Secondary organic aerosol formation from OH-initiated oxidation of m-xylene: effects of relative humidity on yield and chemical composition

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Abstract. The effect of relative humidity (RH) on the secondary organic aerosol (SOA) formation from the photooxidation of m-xylene initiated by OH radicals in the absence of seed particles was investigated in a smog chamber. The SOA yields were determined based on the particle mass concentrations measured with a scanning mobility particle sizer (SMPS) and reacted m-xylene concentrations measured with a gas chromatograph-mass spectrometer (GC-MS). The SOA components were analysed using Fourier transform infrared spectrometer (FTIR) and ultrahigh performance liquid chromatograph-electrospray ionization-high-resolution mass spectrometer (UPLC-ESI-HRMS). A significant discrepancy was observed in SOA mass concentration and yield variation with the RH conditions. The SOA yield is 13.8\% and 0.8\% at low RH (13.7\%) and high RH (79.1\%), respectively, with the difference being over an order of magnitude. The relative increase of C-O-C at high RH from the FTIR analysis of functional groups indicates that the oligomers from carbonyl compounds cannot well explain the suppression of SOA yield. Highly oxygenated molecules (HOMs) were observed to be suppressed in the HRMS spectra. The chemical mechanism for explaining the RH effects on SOA formation from m-xylene-OH system is proposed based on the analysis of both FTIR and HRMS measurements as well as Master Chemical Mechanism (MCM) simulations. The reduced SOA at high RH is mainly ascribed to the less formation of oligomers and the suppression of RO$_2$ autoxidation. As a result, high RH can obstruct the oligomerization and autoxidation that contribute to the SOA formation.

1 Introduction

Secondary organic aerosol (SOA) is a significant component of atmospheric fine particulate matter in the troposphere (Hallquist et al., 2009; Spracklen et al., 2011; Huang et al., 2014), leading to serious concerns as it has a significant influence on the air quality, oxidative capacity of the troposphere, global climate change and human health (Jacobson et al., 2000; Hansen and Sato, 2001; Kanakidou et al., 2005; Zhang et al., 2014). In a previous study from a global model simulation, it has been found that SOA represents a large fraction, approximately 80\% of the total organic aerosol sources (Spracklen et al., 2011).
The formation of SOA in the atmosphere is principally via the oxidation of volatile organic compounds (VOCs) by common atmospheric oxidants such as O₃, OH and NO₃ radicals (Seinfeld and Pandis, 2016). Aromatic compounds mainly from anthropogenic source, including solvent usage, oil-fired vehicles and industrial emissions, contribute 20-30% to the total VOCs in urban atmosphere, which play a significant role in the formation of ozone and SOA in the urban troposphere (Forstner et al., 1997; Odum et al., 1997; Calvert et al., 2002; Bloss et al., 2005; Offenberg et al., 2007; Ding et al., 2012; Zhao et al., 2017). Amongst aromatics, m-xylene is significant, of which mean concentration together with p-xylene in daytime was determined up to 140.8 µg m⁻³ in atmosphere of urban areas in developing countries (Khoder, 2007). The oxidation of aromatics in the troposphere is mainly initiated through OH radicals, which is affected by many chemical and physical factors. The concentrations of oxidant species, VOCs and NOₓ concentrations, as well as the ratio of VOCs to NOₓ (Ge et al., 2017a) determine the main chemical mechanism. Light intensity (Warren et al., 2008), temperature (Qi et al., 2010) and relative humidity (RH) are the most significant physical parameters that affect the chemical process. RH governs the water concentration in the gas phase and the liquid water content (LWC) in the particle phase. Water plays a significant role that can serve as reactant, product and solvent to directly participate in chemistry (Finlayson-Pitts and Pitts Jr., 2000) and indirectly affect the reaction environment such as acidity of particles (Jang et al., 2002).

Investigations of RH effects on aromatics SOA have been conducted in many previous works. In the presence of NOₓ, it was observed that RH significantly enhanced the yield of SOA from benzene, toluene, ethylbenzene and xylenes photooxidation, which was explained by a higher formation of HONO, particle water, aqueous radical reactions and the hydration from glyoxal (Healy et al., 2009; Kamens et al., 2011; Zhou et al., 2011; Jia and Xu, 2014, 2018; Wang et al., 2016). Meanwhile, under low NOₓ condition that no NOₓ were introduced artificially and photolysis of H₂O₂ was as an OH radical source, it has been observed that deliquesced seed contributed to the enhancement of SOA yield from toluene (Faust et al., 2017; Liu et al., 2018). However, under low NOₓ level, it has been found that, in the study on toluene SOA formation, moderate RH level (48%) leads to a lower SOA yield than low RH level (17-18%) (Cao and Jang, 2010). In a most recent study on SOA formation of toluene (Hinks et al., 2018), high RH led to a much lower SOA yield than low RH under low NOₓ level, which is attributed to condensation reactions that remove water, leading to the less oligomerization at high RH. In a study on chemical oxidative potential of SOA (Tuet et al., 2017) under low NOₓ conditions, it was observed that the mass concentration of SOA from m-xylene irradiation under the dry condition was much larger than that under the humid condition, whereas the study did not focus on the RH effect on m-xylene SOA formation. These demonstrate that the RH effects on aromatics SOA yields, especially m-xylene, have not been fully understood and the RH effects are controversial under various NOₓ levels and seed particle conditions.

Chemical components of SOA are important, on which climate- and health-relevant properties of particles are dependent. Chemical compositions of SOA from aromatics-NOₓ photooxidation have been investigated by GC/MS analysis (Forstner et al., 1997). Nevertheless, this study was only performed at 15-25% RH and high temperature at GC injection ports can easily decompose some low-volatile substances in SOA. FTIR was also used to study chemical compositions of SOA from aromatics-NOₓ photooxidation under different RH conditions, in which the information of functional groups in SOA was...
provided (Jia and Xu, 2014, 2018). The recent study on SOA components from toluene-OH system under both dry and humid conditions were analysed via HRMS (Hinks et al., 2018). Although the information of chemical compositions in SOA has been given, the analysis and the mechanism of RH effects still need to be further studied.

