1	Synergetic effects of NH <sub>3</sub> and NO <sub>X</sub> on the production
2	and optical absorption of secondary organic aerosol
3	formation from toluene photooxidation
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### 22 Abstract

NH<sub>3</sub> is the most important alkaline gas in the atmosphere and one of the key 23 species affecting the behaviors of atmospheric aerosols. However, the impact of NH<sub>3</sub> 24 on secondary organic aerosol (SOA) formation remains poorly understood, especially 25 the dynamic evolution of chemical compositions in the SOA formation process. In this 26 study, a series of chamber experiments were performed to probe the individual and 27 common effects of NH3 and NOX on toluene SOA formation through OH-28 photooxidation. The chemical compositions of toluene SOA were characterized using 29 30 the Aerodyne high-resolution time-of-flight aerosol mass spectrometer (AMS). The SOA yield increased from 28.1% in the absence of NH<sub>3</sub> to 34.7% in the presence of 31 NH<sub>3</sub> but decreased to 19.5% in the presence of NO<sub>X</sub>. However, the highest SOA yield 32 of 42.7% and the lowest carbon oxidation state (OS<sub>C</sub>) occurred in the presence of both 33 NH<sub>3</sub> and NO<sub>X</sub>, indicating that the higher volatility products that formed in the presence 34 of NO<sub>X</sub> could partition into the particle-phase when NH<sub>3</sub> was added. This resulted in a 35 36 synergetic effect on SOA formation when NH<sub>3</sub> and NO<sub>X</sub> co-existed. The heterogeneous reaction was the main pathway by which NH<sub>3</sub> participated in SOA formation in the 37 photooxidation process. The synergetic effect of NH<sub>3</sub> and NO<sub>X</sub> was also observed in 38 SOA optical absorption. A peak at 280 nm, which is characteristic of organonitrogen 39 imidazole compounds, was observed in the presence of NH<sub>3</sub> and its intensity increased 40 when NO<sub>X</sub> was added into the chamber. This work improves our understanding of how 41 42 the synergistic interactions between NH<sub>3</sub> and NO<sub>X</sub> influence SOA formation and offers new insights into mitigating haze pollution. 43

- 44 Keywords: Photooxidation; Toluene; NH<sub>3</sub>; Dynamic characteristics; Synergistic
- 45 effects

## 47 **1 Introduction**

Secondary organic aerosols (SOA) are an important component of atmospheric 48 particulate matter (Moise et al., 2015;Liu et al., 2017), and can significantly affect 49 atmospheric visibility, air quality, and human health (Paciga et al., 2014; Yang et al., 50 2016; Liu et al., 2017). Optical properties of SOA have been directly and indirectly 51 linked to their effects on the climate (Laskin et al., 2015;Xie et al., 2017;Peng et al., 52 2020). Because of the complexity of their chemical components, oxidation processes, 53 and environmental factors, SOA formation mechanisms are very complex and the 54 55 current understanding of SOA formation is incomplete. This limited understanding hampers the ability of models to predict the magnitudes, dynamics, and distributions of 56 atmospheric aerosols from particulate and precursor emissions (Ortiz-Montalvo et al., 57 2014). In the past decades, although our understanding of SOA formation mechanisms 58 has been constantly improving, there is still a gap between the simulated SOA 59 concentration in large-scale atmospheric models and field observations (Volkamer et 60 61 al., 2006; Yang et al., 2018).

Ammonia (NH<sub>3</sub>) is the most abundant and ubiquitous alkaline inorganic gas in the atmosphere and one of the critical factors influencing SOA formation (Wang et al., 2016;Wang et al., 2018b;Chen et al., 2019). Some studies have noted that the presence of NH<sub>3</sub> can contribute to the formation of more aerosol mass through photooxidation (Na et al., 2007;Li et al., 2018). Na et al. (2007) observed that aerosol yields in the  $\alpha$ pinene-ozone oxidation system increased by 8% when NH<sub>3</sub> was added. Li et al. (2018) reported that the presence of NH<sub>3</sub> in the aromatic hydrocarbon photooxidation system

69	increased aerosol size growth potentials (by 7%-108%), and resulted in enhanced SOA
70	formation. Qi et al. (2020) found that the concentration and average diameter of SOA
71	showed an immediate and rapid increase after adding NH <sub>3</sub> . Furthermore, the acid-base
72	reactions between $NH_3/NH_4^+$ and the carboxyl groups in SOA molecules might enhance
73	SOA formation (Qi et al., 2020;Liu et al., 2015). The condensable ammonium salts
74	formed from the reaction between NH3 and organic acids reduce the volatility of the
75	organic acids by several orders of magnitude (Paciga et al., 2014), and act as particle-
76	phase organics that further promote SOA formation (Na et al., 2007;Huang et al.,
77	2012;Chen et al., 2019;Qi et al., 2020;Wu et al., 2020). In addition, carbonyls can
78	undergo nucleophilic attack by NH3 through the Maillard reactions and form the
79	corresponding iminium intermediates (Noziere et al., 2009;Laskin et al., 2015;Liu et
80	al., 2015). The iminium intermediates can continue to react with carbonyls, which
81	activates further transformations such as the formation of heterocyclic compounds and
82	oligomerization reactions and forms condensation (oligomeric) products with more
83	stable secondary imines (Schiff bases) (Laskin et al., 2014). Both Noziere et al. (2009)
84	and Ortiz-Montalvo et al. (2014) reported NH3 is an efficient catalyst for reactions with
85	carbonyl compounds to form nitrogen-containing organic aerosols (NOA). The reaction
86	between carbonyl and NH3 can significantly decrease the volatility of oxidation
87	products, which further increases the yield of SOA (Lee et al., 2013;Zhang et al.,
88	2015a;Qi et al., 2020). Babar et al. (2017) found that the substantial formation of
89	secondary imines in the presence of $NH_3$ was responsible for the higher $\alpha$ -pinene SOA
90	yields. However, not all studies have shown that the presence of NH <sub>3</sub> increases SOA

91 yields. One study observed that NH<sub>3</sub> suppressed SOA formation under certain
92 ozonation conditions (Ma et al., 2018b). Furthermore, the consumption of NH<sub>3</sub> by
93 Criegee intermediates was reported to decrease the secondary ozonide yield and thus
94 affect SOA formation.

