SO\textsubscript{2} enhances aerosol formation from anthropogenic volatile organic compound ozonolysis by producing sulfur-containing compounds

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Abstract. Sulfur dioxide (SO\textsubscript{2}) can affect aerosol formation in the atmosphere, but the underlying mechanisms remain unclear. Here, we investigate aerosol formation and composition from the ozonolysis of cyclooctene with and without SO\textsubscript{2} addition in a smog chamber. Liquid chromatography equipped with high-resolution tandem mass spectrometry measurements indicates that monomer carboxylic acids and corresponding dimers with acid anhydride and aldol structures are important components in particles formed in the absence of SO\textsubscript{2}. A 9.4–12.6-times increase in particle maximum number concentration is observed in the presence of 14–192 ppb SO\textsubscript{2}. This increase is largely attributed to sulfuric acid (H\textsubscript{2}SO\textsubscript{4}) formation from the reactions of stabilized Criegee intermediates with SO\textsubscript{2}. In addition, a number of organosulfates (OSs) are detected in the presence of SO\textsubscript{2}, which are likely products formed from the heterogeneous reactions of oxygenated species with H\textsubscript{2}SO\textsubscript{4}. The molecular structures of OSs are also identified based on tandem mass spectrometry analysis. It should be noted that some of these OSs have been found in previous field studies but were classified as compounds from unknown sources or of unknown structures. The observed OSs are less volatile than their precursors and are therefore more effective contributors to particle formation and growth, partially leading to the increase in particle volume concentration under SO\textsubscript{2}-presence conditions. Our results provide an in-depth molecular-level insight into how SO\textsubscript{2} alters particle formation and composition.

1 Introduction

Secondary organic aerosol (SOA) accounts for a large fraction of the organic aerosol mass. The atmospheric oxidation of anthropogenic volatile organic compounds (AVOCs) can produce low-volatility organic products that contribute to SOA formation and growth (Kelly et al., 2018; Fan et al., 2020). The oxidation of AVOCs can dominate SOA formation under severe haze episodes (Nie et al., 2022; He et al., 2020; Huang et al., 2019; Qiu et al., 2020). Thus, AVOCs have been commonly considered as significant SOA precursors. SOA can negatively impact air quality, global climate, and public health (Nault et al., 2021; Zhu et al., 2017). To better understand air pollution and develop effective particle control strategies, it is necessary to investigate the formation mechanism and molecular composition of anthropogenic SOA.

Recently, the impacts of inorganic gases on aerosol chemistry have received significant attention (Deng et al., 2022). In particular, there is increasing evidence that sulfur dioxide (SO\textsubscript{2}) can modulate SOA formation and composition (Ye et al., 2018; Stangl et al., 2019; Liu et al., 2017). Liu et al. (2017) reported that SOA formation from cyclohexene photooxidation was inhibited by atmospherically relevant concentrations of SO\textsubscript{2}, as a result of the reaction of hydroxyl radical (\textsuperscript{-OH}) with SO\textsubscript{2} (to form sulfuric acid (H\textsubscript{2}SO\textsubscript{4})) competing with the \textsuperscript{-OH} reaction with cyclohexene. They demonstrated that H\textsubscript{2}SO\textsubscript{4}-catalyzed SOA enhancement was not sufficient to compensate for the loss of \textsuperscript{-OH} reactivity towards cyclohexene, leading to the suppression in cyclohexene SOA formation.
formation. On the other hand, SO$_2$ can enhance SOA formation and alter SOA composition by interacting with organic peroxides or stabilized Criegee intermediate (sCl) during the ozonolysis of alkenes (Stangl et al., 2019; Ye et al., 2018). For instance, under humid conditions, the reactive uptake of SO$_2$ into organic aerosols was obvious and reactions of SO$_2$ with organic peroxides could contribute to organosulfate (OS) formation (S. Wang et al., 2021; Ye et al., 2018). H$_2$SO$_4$ originating from sCl-induced oxidation of SO$_2$ is also linked to OS production (Stangl et al., 2019). OSs have been detected in different SO$_2$-alkene interaction areas (Hettiyadura et al., 2019; Wang et al., 2018; Bruggemann et al., 2020). Ubiquitous OSs may be used as tracers of SOA influenced by SO$_2$ emissions (Bruggemann et al., 2020). To gain further mechanistic insights into the complex roles of SO$_2$ in SOA formation, it is important to explore the chemical nature and formation mechanism of OSs.