The reaction of \textit{m}-xylene-OH system forms peroxy radicals (RO\textsubscript{2}) that can rapidly react with HO\textsubscript{2} to form RO (R2a) and ROOH (R2b). RO finally forms carbonylic products, such as (methyl) glyoxal and other SOA precursors (Jenkin et al., 2003; Hallquist et al., 2009; Carlton et al., 2010; Carter and Heo, 2013). In most recent studies, RO\textsubscript{2} autoxidation has been observed that leads to the formation of higher-functionalized RO\textsubscript{2} radicals and the further formation of highly oxygenated molecules (HOMs) (Wang et al., 2017; Molteni et al., 2018). Self- or cross-reactions of RO\textsubscript{2} radicals were also found, leading to the formation of accretion products (R1c), especially HOM dimers, as well as RO, ROH and R\textsubscript{2}H\textsubscript{2}O (R1a and b) (Berndt et al., 2018). However, the RH effects on the mechanism of \textit{m}-xylene SOA formation are not well studied.

\begin{align*}
\text{ROO}^- + \text{R}'\text{OO}^- & \rightarrow \text{RO}^- + \text{R}'\text{O}^- + \text{O}_2 \quad \text{(R1a)} \\
& \rightarrow \text{ROH} + \text{R}'\text{H} + \text{O}_2 \quad \text{(R1b)} \\
& \rightarrow \text{ROOR}^- + \text{O}_2 \quad \text{(R1c)} \\
\text{ROO}^- + \text{HOO}^- & \rightarrow \text{RO}^- + \text{HO}^- + \text{O}_2 \quad \text{(R2a)} \\
& \rightarrow \text{ROOH} + \text{O}_2 \quad \text{(R2b)} 
\end{align*}

RH effects on SOA formation from \textit{m}-xylene under low NO\textsubscript{x} condition have not been studied well. In the present study, we present the results from the experiments about the SOA formation from the OH-initiated oxidation of \textit{m}-xylene in the absence of seed particles in a smog chamber. Both the SOA yields and chemical components under both low and high RH conditions will be reported. The underlying mechanism of SOA formation for these different conditions will be also discussed.

### 2 Experimental materials and methods

#### 2.1 Equipment and reagents

Experiments of \textit{m}-xylene photooxidation were performed in a 1 m\textsuperscript{3} Teflon FEP film reactor (DuPont 500A, USA) in an indoor smog chamber, which is similar to our previous works (Jia and Xu, 2014, 2016, 2018; Ge et al., 2016, 2017c, a, b). So only a brief introduction is presented here. A light source was provided by 96 lamps (F40BLB, GE; UVA-340, Q-Lab, USA) inside the chamber to simulate the UV band of solar spectrum in the troposphere. The NO\textsubscript{2} photolysis rate was determined to be 0.23 min\textsuperscript{-1}, which was used to reflect the light intensity in the reactor. To remove the electric charge on the surface of the FEP reactor, two ionizing air blowers were equipped in the chamber and were used throughout each experiment.

The background gas was zero air, which was generated from Zero Air Supply and CO Reactor (Model 111 and 1150, Thermo Scientific, USA) and further purified by hydrocarbon traps (BHT-4, Agilent, USA). The desired RH in the reactor was obtained by bubbling dry zero air through ultrapure water (Milli Q, 18MU, Millipore Ltd., USA). The RH and temperature in the reactor were measured by a hygrometer (Model 645, Testo AG, Germany).
Throughout each experiment, the background NO\textsubscript{x} concentration in the reactor was lower than 1 ppb and OH radicals were provided from H\textsubscript{2}O\textsubscript{2} photolysis. Hydrogen peroxide was introduced to the reactor by evaporating a measured volume of H\textsubscript{2}O\textsubscript{2} solution (30 wt %) into a zero air stream over a period of 30 min to the desired concentration of 20 ppm. Though the H\textsubscript{2}O\textsubscript{2} level was not measured, it was estimated through the measured volume of H\textsubscript{2}O\textsubscript{2} solution evaporated. m-Xylene (99%, Alfa Aesar) was introduced to the reactor subsequently using the same approach. No seed particles were introduced artificially. The experiments were conducted for 4 h.

2.2 Monitoring and analysis

The concentration of m-xylene in the reactor was measured with a gas chromatograph-mass spectrometer (GC-MS, Model 7890A GC and Model 5975C mass selective detector, Agilent, USA), which was equipped with a thermal desorber (Master TD, Dani, Italy). The size distribution and concentrations of particles were monitored with a scanning mobility particle sizer (SMPS, Model 3936, TSI, USA). The particle wall loss constant has been determined to be $3.0 \times 10^{-5}$ s\textsuperscript{-1} at low RH and $6.0 \times 10^{-5}$ s\textsuperscript{-1} at high RH conditions. The particles measured by SMPS consisted of liquid water content (LWC) and SOA at certain RH conditions. To determine the SOA concentration, LWC was determined at the end of each high RH experiment, which was based on the method in a previous study (Jia and Xu, 2018). Thus, here a brief introduction is only given. After the lights were turned off in high RH experiments, the SMPS was modified to the dry mode through adding a Nafion dryer (Perma Pure MD-700-12F-3) to the sampling flow and a Nafion dryer (Perma Pure PD-200T-24MPS) to the sheath flow, leading to the reduction of RH in the sample air to 10 % and that in the sheath to 7 %. After modifying to the dry mode, the humid air in SMPS was quickly replaced by dry air through venting the sheath air at 5 L min\textsuperscript{-1}, and then the dry aerosol was measured by SMPS. The LWC was determined by the difference of the particle mass concentrations before and after the modification of the dry mode.

