1 The shifting of secondary inorganic aerosols formation mechanism 2 during haze aggravation: The decisive role of aerosol liquid water

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13 Abstract

14 Although many considerable efforts have been done to reveal the driving factors on haze aggravation, however, the roles of aerosol liquid water (ALW) in SIAs formation were mainly focused on the condition 15 16 of aerosol liquid water content (ALWC)<100 µg/m³. Based on the in-situ high-resolution field observation, 17 this work studied the decisive roles and the shifting of secondary inorganic aerosols formation mechanism during haze aggravation, revealing the different roles of ALWC in a broader scale (~ 500 μ g/m³) in nitrate 18 19 and sulfate formation induced by aqueous chemistry in ammonia-rich atmosphere. The results showed that 20 chemical domains of perturbation gas limiting the generation of secondary particulate matters presented obvious shifts from HNO₃ sensitive to HNO₃ and NH₃ co-sensitive regime with the haze aggravation, 21 22 indicating the powerful driving effects of ammonia in ammonia-rich atmosphere. When ALWC<75 µg/m³, the sulfate generation was preferentially triggered by the high ammonia utilization, then accelerated by 23 24 nitrogen oxide oxidation from Clean to Moderate pollution stages, characterizing as nitrogen oxidation ratio 25 (NOR)<0.3, sulfur oxidation ratio (SOR)<0.4, ammonia transition ratio (NTR)<0.7 and the moral ratio of NO₃⁻/SO₄²⁻=2:1. While ALWC>75 μg/m³, aqueous-phase chemistry reaction of SO₂ and NH₃ in ALW 26 became the prerequisite for SIAs formation driven by Henry's law in the ammonia-rich atmosphere during 27 28 Heavy and Serious stages, characterizing as high SOR (0.5-0.9), NOR (0.3-0.5), NTR (>0.7) and the moral ratio of NO₃^{-/}SO₄²⁻=1:1. A positive feedback of sulfate on nitrate production was also observed in this work 29 due to the shift of ammonia partition induced by the ALWC variation during haze aggravation. It implies the 30 target controlling of haze should not simply focus on SO2 and NO2, more attention should be paid on gaseous 31 precursors (e.g., SO₂, NO₂, NH₃) and aerosol chemical constitution during different haze stages. 32

33 Keywords: Mechanism shifting, Aerosol liquid water, Secondary inorganic aerosols, Haze

34 aggravation, In-situ observation

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35 **1 Introduction**

Fine particulate matter (PM_{2.5}) presented close link with several environmental issues, such as 36 37 visibility reduction and climate change (Zhang et al., 2015; Shang et al., 2020; Wang et al., 2020; Wang et al., 2016; Nozière et al., 2010). Epidemiological studies have stated the 38 association of PMs with various public health, even adverse birth outcomes (Gwynn et al., 39 40 2000; Lavigne et al., 2016; Zhao et al., 2020). As the most abundant secondary inorganic aerosols (SIAs) in PM2.5 during Chinese winter haze episodes (Fu and Chen, 2017; Liu et al., 41 2019), the formations of sulfate and nitrate play the key roles during haze aggravation, as well 42 as the impacting factors of the oxidants in gas and aqueous phases, the characteristics of pre-43 existing aerosols/fog/cloud, and meteorological conditions. Recently, aerosol liquid water 44 content (ALWC) was reported associating with the SIAs formation, especially sulfates and 45 46 nitrates, during the haze periods (Wu et al., 2018; Zheng et al., 2015a; Wang et al., 2016; Cheng et al., 2016; Carlton and Turpin, 2013; Nguyen et al., 2014; Xue et al., 2014; Tan et al., 2017; 47 Liu et al., 2017b). Atmospheric aerosol liquid water (ALW), which determined by ambient 48 relative humidity (RH), has been proposed as a container since it could provide the reaction 49 medium for the multiphase chemistry during the haze process (Ansari and Pandis, 2000; 50 Shiraiwa et al., 2012; Davies and Wilson, 2015). The roles of ALWC on the generations of 51 particulate sulfate generations (Wang et al., 2016; Cheng et al., 2016) and global secondary 52 53 organic aerosols (Hodas et al., 2014; Mcneill, 2015; Wong et al., 2015) were reported. Thus, 54 fully understanding ALW and its roles during haze aggravation is fundamentally important on atmospheric physicochemical processes, especially the liquid chemical transformation of SO2 55 and NO_x in ALW. 56

57 Ammonia is the most important alkaline gas, neutralizing with acidic species to form 58 ammonium salts. Due to little attention has been paid to NH3 emissions by Chinese government, atmospheric NH₃ experienced a significant increasing trend (Ge et al., 2019; Tan et al., 2017). 59 60 Although the increase in atmospheric NH₃ is beneficial to reduce atmospheric acidity (Liu et al., 2019), its chemical behavior on regional haze formation is still debating. Cheng et al. (2016) 61 indicated that the fast transform of gaseous SO₂ to particle sulfate under polluted conditions is 62 attributed to the neutralization of NH₃, which raises particle pH and thereby facilitated the 63 aqueous oxidation of S (VI) by NO₂. Fang et al. (2017) stated that NH₃ partition significantly 64 modified aerosol pH and thereby adjusting the partition of SO₂ and NO₂. Although the role of 65 NH₃ has been identified from a theoretical perspective, the lack of NH₃ emission control sets 66 barriers for more effective reduction of PM2.5. Therefore, it is urgent to fully understand the 67 reactive gases behavior and the chemical mechanism of SIAs formation during different 68

69 pollution stages, which will be helpful to propose reasonable strategies for each stage.