Nitrogen oxides (NO<sub>X</sub> = NO + NO<sub>2</sub>), which are mainly emitted from the 95 combustion of fossil-fuels, have received significant attention due to their effects on the 96 photooxidation process of volatile organic compounds (VOCs) and SOA formation 97 (Surratt et al., 2006;Ng et al., 2007b;Draper et al., 2015;Berkemeier et al., 98 99 2016;Sarrafzadeh et al., 2016;Zhao et al., 2018). A clear increase at first and then a 100 decrease in the SOA yield was found with increasing NO<sub>X</sub> concentration from the laboratory experiments with both anthropogenic (trimethylbenzene) and biogenic (β-101 pinene) VOCs (Sarrafzadeh et al., 2016; Yang et al., 2020). The competitive chemistry 102 of organic peroxyl radicals (RO<sub>2</sub>) with hydroperoxyl radicals (HO<sub>2</sub>) and NO was 103 responsible for the variability in SOA formation (Ng et al., 2007a;Xu et al., 2014;Jiang 104 105 et al., 2020). RO<sub>2</sub> mainly reacts with HO<sub>2</sub> under low-NO<sub>X</sub> conditions to form oxidation products with lower volatility, which may enable it to partition into the particle-phase 106 and contribute to the SOA mass (Ng et al., 2007a). While the  $RO_2 + NO$  reaction is 107 dominant in high-NO<sub>X</sub> conditions, the increase in volatile products formed through 108 fragmentation was responsible for the decrease in SOA yield with increasing NO<sub>X</sub> 109 (Zhao et al., 2018;Liu et al., 2019a;Xu et al., 2020). In addition, the increase of OH 110 concentration formed through NO + HO<sub>2</sub>  $\rightarrow$  NO<sub>2</sub> + OH reaction at low-NO<sub>X</sub> conditions, 111 and a suppressing effect of NO<sub>X</sub> on OH formation under high-NO<sub>X</sub> conditions was 112

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partly responsible for the first increasing and then decreasing trend of SOA yield with NO<sub>X</sub> concentration (Sarrafzadeh et al., 2016;Bates et al., 2021).

In the last decade, atmospheric pollutants in China have changed significantly in 115 their concentrations and composition (Wang et al., 2015;Xia et al., 2016) with the 116 emissions of SO<sub>2</sub> and NO<sub>X</sub> decreased by 75% from 2007–2015 and 10% from 2011– 117 2015, respectively (de Foy et al., 2016; Vu et al., 2019; Wang et al., 2020). However, 118 owing to the lack of regulation regarding NH<sub>3</sub> emissions, NH<sub>3</sub> emissions increased by 119 ~30% from 2008–2016 over the North China Plain (Liu et al., 2018). As has been 120 121 pointed out in previous research, the promoting effect of NH<sub>3</sub> on the formation of SOA may counteract the decreases in aerosol formation due to reductions in SO<sub>2</sub> and NO<sub>X</sub> 122 (Zhang et al., 2021a). Indeed, field observation and model simulation have pointed out 123 that the simultaneous control of NH<sub>3</sub> emissions in conjunction with SO<sub>2</sub> emission is 124 more effective in reducing PM<sub>2.5</sub> than the process without NH<sub>3</sub> emissions control, and 125 PM<sub>2.5</sub> concentration can be more effectively reduced if NH<sub>3</sub> emission is decreased as 126 127 much as that of SO<sub>2</sub> (Erisman and Schaap, 2004). Hence, the mechanism by which NH<sub>3</sub> affects SOA formation has attracted more and more attention (Wang et al., 2018a;Ge et 128 al., 2019;Zhang et al., 2021b). However, previous studies have not paid sufficient 129 attention to the joint impacts of NH<sub>3</sub> and NO<sub>X</sub> on the formation of SOA and its 130 corresponding optical properties. Due to the lack of real time detection methods for 131 SOA chemical composition, the dynamic characteristics of how NH<sub>3</sub> participates in 132 SOA formation via photooxidation have not been extensively studied. 133

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Toluene is one of the most abundant aromatic VOCs in the urban atmosphere,

which is also an important source of brown carbon (Laskin et al., 2010;Ma et al., 2018a). 135 The effects of NH<sub>3</sub> and NO<sub>X</sub> on SOA formation through the toluene photooxidation 136 process were investigated in this study. The chemical composition of toluene SOA was 137 characterized on-line with an aerosol mass spectrometer and the characteristics of SOA 138 chemical composition under different conditions were further explored by applying a 139 positive matrix factorization (PMF) analysis. The optical properties of toluene SOA 140 particles were determined based on a UV-vis spectrum analysis. Possible mechanisms 141 of the effects of both NH<sub>3</sub> and NO<sub>X</sub> on SOA formation were discussed. The results will 142 143 help us to better understand SOA formation mechanisms under complex pollution conditions with elevated NH<sub>3</sub> and NO<sub>X</sub> concentrations in an urban atmospheric 144 environment. 145

146 **2 Materials and Methods** 

#### 147 **2.1 Photooxidation chamber experiments**

All toluene photooxidation experiments were performed in a 4 m<sup>3</sup> chamber. The chamber has been described in detail in our previous study (Liu et al., 2021). Briefly, the chamber was constructed with a 0.08 mm-thick FEP-Teflon film. 50 UV-B lamps (TUV36W, Philips) with peak wavelengths of 254 nm were set up around the chamber and used as the light source to drive OH radical formation through hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>) photolysis. Mirror surfaced stainless steel was used as the interior wall of the enclosure to maximize and homogenize the interior light intensity. All experiments were performed at room temperature (293~298 K) and atmospheric pressure was
maintained in the chamber at all times.

Before each experiment, the chamber was flushed with zero air for at least 18 hours, 157 after which the concentration of particles was less than 1 cm<sup>-3</sup>. Zero air was generated 158 by a zero air supply (111-D3N, Thermo Scientific<sup>TM</sup>, USA). The flow rate of zero air 159 was controlled at 20 L min<sup>-1</sup> by a mass flow controller (D088C/ZM, Beijing Sevenstar 160 Electron Corporation) during the process of chamber reaction bag inflating. The relative 161 humidity (RH) of zero air was about 20%. For each experiment, measured amounts of 162 163 toluene (Sigma-Aldrich, analytically pure) and H<sub>2</sub>O<sub>2</sub> solution (Sigma-Aldrich, 30 wt% in H<sub>2</sub>O) were injected into a Teflon bulb with micro syringes. Zero air was passed 164 through the injection tube to make sure all the liquids had evaporated to the gas-phase 165 and were blown into the chamber. Toluene concentration was measured with a Proton 166 Transfer Reaction-Mass Spectrometry (PTR-ToF-MS, Ionicon Analytik, Austria). The 167 evolution of toluene concentration for different experiments was shown in Fig. S1. The 168 169 OH concentration in the chamber was calculated based on the first order decay of toluene concentration. There was no obvious difference of OH concentrations in the 170 171 different NO<sub>X</sub> and NH<sub>3</sub> levels (Fig. S2). The average OH concentration over the entire reaction period was  $5.87 \times 10^7$  molecule cm<sup>-3</sup>. NO<sub>X</sub> (Air Liquid Shanghai, 510 ppm) 172 NO<sub>2</sub> in N<sub>2</sub>) and/or NH<sub>3</sub> (Air Liquid Shanghai, 502 ppm NH<sub>3</sub> in N<sub>2</sub>) were introduced 173 directly into the chamber to reach the required concentrations. For experiments with 174 NO<sub>X</sub>, although only NO<sub>2</sub> was introduced into the chamber before photooxidation, NO 175 could be formed through NO<sub>2</sub> photolysis under the UV light irradiation, so NO always 176

177 coexisted with NO<sub>2</sub> in the photooxidation system (Zhao et al., 2018). NOx 178 concentration was measured online by the NO-NO<sub>2</sub>-NOx analyzer (Model 42C, 179 Thermo Electron Corporation, USA). Each experiment was performed without seed 180 aerosols. After all the reactants were added, the chamber stood quietly for 10 min 181 without turning on the light to ensure that the reactant gases in the chamber were evenly 182 mixed. The photooxidation started when the UV light was turn on.