For the analysis of functional groups of the chemical composition in SOA from m-xylene-OH irradiation, the SOA samples were collected and determined by FTIR (Fourier transform infrared spectrometer). The particles were collected on a ZnSe disk using a Dekati low-pressure impactor (DLPI, Dekati Ltd., Finland) at the end of each experiment (Ge et al., 2016; Jia and Xu, 2016). Then, the ZnSe disk was put in a FTIR (Nicolet iS10, Thermos Fisher, USA) for the measurement of functional groups of the chemical composition in SOA samples.

To obtain the detailed information of chemical composition, SOA particles were sampled using the Particle into Liquid Sampler (PILS, model 4001, BMI, USA). The flow rate of sample gas was around 11 L min\textsuperscript{-1}, and the output flow rate of liquid sample was 0.05 mL min\textsuperscript{-1}. Two denuders were used to remove the VOCs and acids in the sample gas. SOA liquid samples collected by PILS were finally transferred into vials for subsequent analysis. Operatively, the blank measurements were obtained by replacing the sample gas with zero air collected in vials. It is well known that the PILS samples watersoluble species in the SOA with high efficiency. In addition, it is reported that the PILS can also samples slightly watersoluble organic compounds with average O:C ratios higher than 0.26 instead of the total SOA composition and the collection
efficiency could exceed 0.6 (Zhang et al., 2016). Thus, the PILS can sample the overwhelming majority of the SOA system in our study, though PILS cannot sample water-insoluble species in the SOA.

The accurate mass of organic compounds in SOA and their MS/MS fragmentations were measured by the ultrahigh performance liquid chromatograph (UPLC, Ultimate 3000, Thermo Scientific, USA)-heated-electrospray ionization-high-resolution orbitrap mass spectrometer (HESI-HRMS, Q Exactive, Thermo Scientific, USA). Methanol (Optima™ LC/MS Grade, Fisher Chemical, USA) was used as the eluent in UPLC system. The elution flow rate was 0.2 mL min⁻¹, and the overall run time was 5 minutes. The injection volume was 20 μL. The acquired mass spectrum of SOA was in the range of 80-1000 Da. The HESI source was conducted in both positive and negative ion modes using the optimum method for characterization of organic compounds. We used the Thermo Scientific Xcalibur software (Thermo Fisher Scientific Inc., USA) to analyse the data from HRMS. To calculate the elemental compositions of compounds, the accurate mass measurements were used. For further analysis of the data from the second stage of data-dependent mass spectrometry (ddMS²), the Mass Frontier program (Version 7.0, Thermo Fisher Scientific Inc., USA) was used in order to simulate breaking the ions into fragments for comparison with the measured fragments to assist in identifying the structures. The reaction pathways and products of m-xylene-OH photooxidation in Master Chemical Mechanism (MCM v3.3.1, the website at http://mcm.leeds.ac.uk/MCM; last accessed October 16, 2017) was used for analysis of the products measured by HRMS (Jenkin et al., 2003; Jia and Xu, 2014).

3. Results and discussion

3.1 RH effects on SOA yields

Figure 1 shows the wall-loss-corrected particle mass concentration as a function of photooxidation reaction time for m-xylene-OH systems under both the low-RH and the high-RH conditions. It can be clearly seen that there is a large difference in the maximum mass concentration between low and high RHs. The maximum mass concentrations fitted are 150.3 and 95.5 μg m⁻³ at low RHs, whereas they are 21.0 and 7.5 μg m⁻³ at high RHs, with the largest difference being over ten times. The RH effect was reproducible when the initial m-xylene concentration was changed under similar conditions. To obtain the particle mass concentrations and SOA yield, an SOA density of 1.4 g cm⁻³ was used (Song et al., 2007). It should be noted that the fairly large scatter in the mass concentrations of SOA in Fig. 1 was observed, which mainly results from the uncertainty of SOA measurement by SMPS instrument. The interval of SOA data sampled by SMPS was 5 minutes, for which the sampling frequency was relatively low. SMPS measurement uncertainty is mainly dominated by size-dependent aerosol charging efficiency uncertainties and CPC sampling flow rate variability. The size-dependent aerosol charging efficiency is typically characterized by an accuracy of ± 10% (Jiang et al., 2014). The combination of various uncertainties, including SMPS measurement, sampling and even conversion of mass concentration from number concentration leads to the fairly large scatter in Fig. 1.
We used the definition of the ratio of the SOA mass to the consumed \( m \)-xylene mass to calculate the SOA yield at the end of each experiment. Under high RH condition, LWC accounts for a large proportion of particles (Jia and Xu, 2018). So, when SOA yield is calculated at high RH, the LWC has to be subtracted. The removal of aerosol water during the LWC measurement may cause the dissolved species that are probably volatile/semi-volatile compounds to evaporate back into the gas phase. Thus, SOA concentrations for high RH conditions were slightly underestimated, but the underestimation is extremely low and can be negligible.

Experimental conditions and SOA concentrations at the end of the experiments in \( m \)-xylene-OH oxidation system are summarized in Table 1. The SOA yields at low RH are 14.0-14.6\%, while those at high RH are only around 0.8-2.2\%. Both mass concentrations and SOA yields at low RH are an order of magnitude larger than those at high RH. It should be noticed that temperatures at high RH are slightly higher than those at low RH. Though low temperature can lead to a high SOA yield, the difference of temperature between low and high RH conditions in this study is lower than two degree, which cannot lead to a significantly different SOA yields to affect the result (Qi et al., 2010).