- So far, the SIAs formation has been extensively studied during short-term, continuous, or 70 persistent haze episodes, proposing several heterogeneous and homogeneous oxidation 71 pathways on sulfate and nitrate formation (Guo et al., 2014; Guo et al., 2017; Zheng et al., 72 2015b; Huang et al., 2014; Liu et al., 2021; Yao et al., 2020; Zhou et al., 2018; Liu et al., 2019). 73 74 In ammonia-rich atmosphere, NH₃ partition significantly modified aerosol pH, adjusted the partition of SO₂ and NO₂ (Fang et al., 2017) and promotes the aqueous oxidation of S (VI) by 75 NO₂ (Wang et al., 2016; Cheng et al., 2016). Although many considerable efforts have been 76 done to reveal the driving factors on haze aggravation, however, the roles of ALW in SIAs 77 formation were mainly focused on the condition of ALWC<100 μ g/m³ (Nenes et al., 2020; Wu 78 et al., 2018; Bian et al., 2014; Jin et al., 2020). Therefore, the roles of ALWC in a broader scale 79 80 and the mechanism shifting of secondary inorganic aerosols formation during haze aggravation in ammonia-rich atmosphere need to be understood in depth. Based on a continuous 81 82 observation with 1-hour resolution from December 2019 to January 2020, this work discussed the shift of dominant mechanism with ALWC variation during the time window of haze 83 aggravation processes, which will be helpful to propose more effective PM2.5 control strategies 84 for each pollution stage. 85
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2 Sampling and Experiment Methods

88 2.1 Description of Sampling Site

Hohhot, the capital city of Inner Mongolia Autonomous Region, is the central city of Hohhot-89 Baotou-Ordos group, as well as an important northern China city with a population of more 90 than 3.126 million and an area of 17224 km² (Fig. 1). This region is featured as continental 91 92 climate with marked seasonality changes, which characterized as long-lasting cold humid winter and short-time other seasons. Thereby, to survive the cold season, approximately half 93 year of coal-fired heating events (Oct. 15-the following Apr. 15) were introduced, which 94 emitting gaseous pollutants as well as PMs around-the-clock. The main industries include 95 thermal power plants, coal-energy based biochemical industry, dairy industry and 96 97 petrochemical industry, etc., which also emit atmospheric pollutants ceaselessly. Thus, high 98 concentrations of PMs pollution cases dominated the major contamination cases during winter season (data obtained from Department of Ecology and Environment of Inner Mongolia 99 Autonomous Region, http://sthjt.nmg.gov.cn/) and gradually emerging as the limiting factor on 100 regional ambient air quality and human health. 101

102 In this study, the observation was conducted at the Inner Mongolia Environmental

Monitoring Center (40°49'22"N, 111°45'2"E) on a top of a sixteen-story building (~40m above
the ground level) located at the eastern part of the downtown near the People's Government of
Inner Mongolia Autonomous Region near the 2nd ring road from December 1, 2019 to January
31, 2020. Residential and administrative regions were characterized as the major functional

107 domain near the sampling site, with no direct industrial regions nearby.



108

109 2.2 Data acquisition and analysis methods

110 2.2.1 Data acquisition

111 On-line ion-chromatograph instrument (MARGA ADI 2080, Metrohm Applikon, Switzerland) was employed to simultaneously determine the water-soluble inorganic ions (Na⁺, NH4⁺, Mg²⁺, 112 Ca²⁺, K⁺, Cl⁻, F⁻, SO4²⁻, NO3⁻) in PM_{2.5} and corresponding trace gases (SO₂, HNO₂, HNO₃, HCl, 113 NH₃). This instrument has been widely used in previous work (Rumsey et al., 2014; Nie et al., 114 2015; Huang et al., 2020) and the details were listed in Supplement (S1.1). Correspondingly, 115 gaseous pollutants (e.g., NOx, CO, PM1, PM2.5, PM10) and meteorological datasets (e.g., wind 116 117 speed, wind direction, RH, temperature, etc.), as well as the adopted models could be found in our previous work (Xie et al., 2021). In addition, peroxyacetyl nitrates (PANs), nitrous oxide 118 (N₂O) and solar spectrophotometry were measured by PANs-100 (Focused Photonics Inc.), 119

120 N₂O Monitor (LSE, Monitors) and CE-318T (CIMEL), respectively.

121

122 2.2.2 Analysis methods

Generally, sulfur oxidation ratio (SOR) and nitrogen oxidation ratio (NOR) were calculated as follows, which were used to indicate the contribution of secondary transformation during the haze events (Song et al., 2007; Zhou et al., 2018).

126
$$SOR = \frac{n(SO_4^{2^-})}{n(SO_2) + n(SO_4^{2^-})}$$

127
$$NOR = \frac{n(HNO_3) + n(NO_3^-)}{n(NO_2) + n(HNO_3) + n(NO_3^-)}$$

128 Meanwhile, as an indicator of ammonia conversion efficient, ammonia transition ratio 129 (NTR), was calculated as the following equation (All units were $\mu g/m^3$).

130
$$NTR = \frac{NH_4^+/18}{NH_4^+/18 + NH_3/22.4}$$

131 In addition, as the fractions of ammonia, nitrate and sulfate in deliquesced aerosol, ϵ 132 (NO₃⁻), ϵ (NH₄⁺) and ϵ (SO₄²⁻) were expressed as follows.

