentration, such a decrease in aerosol acidity was not observed in the present study (Fig. 5c, d). Alternatively, the increased ALWC likely promoted the mass transfer of oxidized organics into the aerosol phase, thereby enhancing OS formation. We note that the observations with moderate to high ALWC but relatively low OS concentration (data points in the triangle in Fig. 5c, d) were associated with low O_x levels (<50 ppb) that significantly limited the oxidation of VOC precursors and hence the formation of OS.

As seen in Fig. 6, the OS production in summer increased dramatically with rising O_x concentration. In addition, high OS concentrations were associated with high ambient temperatures, which can enhance emissions of biogenic precursors and the production of O_x . While the aerosol acidity effect on OS production in summer was still evident, the influence of sulfate and ALWC was not as obvious as in other seasons. This is likely because the OS production in summer was driven by the strong emission and fast photochemistry of VOC precursors. It is noteworthy that the sulfate concentrations, ALWC, and aerosol acidities were overall higher in 2015/2016 than in 2018/2019, but the OS concentrations were similar in two years. This implies that the O_x level is a driving factor for OS formation in ambient aerosols in Shanghai. Very recently, a similar oxidant effect on OS formation was also observed in urban Beijing (Bryant et al., 2020). Therefore, mitigation of O_x pollution may effectively reduce the production of OS and SOA in Chinese megacities.

4 Conclusions

 In this study, we collected ambient PM_{2.5} filter samples over four seasons in 2015/2016 and 2018/2019 in urban Shanghai, China, and comprehensively characterized the sulfur-containing organic compounds (CHOS and CHONS) in these PM_{2.5} samples using UPLC-ESI(-)-QToF-MS. The CHOS and CHONS species accounted for a large fraction of the ion signals for organic compounds in ambient PM_{2.5}. Using a set of authentic and surrogate OS standards, we quantified the abundance of 29 OS and 6 NOS species in ambient aerosols. We find that there was no strong change in the OS concentration in 2018/2019 (59.4±79.7 ng m⁻³) compared to that in 2015/2016 (65.5±77.5 ng m⁻³), though the OM concentration decreased by 29% between 2015/2016 (12.7±8 μg m⁻³) and 2018/2019 (9±5.5 μg m⁻³). As a result, the annual average contribution of quantified OS to OM increased from 0.57% in 2015/2016 to 0.66% in 2018/2019, suggesting an enhanced contribution of SOA to OM in Shanghai in recent years. The OS_i and OS_m accounted, on average, for 36.3% and 31.0% of the quantified OS concentrations, respectively, during the two sampling years, indicating a significant contribution of biogenic emissions to SOA in Shanghai. The abundance of OS_i had strong seasonality and was significantly higher in summer (76.5±93.4 ng m⁻³ in 2015/2016 and 68.4 ± 102.2 ng m⁻³ in 2018/2019) than in other seasons (10.4-17.1 ng m⁻³). The OS_m concentration showed a weaker seasonal variation and was relatively higher in spring and summer than in autumn and winter. In contrast, the quantified OS_a had little seasonal variations and decreased by 32% from 2015/2016 to 2018/2019. The C₂/C₃ OS species that were more abundant in summer than in other seasons, contributed, on average, to 19% of the concentration of the quantified OS species. The C₂H₃O₆S⁻ (GAS), C₃H₅O₅S⁻ (HAS), and C₃H₅O₆S⁻ (LAS), which were derived mainly from isoprene chemistry, were the most abundant C_2/C_3 OS species and together accounted for 76% of the C_2/C_3 OS concentrations.

2-MT-OS was the most abundant OS species in summer. The dramatic larger 2-MT-OS/2-MA-OS ratios in summer (6.8-7.8) versus other seasons (0.31-0.78) imply that the reaction pathways prevalent under low-NO_x conditions (e.g., reactive uptake of IEPOX and photooxidation of ISOPOOH) dominated the production of OS_i and isoprene-derived SOA in summer, while the processes favorable under high-NO_x conditions play an important role in OS_i and SOA formation in other seasons. The $C_{10}H_{16}NO_7S^-$ species derived from monoterpenes was the most abundant NOS species, with an annual average concentration of 6.2±6.5 ng m⁻³ in 2015/2016 and 5.5±6.2 ng m⁻³ in 2018/2019. This agrees well with previous observations that monoterpenes depleted about 60% of nighttime NO₃ radicals in the YRD region of China (Wang et al., 2020a).