The experimental conditions for the toluene photooxidation are listed in Table 1. 183 In our work, the OH and toluene concentrations were higher than those of urban 184 185 conditions. The purpose of the high OH and toluene concentrations is to obtain enough particle production samples for off-line collections and accurate measurements. The 186 toluene concentrations remained stable under the different experimental conditions, the 187 variations of toluene-derived SOA mass concentration and yield were only affected by 188 the different NO<sub>2</sub> and/or NH<sub>3</sub> concentrations in this study. Toluene was studied here as 189 the representative of total aromatic VOCs in the urban atmosphere, thus its 190 191 concentration used in this study was much higher than that in the real atmosphere. However, the concentration ratio of toluene to OH in this study is similar to that under 192 193 the real atmospheric conditions (Prinn et al., 1995; Zou et al., 2015).

### **2.2 Particle concentration measurements**

For each experiment, a scanning mobility particle sizer (SMPS) was used to record the particle size distribution and volume concentration of the toluene-derived SOA. The SMPS was composed of a differential mobility analyzer (DMA model 3081, TSI Inc.,

USA) and a condensation particle counter (CPC model 3776, TSI Inc., USA) which 198 were used for screening particles with specific electrical mobility diameter (from 14.1 199 200 nm to 736.5 nm) and for counting the number of the selected particles, respectively. The sheath gas velocity was 3 L min<sup>-1</sup> and the sample gas velocity was 0.3 L min<sup>-1</sup>. The 201 202 size scan was repeated every 5 min. During each scan circle, the scan time was 240 s, and the particle sizes ranged from 13.6 nm to 726.5 nm. A density of 1.4 g m<sup>-3</sup>, which 203 was measured by Ng et al. (2007), was used for the calculation of toluene SOA mass 204 concentration from the particle volume concentration (Ng et al., 2007b). 205

206 **2.3 Chemical characterization** 

In this study, the toluene SOA chemical compositions were characterized with an 207 on-line high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS, 208 Aerodyne Research Inc. USA). The sample flow passed through a Nafion dryer and the 209 RH of the sample gas was reduced to below 20% before entering the AMS. In the 210 injection port, an aerodynamic lens focused particles with a vacuum aerodynamic 211 diameter below 1 µm into a narrow beam. Particles impacted a flash vaporizer (600°C) 212 at the rear of the sizing region under high vacuum ( $\sim 10^{-7}$  Torr) and were subsequently 213 ionized by electron impact ionization (70 eV). Then, the positively charged ions entered 214 the ToF section and were separated. V-mode (m/ $\Delta m = \sim 2000$ ) was used in the AMS 215 ToF section to achieve the high signal-to-noise ratio. The separated ion fragments were 216 analyzed by a mass spectrometer with scans from 1 to 300 m/z. The composition-217 dependent collection efficiency (CE) was applied to the data based on the methods 218

established by Middlebrook et al. (2012). For mass concentration calculations, 1.1, 1.2, 219 and 1.4 were applied as the default relative ionization efficiency (RIE) values of nitrate, 220 221 sulfate, and organic compounds, respectively. The standard AMS data analysis software SQUIRREL 1.63B coupled with PIKA 1.23B in the Igor Pro (WaveMetrics, Inc., 222 223 Portland, Oregon), which were retrieved from http://cires1.colorado.edu/jimenezgroup/ToFAMSResources/ToFSoftware/, were used for the analysis of elemental ratios 224 and the ion speciated compositions of toluene SOA in the chamber. Note that the 225 elemental ratios (i.e., O/C, H/C, and N/C) and mass-to-carbon ratio (OM/OC) were all 226 227 calculated using the Aiken-Ambient method for comparability with previous studies (Aiken et al., 2008). In order to further explore the changes in SOA chemical 228 composition, a PMF of the high-resolution mass spectra was performed to determine 229 230 the different organic aerosol (OA) factors during the toluene photooxidation process. We preformed the PMF analysis in the same way as Zhang et al. (2011), the details of 231 which are provided in the Supporting Information. 232

### 233 **2.4 Absorption measurements**

The changes of absorption spectra and the absorbance of the toluene derived SOA under different conditions were determined using a UV spectrophotometer (UV-3600, Shimadzu, Japan) with a 1 cm cuvette. The SOA was collected from a 3 m<sup>3</sup> sample gas onto the 46.2 mm PTFE filter (Whatman<sup>TM</sup>, UK). The collected SOA sample was dissolved in 5 mL of methanol (HPLC grade, > 99.8%) with 30 min of sonication. As reported by Chen and Bond (2010), > 92 % of SOA is extractable by organic solvents 240 (e.g., methanol), which means that almost all organic matter was extracted in this study. 241 The filter extracts were filtered through 0.2  $\mu$ m PTFE syringe filters to remove 242 suspended insoluble particles. Before detection of the optical absorbance, a cuvette 243 filled with pure methanol was scanned as a blank to provide a spectral background. The 244 absorption was detected over the range of 200 to 800 nm with a resolution of 0.5 nm<sup>-1</sup>. 245 The light absorption coefficient of the particles at a specific wavelength  $\lambda$  (Abs $_{\lambda}$ , M/m) 246 was calculated according to Eq. R1:

$$Abs_{\lambda} = (A_{\lambda} - A_{700}) \cdot \frac{V_1}{V_a \cdot L} \cdot \ln(10)$$
(R1)

where,  $A_{700}$  is the background value of light absorption intensity, calculated as the average value of light absorption intensity from 695–705 nm to reduce the limits of error in measurement; V<sub>1</sub> and V<sub>a</sub> are the volumes of methanol with dissolved particles and sampled air, respectively; and L is the optical path length. Because Abs<sub> $\lambda$ </sub> was strongly dependent on the amount of SOA, all Abs<sub> $\lambda$ </sub> results were normalized based on the SOA mass collected on the filter. The normalized result was defined as the mass absorption coefficient (MAC, m<sup>2</sup> g<sup>-1</sup>), calculated using Eq. R2:

$$MAC_{\lambda} = \frac{Abs_{\lambda}}{M}$$
(R2)

where, M ( $\mu$ g m<sup>-3</sup>) represents the concentration of methanol-soluble organic carbon.