133
$$\varepsilon(NO_3^-) = \frac{n(NO_3^-)}{n(HNO_3) + n(NO_3^-)}$$

134
$$\varepsilon(NH_4^+) = \frac{n(NH_4^+)}{n(NH_3) + n(NH_4^+)}$$

135
$$\varepsilon(SO_4^{2-}) = \frac{n(SO_4^{2-})}{n(SO_2) + n(SO_4^{2-})}$$

136 2.2.3 Aerosol pH

In this work, a widely used thermodynamic model, ISORROPIA-II (Song et al., 2018; Gao et 137 138 al., 2020), was employed to establish aerosol acidity. Including the concentrations of WSIs in PM_{2.5} and gaseous pollutions (e.g., NH₃, HCl), the simultaneously measured temperature and 139 RH data were imported into its Na⁺-K⁺-Ca²⁺-Mg²⁺-NH4⁺-SO4²⁻-NO3⁻-Cl⁻-H₂O aerosol system. 140 According to previous study (Song et al., 2018) and our data profiles, "Forward Mode" and 141 "Metastable State" were selected in the model of ISORROPIA-II to calculate aerosol acidity 142 $(H_{air}^+, H^+ \text{ loading per volume air } (\mu g/m^3))$ and aerosol liquid water content (ALWC). Then the 143 144 aerosol pH was calculated by the following equation.

145
$$pH = -\log_{10} \frac{1000H_{air}^+}{ALWC}$$

146 The concentrations of NH₃, NH_4^+ , NO_3^- and SO_4^{2-} modeled by this model significantly

147 correlated with their measured values with correlation coefficients of 0.971-0.999, indicating

the accuracy and acceptability of the model in this work (Fig. S1).

149 2.2.4 Heterogeneous sulfate production

Due to the necessity of precise SO4²⁻ generation, heterogeneous sulfate production (P_{het}) was
parameterized and calculated according to the following equation(Jacob, 2000; Zheng et al.,
2015a),

153
$$P_{het} = \frac{3600 sh^{-1} \times 96 gmol^{-1} \times P}{R \times T} \left(\frac{R_p}{D_g} + \frac{4}{v\gamma}\right)^{-1} S_p[SO_2(g)]$$

Where P_{het} was presented in $\mu g \cdot m^{-3} \cdot h^{-1}$, 3600sh⁻¹ is time conversion factor, 96 g/mol is the 154 molar mass of SO4²⁻, P is atmospheric pressure in kPa, R is the gas constant with the value of 155 8.31 Pa·m³·mol⁻¹·K⁻¹·, T is the temperature with the unit of K, R_p represented the radius of 156 aerosol particles (m), D_g is the SO₂ molecular diffusion coefficient and v is the mean molecular 157 speed of SO₂ with the typical tropospheric value of $2 \times 10^{-5} \text{m}^2 \cdot \text{s}^{-1}$ and 300 m·s⁻¹, respectively. γ 158 is the uptake coefficient of SO₂ on aerosols, S_p is the aerosol surface area per unit volume of 159 air (m²·m⁻³) (Jacob, 2000). PM_{2.5} mass concentrations (µg·m⁻³) and mean radius (m) during 160 campaign were roughly calculated utilizing the following empirical formula published by Guo 161 et al. (2014): 162

163
$$R_p = (0.254 \times C_{(PM_{2.5})} + 10.259) \times 10^{-9}$$

164 mean density of particles ρ was calculated and showed as 1.5×10^6 g·m⁻³ using the volume 165 and surface area formulas of a sphere (Guo et al., 2014). S_p was estimated from the following 166 formula:

167
$$S_p = \frac{C_{(PM_{2.5})} \times 10^{-6} g \cdot \mu g^{-1}}{4/3 \cdot \pi R_p^3 \cdot \rho} \cdot 4\pi R_p^2$$

168 relative humidity-dependent γ were derived according to Zheng et al. (2015a) during the 169 campaign in this work and showed as the following formular:

170
$$\gamma = \begin{cases} 2 \times 10^{-5}, & \Psi \le 50\%, \\ 2 \times 10^{-5} + \frac{5 \times 10^{-5} - 2 \times 10^{-5}}{100 - 50\%} \times (\Psi - 50\%), & 50\% \le \Psi \le 100\% \end{cases}$$

171 where ψ referred to RH with the unit of %.

172 **3 Results and Discussion**

Based on National Ambient Air Quality Standards of China (HJ633-2012)
(<u>https://www.mee.gov.cn/ywgz/fgbz/bz/bzwb/jcffbz/201203/t20120302_224166.shtml</u>), air
quality index (AQI) was introduced in this work to classify pollution levels (Wang et al., 2015;

- 176 Kanchan et al., 2015; Xu et al., 2017) and discuss the characteristics of atmospheric pollutants.
- 177 Briefly, daily concentrations of PM_{2.5} ranged from 0-75, 75-115, 115-150, 150-250 and >250
- 178 µg/m³ were classified as clean (C), light polluted (L), moderate polluted (M), heavy polluted
- 179 (H) and serious polluted (S) periods, respectively.