In addition, we find that the abundance of OS was overall positively correlated with the O_x level, sulfate concentration, aerosol acidity, as well as ALWC in spring, autumn, and winter, suggesting the production of OS via acid-catalyzed aqueous-phase reactions of oxidized organic compounds on sulfate. However, OS production in summer was strongly driven by rising O_x and temperature that could enhance the photochemistry and emissions of biogenic precursors. We further find that although sulfate concentrations, aerosol acidities, and ALWC were significantly lower in 2018/2019 than in 2015/2016, the production of OS was largely sustained in 2018/2019 by the nearly unchanged O_x level that maintained the fast oxidation of VOC precursors. These results imply that controlling O_x pollution may also effectively mitigate particulate organic matter pollution in eastern China.

It should be pointed out that GAS was likely underestimated by a factor of 2-6 as a result of the matrix effect during the analysis in our study. If accounting for this effect, it would be the second most abundant OS species after MT-OS. In addition,

- 497 a large fraction of the CHOS signals that arose mainly from anthropogenic sources was not quantified due to the lack of proper
- 498 OS standards in this study. Therefore, the OS concentration and its contribution to organic aerosol in Shanghai could be
- significantly greater. Future studies on the abundance, origin, and formation mechanisms of these unquantified OS are
- warranted for a better understanding of the formation and evolution of OS and SOA in the atmosphere.

- 502 Data availability. The data presented in this work are available upon request from the corresponding author
- 503 (yuezhao20@sjtu.edu.cn).
- 504 Author contributions. YZ designed and led the research, YW, WFZ, and ZC collected ambient samples, YCW and JZY
- 505 provided OS standards, JYS conducted ISORROPIA-II model calculation, YW conducted sample analysis, and YZ and YW
 - processed the data and wrote the paper with contributions from all of the authors.
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G. A	m/z,	Formula,	.	Standards for		F . 6	Average concentration	
Category	[M-H] ⁻	[M-H] ⁻	Retention time	quantification	Proposed structure	Ref.	2015/2016	2018/2019
	167.0014	C ₄ H ₇ O ₅ S ⁻	0.85,1.36	Lactic acid sulfate (LAS)	osō ₃	a	1.13	1.45
Isoprene OS	182.9963	$\mathrm{C_4H_7O_6S^-}$	0.78,2.23	LAS	HO OSO₃	b	2.84	2.19
	197.0120	$C_5H_9O_6S^-$	0.85,1.55,3.11,3.99	LAS	$HO \longrightarrow OSO_3$	c	1.87	1.72
	198.9912	$C_4H_7O_7S^-$	0.74,3.86	LAS	HO OSO 3	d	2.28	2.50
	199.0276	$C_5H_{11}O_6S^{-}$	0.85,8.08	LAS	HO → OSO₃	c	0.57	0.65
	210.9912	C ₅ H ₇ O ₇ S ⁻	0.77	LAS	On oso3	d	6.09	4.81
	213.0069	$C_5H_9O_7S^-$	0.84	LAS	OH OSO ₃	d	3.81	3.82
	215.0226	$C_5H_{11}O_7S^-$	0.72	LAS	HO OSO3 OH	e	11.35	8.92
	237.0069	$C_7H_9O_7S^-$	0.87,1.5	LAS		f	0.58	0.50
	260.0076	$C_5H_{10}NO_9S^-$	1.19	LAS	HO OSO_3 ONO_2	g	2.28	2.96
	273.9869	C ₅ H ₈ NO ₁₀ S ⁻	1.3	LAS	OSO_3 $OOOO_2$ $OOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOO$	h	1.89	4.87
Monoterpene OS	223.0276	$C_7H_{11}O_6S^-$	2.52,2.87,3.19,3.42	Glycolic acid sulfate (GAS)	osō ₃	i	1.37	1.12
	239.0225	$C_7H_{11}O_7S^-$	0.85,1.77,2.03,3.08	GAS	OH O	f	1.91	2.01
	249.0797	$C_{10}H_{17}O_5S^-$	5.34,5.60,6.85,7.04, 7.12,7.36	α-Pinene sulfate	HO OSO3	j	0.33	0.17
	251.0589	$C_9H_{15}O_6S^-$	3.63,4.14,4.82,7.35	Limonaketone sulfate	HO oso₃	j	1.32	1.02
	253.0382	$C_8H_{13}O_7S^-$	3.35,3.63,4.45	GAS	OHO	a	1.39	1.73
	279.0538	$C_{10}H_{15}O_7S^-$	3.42,4.34,4.55,4.71	GAS		g	2.99	4.00
	281.0695	$C_{10}H_{17}O_{7}S^{-}$	4.93,5.14	α-Pinene sulfate	HO OSO ₃ OH	f	0.34	0.21
	294.0648	$C_{10}H_{16}NO_7S^{\scriptscriptstyle -}$	8.14,9.02,9.16,9.35	α-Pinene sulfate	O_2NO	k	6.21	5.55