Cycloalkenes emitted from diesel vehicles and industrial processes are a crucial class of AVOCs in the atmosphere. They can be used to explore key chemical processes involved in atmospheric oxidation and SOA formation (Räty et al., 2021). However, SOA formation chemistry from cycloalkenes has received less attention than that from linear or branched alkenes, leading to significant uncertainties in our understanding of SOA. Recent studies have reported that ozonolysis of cycloalkenes could form highly oxidized products and have considerable SOA yield (Räty et al., 2021; Rissanen, 2018). Among the most common cycloalkenes (with 5 to 8 carbon atoms), cyclooctene has the largest potential for SOA formation (Keywood et al., 2004). Ozonolysis is the dominant oxidation pathway of cyclooctene, with a reaction rate constant of $4.51 \times 10^{-16}$ cm$^3$ molec.$^{-1}$ s$^{-1}$ (298 K). Urban atmosphere is highly complex and may contain various concentrations of cycloalkenes and SO$_2$, which complicates SOA formation and composition. While previous studies have identified compounds containing carbon, hydrogen, and oxygen atoms (CHO compounds) as important contributors to cycloalkene SOA (Hamilton et al., 2006; Gao et al., 2004; Räty et al., 2021), the potential of OS formation from the ozonolysis of cyclooctene in the presence of SO$_2$ and the chemical processes behind OS formation remain unclear.

Given the significance of cycloalkene and SO$_2$ emissions in aerosol formation, we investigated the effects of SO$_2$ on the formation and chemical composition of cyclooctene SOA. Aerosol particles were formed from the ozonolysis of cyclooctene in the absence and presence of SO$_2$ in a smog chamber. Structural identifications of the observed products were reported and corresponding formation mechanisms were proposed. We report the mechanism, showing how SO$_2$ impacts particle formation and growth based on the observation of sulfuring-containing compounds. Our results provide a more comprehensive mechanistic understanding of the roles of SO$_2$ in modulating SOA formation and composition.

## 2 Experimental methods

### 2.1 Particle production

Particle formation from the ozonolysis of cyclooctene ($k_{298K} = 4.51 \times 10^{-16}$ cm$^3$ molec.$^{-1}$ s$^{-1}$) was carried out under dark conditions in a 1.2 m$^3$ Teflon chamber housed in a temperature-controlled room. A summary of experimental conditions and results is listed in Table 1. Detailed experimental equipment and methods have been described in our previous studies (Yang et al., 2022, 2021). Particle formation experiments were operated in a batch mode. Briefly, cyclooctene was introduced into the chamber by passing zero air through a tube containing a known volume of cyclooctene (95 %, Alfa). Then, cyclohexane (99.5 %, Aladdin) was injected in excess (∼ 130 ppm) into the chamber so that more than 98 % of OH generated during the ozonolysis of cyclooctene was scavenged. Control experiments showed that the presence of cyclohexane could lead to a significant decrease in particle volume concentration (Fig. S1 in the Supplement).

When desired, SO$_2$ was added to the chamber from an SO$_2$ calibration cylinder. Initial concentration ratios of SO$_2$ to cyclooctene were in the range of ∼ 0.07–1 ppb ppb$^{-1}$ to simulate different polluted atmospheric conditions. The reactor was stabilized for 20 min under dark conditions to allow for mixing of species. Finally, ozonolysis of cyclooctene was initiated by introducing ozone (O$_3$) produced via a commercial ozone generator (WH-H-Y5Y, Wohuan, China). All experiments were performed at room temperature (∼ 295 K) and atmospheric pressure (∼ 1 atm) without seed particles. Temperature and relative humidity (RH) inside the chamber were measured with a hygrometer (Model 645, Testo AG, Germany). O$_3$ and SO$_2$ concentrations over the course of ozonolysis were monitored by a Thermo Scientific model 491 O$_3$ analyzer and a Thermo Scientific model 43i-TLE SO$_2$ analyzer, respectively. The detection limits of O$_3$ analyzer and SO$_2$ analyzer were 0.5 and 0.05 ppb, respectively. Size distributions and volume concentrations of particles were continuously recorded using a scanning mobility particle sizer (SMPS), which consisted of a differential mobility analyzer (Model 3082, TSI, USA) and an ultrafine condensation particle counter (Model 3776, TSI, USA). The particle volume concentration was measured continuously until we observed a decrease. The particle formation experiments proceeded for 300 min before the collection of aerosol particles.