# 255 **3 Result and Discussion**

### 256 **3.1 SOA formation**

In order to investigate the effect of  $NH_3$  and  $NO_X$  on SOA formation from toluene photooxidation, a control test was carried out. SOA yield (Y) is defined as  $Y = \frac{12}{12}$ 

259	$\Delta M_0/\Delta HC$ , where $\Delta M_0$ is the produced organic aerosol mass concentration (µg m <sup>-3</sup> ),
260	and $\Delta HC$ is the mass concentration of reacted toluene (µg m <sup>-3</sup> ). The evolution of SOA
261	mass concentrations and SOA yield at different conditions during the photooxidation
262	process were shown in Fig. 1. Recent experiments showed that the wall loss of organic
263	vapors to the Teflon walls should not be ignored (Zhang et al., 2014;Zhang et al.,
264	2015b), and represented a major challenge in investigating SOA formation with
265	environmental chambers (Krechmer et al., 2020). The formation of SOA in laboratory
266	chambers may be substantially suppressed due to losses of SOA-forming vapors to
267	chamber walls, but this effect on SOA formation have not yet been quantitatively
268	established. However, the particle wall loss rates were detected at the end of the
269	chamber experiment after the UV-lamps were turned off, and the mass concentration
270	was corrected with the same way of Jiang et al. (2020) and Pathak et al. (2007). After
271	the wall loss correction, the particle mass concentration was almost constant, the
272	different wall loss effect caused by gaseous oxidation products formed in the different
273	experiment conditions was considered remedied. Interestingly, wall loss is increased
274	66% and 205% in Exp.2 (in the presence of NH3) and Exp.3 (in the mixed condition of
275	$NH_3$ and $NO_X$ ), respectively, when compared with the experiments with no $NH_3$ (Exp.1
276	and 4). The larger particle wall loss in the presence of NH <sub>3</sub> could be explained by
277	increasing condensation process of oxidized organic vapors onto the Teflon chamber
278	wall via oligomerization (for dicarbonyls) and ionic dissociation/acid-base reaction (for
279	organic acids). There was a noticeable increase in the SOA mass concentration in the
280	presence of NH <sub>3</sub> , which was consistent with previous studies (Qi et al., 2020;Chu et al.,

2016). The mass concentration of SOA increased from  $637 \pm 14.6 \ \mu g \ m^{-3}$  without NH<sub>3</sub> 281 to a maximum of 867  $\pm$  12.7  $\mu g~m^{\text{-3}}$  with 200 ppb NH3. Previous studies attributed the 282 283 enhancement of SOA to the formation of NOA from acid-base reactions between NH<sub>3</sub>/NH<sub>4</sub><sup>+</sup> and carboxyl groups, or Maillard reactions of NH<sub>3</sub>/NH<sub>4</sub><sup>+</sup> with carbonyl 284 functional groups (Noziere et al., 2009;Ortiz-Montalvo et al., 2014;Liu et al., 2015;Qi 285 et al., 2020). In contrast, SOA concentrations were lower in the presence of NO<sub>X</sub>, and 286 the maximum mass concentration of toluene SOA was only  $452 \pm 18.9 \ \mu g \ m^{-3}$  with 63 287 ppb initial NO<sub>X</sub>. The branching of RO<sub>2</sub> loss among different pathways has an important 288 289 influence on the product distribution and SOA formation. The fate of RO<sub>2</sub> mainly depends on the concentrations of NO<sub>X</sub> (Zhao et al., 2018;Liu et al., 2019a;Xu et al., 290 2020). Several studies have shown that, instead of RO<sub>2</sub> reacting with RO<sub>2</sub>/HO<sub>2</sub>, NO 291 292 would react with RO2 to form the RO intermediate and produces more oxidation products with higher volatilities through fragmentation in the presence of NO<sub>X</sub> (Zhao 293 et al., 2018;Liu et al., 2019a;Xu et al., 2020). Highly volatile compounds cannot readily 294 295 partition into the particle-phase, so this substantially suppresses the formation of SOA.

The NO<sub>X</sub> and NH<sub>3</sub> had opposite effects on toluene SOA formation in this study. Interestingly, however, the highest SOA mass concentration  $(1020 \pm 10.6 \ \mu g \ m^{-3})$ occurred in the presence of both NO<sub>X</sub> and NH<sub>3</sub>, which was nearly 1.6 times higher than that observed in the absence of NO<sub>X</sub> or NH<sub>3</sub>. Although inorganic aerosol (i.e., NH<sub>4</sub>NO<sub>3</sub>) was formed from the interaction of NH<sub>3</sub> and NO<sub>X</sub> in the chamber, the upper limit of the inorganic matter only accounted for 6.6% of the total mass of particulate matter (Table S1) in the NH<sub>3</sub> + NO<sub>X</sub> experiment. Therefore, NH<sub>4</sub>NO<sub>3</sub> was not the main cause of the

increase in particulate matter; the co-occurrence of NH<sub>3</sub> and NO<sub>X</sub> had a synergistic 303 effect on the SOA formation, because their combined effect on SOA formation was 304 305 greater than the sum of their separate effects. Qi et al. (2020) found that the promotion of NH<sub>3</sub> on toluene SOA formation was more obviously under high NO<sub>X</sub> concentration, 306 SOA yield increased 3.7% and 4.6% for 70 ppb and 160 ppb initial NO<sub>X</sub> concentration, 307 respectively, when 200 ppb NH<sub>3</sub> was added into the chamber. Li et al. (2018) showed 308 that the presence NH<sub>3</sub> can promote the particle size growth of SOA; at the same time, 309 this particle growth rate was higher under low VOC/NO<sub>X</sub> (or high NO<sub>X</sub>) conditions. All 310 311 in all, the joint effect of multiple environmental factors on SOA formation is not the simple summation of the influences of various factors on SOA formation. This may at 312 least partly explain why predictions of SOA concentrations in large-scale atmospheric 313 314 models, which typically describe SOA formation from data derived from chamber experiments, are frequently lower than field observations (Volkamer et al., 2006). 315

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#### 3.2 SOA chemical composition

317 The traditional SOA formation mechanism is based on the chemical compositions obtained through off-line detection of the chemical composition of SOA (Jang et al., 318 2002;Liu et al., 2019a;Xu et al., 2020). SOA is continually evolving in the atmosphere 319 and the ageing process of SOA co-occurs with its formation process, resulting in the 320 transformation of SOA chemical composition continuously proceeding during the 321 photooxidation process, but little attention has been paid to the evolution of SOA 322 chemical composition in previous studies. Therefore, the AMS was used for on-line 323

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measurement of the SOA chemical composition and how the chemical composition evolved in the photooxidation process will be discussed in the following section.