In the most recent study on toluene SOA formation conducted without seed particles (Hinks et al., 2018), the SOA yield at low NO\(_x\) level was 15\% under dry conditions (< 2\% RH) and 1.9\% under humid conditions (89\% RH), with the ratio of two yields between dry and humid conditions being over 7.5. The toluene SOA produced under high RH conditions were significantly suppressed, in which the tendency of RH effects on SOA yield was very similar with our study, though the difference of SOA yield between low and high RH conditions in Hinks et al (2018) was slightly smaller than that in this study. The small difference of RH effects between Hinks et al. and our study is likely associated with the difference in experimental conditions, including RHs and initial VOCs and H\(_2\)O\(_2\) concentrations, in addition to different species. This comparison demonstrates that different species of toluene and \( m \)-xylene of aromatics pose very similar RH effects under low-NO\(_x\) conditions. Hinks et al. attributed the suppression of SOA yields by elevated RH to the lower level of oligomers generated by condensation reactions and the reduced mass loading at high RH. In a study on an SOA model for toluene oxidation, it could be found that the SOA yield at low NO\(_x\) level was 28-30\% under low RH conditions (17-18\% RH) and 20-25\% under moderate RH conditions (48\% RH) (Cao and Jang, 2010), but they did not focus on the RH effect to give an explanation. As the difference of RH was only ~30\%, the RH effect on SOA yields was not as significant as those in Hinks et al and this study. Ng et al. have investigated the yields of SOA formed from \( m \)-xylene-OH system at low RH (4-6\%) under low NO\(_x\) conditions (Ng et al., 2007). They obtained that the SOA yields were in the range of 35.2-40.4\% in the presence of seed particles. The SOA yields were larger than that of this study, as they conducted the experiments under different irradiation time and with inorganic seed particles. These seed particles can provide not only surface for chemical reactions, but also acidic and aqueous environments that can promote the SOA formation (Jang et al., 2002; Liu et al., 2018; Faust et al., 2017). In a study on chemical oxidative potential of SOA (Tuet et al., 2017), it was observed that the concentration of SOA from \( m \)-xylene irradiation at low NO\(_x\) level under dry condition was much larger than that under humid condition (89.3 \( \mu \)g m\(^{-3}\) at < 5\% RH and 13.9 \( \mu \)g m\(^{-3}\) at 45\% RH), but they did not calculate the \( m \)-xylene SOA yields or give an explanation for the RH effect. However, a previous study under high NO\(_x\) conditions (Zhou et al., 2011) found that the effect of RH on
SOA yields from aromatics photooxidation were positive, which obtained opposite results to our study. LWC was used to account for this positive effect as it could promote SOA formation through aqueous chemistry (Jia and Xu, 2014; Wang et al., 2016; Faust et al., 2017; Liu et al., 2018).

It should be noted that seed aerosols were not artificially introduced throughout all the experiments, which could lead to the underestimation of SOA, as SOA-forming vapours partly condense to the chamber walls instead of particles (Matsunaga and Ziemann, 2010; Zhang et al., 2014). The extent to which vapor wall deposition affects SOA mass yields depends on the specific parent hydrocarbon system (Zhang et al., 2014; Zhang et al., 2015; Nah et al., 2016; Nah et al., 2017). Zhang et al (2014) have estimated two m-xylene systems under low NO\textsubscript{x} conditions and concluded that SOA mass yields were underestimated by factors of 1.8 (Ng et al., 2007) and 1.6 (Loza et al., 2012) under low RH conditions. In addition, the excess use of H\textsubscript{2}O\textsubscript{2} can lead to an excess OH radicals, leading to a less underestimation of SOA formation as the losses of SOA-forming vapours can be mitigated via the use of excess oxidant concentrations (Nah et al., 2016). Thus, the underestimation of SOA formation can be limited. In fact, the wall loss of m-xylene was not taken into consideration of calculation of mass yields, which generally overestimates the mass yields.

The wall loss of chemical species that is sensitive to humidity may affect the RH effect on SOA yields, as the reduction of SOA yields at the high humidity may be due to the chemical loss to the wet chamber wall. To estimate the extent of how much the wall loss of chemical species affects the SOA formation at different RHs, we take glyoxal and acetone as reference compounds. Glyoxal, a typical compound that can form SOA, can easily dissolve in the aqueous phase due to the large Henry’s law constant of $4.19 \times 10^5$ M atm\textsuperscript{-1} (Ip et al., 2009), very sensitive to humidity. Loza et al. (2010) found that the wall loss of glyoxal was minimal at 5% RH, with $k_W = 9.6 \times 10^{-7}$ s\textsuperscript{-1}, whereas $k_W$ was $4.7 \times 10^{-5}$ s\textsuperscript{-1} at 61% RH. We assume that $k_W$ linearly increases with RH, and the $k_W$ value is estimated to be $6.1 \times 10^{-5}$ s\textsuperscript{-1} at 80% and $7.4 \times 10^{-6}$ at 13% RH, with the difference being 8.2 times. According to the wall loss of glyoxal, glyoxal only decreased by 10% at the end of our experiment at low RH, while glyoxal decreased by 59% at high RH. Acetone can hardly dissolve in the aqueous phase due to the small Henry’s law constant of 29 M atm\textsuperscript{-1} (Poulain et al., 2010), which is 4 orders of magnitude less than that of glyoxal. Ge et al. (2017) obtained that the wall loss of acetone was $5.0 \times 10^{-6}$ s\textsuperscript{-1} at 87% RH and $3.3 \times 10^{-6}$ s\textsuperscript{-1} at 5% RH, with a factor of 1.5. The difference of wall loss between glyoxal and acetone at low RH is about 2 times, while it becomes about 12 times at high RH. Thus, it can be considered that the wall loss among different species at low RH is less affected by the Henry’s law constant, but it is greatly affected at high RH. In our study glycolaldehyde (See the Sec. 3.3) is found to be an important SOA precursor that can form a large fraction of oligomers in our experiments, but the wall loss of glycolaldehyde is not available. The Henry’s law constant of glycolaldehyde was obtained to be $4.14 \times 10^4$ M atm\textsuperscript{-1} (Betterton and Hoffmann, 1988), an order of magnitude lower than glyoxal, indicating that glycolaldehyde is less sensitive to humidity than glyoxal but much more sensitive to humidity than acetone. Based on the data of these two reference species, the wall loss of glycolaldehyde at low RH is taken to be $5 \times 10^{-6}$ s\textsuperscript{-1}, and the difference in wall loss between high and low RHs is about 6 times. Then, the wall loss of glycolaldehyde at high RH can be $3 \times 10^{-5}$ s\textsuperscript{-1}. Then, it is estimated that glycolaldehyde would decrease by 7% at low RH and by 35% at high RH at the end of our experiment, respectively. This means that SOA yield
would be underestimated by 35% at high RH and by 7% at low RH if glycolaldehyde lost to the wall was completely transformed to SOA. If this wall effect of SOA precursors was taken into consideration, the SOA yields at high (Exp. 3) and low (Exp. 2) RHs would be 3.4% and 15.1%, respectively. Alternatively, the SOA yield at high RH was underestimated to be 42% relative to that at low RH. Even the sensitivity of the wall loss to RH was taken to be 8 times, the SOA yield at high RH would be underestimated to be 62% compared to that at low RH. In fact, there were many different SOA precursors from the \textit{m}-xylene oxidation system that probably have much smaller Henry’s law constant relative to that of glycolaldehyde. Thus, it is concluded that the RH effect on SOA formation from \textit{m}-xylene oxidation by H\textsubscript{2}O\textsubscript{2} without NO\textsubscript{x} is negative.