### 2.2 Particle collection and chemical characterization

Aerosol particles were collected on aluminum foils using a 14-stage low-pressure impactor (DLPI+, Dekati Ltd, Finland). All samples were stored in a −20 °C freezer until analysis. Offline functional group measurements of aerosol particles were performed using an attenuated total reflectance-Fourier transform infrared spectrometer (ATR-FTIR, Vertex 70, Bruker, Germany). Before each measurement, the dia-
The LC gradient employed was as follows: 0–3 min at 3 % acetic acid (A) and methanol with 0.1 % (v/v) formic acid (B). The injection volume of samples and flow rate were 2 µL and 200 µL min\(^{-1}\), respectively. The ESI source was operated in both positive (+) and negative (−) ion modes to ionize analyte components with a scan range of mass-to-charge (m/z) of 50 to 750. LC/ESI-MS parameter settings were as follows: 3.5 kV spray voltage (+), −3.0 kV spray voltage (−), 50 V S-lens radiofrequency (RF) level (+), 50 V S-lens RF level (−), 320 °C capillary temperature, 2.76 × 10^5 Pa sheath gas (nitrogen) pressure, and 3.33 L min\(^{-1}\) auxiliary gas (nitrogen) flow. Data-dependent tandem mass spectrometry (MS/MS) analysis was also carried out by high-energy collision-induced dissociation (CID) with stepped collision energies of 20, 40, and 60 eV. For MS/MS experiments, an isolation width of 2 m/z units was applied. Other parameters were also selected in MS/MS experiments as follows: 2 × 10^5 automatic gain control (AGC) target, 50 ms maximum IT, 3 loop count, 1 × 10^6 minimum AGC target, 2–6 s apex trigger, and 6 s dynamic exclusion. The mass resolution of MS and MS/MS were 70 000 (full width at half maximum, FWHM, at m/z 200) and 17 500, respectively. Detailed data processes are reported elsewhere (Yang et al., 2021, 2022). The double bond equivalent (DBE) value is the combined number of rings and double bonds in the product C\(_{x}\)H\(_{y}\)O\(_{z}\)N\(_{p}\)S\(_{q}\). Here, subscripts \(c\), \(h\), \(o\), \(n\), and \(s\) represent the number of carbon, hydrogen, oxygen, nitrogen, and sulfur atom in the product C\(_{x}\)H\(_{y}\)O\(_{z}\)N\(_{p}\)S\(_{q}\) and could be calculated according to Eq. (1). For organosulfate, the two S = O bonds in the sulfate group were not considered based on calculations in previous studies (Wang et al., 2016; Riva et al., 2016b; Kuang et al., 2016). The DBE value of organosulfate reflects the unsaturation degree of its side carbon chain.

\[
\text{DBE} = 1 + c + \frac{n - h}{2} .
\]  

Kendrick mass defect (KMD) analysis could provide chemical insights into the chemical compositions of complex organic mixtures (Kundu et al., 2017; Kenseth et al., 2020). The KMD value is the same for homologous species that differ from each other only by their base units. CH\(_{2}\) and the oxygen atom (O) are usually chosen as base units for Kendrick analysis of complex organic mass spectra. Kendrick mass (KM) could be converted into a new mass scale from the IUPAC mass (Eqs. 2 and 4). KMD is determined as the difference between the nominal mass of a compound (the rounded inte-
The wall loss rate of O\textsubscript{2}SO\textsubscript{4} was determined to be 2.05\(\times\)10\textsuperscript{-3} min\textsuperscript{-1} (Fig. S2), indicating that the losses of these two gas-phase species to the chamber walls were negligible over the course of experiments. The wall loss of cyclooctene (5.23\(\times\)10\textsuperscript{-6} min\textsuperscript{-1}) was also negligible, while its oxidation products may deposit to the inner walls. However, wall losses of gas-phase products could be mitigated due to excess O\textsubscript{3} concentration (Sect. S1 in the Supplement). The quick oxidation and nucleation could provide attractive condensation surfaces for oxidation products, thereby reducing the product wall losses to some extent (Stirnweis et al., 2017). Although wall losses of organic vapors may underestimate the particle mass, this work mainly focuses on the characterization of particle composition rather than the absolute SOA yield.

Independent wall-loss experiments of ammonium sulfate ((NH\textsubscript{4})\textsubscript{2}SO\textsubscript{4}) particles were also performed to determine the size-dependent wall-loss rate constants of particles inside the chamber. An aqueous (NH\textsubscript{4})\textsubscript{2}SO\textsubscript{4} solution was added to a TSI Model 3076 atomizer to produce droplets. The droplets were passed through a silica gel diffusion dryer to get dry (NH\textsubscript{4})\textsubscript{2}SO\textsubscript{4} particles and then these were injected into the chamber. The size distributions of (NH\textsubscript{4})\textsubscript{2}SO\textsubscript{4} particles were characterized using the SMPS for 6 h. The relationship between the wall-loss rate (\(k\), h\textsuperscript{-1}) of particles and their size (\(d_p\), nm) can be expressed as \(k(d_p) = 1.20 \times 10^{-7} \times d_p^{2.32} + 20.59 \times d_p^{1.39}\) based on a size-dependent particle wall-loss correction method.