Chemical composition of SOA is very complex. The average carbon oxidation 326 state (OS<sub>C</sub>) has been shown to be an ideal conceptual framework to describe changes 327 in the degree of oxidation undergone by SOA (Kroll et al., 2011), and has been widely 328 applied in field and laboratory studies (Chen et al., 2018; Mandariya et al., 2019). 329 Average OS<sub>C</sub> calculation was shown in Supporting Information. Fig. 2 shows the 330 changes in the OS<sub>C</sub> of toluene SOA formed in different experiments. Notably, toluene 331 332  $SOA OS_C$  values was in the range between -0.5 and 0, which is consistent with that of semi-volatile oxygenated organic aerosols (SV-OOA). However, the different OS<sub>C</sub> 333 values and the change trends observed for the toluene SOA formed in different 334 conditions (with and without NH<sub>3</sub>/NO<sub>X</sub>) in Fig. 2 indicated that there was a 335 photooxidation mechanism active during SOA formation, which ultimately changed the 336 SOA chemical compositions. 337

The OS<sub>C</sub> increased over time for all SOAs that were formed in the absence of NH<sub>3</sub>. 338 There are several possible reasons for the increasing trend of OS<sub>C</sub> values. Firstly, a 339 dynamic equilibrium of semi-volatile vapors may have been achieved between the 340 particle-phase and gas-phase during the earlier toluene oxidation process. The increase 341 of SOA led to a reduction in the concentration of gas-phase semi-volatile organic 342 products. A decreasing concentration of gas-phase semi-volatile organic compound 343 products would suppress their transformation from gas-phase to particulate-phase. 344 More lower volatility gas-phase oxidation products with higher OS<sub>C</sub> values would then 345

be shifted to the particle phase, which would be responsible for the continuing increase 346 of SOA and its OSc. Secondly, the formed SOA could have further been oxidized by 347 348 OH radical through heterogeneous reactions (Kourtchev et al., 2015;Liu et al., 2019b). This could be the main reason for the increase in the OS<sub>C</sub> when the SOA concentration 349 350 was no longer increasing. Finally, as pointed by Malecha and Nizkorodov (2016), even if there was no OH radical in the chamber, the photodegradation of SOA can produce 351 small oxygenated volatile organic compounds (e.g. acetaldehyde OS<sub>C</sub>=-1, and acetone 352  $OS_C \approx -1.3$ ) under UV light irradiation. The photoproduction of OVOCs from SOA had 353 a lower OS<sub>C</sub> value than that of SOA. Although the loss of SOA through 354 photodegradation is small, the OS<sub>C</sub> value of SOA still had increased to a certain extent 355 (Malecha and Nizkorodov, 2016). 356

The fact that additional photochemical processing results in the dynamic evolution 357 of the OS<sub>C</sub> over time has been demonstrated in both field and laboratory experiments 358 (Jimenez et al., 2009). The atmospheric oxidation of OA tends towards higher  $OS_C$ 359 360 regardless of the original OA source (Herndon et al., 2008). However, when NH<sub>3</sub> was present, the  $OS_{C}$  of total SOA was almost unchanged for the whole photooxidation 361 362 period. Carboxyl and carbonyl are the main oxygen-containing functional groups responsible for the toluene photooxidation production (Ji et al., 2017). An organic 363 ammonium salt with four H atoms can offset an increase in OS<sub>C</sub> value caused by the 364 formation of organic acids/carboxy group with two O atoms through acid-base reactions 365 (Kuwata and Martin, 2012;Liu et al., 2015). In addition, NH<sub>3</sub>/NH<sub>4</sub><sup>+</sup> may react with 366 carbonyl functional groups through Maillard reactions, consuming the oxygen atom in 367

368	the carbonyl group and leading to the formation of species with covalent carbon-
369	nitrogen bonds (Lee et al., 2013;Zhang et al., 2015a;Qi et al., 2020). Xu et al. (2018)
370	showed that imidazole compounds (OSc $\approx$ -1.3) generated through heterogeneous
371	reaction between NH <sub>3</sub> and carbonyl compounds might contribute to the decrease in the
372	$OS_C$ of SOA. It is clear that an increase in $OS_C$ caused by the formation of oxygen-
373	containing functional groups (e.g., carboxyl, carbonyl, etc.) would be counteracted
374	through acid-base reactions or Maillard reactions in the presence of NH <sub>3</sub> . After 60 min
375	of UV light irradiation, there was no more SOA formation; however, the $OS_{\rm C}\xspace$ did
376	decrease slightly in Exp.2 and 3, illustrating that the NH <sub>3</sub> could continue to react with
377	SOA through heterogeneous processes. Huang et al. (2016) also pointed out that the
378	portion of semi-volatile products with low OS <sub>C</sub> formed at the later stage of
379	photooxidation also contributed to the decreased OS <sub>C</sub> .

The OS<sub>C</sub> of the toluene SOA formed with NO<sub>X</sub> was lower than that formed in the 380 absence of NO<sub>X</sub>, no matter whether NH<sub>3</sub> was present in the chamber or not. This 381 382 indicated that an increased NO<sub>X</sub> concentration benefits the formation of high volatility oxidation products with lower OS<sub>C</sub> values (Kroll et al., 2011; Jimenez et al., 2009). 383 384 However, the relationships between OS<sub>C</sub> and SOA mass concentration with and without 385 NH<sub>3</sub> were the opposite to each other. Predictably, the SOA formation mechanism in the presence of NO<sub>X</sub> is different from that with NO<sub>X</sub> + NH<sub>3</sub>. In the absence of NH<sub>3</sub>, the RO 386 intermediate, which is easily fragmented to produce relatively high-volatility 387 compounds, was the dominant product of the NO<sub>X</sub> + RO<sub>2</sub> reaction (Zhao et al., 2018;Liu 388 et al., 2019a;Xu et al., 2020). Highly volatile compounds cannot readily partition into 389

the particle-phase, which subsequently results in a lower SOA yield in the presence of 390 NO<sub>X</sub> (Yang et al., 2020). Thereby, both OS<sub>C</sub> and the SOA mass concentration were 391 392 lower when 60 ppb NO<sub>X</sub> was added into the chamber. However, when both NO<sub>X</sub> and NH<sub>3</sub> were present, the toluene derived SOA had the lowest OS<sub>C</sub> value, but the highest 393 mass concentration. This result suggested that although NO<sub>X</sub> promotes the formation 394 of higher volatility compounds, these higher volatility compounds (e.g. glyoxal) can 395 react with NH<sub>3</sub> and partition into the particle-phase, which could contribute to the 396 increase in SOA formation. Huffman et al. (2009) observed that aerosol volatility was 397 398 inversely correlated with the extent of oxidation of OA components. The low value of OS<sub>C</sub> in the presence of NO<sub>X</sub> indicated that NO<sub>X</sub> would promote the formation of the 399 relatively high-volatility compounds. However, the lower OS<sub>C</sub> value in the presence of 400 401 NH<sub>3</sub> indicated that the high-volatility compounds would promote partitioning into the particle-phase when reacting with NH<sub>3</sub>. 402

