1	Role of ammonia on fine-particle pH in agricultural regions of China:
2	Comparison between urban and rural sites
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22	Abstract: Particle acidity is a fundamental property that affects atmospheric particulate chemistry.
23	Synchronous online monitoring was performed in two urban sites (e.g., Zhengzhou (U-ZZ) and
24	Anyang (U-AY)) and three rural sites (e.g., Anyang (R-AY), Xinxiang (R-XX), and Puyang (R-PY))
25	in Henan Province during a haze episode to investigate the pH value and its driving factors in the
26	agricultural regions of China. The pH values of particles calculated by ISORROPIA-II model at rural
27	sites were slightly higher than those at urban sites, with the median (min-max) values of 5.2 (4.8-6.9,
28	R-PY), 5.1 (4.7–6.5, R-AY), 4.9 (4.1–6.8, R-XX), 4.8 (3.9–5.9, U-AY), and 4.5 (3.8–5.2, U-ZZ).
29	Sensitivity tests show that $TNH_x$ (total ammonium (gas + aerosol)), followed by total sulfate, were the
30	important factors that influenced the predicted pH. Generally, particle pH increased with the cation
31	increasing and the decrease in the anion, temperature, and relative humidity. Similar pH values (~3.0)
32	at the required $NH_x$ concentrations for the five sites indicated that the presence of excess $NH_x$ was
33	likely important for the less acidic of PM <sub>2.5</sub> during the severe haze episodes in this region. Moreover,
34	the concentrations of excess NH <sub>x</sub> may drive the higher pH values at rural sites, because of the higher
35	excess NH <sub>x</sub> concentrations at rural sites than those at urban sites. The underlying influence of regional
36	transport on local particle pH cannot be neglected by differing the chemical components of PM2.5 and
37	meteorological conditions. Air masses transported from rural and agricultural regions may enhance the
38	particle pH value in urban aerosols given the high pH of particles and high ammonia levels. These
39	results suggest that ammonia is urgently needed to be involved in the regional strategy for the
40	improvement of air quality in China.

41 Keywords: ISORROPIA-II model, Particle acidity, Ammonia, Sensitivity test, Regional transport.

#### 42 1 Introduction

43 High concentrations of acids and bases contained in the aqueous phase define the acidity of aerosols (Spurny, 1990). Particle acidity or pH value is an important parameter for atmospheric 44 45 particulate chemistry, such as the gas-particle portioning of semi-volatile and volatile species (e.g.,  $NH_{3(g)}/NH_{4(aq)}^+$ ,  $HCl_{(g)}/Cl_{(aq)}^-$ , and  $HNO_{3(g)}/NO_{3(aq)}^-$ ), the formation of secondary inorganic and organic 46 47 aerosols, and the dissolution of metallic elements (Bougiatioti et al., 2016; Meskhidze et al., 2003; 48 Seinfeld and Pandis, 2006; Shi et al., 2019; Shi et al., 2010; Surratt et al., 2010; Wang et al., 2018b). 49 Particle acidity can affect the ecosystem through its influence on wet/dry deposition, atmospheric 50 visibility, and radiative balance (Boucher and Anderson, 1995; Larssen et al., 2006; Pye et al., 2019; 51 Watson, 2002). In addition, high particle acidity has an adverse impact on public health, especially for 52 the cardiopulmonary and respiration system of humans (Dockery et al., 1996; Ostro et al., 1991).

53 Direct measurements on particle pH are challenging because of the small size and nonideality of 54 chemical species in solvated aerosols. Therefore, thermodynamic models, such as E-AIM 55 (http://www.aim.env.uea.ac.uk/aim/aim.php) and ISORROPIA-II (http://isorropia.eas.gatech.edu) 56 (Clegg et al., 1998; Nenes et al., 1998), which rely on the measurements of particulate and gaseous 57 species, are widely used in estimating particle pH. Generally, the global distribution of fine particulate 58 matter (PM<sub>2.5</sub>, aerodynamic diameter  $\leq 2.5 \,\mu$ m) pH is bimodal with a population of particles having a 59 mean pH of 1–3 and another population having an average pH closer to 4–5 (Pye et al., 2019). Table 60 1 shows that PM<sub>2.5</sub> in mainland China were moderately acidic with pH values that ranged from 3.4-61 5.7 (Ding et al., 2019; Guo et al., 2017; Liu et al., 2017; Shi et al., 2017, 2019; Song et al., 2018; Wang 62 et al., 2019a), which were 3-5 units higher than those reported in other regions, such as Hong Kong,

63 Singapore, USA, and Greece (Behera et al., 2013; Bougiatioti et al., 2016; Guo et al., 2016; Guo et al.,
64 2014; Pathak et al., 2004).

65	High atmospheric ammonia (TNH <sub>x</sub> , gas NH <sub>3</sub> plus particle $NH_4^+$ ) is a dominant factor that drives
66	the high PM <sub>2.5</sub> pH in megacities of China because it suppresses the production of particle hydronium
67	(Cheng et al., 2016; Wang et al., 2016). Liu et al. (2017) argued that excess NH <sub>3</sub> and elevated aerosol
68	water content (AWC) were responsible for the high pH in Beijing. Nevertheless, Guo et al. (2017) and
69	Song et al. (2018) demonstrated that high levels of ammonia did not increase the PM <sub>2.5</sub> pH into a fully
70	neutralized condition in Beijing and Xi'an, China. Weber et al. (2016) calculated that only large
71	increases in NH3 together with sulfate reductions can lead to an increase in pH. Correspondingly,
72	higher pH by about 1 unit than urban aerosols was observed in locations of intensive agriculture with
73	high NH <sub>3</sub> concentrations in the southeastern US (Nah et al., 2018). The primary sources for ammonia
74	include agricultural emissions, such as livestock waste, N-fertilizer application, and biomass burning,
75	as well as traffic and industrial emissions (Huang et al., 2012; Shen et al., 2011; Wang et al., 2018a).
76	Field studies have reported that elevated $NH_3$ and $NH_4^+$ concentrations were recorded at rural sites in
77	the North China Plain (Meng et al., 2011; Meng et al., 2017; Shen et al., 2011; Wen et al., 2015).
78	Therefore, studying in an agriculturally developed region is needed to obtain insight into the role of
79	ammonia in particle pH. Unfortunately, previous studies mainly concentrated on calculating the
80	particle pH in the megacity of China, and few studies focused on the agricultural regions of China. In
81	addition to ammonia, particle pH can also be influenced by other chemical compositions and
82	meteorological conditions, such as sulfate, AWC, temperature (T), and relative humidity (RH). Sulfate
83	is often the main acid component of aerosols, and largely determines the acidity of $PM_{2.5}$ (Weber et al., 4

2016). Sensitivity tests in Beijing suggested that sulfate,  $TNH_x$ , and T were the common driving factors, and  $Ca^{2+}$  and RH were the unique factors in special seasons (Ding et al., 2019). The pH sensitivity based on the 10-year record in Canada showed that chemical compositions had various effects on particle pH under different meteorological conditions; moreover, careful examination for any particular region is needed (Tao and Murphy, 2019).

89 Henan Province is situated in Central China; it has a dense rural population and is a top-ranking 90 province in China in terms of agricultural production and chemical fertilizer consumption (NBS, 2016). 91 NH<sub>3</sub> emission inventory for Henan Province reported that approximately 1031.6 Gg NH<sub>3</sub> was released 92 in Henan in 2015, thereby contributing to approximately 10 % of China's total emissions 93 (approximately 10 Tg) (Huang et al., 2012; Wang et al., 2018a). Livestock waste and N-fertilizer 94 application were major sources for ammonia emissions, which may increase TNH<sub>x</sub> concentrations at 95 rural sites than in urban sites. Furthermore, Henan Province is a severely PM<sub>2.5</sub>-polluted region in 96 China. In January 2018, a large-scale and long-lasting haze episode that caused PM<sub>2.5</sub> concentration to 97 reach 400 µg/m<sup>3</sup> occurred in this region (Wang et al., 2019c). An experiment was performed in two 98 urban and three rural sites in Henan Province to investigate the particle acidity and its driving factor. 99 ISORROPIA-II model was utilized to estimate PM<sub>2.5</sub> pH using a high-time-resolution dataset. The novelty of the work addressed in the present study is that this study is the first on PM<sub>2.5</sub> acidity by 100 101 comparing urban and rural sites in the agricultural regions of China. The specific objectives of the 102 study were presented as follows: (1) Estimation and comparison of the PM<sub>2.5</sub> pH at the five monitoring 103 sites, (2) identification of the factors that determine the pH, and (3) discussion of the role of ammonia 104 on pH. Our results are helpful to understand the factors that determine particle acidity better.

#### 105 **2** Experiment and methods

#### 106 **2.1 Site descriptions**

107 Field sampling was synchronously conducted from January 12 to 24, 2018 at five sites (i.e., two 108 urban sites located in the center of Zhengzhou (U-ZZ) and Anyang (U-AY), and three sites located in 109 the rural areas of Anyang (R-AY), Xinxiang (R-XX), and Puyang (R-PY)). Locations of the five sites 110 (Fig. S1) with brief descriptions are listed in Table 2. U-ZZ site is surrounded by busy roads, and two 111 freeways are located 3 km to the south and 7 km to the east. Moreover, this site is near a coal-fired 112 power plant; a gas-fired power plant; and several small-scale industries, such as pharmaceutical 113 companies, electronic factories, and equipment manufacturing. U-AY site is surrounded by busy roads, 114 and Anyang steelwork is located 8 km to the west. R-AY site is surrounded by farmland and is 1 km 115 west of the Jing-Gang-Ao freeway and 1 km north of a belt freeway. R-XX and R-PY sites are 116 surrounded by farmland without other prominent anthropogenic emission sources.

#### 117 **2.2 Instrumentations**

118 The hourly mass concentrations of water-soluble inorganic ions (WSIIs) in PM<sub>2.5</sub>, such as NH<sub>4</sub><sup>+</sup>, SO<sub>4</sub><sup>2-</sup>, NO<sub>3</sub><sup>-</sup>, Cl<sup>-</sup>, Na<sup>+</sup>, Mg<sup>2+</sup>, Ca<sup>2+</sup>, and K<sup>+</sup> and their gaseous precursors (i.e., NH<sub>3</sub>, HNO<sub>3</sub>, and HCl) 119 120 were measured using an ambient ion monitor (URG-9000D, Thermal Fisher Scientific, USA) in U-ZZ site and the monitor for aerosols and gases (MARGA, Metrohm, Switzerland) in other sites. Both 121 122 instruments have been successfully deployed in several other field experiments (Li et al., 2017; Shi et 123 al., 2017; Wang 2019b), and detailed information is available elsewhere (Markovic et al., 2012; 124 Rumsey et al., 2014). As a brief summary, ambient air is drawn into the systems at a flow rate of 16.7 125 L/min. Subsequently, particles and gases are collected by two aerosol sample collectors through a wet 126 parallel plate or wet rotating denuder. Aqueous solution samples are quantified by using two ion 127 chromatography analyzers. Detection limits for all species during the sampling periods were less than 0.1 µg/m<sup>3</sup>. Measurement uncertainties were assumed to 10 % for URG-9000D here by reference to 128 129 10 % of Markovic et al. (2012) and 12 % in Hu et al. (2014). For the MARGA instrument, Song et al. 130 (2018) took 20 % as overall relative uncertainties for major species, and Rumsey et al. (2014) reported that the analytical biases were lower than 10 % for SO<sub>4</sub><sup>2-</sup>, NO<sub>3</sub><sup>-</sup>, and HNO<sub>3</sub>, but higher than 15 % for 131 NH<sub>3</sub> and NH<sub>4</sub><sup>+</sup>. Therefore, overall uncertainties were estimated to be 20 % for NH<sub>3</sub> and NH<sub>4</sub><sup>+</sup> and 10 % 132 133 for other species. The combined uncertainties for TNH<sub>x</sub> were calculated to be 14 % and 23 % for U-ZZ and other sites, respectively, and 14 % uncertainties of TNO<sub>3</sub> (NO<sub>3(aq)</sub><sup>-</sup> + HNO<sub>3(g)</sub>) and TCl (Cl<sub>(aq)</sub><sup>-</sup> 134 + HCl<sub>(g)</sub>) for all sites. Because of the complex interference during sampling periods at the five sites, 135 136 such as the slope from the internal calibration, the solution volume, temperature, pressure, and airflow 137 rate, the actual uncertainties might be different from the estimated value. Hourly elemental carbon (EC) 138 and organic carbon (OC) concentrations in PM2.5 were simultaneously obtained by the semi-139 continuous carbon analyzers (Model 4, Sunset Laboratory Inc, USA) in the same place, except for the 140 U-ZZ site, of which data were provided by the Department of Environmental Protection of Henan 141 Province. Detailed information on this instrument can be found in Panteliadis et al. (2014). Briefly, 142 PM<sub>2.5</sub> collected by the device was oxidized to carbon dioxide and analyzed by a nondispersive infrared 143 detector. The relative measurement uncertainty for this instrument was estimated to be 20 % (Liu et 144 al., 2013; Healy et al., 2013). Meteorological parameters, including T, RH, wind direction, and wind 145 speed, were recorded by smart weather sensor (LUFFT-WS500, Sutron Corporation, Germany), with the accuracies of  $\pm$  0.2 °C,  $\pm$  2 %, < 3 °, and 0.1 m/s, respectively. 146