3 Results and discussion

3.1 SO\textsubscript{2} effects on aerosol formation

Insights into SO\textsubscript{2} effects on particle formation could be gained through investigating the number and volume concentration as well as the size distribution of particles under various SO\textsubscript{2} level conditions. In the absence of SO\textsubscript{2}, the particle number concentration increased in a burst within the first 20 min of cyclooctene ozonolysis and then decreased because of their coagulation and wall deposition, while the particle volume concentration increased gradually and reached its maximum within 240 min (Fig. S3). An elevating SO\textsubscript{2} level can result in significant increases in the number and volume concentration of particles (Fig. 1a), which is consistent with observations from previous studies (Ye et al., 2018; Yang et al., 2021). We observed a 9.4–12.6-times increase in particle maximum number concentration in the presence of 14–192 ppb SO\textsubscript{2} (Table 1). The promoted effect of SO\textsubscript{2} is shown more clearly in Fig. 1b, where SO\textsubscript{2} was seen to be consumed on a similar timescale to particle formation. Specifically, upon initiation of cyclooctene ozonolysis, SO\textsubscript{2} concentration decreased and the particle volume concentration increased simultaneously. After cyclooctene was completely consumed, both SO\textsubscript{2} consumption and particle production slowed down. SO\textsubscript{2} consumption and particle formation resumed when more cyclooctene was introduced into the reactor. This result indicates that SO\textsubscript{2} may react with certain highly reactive species produced from cyclooctene ozonolysis. For instance, reactions of SO\textsubscript{2} with sCl could form H\textsubscript{2}SO\textsubscript{4} (Boy et al., 2013), which is a key species for new particle formation (Lehtipalo et al., 2018; Yao et al., 2018). Inorganic sulfate absorption at 617 cm\textsuperscript{-1} was observed in the ATR-FTIR spectra of particles formed in cyclooctene–O\textsubscript{3}–SO\textsubscript{2} systems (Fig. 2) (Hawkins et al., 2010; Coury and Dillner, 2008), indicating the formation of H\textsubscript{2}SO\textsubscript{4}. We assumed that all consumed SO\textsubscript{2} was converted to particle-phase H\textsubscript{2}SO\textsubscript{4}, which represents an upper limit of H\textsubscript{2}SO\textsubscript{4} formation (Wyche et al., 2009; Ye et al., 2018). The amount of H\textsubscript{2}SO\textsubscript{4} produced could not fully account for the enhancement of particle volume concentration (Table 1). H\textsubscript{2}SO\textsubscript{4} has been considered as an important driver of particle acidity (Tilgner et al., 2021). Acid catalysis induced by H\textsubscript{2}SO\textsubscript{4} may also promote the formation of additional organic products, leading to the increase in particle volume concentration (Deng et al., 2021).

SO\textsubscript{2} can also affect the growth of new aerosol particles (Fig. 1c). Once O\textsubscript{3} was introduced into the reactor, aerosol particles were produced rapidly. After cyclooctene was depleted, the aerosol particle mass increased slowly. The initial stage of particle formation was then defined as the time from reaction initiation to the complete consumption of cyclooctene (\(\sim\)10 min). From Fig. 1c, in the initial stage of ozonolysis (10 min), particles formed in cyclooctene–O\textsubscript{3}–SO\textsubscript{2} systems had a smaller size mode than those formed in
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The cyclooctene–O\textsubscript{3} system, which may be attributed to the following two factors: first, oligomers formed from sCI reactions with organic species could partition into the condensed phase to contribute to particle growth (Riva et al., 2017). SO\textsubscript{2} presence may lead to a decrease in these oligomers because SO\textsubscript{2} can compete with organic species to react with sCI. Second, counterbalancing the reduction of oligomers via s\textsubscript{CI} + SO\textsubscript{2} reactions is the production of H\textsubscript{2}SO\textsubscript{4}. The production of more new particles in cyclooctene–O\textsubscript{3}–SO\textsubscript{2} systems could provide more condensation sinks. Organic vapors that can condense onto particles are dispersed via new particles, resulting in small particle size at the initial phase of cyclooctene–O\textsubscript{3}–SO\textsubscript{2} systems (Stangl et al., 2019). Interestingly, particles could grow quickly in the presence of SO\textsubscript{2}. At 300 min reaction time, particles formed in the presence of SO\textsubscript{2} even had slightly larger sizes than those formed in the absence of SO\textsubscript{2}. H\textsubscript{2}SO\textsubscript{4}-catalyzed heterogeneous reactions could produce lower-volatility organic species from higher-volatility reactants in the aerosol phase (Yang et al., 2020; Han et al., 2016). Semi-volatile species could undergo evaporation after partitioning to the aerosol phase while low-volatility products generally have a negligible evaporation rate from the aerosol phase. Low-volatility products formed via H\textsubscript{2}SO\textsubscript{4}-catalyzed heterogeneous reactions could build particle mass at a rate almost equal to the condensation rate and, thus, effectively facilitate particle growth in cyclooctene–O\textsubscript{3}–SO\textsubscript{2} systems (Apsokardu and Johnston, 2018).

3.2 Aerosol chemical composition under SO\textsubscript{2}-free conditions

Figure 3 shows the base peak chromatograms (BPCs) of cyclooctene-derived particles in the absence of SO\textsubscript{2}. The chromatograms of blank filter clearly showed no peaks eluted at retention times (RTs) between 0 and 30 min, while there were several significant peaks for cyclooctene SOA chromatograms in both positive and negative ion modes. Each chromatogram peak of cyclooctene SOA represents at least one ion, and major peaks are only labeled with the mass of the most abundant single ion. Compared to the negative chromatogram of cyclooctene SOA, the corresponding label ions in the positive chromatogram were 24 Da higher in mass. This is consistent with the fact that many ions produce adducts with sodium ion ([M + Na]\textsuperscript{+}) in positive ion mode, while negative ion mode leads to the production of deprotonated ions ([M - H]\textsuperscript{-}) (Mackenzie-Rae et al., 2018).