Fragments derived from the AMS data have also been widely used to explore the 403 bulk compositions and properties of SOA (Ng et al., 2010;Ng et al., 2017). The m/z 43 404 (f43) frequency was dominated by ion  $C_2H_3O^+$ , which is the tracer for organic 405 406 compounds with alcohol and carbonyl functional groups (Alfarra et al., 2006). 407 Meanwhile, the m/z 44 (f44) signal was dominated by  $CO_2^+$  ions, which is the tracer for organic compounds with carboxyl functional groups and an indicator of highly 408 oxygenated organic aerosols (Ng et al., 2010). Here, we use the approach of Ng et al. 409 (2010) by plotting the fractions of the total organic signal at m/z 43 vs. m/z 44 (f43 vs. 410 f44). The change of f43 vs. f44, which has an inflection point during the photooxidation 411

process, is shown in Fig. 3 and Fig. S3. In our study, the change before the inflection point was defined as the formation stage, and the linear fit of f43 *vs.* f44 for the formation stage is shown by the dashed lines. The change in f43 *vs.* f44 after the inflection point was defined as the stable stage, and the linear fit of f43 *vs.* f44 in this stage is shown by the solid lines. The formation and stable stages of the f43 *vs.* f44

In the stable stage, the increase in f44 and decrease in f43 with increasing OH 418 exposure indicated that the carbonyl groups in toluene SOA were oxidized to carboxyl 419 420 groups by the ageing process. For the experiments without NH<sub>3</sub> and NO<sub>X</sub>, the slope 421 ratio of f43 vs. f44 was -3.9. When there was 60 ppb initial NO<sub>X</sub>, the f43 was almost stable while the f44 increased with the oxidation process. There was a lower slope ratio 422 of f43 vs. f44, indicating that organic compounds with more alcohol and carbonyl 423 functional groups had formed in the presence of NO<sub>X</sub>. But for the experiments with 200 424 ppb initial NH<sub>3</sub>, the slope ratios of f43 vs. f44 were only -1.1 and -1.3 in the presence 425 426 and absence of NO<sub>X</sub>, respectively. According to the above results, we can see that more carbonyl groups are consumed as carboxyl groups are formed in the presence of NH<sub>3</sub>. 427 428 The carbonyls can be oxidized to organic acids (Kawamura and Bikkina, 2016), but 429 unreacted carbonyl can be nucleophilically attacked by NH<sub>3</sub>/NH<sub>4</sub><sup>+</sup> to form nitrogenheterocyclic compounds, e.g., imidazole (Grace et al., 2019;Lian et al., 2020). 430 Meanwhile, the peak f44 value decreased from 0.13 to 0.10 when NH<sub>3</sub> was added into 431 the chamber. This suggested that the heterogeneous reaction of NH<sub>3</sub>/NH<sub>4</sub><sup>+</sup> could 432 promote the consumption of particle-phase carbonyl groups (Xu et al., 2018), and must 433

inhibit the formation of carboxyl groups in the SOA ageing process. According to the 434 changing trend of SOA concentration over time, the photooxidation process was 435 divided into formation stage and stable stage. As shown in Fig. 1, the first half hour of 436 photooxidation when SOA concentration increased linearly with time was defined as 437 SOA formation stage. After 60 min of photooxidation, SOA concentration was not 438 changed with reaction time, and it was defined as stable stage. The differences in spectra 439 of toluene SOA in the formation stage and stable stage are shown in Fig. 4. A lower 440 signal intensity variation of CO<sub>2</sub><sup>+</sup> in the presence of NH<sub>3</sub> also illustrated that NH<sub>3</sub> would 441 442 inhibit heterogeneous reactions that form carboxyl groups.

In the formation stage, the slope ratios of f43 vs. f44 were almost the same for both 443 experiments without NO<sub>X</sub>. It can thus be seen that the presence or absence of NH<sub>3</sub> does 444 not affect the change trend of f43 vs. f44 in the SOA formation stage. Therefore, the 445 gas-phase homogeneous reaction of NH<sub>3</sub> on SOA formation is not important. Clearly, 446 the particle-phase heterogeneous reaction was the main reaction pathway by which NH3 447 448 participated in the photooxidation process and toluene SOA formation. However, negative correlations were observed between f43 and f44 in the presence of NO<sub>X</sub>. Based 449 450 on this, we concluded that NO<sub>X</sub> not only affects the SOA formation through the particlephase heterogeneous reactions, but also through gas-phase homogeneous reactions. 451

#### 452 **3.3 PMF results**

453 A temporal evaluation of the toluene SOA chemical composition during 454 photooxidation is vital to the analysis of the NOA formation mechanism in the presence

455	of NH <sub>3</sub> and/or NO <sub>X</sub> . Therefore, this study further compared the chemical properties of
456	the SOA generated under different experimental conditions by applying a PMF analysis
457	to the HR-ToF-AMS data (Chen et al., 2019). A summary of the PMF results is
458	presented in Fig. S4-S7. For the toluene OH-photooxidation experiments with NH <sub>3</sub>
459	and/or NO <sub>X</sub> presence, two factors were identified from the PMF analysis in the same
460	way of Chen et al. (2019). The H/C, O/C, and N/C values of these two factors are shown
461	in Fig. 5. The factor with higher N/C values was defined as high-nitrogen OA (Hi-NOA)
462	In contrast, the factor with lower N/C values was defined as low-nitrogen OA (Lo-
463	NOA). Fig. 6 exhibits the evolution of Hi-NOA and Lo-NOA during the photooxidation
464	process as resolved from the PMF analysis of different initial NO <sub>X</sub> /NH <sub>3</sub> concentrations.
465	While similar evolutionary trends were observed under different conditions, the relative
466	intensities and the chemical compositions of these two factors in each experiment were
467	not consistent.

For the toluene SOA formed under NH<sub>3</sub> conditions, both Lo-NOA and Hi-NOA 468 had similar O/C values, which were fully oxygenated with an average of  $0.74 \pm 0.04$ 469 (Fig. 5a). These O/C values were comparable to the low-volatility oxygenated organic 470 aerosols (LV-OOA) with an O/C value ranging from 0.6 to 1 (Jimenez et al., 2009). The 471 472 main difference between these two OA sources was the N/C ratio. The N/C ratio of Hi-NOA (N/C = 0.032) was about three times higher than that of Lo-NOA (N/C = 0.010) 473 (Fig. 5a). The evolution of these two OA sources during the photooxidation process is 474 shown in Fig. 6a. The components of toluene SOA were mostly Lo-NOA during the 475 initial phase of SOA formation, but Hi-NOA toluene SOA started forming after 10 476

minutes and continued to increase. The Lo-NOA reached the maximum mass 477 concentration after 30 min of the photooxidation, and then decreased. Such a declining 478 trend of Lo-NOA at longer reaction times reflected the conversion of Lo-NOA into 479 something else in the particle-phase. As the Lo-NOA decreased, the mass concentration 480 of Hi-NOA gradually increased. Thus, the Hi-NOA should be derived from the 481 heterogeneous reaction of Lo-NOA with NH<sub>3</sub>/NH<sub>4</sub><sup>+</sup>. At the same time, it was proved 482 that the formation pathway of Hi-NOA was not through reaction of NH<sub>3</sub> with later-483 generation gas-phase products in the homogeneous gas phase. With the gradual 484 485 replacement of Lo-NOA by Hi-NOA, the ratio of [Hi-NOA]/[Lo-NOA] stabilized at 5~6. 486

For the toluene SOA formed under NO<sub>X</sub> conditions, there was not a large difference between the N/C ratios of Hi-NOA (N/C = 0.019) and Lo-NOA (N/C = 0.014) (Fig. 5c). At the end of the NO<sub>X</sub> experiment, the ratio of [Hi-NOA]/[Lo-NOA] was only 3:2 (Fig. 6c). It follows that the contribution of the heterogeneous NO<sub>X</sub> reaction to the N/C ratio of toluene SOA was not obvious. Therefore, the formation of NOA in the presence of NO<sub>X</sub> mainly occurred through gas-phase homogeneous reactions, which was consistent with the results in section 3.3.