### 147 **2.3 pH prediction**

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Particle pH was estimated using the ISORROPIA-II thermodynamic model. Input data, including 148 RH, T, concentrations of  $K^+$ ,  $Ca^{2+}$ ,  $Mg^{2+}$ ,  $TNH_x$ , total sulfate (TH<sub>2</sub>SO<sub>4</sub>, replaced by observed SO<sub>4</sub><sup>2-</sup>), 149 150 total sodium (TNa, replaced by observed Na<sup>+</sup>), TCl, and TNO<sub>3</sub>, were used to calculate the particle 151 hydronium ion concentration per volume of air (H<sup>+</sup><sub>air</sub>) and particle water associated with inorganics (AWC<sub>inorg</sub>) by computing the equilibrium composition for the Na<sup>+</sup>-K<sup>+</sup>-Ca<sup>2+</sup>-Mg<sup>2+</sup>-NH<sub>4</sub><sup>+</sup>-SO<sub>4</sub><sup>2-</sup>-NO<sub>3</sub><sup>-</sup>-152 153 Cl<sup>-</sup>-H<sub>2</sub>O aerosol system. Considering that forward mode is less sensitive to measurement error than 154 the reverse mode and high RH levels were recorded in sampling periods, ISORROPIA-II was run in 155 the forward model for the aerosol system in the metastable condition (Ding et al, 2019; Guo et al., 156 2015; Hennigan et al., 2015). Moreover, we excluded data with RH < 30 %, a condition where predicted to observed partitioning of nitrate between the gas and particle phase are in weak correlation, 157 158 and thus uncertainties in the pH prediction are expected to be high (Ding et al., 2019; Guo et al., 2016). 159 Aerosol pH was calculated according to the formula (Bougiatioti et al., 2016):

$$\mathrm{pH}=-\log_{10}H_{aq}^{+}=-\log_{10}rac{1000H_{air}^{+}}{AWC_{inorg}+AWC_{org}},$$
 (1)

161 where the modeled concentrations for  $AWC_{inorg}$  and  $H^+_{air}$  are  $\mu g/m^3$ , and  $AWC_{org}$  is the particle water 162 associated with the organics predicted using the method:

163 
$$AWC_{org} = \frac{m_s}{\rho_s} \frac{k_{org}}{\left(\frac{1}{\text{RH}} - 1\right)},$$
 (2)

where  $m_s$  is the mass concentrations of organic matter (OC × 1.6),  $\rho_s$  is the organic density (1.35 g/cm<sup>3</sup>), and  $k_{org}$  is the organic hygroscopicity parameter (0.06) (Liu et al., 2017).

#### 166 **2.4 pH uncertainty**

167 Two extreme scenarios were evaluated to estimate the pH uncertainty based on the measurement 168 uncertainties of input data (Murphy et al., 2017). Sensitivity tests of pH to input data (see Section 3.3) suggest that simulated pH increase with the cation concentrations (i.e.,  $TNH_x$ , TNa,  $K^+$ ,  $Ca^{2+}$ , and  $Mg^{2+}$ ) 169 170 increasing as well as the anion concentrations (i.e., TH<sub>2</sub>SO<sub>4</sub>, TNO<sub>3</sub>, and TCl), T and RH decreasing. 171 Moreover, the pH value has a positive correlation with OC concentrations via Eqs. (1 and 2). Therefore, 172 cation and OC concentrations were adjusted up to within their maximum positive uncertainties, and 173 anions, RH and T were adjusted down within their maximum negative uncertainties, which represented 174 the pH<sub>max</sub> case; for pH<sub>min</sub> case, cations and OC concentrations were adjusted down, and anions, RH 175 and T were adjusted up. Figure 1 suggests that  $pH_{max}$  cases lead to 6 % (slope - 1) errors for all sites, 176 and pH<sub>min</sub> cases can result in approximately 10 % deviations. Thus, pH uncertainties were set to be -9-6 %, -13-6 %, -12-6 %, -11-6 %, and -10-5 % for U-ZZ, U-AY, R-AY, R-XX, and R-PY sites, 177 178 respectively.

## 179 **2.5 Model validation**

The reliability of pH estimation using the ISORROPIA-II thermodynamic model depends on several assumptions, such as the equilibrated gas and particle phases (Pye et al., 2019). Thus, the predicted and observed semi-volatile species (e.g.,  $NH_{3(g)}/NH_{4(aq)}^+$ ,  $HCl_{(g)}/Cl_{(aq)}^-$ , and  $HNO_{3(g)}/NO_{3(aq)}^-$ ) are compared in Fig. S2. Observed and predicted  $NH_4^+$ ,  $Cl^-$ , and  $NO_3^-$  exhibit significant correlations, have correlation coefficients (*r*) above 0.95 and slopes near 1 at the five sites. In addition,  $NH_3$ concentrations are in good agreement (r > 0.95) with slopes between 0.89 (U-ZZ) and 1.13 (R-PY). These results suggest the excellent performance of ISORROPIA-II for modeling these species. The 187 differences in the slopes of NH<sub>4</sub><sup>+</sup> (Fig. S2b) and NH<sub>3</sub> (Fig. S2e) between U-ZZ and R-PY sites were 188 probably attributed to the unbalanced charge of input WSIIs (see Text S1 for more details of calculation) 189 with average equivalent ratios (Fig. S3) of  $0.99 \pm 0.13$  (U-ZZ) and  $1.20 \pm 0.12$  (R-PY). However, Song 190 et al. (2018) argued that ion balance was not a key factor for pH calculation in the forward mode, 191 because the forward mode calculations account for additional constraints imposed by the partitioning of semi-volatile species. To verify this, we adjusted the measured NH4<sup>+</sup> concentration, which was most 192 193 sensitive to pH modeling (see Section 3.3), to fit the ion balance, and reran the pH calculation. As 194 shown in Fig. S4, re-predicted NH<sub>4</sub><sup>+</sup> and NH<sub>3</sub> concentrations have better performance (r = 0.99, 0.94195 < slopes < 1.06) with input concentrations at two sites than before. However, re-calculated pH values 196 change slightly, and these deviations are included in the range of pH uncertainty discussed in Section 197 2.4. Therefore, the little differences in NH<sub>x</sub> partitioning between the model and measurement among 198 the five sites were acceptable for pH calculation.

199 Correlations between the predicted and measured HNO<sub>3</sub> and HCl are weak. Similar problems 200 were found in the northeast U.S. and Beijing. These discrepancies were potentially due to measurement 201 uncertainties brought about by low gas concentrations, the interference of coarse-mode particles, non-202 volatile cation measurement artifacts, uncertainties in the thermodynamic constants, and kinetic 203 limitations to mass transfer (Ding et al., 2019, Haskins et al., 2018; Pye et al., 2019; Liu et al., 2017). 204 Pye et al. (2019) suggested that ISORROPIA-II yields a mean activity coefficient of  $(H^+, Cl^-)$  that may 205 result in the higher predicted HCl concentration. HNO<sub>3</sub> can be partition to both fine and coarse modes, 206 thereby affecting predicted fine-mode nitrate concentrations (Nah et al., 2018). The best semi-volatile 207 species for evaluation of pH modeling depend on the fraction of the gas phase. In this work, most of HNO<sub>3</sub> and HCl concentrated in the particle phase, and thus they are not suited to test the model (Guo
et al., 2016).

The PM<sub>2.5</sub> pH was also calculated by the E-AIM (Version IV) model to evaluate the performance of ISORROPIA-II using the observed data (RH > 60 %) of the U-ZZ site as an example. Close correlation (r = 0.89) is found between two models with a slope of 0.95 (Fig. S5). The pH values in ISORROPIA-II are 0.46 ± 0.15 units higher than those in E-AIM. These values are comparable to the founding by Liu et al. (2017) and Song et al. (2018), which is possibly due to the differences of activity coefficient values between the ISORROPIA-II and E-AIM models (Pye et al., 2019). Overall, the predicted pH values using ISORROPIA-II are effective in this work.

#### 217 **3 Results and discussion**

#### 218 **3.1 Haze episodes**

219 During the sampling periods, five monitoring sites simultaneously experienced a long-lasting and large-scale haze episode. Time series of the concentrations of major species (i.e., NH<sub>3</sub>, NH<sub>4</sub><sup>+</sup>, SO<sub>4</sub><sup>2-</sup>, 220 221 and NO<sub>3</sub><sup>-</sup>) and meteorological parameters at the U-ZZ site as an example are presented in Fig. 2, and 222 other sites are integrated into Fig S6 with the mean values listed in Table 3. Three study cases were 223 classified on the basis of similar meteorological conditions and WSIIs levels at the five sites. The 224 WSIIs concentration, T, and RH gradually increased in the southern wind during Case 1 (January 225 12-14). Case 3 (January 21-25) was characterized by decreased WSII concentrations, T, and RH with 226 the northeastern wind. Even though the total WSIIs (TWSIIs) concentrations were comparable, the 227 chemical components of WSIIs were various between Cases 1 and 3 (Table 3). In particular, NO<sub>3</sub><sup>-</sup>, NH4<sup>+</sup>, and NH<sub>3</sub> concentrations in Case 1 were higher than those in Case 3 at all sites. Conversely, 228

229	$SO_4^{2-}$ concentrations in Case 1 were lower than those in Case 3. Wang et al. (2018a) reported that the
230	southern cities of Henan Province (e.g., Nanyang, Shangqiu, Zhoukou, and Zhumadian) had relatively
231	higher ammonia emissions than the cities involved in this study. Moreover, the northeastern air masses
232	from the Jing–Jin–Ji regions were easily enriched with sulfate (Wang et al., 2019b; Wang et al., 2018c).
233	Evidently, back trajectory frequency analysis (see Text S2 for more details of the discussion) confirmed
234	that sampling sites during Cases 1 and 3 (Figs. S7a and c) were predominantly influenced by the
235	southern and northeastern air masses, respectively.
236	Elevated WSIIs concentrations during Case 2 (January 14-21) were under high T and RH
237	conditions with variable wind directions. In this case, local emissions played a key role in WSIIs (Fig.
238	S7b) with average NH <sub>4</sub> <sup>+</sup> , SO <sub>4</sub> <sup>2-</sup> , and NO <sub>3</sub> <sup>-</sup> concentrations that ranged from $31.9 \pm 12.5 \ \mu g/m^3 (U-ZZ)$
239	to $47.6 \pm 13.1 \ \mu\text{g/m}^3$ (R-AY), $32.9 \pm 12.5 \ \mu\text{g/m}^3$ (R-XX) to $46.8 \pm 16.9 \ \mu\text{g/m}^3$ (R-AY), and $58.0 \pm 18.3 \ \mu\text{g/m}^3$ (R-AY), $46.8 \pm 16.9 \ \mu\text{g/m}^3$ (R-A
240	$\mu$ g/m <sup>3</sup> (U-AY) to 76.8 $\pm$ 21.1 $\mu$ g/m <sup>3</sup> (R-AY), respectively. Note that higher concentrations of NH <sub>4</sub> <sup>+</sup> ,
241	$\rm NH_3,$ and $\rm TNH_x$ during Case 2 were recorded at rural sites than those at urban sites. Moreover, $\rm NH_3$
242	levels in this work were higher than those in other studies summarized in Table S1. Agricultural
243	emissions, including livestock waste, N-fertilizer application, and humans, were the top three ammonia
244	contributors in Henan Province (Wang et al., 2018a), which may result in elevated ammonia
245	concentrations at rural sites.