From Fig. 3, products with molecular weight (MW) <200 Da eluted from the column at shorter RTs than those with MW >200 Da. Low-molecular-weight products (MW <200 Da) likely correspond to small monomer type compounds (hereafter termed monomeric products), which directly originate from the ozonolysis of cyclooctene. Compounds with MW >200 Da mainly dominate the later part of the chromatogram, and they may be homo- or heterodimeric products.
species (hereafter denoted dimeric products) formed using two monomeric products as building blocks.

Possible structures of major monomeric products were proposed based on their accurate \( m/z \), fragmentation mass spectra, and previous mechanistic insights. Note that the fragmentation of [M + Na]\(^+\) is relatively difficult (Zhao et al., 2016) and, thus, the positive ion mode was not further analyzed in providing structural insights in the current study. The negative chromatogram peaks with RT at 11.85 min (N-145), 16.13 min (N-159), and 20.41 min (N-173) were significant peaks for cyclooctene SOA (Fig. 3b), and they were assigned neutral formulas of \( \text{C}_8\text{H}_{10}\text{O}_4 \), \( \text{C}_7\text{H}_{12}\text{O}_4 \), and \( \text{C}_8\text{H}_{14}\text{O}_4 \), respectively. As shown in Fig. 4, MS/MS spectra of monomer \( \text{C}_8\text{H}_{10}\text{O}_4 \), \( \text{C}_7\text{H}_{12}\text{O}_4 \), and \( \text{C}_8\text{H}_{14}\text{O}_4 \) were similar. Taking \( \text{C}_8\text{H}_{14}\text{O}_4 \) as an example (Fig. 4c), its fragmentation mass spectrum was characterized by a loss of 44 Da (\( \text{CO}_2 \)), suggesting the presence of a carboxyl group. The neutral loss of 18 Da (\( \text{H}_2\text{O} \)) upon fragmentation of the parent ion (\( \text{C}_8\text{H}_{13}\text{O}_4^- \), \( m/z = 173.08209 \)) led to the production of an ion with \( m/z = 155.07143 \). The loss of \( \text{H}_2\text{O} \) is an unspecific fragmentation mechanism, which likely originates from a carboxyl or hydroxyl group (Noziere et al., 2015). The fragment ion (\( m/z = 111.08166 \)) representing the simultaneous neutral losses of \( \text{CO}_2 \) and \( \text{H}_2\text{O} \) was also formed. MS/MS spectra can result from multiple isomeric structures in many cases (Wang et al., 2019). Yasmeen et al. (2011) showed the detailed fragmentation spectrum for the dicarboxylic acid standard (azelaic acid) and indicated that deprotonated azelaic acid also showed losses of \( \text{H}_2\text{O} \), \( \text{CO}_2 \), and \( \text{CO}_2+\text{H}_2\text{O} \). In addition, Noziere et al. (2015) showed that the neutral losses of \( \text{CO}_2 \) and \( \text{H}_2\text{O} \) indicates two carboxyl groups. Thus, monomer \( \text{C}_8\text{H}_{14}\text{O}_4 \) was tentatively assigned to suberic acid and the corresponding fragmentation pathway for \( \text{C}_8\text{H}_{13}\text{O}_4^- \) is proposed in Fig. S4. The fragment ions originating from losses of \( \text{H}_2\text{O} \), \( \text{CO}_2 \), and \( \text{CO}_2+\text{H}_2\text{O} \) were also observed in MS/MS spectra of \( \text{C}_6\text{H}_{10}\text{O}_4 \) and \( \text{C}_7\text{H}_{12}\text{O}_4 \), indicative of adipic acid and pimelic acid, respectively. Carboxylic acids have also been observed in SOA produced from previous alkene ozonolysis (Hamilton et al., 2006; Kenseth et al., 2020; Mackenzie-Rae et al., 2018; Zhang et al., 2015). Carboxylic acids represent a significant class of aerosol components, and they play a significant role in particle chemistry via their influences on particle acidity and through direct involvement in certain heterogeneous reactions to produce low-volatility species (Millet et al., 2015). More experiments using available authentic standards are necessary to better understand their structures, sources, and formation mechanism. Other prominent monomer peaks at RTs of 8.30 min (N-175) and 14.18 min (N-189) corresponded to compounds with neutral formula, namely \( \text{C}_7\text{H}_{12}\text{O}_3 \) and \( \text{C}_8\text{H}_{14}\text{O}_5 \). The losses of \( \text{H}_2\text{O} \), CO, and \( \text{CO}_2 \) in the MS/MS spectrum of \( \text{C}_7\text{H}_{12}\text{O}_5 \) indicated hydroxyl, terminal carbonyl, and carboxyl groups, respectively (Mackenzie-Rae et al., 2018; Riva et al., 2016a), and \( \text{C}_7\text{H}_{12}\text{O}_3 \) was identified as hydroxy-containing oxoheptanoic acid (Fig. S5a and c). Monomer \( \text{C}_8\text{H}_{14}\text{O}_5 \) only showed losses of \( \text{H}_2\text{O} \) and CO (Fig. S5b), and it is difficult to determine the specific type and positioning of oxygen-containing functionalities within \( \text{C}_8\text{H}_{14}\text{O}_5 \) with five oxygen atoms based on its MS/MS spectrum. The labeled dimer peaks in negative ion mode corresponded to [M – H]\(^-\) ion masses of 271, 285, and 331 (Fig. 3b), which were assigned neutral formulas of \( \text{C}_{13}\text{H}_{26}\text{O}_5 \), \( \text{C}_{15}\text{H}_{26}\text{O}_5 \), and \( \text{C}_{15}\text{H}_{26}\text{O}_8 \), respectively. The number of fragment ions of dimers are generally limited, and determining the exact structure of dimers is less certain compared to monomers (Witkowski and Gierczak, 2017). Therefore, only a decrease in molecular structure possibilities is provided. For dimer \( \text{C}_{15}\text{H}_{26}\text{O}_8 \), fragment ions \( m/z = 159.06642 \left( \text{C}_7\text{H}_{11}\text{O}_7^- \right) \) and \( m/z = 189.07722 \left( \text{C}_8\text{H}_{12}\text{O}_5^- \right) \) were detected in the MS/MS spectrum (Fig. 4d). When dimers are subjected to CID, fragment ions corresponding to their building blocks are commonly observed (Witkowski and Gierczak, 2017; Hall and Johnston, 2012). Based on this
rule, it could be concluded that dimer \( C_{15}H_{24}O_7 \) was an association product of \( C_7H_{12}O_4 \) and \( C_8H_{14}O_4 \). Similarly, for dimer \( C_{15}H_{24}O_7 \), there were two significant product ions of \( C_{15}H_{23}O_7^- \), with accurate masses of \( m/z \) 159.06651 (\( C_7H_{11}O_4^- \)) and 173.08217 (\( C_8H_{13}O_3^- \)) (Fig. 4e). Furthermore, fragment ions corresponding to secondary loss of \( \text{CO}_2 + \text{H}_2\text{O} \) from product ions \( C_7H_{12}O_4^- \) and \( C_8H_{13}O_3^- \) were also observed. The fragmentation spectrum of \( C_{15}H_{24}O_7 \) was similar to the MS/MS spectra of \( C_7H_{12}O_4 \) and \( C_8H_{14}O_4 \) (Fig. 4b and c), suggesting again that \( C_7H_{12}O_4 \) and \( C_8H_{14}O_4 \) may be the building blocks of \( C_{15}H_{24}O_7 \). Acid-catalyzed heterogeneous processes can result in the formation of high-molecular-weight dimers in both biogenic and anthropogenic systems (Barsanti et al., 2017). Carboxylic acid monomers formed could be important sources of particle acidity in the absence of \( \text{SO}_2 \). Dimers \( C_{15}H_{24}O_7 \) and \( C_{15}H_{24}O_8 \) may be produced by heterogeneous reactions involving the loss of a water molecule, and the linkage between building blocks is an acid anhydride (Fig. S6) (Hamilton et al., 2006). Another abundant dimer peak (N-271) in the negative chromatogram was identified as \( C_{14}H_{23}O_5^- \) with mass accuracy of \(-0.02492\) ppm. \( C_{14}H_{23}O_5^- \) could dissociate to the product ions of \( C_{14}H_{21}O_4^- \), \( C_{13}H_{21}O_5^- \), and \( C_7H_{11}O^- \) (Fig. 4f). Both secondary ozonide and aldol structures shown in Fig. 4f could match the assigned elemental formula of \( C_{14}H_{24}O_5 \). However, the neutral losses of \( \text{H}_2\text{O} \) and \( \text{CO}_2 \) were not easily produced by secondary ozonide, but were more likely for the aldol structure (Hall and Johnston, 2012). Aldol condensation products were also among the most commonly observed species in previous ozonolysis of alkenes (Zhao et al., 2016; Kenseth et al., 2018; Kristensen et al., 2016). Therefore, \( C_{14}H_{24}O_5 \) as shown in Fig. 4f is likely an aldol condensation product.