					√osō₃			
	296.0440	C ₉ H ₁₄ NO ₈ S ⁻	4.72,4.86,5.24,6.44	Limonaketone sulfate	ONO ₂	k	1.62	2.29
	342.0495	$C_{10}H_{16}NO_{10}S^{-}$	4.57,4.71,5.17,7.35	Limonaketone sulfate	O SO3 OH HO ONO2	i	1.32	1.59
	151.0065	C ₄ H ₇ O ₄ S ⁻	0.92,2.87,4.11,5.23, 5.39,8.21	Methyl sulfate	unknown	-	2.04	1.80
	194.9963	$C_5H_7O_6S^-$	0.82,2.86	GAS	unknown	_	0.83	0.76
	209.0120	$C_6H_9O_6S^-$	0.87,2.09,3.27,4.06	GAS	unknown	_	1.44	0.63
Anthropogenic OS	209.0845	$C_8H_{17}O_4S^-$	9.64	Sodium octyl sulfate	$\sim\sim\sim$	-	1.04	0.86
	172.9909	$C_6H_5O_4S^-$	2.6	Phenyl sulfate	$os\bar{o}_3$	1	0.36	0.12
	187.0065	C ₇ H ₇ O ₄ S ⁻	4.54	Phenyl sulfate	$\cos \bar{o}_3$	1	0.35	0.17
	136.9909	C ₃ H ₅ O ₄ S ⁻	0.78,3.93	GAS	setone tite ONO_2 setone ONO_2 sulfate ONO_2 sulfate ONO_2 sulfate ONO_2 sulfate ONO_2 sulfate ONO_2 sulfate OSO_3 sulfate $OSOO_3$ sulfate $OSOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOO$	_	0.62	0.50
	138.9701	$C_2H_3O_5S^-$	0.69	GAS	o Soso₃	g	0.58	0.57
C ₂ /C ₃ OS	152.9858	$C_3H_5O_5S^-$	0.81	GAS	osō₃	d	2.30	1.79
	154.9650	$C_2H_3O_6S^-$	0.69	GAS	Ho Losō₃	m	2.91	2.25
	155.0014	$C_3H_7O_5S^-$	0.79	GAS	OH OSO3	n	1.21	0.70
	168.9807	$C_3H_5O_6S^-$	0.82	LAS	HO COSO₃	m	2.24	1.94
Unknown	164.9858	C ₄ H ₅ O ₅ S ⁻	0.79	Methyl sulfate	©osō₃	n	1.20	0.78
source OS	241.9971	C ₅ H ₈ NO ₈ S ⁻	3.39,3.57,3.93,4.21	Methyl sulfate	$O_2NO \longrightarrow OSO_3$	n	1.67	1.21
SUM					<u> </u>		65.48	59.04

*Retention times of each OS isomer considered in the quantification. References for proposed OS structures: ^a Schindelka et al. (2013)), ^b Shalamzari et al. (2013), ^c Riva et al. (2016a), ^d Hettiyadura et al. (2015), ^e Surratt et al. (2010), ^f Nozière et al. (2010), ^g (Surratt et al., 2007a), ^h Nestorowicz et al. (2018), ⁱ Yassine et al. (2012), ^j Wang et al. (2017), ^k Surratt et al. (2008), ^l Huang et al. (2018), ^m Olson et al. (2011), ⁿ Hettiyadura et al. (2019).

Table 2. A summary of OS concentration (in ng m⁻³) and its contribution to OM (OS/OM) in four seasons in 2015/2016 and 2018/2019.