246 **3.2 pH of PM<sub>2.5</sub> at the urban and rural sites** 

Figure 3 exhibits the predicted  $PM_{2.5}$  pH values,  $H^+_{air}$ , and AWC at the five sites, pH uncertainties based on the pH<sub>max</sub> and pH<sub>min</sub> calculations are presented in Fig. S8. PM<sub>2.5</sub> have consistent moderate acidity during this haze episode, with median (min–max) pH values of 4.5 (3.8–5.2), 4.8 (3.9–5.8), 4.9 (4.1–6.2), 5.1 (4.7–6.3), and 5.2 (4.8–6.5,) at U-ZZ, U-AY, R-XX, R-AY, and R-PY, respectively. As
summarized in Table 1, PM<sub>2.5</sub> pH values were close to the results of other cities in China (e.g., Beijing,
Xi'an, and Tianjin), but higher than other countries (e.g., Singapore, USA, and Greece). Compared to
the rural sites in the USA, pH values were roughly 3 units higher at the R-AY, R-XX, and R-PY sites.
Moreover, statistical values of pH (Table S2) during three cases show higher pH values at rural sites
than those at urban sites, especially for the U-ZZ and R-PY sites during Case 1 despite their pH
uncertainties (Section 2.4).

257 The pH values (Table S2) of the five sites during Cases 2 and 3 were comparable but slightly 258 lower than those during Case 1. As shown in Fig. S9, predicted pH values present better correlations 259 with H<sup>+</sup><sub>air</sub> concentrations rather than AWC. Particle hydronium ion aqueous concentration depends on 260 both the presence of ions and the amount of particle AWC (Guo et al., 2015). Moreover, H<sup>+</sup><sub>air</sub> was 261 closely associated with the NH<sub>3</sub> mixing ratios, and higher NH<sub>3</sub> always corresponded to lower H<sup>+</sup><sub>air</sub> (Liu et al., 2017). High correlations (r > 0.5) between H<sup>+</sup><sub>air</sub> and TWSIIs (Table S3) imply that local pH was 262 263 significantly affected by the TWSIIs levels, and probably resulting in the high acidity during Case 2. 264 H<sup>+</sup><sub>air</sub> was also correlated with individual chemical species (e.g., TNH<sub>x</sub>, TH<sub>2</sub>SO<sub>4</sub>, TNO<sub>3</sub>, and TCl). 265 Therefore, the diversity of pH in Cases 1 and 3 may be partly owing to the different proportions of 266 particle- and gas-phase constituents that will be discussed in detail below. In addition, high correlations 267 between RH and H<sup>+</sup><sub>air</sub> (Table S3) suggest the major role of meteorological conditions in particle pH. 268 Given the Case 2 was less affected by regional transport, the diurnal patterns of median pH values of 269 the five sites (Fig. 4) indicate that pH values during nighttime were 0.3 (R-PY)-0.5 (U-ZZ) units higher 270 than those during the daytime. Similar results were also found in other cities (e.g., Beijing, Tianjin,

271 southern Canada, and the USA) (Battaglia et al., 2017; Ding et al., 2019; Guo et al., 2015; Murphy et 272 al., 2017; Shi et al., 2019), resulting from the diurnal trends of T and RH (Fig. 4f). After sunrise, high 273 T facilitated the dissociation of particle-phase ammonium (e.g., NH<sub>4</sub>NO<sub>3</sub>) and led to a rapid loss of 274 AWC (Guo et al., 2015; Saraswati et al., 2019).

#### 275 3.3 Sensitivity tests of pH

276 To explore the dominant factors that determine the local particle pH level and resulting in the high pH at rural sites, sensitivity tests of pH to chemical species (i.e., TNH<sub>x</sub>, TH<sub>2</sub>SO<sub>4</sub>, TNO<sub>3</sub>, TCl, TNa, K<sup>+</sup>, 277 Ca<sup>2+</sup>, and Mg<sup>2+</sup>) and meteorological parameters (i.e., T and RH) were performed. Firstly, the real-time 278 279 measured values of a variable (e.g., TNH<sub>x</sub>) and average values of other parameters (i.e., TH<sub>2</sub>SO<sub>4</sub>, TNO<sub>3</sub>, TCl, TNa, K<sup>+</sup>, Ca<sup>2+</sup>, Mg<sup>2+</sup>, T and RH) during Case 2 were input into ISORROPIA-II to investigate the 280 281 sensitivity of local pH to this variable (i.e., TNH<sub>x</sub>) (Ding et al., 2019), and results are listed in Table 4 282 and Fig. S10. In addition, a given range for a variable for all sites with corresponding average values of other parameters was simulated to compare its effects on pH among five sites (Figs. 5 and S11). The 283 284 chosen variation range for each variable was close to the observed minimum and maximum values (Table S4), which aims to better reflect the actual observation conditions. The degree of sensitivity 285 286 was represented as the relative standard deviation (%RSD) of the re-calculated pH values. RSD 287 calculates the absolute value of the coefficient of variation, which helps us to determine how small or large is the standard deviation when compared to the mean of the re-calculated pH data set. Therefore, 288 289 when the re-calculated pH data set has a higher RSD value, this variable can change the pH easier and 290 is more important for local pH.

291 As shown in Table 4, the most important factor that influenced particle pH during Case 2 was

292	$TNH_x$ , followed by $TH_2SO_4$ at the five sites, and the U-ZZ site was also affected by $TNO_3$ . The acidity
293	of $PM_{2.5}$ is governed by the phase partitioning of semi-volatile gases such as $NH_3$ , $HNO_3$ , and $HCl$
294	(Pye et al., 2019). (NH <sub>4</sub> ) <sub>2</sub> SO <sub>4</sub> , NH <sub>4</sub> HSO <sub>4</sub> , and NH <sub>4</sub> NO <sub>3</sub> , which are mainly formed through
295	neutralization reactions between $NH_3$ with $SO_4^{2-}$ and $HNO_3$ orderly according to the regime of local
296	ammonia, are the most abundant components of PM <sub>2.5</sub> in winter haze in this region (Wang et al., 2019c;
297	Pathak et al., 2008). Therefore, particle pH was driven by TNH <sub>x</sub> , TH <sub>2</sub> SO <sub>4</sub> , and TNO <sub>3</sub> . TNa, TCl, and
298	crustal ions (i.e., $K^+$ , $Ca^{2+}$ , and $Mg^{2+}$ ) have less influence on the predicted pH values, because these
299	species were at low concentrations, together accounting for lower than 10 % of TWSIIs. However,
300	since the low volatility of these cations allows them to preferentially neutralize sulfates over NH <sub>3</sub> , the
301	role of crustal dust and sea spray in particle pH cannot be ignored when the mass fraction of these ions
302	are high in typical pollution events (e.g., sandstorm) or areas (e.g., coastland) (Allen et al., 2015; Guo
303	et al., 2018; Vasilakos et al., 2018). In addition to chemical species, T was more sensitive to local pH
304	than RH at the five sites, which is similar to the finding in Beijing and Canada (Ding et al., 2019; Tao
305	and Murphy, 2019). This result can be explained by the remarkable influence of T on the solubility and
306	dissociation constants of the partitioning of $NH_{3(g)}/NH_{4(aq)}^{+}$ (Hennigan et al., 2015).
307	Figures 5 and S11 compare the sensitive degrees of pH to input data among different sites. Except

for TNH<sub>x</sub>, the sensitivities of pH to TH<sub>2</sub>SO<sub>4</sub>, TNO<sub>3</sub>, and other ions at urban sites were more significant than those at rural sites, particularly at U-ZZ sites of 7.2 % and 14.8 % of RSD to TH<sub>2</sub>SO<sub>4</sub> and TNO<sub>3</sub>, respectively. To gain insight into the differences of pH sensitivity among the five sites, sensitivity tests of pH to TH<sub>2</sub>SO<sub>4</sub>, TNO<sub>3</sub>, and TNH<sub>x</sub> were calculated using the fixed TNO<sub>3</sub> (67.5  $\mu$ g/m<sup>3</sup>) and TH<sub>2</sub>SO<sub>4</sub> (36.5  $\mu$ g/m<sup>3</sup>) concentrations under the average meteorological conditions of five sites (i.e., T = 2.5 °C

313	and $RH = 60$ %). These chosen values were close to the average values of the five sites during Case 2
314	(i.e., $36.4 \pm 15.4 \ \mu g/m^3$ for $SO_4^{2-}$ , $67.5 \pm 23.5 \ \mu g/m^3$ for $TNO_3$ , $2.5 \pm 1.5$ °C for T, and $59.3 \pm 14.0 \ \%$
315	for RH), and other chemical species were set to be zero. As shown in Fig. 6, sensitivities of pH to
316	$\mathrm{TH}_2\mathrm{SO}_4$ and $\mathrm{TNO}_3$ increase with the decrease in $\mathrm{TNH}_x$ concentration, particularly when the $\mathrm{TNH}_x$
317	concentrations are lower than 60 $\mu$ g/m <sup>3</sup> and 40 $\mu$ g/m <sup>3</sup> , respectively. Therefore, the pH changes tended
318	to become more sensitive to $TH_2SO_4$ and $TNO_3$ at the U-ZZ site (Fig. 5e, f) with the lowest $TNH_x$
319	concentrations (46.8 $\pm$ 14.7 $\mu\text{g/m}^3$ ). All these results may be explained by the presence of excess
320	ammonia that will be discussed below.
321	Sensitivity tests suggest (Fig. 5, S10, and S11) that particle pH gradually grows with increased