### 3.2 RH effects on functional groups of SOA

Figure 2 shows the FTIR spectra of particles from the photooxidation of \textit{m}-xylene-OH experiments under both low (Exp. 2) and high (Exp. 3) RH conditions. The DLPI sample flow rate was 10 L min\textsuperscript{-1}, and the sampling duration was 15 min. We used same sampling flow rate and duration for both RH conditions. DLPI has 13 stages, and it can collect particles in the size range of 30 nm - 10 mm. When we sampled using DLPI, the four plates for stages 4-7 were removed, so that particles in the range of 108-650 nm were collected on the third plate. As shown in Fig. S2 in the supplementary information, the particles in the range of 108-650 nm can represent the total SOA from \textit{m}-xylene oxidation in this study. The mean collection efficiency of the DLPI was 83\% for stages 4-7 (Durand et al., 2014). Thus, the SOA mass collected on the ZnSe window was 10.3 and 3.0 µg at low RH (Exp. 2) and high RH (Exp. 3), based on the SMPS measurement and the DLPI collection efficiency. As shown in Fig. 2, the SOA from \textit{m}-xylene-OH experiments can be obviously observed under both two RH conditions. The intensities of all functional groups from the low RH experiment are much higher than those from the high RH experiment, which is consistent with the reduced SOA yields under elevated RH conditions.

The assignment and the intensity of the FTIR absorption frequencies is summarized in Table 2. The broad absorption at 3600-2400 cm\textsuperscript{-1} is O-H stretching vibration in phenol, hydroxyl and carboxyl groups. The sharp absorption at 1720 cm\textsuperscript{-1} is the C=O stretching vibration in carboxylic acids, formate esters, aldehydes and ketones. The absorptions at 1605 cm\textsuperscript{-1} match the bending vibration of liquid water. The absorptions at 1415 cm\textsuperscript{-1} match the deformation of CO-H, phenolic O-H and C-O. The absorptions at 1180 cm\textsuperscript{-1} match the C-O-C stretching of polymers, C-O and OH of COOH groups. The absorptions at 1080 cm\textsuperscript{-1} match the C-C-OH stretching of alcohols.

The absorption intensity at ~3200 cm\textsuperscript{-1} that is identified as the hydroxyl group is used to be a representative for reflection of the SOA formation. As well, Table 2 gives the ratio of intensities at high RH to those at low RH to compare the difference of relative intensities of functional groups. The intensities of functional groups are obviously suppressed at high RH, but the extents of the suppression for different functional groups are basically divided into two types. The ratios of O-H, C-H, C=O and C-C-OH groups are 0.29 to 0.34, which is close to the ratio of SOA mass at high RH to that at low RH collected on the ZnSe disk, whereas the ratios of CO-H, C-O-C, C-O-H in COOH are above 0.48. The relative intensity of the C-O-C group is significantly higher than the C=O group, which can be explained by more oligomerization with the formation of C-O-C than other reactions at high RH. Nevertheless, the FTIR results cannot provide further information to well explain the
the differences of SOA yields between low and high RH, which will be further discussed in terms of mass spectra of SOA in the next section.

### 3.3 RH effects on mass spectra of SOA

We selected the sample mass spectra whose intensities are larger than $10^5$ under the low RH condition and corresponding mass spectra under the high RH condition, followed by the blank mass spectra deduction. The blank-deducting mass spectra of SOA formed from $m$-xylene-OH photooxidation under low and high RH conditions in both positive and negative ion modes are presented in Fig. 3, which is plotted as a function of the mass-to-charge ratio. It should be noted that the Y-axis scales for low and high RH are largely different, $10^6$ at low RH and $10^5$ at high RH. As shown in Fig. 3, a visible decrease in the overall peak intensities for both positive and negative ion modes can be obviously observed as the RH elevates, which is consistent with the result that the SOA mass concentration is lower at high RH. In addition, it is obvious that the number of peaks is less under the high RH condition. As shown in Fig. 3, where the m/z values of SOA samples are close for both low and high conditions, the absolute and relative intensities of the peaks are much different, indicating that RH significantly affects the concentration of SOA components.