Concer	2015/	2016	2018/2019			
Season —	OS	OS/OM	OS	OS/OM		
All year	65.5±77.5	$0.57\% \pm 0.56\%$	59.4±79.7	0.66%±0.56%		
Spring	51.1±24.4	$0.34\% \pm 0.10\%$	51.5±28.8	$0.48\% \pm 0.15\%$		
Summer	114.1±128.4	$1.13\% \pm 0.78\%$	102.1±137.7	$1.18\% \pm 0.81\%$		
Autumn	38.2±21.7	$0.36\% \pm 0.11\%$	38.0 ± 20.0	$0.54\% \pm 0.24\%$		
Winter	44.5±17.5	$0.32\% \pm 0.12\%$	37.3±18.4	0.36%±0.13%		

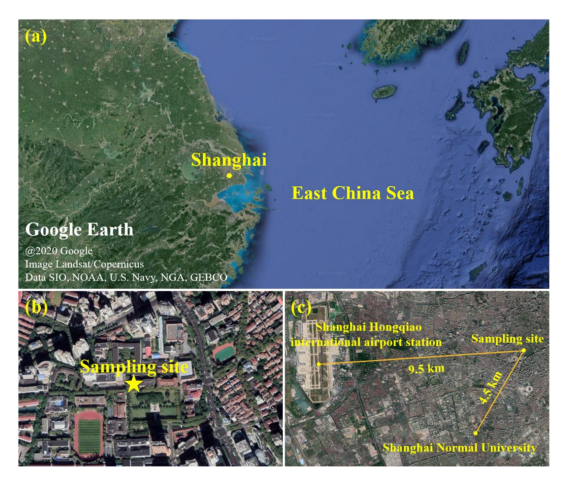


Figure 1. (a) Map of Shanghai. **(b)** Map of the sampling site on the Xuhui Campus of Shanghai Jiao Tong University in downtown at 31.201°N, 121.429°E. **(c)** Map of the PM_{2.5} sampling site, the meteorological station at Shanghai Hongqiao international airport, the air quality monitoring station at Shanghai Normal University, and distances between them.

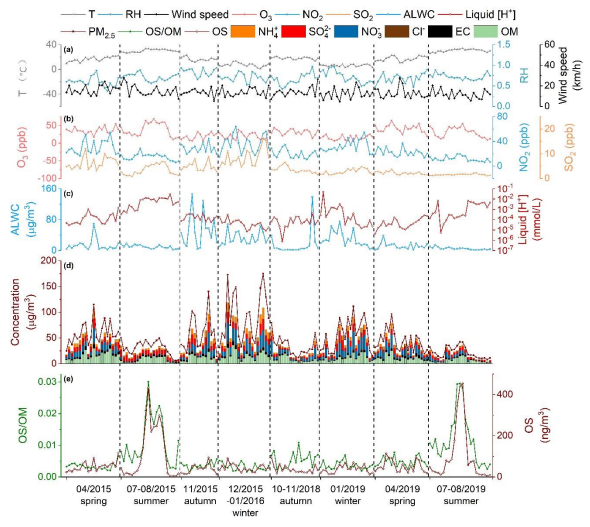


Figure 2. Time series of temperature, relative humidity (RH), wind speed, O₃, NO₂, SO₂, aerosol liquid water content (ALWC) and liquid [H⁺], concentrations of particulate organic matter (OM), elemental carbon (EC), sulfate, nitrate, chloride, ammonium, as well as the abundance of OS and its contribution to OM in 2015/2016 and 2018/2019 in Shanghai.

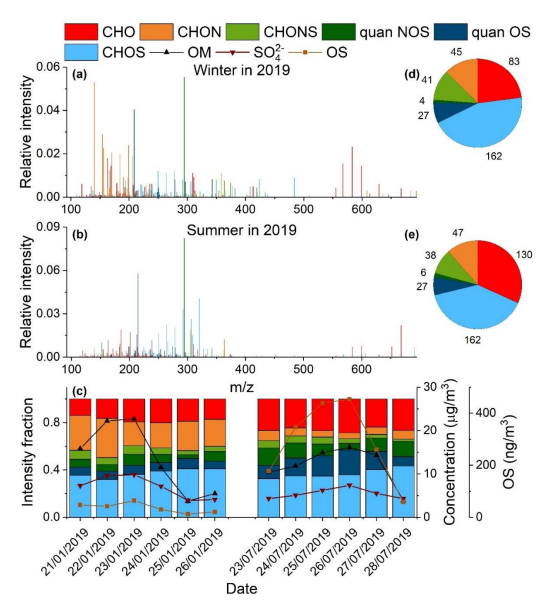


Figure 3. Average mass spectra of detected compounds in ambient aerosols during typical (**a**) wintertime (21-26 January 2019) and (**b**) summertime (23-28 July 2019) pollution episodes in Shanghai. The detected compounds were classified into six categories, i.e., CHO, CHON, CHOS, CHONS, quantified NOS, and quantified OS. The CHOS and CHONS categories excluded quantified OS and NOS, respectively. (**c**) Intensity fraction of different compound categories, as well as the time series of OM, SO₄²⁻, and OS concentrations during two pollution episodes in 2019. (**d**) (**e**) Number of compounds detected in each category during the pollution episodes in winter and summer, respectively.