322 cation and decreased anion concentrations. Specifically, the TNH<sub>x</sub> concentration that increased from 25  $\mu$ g/m<sup>3</sup> to 90  $\mu$ g/m<sup>3</sup> can promote particle pH by 3.5 (U-ZZ)-4.5 (R-AY) units. The crustal cations 323 324 and sea spray have limited effects on pH, but a 10-fold increase in these species still can increase predicted pH values by about 0.1 units. TH<sub>2</sub>SO<sub>4</sub>, and TNO<sub>3</sub> that increased from 10  $\mu$ g/m<sup>3</sup> to 80  $\mu$ g/m<sup>3</sup>, 325 and 1  $\mu$ g/m<sup>3</sup> to 125  $\mu$ g/m<sup>3</sup> can reduce the pH values by 1.5 (R-PY)-4.0 (U-ZZ), and 0.2 (R-AY)-1.4 326 (U-ZZ) units, respectively. In addition, a 20 °C (-5 °C to 15 °C) and 65 % (30 % to 95 %) increase 327 328 drops the pH by approximately 1.3 and 2.7 units at the five sites, respectively. Corresponding to the 329 observed data during Case 2 (Table 3), lower TNH<sub>x</sub> concentrations and higher T values were probably 330 responsible for the higher acidity at urban sites than those at rural sites. Moreover, Figure 6 shows that 331 particle pH increase with decreasing TH<sub>2</sub>SO<sub>4</sub> (Fig. 6a) or increasing TNO<sub>3</sub> (Fig. 6b), particularly when the TNH<sub>x</sub> is at high concentration. These results indicate that predicted pH will be enhanced by 332 increasing the TNO<sub>3</sub>/TH<sub>2</sub>SO<sub>4</sub> ratio, which is consistent with the observation in Beijing (Xie et al., 333

334 2019). All these may be due to the elevated ammonia partitioning with nitrate fraction increase, and 335 nitrate-rich particles would absorb more water compared to the sulfate-rich particles (Xie et al., 2019). 336 Based on the above, higher TNH<sub>x</sub> (Table 3) and TNO<sub>3</sub>/TH<sub>2</sub>SO<sub>4</sub> ratios ( $2.9 \pm 0.6$  and  $1.7 \pm 0.6$  for Cases 337 1 and 3 respectively) during Case 1 may lead to the higher pH values than those during Case 3, even 338 though the TWSIIs levels were comparable. In the long run, the decreasing sulfate concentration in 339 PM<sub>2.5</sub> accompanied with increasing nitrate concentration has been recorded during haze episodes in 340 China, because strong actions were taken to reduce the coal consumption in recent years (Tian et al., 341 2017; Wang et al., 2017). Considering the ammonia in the atmosphere of North China might still be 342 increasing (Liu et al., 2018), future acidity of PM2.5 during winter haze episodes is expected to become 343 increasingly less acidic in this area. A more careful analysis is needed to test this inference, however, 344 because fine-mode particle remains a relatively constant pH of 0-2 despite a 70 % reduction in sulfates 345 in the USA (Vasilakos et al., 2018; Weber et al., 2016).

#### 346 **3.4 Role of ammonia on pH**

347 It has been suggested that unusually high levels of NH<sub>3</sub> can increase pH, on average, a 5-fold to 348 10-fold increase in the NH<sub>3</sub> levels leads to a one-unit change in pH (Nah et al., 2018; Weber et al., 349 2016; Guo et al., 2017). To examine the effects of major indicators of ammonia (i.e., TNH<sub>x</sub>, Required-350 NH<sub>x</sub>, and Excess-NH<sub>x</sub>, see Text S3 for more details of calculation) on aerosol acidity, particle pH was calculated by using a wide range of  $TNH_x$  (25–130  $\mu$ g/m<sup>3</sup>) and average values of other parameters 351 352 during Case 2 (Table 3) of each site. Simultaneously, the concentrations of TNH<sub>x</sub>, Required-NH<sub>x</sub>, 353 Excess-NH<sub>x</sub>, and corresponding pH values are illustrated in Fig. 7. An "S-curve" growth trend of pH 354 to increasing TNH<sub>x</sub> appears with the inflection point around the Required-NH<sub>x</sub> concentration. In the

355	case when the input $\text{TNH}_x$ concentration is lower than the Required-NH <sub>x</sub> , the growth rate of pH
356	increase with TNH <sub>x</sub> increasing, otherwise, the pH growth flattens out, which is similar to the findings
357	of Weber et al. (2016) and Bougiatioti et al. (2016). In the process of increasing NH <sub>3</sub> concentration,
358	$\mathrm{NH_3}$ reacted with $\mathrm{SO_4}^{2-}$ and $\mathrm{HNO_3}$ orderly, during when large amounts of $\mathrm{H^+}$ were consumed and pH
359	values rapidly increased. Theoretically, the Required-NH <sub>x</sub> plus nonvolatile cations can neutralize all
360	anions regardless of the phase partitioning and bisulfate formation, and thus changes in $\text{TNH}_x$ around
361	the Required-NH <sub>x</sub> concentrations have a significant impact on particle pH. Subsequently, dissolving
362	Excess-NH <sub>x</sub> into the particles became difficult, and pH values increased slowly (Ding et al., 2019;
363	Seinfeld and Pandis, 2016). Therefore, the underlying reason why TNH <sub>x</sub> concentrations lower than 60
364	$\mu g/m^3$ and 40 $\mu g/m^3$ elevated the pH sensitivities (Fig. 6) to $TH_2SO_4$ and $TNO_3$ is that the Excess-NH_x
365	will be exhausted under the case of $TH_2SO_4$ and $TNO_3$ that ranged from 10 $\mu$ g/m <sup>3</sup> to 100 $\mu$ g/m <sup>3</sup> ,
366	respectively. On the other hand, the pH values of the five sites at the Required- $NH_x$ concentrations are
367	near 3.0, which is in coincidence with the pH value of Beijing in the no-excess- $NH_x$ cases (Liu et al.,
368	2017). Therefore, the presence of Excess- $NH_x$ in the aerosol was likely important for the less acidic of
369	$PM_{2.5}$ and can enhance the pH values by 1.5 (U-ZZ)–2 (R-PY) units. Considering that the Excess-NH <sub>x</sub>
370	concentrations at rural sites (e.g., $30.1 \pm 6.2 \ \mu\text{g/m}^3$ at R-PY) were higher than those at urban sites (e.g.,
371	$14.8 \pm 4.1 \ \mu g/m^3$ at U-ZZ), and thus Excess-NH <sub>x</sub> concentrations may drive the pH values of rural sites
372	higher than those of urban sites.

373 **3.5 Implications of regional transport** 

From the above discussion, the differences of pH among three cases indicate that the underlying
 influence of regional transport on local particle pH cannot be neglected by differing the chemical 18

376	components of $PM_{2.5}$ and meteorological conditions. In particular, the median pH values (Table S2) of
377	Case 1, during when the air masses transported from the south of sampling regions, increased by 0.2–
378	0.9 units than those during Cases 2 and 3. Aqueous formations of sulfate are strongly dependent on
379	particle pH. Chen et al. (2016) reported that the aqueous-phase sulfate production rates from NO <sub>2</sub> and
380	O <sub>3</sub> oxidation of SO <sub>2</sub> had a positive correlation with particle pH during the Beijing haze events. When
381	pH exceeded approximately 4.5 (higher than this value at rural sites in this work), NO <sub>2</sub> -oxidation
382	dominated the sulfate formation, and its reaction rate increased by one order of magnitude with the
383	rise of pH by one unit. Thus, air masses transported from rural and agricultural regions may promote
384	the sulfate formation in urban aerosols. In addition to sulfate, higher aerosol pH favors partitioning of
385	TNO <sub>3</sub> toward aerosol NO <sub>3</sub> rather than gaseous HNO <sub>3</sub> and thus elevating the particle mass
386	concentration (Nenes et al., 2019; Weber et al., 2016). Sampling regions in this study are located in the
387	transport route for Beijing (MEP, 2017), thereby frequently affecting its local particle pH. The lifetimes
388	of NH <sub>3</sub> (1–5 days or less) and NH <sub>4</sub> <sup>+</sup> (1–15 days) in the atmosphere are sufficient for transporting to
389	Beijing during a haze episode (Aneja, 2000; Lefer et al., 1999; Warneck, 1988). Ding et al. (2019)
390	have also observed that PM2.5 pH from the southwest direction was generally higher than that from the
391	northern direction in Beijing. Therefore, the particle pH in Beijing may be enhanced when southern
392	air masses from this region accompany elevated-pH particles and high ammonia levels.

# 393 4 Conclusions

An experiment was performed using a series of high-time-resolution instruments in two urban (i.e., U-ZZ and U-AY) and three rural sites (i.e., R-AY, R-XX, and R-PY) in Henan Province during a large-scale and long-lasting haze episode. The ISORROPIA-II model was used to investigate the pH value and its driving factors. PM<sub>2.5</sub> exhibited moderate acidity with median pH values of 4.5 (3.8–5.2),
4.8 (3.9–5.8), 4.9 (4.1–6.2), 5.1 (4.7–6.3), and 5.2 (4.8–6.5,) at U-ZZ, U-AY, R-XX, R-AY, and R-PY,
respectively. The pH values at rural sites were slightly higher than those at urban sites.

400 The predicted pH values of PM<sub>2.5</sub> were significantly affected by the WSIIs levels, different 401 proportions of particle- and gas-phase constituents, and meteorological parameters. Sensitivity tests 402 show that TNH<sub>x</sub>, followed by TH<sub>2</sub>SO<sub>4</sub>, were the important factors that influenced the predicted pH at 403 the five sites. In addition, T was more sensitive to local pH than RH. Generally, particle pH rose with 404 the increase in cation and the decrease in the anion, T, and RH. Further study suggests that predicted 405 pH will be enhanced by increasing the TNO<sub>3</sub>/TH<sub>2</sub>SO<sub>4</sub> ratio. Therefore, elevated TNH<sub>x</sub> concentrations 406 and lower T were probably responsible for the higher pH values at rural sites than those at urban sites, 407 and higher TNH<sub>x</sub> and TNO<sub>3</sub>/TH<sub>2</sub>SO<sub>4</sub> ratios during Case 1 may lead to the higher pH values than those 408 during Case 3.

An "S-curve" growth trend of pH to increasing TNH<sub>x</sub> was found with the inflection point around the Required-NH<sub>x</sub> concentration. Moreover, the pH values of the five sites at the Required-NH<sub>x</sub> concentrations are near 3.0, and the presence of Excess-NH<sub>x</sub> in the aerosol can enhance the pH values by 1.5 (U-ZZ)–2 (R-PY) units, and thus Excess-NH<sub>x</sub> was likely important for the less acidic of PM<sub>2.5</sub> during the severe haze episodes in this region. Considering that the Excess-NH<sub>x</sub> concentrations at rural sites were higher than those at urban sites, Excess-NH<sub>x</sub> concentrations may drive the pH values of rural sites higher than those of urban sites.

The underlying influence of regional transport on local particle pH cannot be neglected by differing the chemical components of PM<sub>2.5</sub> and meteorological conditions. Air masses transported 20

418	from rural and agricultural regions with elevated pH particles and high ammonia levels may promote
419	the secondary particle formation in urban aerosols. Therefore, ammonia should be involved in the
420	regional strategy for improving the air quality in China.
421	
422	Data availability. All data in this work are available by contacting the corresponding author Shasha
423	Yin ( <u>shashayin@zzu.edu.cn</u> )
424	
425	Author contributions. Shasha Yin and Ruiqin Zhang designed and led this study. Shasha Yin was
426	responsible for all observations and data collection. Lingling Wang, Yuqing Li, Chen Wang, and Weisi
427	Wang interpreted the data and discussed the results. Shenbo Wang wrote the paper.
428	
429	Competing interests. The authors declare that they have no conflict of interest.
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434	References
435	Allen, H. M., Draper, D. C., Ayres, B. R., Ault, A., Bondy, A., Takahama, S., Modini, R. L., Baumann,
436	K., Edgerton, E., Knote, C., Laskin, A., Wang, B., and Fry, J. L.: Influence of crustal dust and sea
437	spray supermicron particle concentrations and acidity on inorganic NO3 <sup>-</sup> aerosol during the 2013
438	Southern Oxidant and Aerosol Study. Atmos. Chem. Phy., 15, 10669–10685, 2015.