Table 3 lists the peaks whose intensities are larger than $10^6$ of low RH samples and the structure can be proposed according to the gas-phase chemical mechanism of $m$-xylene-OH photooxidation included in MCM and the fragments from MS/MS analysed with Mass Frontier. In the positive ion mode, an [M+H]$^+$ ion of m/z = 137.05962 at low RH and 137.05931 at high RH is assigned as a molecular ion formula of C$_8$H$_9$O$_2$ that has a mass difference of $\Delta = 0.6$ and 1.0 mDa for low and high RH, respectively. The structure of identified compound C$_8$H$_8$O$_2$ is proposed to be 2,6-dimethyl-1,4-benzoquinone, the fragments of which from MS/MS match those from simulation of the Mass Frontier program. This compound was also identified and quantified in a previous study on SOA compositions from $m$-xylene-NO$_x$ irradiation using the method of GC-MS analysis with authentic standards (Forstner et al., 1997). Thus, 2,6-dimethyl-1,4-benzoquinone was the SOA component partitioning into particle phase from the gas phase. The measured ion of m/z = 155.07013 at low RH and 155.06985 at high RH is assigned as a molecular ion formula of C$_8$H$_{11}$O$_3$ that has $\Delta = 1.2$ and 1.5 mDa, and its structure is proposed to be O=CC1(C)OC1C=CC(=O)C, an oxidized unsaturated epoxide. The measured ion of m/z = 171.06509 at low RH and 171.06488 at high RH is assigned as a molecular ion formula of C$_8$H$_{11}$O$_4$ that has $\Delta = 1.2$ and 1.4 mDa, the structure of which is proposed to be a bicyclic peroxide. The measured ion of m/z = 187.06003 at low RH and 187.05678 at high RH is assigned as a molecular ion formula of C$_8$H$_{11}$O$_5$ that has $\Delta = 1.2$ and 4.4 mDa, whose structure is proposed to be O=CC1(C)OC1C(=O)C(=O)C. All these SOA components are suppressed to almost disappear at high RH, except for 2,6-dimethyl-1,4-benzoquinone.

For rough quantification of the RH effect, the peaks in Figure 3 were assigned with the number of carbon atoms. The intensities of the peaks with the same number of carbon atoms (nC) are summed, which are presented in Figure 4. It should be noted that the Y-axis scales at low and high RHs are largely different, with a label step of $4.0 \times 10^6$ at low RH and $4.0 \times 10^5$ at high RH in the positive ion mode, $5.0 \times 10^6$ at low RH and $1.0 \times 10^5$ at high RH in the negative ion mode. The
compounds with nC > 8, larger number of carbon atoms than m-xylene, are proposed to be oligomers that account for a large mass fraction of SOA due to their large molecular weights and lower volatilities, though their peak intensities are lower. As a result, the processes for formation of such compounds play an important role in the formation of SOA. It can be obviously observed that the peak intensities are much lower at high RH in the negative ion mode than that in the positive mode, indicating that the decrease of the compounds obtained in the negative ion mode account for a larger decrease at high RH. Moreover, in the positive ion mode, compounds with nC > 8, especially those with nC > 10, account for more SOA mass at high RH than at low RH, which can be observed by the increase of relative intensities of oligomers. However, in the negative ion mode, compounds with nC > 8 account for less SOA mass at high RH. In the positive ion mode, 15 compounds with nC = 10 identified with HRMS dominate the spectrum. To get more information, the MS/MS data analysis was further carried out. We found that the fragment with m/z = 61.03 obtained by the MS/MS was observed in nine out of the fifteen compounds, which is assigned as a molecular ion formula of C$_2$H$_5$O$_2^+$. The sum of peak intensities of these nine compounds accounts for about 70% of the sum of peak intensities of these fifteen nC = 10 compounds. The structure of this fragment is possibly proposed to be glycolaldehyde (C$_2$H$_4$O$_2$), which has been observed previously in the oxidation of m-xylene (Cocker et al., 2001). It has been implied that the oligomerization of glycolaldehyde took place in the study of formation of toluene SOA (Hinks et al., 2018). This demonstrates that the nC = 10 compounds are mainly from the oligomerization of glycolaldehyde and nC = 8 monomers.

### 3.4 Proposed mechanism of RH effects on SOA formation

It has been confirmed that glyoxal is an important SOA precursor via oligomerization (Jang et al., 2002; Kamens et al., 2011; Zhou et al., 2011). An explanation for the large difference of SOA yields and composition between low and high RH is proposed that water is directly involved in the chemical mechanism and further affects the SOA growth. However, in some particle-phase accretion equilibrium reactions, water is involved as a by-product. The elevated RH alters the equilibrium of reaction toward the decrease of products (Nguyen et al., 2011; Hinks et al., 2018). The hydrolysis of oligomers would be facilitated by the elevated RH after the oligomers are generated in gas phase and partition into particles with certain LWC. In this study and the previous study on toluene SOA formation, C$_2$H$_5$O was one of the most frequently mass difference at both RH, but the peak intensities of its relative compounds were much lower under elevated RH conditions (Hinks et al., 2018). C$_2$H$_5$O was proposed to be from the oligomerization reaction of glycolaldehyde (C$_3$H$_5$O$_2$), which can react with carbonyl compounds by aldol condensation reactions with water as the by-product. However, this mechanism cannot well explain the large different SOA yields at low and high RH.

Another possible explanation for RH effects on SOA chemical components is that RH influences the HOMs formation from m-xylene-OH system (Fig. 5). Reactions between m-xylene (C$_8$H$_{10}$) and OH radicals have two pathways, the H-abstraction from the methyl group and OH-addition to the aromatic ring, which generates products such as methylbenzaldehyde (C$_8$H$_8$O) and methylbenzyl alcohol (C$_8$H$_{10}$O), as shown in Scheme 1. OH-addition is the dominant pathway, as the branching ratio of H-abstraction only accounts for 4% based on the MCM simulation. OH-addition to the aromatic ring is followed by O$_2$-
adduct and isomerization to form a carbon-centered radical, which can form a dimethylphenol (C₉H₁₀O) or is adducted by an O₂ molecule forming a bicyclic peroxyl radical (BPR, C₈H₁₁O₅) (Calvert et al., 2002; Birdsall et al., 2010; Wu et al., 2014), which is called the RO₂ autoxidation. The BPR reacts with other RO₂ radicals or HO₂ forming the bicyclic oxy radical (C₈H₁₁O₄). This RO radical can get further reaction and finally form carboxylic products (R1a and R2a), such as (methyl)glyoxal and other SOA precursors (Jenkin et al., 2003; Hallquist et al., 2009; Carlton et al., 2010; Carter and Heo, 2013), or react with HO₂ radicals forming bicyclic hydroxyhydroperoxides (ROOH, C₈H₁₂O₅) (R2b), or react with other RO₂ radicals forming ROH (C₈H₁₂O₄) and R₄O (C₈H₁₀O₄) (R1b). The self- and cross-reactions of RO₂ radicals also form ROOR (C₁₆H₂₂O₁₀) or ROOR’ that is the accretion products (Berndt et al., 2018; Molteni et al., 2018). The further autoxidation of BPR can form a highly-functionalized RO₂ radicals and further get reacted and finally form HOMs (Types 1 and 2 in Scheme 1) (Wang et al., 2017; Crounse et al., 2013; Ehn et al., 2014; Jokinen et al., 2015; Berndt et al., 2016). Dimethylphenol (C₈H₁₀O) as well as other products from termination reaction with benzene ring or double bond can react with OH radicals and get further reacted to form HOMs as well. The compounds with the number below their formulae in scheme 1 were determined by HRMS to be present in the particle phase.