The changing trend of N/C with time in the presence of  $NH_3$  was different to that with  $NO_X$  present. The evolutions of the N/C of SOA in different experiments are shown in Fig. 7. In the presence of  $NH_3$ , the N/C value gradually increased throughout the photooxidation process. The increased N/C value in the photooxidation process was attributed to the heterogeneous  $NH_3$  reaction with SOA. But in the presence of  $NO_X$ , the N/C increased rapidly to its maximum value where it was stable for the rest of the reaction. This could mean that the heterogeneous reaction of toluene SOA with NO<sub>X</sub> to form NOA was not as important as the gas-phase homogeneous reaction.

When both NO<sub>X</sub> and NH<sub>3</sub> were added into the chamber, the N/C ratios of Hi-NOA 502 and Lo-NOA were 0.062 and 0.029, respectively (Fig. 5b). The N/C ratio of Hi-NOA, 503 which was comparable to the recently isolated nitrogen-enriched OA value (0.053) 504 observed by Sun et al. (2011), was much higher than that observed in the experiments 505 with only  $NH_3$  or  $NO_X$ . It was even higher than the sum of the N/C ratios from both 506 507 Exp. 2 with  $NH_3$  and Exp. 4 with  $NO_X$ . In order to calculate the relative contributions of NH<sub>3</sub> and NO<sub>X</sub> to N/C, it was assumed that the effects of NH<sub>3</sub> or NO<sub>X</sub> on the N/C 508 ratio in the Hi-NOA and Lo-NOA factors did not change among different experimental 509 conditions. For Lo-NOA, the contributions of NH<sub>3</sub> and NO<sub>X</sub> to the N/C value were 510 0.0126 and 0.0164, and their relative intensities were 43% and 57%, respectively. While 511 for the Hi-NOA, the contributions of NH<sub>3</sub> and NO<sub>X</sub> to the N/C values were 0.0404 and 512 513 0.0216, and their relative intensities were 65% and 35%, respectively. For the experiment with both  $NH_3$  and  $NO_X$ , the contribution of  $NH_3$  to N/C was higher by 26%, 514 515 and the contribution of NO<sub>X</sub> to N/C was higher by 17% compared to the experiments with the single pollutants. The co-existence of NH<sub>3</sub> and NO<sub>X</sub> further enhanced the N/C 516 value of toluene SOA, indicating that a synergetic interaction between NH<sub>3</sub> and NO<sub>X</sub> 517 further enhanced organic nitrogen formation. 518

#### 519 **3.4 Optical absorption**

The optical characteristics of toluene SOA formed from different  $NH_3$  and  $NO_X$ conditions were investigated. The MAC of toluene derived SOA detected over the range of 230–600 nm is displayed in Fig. 8. Over the entire UV detection range, an increase in light absorption was observed when the toluene SOA formed in the presence of  $NO_X$ or  $NH_3$ .

By looking at Fig. 8 in detail, one can see that the MAC of toluene SOA formed 525 with (red line) and without (black line) NH<sub>3</sub> overlapped at 250 nm, but when the UV 526 wavelength exceeded 250 nm the MAC of the toluene SOA formed in the presence of 527 NH<sub>3</sub> was higher. The red line reflects an obvious characteristic absorption peak at 528 270~280 nm, which was mainly due to the absorption of the  $n \rightarrow \pi^*$  electronic 529 transitions. The imidazole compounds were formed through the Maillard reactions 530 between NH<sub>3</sub>/NH<sub>4</sub><sup>+</sup> with carbonyl functional groups (Zhang et al., 2015a). The C=N 531 double bonds in the organonitrogen imidazole compounds can act as effective 532 chromophores since both  $\pi \rightarrow \pi^*$  and  $n \rightarrow \pi^*$  transitions are chromatically active 533 (Nguyen et al., 2013). The UV/visible spectrum of imine and pyrrole show broad bands 534 at 270 nm (NIST, 2020), which was consistent with the UV absorption peak of the 535  $n \rightarrow \pi^*$  band observed here. According to the AMS results, carbonyl was the main 536 functional group of toluene SOA. The emergence of absorption peaks at 270~280 nm 537 demonstrated that some organonitrogen imidazole compounds (e.g. imines and pyrrole) 538 were formed through the heterogeneous reaction of toluene with NH<sub>3</sub>. Meanwhile, the 539

high-molecular weight nitrogen-containing organic species might have formed through
Maillard reactions in the particle-phase (Wang et al., 2010). This was also a reason for
the increase in SOA mass concentration in the presence of NH<sub>3</sub>.

The green line in Fig. 8 represents the MAC value of toluene-derived SOA in the presence of NO<sub>X</sub>, which was also higher than the black line (control) throughout the UV detection range. When compared with the red line, the green line had no obvious characteristic peak at 280 nm, but it had higher absorbance in the range between 240 and 280 nm. This indicated that both NO<sub>X</sub> and NH<sub>3</sub> increased the absorbance of toluene SOA, while the chromophores generated from the reactions between toluene-derived SOA with NH<sub>3</sub> or NO<sub>X</sub> did not behave in the same way.

The blue line in Fig. 8 represents the absorbance of toluene SOA formed in the 550 presence of both NO<sub>X</sub> and NH<sub>3</sub>. The MAC of toluene SOA formed in the presence of 551 both NO<sub>X</sub> and NH<sub>3</sub> was higher than the toluene SOA formed in the presence of either 552 NH<sub>3</sub> or NO<sub>X</sub>. There might have been a synergetic effect between NO<sub>X</sub> and NH<sub>3</sub> on the 553 absorbance of toluene SOA. Considering that the mass concentration of toluene SOA 554 formed in the presence of both NH<sub>3</sub> and NO<sub>X</sub> was the highest, as described in section 555 3.1, the co-existence of NH<sub>3</sub> and NO<sub>X</sub> may also result in the toluene SOA having 556 stronger light absorption and atmospheric radiative forcing. We also noted a higher 557 MAC value at 280 nm, which illustrated that the presence of NO<sub>X</sub> could promote the 558 formation of imines and pyrrole in the photooxidation system of toluene with NH<sub>3</sub>. 559