To examine the overall composition of particles, average mass spectra (Fig. S7) corresponding to the chromatogram where particle components eluted were also analyzed. Figure 5 summarizes the oxidation products observed in particles mapped in O-KMD and van Krevelen plot. The molecular formulas of identified oxidation products could be largely classified into homologous series of monomers and dimers (Fig. 5a and b). The elemental composition distribution of products measured in positive and negative ion modes was
similar, with most monomers and dimers having O/C ratios ranging from 0.2 to 0.8, and H/C ratios ranging from 1.2 to 1.8 (Fig. 5c). Lines with slopes of 0, −0.5, −1, and −2 in Fig. 5c can be used to illustrate the addition of hydroxyl/peroxide, carboxylic acid (with fragmentation), carboxylic acid (without fragmentation), and carbonyl groups to a saturated carbon chain, respectively (Heald et al., 2010). As shown in Fig. 5c, cyclooctene SOA occupied a relatively wide range in the van Krevelen diagram, and there are a large number of points scattered in the space between lines with slopes of −0.5 and −2. This behavior is consistent with the importance of high-abundance carboxylic acids in the above analysis.

3.3 SO2 effects on aerosol chemical composition

To obtain further detailed mechanisms regarding SO2 effects and determine whether heterogeneous processes occur, aerosol samples were analyzed using ATR-FTIR and LC/ESI-MS. Both IR and MS analysis of particles revealed changes in aerosol chemical composition induced by SO2 addition.

3.3.1 Characteristics of functional group in aerosol-phase products

Figure 2 shows ATR-FTIR spectra of aerosol particles. Hydroxy (3600–3200 cm\(^{-1}\)), alkyl (2935 and 2864 cm\(^{-1}\)), and carbonyl (1702 cm\(^{-1}\)) groups were identified in particles collected from the cyclooctene–O3 system (Table S1 in the Supplement). These particles also had a broad absorption across the 1500–800 cm\(^{-1}\) region, which may arise from C–H deformation in 1480–1350 cm\(^{-1}\), C–C stretching in 1250–1120 cm\(^{-1}\), and C–O stretching in different regions for various oxygenated species (Hung et al., 2013). Three additional absorption bands at 1413, 1095, and 617 cm\(^{-1}\) were observed in ATR-FTIR spectra of particles formed with the introduction of SO2 (Tammer, 2004; Lal et al., 2012). Absorption bands at 1413 and 1095 cm\(^{-1}\) may be associated with asymmetric and symmetric stretching of −SO2− while inorganic sulfates could give rise to strong absorption at 617 cm\(^{-1}\). The presence of absorption bands of sulfur-containing groups suggests that SO2 addition can result in the production of sulfur-containing compounds.

3.3.2 Organosulfate formation in the presence of SO2

In addition to CHO compounds, products with C\(_n\)H\(_{2n}\)O\(_s\)S\(_r\) elemental formulas were identified in the presence of SO2 (Fig. S8). OS could undergo highly efficient ionization to give deprotonated molecular ions in negative ion mode. Based on MS/MS analysis, unambiguous identification of OS can be achieved, since OSs could give characteristic fragment ions at m/z 80 (SO\(_3^-\)), 81 (HSO\(_3^-\)), and/or 97 (HSO\(_4^-\)) in their MS/MS spectra (Figs. S9–S17). Accurate mass measurements of OSs as well as their retention times and DBE values are provided in Table S2. The proposed structure and fragmentation scheme of each OS and the corresponding precursor are presented in Figs. S9–S17. For instance, OS-209 and OS-223 showed prominent product ions for losses of HSO\(_3^-\) and SO\(_3^-\) (Figs. S11–S12), confirming the organosulfate moiety. Neither a hydroxyl nor a carboxyl group fragment ion (i.e., −H\(_2\)O or −CO\(_2\)) was observed in their MS/MS spectra. C\(_6\)H\(_{10}\)O\(_3\) and C\(_7\)H\(_{12}\)O\(_3\) were proposed as the precursors of OS-209 and OS-223, respectively. MS/MS spectra of C\(_6\)H\(_{10}\)O\(_3\) and C\(_7\)H\(_{12}\)O\(_3\) were character-
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Figure 6. Simplified formation schemes for the selected organosulfates formed from the ozonolysis of cyclooctene.

Figure 7. (a) Two-dimensional volatility–oxidation space of the identified organosulfurs and their precursors. (b) Carbon atom number distribution of organosulfurs observed in the current work and in the studies of Cai et al. (2020), Boris et al. (2016), and Y. Wang et al. (2021). Detailed formulae of these OSs can be found in Table S3. Organosulfurs from previous studies are of unknown origin or unknown structure.

ized by loss of CO, indicating a terminal carbonyl group (Figs. S11–S12). Considering structural features of OS precursor measurements as well as OS-209 and OS-223 all corresponding to DBE = 2, two carbonyl groups could well explain the observed MS/MS spectra of OS-209 and OS-223. The organosulfate substituent was expected to attach to a terminal carbon atom. Although the carbonyl group is more readily observed in positive ion mode, ESI-MS is also highly sensitive to carbonyl compounds containing sulfate substituents and thereby gives intense [M − H]⁻ ions in negative ion mode (Riva et al., 2016b).