180 *3.1 The observed evidence for ammonia-rich atmosphere*

- 181 The characteristics of atmospheric pollutants and meteorological parameters during the studied period were summarized in Supplement (S2.1). In this work, molar ratios of NH4⁺ vs. anions 182 was used to identify the chemical species of ammonium salts (Zhou et al., 2018; Wang et al., 183 2021; Liu et al., 2017b; Shi et al., 2019). The calculated results (Supplement, S2.2) showed the 184 predominant chemical species of ammonium gradually varied from the coexistence of 185 ammonium sulfate ((NH₄)₂SO₄) and ammonium nitrate (NH₄NO₃) to the coexistence of 186 187 ((NH₄)₂SO₄), NH₄NO₃ and ammonium chloride (NH₄Cl) with haze aggravation (Fig. S5). Further, the slope of fitted equation between excess-NH4⁺ and anions were still lower than 1:1 188 189 line after neutralized all the measured anions, indicating the ammonia-rich atmosphere (Fig. S5c). To meet the national demand of ultra-low emissions activities (nearly two times lower 190 than former national standard) on gaseous pollutants, heavy usage of ammonia-containing 191 compounds in the process of desulfurization and denitrification (Solera García et al., 2017; Tan 192 et al., 2017) at broadly distributed thermal power plants (>300,000kWh) and the close-set coal-193 fired heating stations (Fig. 1) resulted ammonia fugitive provided a reasonable explanation on 194 195 this ammonia-rich atmosphere. Although the retrofit of national demand of ultra-low emissions activities on gaseous pollutants (nearly two times lower than former national standard) has 196 been completed, distributed coal-based enterprises could also emit substantial SO₂ and NO₂ 197 and subjecting to heterogeneous reactions to further generate sulfate and nitrate and aggravated 198 199 the haze events (Fig. S7a, S7b).
- To show the reaction between ammonia and nitric acid and the other formation processes 200 201 of nitrate in different (relative) concentrations of sulfate, the data of previous studies and different pollution levels (C, L, M, H, S) in this work were plotted in Fig. 2. When 202 $[NH_4^+]/[SO_4^{2-}] \le 1.5$, the nitrate formation associated with crustal elements rather than 203 ammonium; while [NH4⁺]/[SO4²⁻] >1.5, the homogeneous gas-phase reactions between NH3 204 and HNO₃ became the major pathway for atmospheric ammonia to form NH₄NO₃ (Pathak et 205 al., 2009; Liu et al., 2019). The results illustrated that the ammonia-rich regimes were not only 206 207 found in Hohhot, but also observed in Guangzhou (Huang et al., 2011), Chengdu (Huang et al., 2018), Lanzhou USA West and East, India, Ireland, Europe, Qingdao, Italy, Lin'an (Pathak et 208 al., 2009) in recent decades (Fig. 2). It suggested that atmospheric oxidative modifications in 209

ammonia-rich atmosphere should be a widespread atmospheric issue with significant contributions on SIA generation. It was worth noting that the slopes of our data were becoming steeper, coupling with the NO_3^{-}/SO_4^{2-} ratios change from ~4 to about 1, as the increasing pollution levels. The high $PM_{2.5}$ nitrate concentration during Heavy and Serious stages cannot be explained by the homogeneous gas-phase reaction involving ammonia and nitric acid, which may be associating with the heterogeneous reaction in ALW on the surface of the preexisting aerosols.





218 **3.2 Driving mechanism of SIAs formation**

219 3.2.1 Aerosol liquid water

Our results showed that SOR, NOR and SIAs in PM_{2.5} presented increasing trends with the increasing ALWC during the five pollution levels. The variation of predominant chemical species of ammonium (Fig. 2) indicated more SIAs will be generated on particles with the simultaneous increase of ALWC and PM_{2.5} (Fig.3b). Theoretically, the inorganic compounds conversion was enhanced via aqueous phase chemistry on moist particles owing to the continuous partition of gaseous pollutants (e.g., SO₂, NO₂, N₂O₅) in ALW, then disrupted the equilibrium between the gaseous and condensed phases, resulting in the aggravation of haze

events (Xue et al., 2014; Wu et al., 2018; Zheng et al., 2015b; Wang et al., 2016). Considering 227 seasonal heating characteristics, the shift of the equilibrium between gaseous and condensed 228 phases was enhanced with the increasing atmospheric pollutants concentrations due to the coal-229 fired combustion events in winter. Detailly, owing to hygroscopic nature, the particles must 230 increase their water contents via ALW along with RH (Fig. S8a) to maintain thermodynamic 231 232 equilibrium and water vapor and simultaneously enhance the oxidation and dissolution of precursors in the micro-solution (ALW) of the particulates. This process elevated the inorganic 233 mass fraction as well as particulate mass concentrations during different pollution stages (Fig. 234 S8b) (Bertram et al., 2009; Wang et al., 2016; Zheng et al., 2015a; Cheng et al., 2016). Due to 235 the larger affinity of H₂SO₄ for NH₃ (aq), sulfate was preferentially and fully neutralized by 236 ammonium in the ammonia-rich atmosphere to generate non-volatile nature of (NH4)2SO4 (Liu 237 238 et al., 2017b; Zhou et al., 2018; Wang et al., 2021). Thus, SOR presented higher exponential growth with the elevated AWLC coupling with more sulfate production (Fig. 3b). 239 240 Concomitantly, the preferentially generated (NH₄)₂SO₄ further enhanced the hygroscopicity of particulate matter, in turn, helped more ammonia partitioning into moist particulate matter and 241 generating ammonium salts accelerating haze aggravation (Supplement, Fig. S6, Fig. S8c). 242 Thus, most important of all, the sharp increase of inorganic compounds associating with the 243 elevated ALWC significantly modified the specific surface area of particulates and further 244 accelerated the hygroscopic aerosol growth, which simultaneously provided a substrate for the 245 246 ensuing heterogenous reaction and accelerated the evolution of haze events. Previous work reported that particles of different modes made different contributions to ALWC with the 247 contributions of nuclear, Aitken, accumulation and coarse modes assessed at <1%, 3%, 85% 248 and 12%, respectively, indicating that the contribution of accumulation mode particles to 249 250 ALWC dominated among all the aerosol particle modes (Tan et al., 2017). Meanwhile, the observed significant correlations of ALWC with the ratios (PM_{1.0}/PM_{2.5} and PM_{2.5}/PM₁₀) in 251 this work also indicated that the hygroscopic growth of fine particulate matter ($D_p \leq 2.5$ um) 252 strongly associated with ALWC (Fig. 3a). Accordingly, both the previous work and our 253 monitoring results suggested that the ratios of PM1.0/PM2.5 and PM2.5/PM10 could be used as 254 the proxy of the hygroscopic growth of particulate matter. 255