The distribution of relative intensity of SOA products with same carbon number was shown in Fig. 5 to investigate the potential RH effect on HOMs. Oxidation compounds with the same carbon number but different oxygen number were observed. In the mass-to-charge (m/z) range 135-240 Da in both positive and negative ion modes, the oxidation products contain the carbon skeleton of m-xylene (monomer region, nC = 8), while in the m/z range 320-410 Da the number of carbon atoms is doubled (dimer region, nC = 16). The compounds with high O/C ratios are observed, which are generally called HOMs. In a previous study on m-xylene oxidation, HOM monomers including C₅H₁₀O₅, C₈H₁₂O₅ and C₈H₁₄O₆, and HOM dimers including C₁₆H₂₂O₉ and C₁₆H₂₄O₁₁ were obtained using CI-API-TOF-MS (chemical ionization atmospheric pressure interface time of flight mass spectrometer) (Molteni et al., 2018), which are also determined in our study. More HOMs can be obtained in the negative ion mode where oxygen atom numbers can reach up to 8 for monomers and 12 for dimers, when compared with those in the positive ion mode. In addition, here the O/C ratio was calculated based on the O and C atom numbers in structures and peak intensities, which is 0.57 and 0.71 in the positive and negative ion modes, respectively. This also demonstrates that the compounds in the negative ion mode are much more oxygenated than those in the positive ion mode. As shown in Fig. 4, the peak intensities at high RH are much lower in the negative ion mode than in the positive mode, indicating that the decrease of the more oxygenated compounds account for the larger fraction at high RH. These high O/C ratios cannot be explained by any of the formerly known oxidation pathways except that RO₂ autoxidation is taken into consideration (Crounse et al., 2013; Barsanti et al., 2017). Both HOM monomers and dimers obviously decrease with increasing RH. RH significantly affect the formation of the HOMs. The relative intensities of compounds with same nC but larger oxygen number decrease more significantly. High RH significantly suppresses the more oxidized compounds whose volatility is lower and molecular weights are larger. Therefore, it is considered that RH suppresses the RO₂ autoxidation and further affects the SOA yield.
The wall process of the reactor enlarges the difference of SOA mass between low and high RH. The wall loss of some chemical species is faster at high RH, which leads to the reduction of SOA yield. In addition, the difference of SOA mass can be also enhanced based on the gas to particle partitioning rule (Li et al., 2018).

4. Conclusion and atmospheric implication

The current study investigates the effect of RH on SOA formation from the oxidation of \(m\)-xylene under low \(\text{NO}_x\) conditions in the absence of seed particles. The elevated RH can significantly obstruct the SOA formation from the \(m\)-xylene-OH system, so that the SOA yield decrease from 13.8% at low RH to 0.8% at high RH, with a significant discrepancy of higher than one order of magnitude. The FTIR results of functional groups show the relative increase of the C-O-C group at high RH as compared with low RH, indicating that the oligomers from carbonyl compounds cannot well explain the suppression of SOA yield. HOMs were observed to be suppressed in the HRMS spectra. The chemical mechanism for explaining the obvious difference of RH effects on SOA formation from \(m\)-xylene-OH system has been proposed based on the analysis of both FTIR and HRMS measurements as well as MCM simulations. The reduced SOA at high RH is mainly ascribed to the less formation of oligomers and the suppression of \(\text{RO}_2\) autoxidation. Together with the previous study on toluene SOA, it is conceivable that the effect of RH on SOA yield is a common feature of SOA formation from aromatics oxidation under low \(\text{NO}_x\) conditions and using \(\text{H}_2\text{O}_2\) as the OH radical source. Our results obviously indicate that the production of SOA from aromatics in low-\(\text{NO}_x\) environments can be strongly modulated by the ambient RH probably due to the influence of \(\text{H}_2\text{O}\) on the formation of HOMs and oligomers. Our study highlights the role of water in the SOA formation, which is particularly related to chemical mechanisms used to explain observed air quality and to predict chemistry in air quality models and climate models. The clear pathway of the influence of \(\text{H}_2\text{O}\) on the formation of HOMs needs to be further studied in the future.

Author contribution

Qun Zhang and Yongfu Xu designed the research. Qun Zhang carried out the experiments and analyzed the data. Long Jia provided valuable advices on the experiment operations. Yongfu Xu and Long Jia provided advices on the analysis of results. Qun Zhang prepared the manuscript with contributions from all co-authors.

Acknowledgments

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References


Figure 1. SOA mass concentrations as a function of irradiation time (corrected by particle wall loss).