439	Aneja, V. P., Chauhan, J. P., Walker, J. T.: Characterization of atmospheric ammonia emissions from
440	swine waste storage and treatment lagoons. J. Geophy. ResAtmos., 105, 11535-11545, 2000.

- 441 Battaglia, M. A., Douglas, S., and Hennigan, C. J.: Effect of the urban heat island on aerosol pH,
- 442 Environ. Sci. Technol., 51, 13095–13103, 2017.
- 443 Behera, S. N., Betha, R., Liu, P., Balasubramanian, R.: A study of diurnal variations of PM<sub>2.5</sub> acidity
- 444 and related chemical species using a new thermodynamic equilibrium model. Sci. Total Environ.,
  445 452, 286–295, 2013.
- 446 Boucher, O., Anderson, T. L.: General circulation model assessment of the sensitivity of direct climate
- forcing by anthropogenic sulfate aerosols to aerosol size and chemistry. J. Geophy. Res.-Atmos.,
  100, 26117–26134, 1995.
- 449 Bougiatioti, A., Nikolaou, P., Stavroulas, I., Kouvarakis, G., Weber, R., Nenes, A., Kanakidou, M.,
- 450 Mihalopoulos, N.: Particle water and pH in the eastern Mediterranean: source variability and
- 451 implications for nutrient availability. Atmos. Chem. Phy., 16, 4579–4591, 2016.
- 452 Bureau of Statistics of Henan (NBH), 2018. Henan Statistical Yearbook. China Statistics Press, Beijing.
  453 Accessed date: Dec. 2019.
- 454 Cheng, Y., Zheng, G., Wei, C., Mu, Q., Zheng, B., Wang, Z., Gao, M., Zhang, Q., He, K., Carmichael,
- 455 G.: Reactive nitrogen chemistry in aerosol water as a source of sulfate during haze events in China.
- 456 Sci. Adv., 2, e1601530–e1601530, 2016.
- 457 Clegg, S. L., Brimblecombe, P., Wexler, A. S.: The thermodynamic model of the system
- 458  $H^+-NH_4^+-SO_4^{2-}-NO_3^--H_2O$  at tropospheric temperatures. J. Phy. Chem. A, 102, 2137–2154,
- 459 1998.

- Ding, J., Zhao, P., Su, J., Dong, Q., Du, X., Zhang, Y.: Aerosol pH and its driving factors in Beijing.
  Atmos. Chem. Phy., 19, 7939–7954, 2019.
- 462 Dockery, D. W., Cunningham, J., Damokosh, A. L., Neas, L. M., Spengler, J. D., Koutrakis, P., Ware,
- 463 J. H., Raizenne, M., Speizer, F. E.: Health effects of acid aerosols on North American children:
- 464 respiratory symptoms. Environ. Health Persp., 104, 500–505, 1996.
- 465 Guo, H., Sullivan, A. P., Campuzanojost, P., Schroder, J. C., Lopezhilfiker, F. D., Dibb, J. E., Jimenez,
- 466 J. L., Thornton, J. A., Brown, S. S., Nenes, A.: Fine particle pH and the partitioning of nitric acid
- 467 during winter in the northeastern United States. J. Geophy. Res.-Atmos., 121, 10–355, 2016.
- Guo, H., Weber, R. J., Nenes, A.: High levels of ammonia do not raise fine particle pH sufficiently to
  yield nitrogen oxide-dominated sulfate production. Sci. Rep., 7, 12109, 2017.
- 470 Guo, H., Xu, L., Bougiatioti, A., Cerully, K. M., Capps, S. L., Hite, J. R., Carlton, A. G., Lee, S., Bergin,
- 471 M. H., Ng, N. L.: Fine-particle water and pH in the southeastern United States. Atmos. Chem.
- 472 Phy., 15, 5211–5228, 2015.
- Guo, H., Nenes, A., and Weber, R. J.: The underappreciated role of nonvolatile cations on aerosol
  ammonium-sulfate molar ratios, Atmos. Chem. Phys., 18, 17307–17323, 2018.
- 475 Haskins, J. D., Jaegle, L., Shah, V., Lee, B. H., Lopez-Hilfiker, F. D., Campuzano-Jost, P., Schroder, J.
- 476 C., Day, D. A., Guo, H. Y., Sullivan, A. P., Weber, R., Dibb, J., Campos, T., Jimenez, J. L., Brown,
- 477 S. S., and Thornton, J. A.: Wintertime gas-particle partitioning and speciation of inorganic
- 478 chlorine in the lower troposphere over the northeast United States and coastal ocean, J. Geophys.
- 479 Res.-Atmos., 123, 12897–12916, 2018
- 480 Healy, R. M., Sciare, J., Poulain, L., Crippa, M., Wiedensohler, A., Prévôt, A. S. H., Baltensperger, U., 23

482	Evans, G. J., and Wenger, J. C.: Quantitative determination of carbonaceous particle mixing state
483	in Paris using single-particle mass spectrometer and aerosol mass spectrometer measurements,
484	Atmos. Chem. Phys., 13, 9479–9496, 2013.
485	Hennigan, C. J., Izumi, J., Sullivan, A. P., Weber, R. J., and Nenes, A.: A critical evaluation of proxy
486	methods used to estimate the acidity of atmospheric particles, Atmos. Chem. Phys., 15, 2775-
487	2790, 2015.
488	Hu, G., Zhang, Y., Sun, J., Zhang, L., Shen, X., Lin, W., Yang, Y.: Variability, formation and acidity of
489	water-soluble ions in PM2.5 in Beijing based on the semi-continuous observations. Atmos. Res.,
490	145, 1–11, 2014.
491	Huang, X., Song, Y., Li, M., Li, J., Huo, Q., Cai, X., Zhu, T., Hu, M., Zhang, H.: A high-resolution
492	ammonia emission inventory in China. Global Biogeochem. Cy., 26, GB1030, 2012.
493	Larssen, T., Lydersen, E., Tang, D., He, Y., Gao, J., Liu, H., Duan, L., Seip, H.M., Vogt, R. D., Mulder,
494	J.: Acid rain in China. Environ. Sci. Technol., 40, 418–425, 2006.
495	Lefer, B. L., Talbot, R. W., Munger, J. W.: Nitric acid and ammonia at a rural northeastern US site. J.
496	Geophy. ResAtmos., 104, 1645–1661, 1999.
497	Li, Y. J., Sun, Y., Zhang, Q., Li, X., Li, M., Zhou, Z., Chan, C. K.: Real-time chemical characterization
498	of atmospheric particulate matter in China: A review. Atmos. Environ., 158, 270-304, 2006.
499	Liu, J., Bergin, M., Guo, H., King, L., Kotra, N., Edgerton, E., Weber, R. J.: Size-resolved
500	measurements of brown carbon in water and methanol extracts and estimates of their contribution
501	to ambient fine-particle light absorption. Atmos. Chem. Phy., 13, 12389–12404, 2013.

Sarda-Estève, R., McGuire, M. L., Jeong, C.-H., McGillicuddy, E., O'Connor, I. P., Sodeau, J. R.,

502	Liu, M., Song, Y., Zhou, T., Xu, Z., Yan, C., Zheng, M., Wu, Z., Hu, M., Wu, Y., Zhu, T.: Fine particle
503	pH during severe haze episodes in northern China. Geophys. Res. Lett., 44, 5213-5221, 2017.
504	Liu, W., Lin, S., and Hu, G.: Characterizing remarkable changes of severe haze events and chemical
505	compositions in multi-size airborne particles ( $PM_1$ , $PM_{2.5}$ and $PM_{10}$ ) from January 2013 to
506	2016–2017 winter in Beijing, China, Atmos. Environ., 189, 133–144, 2018.
507	Markovic, M. Z., Vandenboer, T. C., Murphy, J. G.: Characterization and optimization of an online
508	system for the simultaneous measurement of atmospheric water-soluble constituents in the gas
509	and particle phases. J Environ. Monitor., 14, 1872–1884, 2012.
510	Meng, Z., Lin, W., Jiang, X., Yan, P., Wang, Y., Zhang, Y. M., Jia, X. F., Yu, X. L.: Characteristics of
511	atmospheric ammonia over Beijing, China. Atmos. Chem. Phy., 11, 6139-6151, 2011.
512	Meng, Z., Xu, X., Lin, W., Ge, B., Xie, Y., Song, B., Jia, S., Zhang, R., Peng, W., Wang, Y.: Role of
513	ambient ammonia in particulate ammonium formation at a rural site in the North China Plain.
514	Atmos. Chem. Phy., 18, 167–184, 2017.
515	MEP: 2017 air pollution prevention and management plan for the Beijing-Tianjin-Hebei region and its
516	surrounding areas, http://dqhj.mee.gov.cn/dtxx/201703/t20170323_408663.shtml (last access: 18
517	August 2019), 2017.
518	Meskhidze, N., Chameides, W. L., Nenes, A., Chen, G.: Iron mobilization in mineral dust: Can
519	anthropogenic SO <sub>2</sub> emissions affect ocean productivity? Geophys. Res. Lett., 30, 2003.
520	Murphy, J. G., Gregoire, P. K., Tevlin, A. G., Wentworth, G. R., Ellis, R. A., Markovic, M. Z.,
521	VandenBoer, T. C.: Observational constraints on particle acidity using measurements and
522	modelling of particles and gases, Faraday Discuss., 200, 379–395, 2017.
	25

523	Nah, T., Guo, H., Sullivan, A. P., Chen, Y., Tanner, D. J., Nenes, A., Russell, A., Ng, L. N., Huey, L.
524	G., Weber, R. J.: Characterization of aerosol composition, aerosol acidity, and organic acid
525	partitioning at an agriculturally intensive rural southeastern US site. Atmos. Chem. Phy. 18,
526	11471–11491. 2018.

- 527 National Bureau of Statistics (NBS), 2016. China Statistical Yearbook. China Statistics Press, Beijing.
  528 Accessed date: Sept. 2019.
- Nenes, A., Pandis, S. N., Pilinis, C.: ISORROPIA: A new thermodynamic equilibrium model for
  multiphase multicomponent inorganic aerosols. Aquat. Geochem., 4, 123–152, 1998.
- 531 Nenes, A., Pandis, S. N., Weber, R. J., and Russell, A.: Aerosol pH and liquid water content determine
- when particulate matter is sensitive to ammonia and nitrate availability, Atmos. Chem. Phys.Discuss., in review, 2019.
- Ostro, B., Lipsett, M., Wiener, M.B., Selner, J.C.: Asthmatic responses to airborne acid aerosols. Am.
  J. Public Health, 81, 694–702, 1991.
- 536 Panteliadis, P., Hafkenscheid, T., Cary, B., Diapouli, E., Fischer, A., Favez, O., Quincey, P., Viana, M.,
- 537 Hitzenberger, R., Vecchi, R.: ECOC comparison exercise with identical thermal protocols after
- temperature offset correction: instrument diagnostics by in-depth evaluation of operational
  parameters. Atmos. Meas. Tech., 8, 779–792, 2014.
- 540 Pathak, R. K., Louie, P. K., Chan, C. K.: Characteristics of aerosol acidity in Hong Kong. Atmos.
- 541 Environ., 38, 2965–2974, 2004.
- 542 Pathak, R.K., Wu, W.S., Wang, T.: Summertime PM<sub>2.5</sub> ionic species in four major cities of China:
- 543 nitrate formation in an ammonia-deficient atmosphere. Atmos. Chem. Phys. 9, 1711–1722, 2008.