256



of relative humidity

257 *3.2.2 Perturbation gases*

Due to the strict control of SO₂, atmospheric concentrations of NO₂ and NH₃ gradually became 258 as the decisive reactive precursors on regional atmospheric secondary particulate matter 259 generation. Thus, the state-of-the-art flamework proposed by Nenes et al. (2020) was carried 260 out to exam the chemical domain classifications and the decisive precursor based on the data 261 sets of previous studies (Nenes et al., 2020) and this work (Fig. 4). Due to the 262 thermodynamically stable property of the preferentially generated (NH₄)₂SO₄, the semi-263 volatile NH₄NO₃ dominate the partitioning of NH₃^T (sum of NH₃ and NH₄⁺, same to NO₃^T) 264 and NO₃^T. Although aqueous NO₃⁻ concentrations varied with haze processes, the calculated ε 265 (NO_3^T) (detailed calculated method could be found in S1.2), which was an equilibrium 266 parameter between gaseous HNO₃ and particle-phase NO₃⁻ (Guo et al., 2016; Fang et al., 2017), 267 presented consistently full loadings of nitrate on the existing particulates during the studied 268 period (Fig. S9a, Fig. S9b). This could provide clear evidence for the initial HNO₃ sensitive 269 270 area and continuous control of HNO3 during the studied periods. However, with haze aggravation, significant elevated ALWC resulted in more precursors partitioned in micro-271 droplets to maintain water vapor. This process induced a positive shift of HNO₃ dissolution 272

equilibrium and leading more HNO₃ partitioned on particles driven by the Henry's law (e.g., 273 HNO_{3(g)} \leftrightarrow HNO_{3(aq)}, $K_H = 2.07$ mol/(L·Pa)). Meanwhile, HNO₃ and HONO could also produce 274 through the reactions of $NO_2 + H_2O \xrightarrow{Het} HNO_3 + HONO$ (Huang et al., 2018). Accordingly, 275 the OH radicals generated by HONO photolysis also contributed to this oxidation processes 276 (Yue et al., 2020; Zhu et al., 2020). These aqueous oxidations processes were evidenced by the 277 278 observation of significantly elevated HONO and PANs during the haze aggravation (Supplement, Fig. S7c, Fig. S7d). Accordingly, the equations of $NH_4^+ + NO_3^- + H^+ + OH^- \rightleftharpoons$ 279 $NH_4NO_3 + H_2O$ and $NH_4^+ + SO_4^{2-} + H^+ + OH^- \rightarrow (NH_4)_2SO_4 + H_2O$ were shifted to 280 generate more NH4NO3 and (NH4)2SO4 (Nenes et al., 2020; Xie et al., 2020) due to the driving 281 force of more ammonia partitioned in elevated ALWC (NH₃+H₂O ≠ NH₃·H₂O, NH₃·H₂O ≠ 282 $NH_4^++OH^-$). Therefore, NH_3 and NO_x became as the decisive factors on regional atmospheric 283 oxidability in the ammonia-rich regime (Zhai et al., 2021; Tan et al., 2017; Liu et al., 2019; Li 284 et al., 2019). 285

Generally, both NH₃ and HNO₃ were the limiting factors governing the aerosol generations 286 for cities of North China due to high loadings of atmospheric ammonia, while NH₃ governed 287 PM formation for the southeast US (SAS) (Zhao et al., 2020). Thanks to the raw data of 288 Shenzhen (SZ) (Wang et al., 2022), we also calculated the ALWC and aerosol pH using 289 ISORROPIA-II and the scatters of SZ suggested obvious chemical transition from HNO₃-NH₃ 290 regime to NH₃ sensitive regime due to the differently originated air masses. Although both 291 292 cities located in US, the findings in California (CNX) were quite interesting and distributed in the insensitive region and the combined NH₃-HNO₃ sensitive region due to the moderate NH₃ 293 levels and the complicated atmospheric conditions during the observation (Nenes et al., 2020). 294 In our work, the data points (541/744) in summer (pH= 3.47 ± 1.29) mostly lied in HNO₃ 295 sensitive region, while chemical domains of perturbation gas limiting the generation of 296 secondary particulate matters presented obvious shifts from HNO3 sensitive to HNO3 and NH3 297 co-sensitive regime with the haze aggravation in winter. Some data points of this work lied in 298 the combined NH₃-HNO₃ region in winter owing to the more acidic condition. Under the stable 299 pH of aerosols in winter at Hohhot (pH=4-5), the more important is that a fraction of points 300 will distribute in the combined NH3-HNO3 region when ALWC>75 µg/m³, which may be 301 attributed to the aqueous chemical transformation driven by Henry's law mentioned above due 302 to the elevating ALWC. Comparatively, the aerosols pH in summer was significantly lower 303 304 than those in winter in Hohhot. Compared to TJ and SZ, the aerosols pH of Hohhot in winter was also significantly higher (Fig. 4) due to the acidity of atmospheric PM is largely depended 305 on the alkaline material in surface soils in arid and semi-arid region and the elevated 306