Figure 2. FTIR spectra of particles from photooxidation of $m$-xylene-OH experiments under low and high RH conditions.
Figure 3. Selected background-subtraction HESI-Q Exactive-Orbitrap MS results of SOA in both positive and negative ion modes from the photooxidation of \textit{m}-xylene-OH under both low and high RH conditions (Note that the Y-axis scales for low and high RH are largely different, $10^6$ at low RH and $10^5$ at high RH).
Figure 4. Sum of peak intensities based on peaks selected in Figure 3 as a function of the number of carbon atoms under the positive ion mode and negative ion mode (Note that the Y-axis scale at low and high RH are largely different, with a label step of $4.0 \times 10^6$ at low RH and $4.0 \times 10^5$ at high RH in the positive ion mode, $5.0 \times 10^6$ at low RH and $1.0 \times 10^5$ at high RH in the negative ion mode).
Figure 5. Mass spectra of SOA from m-xylene at both low (red) and high (blue) RH in the positive (+) and negative (-) ion modes, grouped with the same number of carbon atoms (from nC =8 to 16). On (n = 2, 3, ……, 12) means the number of oxygen atoms in the formula of the peak.
Scheme 1. The route of OH-initiated m-xylene oxidation. The red number below the molecular formula is its molecular weight, which is determined by HRMS to exist in the particle phase.
Table 1. Experimental conditions, SOA concentrations and yields at the end of the experiments in \textit{m}-xylene-OH oxidation system.

<table>
<thead>
<tr>
<th>Exp. No.</th>
<th>([m\text{-xylene}]_0) (μg m(^{-3}))</th>
<th>([m\text{-xylene}]_\text{reacted}) (μg m(^{-3}))</th>
<th>RH (%)</th>
<th>T (°C)</th>
<th>([\text{SOA}]_e) (μg m(^{-3}))</th>
<th>SOA yield (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2287.9</td>
<td>1026.3</td>
<td>13.6</td>
<td>25.9</td>
<td>150.3 ± 15.0</td>
<td>14.6 ± 1.5</td>
</tr>
<tr>
<td>2</td>
<td>1855.5</td>
<td>682.0</td>
<td>13.7</td>
<td>25.3</td>
<td>95.5 ± 9.5</td>
<td>14.0 ± 1.4</td>
</tr>
<tr>
<td>3</td>
<td>2410.8</td>
<td>941.4</td>
<td>73.6</td>
<td>27.5</td>
<td>21.0 ± 2.1</td>
<td>2.2 ± 0.2</td>
</tr>
<tr>
<td>4</td>
<td>2029.1</td>
<td>946.9</td>
<td>79.1</td>
<td>27.4</td>
<td>7.5 ± 0.7</td>
<td>0.8 ± 0.1</td>
</tr>
</tbody>
</table>

\([\text{SOA}]_e\) indicates the mass concentration of SOA at the end of each experiment with particle wall loss corrected.

Table 2. Absorbance positions of functional groups and the intensities at low and high RHs.

<table>
<thead>
<tr>
<th>Absorption frequencies</th>
<th>Functionality</th>
<th>Intensity ((\times 10^{-3}))</th>
<th>Ratio (^a)</th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>low RH</td>
<td>high RH</td>
</tr>
<tr>
<td>3235</td>
<td>O-H</td>
<td>5.9</td>
<td>1.9</td>
</tr>
<tr>
<td>3000</td>
<td>C-H</td>
<td>4.5</td>
<td>1.4</td>
</tr>
<tr>
<td>1720</td>
<td>C=O</td>
<td>5.1</td>
<td>1.5</td>
</tr>
<tr>
<td>1415</td>
<td>CO-H</td>
<td>4.8</td>
<td>2.4</td>
</tr>
<tr>
<td>1180</td>
<td>C-O-C, C-O and OH of COOH</td>
<td>2.9</td>
<td>1.4</td>
</tr>
<tr>
<td>1080</td>
<td>C-C-OH</td>
<td>5.3</td>
<td>1.8</td>
</tr>
</tbody>
</table>

\(^a\) Ratio of the intensity at high RH to that at low RH.
Table 3. Plausibility of different types of compounds with elemental formulae measured by HRMS in the positive ion mode.

<table>
<thead>
<tr>
<th>Ion formula</th>
<th>Proposed structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>[C₈H₉O₂]⁺</td>
<td><img src="image1.png" alt="Proposed structure" /></td>
</tr>
<tr>
<td>[C₇H₉O₃]⁺</td>
<td><img src="image2.png" alt="Proposed structure" /></td>
</tr>
<tr>
<td>[C₈H₁₁O₄]⁺</td>
<td><img src="image3.png" alt="Proposed structure" /></td>
</tr>
<tr>
<td>[C₈H₁₁O₅]⁺</td>
<td><img src="image4.png" alt="Proposed structure" /></td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Measured (m/z)</th>
<th>Intensity</th>
<th>Error (mDa)</th>
<th>Measured (m/z)</th>
<th>Intensity</th>
<th>Error (mDa)</th>
<th>Ion formula</th>
<th>Proposed structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>137.0596</td>
<td>1.7 × 10⁶</td>
<td>0.6</td>
<td>137.0593</td>
<td>1.4 × 10⁵</td>
<td>1</td>
<td>[C₈H₉O₂]⁺</td>
<td><img src="image1.png" alt="Proposed structure" /></td>
</tr>
<tr>
<td>141.0545</td>
<td>5.6 × 10⁶</td>
<td>1.3</td>
<td>141.0542</td>
<td>-</td>
<td>1.5</td>
<td>[C₇H₉O₃]⁺</td>
<td><img src="image2.png" alt="Proposed structure" /></td>
</tr>
<tr>
<td>155.0701</td>
<td>1.0 × 10⁶</td>
<td>1.2</td>
<td>155.0699</td>
<td>-</td>
<td>1.5</td>
<td>[C₈H₁₁O₃]⁺</td>
<td><img src="image3.png" alt="Proposed structure" /></td>
</tr>
<tr>
<td>171.0651</td>
<td>1.0 × 10⁶</td>
<td>1.2</td>
<td>171.0649</td>
<td>-</td>
<td>1.4</td>
<td>[C₈H₁₁O₄]⁺</td>
<td><img src="image4.png" alt="Proposed structure" /></td>
</tr>
<tr>
<td>187.06</td>
<td>1.1 × 10⁶</td>
<td>1.2</td>
<td>187.0568</td>
<td>-</td>
<td>4.4</td>
<td>[C₈H₁₁O₅]⁺</td>
<td><img src="image5.png" alt="Proposed structure" /></td>
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