544	Pye, H. O. T., Nenes, A., Alexander, B., Ault, A. P., Barth, M. C., Clegg, S. L., Collett Jr., J. L., Fahey,
545	K. M., Hennigan, C. J., Herrmann, H., Kanakidou, M., Kelly, J. T., Ku, IT., McNeill, V. F.,
546	Riemer, N., Schaefer, T., Shi, G., Tilgner, A., Walker, J. T., Wang, T., Weber, R., Xing, J., Zaveri,
547	R. A., and Zuend, A.: The acidity of atmospheric particles and clouds, Atmos. Chem. Phys.
548	Discuss., in review, 2019.
549	Rumsey, I. C., Cowen, K. A., Walker, J. T., Kelly, T. J., Hanft, E. A., Mishoe, K., Rogers, C., Proost,
550	R., Beachley, G. M., Lear, G.: An assessment of the performance of the Monitor for AeRosols
551	and GAses in ambient air (MARGA): a semi-continuous method for soluble compounds. Atmos.
552	Chem. Phy., 14, 5639–5658, 2014.
553	Saraswati, Sharma, S. K., Saxena, M., Mandal, T. K.: Characteristics of gaseous and particulate
554	ammonia and their role in the formation of secondary inorganic particulate matter at Delhi, India.
555	Atmos. Res., 218, 34–49, 2019.
556	Seinfeld, J. H. and Pandis, S. N.: Atmospheric Chemistry and Physics: From Air Pollution to Climate
557	Change, 2nd Edition, John Wiley and Sons, Inc., Hoboken, New Jersey, USA, 2006.

- 558 Seinfeld, J. H. and Pandis, S. N.: Atmospheric Chemistry and Physics: From Air Pollution to Climate 559 Change (3rd edition), John Wiley and Sons, Inc., Hoboken, New Jersey, USA, 2016.
- 560 Shen, J., Liu, X., Ying, Z., Fangmeier, A., Goulding, K., Zhang, F.: Atmospheric ammonia and 561 particulate ammonium from agricultural sources in the North China Plain. Atmos. Environ., 45, 5033-5041, 2011. 562
- 563 Shi, G., Xu, J., Peng, X., Xiao, Z., Chen, K., Tian, Y., Guan, X., Feng, Y., Yu, H., Nenes, A.: pH of
- aerosols in a polluted atmosphere: source contributions to highly acidic aerosol. Environ. Sci. 564 27

565 Technol., 51, 4289–4296, 2017.

566	Shi, X.,	, Nenes, A.	, Xiao, Z	, Song,	S., Yı	ı, H.,	Shi, G	., Zhao,	Q.,	Chen,	K.,	Feng,	Y.,	Russell, A	<u>ь.</u> С	Э.:
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- 567 High-resolution data sets unravel the effects of sources and meteorological conditions on nitrate
  568 and its gas-particle partitioning. Environ. Sci. Technol., 53, 3048–3057, 2019.
- Shi, Z., Bonneville, S., Krom, M. D., Carslaw, K. S., Jickells, T. D., Baker, A. R., Benning, L. G.: Iron
  dissolution kinetics of mineral dust at low pH during simulated atmospheric processing. Atmos.
  Chem. Phy., 11, 995–1007, 2010.
- 572 Song, S., Gao, M., Xu, W., Shao, J., Shi, G., Wang, S., Wang, Y., Sun, Y., Mcelroy, M. B.: Fine particle
- 573 pH for Beijing winter haze as inferred from different thermodynamic equilibrium models. Atmos.
  574 Chem. Phy., 18, 7423–7438, 2018.
- 575 Spurny, K. R.: Atmospheric acidic aerosols (review). J Aerosol Sci., 21, 1990.
- 576 Surratt, J. D., Chan, A. W. H., Eddingsaas, N. C., Chan, M. N., Loza, C. L., Kwan, A. J., Hersey, S.,
- 577 Flagan, R. C., Wennberg, P. O., Seinfeld, J. H.: Reactive intermediates revealed in secondary 578 organic aerosol formation from isoprene. P. Natl. Acad. Sci. USA., 107, 6640–6645, 2010.
- 579 Tao, Y. and Murphy, J. G.: The sensitivity of PM<sub>2.5</sub> acidity to meteorological parameters and chemical
- 580 composition changes: 10-year records from six Canadian monitoring sites. Atmos. Chem. Phy..
- 581 19, 9309–9320, 2019.
- 582 Tian, M., Wang, H., Chen, Y., Zhang, L., Shi, G., Liu, Y., Yu, J., Zhai, C., Wang, J., Yang, F.: Highly
- time-resolved characterization of water-soluble inorganic ions in PM<sub>2.5</sub> in a humid and acidic
  mega city in Sichuan Basin, China. Sci. Total Environ., 580, 224–234, 2017.
- 585 Vasilakos, P., Russell, A., Weber, R., and Nenes, A.: Understanding nitrate formation in a world with 28

less sulfate, Atmos. Chem. Phys., 18, 12765–12775, 2018.

587	Wang, C.,	Yin, S.	, Bai, L.	, Zhang, X.,	, Gu, X	., Zhang,	H., Lu,	Q., Zhang	, R.: Higł	n-resolution	ammonia
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- 588 emission inventories with comprehensive analysis and evaluation in Henan, China, 2006–2016.
- 589 Atmos. Environ., 193, 11–23, 2018a.
- 590 Wang, G., Zhang, F., Peng, J., Duan, L., Ji, Y., Marreroortiz, W., Wang, J., Li, J., Wu, C., Cao, C.:
- 591 Particle acidity and sulfate production during severe haze events in China cannot be reliably 592 inferred by assuming a mixture of inorganic salts. Atmos. Chem. Phy., 18, 1–23, 2018b.
- 593 Wang, G., Zhang, R., Gomez, M. E., Yang, L., Levy, Z. M., Hu, M., Lin, Y., Peng, J., Guo, S., Meng,
- J.: Persistent sulfate formation from London Fog to Chinese haze. P. Natl. Acad. Sci. USA., 113,
  13630–13635, 2016.
- 596 Wang, H., Ding, J., Xu, J., Wen, J., Han, J., Wang, K., Shi, G., Feng, Y., Ivey, C., Wang, Y.: Aerosols
- 597 in an arid environment: The role of aerosol water content, particulate acidity, precursors, and
- relative humidity on secondary inorganic aerosols. Sci. Total Environ., 646, 564–572, 2019a.
- 599 Wang, J., Zhao, B., Wang, S., Yang, F., Xing, J., Morawska, L., Ding, A., Kulmala, M., Kerminen, V.,
- Kujansuu, J.: Particulate matter pollution over China and the effects of control policies. Sci. Total
  Environ., 584, 426–447, 2017.
- Wang, S., He, B., Yuan, M., Su, F., Yin, S., Yan, Q., Jiang, N., Zhang, R., Tang, X.: Characterization
- of individual particles and meteorological conditions during the cold season in Zhengzhou using
  a single particle aerosol mass spectrometer. Atmos. Res., 219, 13–23, 2019b.
- Wang, S., Yan, Q., Yu, F., Wang, Q., Yang, L., Zhang, R., Yin, S., Wang, S., Yan, Q., Yu, F.: Distribution
- and source of chemical elements in size-resolved particles in Zhengzhou, China: Effects of 29

607	regional transport. Aerosol Air Qual. Res., 18, 371–385, 2018c.
608	Wang, S., Yin, S., Zhang, R., Yang, L., Zhao, Q., Zhang, L., Yan, Q., Jiang, N., Tang, X.: Insight into
609	the formation of secondary inorganic aerosol based on high-time-resolution data during haze
610	episodes and snowfall periods in Zhengzhou, China. Sci. Total Environ., 660, 47–56, 2019c.
611	Warneck, P.: Chemistry of the Natural Atmosphere, Academic Press, San Diego, CA, 1988.
612	Watson, J. G.: Visibility: science and regulation. J Air Waste Manage., 52, 973-999, 2002.
613	Weber, R. J., Guo, H., Russell, A. G., and Nenes, A.: High aerosol acidity despite declining
614	atmospheric sulfate concentrations over the past 15 years, Nat. Geosci., 9, 282-285, 2016.
615	Wen, L., Chen, J., Yang, L., Wang, X., Xu, C., Sui, X., Yao, L., Zhu, Y., Zhang, J., Zhu, T.: Enhanced
616	formation of fine particulate nitrate at a rural site on the North China Plain in summer: The
617	important roles of ammonia and ozone. Atmos. Environ., 101, 294-302, 2015.
618	Xie, Y., Wang, G., Wang, X., Chen, J., Chen, Y., Tang, G., Wang, L., Ge, S., Xue, G., Wang, Y., and
619	Gao, J.: Observation of nitrate dominant PM2.5 and particle pH elevation in urban Beijing during
620	the winter of 2017, Atmos. Chem. Phys. Discuss., in review, 2019.

622	Figure	lists:

Fig. 1 pH uncertainties of the five sites based on two extreme scenarios of measurement uncertainty. 623 624 Cations and organic carbon concentrations (OC) were adjusted up to within their maximum positive 625 uncertainties, anions, relative humidity (RH) and temperature (T) were adjusted down within their 626 maximum negative uncertainties, which represented the pH<sub>max</sub> case; for pH<sub>min</sub> case, cations and OC concentrations were adjusted down, and anions, RH and T were adjusted up. The color scale bar 627 628 represents adjusted RH. 629 Fig. 2 Temporal variations of T, RH, wind speed (WS), wind direction (WD), and concentrations of NH<sub>3</sub>, NH<sub>4</sub><sup>+</sup>, SO<sub>4</sub><sup>2-</sup>, and NO<sub>3</sub><sup>-</sup> during three cases at the Zhengzhou (U-ZZ) site. The shaded areas 630 631 represent the measurement uncertainties. Fig. 3 Time series and box plot of predicted PM<sub>2.5</sub> pH, H<sup>+</sup><sub>air</sub>, and aerosol water content (AWC) at the 632 633 five sites. In each box, the top, middle and bottom lines represent the 75th, 50th, and 25th percentile 634 of statistical data, respectively; the upper and lower whiskers represent the maximum and minimum 635 values, respectively. 636 Fig. 4 (a)–(e) Diurnal patterns of median pH values, (f) average RH, and T of the five sites in Case 2. 637 The upper and lower ends of the line represent the maximum and minimum values of pH, 638 respectively. The color scale bar represents AWC concentration. 639 Fig. 5 Comparison of the sensitivities of PM<sub>2.5</sub> pH to T, RH, TCl (HCl<sub>(g)</sub> + Cl<sub>(aq)</sub>), TNH<sub>x</sub> (NH<sub>3(g)</sub> + 640  $NH_{4(aq)}^{+}$ ), TNO<sub>3</sub> (HNO<sub>3(g)</sub> + NO<sub>3(aq)</sub><sup>-</sup>), and TH<sub>2</sub>SO<sub>4</sub> (replaced by observed SO<sub>4</sub><sup>2-</sup>) among the five 641 sites. A given range for a variable for all sites with corresponding average values of other parameters 642 was used as input to the ISORROPIA-II model. The range of the x-axis is close to the observed 31

643 minimum and maximum values in Case 2 (Table S4). The color scale bar represents the pH values.
644 The relative standard deviation (RSD) and range (Range) represent the sensitivity degree of pH to
645 this variable and range (min–max) of the re-predicted pH value in the test, respectively. The square
646 plots on the graph represent the average values of each variable observed in Case 2 with standard
647 deviation as an error bar.

648 Fig. 6 Particle pH calculated with fixed meteorological parameters (T = 275.5 K and RH = 60 %)

under different combinations of  $\text{TNH}_x$  and (a)  $\text{TH}_2\text{SO}_4$  (Fixed  $\text{TNO}_3 = 67.5 \ \mu\text{g/m}^3$ ) and (b)  $\text{TNO}_3$ 

650 (Fixed TH<sub>2</sub>SO<sub>4</sub> = 36.5  $\mu$ g/m<sup>3</sup>). The color scale bar represents the pH values. The markers on the

651 graph represent the average concentrations of TNH<sub>x</sub>, TH<sub>2</sub>SO<sub>4</sub>, and TNO<sub>3</sub> at the five sites during Case

652 2 with standard deviation as error bar.