atmospheric ammonia. In terms of seasonal characteristics, the higher temperature in summer 307 elevates the volatility of NH4NO3 and dominates the partitioning of NH3^T in atmospheric phase 308 to decrease the pH of aerosols. Therefore, as can be seen from Fig. 4, the data points measured 309 in winter Hohhot characterized as higher pH and low ALWC than those in summer (Hohhot, 310 SAS, CNX, SZ). According to the framework of Nenes et al. (2020), the transition points of 311 312 Hohhot (whether winter or summer) between NH3-dominated and HNO3-dominated sensitivity also occurs at a pH around 2 but at lower levels of ALWC. Theoretically, it should be associated 313 with the more aridity of Hohhot locating in the arid and semi-arid region of China. Our results 314 provided the evidence for "the additional insight" proposed by Nenes et al. (2020) that the 315 transition ALWC varies with season change and the aridity of sites, in response to seasonal 316 variability and climate change. Although this effort could provide sound explanation for 317 limiting gaseous pollutants on PM formation, mechanisms on their chemical domains, 318 especially the roles of ALW in different locations with various conditions need further study in 319 320 the future.



321 3.2.3 The shifting of SIAs formation mechanism driven by ALW

322 It's worth noting that two independent correlations were found between SOR and odd oxygen

mechanisms of atmospheric oxidability on sulfate generations at different stages (Fig. 5a). Different to inefficient homogeneous sulfate oxidation efficiency (Supplement, Fig. S10), significant correlations pairs of NO₂ with SOR (Fig.5b) and NOR with SO_4^{2-} (Fig.5c) suggested the haze aggravation was largely related to the regional NO₂ levels due to the regulating effects on atmospheric oxidizability. Thus, the aqueous-phase oxidation of S(IV) by NO₂ (aq) was triggered and accelerated by the increasing ALWC and the following equation (Yao et al., 2020; Wang et al., 2016) (Supplement, Fig. S11a):

331
$$S(IV) + NO_2(aq) + H_2O \rightarrow S(VI) + H^+ + NO_2^-$$
 (R1)

Meanwhile, sharp logarithmic increase between SOR and NH4⁺ were also observed from Clean to Moderate pollution stages (Supplement, Fig. S12). Due to the joint effects of ammonia-rich atmosphere and ammonia's extremely water-soluble property, sufficient hydroxide generated by ammonia dissolution forced the NO₂ partitioned in ALW to maintain pH through neutralization and producing sulfate via R1. Thus, the following equation (R2) was derived with considering the processes of ammonia hydrolysis, which was evidenced by Fig. S11b.

338 $S(IV) + NO_2(aq) + NH_3(aq) + H_2O \rightarrow S(VI) + NH_4^+ + NO_2^- + H^+ + NO_3^-$ (R2)

Generally, NOR<0.1 means insignificant nitrogen oxide oxidation, therefore the observed regime shift of nitrate and ammonia chemical behavior on sulfate generation suggested the sulfate generation was preferentially triggered by the high ammonia utilization, then accelerated by the co-effects of ammonia utilization and nitrogen oxide oxidation (Fig. 5c).



Fig. 5 Correlations between (a) O_x and SOR, (b) NO₂ and SOR, (c) NOR and SO₄²⁻

Accordingly, the reaction R2 was activated due to the increased ALWC forced more ammonia to partition into moist particulate matter driven by the Henry's law in the ammoniarich atmosphere ($NH_{3(g)} \rightarrow NH_{3(aq)}$) (Supplement, Fig. S9c) (Clegg et al., 1998; Wu et al., 2018;

Xie et al., 2020). Meanwhile, our calculated aqueous generated NO₃⁻ nicely matched theoretical 346 nitrate aqueous generation curve (the solid blue line in Fig. S9b) proposed by Guo et al. (2017), 347 suggesting the pathway of fast sulfate formation from oxidation of S(IV) by NO₂ to generate 348 HONO (Wang et al., 2020) (Supplement, Fig. S11) via the reaction R2. As a result, the 349 thermodynamically stable (NH4)₂SO₄ would be preferentially formed to maintain its water 350 351 vapor pressure and thermodynamic equilibrium, then trigged the haze formation. Thus, the 352 mentioned effects resulted in a pronounced increase of NH₃ partitioning with the haze aggravation, suggesting the importance of ammonia partition on sulfate generations, namely, 353 NTR-controlled regime with ALWC<75 μ g/m³. In summary, when ALWC<75 μ g/m³, the 354 sulfate generation was preferentially triggered by high ammonia utilization, then accelerated 355 by nitrogen oxide oxidation from Clean to Light pollution stages (Fig. 5c) with NOR<0.3, 356 357 SOR<0.4 and NTR<0.7. In this period, the chemical composition of SIAs characterized as the moral ratio of NO_3 :SO₄²⁻=2:1 (Fig. 6). 358