## 560 4 Conclusion

Here we present the results of a study in which we characterized the mass 561 concentrations, chemical compositions, and optical properties of SOA formed from the 562 photooxidation of toluene under different NH3 and NOx concentrations. When 563 compared with the control experiment, the SOA mass concentration data showed that 564 the formation of toluene-derived SOA was enhanced in the presence of NH<sub>3</sub>, through 565 acid-base reactions between carboxyl groups or Maillard reactions with carbonyls, but 566 inhibited in the presence of NO<sub>X</sub>. Meanwhile, the mass concentration of toluene SOA 567 568 formed in the presence of both NO<sub>X</sub> and NH<sub>3</sub> was higher than those formed under either NH<sub>3</sub> or NO<sub>X</sub> alone. This result indicated that there was a synergistic interaction between 569 NH<sub>3</sub> and NO<sub>X</sub> that further enhanced toluene-derived SOA formation. At the same time, 570 the lowest OS<sub>C</sub> value was obtained when both NH<sub>3</sub> and NO<sub>X</sub> were present. We 571 concluded that highly volatile compounds, which were formed from toluene 572 photooxidation in the presence of NO<sub>X</sub>, could react with NH<sub>3</sub> to form products with 573 574 lower volatilities, and promoted the partitioning of these products into the particlephase. 575

576 Synergetic effects of  $NH_3$  and  $NO_X$  on the formation of NOA and the optical 577 properties of SOA were also observed in this study. The heterogeneous reaction was 578 responsible for the formation of NOA in the presence of  $NH_3$ . Meanwhile, an absorption 579 peak at 270~280 nm, which is characteristic of imine and pyrrole, was observed. In 580 contrast, the formation of NOA caused by  $NO_X$  alone was mainly due to a gas phase 581 homogeneous reaction.

In the actual atmosphere especially in Chinese urban atmosphere, NO<sub>X</sub> and NH<sub>3</sub> 582 abundantly co-exist. Therefore, the findings presented here clearly show that the 583 584 synergetic effects of NO<sub>X</sub> and NH<sub>3</sub> should not be neglected. In the meantime, our work provides a scientific basis for the consideration of synergistic emission reductions of 585 NH<sub>3</sub> and NO<sub>X</sub> under the compound pollution conditions, which will contribute to 586 reducing the burden of aerosols in the atmosphere. It has to be noted that the 587 concentration of reactants used for the experiments is much higher than that observed 588 in polluted areas. Although the reactant concentrations including NH<sub>3</sub> used in this work 589 590 are much higher than those in the real urban environment, our results are applicable for the polluted urban atmosphere. In the urban atmosphere aromatic VOCs consist of 591 numerous species and their total concentration is much higher than a single species such 592 as toluene. On the other hand, carboxylic acids and carbonyls in the urban polluted 593 atmosphere can be produced from aromatics and many other species. Therefore, it is 594 reasonable for our smog chamber experiments to use toluene as a single precursor with 595 596 a concentration much higher than that in the real atmosphere. Although the mechanisms of SOA formed under high precursor concentrations is expected to be the same as that 597 under low concentrations, the kinetics are probably different. Thus, the effect of NH<sub>3</sub> 598 and NO<sub>X</sub> on the photooxidation of toluene with lower concentrations should be checked 599 in the further study. 600

601 **Data availability** 

The datasets are available from https://doi.org/10.6084/m9.figshare.16910953.

### 604 Author contributions

605 SL and GW designed the experiment. SL, DH and YW conducted the experiments. SL,

606 DH, YW and GW performed the data interpretation. SL and GW wrote the paper. YW,

607 SZ, XL, CW, WD contributed to the paper with useful scientific discussions or 608 comments.

609

### 610 **Competing interests**

611 The authors declare that they have no conflict of interest.

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# 910 **Tables**

### 911 912

#### Table 1. Summary of experimental conditions in this study

No.	Tol <sub>0</sub> (ppb)	∆ Tol (ppb)	NH3 <sup>a</sup> (ppb)	NO <sub>2</sub> (ppb)	RH (%)	SOA mass conc. <sup>b,c</sup> (µg m <sup>-3</sup> )	SOA yield <sup>b</sup> (%)
Exp.1	664.1	551.2	-	-	$25\pm1$	$637 \pm 14.6$	28.1
Exp.2	618.7	499.4	~200	-	$23\pm1$	$867\pm12.7$	34.7
Exp.3	620.9	526.1	~200	62	$26\pm1$	$1020\pm10.6$	42.7
Exp.4	645.7	539.5	-	63	$25\pm1$	$452\pm18.9$	19.5

 $^{a}$  The concentration of NH<sub>3</sub> is estimated by the amount of NH<sub>3</sub> added and the volume of the smog

914 chamber. <sup>b</sup> SOA concentration and yield were calculated after taking into account the wall loss. <sup>c</sup>

915 The reported SOA mass concentrations was the peak values after the wall loss correction.

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## 917 **Figures**



918



920 Fig. 1. The evolution of mass concentration (a) and yield (b) of toluene-derived SOA in different

921 experiments. All the mass concentrations were wall-loss corrected. The error bars were calculated922 by the fluctuation of measured SOA concentration after the UV light was turn off at the end of

923 each experiment



**Fig. 2.** The OS<sub>C</sub> values for the toluene SOA formed under different NH<sub>3</sub>/NO<sub>X</sub> conditions.



Fig. 3. The relationship between total organic signals at 43 m/z (f43) vs. 44 m/z (f44) from SOA
data during the photooxidation process. The f43 vs. f44 plots exhibited inflection points during the
photooxidation process. The dashed lines indicate the trends of f43 vs. f44 for the SOA formation
stage (before the inflection point) and the solid lines for the stable stage.



Fig. 4. The differential spectra of toluene SOA in the formation and stable stages. Data were taken
and analyzed at a high resolution but were summarized to a unit mass resolution for display. Only
minimal N-containing fragments could be observed in the Exp.1 without added NH<sub>3</sub> and NO<sub>X</sub>.
These N-containing fragments could be attributed to the background NH<sub>3</sub> and NO<sub>X</sub> in the chamber
or the systematic errors from AMS.





**Fig. 5.** The H/C, O/C, and N/C values of Hi-NOA and Lo-NOA for each experiment. (a) Exp. 2

with 200 ppb NH<sub>3</sub>, (b) Exp. 3 with 200 ppb NH<sub>3</sub> and 62 ppb NO<sub>2</sub>, and (c) Exp. 4 with 63 ppb
NO<sub>2</sub>.



Fig. 6. The evolution of high-nitrogen OA (Hi-NOA) and low-nitrogen OA (Lo-NOA) during the
photooxidation process under different NO<sub>X</sub>/NH<sub>3</sub> concentrations. Hi-NOA and Lo-NOA were not
consistent among experiments. (a) Exp. 2 with 200 ppb NH<sub>3</sub>, (b) Exp. 3 with 200 ppb NH<sub>3</sub> and 62
ppb NO<sub>2</sub>, and (c) Exp. 4 with 63 ppb NO<sub>2</sub>.





**Fig. 7.** The evolution of N/C in different experiments.



960 Fig. 8. The MAC over the range of 230 – 600 nm for the toluene SOA formed under different
961 experiment conditions.