Fig. 7 Particle pH corresponds to increasing  $TNH_x$  at the five sites to examine the effects of major

654 indicators of NH<sub>3</sub> (i.e., TNH<sub>x</sub>, Required-NH<sub>x</sub>, and Excess-NH<sub>x</sub>) on aerosol acidity. Particle pH was

655 calculated by using a wide range of  $TNH_x$  (25–130  $\mu$ g/m<sup>3</sup>) and average values of other parameters

during Case 2 of each site. The concentrations of TNH<sub>x</sub>, Required-NH<sub>x</sub>, and Excess-NH<sub>x</sub> with

657 corresponding pH values are marked by a hollow box, hollow circle, and arrow respectively. The

658 yellow and blue background colors correspond to the NH<sub>x</sub>-poor and NH<sub>x</sub>-rich, respectively.

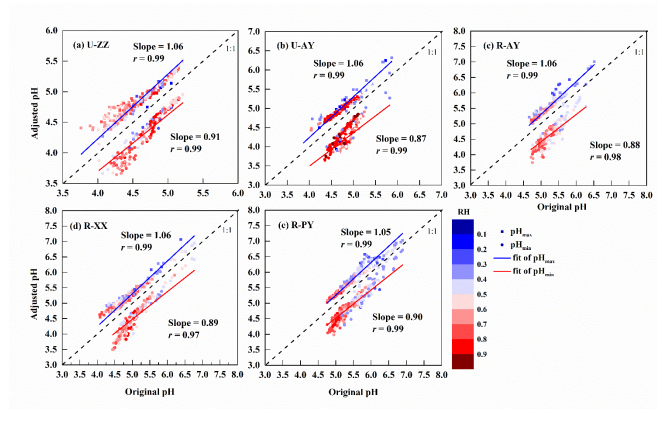


Fig. 1 pH uncertainties of the five sites based on two extreme scenarios of measurement uncertainty. Cations and organic carbon concentrations (OC) were adjusted up to within their maximum positive uncertainties, anions, relative humidity (RH) and temperature (T) were adjusted down within their maximum negative uncertainties, which represented the  $pH_{max}$  case; for  $pH_{min}$  case, cations and OC concentrations were adjusted down, and anions, RH and T were adjusted up. The color scale bar represents adjusted RH.

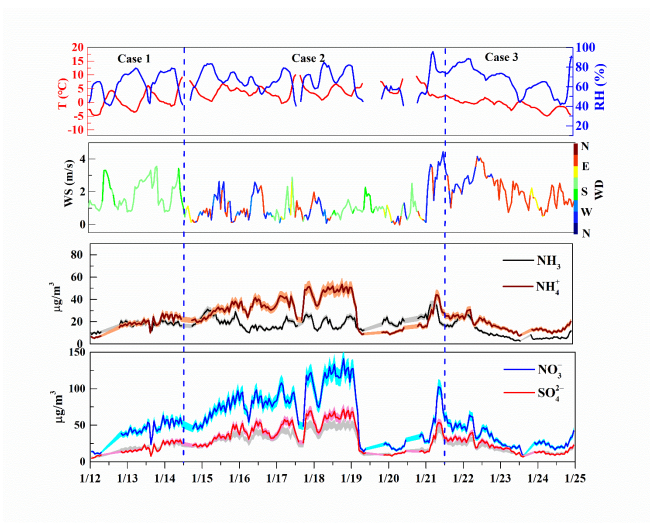


Fig. 2 Temporal variations of T, RH, wind speed (WS), wind direction (WD), and concentrations of NH<sub>3</sub>, NH<sub>4</sub><sup>+</sup>, SO<sub>4</sub><sup>2-</sup>, and NO<sub>3</sub><sup>-</sup> during three cases at the Zhengzhou (U-ZZ) site. The shaded areas represent the measurement uncertainties.

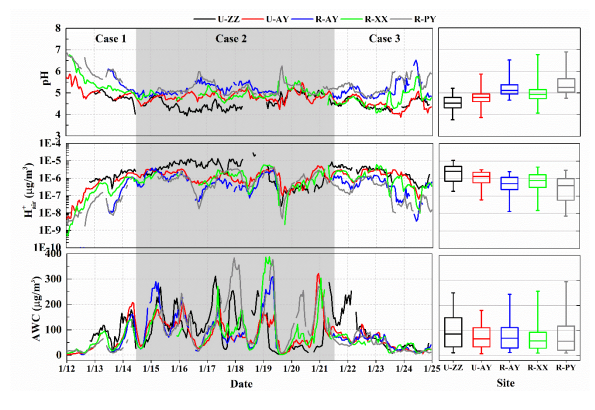


Fig. 3 Time series and box plot of predicted  $PM_{2.5}$  pH,  $H^+_{air}$ , and aerosol water content (AWC) at the five sites. In each box, the top, middle and bottom lines represent the 75th, 50th, and 25th percentile of statistical data, respectively; the upper and lower whiskers represent the maximum and minimum values, respectively.

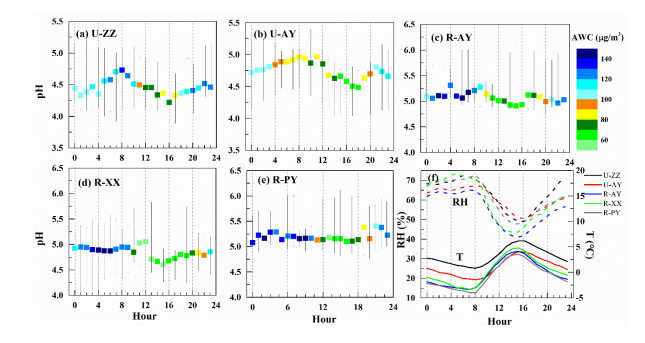


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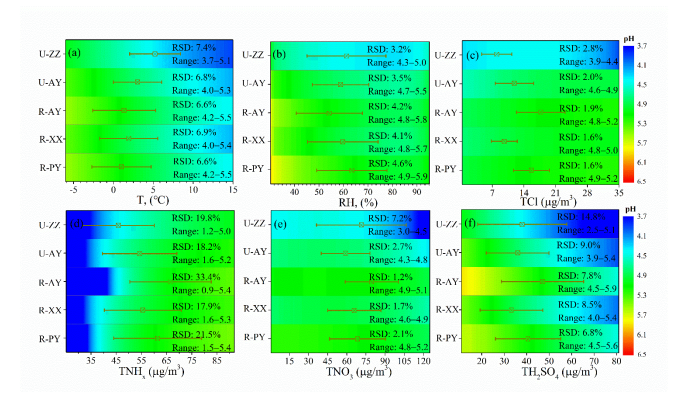


Fig. 5 Comparison of the sensitivities of PM<sub>2.5</sub> pH to T, RH, TCl (HCl<sub>(g)</sub> + Cl<sub>(aq)</sub>), TNH<sub>x</sub> (NH<sub>3(g)</sub> +  $NH_{4(aq)}^{+}$ ), TNO<sub>3</sub> (HNO<sub>3(g)</sub> + NO<sub>3(aq)</sub><sup>-</sup>), and TH<sub>2</sub>SO<sub>4</sub> (replaced by observed SO<sub>4</sub><sup>2-</sup>) among the five sites. A given range for a variable for all sites with corresponding average values of other parameters was used as input to the ISORROPIA-II model. The range of the x-axis is close to the observed minimum and maximum values in Case 2 (Table S4). The color scale bar represents the pH values. The relative standard deviation (RSD) and range (Range) represent the sensitivity degree of pH to this variable and range (min-max) of the re-predicted pH value in the test, respectively. The square plots on the graph represent the average values of each variable observed in Case 2 with standard deviation as an error bar.

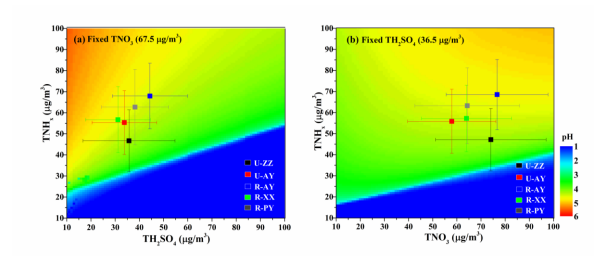




Fig. 6 Particle pH calculated with fixed meteorological parameters (T = 275.5 K and RH = 60 %) under different combinations of TNH<sub>x</sub> and (a) TH<sub>2</sub>SO<sub>4</sub> (Fixed TNO<sub>3</sub> = 67.5  $\mu$ g/m<sup>3</sup>) and (b) TNO<sub>3</sub> (Fixed TH<sub>2</sub>SO<sub>4</sub> = 36.5  $\mu$ g/m<sup>3</sup>). The color scale bar represents the pH values. The markers on the graph represent the average concentrations of TNH<sub>x</sub>, TH<sub>2</sub>SO<sub>4</sub>, and TNO<sub>3</sub> at the five sites during Case 2 with standard deviation as error bar.

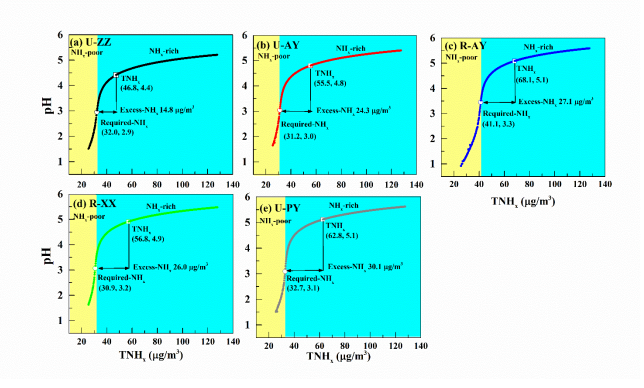




Fig. 7 Particle pH corresponds to increasing  $TNH_x$  at the five sites to examine the effects of major indicators of NH<sub>3</sub> (i.e.,  $TNH_x$ , Required-NH<sub>x</sub>, and Excess-NH<sub>x</sub>) on aerosol acidity. Particle pH was calculated by using a wide range of  $TNH_x$  (25–130 µg/m<sup>3</sup>) and average values of other parameters during Case 2 of each site. The concentrations of  $TNH_x$ , Required-NH<sub>x</sub>, and Excess-NH<sub>x</sub> with corresponding pH values are marked by a hollow box, hollow circle, and arrow respectively. The yellow and blue background colors correspond to the NH<sub>x</sub>-poor and NH<sub>x</sub>-rich, respectively.

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# 719 **Table lists:**

720 Table 1 Comparison of the particle pH values in this study (median, min–max) and other sites (mean ± standard deviation).

- 721 Table 2 Descriptions of the five sampling sites.
- Table 3 Summary (mean  $\pm$  standard deviation) of gaseous precursors ( $\mu g/m^3$ ), water-soluble inorganic ions ( $\mu g/m^3$ ), T (°C), and RH (%) during three cases
- 723 of haze periods at five monitoring sites.

724 Table 4 Sensitivity of pH to input data. The real-time measured values of a variable and the average values of other parameters during Case 2 were input

725 into the ISORROPIA II. The degree of sensitivity was represented as the relative standard deviation (%RSD) of the re-calculated pH values, higher RSD

726 implied higher sensitivity of this factor to pH and thus is more important for local pH.