When ALWC>75µg/m³, the haze was aggravated from Moderate to Serious stages along 359 with the increasing ALWC. As a result of increase in ALW, large amount of H⁺ was dissociated 360 during the generation of ammonium sulfate (Supplement, Fig. S13a). From Light to Moderate 361 pollution stages, the solubility SO₂ driven by Henry's law was self-limiting due to the acidity 362 363 effect in low ALWC (with ALWC<75 µg/m³). Therefore, low sulfate concentrations coupled with low ALWC at the beginning of haze event (Supplement, Fig. S13a). However, due to the 364 365 co-effects of elevated ALWC and hygroscopic nature of pre-generated ammonia sulfate, H⁺ concentrations were diluted and nearly constant in-situ pH with the increase of ALWC during 366 Heavy and Serious pollution stages (Supplement, Fig. S14) (Wang et al., 2016; Clifton et al., 367 1988; Huie and Neta, 1986; Lee and Schwartz, 1982). Hence, the significantly elevated ALWC 368 369 provided more chance for the partition of SO₂, NO₂ and NH₃ in ALW from Moderate to Serious pollution stages. Theoretically, Henry's constants of NO₂ $(9.74 \times 10^{-8} \text{ mol} \cdot (L \cdot Pa)^{-1})$ is 3-4 orders 370 of magnitude lower than those of SO₂ $(1.22 \times 10^{-5} \text{ mol} \cdot (L \cdot Pa)^{-1})$ and NH₃ $(6.12 \times 10^{-4})^{-1}$ 371 $mol \cdot (L \cdot Pa)^{-1}$), however, it is worth noting that the aqueous generated NO₃⁻ from Moderate to 372 Serious stages rapidly increased 2-5 times higher than Clean and Light stages (Supplement, 373 Fig. S9b). Meanwhile, according to our monitoring results, the solar spectrophotometry at 374 375 380nm during Moderate to Serious stages was significantly lower than that in Clean stage (Supplement, Fig. S15), suggesting the aqueous oxidation of NO₂ was the predominant 376 compared to chain photolysis (Huang et al., 2018). Accordingly, it could be deduced that 377 aqueous-phase chemistry reaction of SO2 and NH3 in ALW, driven by Henry law, became the 378 dominant mechanism for sulfate formation due to more NO2 was required to take part in the 379

fast sulfate formation with the increase of ALWC in the ammonia-rich atmosphere by the 380 reaction R2. Thus, with the increasing of ALWC, high concentrations of sulfate and nitrate with 381 high SOR (0.5-0.9), NOR (0.3-0.5) and NTR (>0.7) induced the haze events becoming Heavy 382 and Serious levels (Fig. 5c). Simultaneously, the calculated heterogeneous sulfate production 383 rate (Jacob, 2000; Mcneill, 2015) (Supplement, Fig. S16) presented similar trends with the 384 385 impacts of ammonia on sulfate production during different pollution stages (Xue et al., 2016; Cheng et al., 2016; Liu et al., 2020). It further stated the environmental significance of the 386 partitioning of SO₂ and NH₃ between gas and aqueous (ALW) phases for SIAs formation and 387 haze aggravation. Our results provided the evidence of significant negative correlations 388 between HONO and N₂O (Supplement, Fig. S17) from Moderate to Serious stages and positive 389 correlations between HONO and SOR (Supplement, Fig.S11a), highlighting the recent reported 390 391 secondary aqueous-phase oxidation pathway of SO2 by HONO from moderate pollution period $(2N(III) + 2S(IV) \rightarrow N_2O \uparrow + 2S(VI) + other products)$ (Wang et al., 2020). In summary, 392 when ALWC>75 µg/m³, aqueous-phase chemistry reaction of SO₂ and NH₃ in ALW became 393 the prerequisite for SIAs formation driven by Henry's law in the ammonia-rich atmosphere 394 during Heavy and Serious stages with high SOR (0.5-0.9), NOR (0.3-0.5), NTR (>0.7). In this 395 period, the chemical composition of SIAs characterized as the moral ratio of NO₃⁻:SO₄²⁻=1:1 396 397 (Fig. 6).



Fig. 6 The characteristics and formation mechanism of SIAs during haze aggravation

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399 3.2.4 The positive feedback of sulfate on nitrate production

Previous works suggested that the homogeneous reaction of NO₂ with OH radicals during 400 daylight and heterogeneous hydrolysis of N2O5 at night were the main routes on nitrate 401 formation during haze episodes (He et al., 2018; Liu et al., 2020; Liu et al., 2019). Unsurprising, 402 higher nitrate production rates (ΔNO_3^- , the difference of hour concentrations and matrixing 403 404 afterwards) were frequently observed in ammonia-rich conditions due to that ammonia-rich regime was more conducive on nitrate generation. However, the high level of nitrate production 405 rates (ΔNO_3^{-}) were found in the area characterizing as high ammonium and low sulfate levels, 406 suggesting that highly utilizing ammonium and pre-generated sulfate promoting particle-phase 407 nitrate generations (Fig. 7). 408