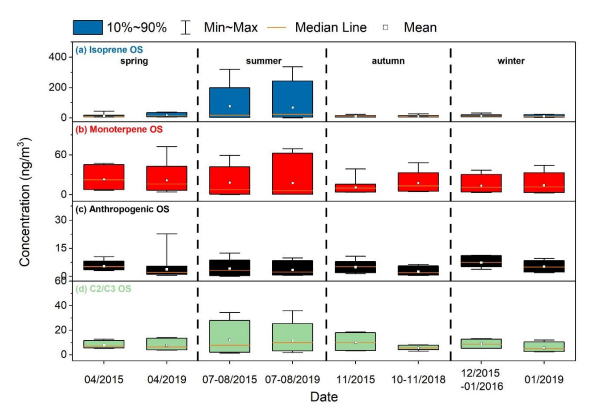


Figure 4. Concentrations of different types of the quantified OS over the four seasons in 2015/2016 and 2018/2019.

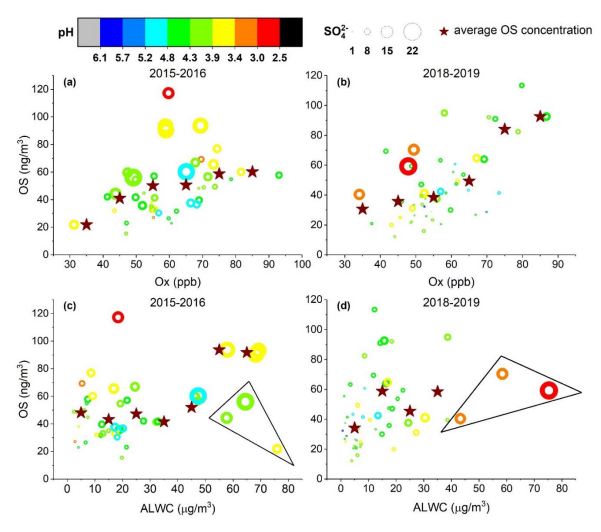


Figure 5. Quantified OS concentrations as a function of (a) (b) the level of O_x ($O_x = O_3 + NO_2$) and (c)(d) aerosol liquid water content (ALWC) in 2015/2016 and 2018/2019 except for summer. The circles are colored according to the aerosol pH, and their size is linearly scaled with the SO_4^{2-} concentration. The markers inside the triangle indicate the observations with low O_x levels (<50 ppb). The measured OS concentrations within regular O_x or ALWC intervals (every ten unit-length) were averaged to more obviously demonstrate the trend (star symbols).

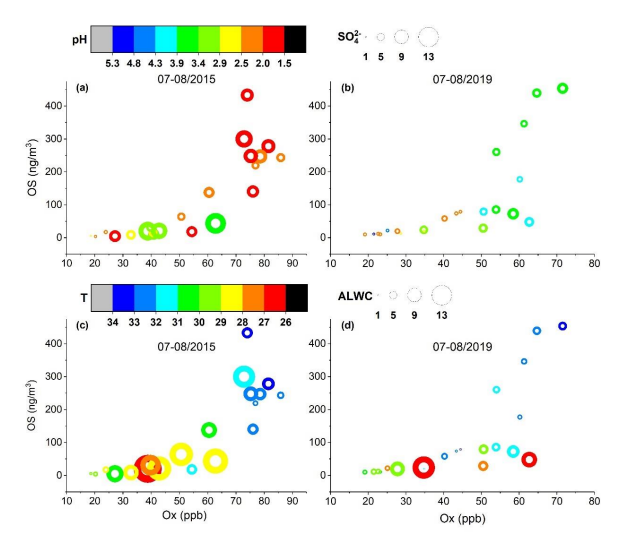


Figure 6. Quantified OS concentrations as a function of the Ox level in the summer of **(a) (c)** 2015 and **(b) (d)** 2019. The color of the circles in **(a) (b)** indicates the aerosol pH, and their size is linearly scaled with the SO₄²⁻ concentration. The color of circles in **(c) (d)** indicates ambient temperature and their size is linearly scaled with the aerosol liquid water content (ALWC).