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	Observation site	Period	pН	Model	Reference
	Zhengzhou, China (Urban)	Jan 2018	4.5 (3.8–5.2)		
	Anyang, China (Urban)	Jan 2018	4.8 (3.9–5.8)		
This study	Anyang, China (Rural)	Jan 2018	4.9 (4.1–6.2)	ISORROPIA-II	
	Xinxiang, China (Rural)	Jan 2018	5.1 (4.7–6.3)		
	Puyang, China (Rural)	Jan 2018	5.2 (4.8–6.5)		
	Beijing, China (Urban)	Feb 2017	$4.5\pm0.7$	ISORROPIA-II	Ding et al., 2019
	Beijing, China (Urban)	Dec 2016	$4.3\pm0.4$	ISORROPIA-II	Liu et al., 2017
	Beijing, China (Urban)	Jan–Feb 2015	4.5	ISORROPIA-II	Guo et al., 2017
	Xi'an, China (Urban)	Nove–Dec 2012	5.0	ISORROPIA-II	Guo et al., 2017
China	Tianjin, China (Urban)	Dec–Jun 2015	$4.9\pm1.4$	ISORROPIA-II	Shi et al., 2017
	Tianjin, China (Urban)	Aug 2015	$3.4\pm0.5$	ISORROPIA-II	Shi et al., 2019
	Hohhot, China	Winter 2015	5.7	ISORROPIA-II	Wang et al., 2019
	PRD, China (Rural)	Fall-winter season 2012	$0.81\pm0.24$	AIM-II model	Fu et al., 2015
	Hong Kong, China (Urban)	2001	0.25	AIM-II model	Pathak et al., 2004
	Singapore (Urban)	Sep-Nov 2011	0.60	AIM-IV model	Sailesh et al., 2013
	Northeastern US (Urban)	Feb-Mar 2015	$0.07\pm0.96$	ISORROPIA-II	Guo et al., 2016
Other	Alabama, USA (Rural)	Jun–Jul 2013	$1.94\pm0.59$	ISORROPIA-II	Guo et al., 2015
countries	Georgia, USA (Rural)	Aug-Oct 2016	$2.2\pm0.6$	ISORROPIA-II	Nah et al., 2018
	Crete, Greece (Background)	Aug–Nov 2012	$1.25 \pm 1.14$	ISORROPIA-II	Boucher et al., 20

Table 1 Comparison of the particle pH values in this study (median, min–max) and other sites (mean ± standard deviation).

						Emissio	n (Gg)	
City	Classification	Site	Coordinate	Location	Surrounding environment	$\mathrm{NH_3}^*$	${\rm SO_2}^{**}$	NO <sub>x</sub> **
Zhengzhou	Urban	U-ZZ	34.82° N 113.54° E	West totheZhengzhoudowntown(ZhengzhouUniversity)	Densely occupied residences, light industry, freeways and roads	39.2	36.7	31.8
Anyang	Urban	U-AY	36.09° N 114.41° E	East to the Anyang downtown (Anyang Environmental Protection Bureau)	Occupied residences, heavy industry and traffic roads	57.8	46.9	37.1
	Rural R-AY		36.22° N 114.39° E	15 km north of Anyang city (Baizhuang town)	High ways, small villages, and cropland			
Xinxiang	Rural	R-XX	35.38° N 114.30° E	35 km northeast of Xinxiang city (Banzao town)	Small villages and cropland	72.1	8.9	19.6
Puyang	Rural	R-PY	36.15° N 115.10° E	44 km north of Puyang city (Liangcun town)	Small villages and cropland	39.6	3.4	3.2

<sup>\*</sup> Data from a 2015-based NH<sub>3</sub> emission inventory (Wang et al., 2018a).

<sup>\*\*</sup> Data from Henan Statistical Yearbook of 2018 (BSH, 2018).

# 741 Table 3 Summary (mean $\pm$ standard deviation) of gaseous precursors ( $\mu g/m^3$ ), water-soluble inorganic ions ( $\mu g/m^3$ ), T (°C), and RH (%) during three cases

742 of haze periods at five sites.

	Case 1 (Janua	ary 12–14)				Case 2 (January 14–21)					Case 3 (January 21–25)				
	U-ZZ	U-AY	R-AY	R-XX	R-PY	U-ZZ	U-AY	R-AY	R-XX	R-PY	U-ZZ	U-AY	R-AY	R-XX	R-PY
HNO <sub>3</sub>	$0.9\pm0.2$	$0.7\pm0.2$	$3.1\pm 0.2$	$3.0\pm 0.1$	$3.9\pm 0.1$	$1.3\pm0.3$	$1.1\pm0.3$	$3.7\pm 0.4$	$3.7\pm 0.5$	$4.2\pm0.2$	$0.9\pm0.3$	$0.7\pm0.2$	$4.9\pm1.0$	$3.3\pm 0.2$	$3.3\pm 0.2$
NH <sub>3</sub>	$17.0\pm3.7$	$19.6\pm8.0$	$22.9\pm 6.3$	$21.6\pm4.1$	$17.8\pm3.7$	$19.5\pm5.2$	$23.6\pm 6.5$	$25.2\pm 6.5$	$24.7\pm9.9$	$26.5\pm 6.7$	$10.5\pm 6.9$	$8.8\pm 4.7$	$10.6\pm4.7$	$8.4\pm 3.5$	$12.1\pm3.5$
HCl	$0.1\pm0.0$	$0.7\pm0.6$	$0.5\pm0.1$	$0.6\pm0.1$	$1.8\pm0.1$	$0.1\pm0.1$	$0.4\pm0.1$	$0.6\pm0.2$	$0.6\pm0.1$	$2.0\pm0.1$	$0.1\pm0.1$	$0.5\pm0.1$	$1.7\pm0.1$	$1.0\pm0.4$	$1.5\pm0.4$
NO <sub>3</sub> -	$41.5\pm14.6$	$28.0\pm 14.6$	$43.0\pm12.5$	$32.8 \pm 12.9$	$25.2\pm9.1$	$74.2\pm32.9$	$58.0 \pm 18.3$	$76.8\pm21.1$	$64.1\pm18.7$	$64.4\pm21.7$	$32.4\pm13.5$	$18.9\pm5.4$	$26.0\pm5.8$	$25.1\pm 6.7$	$18.8\pm4.3$
$\mathrm{NH_4}^+$	$18.6\pm 6.2$	$15.9\pm8.3$	$21.8\pm8.0$	$14.9\pm 6.2$	$12.8\pm4.5$	$31.9\pm12.5$	$35.2\pm12.0$	$47.6\pm13.1$	$35.6\pm10.4$	$39.9\pm 14.9$	$17.4\pm 6.0$	$11.6\pm4.4$	$14.3\pm4.4$	$12.9\pm4.0$	$10.1\pm2.9$
$\mathrm{SO_4}^{2-}$	$17.8\pm7.2$	$14.4\pm9.0$	$13.7\pm10.0$	$10.0\pm5.5$	$8.6\pm2.3$	$\textbf{38.3} \pm \textbf{18.0}$	$34.5\pm13.0$	$46.8\pm16.9$	$32.9 \pm 12.5$	$39.2\pm13.6$	$19.8\pm8.6$	$15.1\pm 6.1$	$15.1\pm7.3$	$14.4\pm4.8$	$13.3\pm4.0$
$Ca^{2+}$	$0.7\pm0.5$	$0.5\pm0.3$	$5.0\pm2.2$	$0.8\pm0.2$	$3.4\pm 0.3$	$0.5\pm0.4$	$0.4\pm0.4$	$2.2\pm1.2$	$1.0\pm0.3$	$3.3\pm 0.6$	$0.1\pm0.1$	$0.2\pm0.2$	$1.8\pm0.7$	$0.5\pm0.1$	$2.4\pm0.5$
$Na^+$	$1.5\pm0.2$	$1.0\pm0.0$	$1.4\pm0.4$	$0.7\pm0.1$	$2.2\pm0.1$	$1.6\pm0.2$	$1.0\pm0.1$	$1.4\pm0.4$	$0.8\pm0.1$	$2.2\pm0.0$	$1.1\pm0.2$	$1.0\pm0.2$	$2.2\pm0.4$	$1.3\pm0.4$	$2.2\pm0.2$
Cl	$7.5\pm 2.5$	$2.7\pm3.4$	$6.6\pm2.5$	$5.4\pm1.5$	$6.3\pm1.2$	$8.5\pm3.2$	$12.0\pm4.2$	$18.5\pm5.0$	$9.7\pm2.6$	$14.4\pm3.8$	$3.3\pm 1.5$	$4.5\pm1.6$	$6.9\pm1.6$	$4.7\pm0.9$	$5.4\pm1.4$
$Mg^{2+}$	$0.2\pm0.0$	$0.1\pm0.0$	$0.4\pm0.1$	$0.1\pm0.0$	$0.5\pm0.0$	$0.2\pm0.0$	$0.1\pm0.0$	$0.4\pm0.4$	$0.1\pm0.0$	$0.5\pm0.1$	$0.2\pm0.0$	$0.1\pm0.0$	$0.4\pm0.1$	$0.1\pm0.1$	$0.4\pm0.1$
$\mathbf{K}^+$	$2.9\pm0.7$	$1.4\pm0.5$	$1.6\pm0.6$	$1.6\pm0.4$	$2.5\pm0.6$	$4.4\pm1.7$	$2.4\pm0.7$	$2.9\pm0.7$	$1.9\pm0.6$	$3.7\pm1.1$	$1.9\pm0.7$	$0.9\pm0.3$	$0.8\pm0.3$	$0.8\pm0.2$	$1.2\pm0.4$
Т	$0.3\pm3.1$	$-0.9\pm3.7$	$-1.8\pm4.4$	$-2.0\pm3.9$	$-1.7\pm4.8$	$4.2\pm2.2$	$2.6\pm2.5$	$0.1\pm2.8$	$1.2\pm2.9$	$0.4\pm3.0$	$-0.8\pm1.8$	$-2.7\pm1.8$	$-3.2\pm1.9$	$-2.8\pm2.6$	$-4 \pm 2.3$
RH	$63.7 \pm 12.1$	$60.3\pm15.4$	$54.0\pm16.0$	$58.5\pm13.5$	$49.7\pm14.5$	$66.0\pm11.4$	$60.1\pm9.9$	$58.6\pm9.9$	$62.7\pm11.6$	$65.8\pm11.7$	$67\pm13.1$	$63.7\pm13.8$	$55.9 \pm 13.6$	$59\pm13.7$	$56.8\pm16$

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Table 4 Sensitivity of pH to input data. The real-time measured values of a variable and the average values of other parameters during Case 2 were input
into the ISORROPIA II. The degree of sensitivity was represented as the relative standard deviation (%RSD) of the re-calculated pH values, higher RSD
implied higher sensitivity of this factor to pH and thus is more important for local pH.

	$\mathrm{TH}_2\mathrm{SO}_4$	TNH <sub>x</sub>	TNO <sub>3</sub>	TNa	TCl	Ca <sup>2+</sup>	$\mathbf{K}^+$	$Mg^{2+}$	RH	Т
U-ZZ	12.1 %	12.8 %	6.1 %	0.1 %	0.3 %	3.3 %	0.3 %	0.2 %	1.3 %	2.7 %
U-AY	5.8 %	7.4 %	1.0 %	0.1 %	1.1 %	0.4 %	0.3 %	0.0 %	1.6 %	2.8 %
R-AY	6.5 %	10.9 %	1.4 %	0.1 %	0.7 %	1.2 %	0.2 %	0.3 %	1.9 %	2.7 %
R-XX	5.4 %	11.1 %	1.1 %	0.1 %	0.7 %	0.4 %	0.2 %	0.1 %	2.0 %	2.9 %
R-PY	4.7 %	9.4 %	1.5 %	0.0 %	0.7 %	0.5 %	0.4 %	0.1 %	2.3 %	3.1 %