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Here, we proposed a hypothesis about the hydrogen ion concentration to respond the above observations. As is known to all, apart from the extremely low levels of crustal elements, ammonia is the only alkaline gas to neutralize the acidic gases in the atmosphere and generate ammonium ions (Xie et al., 2020). Thus, the concentrations of particulate sulfate and nitrate are affected by the partitioning of NH_4^+/NH_3 . Thereby, higher values of ΔNO_3^- and ΔSO_4^{2-}

always occurred in the regions with higher ammonium ions were not confused (Fig. 7, Fig. 415 S18). According to both our results and published laboratory work (Wang et al., 2016), the 416 acidity of the particulate matter could be significantly modified by the bulk aqueous reaction 417 between NO₂ and SO₂, in which this reaction could be further enhanced due to in the presence 418 of NH₃. As a result of the increase in RH, the partitioning of atmospheric ammonia was broken 419 420 in a deep extent, which enhanced the neutralization of S(VI) by ammonia at the particle surface to generate ammonium sulfate and dissociate huge H⁺ (Fig. S13b, red part). Simultaneously, 421 422 the ALWC did not raised significantly (Fig. S14b) at the beginning of haze event with relative low sulfate concentrations. Thus, hydrogen ions generated from sulfate dissociation absorb 423 ammonia more effectively from the ammonia-rich atmosphere at low relative humidity during 424 the early pollution stages, which significantly promotes the net nitrate production. However, 425 426 due to the co-effects of elevated RH and hygroscopic nature of pre-generated ammonia sulfate, H⁺ concentrations were diluted and shown as nearly constant in-situ pH (Fig. S14a). According 427 428 to previous works, the reaction between firstly generated sulfate and bisulfate with ammonia 429 were treated as the determination reaction on particle acidity (Weber et al., 2016; Liu et al., 2017a). This reaction is self-limiting due to the acidity effect, namely that it increases the 430 acidity of aqueous phase and in turn reduces the efficiency of Henry's constant for SO₂ 431 432 solubility and reaction rate and reduced the H⁺ formation rates from moderate periods, compared with clean periods (Fig. S13b, blue) (Wang et al., 2016; Clifton et al., 1988; Huie 433 434 and Neta, 1986; Lee and Schwartz, 1982). Due to the co-effects of RH increase and hygroscopic of sulfate, the ALWC was significantly elevated with the worsen of haze. Although 435 more H⁺ was generated in this process, no significant decrease in pH was found with the haze 436 aggravation due to the dilution effect of ALWC on H⁺. Previous works suggested that in the 437 438 case of ALWC increase, nitrate production is controlled by elevated H⁺ associating with the increase of sulfate, namely, NO3⁻ presented elevating trend with the increases of H⁺ 439 440 concentration (Xie et al., 2020). Thus, although H⁺ from the dissociation of sulfuric acid and full-loaded particle nitrate in conjunction with the haze aggravation generate particle HNO3 441 (Fig. S19a) could forcing more ammonia partitioned on the particles to generate ammonium 442 nitrate (Fig. S19b), net nitrate production (ΔNO_3^-) was nearly consistent. 443

444 **4 Conclusions**

The formation of SIAs, especially sulfates and nitrates, was inherently associated with ALWC during the haze aggravation, in which the roles of ALWC should be more significant in ammonia-rich atmosphere. The novelty of our work is to find the shifting of secondary inorganic aerosols formation mechanism during haze aggravation and explain the different

roles of ALWC in a broader scale (~500 ug/m³) in ammonia-rich atmosphere based on the in-449 situ high-resolution on-line monitoring data sets. The results showed that chemical domains of 450 perturbation gas limiting the generation of secondary particulate matters presented obvious 451 shifts from HNO₃ sensitive to HNO₃ and NH₃ co-sensitive regime with the haze aggravation, 452 indicating the powerful driving effects of ammonia in ammonia-rich atmosphere. When 453 ALWC<75 ug/m³, the sulfate generation was preferentially triggered by the high ammonia 454 utilization, then accelerated by nitrogen oxide oxidation from Clean to Moderate pollution 455 stages, characterizing as NOR<0.3, SOR<0.4, NTR<0.7 and the moral ratio of NO₃⁻:SO₄²⁻=2:1. 456 While ALWC>75 ug/m³, aqueous-phase chemistry reaction of SO₂ and NH₃ in ALW became 457 the prerequisite for SIAs formation driven by Henry's law in the ammonia-rich atmosphere 458 during Heavy and Serious stages, characterizing as high SOR (0.5-0.9), NOR (0.3-0.5), NTR 459 (>0.7) and the moral ratio of NO3⁻:SO4²⁻=1:1. A positive feedback of sulfate on nitrate 460 production was also observed in this work. Our work provides a potential explanation for the 461 462 interactive mechanism and feedback between nitric aqueous chemistry and sulfate formation in ammonia-rich atmosphere based on high-resolution field observation. It implies the target 463 controlling of haze should not simply focus on SO₂ and NO₂, more attention should be paid on 464 gaseous precursors (e.g., SO₂, NO₂, NH₃) and aerosol chemical constitution during different 465 466 haze stages.

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468 *Data availability*. All data of this study are available from the corresponding author upon 469 reasonable request (lcw2008@imu.edu.cn).

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471 Supplement. The Supplement related to this article is available online at

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Author Contributions. FX: Data curation, Formal analysis, Software, Writing-original draft.
YS: Investigation, Formal analysis. YLT: Methodology, Software. YSH: Investigation, Formal
analysis. XJZ: Investigation, Formal analysis, Software. PW: Methodology, Investigation.
RHY: Software, Writing-review & editing. WW: Investigation, Validation, Writing-review &
editing. JH: Investigation, Methodology. JYX: Investigation, Validation, Supervision, Writingreview & editing. CWL: Initiating and leading this research, Supervision, Writing-review &
editing.

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481 *Competing interest.* The authors declared that they have no conflict of interest.

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