A relatively high abundance of OS is helpful for the acquisition of MS/MS data, and therefore high-abundance [M − H]⁻ ions were chosen as representative candidates to clarify the precursors and formation pathways of OSs. A simplified chemical mechanism describing OS production
from the ozonolysis of cyclooctene (C₈H₁₄) is proposed in Fig. 6. The ozonolysis of cyclooctene (C₈H₁₄) can be initiated by O₃ addition to the endocyclic double bond, forming an energy-rich primary ozonide (POZ). POZ can decompose rapidly to an excited CI containing both a terminal carbonyl and carbonyl oxide group. The excited CI could lead to the formation of sCI, vinylhydroperoxide, and dioxirane, illustrating the multiplicity and the complexity of cyclooctene ozonolysis. SCI is mainly capable of involvement in bimolecular reactions to form carboxylic acids and acid esters. Vinylhydroperoxide rapidly decomposes into an alkyl radical (C₈H₁₃O₂•) and an •OH. Molecular oxygen could be subsequently added to C₈H₁₃O₂• to produce an alkyl peroxy radical (RO₂, C₈H₁₃O₄•). Dioxirane intermediate may also undergo decomposition and produce C₇H₁₃O₃•, C₆H₁₃O₄•, and C₅H₁₃O₃•. As the starting point of the RO₂•- and alkoxy radical (RO•) chemistry, resulting in termination CHO compounds with hydroperoxy, carbonyl, or hydroxy groups (Fig. 6). Acid-catalyzed heterogeneous reactions of CHO products have been evidenced to play a major role in OS formation in the atmosphere (Riva et al., 2016a, b). Although acidic seed particles were not directly injected into the reactor during cyclooctene ozonolysis, SO₂-induced H₂SO₄ may create acidic conditions for the occurrence of heterogeneous reactions. In the case of CHO products with a hydroxyl group, H₂SO₄ could protonate the hydroxyl group, leading to the formation of OS and water. The low RH (∼20%) of ozonolysis was helpful for shifting the reaction equilibrium in favor of OS production. Detailed information about the volatility of oxidation products is necessary to evaluate their potential to contribute to aerosol formation. As shown in Fig. 7a, the products could be categorized into intermediate-volatility OCs (IVOCs), semi-volatile OCs (SVOCs), low-volatility OCs (LVOCs), and extremely low-volatility OCs (ELVOC), with C° in the range of 300–3 × 10⁶, 0.3–300, 3 × 10⁻⁴–0.3, and <3 × 10⁻⁴ µg m⁻³, respectively (Donahue et al., 2011). The saturation mass concentration of OSs spanned more than six orders of magnitude (Fig. 7a), suggesting their inherent chemical complexity. A large number of OSs are SVOCs and LVOCs, while their precursors are classified as IVOCs and SVOCs, indicating that the SO₂ presence facilitates the reduction of product volatility (Yang et al., 2020; Han et al., 2016). Lower-volatility OSs generated from acid-catalyzed heterogeneous reactions may build particle mass at a faster rate compared to their higher-volatility precursors, and thereby benefit the formation and growth of particles in the presence of SO₂.

Figure 7b displays the DBE–carbon atom number space for organosulfur compounds. There are some overlaps of organosulfur compounds detected in this work with previous data from field observations (Y. Wang et al., 2021; Boris et al., 2016; Cai et al., 2020). For example, Y. Wang et al. (2021) comprehensively analyzed OS in PM₂.₅ filter samples collected at an urban site in Shanghai, China, and observed the presence of C₆H₁₀O₆S (Fig. 7b, cyan cross). In the absence of chromatographic data such as retention times, C₆H₁₀O₆S was tentatively assigned to diesel vapor-derived OS. Alkenes are important components of diesel and cyclooctene may be also responsible for C₆H₁₀O₆S formation in the atmosphere. The overlaps of organosulfur compounds indicate that the ozonolysis of cycloalkenes in the presence of SO₂ is likely an important source of organosulfur compounds in the ambient atmosphere. In addition, our work further suggests that the sources of OS cannot be determined only based on their elemental formula, and techniques that enable the identification of molecular structures (e.g., MS/MS) are greatly beneficial in field studies. The identified molecular structures of OSs in this study are also helpful in source apportionment in field studies.

4 Conclusion

We have explored O₃-initiated oxidation of cyclooctene in the absence and presence of SO₂, with a focus on the mechanism by which SO₂ impacts particle formation and composition. Cyclooctene can produce a large number of particles upon reacting with O₃. Higher SO₂ concentration led to higher particle number concentration as a result of H₂SO₄ formation from the reactions of sCI with SO₂. Cyclooctene SOA mainly consisted of carboxylic acids and corresponding dimers with acid anhydride and aldo structures when SO₂ was not added. SO₂ addition can induce changes in particle chemical composition through the formation of OSs. Some OSs, classified as compounds of unknown origin or unknown structure in previous field studies, were also observed in this work. The OSs found here are less volatile than their precursors, indicating the stronger ability of OS for particle formation. The formation of OSs can in part lead to the increase in particle volume concentrations in the presence of SO₂.

The results here suggest that SO₂ can influence aerosol particle formation and composition by producing sulfur-containing compounds (i.e., H₂SO₄ and OSs). Nevertheless, the observed number of OSs may be amplified by the high SO₂ concentration used in the present work. In order to determine the actual mass yields of OSs and better quantify roles of SO₂ in particle formation, further experiments using ambient SO₂ levels and authentic standards are warranted.

Data availability. Experimental data are available upon request to the corresponding author.

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